VASCULAR REMODELING IN KAPOSI'S SARCOMA AND AVIAN HEMANGIOMATOSIS: RELATION TO THE VERTEBRATE LYMPHATIC SYSTEM

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Kaposi’s sarcoma (KS) is often defined as a low grade endothelial malignancy with metastatic potential, although Kaposi himself found this last point difficult to reconcile with the occurrence of bilateral lesions on the distal extremities (1). A number of observations in recent decades force a re-evaluation of the Kaposi lesion in terms of constituent cells and its relation to the vascular systems of both blood and lymph. Physicians have assumed that KS lacks an animal counterpart, but new insights suggest the relevance of natural animal models, specifically hemangiomatosis of fowl (2). In addition, certain characteristics of the present epidemic of KS in association with the acquired immunodeficiency syndrome (AIDS), such as the frequency of KS differing by risk group, suggest an independent infectious etiology in Kaposi’s sarcoma.

Classical descriptions of the pathology of Kaposi’s sarcoma regularly include dilated vascular spaces lined by elongated endothelial cells among which is a greater or lesser complement of spindle cells with vascular slits. These elements have been considered to form an amalgam of proliferating tissue possibly harboring other neoplastic mesenchymal elements in the guise of spindle cells.

Recent studies using immunohistochemistry and cell culture have reaffirmed the endothelial cell as the major proliferative constituent in KS (3-5). Prior to the AIDS epidemic, little interest was accorded nascent lesions, which were often said to resemble granulation tissue. Large numbers of biopsies performed on macular lesions in recent years often reveal on histopathology aberrant dilated lymphatic-like channels. These can resemble granulation tissue when sclerosis supervenes, although a component of angiomatoid capillary proliferation may be present. In a histologic review of more than 110 cases of KS of the elderly, the author has seen a number of such nascent lesions, although not as frequently as in AIDS patients. The conclusion is that both clinical forms of KS have similar pathologic beginnings, and patients in the eighth decade of life (the mean age of Swedish patients with the disease) may be less likely to seek medical attention for initially unobtrusive macular lesions of the feet. Also, the rate of histologic stage progression varies among patients. Some may develop a spindle cell nodule in a matter of a few weeks, whereas others may retain the aberrant lymphatic (macular) stage for months. Recognition of the dynamic patterns of development of a single lesion replaces earlier concepts of separate "forms" of KS such as lymphangiomatous (6), angiomatous and sarcomatous (7). A four-stage scheme based on histogenesis is given in reference 8. Stage 1 involves aberrant endothelial growth and coupling between the venous circulation and lymphatic system; stage 2, the spindle cell nodule; stage 3, increased nuclear atypia in spindle cells; and stage...
4, angiosarcoma. The last stage has evolved in only 4% of our patients in Lund.

Immunohistochemistry has been used to define more precisely the endothelial cell of origin but with conflicting results. Monoclonal antibodies purported to be specific for endothelium exclusive of lymphendothelium did not react with cells in KS lesions (5), which was taken as evidence of lymphatic origin. Other monoclonal antibodies specific for blood vessel endothelium only have produced positive staining (4). The apparent contradiction stems from viewing KS as a malignant tumor arising in one or the other of two biologically separate types of endothelium, a premise with other drawbacks.

First, malignant change in lymphatic endothelium does not explain the fact that the lesions carry much blood, often out of proportion to the recognizable lymphatic component. Small areas of new capillary growth are not necessarily congruent with areas of proliferating lymphatics. Congruency would be expected if the blood vessels merely nourished "parenchymal" lymphatic endothelium, analogous to the situation in solid tumors of nonvascular type. Second, the idea of origin in blood vessels ignores the clinical and angiographic lymphaticovenous nature of KS. Finally, veins and lymphatics are biologically intimately related. Lymphatics in man possibly arise from embryonic veins, and the genome may be assumed to ultimately control the dichotomous differentiation of endothelium, at least in the embryo.

