Patient-individual cancer cell lines and tissue analysis delivers no evidence of sequences from DNA viruses in colorectal cancer cells

CURRENT STATUS: UNDER REVIEW

BMC Gastroenterology  BMC Series

Michael Gock (Former Corresponding Author)
Universitätsmedizin Rostock, Abteilung für Allgemein-, Viszeral-, Gefäß- und Transplantationschirurgie
ORCiD: https://orcid.org/0000-0001-6393-1097

Marcel Kordt
Universitätsmedizin Rostock

Stephanie Matschos
Universitätsmedizin Rostock

Christina S. Mullins
Universitätsmedizin Rostock

Michael Linnebacher (New Corresponding Author)

michael.linnebacher@med.uni-rostock.de

ORCiD: https://orcid.org/0000-0001-8054-1402

DOI:
10.21203/rs.2.13448/v2

SUBJECT AREAS
Gastroenterology & Hepatology

KEYWORDS
colorectal carcinoma, oncogenic virus, JC virus, BK virus, Epstein-Barr-Virus, patient-derived carcinoma cell lines
Abstract

Background Several DNA viruses are highly suspicious to have oncogenic effects in humans. This study investigates the presence of potentially oncogenic viruses such as SV40, JCV, BKV and EBV in patient-derived colorectal carcinoma (CRC) cells typifying all molecular subtypes of CRC. Methods Sample material (gDNA and cDNA) of a total of 49 patient-individual CRC cell lines and corresponding primary material from 11 patients, including normal, tumor-derived and metastasis-derived tissue were analyzed for sequences of SV40, JVC, BKV and EBV using endpoint PCR. In addition, the susceptibility of CRC cells to JCV and BKV was examined using a long-term cultivation approach of patient-individual cells in the presence of viruses. Results No virus-specific sequences could be detected in all specimens. Likewise, no morphological changes were observed and no evidence for viral infection or integration could be provided after long term CRC cell cultivation in presence of viral particles. Conclusions In summary, the presented data suggest that there is no direct correlation between tumorigenesis and viral load and consequently no evidence for a functional role of the DNA viruses included into this analysis in CRC development.

Background

The findings gathered in recent years on the relationship between tumorigenesis and several microorganisms, especially viruses, allow a new approach in the prevention and therapy of tumors [1]. For cervical carcinoma associated with human papilloma viruses (especially types 16 and 18, approximately 70% of all cases), there are already worldwide preventive measures and vaccines against the HPV high-risk types in place [2]. If there was a causal relationship between the development of colorectal cancer (CRC) and a viral infection, it would be possible to create new therapeutic and preventive approaches. In 2012 alone, nearly 1.4 million people worldwide developed CRC and 694,000 died as a result. This makes CRC the third leading cause of cancer and the fourth most common cause of cancer death worldwide [3]. In Germany, CRC is the third most common form of cancer among men and the second most among women and lifetime risk to develop a CRC is 5 to 7% [4].

The first oncogenic viruses were discovered in chicken in 1911. Peyton Rous of the Rockefeller
Institute was able to isolate a retrovirus (subsequently named after him Rous sarcoma virus) from tumors of diseased chicken and he could demonstrate their malignant potential by causing the same sarcomas after injecting viral particles into healthy chicken. Rous sarcoma virus belongs to the same family of Retroviridae as avian leukosis virus, which is able to cause leukemia in chicken. Later, it could be shown that murine polyomavirus and simian virus 40 (SV40) could cause tumors in experimental animals. SV40 also has a carcinogenic effect on human cells [1]. It is well known that at least 6 different viruses could cause cancer in humans [1]. For the year 2002 it was estimated, that approximately 10-15% of all cancers worldwide were caused by viral infections [5]. Polyomaviruses are icosahedral, nonenveloped viruses with a size of 45 nm. The capsid is composed of 72 pentameric capsomers and surrounds a circular, double-stranded viral DNA genome of approximately 5.5kbp. The virus particles are taken up by endocytosis into the target cell and transported there to the nucleus [6]. It has been repeatedly confirmed that SV40 is oncogenic both in vitro and in vivo [7]. Also for the human polyomaviruses JC virus (JCV) and BK virus (BKV), direct carcinogenic effects are still under debate [8]. Regarding JCV, several case control studies have been conducted to establish the association between JCV seropositivity and specific human tumor types, including CRC [9, 10, 11, 12, 13, 14, 15]. BK virus DNA has been detected in several human cancers, including cancer of the head and neck, lung, urogenital tract, liver and pancreas [16, 17]. Epstein-Barr-Virus (EBV) is one of the human pathogenic viruses of the family Herpesviridae. The virus particles are enveloped and have a genome of double-stranded DNA. Seroprevalence in adulthood is over 98%. Though, a direct connection between EBV infection and CRC has not yet been established [18].

The aim of our study was to investigate the presence of potentially oncogenic viruses such as SV40, JCV, BKV and EBV in patient-specific CRC cells, selected to represent all molecular subtypes of CRC. Furthermore, the susceptibility of low-passage, patient-derived CRC cells to the human polyomaviruses JCV and BKV was investigated.

