

Review Article

The dysregulation of Cyclin Dependent Kinase Regulators Role in SV40 Related renal cell carcinoma

Comment [RB1]: Is this a research or a review article?

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Abstract:

Objective: This research used PCR and immunohistochemical techniques to evaluate the assistance of SV40 polyomavirus infection to the progression of renal cell carcinoma in patients from the province of Al Najaf.

Method: This present study was planned as cross-section study to detect SV40 with renal cell carcinoma and includes 75 (45 males and 30 females, whose ages ranged from 22 to 70 years) paraffin impeding block tissues of renal cell carcinoma from archives of AL-Sader Medical City and some archives of private histopathology laboratories in Najaf governorate. The data are from January 2016 to the December of the same year by using Polymers Chain Reaction (P.C.R) for detection of DNA SV40 and immunohistochemistry technique (IHC) for detect expression state of Cyclin Dependent Kinase Regulators (KAP or cyclin-dependent kinase inhibitor 3(CDKN3)& Cyclin E1 markers), using Hematoxylin and Eosin stain for diagnosis of RCC.

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Result: Increased positive percentage for KAP or CDKN3 marker and decreased positive percentage of Cyclin E1 marker were seen in the results of the Immunohistochemistry technique (IHC). As well, found that clear cell type was higher with **42 (56%)**, grade I was higher with 31 (41.3%) and tumor stage type I was higher (25). The positive results by PCR techniques in RCC patient showed that 20 (26.7 % out of 75 cases) of block tissues.

Conclusion: The association RCC with SV40 is mostly caused by dysregulation of Cyclin Dependent Kinase regulators (CDK). It is clear from this study that the Simian Virus 40 (SV40), in particular its Large T Antigen (Tag), affects CDK regulators and upsets the delicate equilibrium of cell cycle regulation systems. There may be a connection between renal cell carcinoma development and the SV40 polyomavirus. Renal cell carcinoma patients are thought to undergo routine testing for detection using PCR and IHC methods.

Key wards: SV40, Renal cell carcinoma, Immunohistochemistry, PCR

Introduction:

Polyomaviruses (PyV) is recognized as a small, non-enveloped, double-stranded deoxyribonucleic acid, icosahedral symmetry with 5 kbp genomes, belonging to *polyomaviridae* family. The term polyomavirus (PyV) comes from Greek origin, where poly- indicatenumeros and -oma which denote tumors, was belong to *Papovaviridae* family, an abbreviation suggested via Melnick, as well as gained via combining the names of the following viruses represented by 'Papilloma', 'Polyoma', and 'Vacuolating' (Dalianis and Hirsch, 2013).

The detection of Simian virus 40 SV40, was reported within 1960 when millions of population in Africa, Europe, Canada, Asia and North and South America were

inoculated from both inactivated and a live polio-vaccines, initiate to be infected by Simian virus 40 (Sweet and Hilleman, 1960).

SV40 genome is circular ds DNA, which encodes for 6 proteins: three structural proteins (including VP-1; VP-2 and VP-3, which are structural proteins allow genetic material to be accumulated in SV40 virion (Kawano *et al.*, 2015), 2 proteins important for the life cycle, that induce replication of SV40, gene-expression, in addition to entry of S phase and DNA synthesis, by this means inducing cycle development (large "T" antigen plus small "t" antigen oncoproteins) (Sullivan and Pipas, 2002; Qi *et al.*, 2011) and 2 small proteins of unidentified function (the agnoprotein, which rule the perinuclear localization of "VP-1" throughout virion construction, after that induce assemblage of virion (Saribas *et al.*, 2018), and 17kT, which participate the majority of amino acid sequence with N terminal domain of T-ag, encourage progression of cell cycle in existence of t-Ag, as well as tumorigenic formation (Comerford *et al.*, 2012).

Simian virus 40 return to Polyomaviridae, genus Betapolyomavirus, which is strongly correlated to other types of polyomaviruses including JCPyV and BKPyV (Calvignac-Spencer *et al.*, 2016). SV40 be capable of transmitted by diverse ways like sexual course and faecal-oral ways that are accountable for horizontal virus infection in peoples (Vanchiere *et al.*, 2005).

The infection of cell beginning by attachment capsid of SV40 to the cell surface by binding among VP-1, cell surface receptor ganglioside GM1 and the major histocompatibility complex class-I (MHC-I), which function as coreceptors (Campanero-Rhodes *et al.*, 2007).

