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Genotyping of Polyomavirus BK by Real Time PCR for VP1 gene

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RESEARCH

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2 Genotyping of Polyomavirus BK by Real Time PCR for VP1 Gene

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8 Abstract Polyomavirus BK latently persist in different 9 sites, including the renourinary tract, and may reactivate 10 causing nephropathy in renal transplant recipients or 11 hemorrhagic cystitis in bone marrow recipients. Based on 12 the sequence of the VP1 gene, four genotypes have been 13 described, corresponding to the four serologically differ-14 entiated subtypes I-IV, with different prevalence and 15 geographic distribution. In this study, the development and 16 clinical validation of four different Real-Time PCR assays 17 for the detection and discrimination of BKV genotypes as a 18 substitute of DNA sequencing are described. 379 BK VP1 19 sequences, belonging to the main four genotypes, were 20 aligned and "hot spots" of mutation specific for all the

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strains or isolates were identified. Specific primers and 21 probes for the detection and discrimination of each geno-22 type by four Real-Time PCR assays were designed and 23 technically validated. Subsequently, the four Real-Time 24 PCR assays were used to test 20 BK-positive urine speci-25 mens from renal transplant patients, and evidenced a 26 prevalence of BK genotype I, as previously reported in 27 Europe. Results were confirmed by sequencing. The 28 29 availability of a rapid and simple genotyping method could be useful for the evaluation of BK genotypes prevalence 30 and studies on the impact of the infecting genotype on viral 31 biological behavior, pathogenic role, and immune evasion 32 33 strategies.

Keywords Polyomavirus BK · Genotyping · Real-time PCR · Sequencing · Prevalence

Introduction

BK virus (BKV) belongs to the Polyomaviridae family and 38 has a circular double-stranded DNA genome of about 39 5100 bp [1]. After primary infection that usually occurs in 40 the childhood and, which is mainly asymptomatic, BKV 41 remains latent at different sites, including the renourinary 42 tract, B-cells, and brain [2]. Reactivation with viruria may 43 44 occur, mainly in the context of immunosuppression, and 45 potentially lead to nephropathy (BKV-associated nephropathy-BKVAN) in renal transplant recipients [3] or hem-46 orrhagic cystitis in bone marrow transplant patients [4]. 47

BKV is the only primate polyomavirus with serologi-
cally differentiated subtypes (I-IV) [5]. Jin and colleagues48developed a genotyping method based on the amplification
of the epitope region of the VP1 gene by polymerase chain50reaction (PCR) and classified viral isolates in four different52



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	Article No. : 9386	□ LE	□ TYPESET	
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53 genotypes (BKV I-IV) corresponding to the serological 54 ones [6].

55 The geographical distribution of BKV subtypes has been 56 investigated in different studies [7-10], evidencing a higher 57 prevalence for BKV I worldwide and a minor but consis-58 tent occurrence of BKV IV, in particular in South-East 59 Asia. On the other hand, the prevalence of BKV II and 60 BKV III is very low with a higher presence of BKV III in 61 the African continent. This genotyping method is based on 62 the presence of single nucleotide polymorphisms (SNPs) 63 conserved between the various isolates belonging to each 64 of the four genotypes. Many different biomolecular procedures have been developed to detect and discriminate 65 66 SNPs [11–14]; among these, the ARMS PCR [15] uses the 67 discriminatory power of the terminal 3' nucleotide to obtain 68 a successful amplification only for the DNA strand with a 69 correct complementariness.

In this article, we describe the design of four different Real-Time PCR assays which can detect and discriminate among the four main BKV genotypes (I-IV) as a valid substitute of DNA sequencing.

