Fifty cases of human immunodeficiency virus (HIV) infection: immunoultrastructural study of circulating lymphocytes


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SUMMARY The peripheral lymphocytes of 50 cases of human immunodeficiency virus (HIV) infection (13 of acquired immune deficiency syndrome (AIDS), 17 of AIDS related complex (ARC), and 20 healthy carriers) were studied immunoultrastructurally. The prevalence of “tubuloreticular structures” and “tubular confronting cisternae” increased with the progression of the disease. Numerous tubular confronting cisternae were noted in patients presenting with a high serum acid labile α-interferon values. The patients with depressed natural killer cell activity were characterised by circulating immature natural killer cells with abundant multivesicular bodies that were devoid of “parallel tubular arrays”. With an immunogold staining technique the location of HIV antigen was detected ultrastructurally, both at the surface of “hand-mirror” natural killer cell lymphocytes and inside vacuolised cells, probably corresponding to infected T4 lymphocytes. These findings indicate the usefulness of electron microscopic techniques in evaluating the pathology and the pathogenetic outcome of AIDS.

The existence of ultrastructural markers was first reported in patients with “full-blown” acquired immune deficiency syndrome (AIDS).1 We have described similar findings in cases of AIDS from Central Africa.2 Only a few well documented ultrastructural studies on peripheral lymphocytes in AIDS and AIDS related complex (ARC) have been carried out.3–5 We previously showed the presence of retroviral antigens by immunogold staining in circulating lymphocytes in prodromal AIDS.6 This study aimed to describe and complement our description by investigating the clinical importance of the ultrastructural abnormalities observed in peripheral blood lymphocytes through different clinical stages of human immunodeficiency virus (HIV) infection. Colloidal gold antibody probes were applied for the identification of the membrane phenotype of the abnormal cells and for detecting the ultrastructural location of HIV antigen.

Material and methods

Diagnosis of HIV infection was based on positive HIV serology (ELAVIA, with additional uninfected H9 controls), a low T4:T8 ratio or a positive HIV core antigen detected by a solid phase radioimmunoassay7 on cultured peripheral lymphocytes, or a combination. All cases with positive serology (ELAVIA) but with a T4:T8 ratio of > 1 were confirmed to be positive for IgG antibodies by Western blot analysis or radioimmunoprecipitation. The classification of (“full-blown”) AIDS, ARC, and healthy carriers was made according to the United States criteria of the Center for Disease Control, Atlanta.8,9 AIDS was detected in 13, ARC in 17 patients, and 20 were healthy controls. There were 22 women and 28 men. High risk groups included 11 homosexuals, 25 central Africans, seven heterosexual patients who had had sex with an infected partner, one European missionary living in Zaire, three drug addicts, two Haitians, and one European who had a blood transfusion in Haiti four years ago. Of those with AIDS, one had Kaposi’s

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sarcoma alone and one Kaposi's sarcoma associated with opportunistic infection. The average T4:T8 ratios were 0.24 for the AIDS, 0.34 for ARC, and 1.13 for healthy controls.

Polyclonal hypergammaglobulinaemia was detected in 83% of patients with AIDS, 94% of those with ARC, and in 35% of healthy controls. Serum β-2-microglobulin (Pharmacia radioimmunoassay) was consistently raised above 3.5 mg/l in 75% of AIDS patients, 38% of those with ARC, and in 22% of healthy controls. The simultaneous presence of HIV antibodies and HIV antigens was shown in all the AIDS patients, 75% of those with ARC, and in 35% of healthy controls. The presence of antibodies against HIV without antigen detection was not found in any patients with AIDS, but was present in 25% of patients with ARC, and in 25% of healthy controls. The detection of antigen alone, noted exclusively in 40% of healthy controls, was striking. The determination of serum acid labile α-interferon concentration was performed as previously described by Huygen et al.10 The percentage of circulating large granular lymphocytes and their natural killer function were evaluated by cytofluorimetric analysis (EPICS) with Leu7 and NKH1 monoclonal antibodies, and the release of 31Cr against K-562 cells, respectively. The ability of total blood cells to produce α and γ-interferon was assayed according to the method described by Kirchner.11 Interleukin-2 production isolated by purified lymphocytes was also Ficoll-Hypaque evaluated.

