Viral Genetic Evolution in Host Cells Supports Tumorigenesis

Daniel A Achinko*

University City Science Center, USA

Background

Viruses are considered the smallest organisms and known to be metabolically inert out of a host cell but become active when they integrate and infect a host cell in order to reproduce. In the absence of a host cell, viruses exist as a capsid or a protein coat and at times within a membrane. Viral genetic material could be DNA or RNA enclosed within the capsid and encoding viral elements [1]. Viruses cause mild to severe diseases and sometimes result in epidemic outbreaks observed for Ebola in 2014 and Zika in 2016 [2]. The role of viruses in cancers has undergone several experimental analyses over the years and now, recent evidence shows that at least 6 viruses contribute to human malignancies and they include but not limited to: human papilloma virus (HPV), Epstein-Barr virus (EBV), human T-cell lymphotropic virus (HTLV-1), Hepatitis C virus (HCV), Kaposi's associated sarcoma virus (KSHV) and Hepatitis B virus (HBV) [3]. These viruses contribute to about 10 to 15% of worldwide cancers with about 1.3 million fatalities occurring annually [4]. These human tumor viruses differ in their content of their genetic material with some identified as DNA related viruses (EBV, HPV, HBV, KSHV) while the others are RNA related viruses (HTLV-1 and HCV). The difference in genetic material could suggest different entry patterns used by viruses to penetrate their host and how they regulate host cell transcription machinery for their individual pathogenic gains.

Despite the identification of cancer several years ago, their association to viral infection etiology remained a challenge until experimental analysis was able to show that, a virus was an agent with transmission capacity after filtration [5], hence leading to the identification of Rous sarcoma virus (RSV), and an avian virus that infected chickens [6]. It was considered the first identified tumor retrovirus transmitting chicken leukemia. Human warts were also associated with a viral etiology which only got recognition several years later wherein wart viruses also known as papilloma viruses were associated with cancer [7]. The identification and description of a B-cell malignancy in a Ugandan child was associated with a viral etiology and later called Burkitt’s lymphoma [8]. Epstein-Barr virus was a different kind of herpesvirus, later identified through electron microscopy as the pathogen associated with Burkitt’s lymphoma [9]. Laboratory analysis of several cancer types now links EBV to different cancer types including: gastric carcinoma, nasopharyngeal carcinoma, several cases of Hodgkin’s lymphoma and B-cell lymphoma in patients who are immune compromised [10,11]. Viral etiology to hepatitis went from being a suspicion because of its infectious mode of transmission, to the identification of a blood antigen as a surface protein of DNA viruses [12], shown to react with antibodies from hepatitis patients [13], hence called hepatitis B virus. Studies now associate HBV to hepatocellular carcinoma (HCC) resulting to about three hundred and fifty million infected individuals and three hundred thousand deaths, annually [14]. The advancement of genetic technology and research analysis to further understand viral roles in tumorigenesis led to the discovery of HTLV-1, HCV and KSHV during the 1980s and 1990s.
HCV infected patients were identified to possess an RNA virus agent with a transmission pattern different from HBV and shown to infect more than 270 million people yearly with about 4-7% of them evolving to HCC [15]. HTLV-1 was identified in cultured human T-cell lymphoma cells. Between 10-20% of individuals are infected annually by HTLV-1 and prognosis still remains elusive. KSHV affects less than 10% compared to 90% of individuals infected annually in the USA and Africa respectively. KSHV as a variation of herpesvirus happens to be the most common form associated with several cancer types and its related KS lesions generally occur in connective tissues below the skin further evolving in some cases to angiproliferative tumors [3].

Not all cancers are caused by viruses but from the discovery and evolutionary observation of viruses within cells, it is clear that viral associated cancers could involve several cell types within the body and their targets could be dominantly immune cells as seen for B-cell related viruses with genetic mechanisms associated with immortality of the cell and also T-cell related viruses inhibiting normal immune function. Almost all cancer types do not stimulate an immune response due to the identification of genes as "self" but viruses either stimulate intense immune response leading to cancer cells like HCC or down-regulate major histocompatibility class I and II genes expression and cell transport mechanism required for presenting HLA class I & II viral antigenic peptides to immune cells [2]. The ability of viruses to inhibit host immune response means they have evolutionarily learned host transcription malfunction associated with tumorigenesis, hence using it to successfully proliferate in cells. If the genetic pattern of integration of viral genetic material into host DNA and regulation of transcription and translation mechanism is well understood, then we could be closer to therapies that cut across cancer types.

Conclusion

Viral genetics and cancer formation is an intense area of research and herpesviruses and related cancers have been targeted through several forms of therapy. The ability of a given virus type like EBV, found to be associated with different cancer types, to infect different cell types, suggest different genetic regulatory patterns within the host cell, which is well orchestrated by the virus. These patterns could strongly depend on how the virus integrates into the host genome and taking control of transcription and translation processes. Underlying knowledge of this process will be of great therapeutic importance.

References