Methods

Tissue sampling, cell line establishment and cell culture
Primary CRC resection specimens, specimens of metastases and normal tissue were received fresh from surgery, with informed written patient consent. Tissues were immediately vitally frozen in freezing medium (fetal calf serum (FCS) containing 10% DMSO) at -80°C for subsequent examinations. All cell lines were established from patient derived tumor material as previously described by [19] (used cell lines are listed in Table 1). Cell lines were cultured in T75 culture flasks using Dulbecco’s Modified Eagle Medium (DMEM)/Ham’s F12 medium supplement with 10% FCS and 2mM L-glutamine as described before [20]. All procedures involving patient material were approved by the Ethics Committee of the Medical faculty, University of Rostock (reference number II HV 43/2004) in accordance with generally accepted guidelines for the use of human material.

Altogether 49 cell lines (Table 1) were tested on Polyomaviridae (SV40, JCV and BKV) and Herpesviridae (EBV). Genomic DNA (gDNA) and complementary DNA (cDNA) were used for the study (all virus preparations were a kind gift of the Institute for Medical Microbiology, Virology and Hygiene, Rostock University Medical Center and serve normally as positive controls in diagnostic virus testing). Seven of the investigated established cell lines were present in different passages (Table 1). Of six cell lines only cDNA was tested (Table 1; highlighted in grey). Primary tissue material, including normal, tumor tissue and metastases of 11 CRC patients was analyzed. Altogether 11 tissue samples of tumor material, 10 tissue samples of normal tissue and 3 tissue samples of metastases were included (Table 2).

**gDNA and RNA isolation**

gDNA was isolated from cell pellets using the “Wizard Genomic DNA Purification Kit” from Promega according to the manufacturer’s instructions. The RNA was isolated from cell pellets using the “GeneMATRIX Universal RNA Purification Kit” from EURx according to the manufacturer’s instructions. The measurement of the isolated nucleic acids was then carried out on NanoDrop 1000 from Thermo Scientific.

**cDNA Synthesis**

cDNA was synthesized from RNA using the Reverase Kit (M-MuLV, RNase H minus) from Bioron (Römerberg, Germany) according to the manufacturer’s instructions. The final cDNA concentration
was 12.5 ng/μl after synthesis.

RT-PCR for SV40 and JC/BK

Primers used for SV40 bind to the specific SV40 nucleotide sequence for the large T antigen (PCR product size: 172bp). Used primers for JC/BK virus (Table 3) bind to the common specific JC/BK nucleotide sequence for the large T antigen (PCR product size JC/BK: 178bp and 181bp). As a positive control for SV40 gDNA, served the SV40 immortalized fibroblast cell line LCT 7. For JC/BK a positive control in form of material from an interlaboratory comparison test with proven high viral load and provided by the Institute of Medical Microbiology, Virology and Hygiene from the Rostock University Medical Center was included. 50ng of the sample were used as start material for the PCR. Cycling conditions were as follows: 94°C – 1min followed by 40 cycles of 94°C – 30 sec, 56°C – 30 sec and 72°C – 1 min with a final extension at 72°C for 5 min.

RT-PCR for EBV

Primers used for EBV (Table 4) bind to specific EBV nucleotide sequence (PCR product size: 153bp). As a positive control for EBV gDNA served the EBV immortalized B cell line HROC212. PCR conditions were as described before.

RT-PCR for β-actin

β-actin was used as an unregulated household gene to verify that samples are gDNA free after RNA-isolation (PCR product size: cDNA: 210bp, gDNA: 700bp). gDNA of cell line HROC111Met1 T0 M2 served as positive control for β-actin. The annealing temperature was 58°C. PCR conditions were as described before.

Sanger Sequencing

The initial PCR product was isolated using the „High Pure PCR Product Purification Kit“ from Merck (Darmstadt, Germany) according to the manufacturer’s protocol. Next, 50ng of the isolated product were used as start material for the sequencing PCR. Cycling conditions were as follows: 94°C – 2min followed by 35 cycles of 94°C – 10 sec, 62°C – 20 sec and 72°C – 1 min with a final extension at 72°C for 5min. The sequencing reaction contained 5μl of this PCR product, 2.5μl nuclease-free water and 2.5μl primer. They were sent to a commercial sequence service provider (Eurofins, Ebersberg,
Germany) which delivered sequence data back.

Agarose gel electrophoresis

To illustrate fragment sizes, a 100 bp DNA ladder (Carl Roth) was used. For electrophoresis, a voltage of 80V (15min) and 100V (45min, Elektrophoresis Power Supply, Biometra) was set. Detection took place on the BIO-Print ST4 from VILBER. Separation was performed with 2% Agarose gels in TE-buffer (reagents from Merck).