This virus in nature infects specific species of Asian macaques, especially rhesus monkey. Sequences of SV40 were detected in samples of urine and stool as well as in both children and adults, this representing that the sexual and oro-fecal ways of spread that possible to accountable for horizontal SV40 infection in individuals (Academies, 2003; Vanchiere *et al.*, 2005).

On the other hand, the liberation of SV40 with no exhibit a cytopathic effect (CPE) found in particular types of cell, for instance human epithelial, fibroblasts, mesothelial and embryonic renal cell which points that kidney tissue can function as reservoir for SV40 in humans (Cacciotti *et al.*, 2001).

Expression of both T-Ag and t-Ag can cause elevated cell transformation professionally. In reality, T-ag prohibit the actions of numerous diverse cellular factors concerned in differentiation, cell growth and the cell cycle, for instance p130, p300 and p400. As well as, T-Ag and t-Ag was prohibit the activity of pRb and p53. These interconnection are obligatory so as to accomplish complete cell transformation in human (Khalil *et al.*, 2008).

The oncogenic role of polyomavirus was formerly related with a wide array of tumor types for instance malignant pleural mesothelioma (MPM) and bone (Than *et al.*, 2016), brain (Wang *et al.*, 2017), lung (Ramael and Nagels, 1999), thyroid (Vivaldi *et al.*, 2003), pituitary (Woloschak *et al.*, 1995), and urothelial (Loghavi and Bose, 2011) tumors, pleomorphic adenomas of parotid glands (Martinelli *et al.*, 2002), ependymomas choroid and plexus tumors in youth (Bergsage *et al.*, 1992). Additionally, footprints from DNA of SV40 have been reported in breast (Hachana *et al.*, 2009) and colon carcinoma (Campello *et al.*, 2010).

Also, T-ag of SV40 possibly causes transformation by stimulating mutations to the genome of cellular or numerical/structural variation of chromosomes, like gaps, breaks, ring and dicentric chromosomes, chromatid exchanges, translocations, duplications and deletions (Tognon *et al.*, 1996). The major function of t-Ag in transformation is to link both

subunits, catalytic (36 kDa) and regulatory (63 kDa) of protein phosphatase 2A (PP2A), in-activating role (Garcea and Imperiale, 2003).

Grading Renal Cell Carcinoma:

Patients distributing according grading of The World Health Organization(WHO)/Internationa Society of Urological Pathology.

Table 1 :The world heath organization/International society of urological pathology grading system for clear cell and papillary renal carcinoma

Grade 1	Tumour cell nucleoli absent or inconspicuous and basophilic at 400× magnification
Grade 2	Tumour cell nucleoli conspicuous and eosinophilic at 400× magnification and visible but not prominent at 100× magnification
Grade 3	Tumour cell nucleoli conspicuous and eosinophilic at 100× magnification
Grade 4	Tumours showing extreme nuclear pleomorphism, tumour giant cells and/or the presence of any proportion of tumour showing sarcomatoid and/or rhabdoid dedifferentiation

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Result:

-Clinicopathological analysis:

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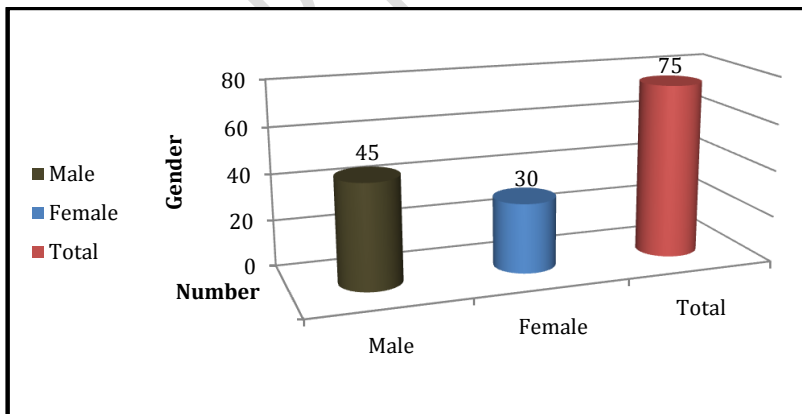


