

Effects of Highly Active Antiretroviral Treatment on Complete Blood Count parameters

ABSTRACT

Aim: This study assesses the effects of HAART on complete blood count parameters among HIV infected participants.

Study design: Case control study (**Cross-sectional study**)

Place and Methods: This study was conducted in Tamale, Ghana from August, 2015 to November 2017.

Methodology: A total of 300 HIV infected participants with ages ranging from 19–79 years, administered with HAART for at least 6 months were recruited. Pre-HAART administration (baseline) demographic and clinical information, with initial full blood count results were retrieved from the medical records of the participants. Post HAART administration blood sample (5mLs) was taken from each participant into an EDTA vacutainer tube and complete blood count (CBC) performed using URIT 5250 haematology analyser. Participants transfused with blood for the last 4 months were excluded from the study.

Results: The study recorded significant decreases in WBC and Neutrophil % post HAART administration. Lymphocyte (%), Haemoglobin, Haematocrit, MCV, MCHC, RDW-SD were all significantly higher post HAART administration. Total Platelets count, MPV, PDW-SD, PCT and P-LCR were significantly lower post-HAART administration. A comparison of the effects of EFV and NVP administered with AZT/3TC backbone yielded the following results. The NVP group recorded a significantly higher HCT compared with the EFV group (**p=0.0073**). A significantly higher mean PCT, MPV, P-LCR, PLCC, PDW-SD were recorded in the EFV group compared to the NVP group respectively.

Conclusion: The administration of HAART is associated with significant improvements in erythroid and lymphoid lineages, reduce anaemia, improves immunity and general patient well-being. NVP improve erythroid cell indices while EFV ameliorate platelet indices. HAART regimen should be chosen based on the pre-HAART laboratory tests conducted on the individual.

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Keywords: Highly Active Antiretroviral Therapy, HIV infection, haematological abnormalities

1. INTRODUCTION

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HIV infection results in severe systemic disorder, with characteristic impairment and progressive damage to both humoral and cellular immune responses [1]. Haematological abnormalities may be the first laboratory findings in HIV infection and may involve all cell lines and noted to be strong independent predictors of morbidity and mortality in HIV infected persons [2, 3]. Calis and colleagues reported a prevalence of 3-38% moderate anaemia (Hb 8.0-9.9g/dL) and 50-91% mild anaemia (Hb10.0-12.0g/dL) among people HIV infected individuals in tropical areas, which increases as the disease progresses and varies with age, gender and the definition used to establish anaemia [2]. The causes of anaemia in HIV infected individuals included decreased erythropoietin production, ineffective erythropoiesis, opportunistic infections, neoplasia and micronutrient deficiency [4]. Iron deficiency has also

26 been reported as the commonest cause of nutritional anaemia among HIV infected
27 individuals [3]. The type of anaemia serves as a guide to the choice of HAART and treatment
28 option for opportunistic infections. Neutropenia was reported in 10% more of HIV infected
29 individual with advanced immunosuppression than in non-advanced HIV infected individuals
30 [5, 6]. Asymptomatic thrombocytopenia was reported in 20%-33% of HIV infected individuals
31 but increased with progression of the disease [6, 7].

32 Kibaru (3) reported that the use of zidovudine, lamivudine and stavudine was associated
33 with significant amelioration in hemoglobin concentration and after 12 months of HAART
34 use, the prevalence of anaemia reduced from 65.5 % to 46% [3]. In another study, after 3
35 months of HAART administration, Huang (8) documented significant increases in mean Hb
36 from 13.9 to 14.1 g/dl [8]. Huang (8) reported improvements in mean cell volume (MCV) from
37 55 to 98.9fl to 105.5fl and 106fl at 3, 6, 9, 12 months respectively after HAART use [8].

38 The current regimens used in Ghana comprise two nucleotide reverse transcriptase
39 inhibitors (NRTI) plus one nonnucleotide reverse transcriptase inhibitor (NNRTI) or two NRTI
40 and one protease inhibitor (PI) [9]. The WHO recommends HAART for all HIV infected
41 individuals since it improves morbidity and mortality associated with HIV infection, and offers
42 better life expectancy [10, 11]. Despite the beneficial effects associated with HAART use,
43 studies have shown that steps must be taken to prevent life threatening side effects and
44 HAART related haematotoxicity [12]. In resource constrained developing countries, based on
45 availability, safety and efficacy, the WHO has made efavirenz, tenofovir in addition to
46 lamivudine or emtricitabine plus nevirapine as the preferred first line antiretroviral
47 medications, though so much information is not available on HAART use in these areas [13].
48 Nevirapine remains the NNRTI of choice when efavirenz cannot be used. Better rates of HIV
49 replication suppression is achieved in patients administered with efavirenz-based HAART
50 who received tenofovir-emtricitabine compared with patients who received zidovudine-
51 lamivudine [14]. In order to avoid the toxicities associated with Zidovudine use, especially in
52 individuals with anaemia, global access to tenofovir has been increased [15]. Not many
53 studies on the haematological parameters abnormalities in HIV infected persons have been
54 conducted in sub Saharan Africa. This study examines the effect of HAART on complete
55 blood count parameters.