74 **Materials and Methods**

75 **BKV** Sequences and Phylogenetic Analyses

76 379 BKV VP1 sequences, belonging to the main four genotypes (I-IV), were obtained from three different arti-77 78 cles [8–10] by extrapolation from the server Nucleotide 79 (www.ncbi.nlm.nih.gov/nucleotide/). Nucleotide positions 80 from 1663 to 1912 (Dunlop strain complete genome, 81 GeneBank Access no. V01108) were considered to obtain 82 250-bp length DNA sequences. Subsequently, the VP1 83 DNA sequences were aligned using the software Clustal X 84 (i.e., the graphical version, with windows interface, of the 85 bioinformatic software Clustal W). This software, that 86 incorporates a novel position-specific scoring scheme and a 87 weighting scheme for down weighting over-represented 88 sequence groups, is used for multiple sequence alignment

and phylogenetic analysis, with the possibility to draw 89 90 phylogenetic trees. The software BioEdit was used to manipulate the alignment for a clearer sight of differences 91 between the aligned sequences. BioEdit software is a user-92 friendly tool for post-alignment modifications, with the 93 possibility to emphasize few sequences, cut and paste 94 them, and make other evaluations, such as restriction 95 mapping. The BKV VP1 sequences were then used for 96 Phylogenetic analyses. Clustal X, that was utilized to make 97 a neighbor-joining phylogenetic tree, and the free software 98 99 NJplot were employed to display the resulting tree. The confidence of branching patterns of the neighbor-joining 100 tree was determined by bootstrap analysis, using a set of 101 1,000 replicates. 102

Primers and Probes Design

The alignment was screened to find "hot spots" of muta-104 tion specific for all the strains or isolates belonging to each 105 genotype (Fig. 1). In the absence of a specific single 106 nucleotide mutation, the occurrence of a different nucleo-107 tide combination between the genotypes was considered. 108 Every mutation was evaluated as a possible target for 109 designing of a set of primers able to recognize all the BKV 110 strains belonging to a specific genotype and to discriminate 111 between the others by a mismatch at the 3' primer ending. 112 The parameters that were taken into consideration for 113 selecting the "hot spots" and discriminating the primers 114 design were the presence of a unique sequence shared 115 between the strains belonging to one genotype but not to 116 the others, the 60°C melting temperature, and the possi-117 bility to design a Real-Time PCR probe in the most con-118 served nucleotide region inside the hypothetic amplicon. 119

Primers and probes were designed with the help of the 120 software Primer Express[®] v3.0 (Applied Biosystem, 121 Cheshire, UK). The primers obtained were then analyzed 122 with the Autodimer Software, an open-source tool that 123 evaluates the hairpin and primer-dimer formation, with the 124 aim of excluding the presence of factors that could alter the 125 126 amplification efficiency. In order to include all the strains

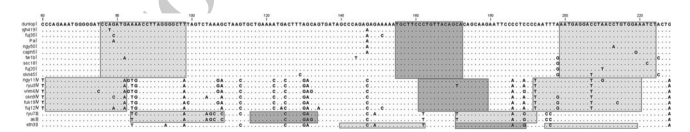


Fig. 1 Regions considered for the primer and probe design. Identical residues are represented as dots. Light squares indicate the primer regions. Dark squares represent the probe regions



