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# Antivirt Antivirt

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#### 9 Abstract

<sup>10</sup> Both bacteria and protozoa require Folic acid for replication and Cotrimoxazole inhibits

<sup>11</sup> synthesis of the vitamin. For its mechanism of inhibiting Folic acid, the medicine has been in

<sup>12</sup> use as antibacterial drug for many decades but it is not being used to treat trypanosomosis

<sup>13</sup> (protozoan disease). To enhance anti-Folic acid activity of the medicine in order to improve its

<sup>14</sup> anti-trypanosome efficacy and make it function as new medicine for sleeping sickness (tropical

<sup>15</sup> disease of man and animals) it was stabilized with Antivirt® (Medicinal synthetic

<sup>16</sup> Aluminum-magnesium silicate). At 100

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#### 20 1 Introduction

rypanosomosis is debilitating and often fatal in both man and animals [1] [2]. The disease is found mainly in subSaharan Africa [3]. It has wide range of animal-hosts [4] [5] and is transmitted by Glossina spp, flies (Tabanids and Stomoxys) and Vampire bats. Dourine (caused by T. equiperdum) is sexually transmitted in Equines.

African trypanosomosis has been reported to be responsible for 55,000 human and 3 million livestock deaths, annually [6]. In Nigeria, Trypanosoma brucei and T. congolense are the most pathogenic species for domestic animals [7].

The disease is a major cause of mortality in animals in Nigeria and contributes greatly to underdevelopment of Sub-Saharan Africa and other socioeconomic consequences despite huge amounts being spent on research to control it [8]. World health organization [9] reported that over 60 million people and 48 million livestock in

30 Africa, are at risk of \trypanosomosis.

Of the different species of Trypanosomes which affect man and animals, Trypanosoma. congolense is strictly

<sup>33</sup> a parasite of the microcirculation, producing primary lesions in blood vessels and lymph nodes [10] [11] [12].

 $_{\rm 34}$   $\,$  Trypanosoma brucei is found in connective tissues, producing inflammation, cellular degeneration and necrosis

which lead to tissue and organ damage ??4] [12].

42 animals at risk [14].

Index terms— cotrimoxazole; folic acid-inhibition; anti-trypanosomes efficacy; antivirt®; sleeping sickness,
 new medicine.

<sup>31</sup> Estimated losses due to trypanosomosis in Africa run into billions of Dollars.

Control measures for trypanosomosis are either by controlling the vectors or by use of chemotherapy or a combination of both. In poor, rural communities, affected by the disease, control is mainly by use of trypanocidal drugs [13]. Drugs currently employed in treatment of trypanosomosis are: Homidium salts (Ethidium-Novidium®:); Quinapyramine sulfate (Antrycide®:); Diminazene aceturate (Berenil®:); Isometamidium (Samorin-Trypamidium®:) and Suramin sodium. These drugs have been in use for more than half a century now [14]. It is estimated that 35 million doses of the drugs are used in Africa each year, with about 50-70 million

Microorganisms exposed to drugs over such a long time usually develop resistance to the drugs. Mechanisms 43 for drug-resistance include, loss of surface specific receptors or transporters for the drugs, increased metabolism 44 of the drugs and alteration (by mutation) of specific targets for the drugs on the organisms. These result in 45 resistance to a small number of related drugs, too. More often, cells express mechanisms of resistance that confer 46 simultaneous resistance to many different structurally and functionally unrelated drugs [15]. For the problem 47 of drug-resistance by pathogens, there is need for constant search for new drugs for treatment of important 48 diseases such as trypanosomosis. Cotrimoxazole is a combination of five parts of sulfamethoxazole and one part 49 of trimethoprim based on dose of each of the two drugs. It is being used for treatment of bacterial, fungal and 50 protozoan infections [16]. The drug-formulation was introduced into clinical use in the late 1960s and it is for 51 treatment of urinary tract infections, respiratory infections, sexually transmitted diseases, enteric infections and 52 typhoid fever. Advantages of Cotrimoxazole include combination of the two components. The drug inhibits 53 synthesis of tetrahydrofolic acid which is needed as a cofactor in synthesis of thymidine and purines which are 54 components of bacterial DNA [17] [18] [19]. Sulfamethoxazole inhibits synthesis of the intermediary dihydrofolic 55 acid from its precursors [20] while Trimethoprim competitively inhibits dihydrofolate reductase and consequently, 56 production of tetrahydrofolic acid from dihydrofolic acid [21]. Potentiating sulfamethoxazole with trimethopriim 57 reduces toxicity and microbial resistance [22]. The synergy between trimethoprim and sulfamethoxazole was first 58 59 described in the late 1960s [23]. Trimethoprim and sulfamethoxazole have a greater effect when given together 60 than when given separately, because they inhibit successive stages in the folate synthesis pathway. They are 61 formulated in a ratio of one-to-five so that when they enter the body their concentrations in the blood and tissues are exact ratio required for a peak synergistic effect between the two [24]. Trimethoprim causes a backlog of 62 dihydrofolate and this backlog can work against inhibitory effect the drug has on tetrahydrofolate biosynthesis. 63 This is where the sulfamethoxazole comes in. Its role is in depleting the excess dihydrofolate by preventing it 64 from being synthesized in the first place. 65 Trimethoprim-sulfamethoxazole (Cotrimoxazole) has proved effective in the treatment of infections of coccidian 66 protozoa parasites, Isospora and Cyclospora [25] [26]. So, by potentiating its anti-folic acid effects its protozoan 67 efficacy may be enhanced to achieve cure for trypanosmosis. 68 Molecules of Aluminum magnesium silicate (AMS: clay) are 0.96 nanometer thick and some hundred 69

