CONTENTS

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ORIGINAL ARTICLES

Haematologic features of the human immunodeficiency virus (HIV) infection in Black children in Harare

JO Adewuyi, I Chitsike ...................................................... 333

An investigation of the schistosomiasis transmission status in Harare

J Ndamba, MG Chidimu, M Zimba, E Gomo, M Munjoma .................................................. 337

Hepatic function tests in children with sickle cell anaemia during vaso occlusive crisis

A Ouowo, MA Adedoyin, D Fagbule ........................................ 342

The Zimbabwe external quality assessment scheme (ZEQAS) in clinical chemistry: results of the pilot programme

WB Mujaji, HN Mazhindu, ZAR Gomo, HT Marima-Matarira, C Samuwai, T Nyamayaro, DG Bullock, JG Ratcliff ........................................... 345

CASE REPORTS

Complete rectal prolapse in adults: a Tanzanian experience

Mr Aziz, NAA Mbembati ............................................. 349

Delayed diagnosis of retinoblastoma

SNN Nwosu, GSC Okoye, TO Ulası .................................... 353

Bilateral fracture of the femoral neck as a direct result of electrocution shock

L Nyoni, CR Saunders, AB Morar ........................................ 355

LETTERS TO THE EDITOR

The gastroscope, labour intensive family planning and incentives

DAA Verkuyl ........................................................ 356

REVIEW ARTICLES

Hydatidiform mole

P Zvandasara .......................................................... 357

BOOK REVIEW

Biological oxidants and antioxidants

YS Naik ........................................................... 362

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Haematologic features of the Human Immuno-deficiency Virus (HIV) Infection in Black children in Harare

*JO ADEWUYI, # I CHITSIKE

SUMMARY

Forty six Black Zimbabwean children aged between three months and seven years who were admitted into Parirenyatwa Central Hospital with serologically positive and symptomatic HIV infection were investigated for their haematologic profiles. Tests done included full blood counts, manual white cell differential counts, coagulation screening tests and bone marrow aspiration in clinically indicated cases.

Anaemia was found in 84 pc, leucocytosis in 60 pc and thrombocytopenia in 30 pc of the cases. In contrast to reports in adults leucopaenia or neutropaenia were not seen. Coagulation profiles were mostly normal but presumptive diagnosis of circulating coagulation inhibitor was made in one case.

Morphological changes suggestive of myeloid dysplasia and in particular dysgranulopoiesis were commonly seen. Bone marrow aspirates examined in eight of the children all showed hyper or normal cellularity with adequate and productive megakaryocytes.

INTRODUCTION

A variety of haematologic abnormalities have been described in patients with HIV infection and AIDS. Few reports however, describe the changes in African children. The aim of this study is to document the haematologic features seen in children with HIV infection admitted into the Parirenyatwa Hospital in Harare.

MATERIALS AND METHODS

Patients: Children were considered infected with the HIV virus if they presented with clinical features suggestive of HIV infection and they had a positive serological test. The clinical features met the Centre for Disease Control criteria for diagnosis of HIV infection. In children below 15 months of age, the added criteria of demonstrating the presence of the virus in blood or evidence of immune dysfunction were not met because of unavailability of resources to perform the tests. Therefore the diagnosis of HIV infection in children was made if they fulfilled both criteria above, irrespective of age. This definition is similar to that used by Nkrumah et al and Topley also in Zimbabwe.

Methods: The presence of anti-HIV-I antibodies was detected by two ELISA methods and in doubtful cases confirmed by Western Blot. Parents were offered pre and post test counselling. Forty six patients aged three months to seven years fulfilled the above criteria and formed the subjects of the study. The patients were all known to have acquired the infection perinatally and none was on AZT or any other antiviral agent nor on cotrimoxazole prophylaxis. Other antimicrobial agents were however, used as indicated.

Methods: All the children had the following tests done: automated full blood counts (Coulter JS), blood film examination, manual 200-cell white cell differential counts, coagulation screening tests comprising prothrombin time (PT), activated partial thromboplas-
tin time (APTT) and thrombin clotting time (TT). Correction tests and where indicated coagulation factor assays were done in cases of prolonged clotting times. Bone marrow aspiration (BMA) was done in cases of clinical bleeding and in those with platelet counts less than 100 x 10⁹/1.