•	Journal : Large 12033	Dispatch : 16-2-2011	Pages : 8
	Article No. : 9386	□ LE	□ TYPESET
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BK I\$'-CCAGATGAAACCTTAGGGGGCTT-3'\$'-CCTGATGAAACCTTAGGGGGCTT-3'\$'-CCAATGAAAACCTTAGGGGGCTT-3'\$'-CCAATGAAAACCTTAGGGGGCTT-3'\$'-CCAGATAAAAACCTTAGGGGGCTT-3'BK II\$'-GCCAGACAAAAAACCTTAGGGGGCTT-3'BK III\$'-GCCAGACAAAAAATGCTTCCT-3'BK III\$'-CCAGAAATGGGGGGTATAGTCCAGATA-3'\$'-CCAGAAATGGGGGGATCCAGATA-3'\$'-CCAGAAATGGGGGGATCCAGATA-3'\$'-CCAGAAATGGGGGGATCCAAATA-3'\$'-CCAGAAATGGGGGGATCCAAATA-3'\$'-CCAGAAATGGGGGGATCCAAATA-3'\$'-CCAGAAATGGGGGGATCCACAATA-3'\$'-CCAGAAATGGGGGGATCCACAATA-3'		Reverse primers	Probes
S	GGGCTT-3′	5'-AGATTTCCACAGGTTAGGTCCTCATT-3'	5'-FAM-TGCTTCCCTGTTACAGCA-MGB-3'
5	3GGCTT-3/	5'-AGATTTCCACATGTTAGGTCCTCATT-3'	
	3GGCTT-3'	5'-AGATTCCCACAGGTTAGGTCCTCATT-3'	
	GGGCTT-3'	5'-AGATTTCCACAGGTTAGATCCTCATT-3'	
-	GGGCTT-3'	5'-AGGTTTCCACAGGTTAGGTCCTCATT-3'	
		5'-AGATTTCCACAGGTAAGGTCCTCATT-3'	
	CTTCCT-3'	5'-TCCACAGGTTAGGTCCTCATTTAGA-3'	5'-FAM-CACAGCAAGAATTCCACTG-MGB-3'
	GTCAGCAC-3/	5'-GGGCAGTGGAATTCTTGCTGTA-3'	5'-FAM-CTGAAAATGCCTTTGASA-MGB-3'
5'-CCAGAAATGGGGGGACCCAGATA-3' 5'-CCAGAAATGGGGGGATCCAAATA-3' 5'-CCAGAAATGGGGGGATCCACATA-3'	AGATA-3'	5'-TTCCACAGGTTAGATCCTCATTTAAATTA-3'	5'-FAM-CTGTTACAGYACAGCAAG-MGB-3'
5'-CCAGAAATGGGGGGATCCAAATA-3' 5'-CCAGAAATGGGGGGATCCACATA-3'	AGATA-3'	5'-TTCCACAGGTTAGATCCTCATTCAAATTA-3'	
5'-CCAGAAATGGGGGGATCCACATA-3'	AAATA-3'	5'-TTCCACAGGTCAGATCCTCATTTAAATTA-3'	
	ACATA-3'	5'-TTCCACGGGTTAGATCCTCATTCAAATTA-3'	
CLONING 5'-GACAATCACATGCCTGGATAATG-3'	ATAATG-3'	5'-TGGTAATGGACTAAGTGTGCGTTATTTTC-3'	

belonging to one genotype, many pairs of primers were designed and produced. Four Taqman[®] probes MGB 128 (minor groove binding) labeled at the 5' end with FAM 129 (6-carboxyfluorescein), each being able to recognize one of the four target regions amplified by the different pairs 131 of primers, were also designed. The list and sequences of primers and probes are reported in Table 1. 133

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Real-Time PCR Optimization

Four different mixes for the amplification of each genotype 135 were prepared. Different concentrations of primers and 136 probe were evaluated: in particular, 200/200, 200/100, 137 100/200, 200/50, and 50/200 nM for primer forward/ 138 reverse, and 250 or 100 nM for probe. The reaction mix-139 ture contained 1× Master Mix (Platinum qPCR super-140 mix-UDG with ROX [Invitrogen, Carlsbad, CA]). For 141 this evaluation, isolates from BKV I (Dunlop strain) and 142 BKV IV (kindly provided by Dr. Andi Krumbholz and Dr. 143 Ellen Krautkramer), and ultramers that simulated BKV II 144 145 and III sequences (comprehending a portion of isolates ETH3 and NEA27, respectively [Tema Ricerca, Bologna, 146 Italy]), were used. Fifteen microliters of amplification mix 147 were added to 5 µl of BKV DNA, obtaining a final volume 148 of 20 µl. The procedure was performed and optimized on 149 the 7300 Real-Time PCR System (Applied Biosystems). 150 The following thermal profile was used: one cycle of 151 decontamination at 50°C for 2 min, one cycle of denatur-152 ation at 95°C for 10 min, and followed by 45 cycles of 153 amplification at: 95°C for 15 s, and 60°C for 60 s. The 154 following parameters were considered for the evaluation: 155 quick observation of the amplification, and good discrim-156 ination between the specific and aspecific amplification. 157