Nanometers across [27] [28] [29]. As a Nanomedicine, AMS helps in delivering drug-molecules to target cells. 70 Drug molecules in "corridors" of AMS "house of cards" are also bound by charged faces and edges of its platelets. 71 72 So, they are protected from degradation by both physical and physiological factors but are released gradually 73 into blood of treated patients. Also, silicates are immune stimulants [30] and AMS is a stabilizing/potentiating agent [31] [32]. By stabilizing drugs, AMS increases potency of the drugs [33]. When drugs are potentiated their 74 doses required for desired effects reduce and using lower doses for treatments reduces side effects of drugs so 75 that immune responses of patients improve. With enhanced efficacy of drugs and improved immune responses 76 of patients both sensitive infections and drug-resistant infections could be effectively treated. So, using The 77 Medicinal synthetic Aluminum-magnesium silicate (Antivirt®) to stabilize Cotrimoxazole may enhance its efficacy 78 against trypanosomes enough so that the medicine being commonly used for treatment of bacterial diseases and 79 amebiasis (protozoan disease) can also function as a medicine for trypanosomosis (sleeping sickness). 80

#### <sup>81</sup> **2 II.**

#### <sup>82</sup> 3 Materials and Methods

Twenty-five mice were assigned to five (5) groups of five (5) each, as follows: Blood samples were collected from each of the mice and examined daily until parasitemia was established in all infected groups. Treatment was started 7 days post-infection and lasted for 5 days while assessment of parasitemia was on day-2 posttreatment.Group 1: Infected/Untreated

#### <sup>87</sup> 4 a) Data analysis

The parasitemia and total WBC were presented as means  $\pm$  SEM and analyzed for statistical differences by one way analysis of variance and the significant differences were accepted at the level of P? 0.05.

#### 90 **5** III.

#### 91 6 Results

Parasitemia was observed in all the groups from four days post infection and it increased steadily until treatment was commenced by day-7 post infection. Zero mean parasitaemia  $(0.00\pm00)$  of the group of trypanosome-infected

mice treated at 100 % dose of Cotrimoxazole with Cotrimoxazole-Antivirt® drug formulation was significantly

(P?0.05) less than  $5.87\pm0.43$  of the group treated with 100 % dose of Cotrimoxazole with Cotrimoxazole alone.

Both the  $0.00\pm00$  parasitaemia of the 100 % -dose of Cotromoxazole in Antivirt® group and  $5.87\pm0.43$  of the

97 100 % -dose of Cotrimoxazole-group were significantly (P<0.05) lower than 11.73±0.86 of the 75 % -dose of

- 98 Cotrimoxazole-group and 11.30±1.01 of the 75 % -dose of Cotrimoxazole in Antivirt®-group but there was no
- $_{99}$  significant difference (P>0.05) in mean parasitemia (11.73 $\pm$ 0.86) of the group of 75 % -dose of Cotrimoxazole and

11.30±1.01 of the 75 %-dose of Cotrimoxazole in Antivirt®-group. Mean parasitemia of the trypanosome-infected
 groups of mice treated with Cotrimoxazole and Cotrimoxazole-Antivirt® are as shown on Table 1.

Mean WBC was highest in the group infected and treated with 100 %-dose of Cotrimoxazole in Antivirt® when compared with the other infected groups. The mean WBC of the group of infected/treated with 100 %-dose of Cotrimoxazole was comparable to that of the group of infected/untreated, that of the group of infected/treated with 75 %-dose of Cotrimoxazole and that of the group of infected/treated with 75 %-dose of Cotrimoxazole in Antivirt® (Table 2).

#### 107 7 Discussion

Significant (P? 0.05) reduction of parasitemia from  $12.76\pm1.20$  to  $5.87\pm0.43$  (54 % infection reduction) in trypanosome-infected mice treated with 100 % antibacterial-dose of Cotrimoxazole is evidence that the drug has anti-trypa mosomal effect. Cotrimoxazole is known to inhibit synthesis of Folic acid and trypanosomes need the vitamin for replication. However, 54 % infection clearance is much lower than the 95 % required for treatments to be effective. This failure to achieve enough level of clearance to terminate trypanosome infections may be reason Cotrimoxazole is not yet being recommended for treatment of trypanosomosis.

