Evaluation of some cellular immune index in HIV infected participants

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Abstract
The study was designed to evaluate some cellular immune index of HIV infected participants
For the study, 80 HIV infected participants were recruited for the study. They were destruction aged 15 – 55 years. 45 of these participants were classified as Symptomatic HIV (Stage 11), while the remaining 35 were Asymptomatic HIV (Stage 1). Similarly, 40 HIV seronegative participants served as Control. Blood samples collected from the participants were used HIV screening and confirmation, CD4+ T cell count, absolute lymphocyte count and percent lymphocyte transformation. The CD4+T cell count and percent Lymphocyte Transformation count were significantly lowered in HIV infected participants compared with the HIV Seronegative participants (p<0.05 in each case). Symptomatic HIV seropositive participants also presented with lowered CD4 and percent lymphocyte transformation, compared with the asymptomatic HIV participants (p<0.05 in each case). The lowered CD4+T cell count suggests possible quantitative of cellular immune cells (mainly Th1 cells). While the lowered percent blast formation in HIV infection indicates functional derangement of the cellular immune cells. Meanwhile no significant difference was observed in absolute lymphocyte count among the symptomatic, asymptomatic and control participants (p> 0.05).

Keywords: Human Immunodeficiency Virus (HIV), CD4+ T Cells, Lymphocyte Transformation and Absolute lymphocyte counts.
**Introduction**

HIV has increased sharply in Sub-Saharan African and Asia, and hence calls for deep concern.

HIV infection is characterized by high rate of viral replication throughout the course of the infection with resulting viral and immune mediated destruction of CD4 cells. Consequently, the infected individual becomes susceptible to opportunistic infections, malignancies and neurological diseases (Uchaikin 1989).

Uchaikin (1989) also observed a dramatic lowering of the absolute numbers of T helper lymphocytes and an appreciable decrease of the T helper/T suppressor ratio, associated with high levels of circulating immune depression at the early stage of the pathological process.

The present study was designed to consider some cellular index of immunity in HIV infected participants at two defined stages of HIV infection. This will allow us insight into possible pattern of these cellular immune indexes as HIV progresses. Through the determination of CD4 count, percent lymphocyte Transformation and absolute lymphocyte count.

**Methods and Selection of Study Population.**

Eighty HIV infected participants were recruited for the study at the VCT and HIV clinic of Nnamdi Azikiwe University Teaching Hospital Nnewi. They were aged 15-55 years. Using WHO classification for HIV infection and CD4 count, 45 of these patients were classified Symptomatic HIV (stage 11), while the remaining 35 were classified as Asymptomatic HIV (stage 1). Forty apparently healthy HIV seronegative participants drawn from students and staff population of Nnamdi Azikiwe University Teaching Hospital Nnewi, served as control.

8ml of blood was collected from the participants and dispensed into EDTA tubes for CD4+Tcell count, by Cyflow Counter method and lymphocyte transformation counts
by phytohaemagglutinin (PHA) stimulation in culture medium absolute lymphocyte by routine manual method.

Informed consent was obtained from those who participated in the study. The Nnamdi Azikiwe University Teaching Hospital Board of Ethical Committee approved the study design.

PROCEDURES
Test for HIV Infection by Immunochromatography as described by Manufacturers of the kit (Acon Laboratories Inc. USA).

The procedure was as described by the manufacturers of the kit (Acon Laboratories Inc. USA). 25ul of serum samples was dispensed into the “specimen pad” of the test strip. 80ul of buffer was added. The reaction was allowed for 5 minutes, the appearance of distinct red lines at test region and control region of the kit suggest positive HIV test while one distinct red line in the region of the control suggest HIV sero negative test. The appearance of the distinct red line of the control region validates the result without which the kit is assumed to be non functional.

HIV Confirmatory Test by Western Blot as described by Popovic et al (1984)

Method
2mls of the reconstituted phosphate buffer PH 8.6 was added to the dish containing the nitrocellulose strip. The cells were incubated for five minutes at room temperature under slow shaking. 20ul of each patient’s serum sample was added into the corresponding cell and incubated for 2 hours, at room temperature under slow shaking. The content of each dish was drained and each strip was washed twice for 5 minutes each time with 2mls of buffer. 2ml of enzyme linked anti IgG antibody was dispensed into each cell and incubated for 1 hour, 5 minutes at room temperature under shaking. 2mls of colour developing reagent (enzyme substrate) was dispensed into each cell after wasting. The colour reaction was stopped by removing reagent and washing 3 times
with buffer as soon as the colour developed. The presence of shades of colour indicates positive results No shades of colour representing gag antigen indicate indeterminant result.