These problems are resolved by taking into account the extensive remodeling of the blood capillary, venular and lymphatic beds during lesional development, which at the outset leads to connections between both vascular systems (Fig. 1) (8). Remodeling begins in any given site at the capillovenular level and tends to continue in the direction of blood flow in veins. This results in lesions developing along lymphatic and venous pathways such as portal zones and fibrous septa of the liver, as seen in Fig. 2.

Skin is rich in small blood vessels and lymphatics, and the broad expanse of dermis allows the spectrum of pathologic changes in KS to be observed more readily than in the crowded confines of hepatic portal zones, for example. Nevertheless, the pathogenetic events may be initiated in other sites, such as the muscularis mucosae of the gastrointestinal canal, septa and portal tracts of the liver, and marginal sinus area of lymph nodes. The first recognizable stage of cutaneous KS shows histologically the dissection of collagen bundles by jagged, aberrant lymphatic clefts. Their lymphatic nature is recognized when in the normal anatomical position around the superficial dermal venular plexus, forming so-called glomeruloid structures (Fig. 3). In addition, ultrastructural, histochemical, and immunohistochemical analysis confirm that the cells lining these clefts show features such as a partly absent basal lamina, lack of surrounding pericytes (9) and the presence of 5' nucleotidase (1), which are more characteristic of lymph-endothelium. The earliest clefts are bloodless in sections, but hemosiderin is frequently found in the lymphatic-like endothelium and adjacent fibroblasts lining collagen bundles. In a slightly later phase, glomeruloid structures diminish in size and number to be replaced by anastomosing blood-filled spaces lined by
Fig. 2. Cut surface of liver shows multicentric KS with concentration about a major septum. The other foot may arise in portal tracts.

Fig. 3. Glomeruloid structure containing central blood capillary and venule within aberrant lymphatic cleft.

cells more closely resembling blood vascular endothelium. The transition between these two states is also marked by disruption of the cleft lining with perifocal hemorrhage and sclerosis. Delineation of these stages is facilitated by examples in which the changes occur in "pure" form, but many specimens show
transitions. In general, spindle cells arise after blood flow is established between the venous side of the circulation and aberrant lymphatic channels. Spindle cells stain variably for Factor VIII-related antigen and fucosyl moieties (by Ulex europaeus I lectin binding) and lack Weibel-Palade bodies. These cells may represent transformed endothelial clones, and Dorfman in this symposium has suggested suitable methods for studying this question.

As the above sequence of events is repeated along progressively larger veins, radial venolymphatics become the hallmarks of lymphaticovenous union. Jagged channels histologically similar to nascent lesions in biopsies of skin penetrate the walls of veins (Fig. 4) affording potential linkage between both vascular systems beyond the capillaryvenular level. This is observed in the hilum of lymph nodes, gastrointestinal submucosa, subcutaneous and perivisceral fat and the fibrous septa of the liver, for example. Hemorrhage tends to be more prominent about these structures, conceivably due to greater pressures acting on the incompetent venolymphatics. That venular glomeruloid structures and radial venolymphatics account for angiographically demonstrated venolymphatic anastomoses is by inference. It is strongly suggested by careful histological, immunohistochemical, and electronmicroscopical analysis of lesional development. Independent proof might require simultaneous histographic and lymphographic/angiographic study of single lesions. Newly devised infiltrative lymphographic techniques could be applied in skin lesions (11).

Our comparison of AIDS-related and non-AIDS KS suggests that the progression upwards along the venous system may be related to the presence of an immune defect. Sections from ten autopsies performed on patients with KS but without AIDS over a 25-year period in Sweden revealed only a solitary focus of venolymphatics in one patient, and that patient had received ACTH. This compares with their presence in all of nine AIDS patients with KS at autopsy (8). The question needs more study, in particular with additional comparison of autopsy material in endemic African KS, since such cases have been reported to show structures corresponding to radial venolymphatics, though presumably outside the context of a clinical immunode-
Leghorn fowl. The clinical features of each are given in *Table 1*. Both entities are related to lymphoma. In the case of the avian tumors, this is due to the lymphoid leukosis group of retroviruses to which belong the agents causing hemangiomatosis and lymphomatosis (13). Hemangiomatosis is apparently most commonly associated with visceral lymphomatosis (14,15) and shows vertical transmission. Avian leukosis viral DNA becomes integrated in the genome of endothelial tumor cells, but oncogenes possibly involved in tumorigenesis have not been definitely identified. However, recently Robinson et al. (16) described rapid induction of angiosarcomas in chickens using a leukosis virus which transduced a portion of the gene (c-erbB) responsible for epidermal growth factor receptor. It remains to be seen whether this oncogene is also involved in naturally occurring hemangiomatosis.