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57 **2. MATERIAL AND METHODS / EXPERIMENTAL DETAILS / METHODOLOGY**

58 **Study Design**

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60 This was a cross sectional study carried out from 12th August 2016 to 21st December 2017.

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62 **Study population**

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64 A total of 300 HIV infected participants with ages ranging from 19 to 79years and have been
65 on HAART for at least 6 months were recruited. Participants who have been transfused with
66 blood for less than 4 months were excluded from the study. (How was the sample-size
67 arrived at? In case sample-size was calculated, indicate the formula used, and provide a
68 Reference for any previous Study used in the calculation). (Was Random Sampling done? If
69 yes, indicate the method of sampling used)

70 **Data Collection**

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72 Pre-HAART administration (baseline) information such as weight, systolic and diastolic blood
73 pressure, age, date of HAART initiation and initial complete blood count results were
74 retrieved from the medical records of the participants. Post HAART information such as age,
75 weight, blood pressure measurement, HAART type and duration of HAART use were
76 recorded using questionnaire which had been pre-tested among 10 HIV infected individuals
77 administered with HAART to clear possible ambiguity and difficulty in answering the

78 questions. Data from the pre-tested questionnaires were however not included in the results
 79 analysis.
 80 Blood samples (5mLs) was taken into an EDTA vacutainer for Complete Blood Count (CBC)
 81 analysis. The CBC was performed using URIT 5250, a 5-part differential, 28 parameters
 82 haematology analyser, from URIT Medical Electronic (group) Co., LTD, China,
 83 <http://www.urit.com/index.aspx>. The HAART comprised two Nucleotide Reverse
 84 Transcriptase Inhibitor (NRTI) plus one Nonnucleotide Reverse Transcriptase Inhibitor
 85 (NNRTI) or two NRTIs and a Protease Inhibitor (PI). The participants were further stratified
 86 into short term (<52 months), medium term (≥52 but ≤104months), long term (>104months)
 87 based on the duration of HAART.

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89 **Statistical analysis**

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91 Data was entered into Microsoft excel 2016 and exported to GraphPad prism version 6.0
 92 (www.graphpad.com) for analysis. Data was presented as number, percentages, means and
 93 standard deviation. Means were compared between groups using Student's paired t-test and
 94 ANOVA and p<0.05 was considered statistically significant.

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97 **3. RESULTS AND DISCUSSION**

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99 Table 1 shows the demographic data of the study population. Out the 300 participants,
 100 majority (80.7%) were females, 176 (58.7%) were short-term HAART users, 96 (32%) were
 101 medium-term HAART users while 28 (9.3%) were long-term HAART users.

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Table 1: Demographic characteristics of the study population

Variables	Pre-HAART (n=300)	Post-HAART (n=300)
Age (years)*	35.4 ± 9.4	39.7 ± 10.0
Weight (Kg)*	58.8 ± 12.8	64.3 ± 25.6
Age Group (years)		
10-19	10 (3.3%)	3 (1%)
20-29	76 (25.3%)	36 (12%)
30-39	125 (41.7%)	120 (40%)
40-49	64 (21.3%)	96 (32%)
50-59	20 (6.7)	34 (11.3%)
60-69	5 (1.7%)	13 (4.3%)
70-79	0	1 (0.3%)
Gender		
Male	58 (19.3%)	58 (19.3%)
Female	242 (80.7%)	242 (80.7%)
HAART Duration (Months)		
Short term	0	176 (58.7%)
Medium term	0	96 (32%)
Long term	0	28 (9.3%)