Fig 1:Distribution of RCC Patients according to Gender

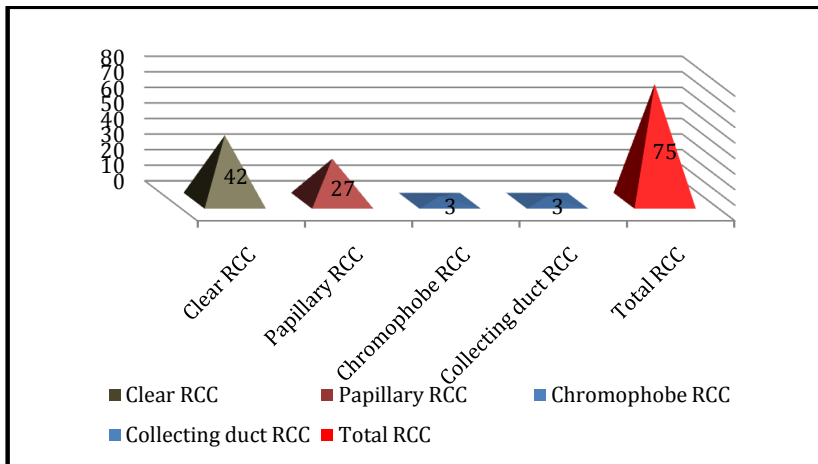


Fig 2: Distribution of RCC Patients according to histological types.

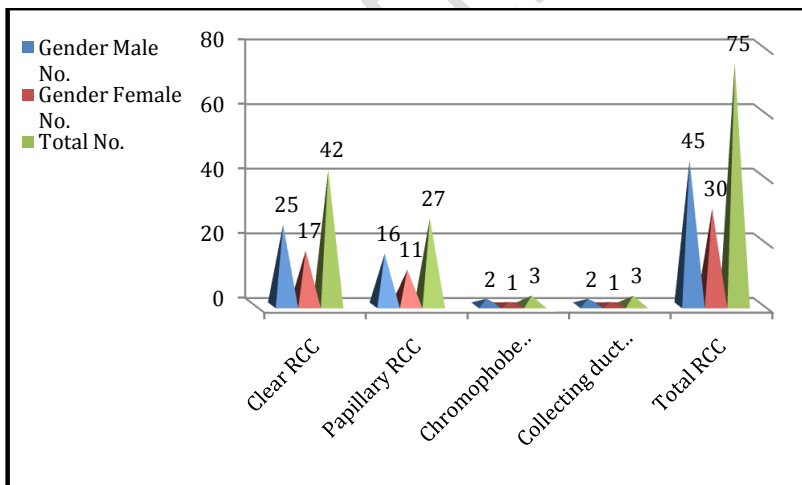


Fig 3: Distribution of RCC patients according to their Histopathological types and Gender