ULTRASTRUCTURAL STUDY
Peripheral mononuclear cells isolated by Ficoll-Hypaque gradient (Lymphoprep) were rapidly fixed for 30 minutes at room temperature by a solution of 4% glutaraldehyde, buffered by 0·1 M phosphate, at pH 7.4.12 The cells washed in the same buffer were fixed with 2% phosphate buffered osmium tetroxide and were dehydrated by graded concentrations of ethanol. The material was embedded in Epon13 and sectioned by a Diatome diamond knife with an LKB Ultratome V microtome. Ultrathin sections stained with uranyl acetate and lead citrate were observed with a Siemens Elmiskop electron microscope. After embedding, the circulating lymphocytes, which had been fixed by 2% glutaraldehyde buffered by 0·1 M sodium cacodylate (pH 7.6) for a period of two hours at room temperature, were stained with immunogold. Postfixation was omitted. Ultrathin sections on nickel grids were then incubated as previously described.4 This post-embedding method was performed for the detection of intracytoplasmic HIV antigen using a rabbit polyclonal antiserum against the p13 core protein of lymphadenopathic virus (LAV) (gift from the Pasteur Institute, Paris). The specificity of this antiserum was established by control experiments performed on HIV infected and non-infected H9 cells (gift from the laboratory of Dr RC Gallo, United States National Cancer Institute, Bethesda). Positive immunogold staining was observed only in HIV infected H9 cells.

The pre-embedding immunogold staining method6 was used for the identification of HIV antigen on the cellular membrane with the same anti-LAV antiserum and for the determination of the membrane phenotype of the cell with a set of the following monoclonal antibodies: OKT4, OKT8 and anti-HLA DR (Ortho), NKH1 (Coulter), and Leu7 (Becton-Dickinson).

Results
A network of branched tubules of 20 to 25 nm called "tubuloreticular structures" (fig 1) was located in the rough and smooth endoplasmic reticulum, in the perinuclear cisterna (rarely), or in the Golgi apparatus. Tubuloreticular structures were seen in all the subpopulations of the circulating mononuclear
cells including monocytes. They were predominant in the OKT8 positive and the NKH1 positive phenotypes (fig 2). Tubuloreticular structures were observed in 13 of 13 patients with AIDS, 11 of 17 patients with ARC (64%), and in only two of 20 healthy controls (10%). The average T4:T8 ratio was 0.25 among the patients with tubuloreticular structures and 1.13 in those without tubuloreticular structures.

Serum acid labile \( \alpha \)-interferon concentration was above 10 U/ml (normal value <6) in 11 of 15 (73%) patients with tubuloreticular structures and in two of 10 (20%) of the patients without: one of these last two patients had recently seroconverted at the time of writing.

"Tubular confronting cisternae"\(^4\) appeared in longitudinal view as cylinders ranging between 1.5 to 2.5 \( \mu \)m in length and about 300 nm in width. In cross sectional aspect they comprised two concentric cisternae with an electron dense layer of about 25 nm between them (fig 3). Tubuloreticular structure were sometimes visible inside tubular confronting cisternae (fig 4). Tubular confronting cisternae were also detected in all the subpopulations of the circulating mononuclear cells. We observed tubular confronting cisternae in six of 13 patients with AIDS (46%), five of 17 patients with ARC (29%), and in one of 20 healthy controls (5%): this last was the only healthy control to have progressed to AIDS during the year of follow up. Tubular confronting cisternae were never shown in the absence of tubuloreticular structures. The average T4:T8 ratio of the patients with tubular confronting cisternae was 0.21. Serum acid labile \( \alpha \)-interferon concentration was raised in four of five patients positive for tubular confronting cisternae. Three of these four had the highest values of \( \alpha \)-interferon observed in our study—28, 56, and 70 U/ml, respectively.

Circulating plasma cells were observed with HLA DR positive phenotype and often showed abnormal mitochondria (fig 5). These cytological characteristics were shown in 57% of our patients and were associated with a polyclonal hypergammaglobulinaemia in 91% of these cases.

In two patients "hand-mirror" cells with an eccentric nucleus and a cytoplasmic uropod bristling with microspikes were noticed (fig 6). After immunogold staining the membrane phenotype of these hand mirror cell disclosed their likely natural killer nature (fig 2). Moreover, a characteristic immunogold staining of the tip of their uropod (fig 7) was observed after