Samples for coagulation tests were separated immediately and the plasma kept on ice and tested within one hour. Samples for blood counts and bone marrow aspirates were processed within two hours.

RESULTS

The age distribution of the children is show in Figure I and the main abnormal peripheral blood findings are summarised in Table I.

Table I: Abnormal blood counts in 46 children with HIV infection.

<table>
<thead>
<tr>
<th>Blood count</th>
<th>No</th>
<th>pc</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaemia</td>
<td>Hb &lt; 11g/dl</td>
<td>39</td>
</tr>
<tr>
<td>Leucocytosis</td>
<td>WBC &gt; 11 x 10⁹/1</td>
<td>27</td>
</tr>
<tr>
<td>Leucopaenia</td>
<td>WBC &lt; 3,0 x 10⁹/1</td>
<td>0</td>
</tr>
<tr>
<td>Neutrophilia</td>
<td>N &gt; 7,5 x 10⁹/1</td>
<td>12</td>
</tr>
<tr>
<td>Neutropaenia</td>
<td>N &lt; 1,5 x 10⁹/1</td>
<td>0</td>
</tr>
<tr>
<td>Lymphocytosis</td>
<td>L &gt; 7,0 x 10⁹/1</td>
<td>18</td>
</tr>
<tr>
<td>Lymphopaenia</td>
<td>L &lt; 2,5 x 10⁹/1</td>
<td>8</td>
</tr>
<tr>
<td>Eosinophilia</td>
<td>Eo &gt; 0,5 x 10⁹/1</td>
<td>2</td>
</tr>
<tr>
<td>Monocytosis</td>
<td>M &gt; 1,0 x 10⁹/1</td>
<td>5</td>
</tr>
<tr>
<td>Thrombocytopaenia</td>
<td>Ptt &lt; 150 x 10⁹/1</td>
<td>14</td>
</tr>
</tbody>
</table>

Figure I: Age distribution of children with HIV infection.

Anaemia: The bulk of the children (84 pc) were anemic with a haemoglobin concentration below 11g/dl. About half the anaemic children had a normochromic and normocytic blood picture while 40 pc had a hypochromic and microcytic anaemia with MCV and MCH below 75fl and 25pg respectively. The remaining 10 pc had significant macrocytosis with MCV above 100fl.

Leucocytosis and leucopaenia: About 60 pc of the children had leucocytosis (total WBC above 11 x 10⁹/1) with about equal proportions showing predominant neutrophilia and lymphocytosis. There was a left shift of neutrophils in about half the cases showing neutrophilia and in about 20 pc of those with normal white cells counts. A common qualitative abnormality of granulocytes was cytoplasmic vacuolation which was found in about 50 pc of the cases (Figure II). Toxoc granulations and Dohle bodies were less commonly seen. Some neutrophils exhibited cytoplasmic budding or extensions which contained no granules and stained pale blue; these could be confluent Dohle bodies.

Monocytes were increased in only about 10 pc of the cases and showed heavy cytoplasmic vacuolation. Only two children showed significant eosinophilia of greater than 0,5 x 10⁹/1. No cases of leucopaenia (wbc less than 3,0 x 10⁹/1) or neutropaenia (less than 1,5 x 10⁹/1) were seen but there was lymphopaenia of less than 2,0 x 10⁹/1 in 17 pc of the children.

Thrombocytopaenia: Fourteen children (30 pc) had platelet counts below 150 x 10⁹/1. Thromboocytopaenia was mild (100 - 149 x 10⁹/1) in eight cases (17 pc); moderate (50-99 x 10⁹/1) in three cases (6,5 pc) and...
severe (less than $50 \times 10^9/\text{l}$) in three cases (6.5 pc) with one of the latter having severe epistaxis.

**Bone marrow findings:** Bone marrow aspiration was performed in the eight children, two of whom were bleeding while six had moderate or severe thrombocytopenia without clinical bleeding. The findings are shown in Table II. All marrows had hyper or normal cellularity with adequate and productive megakaryocytes. In the erythroid series, dysplasia was seen as irregularity or lobulation of nuclei, presence of nuclear remnants in the cytoplasm and basophilic stippling. Myeloid dysplasia manifested as cytoplasmic vacuolation of granulocytic precursors and hypogranularity in a few cases. Megakaryocytes dysplasia was minimal.

