Molecular analysis of clonality in Kaposi's sarcoma

Eric Delabesse, Eric Oksenhendler, Celeste Lebbé, Olivier Vérola, Bruno Varet, Ali G Turhan

Abstract
Background—Kaposi's sarcoma is considered to be an angioproliferative disease associated with a novel herpesvirus (KSHV/HHV8), but the precise pathophysiology of the lesion remains unclear. The study of clonality in Kaposi's sarcoma using X linked DNA polymorphism has been difficult so far, because of a very strong prevalence of the disease in males.

Aims—To study the clonality of Kaposi's sarcoma lesions.

Methods—An assay based on a methyl sensitive restriction digest followed by polymerase chain reaction (PCR) amplification of the highly polymorphic human androgen receptor (HUMARA) gene was used. Tissues from Kaposi's sarcoma lesions and control tissues from the same patients were obtained from seven females, four with classic Kaposi's sarcoma and three with AIDS associated Kaposi's sarcoma. A cutaneous angiosarcoma was also analysed, for comparative purposes, and showed evidence of clonality after HpaII digestion.

Results—All patients were heterozygous for the HUMARA polymorphism and informative for analysis. In all patients, including four with a nodular form of Kaposi's sarcoma and more than 70% spindle cells in the lesion, a polyclonal pattern of inactivation could be demonstrated.

Conclusions—The Kaposi's sarcoma lesion is first of all a polyclonal cell proliferation.

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Keywords: Kaposi's sarcoma; clonality; HUMARA

Kaposi's sarcoma is an angioproliferative disease characterised by proliferation of spindle shaped cells predominantly of vascular endothelial cell origin, neoangiogenesis, inflammatory cell infiltration, and oedema.1 It is the most common neoplasm associated with AIDS but the tumour occurs also in selected human immunodeficiency virus (HIV) negative groups, including immunosuppressed transplant recipients and some African and Mediterranean populations. The demonstration of DNA sequences of a novel herpesvirus (KSHV/HHV8) in virtually all Kaposi's sarcoma specimens has stimulated the hypothesis that the sarcoma has a viral origin.2 This virus is related to two γ herpesviruses, Epstein-Barr virus and Herpesvirus saimiri, which are both transformants. No definite demonstration of a clonal origin has been provided, mainly because of the lack of a cellular or molecular marker of the lesion.

Clonality is an important characteristic of neoplastic lesions, and absence of clonality is indicative of reactive hyperplasia. A useful marker for clonality is the inactivation pattern of the X chromosome. In females, one of the two X chromosomes in each cell is inactivated by methylation and the other one remains active. This methylation occurs in an early stage of embryonic development, at random, and the same methylation pattern is passed to daughter cells in somatic replication.3 Thus, in females, normal tissues are composed of cellular mosaics, differing only in which of the two X chromosomes has been inactivated. In a female heterozygous for an X linked polymorphism, polyclonal tissues will contain approximately equal numbers of cells in which one or the other polymorphic allele is inactivated. In contrast, in a neoplasm derived from a single somatic cell, all of the tumour cells would retain the same X chromosome inactivation pattern.4

The human X linked androgen receptor gene (HUMARA) is used widely to assess for clonal patterns of X chromosome inactivation. A polymorphic short tandem repeat has been identified in the HUMARA gene. More than 80% of females are heterozygous for the number of CAG trinucleotide repeats in the region.5 Polymerase chain reaction (PCR) amplification of a target within the gene that contains flanking HpaII methylation sensitive restriction endonuclease sites has been described. After digestion of the DNA with HpaII, it is possible to distinguish between polyclonal and clonal cell populations. In a polyclonal population (for example, normal tissue) both maternal and paternal alleles are digested equally and there is an equal reduction of the signal from both alleles after PCR amplification. In contrast, tumour cells arise from a single precursor cell and have the same parental allele inactivated. After digestion and amplification, the signal from the inactive allele remains unchanged, while the signal from the active allele nearly disappears.