<sup>114</sup> When the drug was stabilized with Antivirt<sup>®</sup>, the 100 %-dose completely and significantly (P?0.05) cleared <sup>115</sup> the parasitemia ( $00.00\pm0.00$ ). AMS is a stabilizing agent and a Nanomedicine. As stabilizing agent it prolongs <sup>116</sup> time medicines remain at high concentration in blood of treated animals while as a Nanomedicine, it enhances <sup>117</sup> delivery of drugs to targets. Both prolongation of time of high bioavailability of drugs and delivering them to <sup>118</sup> effect-targets enhance efficacy of drugs. So, the Antivirt<sup>®</sup> may have enhanced ability of Cotrimoxazole to inhibit <sup>119</sup> synthesis of Folic acid and so terminated the trypanosome infections.

As a silicate AMS also enhances immune response. So, the increase in WBC count of the mice suggests that the Antivirt® may, in addition to improving efficacy of Cotrimoxazole, have enhanced immune response of the mice. Synergy of the enhanced immune response of patients and enhanced efficacy of medicines may be responsible for the zero parasitemia achieved.

In earlier studies of effects of the Antivirt® on antimicrobial drugs, it made 75 %-dose more effective than 124 recommended (100%) doses of drugs but in this study its effects on 100 %-dose of Cotrimoxazole were the best. 125 Dose used as 100 % in this study is the dose recommended for treatment of bacterial infections (not 126 for trypanosomosis). That the 100 %-dose of Cotrimoxazole without the Antivirt® was able to reduce the 127 trypanosome infection, significantly, suggests that if the dose is increased it may lead to cure of the infection even 128 without the Antivirt<sup>®</sup>. Failure to determine correct dose of Cotrimoxazole for treatment of trypanosomosis may 129 be reason the drug has not been recommended for treatment of the zoonotic/tropical disease. That dose, used 130 as 100 % may be 75 % of dose of the drug needed for treatment of trypanosomosis in absence of the Antivirt<sup>®</sup>. 131 However, reducing doses by stabilizing drugs with the Antivirt® reduces side effects to enhance immune responses 132 133 and also reduces cost of drug formulations.

Trypanosomosis is a very serious disease of both man and animals in the tropics and the causative agents, very often, develop resistance against existing drugs. So, there is need to constantly research for new drugs. Since

the Antivirt® made Cotrimoxazole achieve total clearance of the trypanosome-infections it may cure the disease

137 and may also make development of resistance against the new therapy difficult by achieving total clearance of infections.

#### 1

| Cotrimoxazole in Antivirt® |                    |                     |                          |                   |                  |  |  |  |  |
|----------------------------|--------------------|---------------------|--------------------------|-------------------|------------------|--|--|--|--|
| S/N                        | Infected/Untreated |                     | Infected/Treated         | Infected/Treated  |                  |  |  |  |  |
|                            |                    | Cotri               |                          | Cotri-            |                  |  |  |  |  |
|                            |                    |                     |                          | MSAMS             |                  |  |  |  |  |
|                            |                    | 100%                | 75%                      | 100%              | 75%              |  |  |  |  |
| 1                          | 15.85              | 6.31                | 12.59                    | 0.00              | 12.59            |  |  |  |  |
| 2                          | 12.59              | 5.01                | 10.00                    | 0.00              | 10.00            |  |  |  |  |
| 3                          | 12.59              | 6.30                | 12.59                    | 0.00              | 12.59            |  |  |  |  |
| 4                          | 10.00              |                     | 10.00                    |                   | 10.00            |  |  |  |  |
| MEAN <b>#2576</b> ±1.20 c  |                    | $5.87 {\pm} 0.43$ b | $11.73{\pm}0.86~{\rm c}$ | $0.00{\pm}0.00$ a | $11.30{\pm}1.01$ |  |  |  |  |
|                            |                    |                     |                          |                   | с                |  |  |  |  |

Figure 1: Table 1 :

#### $\mathbf{2}$

|                           |                    | Cotrimoxazole in          | Antivirt®         |                  |            |
|---------------------------|--------------------|---------------------------|-------------------|------------------|------------|
| S/N                       | Infected/Untreated |                           |                   | Infected/Treated |            |
|                           |                    | Cotri                     |                   | Cotri-MSAMS      |            |
|                           |                    | 100%                      | 75%               | 100%             | 75%        |
| 1                         | 1.50               | 1.30                      | 1.50              | 2.90             | 1.28       |
| 2                         | 1.39               | 1.56                      | 1.77              | 2.78             | 1.56       |
| 3                         | 1.94               | 2.01                      | 1.89              | 2.89             | 1.39       |
| 4                         | 1.17               |                           | 10.00             |                  | 1.00       |
| MEAN±SE±0.16 b 1.62±0.21b |                    | $1.62{\pm}0.21\mathrm{b}$ | $1.72 {\pm} 0.11$ | $2.86{\pm}0.38a$ | $1.31\pm0$ |
|                           |                    |                           | ab                |                  |            |

IV.

Figure 2: Table 2 :

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