Herpes Simplex Virus Type 1 Promoted the Possibility of Leukemia Caused by L6565 Murine Leukemia Virus

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Abstract: Herpes Simplex Virus Type 1 (HSV-1) is a virus commonly existent in nature. Others reported that herpesvirus DNA probably activates type C virus in vitro. L6565 Murine Leukemia Virus (L6565 MuLV) is a kind of virus inducing the murine Leukemia. The main aim of the study was to explore the activation of HSV-1 to the pathogenicity of L6565 MuLV in vivo. Suckling mice were inoculated with either L6565 MuLV & inactivated HSV-1, or L6565 MuLV alone. In order to determine the infection of the mice, HE staining and RT-PCR techniques were applied. The result shows that the L6565 MuLV RNA was detected in 8th week in the mice of MuLV plus HSV-1 group, while delayed to 10th week in the L6565 MuLV alone group. The infection rates of the two groups were 74.3% and 41.7%, respectively. Also, L6565 MuLV RNA was detected in the thymus one week after the inoculation, and it was detected in both thymus and spleen two weeks after inoculation. It is obvious that HSV-1 promoted the infection of L6565 MuLV and increased the incidence rate of murine leukemia induced by L6565 MuLV.

Keywords: Mouse leukemia, L6565 MuLV, HSV-1.

INTRODUCTION

Murine Leukemia Virus (MuLVs) is a kind of lentivirus that belongs to the Gammaretrovirus family (i.e., type C virus) [1]. It can induce a number of different types of lymphomas/leukemia in mice, and the main signs of the disease are lymphopenia and immunity descends [2]. Currently, there is some ongoing research on Moloney MuLV, and Friend MuLV [2-5]. L6565 Murine Leukemia Virus (L6565 MuLV) is another kind of MuLVs, which was set up in China by Zhen BF, et al. in Department of Pathophysiology, Shanghai Medical University [6]. It offered a new method to study the pathogenicity of leukemia virus. Then Cheng L, et al. in the same department established a viral L6565 cell clone (L6565 Lymphocytes leukemia cell) [7], which was used extensively in MuLV pathogenicity studies [8, 9]. It was then proven by us that the cell causes Murine Leukemia [10]. Evaluation of the pathogenicity of L6565 MuLV is based on the L6565 cell clone in this study.

Herpes Simplex Virus Type 1 (HSV-1) is a type of Herpes Simplex Virus that belongs to the Herpesvirus family [1]. It is a kind of virus commonly existent in nature. About ninety percent population is in latent infected. Upon infection, the virus is transported by way of axons in the sensory nerve endings to the neurons of sensory ganglia, where it remains latent, and serves as a latent reservoir for recurrent disease in the case of immunity descends or breakdown. Previous studies on the relationship of Leukemia Virus and HSV-1. ANN L, et al. have reported that herpesvirus DNA probably activates type c virus in vitro [11]. Lasky, et al. also proposed that the expression of retroviral proteins in Ramos lymphoma lines is enhanced after conversion with Herpesvirus [12]. It is suggested that the pathogenicity of MuLV is perhaps related to HSV. An attempt was made to study if HSV-1 promotes the development of Leukemia induced by L6565 MuLV, in vivo. Whereas others have not reported anything.

MATERIAL AND METHOD

Cells and Viruses

The L6565 Lymphocytes leukemia cells, which express L6565 MuLV, are obtained from the Department of Pathophysiology, Shanghai Medical University, China. The cell line was maintained in DMEM containing 10% fatal bovine serum, cultured at 37°C in 5% CO2. HSV-1 was propagated in Hep-2 cell line, and inactivated by ultraviolet radiation at 50W, 15cm for 10 min before use. Both viruses are preserved in our institution. The Coxsackie’s B3 virus was used as a negative control in RT-PCR. It was also propagated in Hep-2 cell line and preserved in our institute. All the viruses were filtered by 0.45 μm Millipore before inoculated into mice.

Mice

Kunming suckling mice, born in 24h, were used for all experiments in this study. Pregnant mice were obtained from Experimental Animal Center of Wuhan University, which maintained and observed individual cages in the holding room before delivery. Births were timed to the nearest 1/2 day.

Newborn Kunming mice were randomly divided into 3 groups in this experiment, they are L6565 MuLV and HSV-1 co-inoculated group, group L6565 MuLV and group control. 0.2ml of supernate of L6565 Lymphocytes leukemia cells were injected intraperitoneally, and repeated again the next
day. 100TCID50 HSV-1 was injected with 0.02ml in hypodermis once a week for the first 2 weeks to sustain the HSV-1 antigen in a persistent latent infection.