The study of clonality in Kaposi's sarcoma using X linked DNA polymorphism is difficult because of a very strong prevalence of the disease in males.6,7 Rabkin et al8 have shown that in advanced cases of AIDS related Kaposi's sarcoma, clonal origin could be demonstrated using X linked DNA polymorphism. More recently, the terminal repeat analysis of KSHV/
HHV8 DNA from a Kaposi's sarcoma lesion has also suggested a monoclonal expansion of the virus in the Kaposi's sarcoma tumour. Recently, we have developed a non-radioactive and quantitative clonality assay that allows this analysis to be performed in at least 70% of an unselected female population. We had the opportunity to use this methodology to study seven female patients with a diagnosis of Kaposi's sarcoma.

Materials and methods

Tissues
All seven patients had a skin biopsy (with informed consent) and all analyses were performed with institutional approval. Punch biopsies (6 mm diameter) were performed, using standard procedures, from both a lesion and normal non-adjacent skin. Each biopsy specimen was divided into two samples. One sample was either frozen immediately for further analysis or processed for DNA extraction, and the other was used for confirmation of Kaposi's sarcoma diagnosis by standard histology after staining with haematoxylin and eosin.

In six patients, both normal skin and Kaposi's lesion were analysed in parallel. In one patient (KO), normal skin tissue was not available for analysis, and peripheral blood mononuclear cells were used as a control.

As a control for clonal tissue, a tumour sample from a patient with a diagnosis of cutaneous angiosarcoma or samples of patients with haematological malignancies were used.

Histopathology
The various evolutionary stages of 10 Kaposi's sarcoma lesions obtained from seven patients were determined by haematoxylin and eosin staining. Pathological staging was performed according to Chor. In the macular or patchy phases, small blood vessels dissecting collagen, few extravasated red blood cells, rare plasma, and other inflammatory cells were seen. In the plaque stage, the degree of vascular proliferation involving the whole thickness of the dermis increased with progressive lesional evolution. In the advanced tumoural or nodular phase, the entire lesion presented as the proliferation of long spindle shaped cells associated with slit-like blood vessels.

DNA extraction and enzymatic digestion
DNA was extracted from the tumour and normal skin samples using standard procedures. Briefly, samples were digested in 500 μl Tris NaCl EDTA buffer containing 1% sodium dodecyl sulphate (SDS). Proteinase K was then added to a final concentration of 100 μg/ml and the sample was kept at 37°C overnight. The digestion was pursued for an additional 24 hours with the addition of 40 μl of proteinase K (10 mg/ml) and samples were extracted with TE buffer equilibrated phenol (twice), 1:1 phenol–chloroform (once) and a mixture of 24:1 chloroform–isoamyl alcohol (once). DNA was precipitated in the presence of 2.5 M ammonium acetate and two volumes of ethanol. The pellet was washed with 75% ethanol and dried.

To perform clonal analysis on informative patient samples, normal and Kaposi's sarcoma DNA were divided into two fractions and one of these was digested with an excess amount of HpaII (10 U) overnight, whereas the other half was left undigested. Both fractions were used for PCR amplification.

PCR amplification
This analysis was performed according to a clonality assay that we have developed recently. Briefly, 500 ng of DNA was amplified by PCR with 5’ and 3’ primers (0.5 μM each) encompassing the polymorphic CAG region of the HUMARA gene, HUMARA-C sense which is labelled by fluorescein isothiocyanate (5’ GTG CGC GAA GTG ATG CAG AAC C 3’) and HUMARA-C antisense (5’ TAC GAT GGG CTT GGG GAG AAC C 3’). After migration on an automatic ABI 373A sequence analyser using a 10% denaturing polyacrylamide gel, fluorescent peaks were quantified with the Applied Biosystems software GeneScan Analysis.

Data interpretation
Amplification of each allele generated a set of tightly clustered multiple bands, including two major product bands and several associated minor bands of lesser intensity. This result is frequently observed after PCR amplification of DNA short tandem repeats. Clonality assessment was based on the major product bands generated from each allele.

A screening gel using DNA extracted from peripheral blood mononuclear cells was performed initially to detect informative patients. Patients were considered heterozygous if PCR amplification of undigested DNA showed two major HUMARA product bands.