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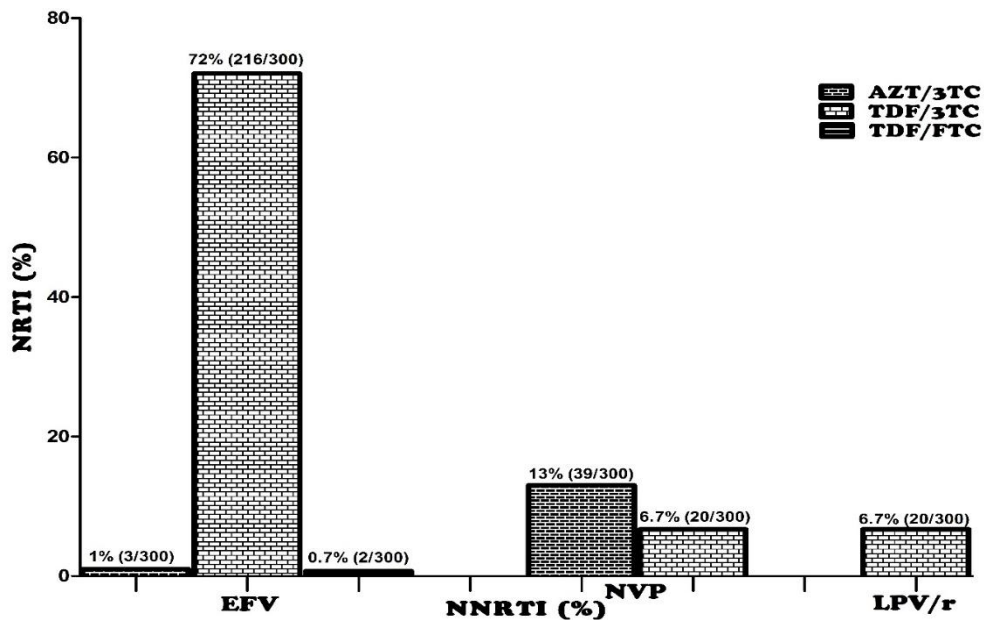
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*Data are presented as frequencies and percentages, * - comparison between pre and post HAART administration, with p-value of paired t-test <0.001. P-values 0.05 were considered significant.*

(Leave out significant difference in mean-age – the mean-age naturally expected to increase after at least 6 months of HAART)

112 **Distribution of HAART in the Study Population**

113 HAART regimen was distributed as follows, TDF+3TC+EFV (72%), AZT+3TC+NVP (13%),
 114 TDF+3TC+NVP (6.7%), TDF+3TC+LPV/r (6.7%), AZT+3TC+EFV (1%) and TDF+FTC+EFV
 115 (0.7%) regimen.



116 **Fig. 1. Percentage distribution of the various types of HAART regimen administered at**
 117 **the ART centers**

120 Table 2 compared CBC parameters in the study population pre and post-HAART
 121 administration. WBC (%) 7.446±5.066 vrs 4.733±2.008, neutrophil (%) 51.52±18.37 vrs
 122 44.26±14.46, basophils (%) 4.0±0.2 vrs 3.9±0.06 were significantly higher in pre-HAART
 123 administration while lymphocyte (%) 33.11±14.05 vrs 43.94±12.87 increased significantly
 124 post-HAART administration.

125 Haemoglobin (g/dL) 10.88±2.255 vrs 11.63±2.401, haematocrit (%) 32.75±6.239 vrs
 126 34.85±6.932, MCV 81.96±10.77 vrs 88.0±12.96, MCHC 33.52±1.651 vrs 34.50±5.368,
 127 RDW-SD 44.26±11.78 vrs 68.69±42.64 were significantly higher post-HAART administration
 128 (Table 2). (Spell out each of these abbreviations first, and briefly define them).

129 Again, thrombocytopenia was significantly pronounced post-HAART administration, with PLT
 130 284.7±148.1 vrs 254.4±145.7, MPV 9.028±1.773 vrs 6.840±1.601, PDW 14.55±6.180 vrs
 131 8.881±4.37, PCT 0.260±0.140 vrs 0.169±0.085 and P-LCR 28.87±9.586 vrs 20.60±15.12
 132 significantly decreased post-HAART administration (Table 2).

133 **Table 2: Complete blood count tests parameters pre and post HAART administration**

Parameters	Pre-HAART (n-300)	Post-HAART (n-300)	P – value
White blood cell count (10 ⁹ /L)	7.5 ± 5.1	4.7 ± 2.0	< .00
Lymphocytes %	33.1 ± 14.1	43.9 ± 12.9	< .00
Monocytes %	10.0 ± 4.9	9.8 ± 5.7	.31
Neutrophils %	51.5 ± 18.4	44.3 ± 14.5	.00
Eosinophils %	4.2 ± 3.7	3.7 ± 3.4	.63
Basophils %	0.4 ± 0.2	0.08 ± 0.06	.00
Red Blood Cell Count (10 ¹² /L)	4.0 ± 0.9	3.9 ± 0.7	.79
Haemoglobin (g/dL)	11.0 ± 2.6	11.6 ± 2.4	.04
Haematocrit (HCT, %)	32.8 ± 6.3	34.9 ± 6.9	.01

Mean Cell Volume (MCV, fL)	82.0 ± 10.8	88.0 ± 13.0	.00
Mean Cell Haemoglobin (MCH, pg)	29.4 ± 18.1	30.5 ± 5.3	.78
MCHC (g/dL)	33.5 ± 1.7	34.5 ± 5.4	.03
RDW-SD (fL)	44.3 ± 11.8	68.7 ± 42.6	.00
RDW-CV (%)	13.9 ± 4.9	15.7 ± 10.7	.13
Platelets (10 ⁹ /L)	284.7 ± 148.1	254.4 ± 145.7	.00
MPV (fL)	9.0 ± 1.8	6.8 ± 1.6	< .00
PDW (fL)	14.6 ± 6.2	8.9 ± 4.4	< .00
Platelet crit (PCT, %)	0.3 ± 0.1	0.2 ± 0.1	< .00
P-LCR (%)	28.9 ± 9.6	20.6 ± 15.1	< .00
P-LCC (10 ⁹ /L)	72.8 ± 45.7	48.4 ± 27.7	0.32