Plasmids and Standards

The BKV I Dunlop standard plasmid was kindly provided 159 by Prof. Tiziana Musso. The BKV II and BKV III standard 160 plasmids were produced starting from the ultramers (Tema 161 Ricerca), then amplified with cloning primers (Table 1). 162 The BKV IV standard plasmid was prepared starting from 163 the BKV IV isolate J/2296/04 (kindly provided by Dr. Andi 164 Krumbholz). The primers were used for producing a PCR 165 product, then cloned using the pTOPO-TA cloning system 166 (Invitrogen) and propagated in competent Escherichia coli 167 TOP10 cells. After overnight culture, selected transformed 168 clones were amplified by culture in liquid LB medium 169 (10 g BACTOTRYPTONE, 5 g yeast extract, 171 mM 170 NaCl, and 15 g/l agar, pH 7.5) containing 50 µg/ml of 171 172 ampicillin. Plasmid DNA was extracted and purified using the Wizard Plus SV Miniprep DNA Purification System 173 (Promega, Madison, WI). The plasmid concentration was 174 175 estimated on spectrophotometric reading at OD260.

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•	Journal : Large 12033	Dispatch : 16-2-2011	Pages : 8
	Article No. : 9386	□ LE	□ TYPESET
•	MS Code : MOBI-D-11-00032	🖌 СР	🖌 disk

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176 *Real-Time PCR Evaluation (Linearity, Efficiency,*177 *Interference Limit, Limit of Detection, Precision)*

178 The four genotyping assays were evaluated using serial 179 10-fold dilutions of the target plasmid (ranging from 10^7 to 180 10 copies per reaction). The linearity was assessed by the 181 correlation coefficient (R^2) of the standard curve obtained by 182 plotting 10^7 -10 standard copies per reaction with the four 183 different tests. The efficiency was evaluated by the slope of 184 the standard curve, using the formula $E = 10^{(-1/slope)}$ -1.

In order to discriminate between specific and aspecific amplifications, three repetitions of 10^7 , 10^5 , and 10^3 , and 10copies per reaction of the four standards were amplified with the four different genotyping mixes. The average threshold cycle (*C*t) values of specific and aspecific amplifications were obtained. The background *C*t was evaluated by the formula $Ct_{\text{lim}} = Ct_a - t_{0.01} \cdot \sigma_a / N$, where *N* is the number of observations, and $t_{0.01}$ is the tabulated value of Student's *t* for the 1% probability level and *N*-1 degree of freedom [16]. For the evaluation of the lowest genotype concentration that could be distinguished from the aspecific amplification of the other three genotypes, the formula $IL = 100/(2^{(Ct_s - Ct_{\text{lim}})} +$ 1 was used, where *C*t_s represents the specific *C*t [17].

198 The limit of detection, defined as the lowest target 199 quantity detectable, was also estimated by serial 10-fold 200 standard dilutions. The precision of each test was calcu-201 lated by the intra- and inter-test coefficients of variation, 202 according to the formula $CV = 100 \bullet (\sigma/MC)$, where MC is the average measured concentration, and σ the correspon-203 204 dent standard deviation. Three repetitions for each standard 205 dilution were considered in the same run or in different 206 experiments, for the intra- and the inter-assay coefficients 207 of variation, respectively.

208 Clinical Specimens

Twenty urine specimens resulting positive to a reference 209 210 amplification assay (BKV Q-PCR Alert Kit [Nanogen, 211 Buttigliera Alta, Turin, Italy]) were tested with the four 212 amplification assays to assess the genotyping procedure. 213 Automated DNA extraction was performed with the NucliSens EasyMAG platform (bioMeriéux, Marcy l'Eto-214 215 ile, France) from 1 ml of urine, according to the manu-216 facturer's instructions, and eluted to a final volume of 217 50 µl. Five microliters of extracted specimen were then 218 added to 15 µl of each genotyping mix. The results were 219 analyzed using the System SDS software.

220 Sequencing Procedure

To confirm the results obtained by the Real-Time ampli-fication, sequencing procedure was performed. The 20