**Tissue Sampling and Preparation**

Two suckling mice of each group were killed every week until MuLV RNA can no longer be detected in the mice (the 14th week according the RNA detection). 16 mice in the L6565 MuLV group and 20 mice in co-infected group were killed, in total. All the main viscera, such as thymus, spleen, lymph node, liver, kidney, and peripheral blood were taken for further studies. Half of them were fixed in 10%formalin, embedded in paraffin, and sections were stained with hematoxylin and eosin, and Masson’s trichrome. While the other half were immediately frozen at −70 °C and later used for RNA detection for the determination of L6565 MuLV.

**RNA Isolation and RT-PCR Assay**

For the determination of the L6565 MuLV genes expression, total RNA was extracted from each tissue of mice with TRIzol Reagent (Invitrogen) as described in the manufacturer’s protocol. RNA quality was assessed by electrophoresis on 1.5% agarose gel. And to synthesize cDNA by reverse transcription, 2 ul total RNA, 200 units M-MuLV reverse transcriptase (Promega, USA) and 0.5uM oligo dT primer ([5'-GGCCACCGTGCTCAGACTGAC(T)16(A/C/G)-3']) were reacted for 1 h at 37 °C in 25ul reaction mixture containing 1mM dNTP in first strand buffer according to the manufacturer’s instruction. PCR amplification was performed with the following described primers: forward primer 5’-GAGACTGTTGGACCAGGGAA-3’ (sense strand correspond to 2834—2854 genome) and a reverse primer 5’-TTGTCTCTGAGATCCCAT-3’ (antisense strand correspond to 3125—3143 genome). RNA of L6565 MuLV suspension and Coxsackie’s B3 virus were used as positive and negative controls. The PCR reaction was carried out for 30 cycles: 45s at 92 °C, 45s at 57 °C and 60 at 72 °C, and last extend at 72 °C for 5 min. PCR products were analyzed on 1.5% agarose gels for 1h at 80V.

**DNA Isolation and PCR Assay**

Genomic DNA was extracted from each mouse tissue as described by the method below. Tissue homogenates were layered over a 10-ml shelf of 35% sucrose (wt/wt) in 10 mM Tris/1 mM EDTA, PH 7.4, and the virus was pelleted at 20,000 rpm for 1 hr in an SW-27 rotor. Wash again and treat with RNase A and proteinase K for 15min at 68°C and then overnight at 37°C. Genomic DNA was extracted with phenol:chloroform(1:1) twice before use. The isolate was detected by NA Analysator (Eppendorf) to make sure there is no RNA carry over. Then the PCR reaction was carried out in the same conditions as cDNA replication above.

**RESULTS**

**Infection Comparing**

The number of infection in mice was determined by RNA detection. Viral RNA in the L6565 MuLV plus HSV-1 group is detected 8 weeks after inoculation, while viral RNA in the L6565 MuLV alone infected group is detected 10 weeks after inoculation. In the 14th wk (when there was no L6565 MuLV RNA detected), 35/36 mice remained in the experimental group. Among these mice, the infection rate of L6565MuLV plus HSV-1 group is higher than that of L6565MuLV group (Table 1). Plus, mice that were infected with L6565 MuLV showed signs of infection such as diminished activity and hair stood up.

**Table 1. L6565 MuLV Infected Condition of Each Group**

<table>
<thead>
<tr>
<th>Group</th>
<th>Mice Number</th>
<th>MuLV Infected Number</th>
<th>The First Time of RNA Detected</th>
<th>Infection Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L6565MuLV</td>
<td>36</td>
<td>15</td>
<td>10wk</td>
<td>41.7</td>
</tr>
<tr>
<td>L6565MuLV+HSV-1</td>
<td>35</td>
<td>26</td>
<td>8wk</td>
<td>74.3*</td>
</tr>
<tr>
<td>normal control</td>
<td>21</td>
<td>0</td>
<td>-</td>
<td>0.0</td>
</tr>
</tbody>
</table>

*Compared two group X2=7.74; P<0.01.

**Morphology Comparing**

One mouse in the L6565 MuLV plus HSV-1 group developed leukemia, and had greatly enlarged thymus, spleen, liver and lymph nodes. The other mice were in the early phase of leukemia. These include all mice in L6565 MuLV group, which showed a slightly/moderately enlarged spleen and whitened liver. Peripheral lymph nodes and thymuses of these mice were slightly enlarged even at the advanced stage of the disease. The heart, lungs and kidneys of all the mice did not show any obvious changes. Fig. (1) shows the changes of the spleens in erythroleukemia.

**Fig. (1). Spleens of a normal mouse (A: HE x 100) and a mouse with erythroleukemia (B: HE x 100): (A) megakaryocyte (Δ) is showed in the erythroleukemia tissue slice (B).**
L6565 MuLV RNA and DNA Detection

In the L6565MuLV plus HSV-1 group, the L6565 MuLV RNA could be detected as early as 1st week after inoculation and as late as 14 weeks after inoculation. Viral RNA could be detected earlier in mice of the L6565MuLV plus HSV-1 group than in mice of the L6565 MuLV group. The data is shown in Table 2 and Fig. (2). Also, the amplification of RNA and DNA reveals the same amplification fragment (310bp), shown in Fig. (3).