135 **Data are presented as mean ± SD. Table shows paired t-test comparison of complete blood**
 136 **count parameters before and after HAART administration. P-values 0.05 were considered**
 137 **significant.**

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 139 Table 3 shows the influence of duration of HAART usage on CBC parameters. Neutrophil
 140 (%) and basophils (%) decreased significantly (p<0.05) from short-term HAART users,
 141 medium-term through to long-term HAART users. The significant reduction in neutrophil (%)
 142 was from 46.36±14.26 in short-term HAART users to 39.53±15.13 in long-term HAART
 143 users while basophils (%) were from 0.065±0.049 to 0.053±0.032 to 0.002±0.001 in short-
 144 term, medium-term to long-term HAART users respectively.
 145 However, lymphocytes (%), haemoglobin concentration, haematocrit, mean cell volume
 146 (MCV) and mean cell haemoglobin (MCH) increased significantly (p<0.05) with duration on
 147 HAART uses. Haemoglobin concentration increased significantly from 11.31±2.394,
 148 12.25±2.172 and 12.93±7.352 in short-term, medium-term and in long-term HAART users
 149 respectively. While for lymphocytes, the significant increased were between 42.13±13.18 in
 150 short-term to 48.74±11.50 in long-term HAART users and from 46.03±12.04 in medium-term
 151 to 48.74±11.50 in long-term HAART users. For haematocrit, there was a significant
 152 increased between 33.93±5.945 in short-term to 36.49±8.026 in medium-term HAART users
 153 only but for MCV and MCH, significant increased were observed between short-term
 154 (86.56±13.02fL and 29.60±5.132ng) and median-term (91.20±12.43fL and 31.64±5.612ng)
 155 HAART users as well as short-term (86.56±13.02fL and 29.60±5.132ng) and long-term
 156 (92.25±14.67fL 33.04±5.172ng) and HAART users respectively.

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Table 3 Complete Blood Count based on the duration of HAART usage

PARAMETER	DURATION OF HAART			F test	P - value
	Short-term n-176	Medium-term n-96	Long-term n-28		
White blood cell count (10 ⁹ /L)	4.7 ± 1.6	4.5 ± 1.7	4.9 ± 2.0	0.8	.46
Lymphocytes %	42.1 ± 13.2	46.0 ± 12.0 ⁺	48.7 ± 11.5 [*]	5.1	.01
Monocytes %	9.4 ± 5.5	10.7 ± 6.4	10.7 ± 8.3	1.7	.18
Neutrophils %	46.4 ± 14.3	41.7 ± 14.0	39.5 ± 15.1 [*]	5.0	.01
Eosinophils %	3.4±2.8	3.4 ± 2.6	3.0±2.8	0.1	.91
Basophils %	0.07 ± 0.05 [@]	0.05 ± 0.03 ⁺	0.002 ± 0.001 [*]	11.4	< .00
Red Blood Cell Count (10 ¹² /L)	4.0 ± 0.7	4.0 ± 0.6	3.748 ± 0.7	1.2	.29
Haemoglobin (g/dL)	11.3 ± 2.4 [@]	12.3 ± 2.2 ⁺	12.9 ± 7.4 [*]	4.9	.01
Haematocrit (HCT, %)	33.9 ± 6.0 [@]	36.5 ± 8.0	35.6 ± 8.1	4.5	.01
Mean Cell Volume (MCV, fL)	86.6 ± 13.0 [@]	91.2 ± 12.4	92.3 ± 14.7 [*]	5.2	.01

Mean Cell Haemoglobin (MCH, pg)	29.6 ± 5.1 [@]	31.6 ± 5.6	33.0 ± 5.2	7.9	.00
MCHC (g/dL)	34.4 ± 6.0	34.8 ± 4.8	35.0 ± 3.7	0.3	.76
RDW-SD (fL)	65.7 ± 13.4	66.9 ± 12.6	68.9 ± 11.8	0.8	.46
RDW-CV (%)	14.7 ± 5.1	14.2 ± 2.8	15.4 ± 7.1	0.5	.78
Platelets (10 ⁹ /L)	251.1 ± 102.9	254.8 ± 213.4	227.8 ± 82.2	0.5	.61
MPV (fL)	6.9 ± 1.6	6.8 ± 1.7	6.8 ± 1.7	0.3	.72
PDW (fL)	9.0 ± 4.5	8.8 ± 4.5	8.0 ± 2.7	0.7	.52
Platelet crit (PCT, %)	0.2 ± 0.1	0.19 ± 0.13	0.16 ± 0.07	0.5	.61
P –LCR (%)	20.3 ± 9.5	19.2 ± 8.2	21.2 ± 9.4	0.7	.49
P –LCC (10 ⁹ /L)	50.6 ± 30.1	44.4 ± 22.2	47.8 ± 26.3	1.6	.20