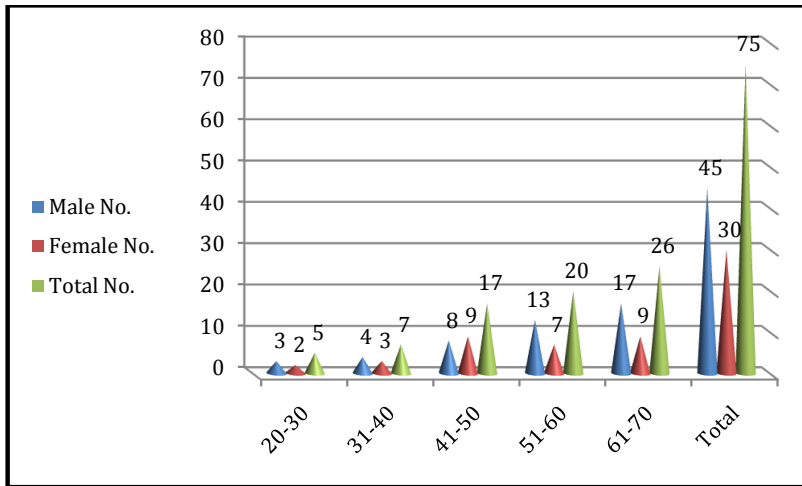


Fig 4: Distribution of RCC patients according to their Gender and Age

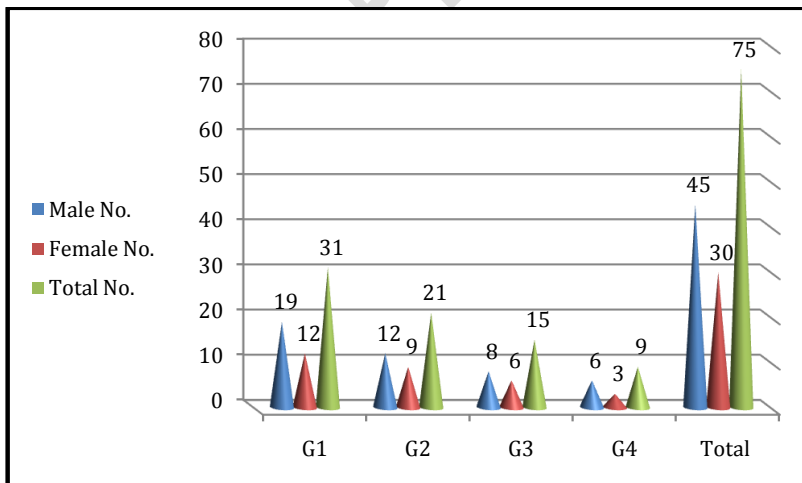


Fig 5: Distribution of RCC patients according to their Gender and Grading Systems

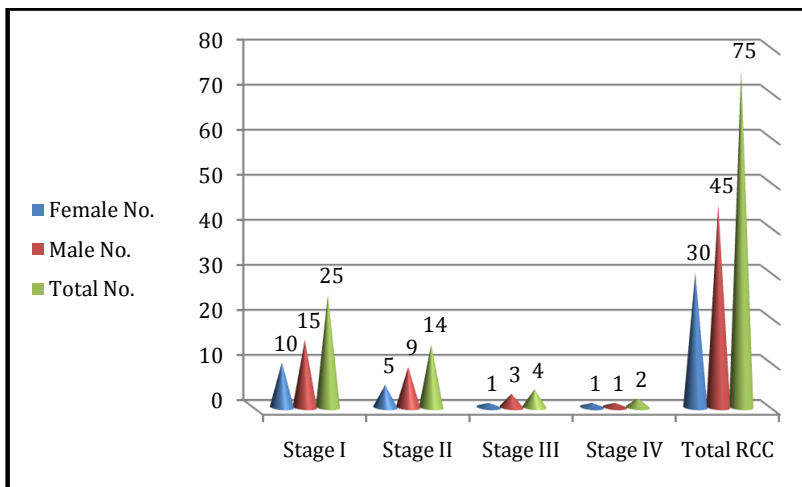


Fig 6: Distribution of RCC patients according to their Gender and Pathological Tumor Stage

- Immunohistochemical Analysis (Cyclin E1 & CDKN3)

In the this study, the results of IHC by utilizing EnVision™ FLEX stain revealed a brownish discolouration of nucleus or nucleoplasm for Cyclin E1 whereas in CDKN3 was staining the cytosol or cytoplasm, as showed in figures (7&8&9).

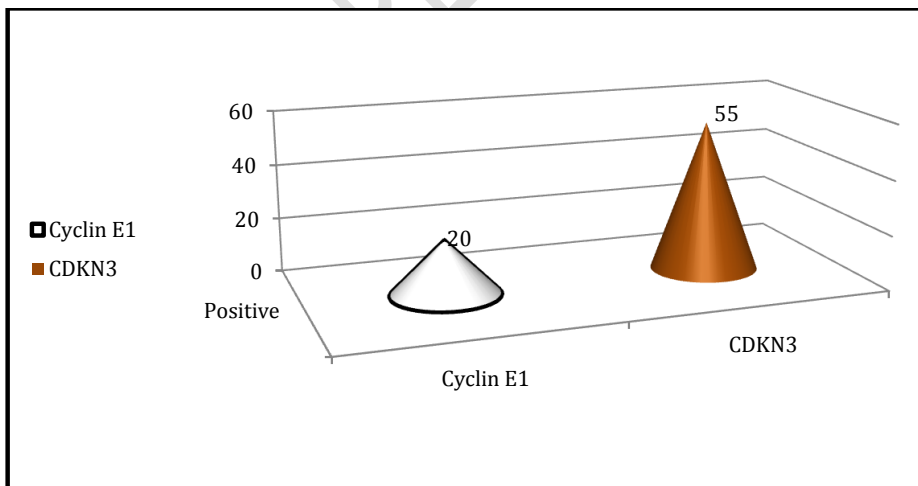


Fig 7: Circulation of Cyclin E1 and CDKN3 by using IHC.