**CD4+ T cell Count by Flow Cytometry**

20ul EDTA whole blood was collected into Partec test tube (Rohren tube). Then 20ul of CD4+T antibody was added into the tube .The contents were mixed and incubated in the dark for 15 minutes at room temperature. 800ul of CD4 buffer was gently added into the mixture and mixed gently. Then the Partec tube was plugged on the Cyflow counter and the CD4+T cells were displayed as peaks and interpreted as figures.

**Lymphocyte Transformation Assay as Described by Boyum (1964)**

**Lymphocyte Isolation**

Fresh blood samples was collected and dispensed into anti-coagulant (EDTA). 2ml each, of blood sample and balanced salt solution were dispensed into a 10ml test tube and the contents of the tube was carefully mixed; using a Pasteur pipette. The diluted blood sample was layered unto 3ml of Ficoll-paque solution in a centrifuge tube. The tube was centrifuged at 400g for 30mins at room temperature. The upper layer of the segments formed was removed, while the lymphocyte layer at the interface was pipette into another centrifuge tube. The isolated lymphocyte was washed in 6ml of the balanced salt solution and finally suspended in 0.5ml of the balanced salt solution. Lymphocyte viability test was done by the trypan blue exclusion test.

**Lymphocyte Transformation Test**

The lymphocytes were prepared into concentrations of 1.0 x10^6 cells/ ml of blood and cultured in the wells of microtitre culture plates, using TC 199 as culture medium and which had been enriched with AB blood group serum. 0.1ml of phytohaemagglutinin was added to each well of the microtitre plate. The culture mixtures were then incubated at 37° c for 72 hrs. At the end of the incubation period, cultures were centrifuged at 1,500 rpm for 7 minutes and the isolated Cells were then fixed in 1:10 glacial acetic
acid/ alcohol mixture and re-centrifuged. The cell pellets were finally resuspended in 0.5ml of culture medium. Thin blood film was prepared from the isolated cells in a microscope slides. The films were allowed to dry, stained with leishmans stain. The percentage of transformed T cells was determined for each sample. HIV negative Control samples were similarly processed. The lymphoblasts (as the transformed cells), were counted in a light microscope using x100 objective and the count expressed in percentage.

Statistical Analysis:
The result of the analysis was statistically analyzed. Student’s t, one way analysis of variance (ANOVA) were used to compare means. The analysis were performed with the use of SPSS statistical software package. P<0.05 was considered statistically significant.

Results

Lymphocyte Transformation and CD4+ T cell Counts.
The mean (±SD) percent lymphocyte transformation was significantly lowered in the asymptomatic 27.5 (±8.7) %, symptomatic 14.47 (±5.8) % HIV infected participants compared with corresponding values in control participants 59 (± 10) (p<0.01). Similarly, the mean (±SD) blood CD4+ T cell count (per ul of blood) of 300 (± 139) in symptomatic HIV infected patient was significantly lower than corresponding values of 590 (±389) in asymptomatic HIV infected patients and controls 943 (±436) (p<0.01 in each case). The mean (±SD) CD4 count in the symptomatic HIV infected patients was
significantly lower than the value in asymptomatic patients \((p<0.05)\). Similarly, the mean \((\pm SD)\) lymphocyte transformation in symptomatic HIV infected patients was significantly lower than the asymptomatic HIV infected patients \((p<0.01)\). See Table 1.

**Table 1: Mean \((\pm SD)\) CD4+ T Cell Count (per ul), Lymphocyte Transformation (percent) and Absolute Lymphocyte count \((x10^9)\) amongst HIV Infected Patients and Control Participants.**

