Expression of Human Herpesvirus-6 Antigens in Benign and Malignant Lymphoproliferative Diseases

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Immunohistochemistry was used to look for the expression of human herpesvirus-6 (HHV-6) antigens in a well characterized series of benign, atypical, and malignant lymphoid lesions, which tested positive for the presence of HHV-6 DNA. A panel of specific antibodies against HHV-6 antigens, characteristic either of the early (p41) or late (p101K, gp106, and gp116) phases of the viral cycle, was applied to the lymphoid tissues from 15 non-Hodgkin’s lymphomas, 14 Hodgkin’s disease cases, 5 angioimmunoblastic lymphadenopathies with dysproteinemia, 14 reactive lymphadenopathies, and 2 cases of sinus histiocytosis with massive lymphadenopathy (Rosai-Dorfman disease). In lymphomatous tissues, the expression of late antigens was documented only in reactive cells, and mainly in plasma cells. Of interest, the expression of the early p41 antigen was detected in the so-called “mummified” Reed-Sternberg cells, in two Hodgkin’s disease cases. In reactive lymphadenopathies, the HHV-6 late antigen-expressing cells were plasma cells, histiocytes, and rare granulocytes distributed in interfollicular areas. In both cases of Rosai-Dorfman disease, the p101K showed an intense staining in follicular dendritic cells of germinal centers, whereas the gp106 exhibited an intense cytoplasmic reaction in the abnormal histiocytes, which represent the histological hallmark of the disease. The expression of HHV-6 antigens is tightly controlled in lymphoid tissues. The lack of HHV-6 antigen expression in neoplastic cells and the limited expression in degenerating Reed-Sternberg cells argue against a major pathogenic role of the virus in human lymphomagenesis. The detection of a rather unique pattern of viral late antigen expression in Rosai-Dorfman disease suggests a possible pathogenic involvement of HHV-6 in some cases of this rare lymphoproliferative disorder. (Am J Pathol 1998, 153:815–823)
HHV-6 antigens by immunohistochemistry has provided a unique and sensitive tool for the identification of infected cells both in fresh cellular populations and in archival tissue sections. Recently, this technical approach has been revealed to be successful in localizing HHV-6-infected cells in brain tissues from patients with multiple sclerosis, providing the first strong evidence toward an etiological relationship. Thus, we judged it appropriate to use immunohistochemistry with a panel of antibodies for different viral antigens to look for the presence and distribution of HHV-6 antigen-expressing cells in the lymphoid tissues positive for the presence of HHV-6 DNA from a well characterized series of patients with benign, atypical, and malignant lymphoproliferative diseases.

**Materials and Methods**

**Patients**

Tissues from 15 non-Hodgkin’s lymphomas (NHL), 14 Hodgkin’s disease (HD) cases, 5 angioimmunoblastic lymphadenopathies with dysproteinemia (AILD), 14 reactive lymphadenopathies, and 2 cases of sinus histiocytosis with massive lymphadenopathy (Rosai-Dorfman disease) were included in the study. In all cases of NHL and HD, the diagnosis was determined on the basis of the histological analysis of a lymph node biopsy, and revised according to the REAL classification. All of the NHL cases were subjected to standard immunophenotyping procedures as well as to Southern blot analysis of immunoglobulin (Ig) heavy and light chain genes and of T-cell receptor β-chain gene, to confirm either the B- or T-cell lineage.

The distribution of NHL cases according to lymphoma subtype was as follows: 4 follicular center, 2 mantle cell, 2 marginal zone B cell, 3 diffuse large B cell, 1 Burkitt’s, 2 peripheral T cell, and 1 anaplastic large cell. The distribution of HD cases according to subtype was as follows: 6 mixed cellularity, 5 nodular sclerosis, 2 lymphocyte predominance, and 1 lymphocyte depletion. The series of 14 reactive lymphadenopathies consisted of 5 cases of florid follicular hyperplasia, 4 cases with a predominantly paracortical lesion, 4 cases with sinus histiocytosis, and 1 histiocytic necrotizing lymphadenitis.