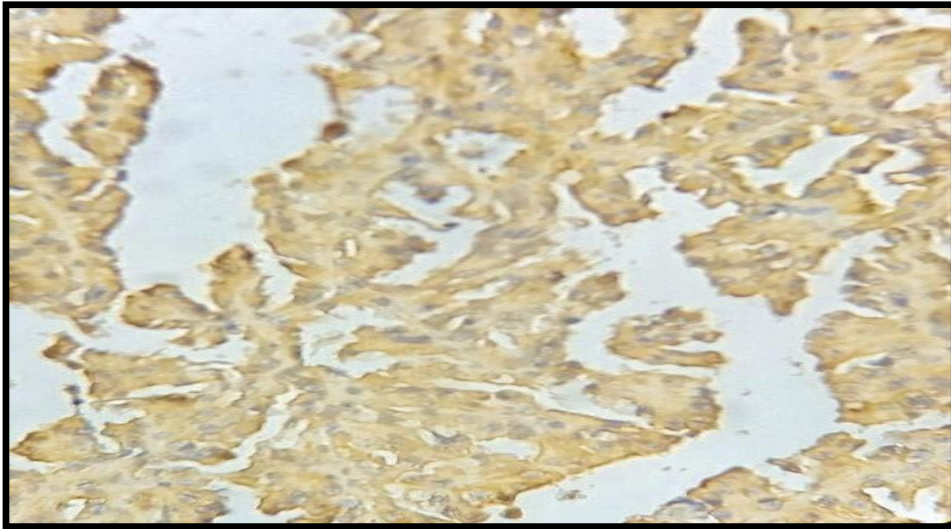


Fig8: Negative cyclinE1 stain of papillary type of RCC patients

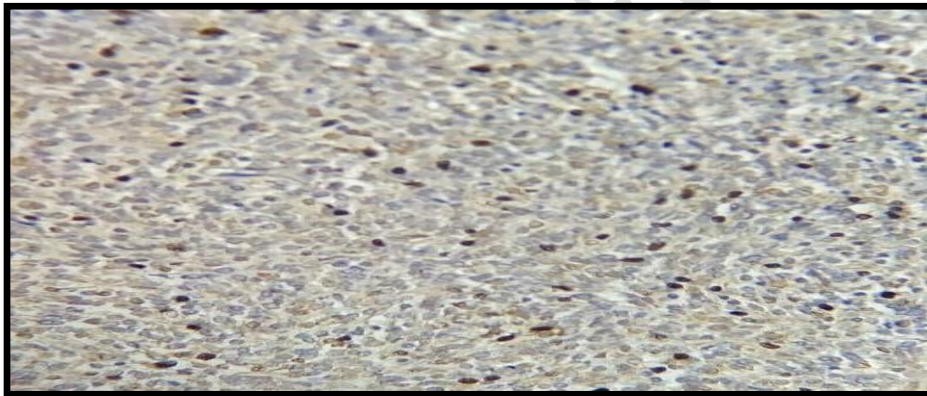


Fig9: Sarcomatoid carcinoma positive strong cyclin E1 stain score 2 (10 X40)

Discussion:

In this study, the existence of SV40 in blocked tissue taken from 75 patients suffering from RCC, it uses molecular technique involving PCR technique for detection of SV40 DNA united with immunohistochemistry technique (IHC) which are significant to verify the existence of Simian virus 40.

Simian virus 40 (SV40) define as a monkey virus which by accident entered to man, in 1955-1963 years, throughout polluted polio-virus vaccines that found the transforming and oncogenicity actions of T-Ag and t-Ag of this virus, which provoked examination of SV40 in human cancer. Generally, it is thought that contamination of polio vaccines were consider the major cause of infection with SV40 in humans, nearly all researches have define exposure of SV40 founded on vaccination (Engels *et al.*, 2003).

Most studies demonstrate that the kidney can function as a reservoir for SV40 in individuals. The sequence of this virus was reported in renal tissue and cells of urine sediments suffering from RCC (Li *et al.*, 2002) like Garcea and Imperiale, (2003) who found that SV40 causes infection in renal cells somewhere might possibly reactivation by immunosuppression. Also, Vanchiere *et al.*, (2005) reported that discovery of SV40 in renal tissue of human which indicates that kidney represented a position of viral latency, similar to in the usual simian host.