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specimens were amplified with primers BKV-1 (5'-GAA 223 GTT CTA GAA GTT AAA ACT GGG-3') and BKV-2 224 (5'-GTG GAA ATT ACT GCC TTG AAT AGG-3') [9], 225 obtaining an amplicon of 354 nucleotides within the VP1 226 gene (from 1663 to 2016, Dunlop reference numbering). 227 The amplification mix contained 6 µl of GoTaq[®] HotStart 228 Polymerase buffer $5 \times$ (Promega), 200 μ M of each dNTP, 229 6 mM of MgCl2. 1 unit of GoTaq[®] HotStart Polymerase 230 (Promega), and 20 pmol of BKV-1 and BKV-2 primers, 231 respectively. The resulting amplicons were run on agarose 232 gel (2% w/v) by electrophoresis, the gel was observed on 233 an UV transilluminator UV and the bands cut. Subse-234 quently, the bands were purified using the Nucleospin[®] 235 Extract II (Macherev-Nagel, Düren, Germany). The puri-236 fied PCR products were sequenced using the BigDye[®] 237 Terminator v1.1 Cycle Sequencing Kit (Applied Biosys-238 tem). In brief, a mix containing 8 µl of Terminator Ready 239 reaction mix, 2 µl of PCR product, 3.2 pmol of inner pri-240 mer (BKV-1 or BKV-2), and deionized water to obtain 241 20 µl final volume was prepared. Then, the Cycle 242 sequencing was carried out on 9800 Fast Thermal Cycler 243 (Applied Biosystem) with an initial denaturation step at 244 96°C for 1 min, followed by 25 cycles of 96°C for 10 s, 245 50°C for 5 s, and 60°C for 4 min. 246

The Cycle sequencing product was then purified by247Ethanol/EDTA/Sodium Acetate precipitation and loaded248onto the ABI PRISM 31 Genetic Analyzer (Applied Bio-
system). Forward and reverse sequences were aligned with
the ClustalX sotware and then compared to BKV
sequences.251

Results

254 The best amplification was obtained with the following 255 concentrations of primers forward/reverse: 100/200, 200/200, 200/200 and 200/100 nM for genotypes I, II, III, 256 and IV, respectively, and with a probe concentration of 257 250 nM; however, the amplification efficiencies were 104, 258 104, 97, and 88%, respectively. The R^2 coefficient was 259 0.996, 0.999, 0.995, and 0.999 for BK I, II, III, and IV 260 genotype amplifications, respectively. 261

Subsequently, the Real-Time PCRs were tested for the 262 discrimination of the four genotypes. Three repetitions of 263 10^7 , 10^5 , and 10^3 , and 10 copies/reaction of the target 264 sequence of the main four BKV genotypes were employed to 265 compare the amplification and distinguish between specific 266 and aspecific amplifications using the four mixes (Table 2). 267 268 As regards the BKV I genotype mix, this resulting product was able to amplify the genotype I-specific target, and only 269 BKV II and IV genotype mixes cross-recognize the specific 270 genotype I target at 10^7 copies/reaction (both), and at 10^5 271 copies/reaction (only the BKV II genotype mix). However, a 272

Journal : Large 12033	Dispatch : 16-2-2011	Pages : 8
Article No. : 9386	□ LE	□ TYPESET
MS Code : MOBI-D-11-00032	🗹 СР	🖌 disk

	Specific mean Ct	Aspec	ific mean Ct	
	BK I	ВК П		BK III BK IV
10 ⁷	17.36 (0.07)	32.16	(0.34) [32.06]	
10 ⁵	24.12 (0.07)	38.8 (0.16) [38.75]	38.13
10^{3}	30.9 (0.51)			
10	36.12			
	BK II	BK I	BK III	BK IV
10 ⁷	16.85 (0.04)		36.57 (0.62) [36.39]	32.2 (0.36) [32.10]
10 ⁵	23.21 (0.04)			35.94 (0.65) [35.75]
10^{3}	29.97 (0.19)	36.14	37.74	36.51 (0.94) [36.24]
10	36.03		40.89	37.15
	BK III	BK I	BK II	BK IV
10 ⁷	17.78 (0.16)			35.62 (0.9) [35.36]
10 ⁵	24.2 (0.16)			
10^{3}	30.84 (0.03)			
10	37.62 (0.63)		36.91	38.61 (0.74) [38.40]
	BK IV	BK I	BK II	BK III
10 ⁷	17.67 (0.03)		36.12 (0.81) [35.89]	35.88 (0.59) [35.71]
10 ⁵	24.74 (0.01)			
10^{3}	32.3 (0.19)	36.78		
10				

Table 2 Aspecific and specific means Ct for each genotyping mix

The values in the round brackets are the standard deviations of the repetitions for each dilution; in the square brackets the Ct_{lim} are reported (see text for explanation)

273 gap of approximately 14.8–14.68 Ct_s separated the specific
274 and aspecific amplifications; moreover, no aspecific ampli275 fication was evidenced for lower dilution.