161 *Data presented as mean ± SD. Shows One-way anova comparison of Complete Blood*
 162 *Count parameters in short, medium and long-term HAART users. @ - comparison between*
 163 *short and medium term, * - comparison between short and long term, + - comparison*
 164 *between medium and long term HAART users. P-values 0.05 were considered significant.*

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 166 Table 4 considered the effects of EFV and NVP administered with a common backbone
 167 AZT/3TC. Haematocrit 32.50±5.328 vrs 33.87±5.559 was significantly higher in the NVP
 168 group compared with EFV group. However, PCT 0.263±0.124 vrs 0.158±0.062, MPV
 169 8.667±1.290 vrs 6.479±1.502, P-LCR 34.70±9.930 vrs 17.32±8.894, P-LCC 111.0±78.89 vrs
 170 41.64±21.54 and PDW-S 12.97±3.083 vrs 7.567±2.255 were significantly higher in the EFV
 171 group compared with the NVP group. The EFV group had a more favourable effect on
 172 platelet indices compared with the NVP group. Comparatively, there were no significant
 173 variations in the effects of EFV, LPV/r and NVP administered with a TDF/3TC back bone on
 174 CBC parameters

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186 **Table 4 Effects of EFV, NVP and LPV/r with common backbones (AZT/3TC and TDF/3TC) (Indicate at this Table the exact stastical**
 187 **test done)**

PARAMETER	AZT/3TC BACKBONE			TDF/3TC BACKBONE			
	EFV GOURP	NVP GROUP	P - value	EFV GROUP	LPV/r GROUP	NVP GROUP	P - value
White blood cell count (10 ⁹ /L)	3.9±1.6	4.6±1.4	.38	4.7±1.7	5.2±2.5	5.1±4.3	.39
Lymphocytes %	35.5±27.0	45.8±13.5	.24	43.8±12.9	44.1±11.3	43.7±10.4	.99
Monocytes %	2.7±1.9	3.6±2.7	.59	3.8±3.7	1.7±0.9	3.7±1.9	.28
Neutrophils %	0.07±0.02	0.06±0.03	.66	0.06±0.04	0.05±0.02	0.08±0.06	.43
Eosinophils %	8.9±2.0	10.1±6.2	.73	9.5±5.2	10.9±7.7	10.8±7.3	.39
Basophils %	55.3±25.7	41.8±17.1	.21	44.6±14.0	43.4±13.1	44.1±13.0	.93
Red Blood Cell Count (10 ¹² /L)	3.5±0.3	3.5±0.7	.98	4.0±0.7	4.0±0.7	4.01±0.6	.97
Haemoglobin (g/dL)	10.5±2.4	11.8±2.08	.33	11.6±2.4	11.86±2.5	11.4±3.5	.86
Haematocrit (HCT, %)	32.5±5.3	33.9±5.6	.01	34.6±5.7	34.9±7.5	39.8±15.45	.01
Mean Cell Volume (MCV, fL)	92.6±13.5	97.4±13.6	.55	87.0±12.1	87.0±13.9	92.0±13.3	.22
Mean Cell Haemoglobin (MCH, pg)	30.1±6.6	34.3±5.4	.21	29.9±4.8	29.4±4.3	31.4±8.1	.39
MCHC (g/dL)	32.1±2.7	35.2±3.7	.17	34.5±5.5	33.6±4.4	33.8±7.8	.68
RDW-SD (fL)	71.9±9.8	72.9±13.9	.90	68.7±49.7	64.7±11.9	65.2±12.2	.89
Platelets (10 ⁹ /L)	321.3±182.0	242.9±71.6	.11	254.4±161.9	267.3±112.5	247.1±112.5	.91
Platelet crit (PCT, %)	0.3±0.1	0.2±0.1	.01	0.17±0.09	0.2±0.1	0.2±0.1	.51
MPV (fL)	8.7±1.3	6.5±1.5	.02	6.8±1.6	7.1±1.7	6.7±2.0	.75
P –LCR (%)	34.7±9.9	17.3±8.9	.00	21.0±17.0	21.7±8.3	18.8±6.9	.81
P –LCC (10 ⁹ /L)	111.0±78.9	41.6±21.5	.00	46.9±23.0	55.3±30.2	46.7±28.1	.33
PDW-S (fL)	13.0±3.1	7.5±2.3	.00	9.0±4.5	10.2±6.9	8.6±2.0	.46

188 *Data are presented as mean ± SD. Shows a comparison of the individual influences of EFV and NVP (with AZT/3TC backbone), EFV and*
 189 *LPV/r (with TDC/3TC backbone) on Complete Blood Count parameters (CBC). P-values 0.05 were considered significant.*