All of these tissue samples harbored HHV-6 DNA sequences, as detected by PCR, which was performed as previously reported. One NHL, two HD, and the five AILD cases have been previously described. In one NHL case and in two HD cases, the viral copy number was so high that it was detectable also by Southern blot analysis. Genotype characterization, performed by a PCR assay, showed the HHV-6 variant B genome in all cases, with the exception of three AILD cases, showing HHV-6 variant A genome in two cases and a mixture of HHV-6 variant A and B genomes in one case, as previously reported. Furthermore, tissues from five NHL, four HD, and five reactive lymphadenopathy cases, which tested negative for HHV-6 sequences by PCR, were also included in the study to serve as negative controls in immunohistochemical experiments.

**Immunohistochemistry**

Sections from formalin-fixed, paraffin-embedded tissues were stained with an avidin-biotin complex immunodetection system in a TechMate instrument (BioTek Solutions, Santa Barbara, CA). Sections (4 μm) mounted on positively charged slides were deparaffinized with xylene and rehydrated, immersed for 10 minutes in 3% hydrogen peroxide/methanol to quench endogenous peroxidase, and microwaved for 10 minutes in citrate buffer. After slow cooling, the sections were incubated successively with primary antibody, biotin-labeled goat anti-mouse or anti-rabbit Ig, streptavidin-biotin peroxidase complex, and 3,3’-diaminobenzidine tetrahydrochloride chromogen. Tissues were counterstained with hematoxylin. Mouse IgG1 monoclonal antibody (mAb) to HHV-6B virion protein p101K (late antigen) was obtained from P. Pellett (Centers for Disease Control and Prevention, Atlanta, GA) and from Chemicon International (Temecula, CA) and used at a dilution of 1:200. Mouse IgG2a mAb C5 to DNA binding protein p41 (early antigen) was used at a dilution of 1:50 (Biodesign International, Kennebunkport, ME). Mouse IgG2b mAb to gp106 (late antigen) and IgG2b mAb to gp116 (late antigen) (both from Advanced Biotechnologies, Inc., Columbia, MD) were used at a dilution of 1:50. The specificity of these antibodies was confirmed by testing isotype-matched control mouse mAbs against human IgG (IgG1, X931; IgG2a, X943; and IgG2b, X944; Dakopatts, Glostrup, Denmark) in selected HHV-6-positive cases. Control and test antibodies were used at the same IgG concentrations. Antibodies to other herpesviruses included the anti-human cytomegalovirus mAbs DDG9 and CCH2 (Dakopatts) to early and immediate-early antigens, rabbit polyclonal anti-herpes simplex virus 1 antibody B114 (Dakopatts), and anti-Epstein-Barr virus (EBV) mAb to latent membrane protein-1 (LMP-1) (Dakopatts). The antibody panel also included CD3, UCHL-1 (CD45RO), L-26 (CD20), KP1 (CD68), S-100, CD21, Ber-H2 (CD30), and anti-Epstein-Barr virus (EBV) mAb to latent membrane protein-1 (LMP-1) (Dakopatts). The antibody panel also included CD3, UCHL-1 (CD45RO), L-26 (CD20), KP1 (CD68), S-100, CD21, Ber-H2 (CD30), and EMA/E29 (all from Dakopatts), and Leu-M1 (CD15; Becton-Dickinson, San Jose, CA). All antibodies were used as recommended by the manufacturers.

**Results**

**Immunohistochemical Localization of HHV-6 Antigens in Lymphoid Tissues**

**NHL**

The expression of HHV-6 antigens was investigated by immunohistochemistry in 15 NHL cases, which tested positive for HHV-6 DNA by PCR. One of these cases was also positive by Southern blot analysis, indicating high copy number latent integration of the HHV-6 genome, as previously reported. Neoplastic cells were consistently negative for HHV-6 antigens in all NHL tissues examined, even in the B-NHL case (follicular center type) with so a
high copy number that it tested positive for HHV-6 DNA by Southern blot analysis. Cytoplasmic staining was observed with the antibody p101K in rare plasma cells interspersed among neoplastic cells and in some isolated spindle-shaped stromal cells located in the lymph node capsule and in the surrounding fibroadipose tissue. No reactivity was observed with antibodies gp106, gp116, and p41.