BKV II genotype mix that resulted was able to amplify 276 277 the specific BKV II target sequence;' however, this was 278 also amplified by the other three BKV discriminating 279 mixes. Only the amplification of the BKV II-specific target 280 with the corresponding mix was linear, and the threshold 281 crossing appeared at least 15 cycles earlier for BKV II mix at 10^7 copies/reaction compared to the other amplification 282 mixes, decreasing to six cycles at 10^3 copies/reaction. The 283 amplification with BKV II mix at lower BKV II genotype-284 specific target concentration was considered not discrimi-285 286 nating, because of the overlapping Ct with BKV III and IV amplification mixes. The BKV III target was recognized by 287 288 the corresponding mix with a linear amplification; how-289 ever, 10⁷ copies/reaction were recognized also by the BKV I mix, although a mean gap of 17.84 Ct_s was observed 290 291 between specific and aspecific amplifications. An undis-292 tinguishable specific amplification was seen with a con-293 centration of BKV III of 10 copies/reaction, because of the 294 simultaneous amplification with BKV II, III, and IV mixes 295 with a similar Ct. The BKV IV target showed a specific recognition using the corresponding mix, although the 296 target was also amplified with BKV II and III mixes at 10^7 copies/reaction, with a mean gap of 18.21 and 18.45 Ct_s 298 between specific and aspecific amplifications for BKV II 299 and III mixes, respectively. However, at a concentration of 10^3 copies/reaction, the target was also recognized by BKV 301 I mix with a mean gap of 4.48 mean Ct_s. 302

Using the same data to evaluate the specific and aspecific amplifications, a Ct_{lim} was calculated for each genotyping amplification, as an interference limit (Table 3). The precision was calculated with the intra- and inter-test coefficients of variation (Table 4). 307

In order to establish the limit of detection for the four dif-308 ferent genotyping amplifications, 10-fold dilutions of the 309 targets (ranging from 10^7 to 10 copies/reaction) were ampli-310 fied with the above four methods. The limit of detection dif-311 fered between the four genotyping amplification procedures 312 and was as follows: 10² copies/reaction for BKV IV and 10 313 copies/reaction for BKV I, II, and III, each. However, only 314 BKV III procedure showed a sensitivity of 10 copies/reaction, 315 while for the others assay, it was of 100 copies/reaction. 316

Results for the clinical specimens are reported in Table 5. 317 In brief, the amplification procedures yielded positive results 318

•••	Journal : Large 12033	Dispatch : 16-2-2011	Pages : 8
	Article No. : 9386	□ LE	□ TYPESET
·	MS Code : MOBI-D-11-00032	🗹 СР	🖌 disk

 Table 3 Interference limit

	107 (%)	10 ⁵ (%)	10 ³ (%)	10 (%)
I–II	0.00375	0.00393		
II–III	0.00013			
II–IV	0.00257	0.01674	1.27874	
III–IV	0.00051			36.83754
IV->II	0.00033			
IV–III	0.00037			

The lowest limit of quantity of BKV-specific genotype, expressed in percentage, distinguishable from the background (aspecific amplification). In the case of no reported percentage, sporadic or no background amplification was observed

Table 4 Intra- and inter-assay coefficients of variation for dilutions of 10^7 , 10^5 , and 10^3 , and 10 copies/reaction

	10 ⁷ (%)	10 ⁵ (%)	10 ³ (%)	10 (%)
Coeffici	ent of intra-assay	variation		
Ι	0.40	0.29	1.65	
II	0.23	0.17	0.63	
III	0.89	0.66	0.10	1.67
IV	0.16	0.04	0.59	
Coeffici	ent of inter-assay	variation		
Ι	0.92	2.70	0.84	1.85
II	1.19	0.15	1.45	0.99
III	1.30	1.51	0.91	2.92
IV	3.34	2.20	2.26	

in 16 cases; in particular: 12716 BKV I, 1/16 BKV II, 1716
BKV III, and 1/16 BKV IV. The DNA sequencing confirmed
the results obtained with the Real-Time PCR assays for all
the samples. One specimen (#4464) showed a concomitant
amplification for BKV I and BKV IV, with overlapping Ct_s;
sequencing identified it as BKV I genotype.