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193 Results in Table 6 shows the effects of AZT and TDF administered with 3TC/EFV or
 194 3TC/NVP. In comparing AZT with TDF (administered with 3TC/EFV), P-LCC 111.0±78.89
 195 vrs 46.88±22.97 was significantly higher in the AZT group compared with the TDF group. In
 196 comparing the AZT group with TDF group (administered with 3TC/NVP), RBC 3.525±0.651
 197 vrs 4.008±0.614 and HCT 33.87±5.559 vrs 39.80±15.48 were significantly higher in the TDF
 198 group compared with the AZT group. However, RDW-SD 33.87±5.559 vrs 39.80±15.48 was
 199 significantly higher in the AZT group compared with the TDF group.

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201 **Table 5 Comparison of the individual effects of AZT and TDF (administered with**
 202 **3TC/EFV and 3TC/NVP) on Complete Blood Count parameters ((Indicate at this Table**
 203 **the exact stastical test done)**

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PARAMETER	3TC/EFV			3TC/NVP		
	AZT	TDF	P = value	AZT	TDF	P = value
WBC	3.9±1.6	4.7±1.7	.42	4.6±1.4	5.1±4.3	.52
Lymphocyte (%)	35.5±27.0	43.8±12.9	.28	45.8±13.5	43.7±10.4	.55
Eosinophil (%)	2.7±1.9	3.8±3.7	.64	3.6±2.7	3.7±2.0	.94
Basophil (%)	0.07±0.03	0.06±0.04	.81	0.06±0.03	0.08±0.05	.22
Monocyte (%)	8.9±2.0	9.6±5.2	.83	10.1±6.2	10.8±7.3	.71
Neutrophil %	55.3±25.7	44.6±14.0	.20	41.8±17.1	44.1±13.0	.61
RBC	3.5±0.3	4.0±0.7	.20	3.5±0.7	4.0±0.6	.01
HBG	10.5±2.4	11.6±2.4	.42	11.8±2.1	11.4±3.5	.65
HCT	32.5±5.3	34.6±5.7	.53	33.9±5.6	39.8±15.5	.04
MCV	92.6±13.5	87.0±12.1	.43	97.4±13.6	92.0±13.3	.16
MCH	30.1±6.6	29.9±4.8	.93	34.3±5.4	31.4±8.1	.11
MCHC	32.1±2.7	34.5±5.5	.44	35.2±3.7	33.8±7.8	.36
RDW-CV	16.2±1.6	14.0±7.5	.61	13.9±3.5	14.4±3.0	.60
RDW-SD	71.9±9.8	68.7±49.7	.91	72.9±13.9	65.2±12.2	.04
PLT	321.3±182.0	254.4±161.9	.48	242.9±71.6	247.1±112.5	.85
PCT	0.3±0.1	0.2±0.1	.05	0.2±0.1	0.2±0.1	.33
MPV	8.7±1.3	6.8±1.6	.05	6.5±1.5	6.7±2.0	.57
P - LCR	34.7±9.9	21.0±17.0	.17	17.3±8.9	18.8±6.9	.52
P - LCC	111.0±78.9	46.9±23.0	< .00	41.6±21.5	46.7±28.1	.45
PDW-S	13.0±3.1	9.0±4.5	.12	7.6±2.3	8.6±2.0	.10

205 **Shows a comparison of the effects of AZT and TDF (administered with 3TC/EFV and**
 206 **3TC/NVP) on Complete Blood Count. Data are presented as mean ± SD. P = values 0.05**
 207 **were considered significant.**

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209 Discussion

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211 The findings in this study show that, the percentage of short, medium and long-term HAART
 212 users increased gradually from 9.3% in long-term HAART users through 32.0% in medium-
 213 term HAART users to 58.7% in short-term users. This reflects a gradual increase in the
 214 number of individuals enrolled on HAART. This shows that more HIV infected individuals are
 215 being recruited unto HAART. This results are in agreement with the findings of Atuyambe
 216 (16) and Nii-Trebi (15) who also recorded an increased number of HIV infected individuals
 217 on HAART.

218 The mean duration of HAART is denoted by the mean difference between Post HAART age
 219 and pre-HAART age. The participants have been on HAART for a long duration as the post-
 220 HAART age is significantly higher than the pre-HAART age. This shows that the participants
 221 have been on HAART long enough for HAART to elicit its effect on CBC parameters.

222 There was a significant increase in the body weight of the study population post HAART
223 administration. This indicates an improvement in the general health status of the individuals
224 or the absence of frequent illness in the study population as a result of immune restoration
225 indicated by improvement in lymphocyte count. Guillén (17) and [18] reported increases in
226 the body weight of people living with HIV post HAART administration.

