

1 Novel human polyomaviruses in pregnancy: higher prevalence of BKPyV, but no WUPyV,
2 KIPyV and HPyV9

3
4 Eszter Csoma ^{a,*}, Tamás Sáy ^b, Beáta Mészáros ^a, Lajos Gergely ^a

5
6 ^a Department of Medical Microbiology, Medical and Health Science Centre, University of
7 Debrecen, Nagyerdei krt. 98., H-4032 Debrecen, Hungary

8 ^b Department of Obstetrics and Gynecology, Medical and Health Science Centre, University
9 of Debrecen, Nagyerdei krt. 98., H-4032 Debrecen, Hungary

10
11 * Corresponding author. Tel.: +36 52 255 425; fax: +36 52 255 424. Email address:
12 csomae@freemail.hu.

13
14 Abbreviations:

15 WU polyomavirus (WUPyV), KI polyomavirus (KIPyV), human polyomavirus 9 (HPyV9),
16 BK polyomavirus (BKPyV), genome equivalent (GEq), polymerase chain reaction (PCR)

17
18
19
20
21
22 Number of words in the Abstract: 248

23 Number of words in the text: 1879

26 **Abstract**

27 Background: Immunosuppression due to pregnancy may lead to higher susceptibility to
28 infections and reactivation of latent infections, such as BK polyomavirus (BKPyV). There is
29 lack of information about the prevalence of novel human polyomavirus 9 (HPyV9), WU
30 (WUPyV) and KI (KIPyV) during pregnancy.

31 Objectives: To study whether pregnancy results in higher prevalence of HPyV9, WUPyV,
32 KIPyV and their correlation with BKPyV.

33 Study design: Plasma, urine and throat swab samples from 100 pregnant and 100 non
34 pregnant women were screened for the presence of WUPyV, KIPyV, HPyV9 and BKPyV by
35 PCR.

36 Results: No WUPyV DNA was detected in plasma, urine and respiratory samples from
37 pregnant and non pregnant women. KIPyV DNA was found in two plasma samples from non
38 pregnant women (2 %) and not detected in other samples from neither pregnant nor non
39 pregnant women. HPyV9 DNA was determined in all sample types of pregnant and non
40 pregnant women, respectively. There were no significant differences between pregnant and
41 non pregnant women in HPyV9 DNA frequencies for plasma (2 % vs. 6 %), urine (3 % vs. 2
42 %) and respiratory samples (2 % vs. 2 %). Prevalence of BKPyV in urine samples was
43 significantly higher ($p=0.039$) in pregnant women (13 %) than in non pregnant women (4 %);
44 co nfection with KIPyV and/or HPyV9 was not detected.

45 Conclusions: In contrast with BKPyV, infection with WUPyV, KIPyV and HPyV9 was not
46 detected more frequently during pregnancy. To our knowledge HPyV9 was detected first in
47 respiratory samples in our study.

48

49 Key words: human polyomaviruses, pregnancy

50

51 **1. Background**

52 Human polyomavirus BK (BKPyV) seroprevalence increases with age reaching high,
53 80-90 % in adult population.¹ Similarly high, 55-90 % adult seropositivities were observed for
54 recently discovered KI² and WU³ polyomaviruses (KIPyV, WUPyV).⁴⁻⁶ Investigation of
55 seropositivity against the newly discovered human polyomavirus 9 (HPyV9)⁷ revealed 47 %
56 positivity for healthy adults.⁸ It is well known that after the childhood primary infection with
57 BKPyV, lifelong persistent infection is established mainly in renal and urinary tract cells.¹
58 Transient immunosuppression due to pregnancy may lead to reactivation of BKPyV resulting
59 in generally asymptomatic viruria with frequency of 3 to 54 %.^{1, 9-11} Beside viruria, BKPyV
60 viraemia was also detected in pregnant women.¹¹ The pathogenic role of the novel WUPyV,
61 KIPyV and HPyV9 is far from clear, only speculative. WU and KI viruses were found in
62 various sample types – respiratory samples, blood, faeces, cerebrospinal fluid, lymphoid
63 tissues, urine – and higher prevalence was observed in children and immunocompromised
64 patients.¹²⁻¹⁶ HPyV9 was described from blood and urine samples of kidney transplant
65 patients, then it was found in skin samples, but no in respiratory and fecal samples.^{7, 17} The
66 higher frequency of these viruses in immunocompromised patients suggests higher
67 susceptibility or reactivation due to immunosuppression. Up to now only four urine samples
68 from pregnant women were investigated for the presence of KIPyV and WUPyV DNA with
69 negative result.¹⁸ The genetic and possible transmission similarities to BKPyV, and the higher
70 PCR prevalence data among immunocompromised patients may suggest that
71 immunosuppression, thus pregnancy may lead to higher susceptibility to infection with
72 WUPyV, KIPyV and HPyV9 or may result in reactivation of possible latent infections.