325 In most of the cases, the viral load found with the devel-326 oped methods differed from that obtained with the reference 327 method, with a percentage of variation ranging from -288.4328 to 89.9% (Table 5). Furthermore, more aspecific amplifica-329 tions were observed in clinical specimens. In fact, the sample #2998 showed two aspecific plots (BKV I and BKV II), while 330 331 no BKV I amplification was observed with a 10⁷ copies/ 332 reaction plasmid dilution. Nevertheless, the occurrence of 333 aspecific amplifications did not impact on the genotyping 334 performance of the Real-Time PCRs, as the gap between the 335 curves of amplifications remained wide.

336 Discussion

In this study, a Real-Time PCR-based method for geno-typing of polyomavirus BK was developed. By

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hemagglutination inhibition tests, four main BKV sero-339 340 types were described [5], based on the differences in a short amino acidic sequence on VP1 protein, probably being 341 responsible for the serotypical differences among BKV 342 variants [18]. Correspondingly, four main BKV VP1 343 genotypes were found, and their nucleotide sequences 344 analyzed [18]. On the basis of this conserved region, 379 345 VP1 sequences were collected from three different articles 346 [8-10] investigating the prevalence of BKV subtypes in 347 348 different countries, including Germany, Japan, and China. 349 These sequences were aligned to determine which nucleotide residues were conserved within each genotype with 350 the aim to design a Real-time PCR assay based on the 351 principle of the Amplification Refractory Mutation Assay 352 (ARMS). 353

A complete list of single nucleotide polymorphisms in
the total BKV genome for each genotype has been reported354[19]. The 379 aligned VP1 sequences presented concordant
residues with those previously listed by Luo and colleagues356[19], confirming the uniqueness and conservation of these
polymorphisms among the genotypes.359

360 It has been reported [19] that genotyping on large T antigen (LTA) is preferable, in particular, because LTA is a 361 larger region with more informative sites compared to the 362 327 bp of VP1 region considered by Jin and colleagues [6]. 363 However, in this study, we chose to use the VP1 region 364 instead of LTA as we aimed to produce powerful dis-365 criminating Real-Time PCRs taking into consideration the 366 data published in the literature. Indeed, a higher number of 367 sequences is banked in public domain servers for VP1 368 369 compared to LTA, thus permitting a more detailed primersand-probes design for the detection of a higher number of 370 sequences and corresponding polymorphic residues. In this 371 study, four Real-Time PCR assays, each one that can 372 amplify specifically a specific genotype, were designed, 373 and four different amplification mixes were produced to 374 avoid a too high concentration of primers in a single well, 375 that could determine dimer formation or an increase in the 376 377 aspecific performance. In fact, in the presence of high viral 378 loads, aspecific amplifications might appear, and these 379 were detectable as a low quantity amplification with other genotype mixes; anyway, the aspecific amplification was 380 easily recognizable by the wide gap between the specific 381 and aspecific plots. Moreover, the introduction of Ctlim and 382 interference limit gave further clues about discrimination. 383 Clearer results were available for $10^7 - 10^5$ copies/reaction 384 dilutions, while 10^3 caused the lowest dilution with an 385 acceptable genotyping performance. For this reason, a 386 result comparable to 10^3 was set as genotyping limit for all 387 the four Real-Time PCRs, and all those results with a lower 388 quantity were not accepted. 389

As regards the clinical validation of the four genotyping 390 assays, the distribution of BKV genotypes was evaluated in 391

Journal : Large 12033	Dispatch : 16-2-2011	Pages : 8
Article No. : 9386	□ LE	□ TYPESET
MS Code : MOBI-D-11-00032	🗹 СР	🖌 DISK