Viral infection of cell lines

To test the susceptibility of CRC cells to the JC and BK virus, HROC257 cells were infected with these viruses. The ratio of virus particles and target cells was calculated on the basis of the MOI (= multiplicity of infection) according to the following formula:

\[
\text{MOI} = \frac{\text{Volume (Virus) x concentration (Virus)}}{\text{Volume (cell culture) x concentration (cell culture)}} = \frac{V_v \times C_v}{V_c \times C_c}
\]

MOI calculation for JC and BK virus:

\[
\begin{align*}
\text{MOI} &= \frac{40\text{nm}^3 \times 2,45 \times 10^4 \text{ copies/ml}}{10\mu\text{m}^3 \times 2,50 \times 10^2 \text{ cells/ml}} \\
&= 0.39
\end{align*}
\]

500 HROC257 cells in 2 ml of DMEM were seeded on a 6-well plate and infected with virus at an MOI of 0.39. One well served as negative control. Cells were subsequently incubated at 37° C and 5% CO₂.

Results

Primary aim of this study was the detection of the potentially oncogenic viruses SV40, JCV, BKV and EBV in patient-individual low passage (<P50) CRC cell lines [21]. Therefore, gDNA and cDNA from established and characterized cell lines and corresponding patient tumor (tu) and normal (n) tissues,
as well as metastases (met) were examined by RT-PCR.

Analysis of SV40 in cell lines and primary tissue samples

For the detection of SV40, altogether 49 individual patient-derived colorectal tumor cell lines were examined. No specific band pattern could be shown for the SV40 virus in all cell lines (Figure 1). The positive control shows a clear specific band at 172bp. In order to clarify whether SV40 sequences are detectable at RNA level, additionally cDNA from the same cell lines was examined. Again, no viral sequences could be amplified with the positive control showing a band at 172bp (not shown).

However, examination of primary tumor material revealed signals in HROC277tu and HROC296tu (Figure 2).

PCR product sizes were about 170bp and 300bp, respectively. To clarify the specificity of the detected bands, a reamplification of the PCR products was carried out with the same primers. However, the initially detected band of HROC296 (300bp) could not be reamplified, while the reamplification product of HROC277 exhibited an unspecific smear and must therefore also be regarded as negative (Figure 3).

Analysis of JC and BK virus in cell lines and primary tissue samples

In 2 of 49 examined cell lines, the PCR with primers designed for the amplification of JCV/BKV sequences delivered a signal at approximately 180bp (HROC348Met P9 and HROC212 P17) (Figure 4) suggesting a viral infection. Again, the positive controls delivered the expected products of 178bp (JCV) and 181bp (BKV), respectively (Figures 4 and 5).

To verify these findings, a highly sensitive diagnostic qPCR (TaqMan®) was performed to detect JCV and BKV at the Institute of Microbiology of the University Medical Center, Rostock as described originally by Watzinger et al., in 2004 with minor modifications [22]. However, no JC or BK viral load could be detected in any of the samples and the initial results could thus not be confirmed. In analogy to the SV40 analysis, cDNA was also assayed for JCV and BKV. All cDNA samples were negative (data not shown).

The examination of primary tumor material revealed a signal at approximately 380bp for the samples HROC107 n/tu, HROC126 n, HROC183 n/tu and HROC212 n (Figure 5). As product size was greater
than the expected 178bp or 181bp for JC/BK virus, a reamplification PCR was performed with the PCR product of the primary materials. Here, the result of the first RT-PCR could be confirmed. All samples of the reamplification showed a product size of approximately 380bp. To check the specificity of this amplified nucleotide sequence of the primary patient materials, the reamplified amplicons were sequenced by Sanger sequencing. This analysis identified a human DNA sequence from clone RP11-203L2 (chromosomes 9q21.11-21.2) with a 99% identification probability. Thus, in none of the investigated primary materials, JCV or BKV were detectable.

**Analysis of EBV in cell lines and primary tissue samples**

RT-PCR revealed several amplification products at about 700bp in size in the examined cell lines and primary tissue samples with no difference between tumor and normal tissue (data not shown). The product size is about 700bp and is thus by far larger than the expected product size of 150bp. A reamplification failed to reproduce the initial 700bp bands in all cases. Similar to the observation for SV40, either an unspecific smear or no signal at all was obtained. In conclusion, it can be assumed that the 700bp amplicons are non-specific.

**Susceptibility of HROC257 to JC and BK virus**

To test the susceptibility of CRC cells to infection with JC and BK virus, the cell line HROC257 was infected with JC and BK viruses with an MOI of 0.39 as explained before. Growth and morphology of the cells were documented photographically (Figure 6). Particular attention was paid to the following parameters: changes in growth behavior / inhibition of proliferation, cell lysis or potential transformatory effects, reflected in the morphology of the cells. During cultivation of the HROC257 cells in the presence of the JCV and BKV, no morphological changes could be observed in comparison to untreated control cells. As shown in Figure 6, the growth behavior of infected and uninfected cells is similar. In order to further analyze susceptibility to JCV and BKV, these HROC257 cells were subjected to the above described RT-PCR, 40 days after viral infection. Both, cell culture supernatant and isolated cellular gDNA were used as templates. In positive controls, the analysis of cell culture supernatant showed bands at 178bp (JCV) and 181bp (BKV). But no products were detectable in supernatant of neither JCV nor BKV treated cells (data not shown). This result suggests that either no
or very few viral particles were present in the supernatant. In addition, isolated cellular gDNA of virus-treated HROC257 cells were tested for JCV and BKV after 40 days of incubation. But again no viral amplicon could be detected (data not shown). These results indicate that HROC257 were not susceptible to JC or BK virus infection.