73

74 **2. Objective**

75 The aim of the present study was to evaluate the prevalence of three new human
76 polyomaviruses (WUPyV, KIPyV and HPyV9) during pregnancy, to study whether
77 immunosuppression due to pregnancy may lead to higher prevalence as it was found in case
78 of BKPyV. The possible correlations of these viruses were also investigated.

79

80 **3. Study design**

81 *3.1. Patients and samples*

82 Urine, plasma (from EDTA blood samples) and throat swab samples were collected on
83 the same day from 100 healthy pregnant women (age 16.5-41.9 years, median 32.1 years;
84 pregnancy 5-39 weeks; median 26 weeks) and 100 non pregnant women (age 18-44.3 years,
85 median 31.6 years) between September 2011 and December 2011. Samples from pregnant
86 women were collected in all three trimesters: first trimester n=28; second trimester n=27;
87 third trimester n=45. The control samples were taken from healthy, non pregnant, fertility
88 exam visitor women.

89 Immediately after collection, nucleic acid was isolated from samples using High Pure
90 Viral Nucleic Acid Kit (Roche, Switzerland) according to the manufacturer's instructions.
91 Briefly, nucleic acids from 200 µl plasma, 200 µl urine specimen and throat swab sample
92 washed in 200 µl buffer were eluted in 50 µl and stored at -20 °C until use.

93 The study was approved by Regional and Institutional Ethics Committee of
94 University of Debrecen. All patients were asked to sign written informed consent.

95

96 *3.2. Nested and real-time PCR for WUPyV, KIPyV, HPyV9 and BKPyV*

97 All PCR methods were carried out with 10 µl nucleic acid in a final volume of 25 µl.
98 For nested PCR AmpliTaq Gold 360 Master Mix, for WUPyV and KIPyV real-time PCR
99 TaqMan Universal PCR Master Mix (Applied Biosystems, USA) were used. The calibrants

100 for quantitative PCRs were serial dilutions of KIPyV plasmid (in which the genome of KI
101 polyomavirus isolate Stockholm 60 was incorporated) and AP-p003 plasmid (containing the
102 2228 bp half genome of WU polyomavirus) kindly provided by Tobias Allander and David
103 Wang. WUKI nested PCR and real-time PCR for WU and KI virus were performed as
104 described previously.¹⁶ HPyV9 PCR was carried out with diagnostic primers and annealing
105 temperature published by Scuda et al.⁷ For the first round of BKV nested PCR, k1 (5'
106 TGAAGCATATGAAGATGGCC 3') and k2 (5' GTTACAGCCTCCCACATC 3') primers
107 were used with 60 °C annealing temperature, while for the second round b1 (5'
108 GATGGCCCCAACCAAAAG 3') and b2 (5' CTAGAACTTCTACTCCTCC 3') primers and
109 56 °C annealing temperature were applied. PCR products were visualized by electrophoresis
110 in 1.5 % agarose gel containing ethidium bromide (0.5 µg/mL). The amplified PCR products
111 from WUKI and HPyV9 nested PCR were cut, purified with QIAquick Gel Extraction Kit
112 (Qiagen) according to the instructions and sequenced by using ABI PRISM 3100 Genetic
113 Analyzer (Applied Biosystems). To determine BKPyV viral load BKV virus R-gene
114 quantification kit was used (Argene, USA) according to the manufacturer's instructions.

115

116 3.3. Statistical analysis

117 Difference in frequency for categorical variables was analysed by Fisher's exact test.
118 For continuous variables Mann-Whitney U test was applied. Difference was considered
119 significant if p value was less than 0.05.