**AILD**

In five cases of AILD, previously shown to be positive for HHV-6 DNA by PCR, the immunostaining was observed with antibody p101K, gp106, and gp116 in scattered plasma cells. In two cases, focal collections of positively staining plasma cells were present (Figure 1A). No reactivity was observed with antibody p41.

**Figure 1.** A: AILD, positive staining with p101K antibody in a focal collection of plasma cells. Immunoperoxidase method; magnification, ×170. B and C: HD, nodular sclerosis type; positive staining with p41 antibody in R-S cells with shrunken cytoplasm and pyknotic nuclei (mummified cells). Immunoperoxidase method, ×330. D: HD, nodular sclerosis type, positive staining with p11 antibody in mummified R-S cells compared with negative staining in a viable R-S cell (right side). Immunoperoxidase method, ×550.
HD

The expression of HHV-6 antigens was investigated by immunohistochemistry in 14 lymph nodes affected by HD that were positive for HHV-6 DNA by PCR. Two of these cases were positive also by Southern blot analysis, as previously reported. In all cases, complete effacement of the lymph node architecture was observed, and no remnants of previous normal structure could be detected. Only rare cells were found to exhibit cytoplasmic reaction for HHV-6 antigens, as evidenced by the p101K and gp116 antibodies. No staining was observed with the gp106 antibody. The morphology of the positive cells was that of reactive histiocytes and plasma cells. Isolated granulocytes also showed a positive reaction. Reed-Sternberg (R-S) and Hodgkin cells were consistently negative for HHV-6 antigens.

The p41 antibody stained rare plasma cells and isolated granulocytes in all cases. Of interest, in the two cases with a high enough copy number of HHV-6 sequences to be detectable by Southern blot analysis, this antibody also showed positive immunohistochemical staining in R-S cells with shrunken cytoplasm and pyknotic nuclei. These were considered to represent degenerating “mummified” cells. No staining was observed in viable R-S and Hodgkin cells with antibody p41 (Figure 1D). In the same cases, there was clear evidence of staining of rare plasma cells and granulocytes in the reactive component, in addition to the positive mummified cells.

The HD cases were also examined with anti-EBV mAb to LMP-1. Positive reactivity was observed in seven cases and restricted only to R-S and Hodgkin cells, whereas reactive lymphoid cells were consistently negative (not shown). Of note, in the two HD cases with mummified cells positive for HHV-6 p41, immunostaining for EBV LMP-1 was negative both in viable R-S and Hodgkin cells and in mummified cells. Moreover, using immunohistochemistry for EBV LMP-1 and for four HHV-6 antigens, we could not demonstrate the co-infection of the two viruses in lymphoid cells, as the only lymphoid cells expressing HHV-6 antigens were rare plasma cells, which invariably stained negative for LMP-1.

Nonneoplastic Lymph Nodes

Fourteen nonneoplastic lymph nodes, including five cases of florid follicular hyperplasia, four cases with a predominantly paracortical lesion, four cases with sinus histiocytosis, and one histiocytic necrotizing lymphadenitis as well as two cases of Rosai-Dorfman disease, were studied for the expression of HHV-6 antigens. All of the examined cases were positive for HHV-6 DNA sequences by PCR.

In most cases, only isolated cells stained positive with the HHV-6-specific antibodies p101K, gp106 and gp116. Antibody p41 showed no reaction. The number of immunopositive cells was always less than 1% of the total lymph node cell burden, being usually lower than 0.5%. No staining was observed in the single case of histiocytic necrotizing lymphadenitis. Positive cells were found to correspond to plasma cells and histiocytes, whereas endothelial cells, follicular dendritic cells, and different types of lymphoid cells (lymphocytes, centrocytes, centroblasts, and immunoblasts) were consistently immunonegative. Isolated granulocytes, present in dilated sinuses in two cases of reactive lymphadenopathy with sinus histiocytoses, showed a positive reaction. Positive cells were not homogeneously distributed in the nodal tissue but appeared to be scattered in interfollicular regions (Figure 2A). Hyperplastic germinal centers were usually negative, showing a bland, doubtful staining only in one case.