N°	ID	Reference copies/ml	BKV I (Ct)	BKV II (Ct)	BKV III (Ct)	BKV IV (Ct)	Genotype	Copies/ml
1	2998	$>5 \times 10^{6}$	31.63	33.82		11.59	IV	$>5 \times 10^{7}$
2	3068	4237203	19.57	36.36		32.78	Ι	$>5 \times 10^{7}$
3	3204	569						
4	3206	$>5 \times 10^{6}$	12.01	26.65		25.34	Ι	$>5 \times 10^{7}$
5	3207	2413	30.60				Ι	5379 (-122.9%)
6	3607	126470	29.38	36.56			Ι	12800 (89.9%)
7	3619	1542						
8	3965	502131	24.72	43.06		38.04	I	351153 (30.1%)
9	4011	868						
10	4063	4158	30.14				Ι	7458 (-79.4%)
11	4326	495	34.78				Т	<500
12	4398	49606	29.73	43.09		38.53	Ι	9981 (79.9%)
13	4405	2472943	34.93	19.80		32.37	П	1220410 (50.6%)
14	4464	8689	36.59			36.69	Ι	<500
15	4674	6441	29.22	42.79		42.64	I	14342 (-122.7%)
16	4712	2393	30.58			37.65	I	5456 (-128%)
17	4714	<59	36.17		4		Ι	<500
18	4723	$>5 \times 10^{6}$	12.19	28.37		26.11	Ι	$>5 \times 10^{7}$
19	4758	<59				Y		
20	4768	44562		31.50	29.14		III	173076 (-288.4%)

Table 5 BK genotyping in clinical specimens

Viral loads (expressed in copies/ml) of positives samples for the reference method, positivity to each amplification procedure (expressed in Ct), genotype and quantities (expressed in copies/ml) are reported. *Bold* numbers shows Ct_s of overlapping curves

392 urine specimens; this type of sample was chosen for the 393 higher mean viral load in comparison to serum samples [2]. 394 However, clinical samples with a low reference viral load were not always detected. Most of the specimens were 395 396 infected by BKV I (80%), while BKV II, III, and IV were 397 infrequent. The prevalence of the four genotypes was 398 concordant with the literature [20]. BKV IV, that is more 399 prevalent in East Asia [8], is found also in Europe [9] in a 400 percentage comparable to the one observed by us; however, 401 notwithstanding the uncommon prevalence, both BKV II 402 and BKV III were detected in two different samples; the 403 results were confirmed by the subsequent VP1 DNA 404 sequencing.

405 Considering viral load of different genotypes, a high divergence was observed between the genotyping assays 406 and the reference method, with a variability ranging from 407 408 -288.4 to 89.9%. These data highlighted the problem 409 exposed by Hoffman and coll. [21]: in the presence of various genotypes, difference in probes and primers for the 410 411 BKV quantification led to a substantial disagreement 412 between assays. This was confirmed in our study, between 413 the reference and the genotyping amplifications.

In conclusion, the four Real-Time PCRs that were produced were able to detect and discriminate the main four
BKV genotypes. Compared to classical DNA sequencing,

417 these procedures are more rapid and simpler, since they are single PCR assays with no purification steps; results are 418 available in approximately 2 h. One drawback could arise 419 because of the need of at least 10³ copies/reaction for the 420 genotyping; in these cases, a PCR assay employing exter-421 nal primers, or other protocols, would be necessary to 422 increase the target sequence amplifiable, although this 423 should also increase the time required for genotyping. 424

The comparison between genotyping procedures on 425 clinical samples and plasmid dilutions evidenced some 426 differences, including the detection of aspecific plots dur-427 ing the testing of clinical samples, and the lack of similarity 428 with the quantities reported by the reference amplification 429 procedure. This could be caused by the primers-and-probes 430 nature, amplifying Large T (reference method) and VP1, 431 432 and by the presence of polymorphism that alter the efficiency of amplification, as previously suggested [21]. 433 Nevertheless, the four Real-Time PCRs developed in this 434 study should not be considered quantitative methods, as the 435 436 procedures were not tested for the quantification of a broad number of isolates. The availability of a rapid and simple 437 438 genotyping test could be useful for the evaluation of BKV genotypes' prevalence. and studies on the impact of the 439 infecting genotype on viral biological behavior, pathogenic 440 role, and immune evasion strategies. 441

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