In informative patients, analysis of undigested DNA from clonal tissues will still produce two major product bands, as both X chromosomes are available for amplification. Predigestion of DNA with the methylation sensitive restriction enzyme HpaII precludes the PCR amplification of the non-methylated allele, permitting amplification of fragments originating from the methylated allele only. Amplification of HpaII digested DNA from polyclonal tissue will continue to show two major product bands.

The integration analysis of the area under the peaks allowed the quantification of the data. For each sample, either from the normal tissue or the Kaposi's sarcoma lesion, the relative index (RI) value is calculated by dividing the ratio of the two peaks of the HpaII digested sample by the ratio of the two peaks of the undigested sample. The relative corrected index (RCI) is the ratio of the RI obtained with the Kaposi's sarcoma sample divided by the RI obtained with the normal skin of the same patient. The final value is inverted if necessary to obtain a value always greater than one. The RCI value is closer to one when the sample is polyclonal. In the case of monoclonality, values tend to be infinite.

In previous studies performed in patients with acute leukaemia and myelodysplasia, we
assessed the sensitivity of our technique, by using mixing experiments with an increasing amount of a known clonal sample (from 5% up to 95%) in a background of a polyclonal sample, both samples being digested with an excess amount of HpaII. In two separate experiments, the value of the RCI increases with the presence of an increasing amount of clonal cells. A 10% to 20% contamination with clonal cells could be detected in a polyclonal background as the RCI value ranged between two and three, while a 40% contamination was associated with a RCI value above 10 (data not shown).

Results

CLINICAL AND PATHOLOGICAL DATA ON SEVEN KAPOSI’S SARCOMA PATIENTS

Table 1 indicates the clinical and pathological characteristics of the patient population studied. Three patients had HIV associated Kaposi’s sarcoma and four patients had classical Kaposi’s sarcoma. Kaposi’s sarcoma was present for six months or less in four patients and for more than 10 years in patient GI.

Only one case was examined in the macular stage. Two cases were examined in the plaque or intermediate phase. In three cases, including one patient for whom three lesions were studied, the lesions were nodular. In the last patient, one lesion was a plaque and another lesion was nodular (Fig 1).

Table 1  Clinical and pathological data on seven Kaposi’s sarcoma (KS) patients

<table>
<thead>
<tr>
<th>Patient</th>
<th>Histopathology of KS</th>
<th>Duration of KS months</th>
<th>HIV status</th>
<th>CD4 cell count (&lt;10^9/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BI</td>
<td>Nodule</td>
<td>6</td>
<td>Negative</td>
<td>800</td>
</tr>
<tr>
<td>GH</td>
<td>Nodule</td>
<td>6</td>
<td>Negative</td>
<td>972</td>
</tr>
<tr>
<td>GI-1</td>
<td>Nodule</td>
<td>170</td>
<td>Negative</td>
<td>1034</td>
</tr>
<tr>
<td>GI-2</td>
<td>Nodule</td>
<td>2</td>
<td>Positive</td>
<td>152</td>
</tr>
<tr>
<td>GI-3</td>
<td>Nodule</td>
<td>2</td>
<td>Positive</td>
<td>30</td>
</tr>
<tr>
<td>KO</td>
<td>Plaque</td>
<td>36</td>
<td>Negative</td>
<td>300</td>
</tr>
<tr>
<td>GU</td>
<td>Macule</td>
<td>2</td>
<td>Negative</td>
<td>152</td>
</tr>
<tr>
<td>NZ</td>
<td>Plaque</td>
<td>24</td>
<td>Positive</td>
<td>30</td>
</tr>
<tr>
<td>MA-1</td>
<td>Plaque</td>
<td>4</td>
<td>Positive</td>
<td>11</td>
</tr>
<tr>
<td>MA-2</td>
<td>Nodule</td>
<td>6</td>
<td>Negative</td>
<td>300</td>
</tr>
</tbody>
</table>

CLONALITY ANALYSIS

All seven patients were found to be informative for the HUMARA gene polymorphism and analysable. In six patients, including one patient (MA) with two separate biopsy specimens, a typical polyclonal pattern was demonstrated in the Kaposi’s sarcoma tissue with a RCI ranging from 1.1 to 1.7 (Fig 2, table 2). In one patient (GI), we had the opportunity to study three separate tumour tissue samples. In all these three biopsy specimens we demonstrated close RCI values ranging from 1.5 to 2.3. However, in this patient the RI in normal skin was 2.3, indicating a skewing of X inactivation, and the interpretation of these results in terms of clonality is difficult.