An interesting reactivity with p101K and gp106 antibodies was observed in both cases of Rosai-Dorfman disease. The p101K antibody exhibited an intense staining in follicular dendritic cells of germinal centers (Figure 2B), both hyperplastic and burned out, present in areas in which the normal nodal architecture was still retained. In the two Rosai-Dorfman disease cases, adjacent tissue sections were stained either with anti-HHV-6 p101K or with CD21 antibodies, respectively. The same distribution of positive cells in the follicles was observed with both antibodies, providing further evidence that the HHV-6-positive cells in the follicles corresponded to follicular dendritic cells. With antibody gp106, an intense granular positive reaction was observed in the cytoplasm of abnormal histiocytes located within distended sinuses (Figure 2, C and D). Emperipolesis was present inside some positive cells (Figure 2, C and D). The distribution of the staining was not uniform throughout the lymph nodes but tended to be localized in discrete regions. A weak positive reaction was detected in isolated plasma cells with both p101K and gp106 antibodies (Figure 2C).

Specificity of Immunohistochemical Localization of HHV-6 Antigens

The authenticity and specificity of the immunohistochemical results was supported by the following series of findings. First, we could document a good concordance between the results obtained by PCR/Southern blot analysis and the immunohistochemical assay. In fact, all of the specimens harboring HHV-6 DNA sequences, as detected by molecular methods, were also positive at least with one anti-HHV-6 antibody. Conversely, we observed the complete unreactivity of the anti-HHV-6 antibodies (p41, p101K, gp 106, and gp 116) on the lymph node sections from control specimens that tested negative for HHV-6 DNA by PCR. Second, the specificity of the staining reaction was confirmed by testing isotype-matched control mouse mAbs against human IgG (IgG1, X931; IgG2a, X943; and IgG2b, X944; Dakopatts) in selected HHV-6-positive cases (Figure 3, A and B). Control and test antibodies were used at the same IgG concentrations. Furthermore, selected HHV-6-positive specimens that stained positively with antibodies to HHV-6 did not show a positive immunohistochemical reaction when reacted with an mAb to human cytomegalovirus, with a polyclonal antibody to herpes simplex virus 1, and with mAb to EBV LMP-1. On the other hand, it should be noted
that controlled studies have ascertained that these antibodies do not display cross-reactivity to other herpes or common human viruses. Finally, the specificity and suitability of p101K and p41 antibodies for in situ detection of HHV-6 antigens in human tissues have been already successfully reported by one of us (RG).

**Discussion**

A pathogenetic association between HHV-6 infection and NHLs has been previously suggested on the basis of the following findings: 1) the first isolation of the virus from the peripheral blood of patients with NHL, related to acquired immune deficiency syndrome or not; 2) the detection of HHV-6 sequences in the lymphomatous tissues by PCR and, although rarely, also by Southern blot analysis; and 3) the very recent establishment of an HHV-6 latently infected cell line from the pathological tissue of one case of HHV-6-positive and EBV-negative Burkitt’s lymphoma. Identification of the viral genome in the neoplastic cells certainly represents one of the most important and classical parameters to suggest an etiological relationship between an oncogenic DNA virus and...
a human tumor. Because DNA *in situ* hybridization studies have invariably failed to document the presence of the viral genome in the lymphomatous cells, we judged it appropriate to apply a sensitive immunohistochemical technique, looking for the possible expression of viral antigens, occurring during the early or late phase of the viral cycle, in a series of HHV-6-infected lymphoma tissues. The expression of viral antigens is restricted to reactive cells and is invariably absent in neoplastic cells in all cases examined. A hit-and-run mechanism of HHV-6, in which HHV-6 might have contributed to the initial transformation of lymphoid cells and then is released by the cells themselves once they become fully malignant, is always possible but cannot be proven. Similarly, indirect effects of herpesvirus-infected reactive cells on the neoplastic clone, within the lymphomatous lesion, is suggestive but still speculative, in the case of HHV-6 infection. The negative results of our immunohistochemical study, performed with the largest panel of antibodies to HHV-6 antigens now available, argues against a major "orthodox" role for HHV-6 infection in the occurrence of human NHLs.