120

121 4. Results

122 4.1. Detection of WUPyV, KIPyV and HPyV9 DNA in plasma, urine and respiratory samples

123 Table 1 shows the results of PCR detections for the various samples. WUPyV DNA
124 was not detected in plasma, urine and respiratory samples neither from pregnant nor from non

125 pregnant women. KIPyV was found in two plasma samples of non pregnant women, but was
126 not determined in any other samples. To confirm the positive PCR results and to determine KI
127 or WU virus DNA was detected, PCR products were sequenced. The viral loads were below
128 the limit of detection (< 250 GEq/mL; genome equivalent/mL) by real-time PCR. HPyV9
129 DNA was detected in urine, plasma and respiratory samples from both studied groups. To
130 prove the results from PCR, all PCR products were sequenced. In details, the prevalence of
131 HPyV9 DNA in plasma samples was higher in control, non pregnant group than in pregnant
132 women (6/100; 6 % vs. 2/100; 2%), but the difference was not statistically significant. The
133 two positive samples were taken in the second trimester of pregnancy. Two samples from
134 control, non pregnant women with HPyV9 viraemia were also positive for KIPyV DNA. In
135 respiratory samples the frequency of HPyV9 DNA was the same in both studied groups
136 (2/100; 2% and 2/100; 2%). Both of the positive samples in pregnant women group were
137 collected in the first trimester. Three urine samples from pregnant women were HPyV9 PCR
138 positive (3/100; 3%), while in control group 2 samples were positive (2/100; 2%) which is not
139 statistically significant difference.

140 *4.2. Prevalence of BKPyV in urine and plasma samples*

141 BKPyV was not detected in plasma samples. Frequency for BKPyV viruria was 13 %
142 (13/100) in pregnant women and 4 % (4/100) in non pregnant, control group (Table 1.). The
143 difference is statistically significant ($p=0.039$). The BKPyV viral load in samples from
144 pregnant women (range $50-1.86 \times 10^8$; median 11.82×10^3 GEq/mL) did not show statistically
145 significant difference from the viral load in control samples (range $2.25 \times 10^2-3.58 \times 10^5$;
146 median 2.98×10^2). BKPyV presence in urine samples was found in all trimesters.

147

148 **5. Discussion**

149 In our study significantly higher prevalence of BK viruria was observed in pregnant
150 women in contrast with non pregnant women. Human polyomavirus 9 was found in plasma,
151 urine and respiratory samples from pregnant women but not more frequently than in samples
152 from non pregnant women. WU and KI viruses were not detected in any of the studied
153 samples from pregnant women.

154 BK polyomavirus is ubiquitous in the human population, the primary infection
155 generally occurs during childhood without significant clinical consequences, respiratory
156 diseases might occur. Transmission of the viruses is not well clarified, but it is suggested that
157 these viruses are acquired mainly through respiratory, faecal-oral and urinary routes,
158 alternatively by blood transfusion and organ transplantation.¹ After the primary infection,
159 lifelong persistence of the virus is established mainly in kidney and urinary tract.^{19, 20} Lytic
160 infection with viruria occurs in 5-10 % of immunocompetent individuals²¹, but more
161 frequently in immunocompromised patients.²² During pregnancy immunologic changes
162 together with hormonal effects may result in viral infections, reactivations. Viruria was
163 detected for 3-54 % of pregnant women, while viraemia was found to be less frequent.^{1, 10, 11,}
164 ²³ In accordance with literature, in this study 13 % of pregnant women had active BKPyV
165 replication resulting in viruria, but no viraemia. The possible effect of BK virus replication
166 during pregnancy is not clarified. Although viral DNA was demonstrated in fetal tissues²⁴ the
167 hypothesis of transplacental transmission was not confirmed^{10, 11}. Recently serological
168 evidence for vertical transmission of BKPyV was published.⁹

169 Hitherto, there are no prevalence data about the novel WU, KI and human
170 polyomavirus 9 during pregnancy. Bofill-Mas et al. investigated 4 urine samples from
171 pregnant women, but WU and KI viruses were not found.¹⁸ Foetal tissues were also negative
172 for WU and KI viruses.²⁵ In this study WU and KI viruses were not found in urine, plasma
173 and respiratory samples collected during pregnancy. KIPyV DNA was detected in two plasma

174 samples, but not in urine and respiratory samples from control, non pregnant women. The
175 high, 55-90 % seropositivity in adult population, and the higher PCR prevalence in samples
176 from children suggest childhood primary KI and WU virus infection.^{6, 15} Viruses were found
177 with frequency 0.4-14 % in various samples types including respiratory samples, blood,
178 faeces, cerebrospinal fluid, lymphoid tissues and urine samples, with generally higher
179 frequency in immunocompromised patients. The possible way of transmission might be
180 respiratory and/or faecal-oral.^{2, 3, 12-16} The higher PCR prevalence data of
181 immunocompromised patients suggests that immunosuppression might result in reactivation
182 of these viruses, or might establish higher susceptibility to KIPyV and WUPyV infection.¹ It
183 was hypothesized that similarly to BKPyV, transient immunosuppression due to pregnancy
184 might result in higher frequency of WU and/or KI viral infections, but no evidence for it was
185 found during this study. However it is important to note, that it was not a follow up study,
186 samples were collected once randomly during pregnancy.