Spontaneous regression is often noted in both diseases. For fowl, this may be a result of natural immunity (13). KS appears to be modulated by immune function and in many patients with acquired immunodeficiency lesions regress as immune function improves.

Avian hemangiomatosis has been the focus of few pathologic studies. Yet the salient pathologic features presented in *Table 2* also correspond in a striking manner to Kaposi’s sarcoma. Most importantly, both lesions initially resemble an innocuous dissecting proliferation of endothelium and evolve into sarcoma-like tumors or even frank angiosarcoma, as shown in *Fig. 6*.

It is not known whether the avian tumors involve lymphatics. However, intravenous extension has been illustrated (17), and this is typical of endemic African KS, in particular. Studies should be undertaken to map the progression of these vascular avian tumors from their inception, such as has been done in KS. This would include the use of immunohistochemistry and electron microscopy to distinguish between lymphatic and blood vascular endothelium. Lymphaticovenous connections might also be sought by
### Table 1
Comparison of Clinical Features in Kaposi's Sarcoma and Avian Hemangiomatosis

<table>
<thead>
<tr>
<th>Etiology</th>
<th>Kaposi's Sarcoma</th>
<th>Avian Hemangiomatosis</th>
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<tbody>
<tr>
<td>Probable virus</td>
<td>Retrovirus in lymphoid leukemia group</td>
<td></td>
</tr>
<tr>
<td>Epidemiology</td>
<td>Sporadic &amp; epidemic; increased incidence in some immunodeficiency states, increased lymphoma</td>
<td></td>
</tr>
<tr>
<td>Sites</td>
<td>Distal skin initially; multicentric, internal organs</td>
<td></td>
</tr>
<tr>
<td>Regression</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Metastasis</td>
<td>Rare</td>
<td>Not documented</td>
</tr>
<tr>
<td>Malignant Progress</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

### Table 2
Comparison of Salient Histologic Features in Kaposi's Sarcoma and Avian Hemangiomatosis

<table>
<thead>
<tr>
<th>Stage</th>
<th>Kaposi's Sarcoma</th>
<th>Avian Hemangiomatosis</th>
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<tbody>
<tr>
<td>1st stage</td>
<td>Proliferating lymphatic &amp; blood vessel endothelium</td>
<td>Proliferating endothelium</td>
</tr>
<tr>
<td>2nd stage</td>
<td>Spindle cell growth with slits</td>
<td>Spindle cell growth with slits</td>
</tr>
<tr>
<td>3rd stage</td>
<td>Fibrosarcoma-like</td>
<td>Fibrosarcoma-like</td>
</tr>
<tr>
<td>Dilated spaces lined by benign endothelium</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Spindle cell growth in veins</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
Fig. 6 (at left). (A) Closely spaced atypical endothelial cells and nucleated red cells in subcutaneous lesion in chicken. [Courtesy of Dr. B. Jarplid, reprinted from J. Comp. Pathol.; ref. 15.] (B) Stage 3 KS lesion with sarcomatoid atypia and loss of vascular slits.

applying angiographic techniques. With continued advances in molecular biology, the involvement of both vascular systems, if found, could eventually shed light on the genetic origins of the vertebrate lymphatic system.

ADDENDUM

Recent transfection studies of KS have tentatively implicated rearrangements of DNA flanking an unusual oncogene which codes for a protein homologous to angiogenic factors (18,19).

ACKNOWLEDGEMENTS

This work is supported by the John och Augusta Perssons stiftelse for vetenskaplig medicinsk forskning, the Royal Physiographical Society of Lund, and the Medical Faculty of the University of Lund.

REFERENCES


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