227 The study found significant decline in total WBC post HAART administration and a
228 consistent decline in neutrophil and basophil counts from short-term through to long-term
229 among post HAART users post HAART administration. This indicates gradual development
230 of severe leucopenia with time. Other studies also found leucopenia in HIV infected
231 individuals who commenced HAART [19-21]. (Describe the Findings of those Studies in
232 some detail here) Leucopenia may be largely due to neutropenia caused by HIV suppression
233 of the bone marrow, leading to ineffective granulopoiesis [21, 22]. Kimura (22) observed
234 neutropenia may also be caused by the presence of antigranulocyte antibodies which attack
235 and destroy granulocytes. Many of the HAART drugs are considered as myeloid
236 suppressive, especially Zidovudin, hence should be reviewed for patients who already have
237 cytopenias [23]. The leucopenia in the present study may possibly be due to the presence of
238 inflammation in the tissues as a result of exposure to the drug. There is direct leucopenic
239 inhibitory effect on matured granulocytes in the peripheral system or on the myeloid
240 progenitor cells in the bone marrow. Consolini (5) reported leucopenia after six months of
241 AZT based HAART [5, 24, 25]. However, the findings of this study contradicted the findings
242 of [3] Kibaru and his colleagues who did not record leucopenia post HAART administration
243 [3]. (Try to offer an explanation, citing References)

244 Lymphocyte count post HAART administration increased significantly in a time dependent
245 manner. This is accounted for by the sturdy steady-state equilibrium in favour of endogenous
246 cell provisions over virus mediated cell killing. In the tap and drain model proposed by Wei
247 (26), decreased viral killing of lymphocytes quickly turn the balance in favour of lymphocyte
248 production and survival which allows at least partial immune reconstitution to occur. This
249 signifies a shrinkage in the myeloid lineage in the favour of the lymphoid lineage [27]. Many
250 other studies also found an increase in lymphoid tissue activity after HAART administration
251 [28-30]. Studies suggest that, the observed increase in lymphoid activity was a response
252 from the immune system following HAART administration [28-30].

253 This study also recorded significant improvements in erythroid cell line and indices. HGB,
254 HCT, MCV and RDW-SD were significantly higher post HAART administration in a time
255 dependent manner. The present study findings are consistent with studies by [4] and [31]
256 who also found increases in red cell indices post-HAART administration. These findings are
257 also consistent with the findings of Ogunbusuyi (32) who reported the prevalence of
258 anaemia as higher in treatment naïve patients compared with those on HAART. Other
259 studies reported a decline in the erythroid lineage due to the toxic effect of medications on
260 the bone marrow, nutritional deficiency especially B12, iron deficiency, decreased
261 erythropoietin release, gastrointestinal bleeding, malabsorption, autoimmune antibodies to
262 haemopoietic precursors etc [33] while other studies attribute the increased incidence of
263 anaemia to the direct effect of HIV on bone marrow stroma as a prelude to bone marrow
264 failure [23].

265 Thrombocytopenia was recorded as an early haematological abnormality in HIV infected
266 individuals in sub-Saharan Africa [19]. The incidence of thrombocytopenia post HAART
267 administration in this study was significant. This finding is consistent with the findings of
268 Akinbami (23) who also recorded higher incidence of thrombocytopenia post HAART
269 administration. The possible explanation may be due to increased platelet destruction by the
270 deposition of immune complexes on platelets and decreased platelet production, which
271 results in decreased MPV and plateletcrit [27]. Thrombocytopenia is also a consequence of
272 HIV's direct infection and destruction of megakaryocytes [34]. Amegor (27) reported that
273 thrombocytopenia increased as immunological incompetence of the participants worsens.
274 The presence of anti-platelet antibodies which leads to an increased destruction of platelets

275 has also been hypothesized [4]. However, the results of this study is not in line with the
276 findings of a study conducted in Kenya which recorded a decrease in thrombocytopenia from
277 20% to 6.5% after 6 months of ART [3, 35].

278 The predominant HAART regimen in this study was the TDF+3TC+EFV which is the WHO
279 recommended alternate first line HAART regimen in some resource limited areas [36, 37].
280 Majority, (72%) of the study population were on TDF+3TC+EFV, 13% were on
281 AZT+3TC+NVP (which is the standard WHO recommended first line HAART regimen in
282 resource limited settings), while 6.7% were on TDF+3TC+NVP which is also an alternate
283 first line regimen [36, 37]. TDF+3TC+LPV/r was the commonest second line HAART
284 regimen recorded in this study. This implies that 6.7% of the HIV infected individuals
285 experienced treatment failure and have switched to a second line therapy. Also, 0.7% of the
286 study population were administered with TDF/FTC/EFV, which is an alternate first line
287 HAART regimen in resource rich areas. According to the WHO guidelines on HAART issued
288 in 2003, the formulation of first line HAART includes the combination of AZT/3TC plus
289 stavudine (d4T) or NVP or EFV [37]. In 2006, this recommendation was revised to include
290 TDF or ABC as alternative first-line NRTIs while encouraging health care givers not to
291 include (d4T) base combinations **in-order** to minimize its possible accumulation in the
292 mitochondria leading to d4T associated complications [36]. Majority of the regimen in this
293 study included AZT, 3TC and either EFV or NVP or LPV/r. **(You have not discussed the**
294 **differences, whether significant or not, in the cell-counts in Table 4)**