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**Fig 2** "Tubuloreticular structure" (\( T \)) of a natural killer cell of amoeboid configuration. Cellular membrane is stained (\( \rightarrow \)) with immunogold with NKH1 monoclonal antiserum. \( N \) = nucleus.
A striking finding was that in two thirds of the 50 cases about 1% of large multinucleated circulating cells were observed. Their cytoplasm showed cytoplasmic vacuoles, ranging from 0.8 to 1.5 μm in average diameter, and containing a granular material with occasional double outlined vesicles (fig 10). These cells did not display the ultrastructural appearance of monocytes, usually recognisable by their typical granules with a clear peripheral halo. Moreover, these vacuolated cells were neither found in the purified population of non-rosetting cells (B lymphocytes and monocytes) nor in the purified preparation of OKT8 positive cells that we have previously described. An immunospecific reactivity with a polyclonal antiserum directed against the principal core protein p24 of the bovine leukaemia virus was also localised in these vacuoles. Similar positive staining was obtained using the polyclonal antiserum directed against LAV in four of four cases of AIDS, one of two cases of ARC, and one of eight healthy controls. Vacuoles containing retroviral core proteins were also observed in fetal lamb kidney cells, in MT2 cells, and in H9 cells infected by BLV, HTLV I, and HTLV III, respectively. To identify the membrane phenotype of these vacuolated circulating cells pre-embedding immunogold staining with anti-HLA DR, OKT4, OKT8, NKH1, and Leu7 monoclonal antibodies was performed. Surprisingly, all of these monoclonal antibodies were unreactive. After a similar experiment performed on cultured peripheral lymphocytes, infected by HIV, we observed a strong membrane reactivity with the OKT8 antiserum in lymphocytes incubation with the antiserum directed against LAV. Control experiments with a normal rabbit serum and a monoclonal antibody against baboon endogenous virus (BaEV) gave negative results.

The cytofluorimetric analysis (EPICS) of the purified Ficoll-Hypaque lymphocytes disclosed a normal number of positive cells for the Leu7 and NKH1 phenotypes in all nine cases tested (table). Whatever the clinical stage and concentrations of α interferon, γ-interferon, or interleukin-2 productions, four of these nine patients had depressed natural killer cell activity. Patients with a normal natural killer cell function against K562 target cells had about 10% of lymphocytes containing the typical “parallel tubular arrays,” which were recognised as the ultrastructural marker of natural killer cell lymphocytes (fig 8). By contrast, patients with a depressed natural killer cell function were identified by the absence of parallel tubular arrays and the presence of about 10% of lymphocytes with numerous cytoplasmic multivesicular bodies (fig 9).

Fig 3 Cross sectional aspect of “tubular confronting cisternae” (F).

Fig 4 “Tubuloreticular structure” (→) inside “tubular confronting cisternae”.
that had not been infected with HIV. By contrast, the membrane of the lymphocytes actively producing HIV particles did not react with OKT8 or OKT4 monoclonal antibodies.

Discussion

"Tubuloreticular structures" and "Tubular confronting cisternae" are not specific for AIDS or ARC, but, nevertheless, they have not been reported in healthy homosexuals. Systemic lupus erythematosus, herpes simplex encephalitis, congenital rubella, cytomegalovirus pneumonia, simian non-A non-B hepatitis, and miscellaneous malignant disease are the other principal conditions in which tubuloreticular structures have been found. Tubuloreticular structures can also be generated in cell culture by α and β interferon but not by γ-interferon. Moreover,

Table  Analysis of natural killer population

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NKH1 = monoclonal antibody against natural killer cells; f NK = natural killer function; IL2 = interleukin-2; PTA = parallel tubular arrays; MVB = multivesicular bodies.
Immunostructural study of circulating lymphocytes in HIV infection

Tubuloreticular structures and tubular confronting cisternae seemed to be related to the progression of the disease, according to the studies of Sidhu and Orenstein. Interestingly, the presence of serum acid labile α-interferon in patients with ARC was also reported as a prognostic marker for progression to AIDS. With Orenstein, we consider that a particularly poor prognosis was associated with the detection of tubular confronting cisternae. This assumption should be further verified through prospective study.

Circulating plasma cells displayed an immature phenotype (HLA DR positive) and showed several ultrastructural features of myeloma cells, especially the presence of elongated mitochondria with poorly developed cristae in a clear matrix. Circulating plasma cells were associated with polyclonal hypergammaglobulinaemia in 91% of our cases. This finding is characteristic of the chronic stimulation, principally by HIV and EBV, of the B cell lineage.

"Hand-mirror cells" were first described by Lewis. Systemic treatment with α-interferon induced the appearance of tubuloreticular structures in circulating mononuclear cells. AIDS and systemic lupus erythematosus have a clinical common denominator—the presence in the serum of an unusual acid labile α-interferon. In the present study we noticed the simultaneous occurrence of tubuloreticular structures and acid labile α-interferon in 73% of the patients examined.