Discussion

Recent findings shed light into the relationship between tumor development and infections by several viruses. This culminates in the development of preventive vaccinations against cervical carcinoma associated with HPV high-risk types [2]. A causal relationship between the development of CRC and viral infections has repeatedly been suggested and this could, if proved, open new avenues for similar therapeutic and preventive approaches.

Our study aimed at deciphering a potential involvement of several oncogenic DNA viruses in CRC development. The starting hypothesis was that if viruses would have an active role in the development of CRC, their footsteps in form of virus-encoded nucleotide sequences should be present in patients’ CRC cells. In order to avoid any bias by non-tumor cells present in the tumor microenvironment, we focused on patient-individual, low passage CRC cell lines; but corresponding patient tissue samples were analyzed in addition. Using highly sensitive PCR approaches, we investigated the natural occurrence of the polyomaviruses SV40, JCV and BKV as well as the herpesvirus EBV – the most frequently discussed viruses with a potential role in CRC development [9, 15, 21, 13, 10, 23, 24, 25, 26]. First, both patient-individual CRC cell lines (n = 49) and corresponding tissue samples from patients (normal and tumor tissue, as well as liver, lung and peritoneal metastases) were analyzed for the presence of viral sequences by end-point PCR. But in none of the tested samples, viral sequence amplification was successful.

Regarding a potential SV40 infection, none of the 49 low-passage patient-derived cell lines was positively tested for SV40. This finding excludes SV40 infection at least as a frequent event in CRC cells. It is well known that SV40 might act as a cofactor in the carcinogenesis of several carcinomas; but its involvement in CRC oncogenesis has been examined poorly up to now [27]. Only one case control study by Campello et al. analyzed the involvement of SV40 in CRC pathogenesis [9]. However,
only six out of 94 analyzed patients’ tumors were tested positive for SV40 and the authors could not
delineate a relationship between viral infection and CRC. An explanation for the differences to the
results of the present study could be the extracellular presence of viral particles – either between the
cells, more likely in infiltrating lymphocytes or macrophages, and possibly even in the lumen of the
gut.

An infection of the analyzed cell lines with the polyomaviruses JCV and BKV could be ruled out
likewise. Hitherto existing data on prevalence of JCV infections in CRC are very heterogeneous.
Several small case control studies were reported. A study from Portugal could show a prevalence of
90% JCV infections in samples of CRC lesions [15] and in another study from Greek about 60% of the
samples tested positive for JCV [28]. A Chinese study of 137 samples described a rate of infection in
CRC of 40.9% [13]. But in contrast, a large American study failed to prove JCV infections in 233
patient samples [29] and a study from Iran described a positive detection rate of only 1.46% (2 of 70
patients) [30]. Our own results are pretty much in line with the latter findings and also indicate that
JCV is unlikely involved in CRC development.

Worldwide ninety percent of adults have been exposed to BKV. BKV DNA has been detected in several
human cancers, including carcinoma of the lung, urogenital tract, liver, head and neck, in
rhabdomyosarcoma and in brain tumors [17, 31]. However, reports on detection of BKV DNA in CRC
tissues are very scarce. A study by Giuliani and colleagues could detect BKV DNA in 9% out of 66
samples [23] and Casini et al. found BKV DNA in several CRC tissues [10]. Again, in our study
population in none of the examined cell lines or tissue samples BKV DNA could be detected.
Finally, all cell lines and primary materials were examined for the presence of EBV. Besides non-
specific PCR products, presence of EBV DNA could not be verified. All analyzed samples must
therefore be considered as EBV negative. Militello et al. described a very low positive detection
frequency for EBV DNA in cancerous and cancer-adjacent mucous samples [32] and a study by Cho et
al. could not detect EBV in 274 CRC specimen [33]. In contrast, several studies reported a high EBV
DNA positivity of CRC tissues [24, 25, 26]. Interestingly, a study by Fiorina et al. demonstrated EBV in
23 of 44 CRC samples but they could prove that this was restricted to latency in the lymphoid
infiltrate of the tumors [34]. This latter observation delivers the best explanation for the inconsistencies between the before mentioned results. EBV most effectively infects B cells and rarely other lymphoid cell populations. In cultured epithelial-derived cells with no contaminating lymphocytes left over from the initial culturing process, a cross-contamination of the PCR results can safely be excluded. Of note, such a large panel of patient-individual cell lines with corresponding tissue samples has never been tested before. The primary CRC tissues included into our analysis were obtained from the invasive margins which are especially rich in (vital) tumor cells – but infiltrating T cells and also B cells are also readily detectable [35]. Thus, we cannot exclude sporadic positive results in EBV testing due to EBV-positive lymphocytes present in the tumor microenvironment, if a larger sample set of our primary CRC tissue collection would be analyzed.