187 Human polyomavirus 9 was described in 2011.⁷ Up to now, viral DNA was found in
188 blood and urine samples from immunocompromised patients and skin samples, but neither in
189 respiratory samples from patients with respiratory failure nor in faeces from children with
190 gastroenteritis.^{7, 17} Based on these data and the recently published 47 % adulthood
191 seropositivity⁸, Van Ghelue et al. hypothesized that HPyV9 is less frequent in the human
192 population⁶. We found HPyV9 DNA in all studied samples from pregnant and non pregnant
193 women with frequency of 2-6 %. There was no or not statistically significant difference
194 between the PCR prevalence in the respiratory (2 vs. 2 %), urine (3 vs. 2%) and plasma
195 samples (2 vs. 6 %) between pregnant and non pregnant women. To our knowledge we
196 published first HPyV9 presence in respiratory samples which may suggest respiratory
197 transmission of this virus. In this study higher prevalence of HPyV9 was not found during
198 pregnancy, but the viral loads were not examined which might have been different. Since

199 mother to foetus transmission of polyomaviruses BK and JC are suggested, this way of
200 transmission cannot be excluded in case of the novel WUPyV, KIPyV and HPyV9. Even if this
201 study could not support evidence for higher susceptibility of infection by these viruses,
202 further, follow up study of pregnant women during the whole period of pregnancy might
203 answer this question.

204 In conclusion, KI and WU viruses were not found in urine, respiratory and blood
205 samples from pregnant women, while HPyV9 was detected in all sample types but with no
206 significantly higher frequency than it was observed for non pregnant women.

207

208 **Conflict of interest**

209 The authors have no conflict of interest.

210

211 **Acknowledgements**

212 We thank Tobias Allander from Karolinska Institute (Sweden), David Wang from
213 Washington University (USA) and Bernhard Ehlers from Robert Koch-Institut (Germany) for
214 providing KIPyV, WUPyV and HPyV9 plasmids, respectively. This work was supported by
215 grants from the Hungarian Scientific Research Found (OTKA 73145) and by the TÁMOP-
216 4.2.2/B-10/1-2010-0024 project which is co-financed by the European Union and the European
217 Social Fund.

218

219 **References**

- 220 1. Jiang M, Abend JR, Johnson SF, Imperiale MJ. The role of polyomaviruses in human
221 disease. *Virology* 2009; **384**(2):266-73.
- 222 2. Allander T, Andreasson K, Gupta S, Bjerkner A, Bogdanovic G, Persson MA, et al.
223 Identification of a third human polyomavirus. *J Virol* 2007; **81**(8):4130-6.

- 224 3. Gaynor AM, Nissen MD, Whiley DM, Mackay IM, Lambert SB, Wu G, et al.
225 Identification of a novel polyomavirus from patients with acute respiratory tract
226 infections. *PLoS Pathog* 2007; **3**(5):e64.
- 227 4. Neske F, Prifert C, Scheiner B, Ewald M, Schubert J, Opitz A, et al. High prevalence
228 of antibodies against polyomavirus WU, polyomavirus KI, and human bocavirus in
229 German blood donors. *BMC Infect Dis* 2010; **10**(215).
- 230 5. Nguyen NL, Le BM, Wang D. Serologic evidence of frequent human infection with
231 WU and KI polyomaviruses. *Emerg Infect Dis* 2009; **15**(8):1199-205.
- 232 6. Van Ghelue M, Khan MT, Ehlers B, Moens U. Genome analysis of the new human
233 polyomaviruses. *Rev Med Virol* 2012;
- 234 7. Scuda N, Hofmann J, Calvignac-Spencer S, Ruprecht K, Liman P, Kuhn J, et al. A
235 novel human polyomavirus closely related to the african green monkey-derived
236 lymphotropic polyomavirus. *J Virol* 2011; **85**(9):4586-90.
- 237 8. Trusch F, Klein M, Finsterbusch T, Kuhn J, Hofmann J, Ehlers B. Seroprevalence of
238 the human polyomavirus 9 (HPyV9) and cross-reactivity to the African green
239 monkey-derived lymphotropic polyomavirus (LPV). *J Gen Virol* 2012;
- 240 9. Boldorini R, Allegrini S, Miglio U, Paganotti A, Cocca N, Zaffaroni M, et al.
241 Serological evidence of vertical transmission of JC and BK polyomaviruses in
242 humans. *J Gen Virol* 2011; **92**(Pt 5):1044-50.
- 243 10. Kalvatchev Z, Slavov S, Shtereva M, Savova S. Reactivation of Polyomavirus hominis
244 1 (BKV) during pregnancy and the risk of mother-to-child transmission. *J Clin Virol*
245 2008; **43**(3):328-9.
- 246 11. Boldorini R, Veggiani C, Amoruso E, Allegrini S, Miglio U, Paganotti A, et al. Latent
247 human polyomavirus infection in pregnancy: investigation of possible transplacental
248 transmission. *Pathology* 2008; **40**(1):72-7.

