Tubuloreticular Structures in Kaposi’s Sarcoma: A Comparison of the Classical and AIDS-Associated Forms

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The ultrastructural morphology of both the classical and the AIDS-associated forms of Kaposi's sarcoma was examined electron-microscopically. Tubuloreticular structures were found only in the AIDS-associated form of Kaposi's sarcoma, and not in the classical variant of the tumor. Moreover, the tubuloreticular structures, 20-30 nm in diameter, were present in very high numbers and in two different forms: the loosely intertwined tubuli, which were clearly predominant; and those forming a more compact pattern. These findings suggest that the presence of tubuloreticular structures may well be an ultrastructural marker for diagnosing AIDS and AIDS-associated disorders. Key words: Acquired immune deficiency syndrome. (Received January 30, 1986.)

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The classical form of Kaposi's sarcoma (KS), a rare malignant tumor with an unusual clinical course and exhibiting interesting epidemiological features, was first described by Kaposi in 1872 (1) and is still not completely understood. The new epidemic form of KS, found in patients with the acquired immune deficiency syndrome (AIDS), has stimulated new interest in this rare vascular neoplasm. It is, unlike the classical form, biologically more aggressive and affects young adults. Further, the disease tends to be more generalized with visceral and lymph node involvement.

Tubuloreticular structures (TRS) are irregularly branching and anastomosing tubules within the endoplasmic reticulum, and have been found in a variety of autoimmune, viral and neoplastic diseases (2, 3).

As we have consistently found TRS in the AIDS-associated form of KS, it seemed essential to re-examine the classical form of KS with regard to the presence of TRS. Thus, in the present paper, we report on the occurrence of TRS in KS cells in three patients with AIDS as compared with the classical form of KS.

MATERIAL AND METHODS
Skin biopsies were taken from classical KS lesions of two patients, and from early plaque lesions and tumor lesions of KS of three patients with AIDS. In the AIDS patients the diagnosis was established by HTLV-III serology, the demonstration of an inverted T-helper/T-suppressor cell ratio, the presence of opportunistic infections and of KS. The excised tissues were immediately processed for light and electron microscopy according to standard methods.

RESULTS
Light microscopically, the tumor lesions in both the classical and the AIDS-associated form of KS revealed typical spindle-shaped KS cells and numerous extravasated erythro-
Fig. 1a, b. Tumor lesion of Kaposi's sarcoma in AIDS. (a) Neoplastic spindle cell containing the loosely intertwined variety of tubuloreticular structures (TRS). N, nucleus; M, mitochondria; ER, endoplasmic reticulum; P, polysomes. (×23 850.) (b) High power magnification of the tubuloreticular structures shown in Fig. 1a. Note the tubular lumen, and apparent contiguity of the tubuloreticular structures and the granular endoplasmic reticulum. (×75 600.)
cytes. Further, there were irregular vascular slits and channels surrounded by infiltrates consisting of lymphocytes and plasma cells.

TRS were found only in AIDS-associated KS and not in the classical form of the tumor. In AIDS-associated KS the TRS were present in almost every vascular endothelial cell and neoplastic spindle cell, but only rarely in lymphocytes and macrophages.

Ultrastructurally, the TRS were numerous and were seen in two distinct forms. One consisted of loosely intertwining branching and anastomosing tubuli (Figs. 1a, b). The other was a more compact reticular network of tubular profiles of the same individual size as the loosely intertwined variety (Figs. 2, 3). The loosely interwined tubuli were far more numerous than those with a more compact form. In very rare instances, both TRS patterns could be observed within one and the same cell (Fig. 4). There were no differences either in the number or morphology of TRS in early plaque lesions or in tumor lesions of AIDS-associated KS.

The individual tubuli measured up to 20-30 nm in diameter, and the TRS as a whole extended over an area of 1.5-2.0 μm. They occurred most frequently within the endoplasmic reticulum (Fig. 1b), or very occasionally, in the perinuclear cisternae of the cells (Fig. 3).
DISCUSSION

This study confirms not only the presence of TRS (4) but also that they occur in very high numbers in KS cells obtained from both early plaque lesions as well as tumor lesions in patients with AIDS. It also shows that in patients with the classical form of KS TRS do not occur in lesional skin. This statement is based on the fact that a very large number of electron micrographs were closely scrutinized and corresponds to earlier reports on the ultrastructure of classical KS (5, 6, 7) which fail to mention the presence of TRS within KS tumor cells.

TRS are not specific to AIDS, although they have been described in different tissue biopsies and peripheral blood samples from patients with AIDS and AIDS-related diseases (8, 9, 10, 11); they may be found in various autoimmune (12), viral (13, 14) and neoplastic (15, 16) disorders. However, TRS are ultrastructurally distinct from any known virus particles. As demonstrated by digestion experiments, TRS are resistant to digestion by trypsin and by bovine pancreatic ribonuclease (RNase) and deoxyribonuclease (DNase) and are thus cytochemically believed to be complexes of phospholipid membranes and glycoprotein components (17, 18).

Fig. 3. Tumor stage of Kaposi's sarcoma in AIDS. Compact tubuloreticular structures (TRS) in perinuclear cisterna (PNC). N, nucleus; P, polymesomes. (x75,600.)

Fig. 4. Tumor stage of Kaposi's sarcoma in AIDS. Both loose (arrow) and compact (double arrow) tubuloreticular structures within endoplasmic reticulum of one and the same Kaposi's sarcoma cell. M, mitochondrion; ER, endoplasmic reticulum. (x75,600.)
The formation of TRS can be induced in lymphoid cells by halogenated pyrimidines, which are known to activate latent virus infections (19, 20, 21). Also leucocyte (α) and fibroblast (β) interferons, but not immune (γ) interferon, can stimulate the development of TRS (22). There seems to be no direct correspondence between the presence of TRS and serum interferon levels in patients with AIDS (23). Further investigation on the role of interferon in TRS formation is thus essential.

The chemical nature and the pathophysiological function of TRS in AIDS is not yet completely understood. It cannot be said with certainty if TRS are merely host cell products, defective viruses or a host cell reaction pattern in response to different stimuli.

There can be little doubt that the presence of innumerable TRS together with the serological demonstration of HTLV-III antibodies and other immunological parameters is of diagnostic relevance. This is especially so in diagnosing the early stages of AIDS and AIDS-related disorders.

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REFERENCES