SHORT COMMUNICATION

Human herpes virus-8 in human immunodeficiency virus/acquired immunodeficiency syndrome - associated oral Kaposi’s sarcoma

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Keywords
Acquired immunodeficiency syndrome, human herpes virus-8, human immunodeficiency virus, oral Kaposi’s sarcoma

Abstract
Human herpes virus-8 (HHV-8), now called Kaposi’s sarcoma-associated herpes virus (KSHV), as a probable causative agent of Kaposi’s sarcoma (KS) was put forth when it was first detected in KS specimens in 1994. Since then, many investigators have confirmed the association of HHV-8 and human immunodeficiency virus/acquired immunodeficiency syndrome (HIV/AIDS) - associated KS. This study was done to review the original research aimed at assessing the presence of HHV-8 in HIV/AIDS-associated oral KS and other oral lesions.

Introduction
Kaposi’s sarcoma (KS), named after the Hungarian physician Moricz Kaposi, is a common vascular tumor seen in human immunodeficiency virus (HIV) infected patients and is one of the acquired immunodeficiency syndrome (AIDS) defining illness. AIDS-associated KS frequently involves the mucocutaneous regions of the head and neck as primary sites, in approximately 10% of all HIV-infected patients and in about 20% of patients in the homosexual/bisexual risk group. \(^1\) The oral cavity is involved in 50-80% of these cases, with the mucosa of the hard and soft palate, gingiva, and tongue being the most common locations. \(^2\)

This study aims to review the original research articles for assessing the prevalence percentage of human herpes virus-8 (HHV-8) in HIV/AIDS-associated oral KS.

Materials and Methods
A MEDLINE search was done using the keywords Kaposi’s sarcoma, oral Kaposi’s sarcoma, HIV-associated oral KS, AIDS-associated oral KS, HHV-8, and Kaposi’s sarcoma-associated herpesvirus (KSHV). The purpose of the search was to obtain original research articles pertaining to this review. A total of 8 articles could be included in the study; case reports being excluded. The details of the articles included for this review are listed in Table 1.

Based on the articles reviewed, the study sample could be categorized into three groups i.e. samples from:
1. HIV-positive (HIV(+) ) patients with oral KS.
2. HIV(+) patients with other oral lesions.
3. HIV-negative (HIV(−) ) patients with other oral lesions.

Data entry and statistical analysis were performed using SPSS version 10.0.5*. Analysis of variance (ANOVA) was done to compare the prevalence percentage of HHV-8 in each group.

Results
The prevalence percentage of HHV-8 (Table 2 and Graph 1):
- 82% in Group 1 i.e., HIV(+) oral KS samples
- 23% in Group 2, i.e., HIV(+) other oral lesion samples
- 0% in Group 3, i.e., HIV(−) other oral lesion samples.
The difference in prevalence percentage between the three groups was statistically significant ($P = 0.001$).

**Discussion**

It is clear that HHV-8 is present in a high proportion of samples from HIV(+)/AIDS-associated oral KS when compared to other lesions (Table 2 and Graph 1). Studies documenting the presence of HHV-8 in AIDS-associated KS have been extended to include the other types of lesions in HIV(+)/AIDS patients and HIV(−) patients. Furthermore, all the oral KS samples have been found to contain HHV-8 DNA sequences, and the HHV-8 copy number is higher in the KS lesions than in other groups. These studies suggest that HHV-8 are not an opportunistic infectious agent related to AIDS. Although some non-KS tissues (other oral lesions) have been found to contain HHV-8 sequences, most of these are from patients who either have cutaneous/systemic KS or are obtained from HIV(+) patients, and are likely to represent disseminated HHV-8 infection or be a result of small undetectable foci of HHV-8 in HIV/AIDS-associated oral Kaposi’s sarcoma.
KS. Furthermore, all the studies have reported a lack of these sequences in HIV(−) other oral lesions, including a wide variety of vascular tumors, as well as inflammatory conditions that resemble KS in their cellular composition. Thus, it appears that HHV-8 sequences are present specifically in oral KS tissues, as well as in some other oral lesional tissues from patients with KS or at high risk for developing KS, such as HIV infection.

Further support for a causal association between HHV-8 and KS comes from the analysis of peripheral blood from patients with KS, or at high risk of developing KS, compared with other AIDS risk groups and HIV(−) individuals. In these studies, the presence of HHV-8 DNA sequences in the peripheral blood of many of these patients has preceded the development of KS, and the presence of HHV-8 DNA sequences in the peripheral or semen of KS(-) patients has predicted the subsequent appearance of KS lesions in some patients. Furthermore, serological assays have been developed and used to analyze many serum samples from various patients’ population with or without KS. These studies have shown that the rate of HHV-8 seropositivity is high in HIV(+) patients with KS, and in HIV(+), KS(-), patients, but is much lower in HIV(−) individuals. An exception is finding of a high percentage HHV-8 seropositive individuals without KS, regardless of HIV status, suggesting that HHV-8 infection rates are higher in populations in whom KS is endemic. Furthermore, seroconversion to positivity for antibodies against HHV-8 related nuclear antigens has been found to occur before the clinical appearance of KS in a significant proportion of patients with AIDS-associated KS. Thus, the distribution of HHV-8 seropositivity indicates that HHV-8 varies among different populations but is not always ubiquitous. Studies so far suggest that HHV-8 is a sexually transmitted virus which closely relates with the risk for KS development, but also that there is a non-sexual route of transmission and that infection with HHV-8 does not necessarily imply the future development of KS.

Regardless of the incidence of HHV-8 infection in the general population, it is possible that the mechanisms operational in pathogenesis of the HHV-8 associated diseases could be explained by viral reactivation rather than primary infection. As in the case of other human herpesviruses, many conditions, including immunosuppression; immune stimulation; viral dose at the time of infection; and multiple genetic, environmental, and behavioral factors may cooperate in variable combinations in the pathogenesis of KS.

A different issue that has raised questions regarding the causal relationship between HHV-8 and KS is the observation that “KS” cell lines are negative for HHV-8, and cells cultured from KS lesions lose HHV-8 DNA after a few passages in vitro. However, in situ hybridization studies have shown the presence of HHV-8 in the spindle and endothelial cells, which are generally considered to represent the KS tumor cells. It remains to be determined whether HHV-8 is “lost” from these cells during culture or whether the cultured cells are different from the HHV-8 infected cells seen in the KS lesions.

Several issues regarding the incidence of HHV-8 infection and its relationship with the KS cells remain controversial and await further studies. However, the almost invariable presence of HHV-8 in all types of KS (virtually 100% of cases in most studies), its absence from other conditions that histologically resemble KS, its presence in the peripheral blood and tissues of patients with KS or at high risk for developing KS, and the patterns of immunoreactivity to HHV-8 antigens suggest an important, and most likely causal role for HHV-8 in the pathogenesis of KS.

Conclusion

An HHV-8 prevalence percentage of 82% in HIV(+) oral KS samples points to a possible causal role of HHV-8 in the multifactorial pathogenesis of HIV/AIDS-associated oral KS and HHV-8 infection as a high risk for the development of oral KS in HIV(+)/AIDS patients.

References