We were the first to document an unusually high frequency of HHV-6 sequences in the pathological tissues of AILD, by PCR. Using immunohistochemistry in the same series of cases, we showed that the expression of viral antigens is absent in T lymphocytes, which are considered the proliferating elements in such a disease, arguing against a direct role for HHV-6 in the development of this atypical lymphoproliferation. The frequent presence of HHV-6 DNA may be also related to a reactivation or to a primary infection of this herpesvirus, favored by the immunesuppression of AILD patients, who, indeed, often die from severe opportunistic infections. However, the fact that the expression of HHV-6 antigens is limited to a small proportion of cells and restricted to plasma cells, suggests that an active disseminated infection with HHV-6 is uncommon, even under conditions of severe immune impairment, which are typical of AILD.

A pathogenetic association between HHV-6 infection and the development of HD is apparently more solid and based on the following epidemiological evidence, obtained by our group and others: 1) higher frequency and higher titers of anti-HHV-6 antibodies in HD patients than in NHL patients or in blood donors, and 2) correlation between anti-HHV-6 antibody titers and the clinical course and prognosis of HD, and 3) higher frequency of HHV-6 sequences by PCR and Southern blot analysis in HD than in NHL cases. Again, so far, *in situ* DNA hybridization studies have failed to identify HHV-6 in the putative neoplastic cells of HD, namely in Hodgkin and R-S cells and have simply documented the presence of viral genomes in normal, reactive lymphocytes. We provide here the first evidence that, although rarely, HHV-6 may infect R-S cells. HHV-6 infection is latent, as only p41, an early antigen, is expressed. Infected R-S cells represent the so-called mummified cells, ie, R-S cells that have undergone apoptosis. It is not possible to assess whether viral infection of R-S cells has been favored by the apoptotic process of the cells or the virus has triggered the programmed cell death of the cells themselves. Relevant to this, HHV-6 has been shown to induce apoptosis in infected T cells *in vitro*. If an apoptotic effect of HHV-6 on R-S cells is confirmed, the virus might have a protective rather than a pathogenetic role in such a disease. Of interest, HHV-6-expressing R-S cells could not be identified in HD cases with a low HHV-6 copy number (ie, PCR positive), but only in the two cases of HD harboring an extraordinarily high amount of viral DNA, sufficient to test positive also by Southern blot analysis. We are well aware of the fact that mummified cells may be prone to nonspecific staining. However, it should be noted that mummi-
fied cells stained positive only with p41, but stained neg-
ative with at least three of three anti-HHV-6 antibodies for late
antigens (p101K, gp106, and gp116), and with antibo-
dies to three herpes viruses (cytomegalovirus, herpes sim-
plex virus, and EBV). Moreover, in two HHV-6 DNA-neg-
ative HD cases examined as controls, mummified cells
were present but did not react with p41. These data
definitely argue against a nonspecific staining of mum-
mified cells at least for a early HHV-6 antigen, recognized
by p41 antibody, in the two HD cases with high viral load.

In conclusion, as R-S cells in the vast majority of HD
cases resulted negative for the expression of viral anti-
gen, a causative role for HHV-6 infection in the devel-
opment of HD cannot be assessed. However, as we
show, for the first time, that HHV-6 may have a tropism for
R-S cells in vivo, the possibility exists that other as-yet-
uncharacterized proteins may be expressed in HD tis-
ues. Relevant to this, we recently identified the first
HHV-6 protein, called ORF-1, which has a transactivating
and transforming activity mediated through the binding to
p53.42 As we already detected the sequences encoding
for ORF-1 in HD tissues in vivo,20 we are currently looking
for the expression of this oncprotein in R-S and Hodgkin
cells. Furthermore, similarly to what occurs in AILD,
HHV-6 infection of HD tissues is mainly a latent infection,
and the expression of viral antigens indicative of a repli-
cative cycle was limited to a small number of cells and
restricted to few cell types, including plasma cells. Simi-
larly, in cases of EBV-associated HD, the expression of
lytic genes of EBV is a very rare phenomenon in R-S
cells.39 Moreover, the expression of HHV-6 antigens in
plasma cells, which seems a constant finding in HHV-6-
associated lymphoproliferative diseases, again reminds
us of the behavior of EBV. For example, in the EBV-
duced lymphomas arising in the severe combined immu-
nodeficiency mouse model, EBV-infected tumor cells
show a plasma cell phenotype and a reduced expression
of viral latent genes.40