Table 2. The Earliest Time of Each Viscera L6565 MuLV RNA Detected (wk)

<table>
<thead>
<tr>
<th>L6565 MuLV RNA</th>
<th>Thymus</th>
<th>Spleen</th>
<th>Peripheral Blood</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>L6565 MuLV group</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>L6565 MuLV plus HSV-1 group</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>Control group</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

DISCUSSION

The study reported here not only confirmed and expanded upon a previous study (Zheng et al. [6]), in which inoculation of L6565 MuLV was found to result in persistent retrovirus infection in kunming suckling mice, but also indicated that HSV-1 possess the ability to activate the pathogenicity of L6565 MuLV in vivo under the conditions used. Other studies have reported that HSV-1, HSV-2 and their DNA or TK-gene-containing cloned sequence all can activate C-type murine retrovirus in vitro [14-16]. Moreover, Liu etc. of our institute have proven that HSV-2 can activate L6565 MuLV [13]. All studies demonstrate that the ability to activate the pathophysiology of L6565 MuLV is a common property shared by all Herpesviruses.

Pathogenicity

The L6565 Leukemia mice model was established by Zheng BF etc. in Department of Pathophysiology, Shanghai Medical University, in 1965 [6]. It is a utility model to study the etiopathogenesis and pathogenesis of leukemia induced by L6565 MLV. They defined L6565 MuLV as the pathogenic virus for the model, but yet to study the pathogenicity of the virus. We expanded the L6565 MuLV mice model by using L6565 lymphocyte leukemia cells and reported the pathogenicity of the virus [10].

First, our results proved again that L6565 MuLV is a kind of Gammaretroviruses of Retroviridac homologous with Moloney MuLV. We designed a couple of primers based on the conservative nucleotide sequence of the Pol gene in the Moloney MuLV genome [17]. By RT-PCR, we successfully obtained the amplification fragment of L6565 MuLV RNA in both the leukemia cell lines and the tissues of infected mice.

Second, there are two forms of L6565 MuLV that exist in leukemia cells and infected mice: L6565 MuLV RNA and integrated proviral DNA. The DNA isolated from L6565 MuLV infected mice was also detected by PCR in the same conditions as RNA amplification. Both show the same amplification fragment.

Third, similar to Moloney MuLV, L6565 MuLV RNA was first detected in the thymus and spleen, then followed by peripheral blood, liver, kidney and other viscera. By molecular hybridization and PCR technique, previous research shows that MuLV RNA was first detected in bone marrow tissue, spleen and thymus [2, 3], and bone marrow multipotent stem cell is the target cell of murine leukemic virus infection [18]. Others reported that onc-genes in leukemic mice, such as c-myc and c-fos, are over-expressed [19] and viral nucleate inserts itself into a region near the cellular proto-oncogene [20]. So we propose that L6565MuLV may be infect lymphoid stem cells in thymus and spleen first, and then insert itself into a place near the cellular proto-oncogene in the form of proviral DNA. Consequently, this activates the expression of those onc-genes and turn lymphoid stem cells into leukemic cells. Once reaching a certain number, the leukemic cells flow with blood, encroach main viscera, and Leukemia appears.

Activation by HSV-1

Our result demonstrates that the delitescence of the HSV-1 plus L6565 MuLV co-infected group (10 weeks) is earlier
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than that of the L6565 MuLV infected group (8 weeks); the infection rate increased from 41.7% to 74.3% with a statistical significant difference (P<0.01). From the results, we conclude that HSV-1 can promote the pathogenicity of L6565 MuLV to induce Murine Leukemia.

HSV-1 and HSV-2 can activate type C oncarnovirus in vitro, which was discovered by Hamper etc. [14]. Further research demonstrated that HSV-1, HSV-2 and their DNA and TK-gene-containing cloned sequence may all activate C-type murine retrovirus [10]. Other research demonstrated that HSV-1 and HSV-2 can induce host cell transformation. This effect can also be seen in certain viral DNA or inactivated virus, which damaged cytochromatin, induced rearrangement of cell genome, and changed cellular phenotype [21]. Our studies confirmed the fact that HSV activates L6565 MuLV in vivo, and we also suppose the mechanisms could be the ways below. First, it is may be the synergistic effect of HSV-1 and MuLV to promote the transformation of lymphoid stem cells into leukemic cells. The more leukemic cells proliferated, the higher the possibility of Leukemia to appear. Second, HSV-1 maybe damages the host genome, promotes the insertion of MuLV provirus, activates the expression of onc-gene, and then transforms the normal lymphocytes into leukemic cells. Third, HSV-1 maybe activates the replication of L6565 MuLV, so as to increase the possibility of infection.

REFERENCES


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