As a monoclonal control, and to validate the ability to detect clonal populations from skin tumours with our assay, we have used DNA extracted from a cutaneous angiosarcoma sample. This analysis showed a typical clonal pattern, with an index value of 19.3 (Fig 3).

Discussion

Whether Kaposi’s sarcoma is a neoplastic lesion or a reactive process remains controversial.7 Most data support the hypothesis that the initial lesion is a reactive hyperplasia: the normal chromosome complement of the spindle cells,12 the occasional tumour regression either spontaneously or after reducing the immunosuppression in transplant recipients,13 and the dependence on cytokines such as basic fibroblast growth factor, oncstatin M, and interleukin 6 (IL-6) for the growth of Kaposi’s sarcoma derived cells in vitro.14 However, in vitro derived Kaposi’s sarcoma cells do not have the characteristics of transformed cells. However, at least two Kaposi’s sarcoma derived cell lines, KS Y-15 and SLK16 can induce tumours in nude mice.

Clonality is an important characteristic of neoplastic lesions, and absence of clonality is indicative of reactive hyperplasia. In the absence of a specific clonal chromosomal abnormality, methylation analysis of X linked genes can give important information with

Figure 1  Representative case of Kaposi’s sarcoma in an AIDS patient (MA). MA-1, a plaque lesion (A) exhibiting the association of spindle shaped cells and extravasated red blood cells, plasma and other inflammatory cells. MA-2, a nodular lesion (B) presenting as a dense proliferation of long spindle shaped cells associated with slit-like blood vessels. (Haematoxylin and eosin, original magnification, x200.)

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Clonal analysis of Kaposi’s sarcoma

Figure 2. Evaluation of clonality in Kaposi’s sarcoma using human androgen receptor (HUMARA) polymorphism analysis in seven female patients. After digestion by HpaII, DNA was amplified from normal skin and from a Kaposi’s sarcoma lesion. In most cases the pattern observed after digestion with HpaII was very similar in the Kaposi’s sarcoma lesion(s) and the normal skin of the same patient. For patient KO, in the absence of a normal skin specimen, peripheral blood mononuclear cells were used as a control.

Relevant to the presence of a clonal cell population in a given sample, provided that an embryologically identical normal tissue has been analysed in parallel as a control. Using this assay, we and others have been able to demonstrate the monoclonal nature of human non-lymphoid leukaemic cells, whereas normal

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Table 2: Evaluation of clonality in Kaposi’s sarcoma using human androgen receptor (HUMARA) polymorphism analysis

<table>
<thead>
<tr>
<th>Patient</th>
<th>RI&lt;sub&gt;n&lt;/sub&gt;</th>
<th>RI&lt;sub&gt;m&lt;/sub&gt;</th>
<th>RCI</th>
</tr>
</thead>
<tbody>
<tr>
<td>BI</td>
<td>1.2</td>
<td>1.5</td>
<td>1.2</td>
</tr>
<tr>
<td>GH</td>
<td>0.7</td>
<td>1.3</td>
<td>1.7</td>
</tr>
<tr>
<td>GI-1</td>
<td>2.3</td>
<td>3.5</td>
<td>1.5</td>
</tr>
<tr>
<td>GI-2</td>
<td>2.3</td>
<td>4.3</td>
<td>1.9</td>
</tr>
<tr>
<td>GI-3</td>
<td>2.3</td>
<td>5.4</td>
<td>2.3</td>
</tr>
<tr>
<td>KO</td>
<td>0.9</td>
<td>1.0</td>
<td>1.2</td>
</tr>
<tr>
<td>GI</td>
<td>0.9</td>
<td>0.9</td>
<td>1.2</td>
</tr>
<tr>
<td>NZ</td>
<td>0.3</td>
<td>0.5</td>
<td>1.7</td>
</tr>
<tr>
<td>MA-I</td>
<td>1.3</td>
<td>1.0</td>
<td>1.4</td>
</tr>
<tr>
<td>MA-2</td>
<td>1.3</td>
<td>1.4</td>
<td>1.1</td>
</tr>
</tbody>
</table>