295 Comparison of lymphocyte values in NVP and EFV groups showed higher counts in the NVP
296 group compared with the EFV group. These findings are consistent with findings of a Kenyan
297 study which recorded higher lymphocyte values in NVP group compared to the EFV group
298 [38]. The higher lymphocyte count signifies restoration of immunity following HAART
299 administration. Comparison of AZT and TDF, both administered with a 3TC/NVP shows that
300 RBC and HCT were both higher in the TDF group. AZT/3TC/NVP had better result on
301 erythroid cell indices compared to AZT/3TC/EFV. The worst decrease in both erythroid and
302 myeloid cell lines were recorded in the AZT/3TC/EFV. This demonstrated the compensatory
303 effect of NVP over AZT when administered together, and showed AZT's toxicity on
304 progenitor cells in the bone marrow stroma [23]. This was consistent with a study by Hema,
305 2011 who also found NVP more favourable to erythroid cell indices compared with AZT or
306 EFV.

307 The AZT/3TC/EFV group recorded the highest platelet large cell count (PLCC). This shows
308 increased variation in the sizes of thrombocytes. This corroborates with the findings of a
309 study conducted by Munyazesa (19) in sub-saharan Africa who found increased variation in
310 platelet sizes post HAART administration. From the findings of this study, a comparison of
311 EFV and NPV administered with AZT/3TC backbone showed that EFV improved platelet
312 indices as opposed to NVP which did not improve platelet indices. This may probably be the
313 reason why the 2013 WHO HIV treatment guidelines recommended EFV as the first line
314 choice of NNRTIs (<http://www.who.int/hiv/pub/guidelines/arv2013>). Thrombocytic indices;
315 PLT, PCT, MPV, PLCR, P-LCC and PDW-S were all higher in the EFV group compared with
316 the NVP group. The results of this study show that EFV has an ameliorative influence on
317 platelet indices compared to NVP.

318 There was no significant difference in the effect of AZT and TDF administered with 3TC/EFV
319 on complete blood count parameters. A comparison of AZT and TDF administered with
320 3TC/NVP shows a more positive influence on erythroid cell lines in the TDF group compared
321 with the AZT group. Again, in this study TDF/FTC/EFV which is the gold standard first line
322 regimen in resource rich area had the best effect on platelet indices [36].

323

324 **CONCLUSION AND RECOMMENDATION**

325

326 In conclusion, the administration of HAART is associated with significant improvements in
327 erythroid and lymphoid lines, which reduces anaemia, improves immunity and general

328 patient well-being. Cumulatively, NVP has a much more significant improvement on erythroid
329 cell indices while EFV significantly improves ameliorated platelet indices. As found in this
330 study, NVP favoured lymphoid lineage growth compared with EFV. TDF favoured RBC
331 indices compared with AZT and EFV improved platelet indices as opposed to NVP which did
332 not improve platelet indices. AZT/3TC/NVP, TDF/3TC/LPV/r, TDF/3TC/NVP should be
333 considered for improvement in erythroid cell indices. AZT/3TC/EFV and TDF/FTC/EFV
334 should be considered in **clients patients** with thrombocytopenia. This makes HAART a
335 suitable choice **over its taking into account side effects** in the management of
336 people living with HIV to prevent the development of AIDS especially when
337 started early. This study therefore provides **added basic** information to encourage
338 health workers to intensify the effectiveness of HAART campaign to get HIV
339 infected individuals to initiate HAART early. Also, ART should be chosen base on the
340 results of the initial laboratory tests conducted on the patient.

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342

343 **COMPETING INTERESTS**

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345

Authors have declared that no competing interests exist.

346
347

348 **Consent**

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350

Written informed consent was obtained from all participants for publication of this case
351 control study. A copy of the written consent is available for review by the Editorial office of
352 your journal.

353
354

355 **Ethical considerations**

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357

This study was approved by the Committee for Human Publication and Research Ethics of
358 the Kwame Nkrumah University of Science and Technology, Kumasi Ghana. Written
359 informed consent was obtained from all participants before recruiting them into the study.
360 Consent form was given to each participant to sign or thumb-print and confidentiality was
361 assured.

362
363

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476

477 **DEFINITIONS, ACRONYMS, ABBREVIATIONS**

478 Short term HAART users = Participants who have been on HAART for less than or equal to
479 52 months.

480

481 Medium term HAART users = Participants who have been on HAART for more than 52
482 months but less than or equal to 104 months.

483

484 Long term HAART users = Participants who have been on HAART for more than 104
485 months.

UNDER PEER REVIEW