<table>
<thead>
<tr>
<th></th>
<th>CD4+ T cells count (per ul)</th>
<th>Lymphocyte Transformation (%)</th>
<th>Absolute Lymphocyte count ((x10^9))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control ((n=40))</td>
<td>943 ± 436</td>
<td>59.3 ± 9.9</td>
<td>1.96 ± .38</td>
</tr>
<tr>
<td>Asymptomatic ((n=35))</td>
<td>590 ± 389</td>
<td>27.5 ± 8.8</td>
<td>2.2 ± .65</td>
</tr>
<tr>
<td>Symptomatic ((n=45))</td>
<td>300 ± 139</td>
<td>14.5 ± 5.8</td>
<td>2.08 ± .82</td>
</tr>
<tr>
<td>F(p)</td>
<td>29.6 ((0.000))</td>
<td>230.4 ((0.000))</td>
<td>940 ((0.408))</td>
</tr>
<tr>
<td>tlp(^a)</td>
<td>0.014</td>
<td>0.000</td>
<td>0.167</td>
</tr>
<tr>
<td>tlp(^b)</td>
<td>0.000</td>
<td>0.000</td>
<td>0.695</td>
</tr>
<tr>
<td>tlp(^c)</td>
<td>0.001</td>
<td>0.000</td>
<td>0.692</td>
</tr>
</tbody>
</table>

**Key:**

F \(p\) = symptomatic, asymptomatic and control compared (using ANOVA)

tlp\(^a\) = asymptomatic HIV patients compared with control (using students t)

tlp\(^b\) = symptomatic HIV patients compared with control (using students t)

tlp\(^c\) = asymptomatic HIV patients compared with symptomatic (using students t)
Discussion

This study reveals lowered, CD4+ T cells count in HIV in HIV infected participants. This observation may be attributed to T cell death caused by the HIV virus (Mark et al 2005). This results in the numerical decline of CD4+ T cells; the implication is immunodeficiency state in these individuals. There are some hypotheses to account for the T cell – death in HIV infection; the first is programmed cell death or apoptosis, where even uninfected and therefore unstimulated cells die within first 24 hours. The second mechanism is the activation, associated lymphocyte death, by which cells stimulated by strong mitogenic stimuli such as phytohaemagglutinin (PHA) die after 48 – 72 hours due to hyperactive stimulation (Grossman et al 2002). Antigen can induce special resting cells into activation burst, and this may be due to rapid cells proliferation and differentiation into effector cells over period of days or weeks.

There is also evidence that regulatory T lymphocyte (Tregs) are depleted during the course of HIV infection (Weiss et al 2002), the loss may facilitate the immune hyper activation which may in turn lead to increase CD4+ T cell death associated with HIV infection.

The destruction of the CD4+ T cell leads to derangement of the cellular immune response. Therefore there is derangement in the protective immunity, since the other types of immune response requires cognate help from the cellular immune response for effective immune mediation.

The role of T cell activation or immune activation in the pathogenesis of HIV infections has been highlighted in several reports (Grossman et al 2002, Bofill et al 1995, Hazenberg et al 2003, Sousa et al 2002) Immune activation may be a stronger predictor of the progression of HIV disease than either the CD4+ T cell count or viral load, and

It has been suggested that chronic immune activation arising from endemic pathogenic infections in Africa, can drive CD4+ T cell depletion even in HIV – sero negative individuals Bentwich et al (1996), Borkow et al (2001) and may convincibly contribute to HIV disease progression in co – infection (Eggena et al 2005).

In this study PHA was used to stimulate Blast Formation or lymphocyte transformation. The percent blast formation due to PHA was significantly lowered in participants. Only the T cells that have not been pre stimulated by the virus would be in a state to respond to pre-stimulation. The implication is that there is lowered blast formation (T cell function) in HIV infection. This leads to induced cellular immune derangements.

Significant differences in proliferative responses between HIV infected and control participants has been reported (Shakoor 2003, Bocchieri et al 1995). In Africa, Eggena et al (2005), found high levels of T cell activation in HIV infected Ugandans, when compared to their sero negative counterparts. We also observed that symptomatic HIV participants had lower percentage transformable T cells than asymptomatic group. This finding is most probably attributed to the level of disease progression. It would therefore appear from this observation that T cell activation or blast formation due to PHA, stimulation may correlate with disease progression. Some earlier reports had also associated a diminishing rate or percentage of lymphocyte transformation, with the progression of HIV infection towards the acquired immunodeficiency syndrome (AIDS) (Shakoor 2003, Bansal et al 1993).
The finding in the study leads to conclusion that the degree of T cell activation as determined by PHA – induced transformation decreases in HIV disease and rate of CD4+T cell depletion in HIV infection appear to associate with disease progression.

References


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