RI<sub>n</sub>, relative index; RI<sub>m</sub>, relative index of normal skin; RCI, relative corrected index (ratio of RI<sub>n</sub>:RI<sub>m</sub>). When RCI is close to 1, the sample is polyclonal. In the case of monoclonality, values tend to be infinite. For patient KO, in the absence of a normal skin specimen, peripheral blood mononuclear cells were used as a control.

Figure 3. Evaluation of clonality in a cutaneous angiosarcoma using HUMARA polymorphism analysis. DNA from the lesion was amplified directly (left) and after digestion by HpaII (right). After digestion and amplification, the signal from the inactive allele remains unchanged, while the signal from the active allele nearly disappears.

cells from the same patients exhibited a normal polyclonal pattern.

In Kaposi’s sarcoma, the only available study included three patients with AIDS related nodular Kaposi’s sarcoma who showed a clonal pattern of the sarcoma lesion using HUMARA polymorphism analysis. Eight women had undergone cutaneous biopsy but only three specimens were selected for analysis on the basis of less than 20% contaminating normal cells after examination of the section by light microscopy. This selection may have favoured sarcomatous lesions observed in patients with longer standing Kaposi’s sarcoma lesions. Our patient population was different from that included in this series. First, in four of the seven patients, Kaposi’s sarcoma was not related to HIV infection, and in four cases, the lesions had been present for less than six months. Second, four of 10 lesions studied were macules or plaques, and in these lesions the proportion of spindle cells did not exceed 20%. However, the six other lesions were nodular with a proportion of spindle cells exceeding 70% in four of them.

The present study presents indirect molecular evidence that most cells involved in the Kaposi’s sarcoma lesion are polyclonal. The most likely explanation for a false polyclonal pattern in a clonal tissue is the contamination of the biopsy specimen with normal cells. We attempted to minimise this bias by evaluating the percentage of spindle cells by careful classical morphological examination of adjacent sections. The presence of more than 70% spindle cells in four out of 10 specimens suggests that a clonal proliferation of these cells could have been detected. In these four lesions (BI, GH, GI-2, MA-2) the RCI was below two, indicative of a polyclonal cell population. In our experience, and as demonstrated by Enamoto et al., clonality can be assessed with the HUMARA technique when monoclonal cells constitute >20% of the total cell population. Furthermore, using the same technique, the analysis of a cutaneous angiosarcoma demonstrated unequivocal clonality of the lesion.

It has been demonstrated by PCR in situ hybridisation that KSHV/HHV8 is localised to the endothelial cells that line the irregular vascular slits and to some of the perivascular spindle-shaped cells seen in Kaposi’s sarcoma lesions. These results suggest that the first critical event of Kaposi’s sarcoma pathogenesis could be the infection of the target cell with KSHV/HHV8. The finding by Russo et al. that KSHV/HHV8 is clonal in a Kaposi’s sarcoma specimen may represent the clonal evolution of an advanced lesion or the presence in the lesion of both clonal infected cells and proliferating uninfected cells.

The viral infection could induce proliferation of the spindle cells through autocrine and/or paracrine production of cellular cytokines or through the expression of viral genes, homologues of cellular genes involved in cell proliferation (vIL-6, G-protein coupled receptor, or transformation (vBcl-2, vCyclin), and of viral homologues of chemo-
kines (macrophage inflammatory protein: vMIP-I, vMIP-II) leading to the formation of tumoural lesions. In a later stage or after a new oncogenic event, clonal selection may contribute to the development of more aggressive tumoural nodules. With such a scenario, very similar to that observed in Epstein-Barr virus associated lymphoproliferative disorders, a polyclonal proliferation of the spindle cells could be followed by clonal restriction.

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