Based on the data obtained in this work for SV40, JCV, BKV and EBV, it can be concluded that none of the tested viruses are likely to have an obvious general role in CRC development. Possible reasons for the variation in the detection rate of these viruses in our and several of the aforementioned studies, are beside the lymphocytic contamination of the specimens [36], differences in the detection limit of the applied detection systems but possibly also the genetic background of the different patient populations.

In order to verify the general negative findings on a more functional level, we tested exemplary the susceptibility of the CRC cell line HROC257 T0 M1 to JCV and BKV infection. However, even after an extended cultivation period subsequent to viral inoculation, no evidence for a successful viral infection could be provided. This led us to conclude that CRC cells are not susceptible to JC or BK virus infection and thus further supports the negative screening results.

Conclusions
In conclusion, the presented data suggest that there is no direct correlation between tumorigenesis and viral load of SV40, JVC, BKV and EBV and consequently no evidence for a functional role of the DNA viruses included into this analysis in CRC development.

List Of Abbreviations
colorectal cancer (CRC), human papilloma viruses (HPV), simian virus 40 (SV40), Epstein-Barr-Virus


(EBV), JC virus (JCV), BK virus (BKV), fetal calf serum (FCS), multiplicity of infection (MOI), Dulbecco's Modified Eagle Medium (DMEM)

Declarations

Ethics approval and consent to participate

All procedures involving patient material were approved by the Ethics Committee of the Medical faculty, University of Rostock (reference number II HV 43/2004) in accordance with generally accepted guidelines for the use of human material.

Consent for publication

Not applicable

Availability of data and material

The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests

Funding

This research received no external funding.

Authors’ Contributions: Conceptualization of research: ML and MG; MK, CSM and SM. performed research; MG and ML analyzed and reviewed data; MG and ML wrote the manuscript; CSM and SM contributed parts of the manuscript and critically reviewed the final version of the manuscript.

Acknowledgments

The authors thank Dr. T. Köller from the Institute for Medical Microbiology, Virology and Hygiene for delivery of positive viral controls and for performing the qPCR (TaqMan®) analyses of JCV and BKV.

The authors thank Prof. Dr. R. Jaster from the Division of Gastroenterology, Department of Medicine II, Rostock University Medical Center for providing the LCT 7 cells.

References

1. Martin, D.; Gutkind, J.S. Human tumor-associated viruses and new insights into the molecular mechanisms of cancer. Oncogene. 2008;27:S31-42. doi: 10.1038/onc.2009
2. Lee, S.J.; Yang, A.; Wu, T.C.; Hung, C.F. Immunotherapy for human papillomavirus associated disease and cervical cancer: review of clinical and translational research. J Gynecol Oncol. 2016;31:e51. doi: 10.3802/jgo.2016.27.e51

3. Ferlay, J.; Soerjomataram, I.; Dikshit, R.; Eser, S.; Mathers, C.; Rebelo, M.; Parkin, D.M.; Forman, D.; Bray, F. Cancer incidence and mortality worldwide: sources, methods and major patterns in GLOBOCAN 2012. Int J Cancer. 2015;136(5),E359-86. doi: 10.1002/ijc.29210

4. Kaatsch, P.; Spix, C.; Katalinic, A.; Hentschel, S.; Luttmann, S.; Stegmaier, C.; Caspritz, S.; Christ, M.; Ernst, A.; Folkertsor, J.; Hansmann, J.; Klein, S.; Kranzhöfer, K.; Kunz, B.; Manegold, K.; Penzkofer, A.; Treml, K.; Weg-Remers, S.; Wittenberg, K.; Baras, N.; Barnes, B.; Bertz, J. Krebs in Deutschland 2011/2012, In: Gesellschaft der epidemiologischen Krebsregister in Deutschland e.V.. Berlin: Robert Koch Institut, Germany 2015; pp. 38-41.

5. Parkin, D.M. The global health burden of infection-associated cancers in the year 2002. Int J Cancer. 2005;118:3030-44. doi:10.1002/ijc.21731

6. Ferenczy, M.W.; Marshall, L.J.; Nelson, C.D.; Atwood, W.J.; Nath, A.; Khalili, K. Molecular biology, epidemiology, and pathogenesis of progressive multifocal leukoencephalopathy, the JC virus-induced demyelinating disease of the human brain. Clin Microbiol Rev. 2012;25(3):471-506. http://dx.doi.org/10.1128/CMR.05031-11.