"Tubular confronting cisternae", referred to also as “test-tube and ring-shaped forms”, have a lower prevalence in HIV infection than tubuloreticular structures. They have rarely been reported elsewhere; in sporadic cases of simian non-A non-B hepatitis, HTLV-I related leukaemia, multiple sclerosis, and collagen diseases. The generation of tubular confronting cisternae in vitro and in vivo by α-interferon has not yet been confirmed. Tubular confronting cisternae, however, have never been detected in the absence of tubuloreticular structures, and both structures were closely connected in the lymphocytes of some of the aforementioned patients (fig 4). Moreover, the highest values of acid labile α-interferon in our study occurred in patients with tubular confronting cisternae. The prevalence of systemic lupus erythematosus.
They were subsequently reported in normal lymph nodes, infectious mononucleosis, lymphoma, myeloma, and in acute leukaemia. These uropod-bearing lymphocytes might be induced by virus or by retroviral antigens, or both. In infectious mononucleosis the demonstration of hand mirror cell with parallel tubular arrays and their lytic activity against B lymphocytes infected by Epstein-Barr virus was consistent with their natural killer phenotype. Moreover, uropod-bearing cells were often found among the large granular lymphocytes. In the present study the pre-embedding immunogold staining method showed a NKH1-positive membrane phenotype and LAV antigenic sites at the tip of the uropod. Hand mirror cells were also observed by Sidhu in HIV infection. He described an electrodense layer at the tip of the uropod, probably corresponding to adsorbed circulating immune complexes.

The large granular lymphocytes (LGL), which assume the antibody dependent cellular cytotoxicity and the natural killer function, were easily recognisable by electron microscopy owing to their “parallel tubular arrays”. The role of the multivesicular bodies in the genesis of the dense granules of large granular lymphocytes was clearly shown. Numerous multivesicular bodies were observed in immature forms of large granular lymphocytes devoid of natural killer cell function. We attempted to verify this association between the ultrastructural characteristics of large granular lymphocytes and the natural killer cell function against K562 target cells in nine of our cases (table 1). Four patients with depressed natural killer cell activity showed large granular lymphocytes with numerous multivesicular bodies, five patients with normal natural killer cell activity had large granular lymphocytes with typical parallel tubular arrays. These disturbances of the natural killer cell population seemed to be unrelated to the clinical stage of HIV infection. Despite the importance of interferons and interleukin-2 in the differentiation of natural killer cells no clear association between low natural killer cell activity and an abnormal production of interferon or interleukin-2 was found. In another study normal binding of natural killer cells to their targets but a defective killing of attached tumour cells were shown in HIV infection. Our results were also consistent with defective killing. A low natural killer cell activity corresponded to the existence of immature natural killer cells that were devoid of lysosomal parallel tubular arrays. Occurrence of immature...
natural killer cells could be explained by an increased turnover secondary to chronic HIV infection. A direct effect of HIV infection on natural killer cells was not excluded as natural killer cells have already been incriminated in the production of the abnormal acid labile α interferon.52

A striking feature was the presence of LAV antigen located in intracytoplasmic vacuoles inside large circulating cells that were sometimes multinucleated. The presence of incomplete intracytoplasmic retrovirus-like particles has been reported previously in fresh peripheral blood cells from a haemophiliac with AIDS.53 In this study both anti-p24 against bovine leukaemia virus6 and anti-p13 against LAV gave a positive reaction inside these vacuoles. This cross reaction (observed also in the vacuoles of fetal lamb kidney cells infected by bovine leukaemia virus) was probably promoted by the glutaraldehyde fixation (post-embedding immunogold staining method).

More recently, Pekovic54 also detected HIV antigen by immunofluorescence in peripheral lymphocytes sometimes displaying multinucleated giant forms. Surprisingly, we were unable to identify the membrane phenotype of these vacuolated circulating cells after the pre-embedding immunogold staining technique with anti-HLA DR, OKT4, OKT8, NKH1, and Leu7 monoclonal antibodies. Moreover, in a previous study6 no vacuolated giant cells were found in the non-rosetting subset of peripheral mononuclear cells nor in the purified OKT8 positive subset. The hypothesis of a change in the antigenic CD4 site and in the expression of the CD4 molecule in HIV infected cells has been suggested by Klatzmann et al.55 The role of the CD4 antigen as a receptor for the gp110 of HIV has been shown.56,57 During the initial attachment of HIV, the binding of the OKT4a monoclonal antibody seemed to be inhibited but after the production of viral antigens by HIV infected cells the expression of all the CD4 epitopes including OKT4 was found to be depressed.58,59 This last mechanism could underlie the lack of staining of the vacuolated lymphocytes with OKT4 monoclonal antibody.

In conclusion, this immunolstructural study of non-cultured circulating lymphocytes from patients infected with HIV allowed us to identify changes which suggest functional disturbances in the natural

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**Fig 10** Perinuclear zone of atypical vacuolated lymphocyte. $N =$ nucleus; $M =$ Mitochondria; $V =$ vacuoles.
70 Feremans, Huysgen, Menu, Farber, de Caluwe, van Vooren, Marcelis, Andre, Brasseur, Bondue, Lebon, Cluneck

killer cell population. In addition to the presence of possible prognostic ultrastructural markers (tubuloreticular structures, tubular confronting cisternae), the location of HIV antigen was also shown.

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References


Immunosstructural study of circulating lymphocytes in HIV infection