Although HHV-6 primary infection is a well-recognized
cause of mononucleosis-like illnesses,5 there is not a
histological pattern of the lymph node that has been
considered characteristic of HHV-6 infection, as paracor-
tical expansion is generally typical of EBV-induced infec-
tious mononucleosis,41 or giant germinal center hyper-
plasia with increased vascularity seems to be often
related to human herpesvirus-8 infection of the lymph
nodes.42 That is why we judged it appropriate to investi-
gate a highly heterogeneous series of reactive lympha-
nopathies, with different histological features, selected
simply on the basis of their positivity for HHV-6 DNA by
PCR. EBV infection may have different patterns of infec-
tion in nonneoplastic lymph nodes. Indeed, in typical
EBV-induced infectious mononucleosis, large numbers of
EBV-positive B lymphoid blasts are detectable in extrafol-
licular areas, whereas germinal centers are free of EBV-
positive cells.41 In other reactive lymph nodes harboring
EBV sequences, EBV-positive cells are small lympho-
cytes generally present in extrafollicular areas but, some-
times, also in germinal centers.41 In our extensive immu-
nohistochemical study, we did not observe different
patterns of HHV-6 antigen expression in the various types
of lymphadenopathies examined. The localization of
HHV-6-expressing cells in interfollicular areas and their
absence in germinal centers were a constant feature,
whereas the infected cell types were the same, ie,
plasma cells, histiocytes, and granulocytes in all cases
examined. It should also be noted that lymphocytes,
which tested invariably negative for HHV-6 antigens,
seem to be not permissive for viral replication, in contrast
to what occurs in vitro.12 The possibility still exists that
as-yet-uncharacterized viral proteins may be expressed
in infected lymphocytes, in vivo.

A very peculiar pattern of expression of HHV-6 anti-
gen was documented in both cases of Rosai-Dorfman
disease. The expression of p101K antigen in follicular
dendritic cells seems to be a rather specific phenome-
non, as it tested absent in all of the other lymphadenop-
athy cases examined. It has been previously reported
that EBV is able to infect and transform follicular dendritic
cells in vitro.43 Infection of follicular dendritic cells with
herpesviruses may be an underestimated phenomenon
that merits further investigation. Clearly, the most inter-
esting finding is represented by the demonstration of a
HHV-6 protein in the late phase of the viral cycle in a
significant proportion of the abnormal histiocytes, which
represent the hallmark of the Rosai-Dorfman disease. Our
study extends the results by Levine and colleagues,44
who documented the presence of HHV-6 DNA in such
cells by in situ DNA hybridization. The novelty and the
interest of our finding is that we provide the first evidence
that HHV-6 not only may infect the abnormal histiocytes
of Rosai-Dorfman disease but, more importantly, it is func-
tionally active in such cells. Although the diagnostic cells
of Rosai-Dorfman disease have many histiocyte-associ-
ated morphological features, they also have an uncom-
mon phenotype, as they express monocyte/macrophage-
associated markers and the dendritic cell-associated
marker S-100.45 Of interest, the expression of HHV-6
genes in lymphoid cells/tissues, either normal or patho-
logical, as detected by Northern blot analysis, is a very
rare phenomenon, documented so far only in the neo-
plastic cells of two cases of the rare S-100-positive T-cell
chronic lymphoproliferative disease,46 suggesting that
S-100-positive cells are in some way more permissive for
HHV-6 replication. Rosai-Dorfman disease has been pro-
posed to represent an exaggerated immunological re-
sponse to an infectious agent, and a pathogenetic asso-
ciation with EBV has been previously reported.44,45 Thus,
we suggest that HHV-6 should also be investigated as
another possible trigger of the uncontrolled proliferation
of abnormal histiocytes in such disease.

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