7. Ahuja, D.; Sáenz-Robles, M.T.; Pipas, J.M. SV40 large T-Antigen targets multiple cellular pathways to elicit cellular transformation. Oncogene. 2005;24(52):7729-45. doi: 10.1038/sj.onc.1209046

8. Maginnis, M.S.; Atwood, W.J. JC virus: an oncogenic virus in animals and humans? Semin Cancer Biol. 2009;19(4):261-9. http://dx.doi.org/
9. Campello, C.; Comar, M.; Zanotta, N.; Minicozzi, A.; Rodella, L.; Poli, A. Detection of SV40 in colon cancer: A molecular case-control study from Northeast Italy. J Med Virol. 2010;828(7):1197–200. doi: 10.1002/jmv.21798

10. Casini, B.; Borgese, L.; Del Nonno, F.; Galati, G.; Izzo, L.; Caputo, M.; Perrone Donnorso, R.; Castelli, M.; Risuleo, G.; Visca, P. Presence and incidence of DNA sequences of human polyomaviruses BKV and JCV in colorectal tumor tissues. Anticancer Res. 2005;25(2A):1079-85.

11. Vilkin, A.; Ronen, Z.; Levi, Z.; Morgenstern, S.; Halpern, M.; Niv, Y. Presence of JC virus DNA in the tumor tissue and normal mucosa of patients with sporadic colorectal cancer (CRC) or with positive family history and Bethesda criteria. Dig Dis Sci. 2012;57(1):79-84. http://dx.doi.org/10.1007/s10620-011-1855-z

12. Butcher, L.D.; Garcia, M.; Arnold, M.; Ueno, H.; Goel, A.; Boland, C.R. Immune response to JC virus T antigen in patients with and without colorectal neoplasia. Gut Microbes. 2014;5(4):468-75. http://dx.doi.org/10.4161/gmic.29573

13. Mou, X.; Chen, L.; Liu, F.; Lin, J.; Diao, P.; Wang, H. Prevalence of JC virus in Chinese patients with colorectal cancer. PLoS One. 2012;7(5):e35900. http://dx.doi.org/10.1371/journal.pone.0035900.

14. Samaka, R.M.; Abd El-Wahed, M.M.; Aiad, H.A.; Kandil, M.A.; Al-Sharaky, D.R. Does JC virus have a role in the etiology and prognosis of Egyptian colorectal carcinoma? APMIS. 2013;121(4):316-28. http://dx.doi.org/10.1111/apm.12001.

15. Coelho, T.R.; Gaspar, R.; Figueiredo, P.; Mendonca, C.; Lazo, P.A.; Almeida, L. Human JC polyomavirus in normal colorectal mucosa, hyperplastic polyps, sporadic adenomas, and adenocarcinomas in Portugal. J Med Virol. 2013;85(12):2119-27. http://dx.doi.org/10.1002/jmv.23705.
16. Drop, B.; Strycharz-Dudziak, M.; Kliszczewska, E.; Polz-Dacewicz, M. Coinfection with Epstein-Barr Virus (EBV), Human Papilloma Virus (HPV) and Polyoma BK Virus (BKPyV) in Laryngeal, Oropharyngeal and Oral Cavity Cancer. Int J Mol Sci. 2017;18(12):E2752. http://dx.doi.org/10.3390/ijms18122752.

17. Fioriti, D.; Videtta, M.; Mischitelli, M.; Degener, A.M.; Russo, G.; Giordano, A. The human polyomavirus BK: Potential role in cancer. J Cell Physio. 2005;204(2):402-6. http://dx.doi.org/10.1002/jcp.20300.

18. Bedri, S.; Sultan, A.A.; Alkhalaf, M.; Al Moustafa, A.E.; Vranic, S. Epstein-Barr virus (EBV) status in colorectal cancer: a mini review. Hum Vaccin Immunother. 2018;31. doi: 10.1080/21645515.2018.1543525.

19. Maletzki, C.; Stier, S.; Gruenert, U.; Gock, M.; Ostwald, C.; Prall, F.; Linnebacher, M. Establishment, characterization and chemosensitivity of three mismatch repair deficient cell lines from sporadic and inherited colorectal carcinomas. PLoS One. 2012;7(12):e52485. doi: 10.1371/journal.pone.0052485. Epub 2012 Dec 31

20. Maletzki, C.; Gock, M.; Randow, M.; Klar, E.; Huehns, M.; Prall, F.; Linnebacher, M. Establishment and characterization of cell lines from chromosomal instable colorectal cancer. World J Gastroenterol. 2015;7;21(1):164-76. doi: 10.3748/wjg.v21.i1.164.

21. Mullins CS, Micheel B, Matschos S, Leuchter M, Bürtin F, Krohn M, Hühns M, Klar E, Prall F, Linnebacher M. Integrated Biobanking and Tumor Model Establishment of Human Colorectal Carcinoma Provides Excellent Tools for Preclinical Research. Cancers (Basel). 2019;9;11(10). doi: 10.3390/cancers1101520.