- 249 12. Dalianis T, Ramqvist T, Andreasson K, Kean JM, Garcea RL. KI, WU and Merkel cell
250 polyomaviruses: a new era for human polyomavirus research. *Semin Cancer Biol*
251 2009; **19**(4):270-5.
- 252 13. Bialasiewicz S, Whiley DM, Lambert SB, Nissen MD, Sloots TP. Detection of BK, JC,
253 WU, or KI polyomaviruses in faecal, urine, blood, cerebrospinal fluid and respiratory
254 samples. *J Clin Virol* 2009; **45**(3):249-54.
- 255 14. Mourez T, Bergeron A, Ribaud P, Scieux C, de Latour RP, Tazi A, et al.
256 Polyomaviruses KI and WU in immunocompromised patients with respiratory disease.
257 *Emerg Infect Dis* 2009; **15**(1):107-9.
- 258 15. Babakir-Mina M, Ciccozzi M, Perno CF, Ciotti M. The novel KI, WU, MC
259 polyomaviruses: possible human pathogens? *New Microbiol* 2011; **34**(1):1-8.
- 260 16. Csoma E, Meszaros B, Asztalos L, Konya J, Gergely L. Prevalence of WU and KI
261 polyomaviruses in plasma, urine, and respiratory samples from renal transplant
262 patients. *J Med Virol* 2011; **83**(7):1275-8.
- 263 17. Sauvage V, Foulongne V, Cheval J, Ar Gouilh M, Pariente K, Dereure O, et al.
264 Human polyomavirus related to African green monkey lymphotropic polyomavirus.
265 *Emerg Infect Dis* 2011; **17**(8):1364-70.
- 266 18. Bofill-Mas S, Rodriguez-Manzano J, Calgua B, Carratala A, Girones R. Newly
267 described human polyomaviruses Merkel cell, KI and WU are present in urban sewage
268 and may represent potential environmental contaminants. *Virol J* 2010; **7**(141).
- 269 19. Heritage J, Chesters PM, McCance DJ. The persistence of papovavirus BK DNA
270 sequences in normal human renal tissue. *J Med Virol* 1981; **8**(2):143-50.
- 271 20. Chesters PM, Heritage J, McCance DJ. Persistence of DNA sequences of BK virus and
272 JC virus in normal human tissues and in diseased tissues. *J Infect Dis* 1983;
273 **147**(4):676-84.

- 274 21. Egli A, Infanti L, Dumoulin A, Buser A, Samaridis J, Stebler C, et al. Prevalence of
275 polyomavirus BK and JC infection and replication in 400 healthy blood donors. *J*
276 *Infect Dis* 2009; **199**(6):837-46.
- 277 22. Dharnidharka VR, Abdalnour HA, Araya CE. The BK virus in renal transplant
278 recipients-review of pathogenesis, diagnosis, and treatment. *Pediatr Nephrol*
279 **26**(10):1763-74.
- 280 23. Boothpur R, Brennan DC. Human polyoma viruses and disease with emphasis on
281 clinical BK and JC. *J Clin Virol* 2010; **47**(4):306-12.
- 282 24. Pietropaolo V, Di Taranto C, Degener AM, Jin L, Sinibaldi L, Baiocchi A, et al.
283 Transplacental transmission of human polyomavirus BK. *J Med Virol* 1998;
284 **56**(4):372-6.
- 285 25. Sadeghi M, Riipinen A, Vaisanen E, Chen T, Kantola K, Surcel HM, et al. Newly
286 discovered KI, WU, and Merkel cell polyomaviruses: no evidence of mother-to-fetus
287 transmission. *Virol J* 2010; **7**(251).
- 288
- 289