Bofill-Mas *et al.*, (2000) does not discover SV40 sequences in any tissue of RCC combine in diverse geographic regions of Europe and South Africa, while other types of polyomavirus sequences were detected from the majority of these tissues. In contrast to Manfredi *et al.*, (2005) who have failed to discover the sequences of SV40 in these tumor.

In molecular technique involving PCR, it was found that only 20 of 75 paraffin-embedded block tissue yielded SV40 for the reason that only extremely little amounts of these tissue block were offered for investigation, it was inspiring that DNA of SV40 recognized from 75 renal block tissue. The likelihood of occasional laboratory pollution of tissue block was excluded due to genetic material (DNA) linked with cancer and DNA of SV40 from laboratory progeny diverge sequences both within the viral regulatory area and at the carboxy terminus of T-Ag (Stewart *et al.*, 1998).

Some reports have lacking proof that SV40 was causation significant in the progression of human cancer but, Butel and Lednický, (1999) reported that the presence of the DNA of SV40 will suggest that the opportunity of these virus in the genesis of some RCC in human.

Bergsage *et al.* (1992) have revealed that negative SV40 outcome in renal tumor possibly because of utilize of few technical approaches. Also similarly, Leithner *et al.*, (2002) and Priftakis *et al.*, (2002) have recorded that never detected the sequences of SV40 in both Austria and Turkey, as in Sweden. While the predominance of SV40 DNA that are revealed in these cancers was diverse country for instance in Germany and Hungary (Heinsohn *et al.*, 2009).

Various reports recorded by Lopez-Rios *et al.*, (2004) showing that positive sequences of SV40 DNA by PCR technique as well as Mayall *et al.*, (2003) and Aoe *et al.*, (2006) reported that negative results by using quantitative PCR assay.

In general Iraq is considered as one of various countries in the Middle East regions that have special exciting to renal cell carcinoma and which regarded as the second mainly frequent urological malignancy (Ibrahim, 2013). As a result it is found the elevated proportion of males than females has in agreement with many studies finished by Vikram *et al.*, (2016) and Mahasin *et al.*, (2018).

Renal cell carcinoma is the majority frequent malignancy of kidney, as well as can be classified into five types including ccRCC, pRCC, chRCC, cd RCC and unclassified types. It is found in the presented study the most frequent type was clear cell RCC (42 of 75) which concordance with reports accomplished by Aiman *et al.*, (2013) and Mahasin *et al.*, (2018).

By using TNM classification of malignant tumors of RCC rely on the American Joint Committee on Cancer (AJCC), Stafford *et al.*, (2008) recorded that males patients have higher stage tumors while females patients have lower stage cancers, This is in concordance with our study. When in examination of the Fuhrman nuclear grade, Mukhopadhyay *et al.*, (2015) have discovered higher frequency of Grade 1 and lower frequency of Grade 3 and Grade 4.

The most common age group in their study is 61-70 years followed by 51-60 years. These results are in conformity with numerous reports such as Noroozina *et al.*, (2014) Khafaja *et al.*, (2015) and Hassan *et al.*, (2017) while unlikeliness with Latif *et al.*, (2011) and Takure *et al.*, (2013).

In immunohistochemical technique, the immunohistochemical indicators are significant in identifying RCC patients that are investigated by the EnVision System, This is in agreement with the report done by Lai *et al.*, (2017) who have recorded that a elevated expression of CDKN3 in renal tissues whilst Bisteau *et al.*, (2014) have found that tough expression of cyclin E1 which is related with poor prognosis of patients.

Also, Brousseau *et al.*, (2005) have unsuccessful to discover Tag of SV40 in these tumor by using immunohistochemistry technique with a extremely sensitive technique in spite of actuality that recorded in experienced tissues have DNA sequences of SV40.

The results of analysis of DNA SV40 polyomavirus by PCR in patient with RCC as; the total number of positive results of PCR is 20 (26.7%) whilst the negative results of PCR is 55 (73.3%).

Conclusion:

Finally, the dysregulation of CDK regulators in renal cell carcinoma associated with SV40 highlights the complex molecular pathways involved in the etiology of cancer. In addition to expanding our knowledge of the condition, the findings of this study open the door for the creation of tailored treatments meant to counteract SV40-related renal cell carcinoma by reestablishing the equilibrium of cell cycle regulation.

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Comment [RB7]: Add recent references. 70% References should be of past 5 years and not below that.

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