22. Watzinger F, Suda M, Preuner S, Baumgartinger R, Ebner K, Baskova L, Niesters HG, Lawitschka A, Lion T. Real time quantitative PCR assays for detection and monitoring of pathogenic human viruses in immunosuppressed pediatric patients. J Clin Microbiol. 2004 42(11):5189-98. doi: 1128/JCM.42.11.5189-5198.2004
23. Giuliani, L.; Ronci, C.; Bonifacio, D.; Di Bonito, L.; Favalli, C.; Perno, C.F.; Syrjänen, K.; Ciotti, M. Detection of oncogenic DNA viruses in colorectal cancer. Anticancer Res. 2008;28(2B):1405-10.

24. Liu, H.X.; Ding, Y.Q.; Li, X.; Yao, K.T. Investigation of Epstein-Barr virus in Chinese colorectal tumors. World J Gastroenterol. 2003;9(11):2464-8.

25. Karpinski, P.; Myszka, A.; Ramsey, D.; Kielan, W.; Sasiadek, M.M. Detection of viral DNA sequences in sporadic colorectal cancers in relation to CpG island methylation and methylator phenotype. Tumour Biol. 2011;32(4):653-9. doi: 10.1007/s13277-011-0165-6. Epub 2011 Apr 6

26. Rüschoff, J.; Dietmaier, W.; Lüttges, J.; Seitz, G.; Bocker, T.; Zirngibl, H.; Schlegel, J.; Schackert, H.K.; Jauch, K.W.; Hofstaedter, F. Poorly differentiated colonic adenocarcinoma, medullary type: clinical, phenotypic, and molecular characteristics. Am J Pathol. 1997;150(5):K1815-25.

27. Vilchez, R.A.; Butel, J.S. Emergent human pathogen simian virus 40 and its role in cancer. Clin Microbiol Rev. 2004;17(3):495-508. doi: 10.1128/CMR.17.3.495-508.2004

28. Theodoropoulos, G.; Panoussopoulos, D.; Papaconstantinou, I.; Gazouli, M.; Perdiki, M.; Bramis, J.; Lazaris, AC. Assessment of JC polyoma virus in colon neoplasms. Dis Colon Rectum. 2005;48(1):86-91.

29. Newcomb, P.A.; Bush, A.C.; Stoner, G.L.; Lampe, J.W.; Potter, J.D.; Bigler, J. No evidence of an association of JC virus and colon neoplasia. Cancer Epidemiol Biomarkers Prev. 2004;13(4):662-6.

30. Sarvari, J.; Mahmoudvand, S; Pirbonyeh, N.; Safaei, A.; Hosseini, S.Y. The Very Low Frequency of Epstein-Barr JC and BK Viruses DNA in Colorectal Cancer Tissues in Shiraz, Southwest Iran. Pol J Microbiol. 2018;67(1):73-79. doi: 10.5604/01.3001.0011.6146.
31. Drop, B.; Strycharz-Dudziak, M.; Kliszczewska, E.; Polz-Dacewicz, M. Coinfection with Epstein-Barr Virus (EBV), Human Papilloma Virus (HPV) and Polyoma BK Virus (BKPyV) in Laryngeal, Oropharyngeal and Oral Cavity Cancer. Int J Mol Sci. 2017;18(12):E2752. http://dx.doi.org/10.3390/ijms18122752.

32. Militello, V.; Trevisan, M.; Squarzon, L.; Biasolo, M.A.; Rugge, M.; Militello, C.; Palù, G.; Barzon, L. Investigation on the presence of polyomavirus, herpesvirus, and papillomavirus sequences in colorectal neoplasms and their association with cancer. Int J Cancer. 2009;124(10):2501-3. doi: 10.1002/ijc.24224.

33. Cho, Y.J.; Chang, M.S.; Park, S.H.; Kim, H.S.; Kim, W.H. In situ hybridization of Epstein-Barr virus in tumor cells and tumor-infiltrating lymphocytes of the gastrointestinal tract. Hum Pathol. 2001;32(3):297-301. doi: 10.1053/hupa.2001.22766

34. Fiorina, L.; Ricotti, M.; Vanoli, A.; Luinetti, O.; Dallera, E.; Riboni, R.; Paolucci, S.; Brugnatelli, S.; Paulli, M.; Pedrazzoli, P.; Baldanti, F.; Perfetti, V. Systematic analysis of human oncogenic viruses in colon cancer revealed EBV latency in lymphoid infiltrates. Infect Agent Cancer. 2014;9:18. doi: 10.1186/1750-9378-9-18. eCollection 2014.

35. Maletzki, C.; Jahnke, A.; Ostwald, C.; Klar, E.; Prall, F.; Linnebacher, M. Ex-vivo clonally expanded B lymphocytes infiltrating colorectal carcinoma are of mature immunophenotype and produce functional IgG. PLoS One. 2012;7(2):e32639. doi: 10.1371/journal.pone.0032639.