**Table II: Bone marrow aspiration findings in eight children with HTV infection.**

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>pc</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normo — or hypercellular marrow</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td>Adequate megakaryocytes</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td>Low myeloid/erythroid ratio (&lt; 2:1)</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>High myeloid/erythroid ratio (&gt; 10:1)</td>
<td>2</td>
<td>25</td>
</tr>
<tr>
<td>Myeloid dysplasia/dysgranulopoesis</td>
<td>5</td>
<td>62</td>
</tr>
<tr>
<td>Erythroid dysplasia</td>
<td>2</td>
<td>25</td>
</tr>
<tr>
<td>Lymphocytosis (&gt; 40 pc)</td>
<td>2</td>
<td>25</td>
</tr>
<tr>
<td>Plasmacytosis (&gt; 5 pc)</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Histiocytosis</td>
<td>1</td>
<td>12</td>
</tr>
</tbody>
</table>

Bone marrow lymphocytosis of greater than 40 pc was found in two aspirates and mild plasmacytosis of 5 pc was found in one. The lymphocytes and plasma cells were normal in morphology. The presence or absence of lymphoid or plasma cell aggregates or increased fibrosis could not be determined as only one aspirate was accompanied with a trephine biopsy.

One aspirate showed marked infiltration by histiocytes with eccentric nuclei and copious pale and fibrillar cytoplasm. These were probably pseudo-Gaucher cells.

Coagulation screening tests: Prothrombin time and partial thromboplastin time were normal except for two children who had prolonged APTT. One of the two children was bleeding and also had prolonged PT and TT with other evidence of disseminated intravascular coagulation. The other child had normal PT and the moderately prolonged APTT (60:35 seconds) was not corrected by mixing with 50 pc normal plasma before or after incubation at 37°C for two hours.

This suggested the presence of a circulating inhibitor but further tests were not done to establish whether it was a lupus anticoagulant or antcardiolipin antibody.

**DISCUSSION**

The haematologic findings in this study broadly agree with those reported in other studies. The most common complication seen was anaemia which was most probably multifactorial. It could be nutritional or due to HIV infection per se or other infections such as pneumonia.

Thrombocytopenia was found in 30 pc of patients; non-vital causes such as malaria were excluded. Though tests for specific anti-platelet antibodies or platelet-associated immunoglobulins were not done, thrombocytopenia in our patients was probably not due to auto-immune causes since classic immune thrombocytopenic purpura is usually acute and self limiting and recovery is the rule in the age group of most of our patients.

Our results confirm the finding by others that thrombocytopenia is an early and common feature of HIV infection children. It is therefore advisable in children with unexplained thrombocytopenia to exclude HIV infection before embarking on invasive investigations.

In contrast to reports in adults leucopaenia and neutropaenia were not found as features of HIV infection in these children. On the contrary about 60 pc of them had leucocytosis with half being predominantly neutrophilic. Sandhaus et al also did not observe leucopaenia in their small series. Leucocytosis could be secondary to infection.

The presence of vacuolated monocytes has been described in HIV infection and neutropaenia were not found as features of HIV infection in these children. On the contrary about 60 pc of them had leucocytosis with half being predominantly neutrophilic. Sandhaus et al also did not observe leucopaenia in their small series. Leucocytosis could be secondary to infection.

The presence of vacuolated monocytes has been described in HIV infection and this feature was very conspicuous in our study. Particular attention has, however, not been drawn to vacuulations in peripheral blood neutrophils and immature granulocytes, a phenomenon we observed in nearly 50 pc of our patients. These changes could not have been storage artefacts since samples were processed within two hours and neutrophil vacuolations were only occasionally observed and few in routine samples from non HIV-infected children.
It is not clear whether these vacuolations were the result of intense phagocytic activity or a further sign of myeloid dysplasia. Features of dysgranulopoiesis which have previously been described include hypogranularity\textsuperscript{11} and giant neutrophils with increased peroxidase activity.\textsuperscript{12} Functional impairment of neutrophils in childhood HIV infection has also been demonstrated in at least one study.\textsuperscript{13}

The frequency of prolonged APTT attributable to circulating coagulation inhibitors appeared to be lower in our study (1 in 46) than has been reported\textsuperscript{14} and bleeding or thrombosis due to inhibitors was not encountered.

REFERENCES