36. Rollison, D.E. JC virus infection: a cause of colorectal cancer? J. Clin Gastroenterol. 2010;44(7):466-468. doi: 10.1097/MCG.0b013e3181e0084b

Tables
| Number | Name               | Material |
|--------|-------------------|----------|
| 1      | HROC18 P27        | gDNA     |
| 2      | HROC24 P31        | gDNA     |
| 3      | HROC24 P42        | cDNA     |
| 4      | HROC24 T1 M2 P53  | cDNA     |
| 5      | HROC24 T3 M1 #2 P9| gDNA     |
| 6      | HROC32 P34        | gDNA     |
| 7      | HROC32 T0 M2      | cDNA     |
| 8      | HROC32 T3 M1 P10  | gDNA     |
| 9      | HROC39 P30        | gDNA     |
| 10     | HROC39 T0 M1      | cDNA     |
| 11     | HROC40 P29        | gDNA     |
| 12     | HROC46 T0 M1 P40  | gDNA     |
| 13     | HROC50 T1 M5 P29  | gDNA     |
| 14     | HROC57 P31        | gDNA     |
| 15     | HROC60 P38        | gDNA     |
| 16     | HROC69 P29        | gDNA     |
| 17     | HROC69 T0 M2 P32  | gDNA     |
| 18     | HROC80 T1 M1 P25  | gDNA     |
| 19     | HROC87 T0 M2 P25  | gDNA     |
| 20     | HROC87 T0 M2 P29  | cDNA     |
| 21     | HROC103 T0 M1 P36 | gDNA     |
| 22     | HROC107 cT0 M2 P38| gDNA     |
| 23     | HROC111 Met1 T0 M2 P7 | gDNA |
| 24     | HROC111 Met1 T0 M2 P8 | gDNA |
| 25     | HROC112 Met P30   | gDNA     |
| 26     | HROC113 cT0 M1 P32| gDNA     |
| 27     | HROC126 P26       | gDNA     |
| 28     | HROC131 T0 M3 P21 | gDNA     |

Table 1: 49 cell lines were analyzed. Seven cell lines were present in different passages. Of six cell lines only cDNA was analyzed (grey highlighted), instead of gDNA and cDNA.
Table 2: Used primary material (n= normal epithelial tissue, tu= tumor tissue, met= metastasis).

| Tissue type | Tissue type |
|-------------|-------------|
| HROC32      | n           |
| HROC103     | n, tu       |
| HROC107     | n, tu       |
| HROC126     | n, tu       |
| HROC147     | n, tu, met  |
| HROC183     | n, tu       |
| HROC212     | n           |
| HROC222     | n, tu       |
| HROC277     | n, tu, met  |
| HROC278     | n, tu, met  |
| HROC296     | n, tu       |

Table 3: Primer sequences for SV40 and JC/ BK virus
| Primer Type | Primer Sequence | Supplier Information |
|-------------|-----------------|----------------------|
| SV40 LargeT Forward | 5´-AGA TTC CAA CCT ATG GAA CTG A-3´ | Eurofins MWG Operon LLC; Kentucky Louisville, USA |
| SV40 LargeT Reverse | 5´-GAA AGT CCT TGG GGT CTT CTA CC-3´ | Eurofins MWG Operon LLC; Kentucky Louisville, USA |
| JC/BK Forward | 5´-AGG TGC CAA CCT ATG GAA CAG A-3´ | Metabion International AG, Planegg, Germany |
| JC/BK Reverse | 5´-GAA AGT CTT TAG GGT CTT CTA CC-3´ | Metabion International AG, Planegg, Germany |
| EBV1 Forward | 5´-CAC TTT AGA GCT CTG GAG GA-3´ | Thermo Scientific; USA, Massachusetts |
| EBV2 Reverse | 5´-TAA AGA TAG CAG CAG CGC AG-3 | Thermo Scientific; USA, Massachusetts |

Table 4: Primer sequences for EBV

Figures
Figure 1
Detection of SV40 by endpoint PCR. Shown are exemplary results of examined cell lines.

Material: gDNA, product size 172bp.

Figure 2
Detection of SV40 by endpoint PCR. Shown are the results of the analysis of primary material. n= normal tissue, tu= tumor tissue, met= metastasis. Material: gDNA, product size 172bp.
Figure 3

Reamplification PCR for detection of SV40. Material: PCR product of the primary material HROC277tu and HROC296tu. Product size of the specific amplicon is 172bp.
Detection of JC and BK virus by endpoint PCR. Shown are exemplary results of investigated cell lines. Material: gDNA, product size is 178bp (JCV) and 181bp (BKV).

Detection of JC and BK by endpoint PCR. Shown are the results of the analysis of primary material. Material: gDNA, product size is 178bp (JCV) and 181bp (BKV).
Figure 6

A: Cultivation of HROC257 cells with BK virus after 10, 25 and 38 days of incubation. B:
Cultivation of HROC257 cells with JC virus after 10, 25 and 38 days of incubation. C:
Negative control (untreated HROC257 cells) after 10, 25 and 38 days. 10x magnification for
all cell culture images.