Artemisia Annua L. Infusion Consumed Once a Week Reduces Risk of Multiple Episodes of Malaria: A Randomised Trial in a Ugandan Community

Patrick E Ogwang¹,³, Jasper O Ogwal⁴, Simon Kasasa², Deogratius Olila³, Francis Ejobi³, David Kabasa³ and Celestino Obua⁴*

¹Natural Chemotherapeutics Research Institute, Ministry of Health, P.O Box 4864. ²School of Public Health, College of Health Sciences, Makerere University, PO Box 7072, Kampala. ³Faculty of Veterinary Medicine, Makerere University, PO Box 7072, Kampala; ⁴Department of Pharmacology and Therapeutics, School of Biomedical Sciences, College of Health Sciences, Makerere University, PO Box 7072, Kampala, Uganda

Abstract

Purpose: To evaluate the protective effect of Artemisia annua infusion against malaria in a community that uses it as herbal ‘tea’ for malaria prevention.

Methods: 132 flower farm workers who met the study inclusion criteria and were not yet using A. annua infusion were randomized either to A. annua or placebo groups in the ratio of 1:1. Treatments were administered once a week under direct observation to participants. Malaria episodes were documented over a 9-month period while adverse effects were documented over 12 months.

Results: A. annua herbal ‘tea’ significantly reduced the risk of suffering more than one episode of malaria in nine months by 55 % (12/67 vs 26/65, p = 0.005, No participant experienced any serious adverse effect although bitter taste was the most common side effect of the infusion.

Conclusion: Artemisia annua infusion consumed once a week was effective in preventing multiple episodes of malaria in humans living in malaria endemic areas. However, its bitter taste and the risk of development of malaria parasite resistance to the artemisinin contained in it remain major challenges for its use in the mass control of malaria.

Keywords: Artemisia annua, Herbal tea, Community, Malaria prevention

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*Corresponding author: Email: cobua@chs.mak.ac.ug; Tel: +256 712 210 937
INTRODUCTION

Artemisia annua is a medicinal plant used in traditional Chinese medicine in the treatment of febrile fevers, including malaria [1]. In 1972, artemisinin was isolated from A. annua as its most active antimalarial component [2]. Artemisinin and its synthetic derivatives form the major component of Artemisinin Combination Therapies (ACTs) which are the current WHO recommended first-line drugs for the treatment of uncomplicated malaria caused by Plasmodium falciparum [3-5]. Artemisinin and its metabolite, dihydroartemisinin, are highly active against malaria parasites, but they have very short plasma half-life [6]. The short half-life makes them unsuitable for malaria prevention and are probably less prone to the development of resistance against them by the parasite.

In Uganda, A. annua is being used by some communities as a herbal tea, taken once a week for malaria prevention and for other illnesses such as HIV/AIDS [7]. The effectiveness and safety of the tea when used in malaria prevention remain unknown. This study was therefore, designed to evaluate the protective effect of A. annua against malaria in a community that uses it as a ‘tea’ for malaria prevention, and to predict its mode of action in the claimed prophylaxis.

EXPERIMENTAL

Collection and processing of Artemisia annua and Thea sinensis dry leaf powders

Artemisia annua used by the farm workers in the first 8 months of the study was sourced from the farm garden and when it was out of stock by the 9th month, the farm purchased it from a commercial source. Artemisinin content in A. annua dry leaves from the farm ranged from 0.4 - 0.5 % while for the commercial source, it was 0.8 %. The concentration of artemisinin in the infusion made from farm-derived A. annua was 0.055 mg/ml while for the commercial one, it was 0.100 mg/ml. Thea Sinensis (tea leaves) powder used as placebo in the study was purchased from the local market near the study site. A. annua from the farm garden was collected in February 2009 and identified by M Nusula of Natural Chemotherapeutics Research Institute. A voucher specimen NCJ 257 was deposited at the herbarium of Natural Chemotherapeutics Research Institute, Ministry of Health, Uganda.

Study site

The study was conducted in Wagagai flower farm located on the shores of Lake Victoria in Entebbe, about 35 km south of Kampala, the capital city of Uganda. Wagagai flower farm was founded in 1998 by Dutch investors and it exports flowers to Europe. A. annua use for malaria prevention was initiated by the farm management in 2006 with about 80 workers and by the time of this study about 600 of 1500 workers were using it (Wagagai farm records, unpublished). The flower farm has a well-equipped health centre that serves the farm workers and neighbouring community. The outpatient clinic records indicated that malaria accounted for 30 % of outpatient hospital attendance, which falls within national figures of 25 – 40 % reported by the Ministry of Health [8]. However, by the time this study was initiated among the farm workers, malaria accounted for between 16 - 20 % of outpatient attendance (Wagagai clinic record, 2009 unpublished).

Standardisation of Artemisia annua ‘tea’

We standardised A. annua infusion used for malaria prophylaxis based on the practice in the farm. Standardised A. annua infusion was equivalent to 20 mg/ml (dry leaf powder) prepared by adding boiling water to A. annua powder, stirred and allowed to infuse for 30 min, sieved and served immediately. In our study, honey was added to improve taste. The content of artemisinin in leaf powder and in the infusion of A. annua were determined at three time points - April, June and December 2009 using a standard high
pressure thin layer chromatography method [9].

In administering the tea, each participant in the A. annua group received a dose equivalent of 250 ml infusion (containing 5 g dry powder), which was consumed at once every Wednesday under direct observation by the study nurse. The control group received equivalent volume of T. Sinensis which also had equivalent volume of honey as that used in A. annua infusion.

**Randomisation and blinding**

The volunteers with no malaria parasites in the blood smears (Giemsa technique) who met eligibility (inclusion) criteria were randomised in the ratio of 1:1 to either Artemisia or placebo groups previously assigned as odd and even, respectively. A participant picked, without replacing, from a box a number written on folded paper to determine his or her group (see Table 1 and Figure 1).

**Sample size determination**

The study was designed to have a statistical power of 80 % and to detect a 60 % difference at a significant level of 0.05 based on the national outpatient malaria incidence of 40 %. The design required ninety eight (98) participants randomised to either A. annua or placebo groups in the ratio of 1:1 [10]. We recruited all the 132 participants who turned up after screening in order to take care of the anticipated dropout rate due to high labour turnover in this farm and also lower malaria burden among the farm workers than had been used in sample size determination (Wagagai Farm records, unpublished).

**Study subjects screening**

One hundred and seventy one (171) newly recruited adult farm workers who were not yet on A. annua prophylaxis registered to participate and were screened for the study. Eligibility criteria for a participant were: history of at least 2 episodes of malaria in a year, no use of malaria prevention medication or drugs or herbs before or during the study period other than those in the study, passed physical examination, Giemsa stain blood smears negative for malaria parasites, normal organ functions including renal, hepatic and haematological and human chorionic gonadotropin (pregnancy) test negative for females.

**Monitoring of study participants**

Study participants were monitored for episodes of malaria over nine months (April to December 2009) instead of planned 12 months because from month 8 to 9, no cases of malaria occurred in A. annua group. Giemsa stain (thin and thick) microscopy technique [11] was used to detect malaria parasites in blood while polymerase chain reaction (Gene AMP, PCR system model 9700, Applied Biosystems) was used for speciation and quality control of microscopy. Malaria episode was defined as presence of signs and symptoms of malaria (fever, headache, joint pain or bitter taste) plus blood smear with > 0 parasites. The number of participants with no clinical malaria but carrying parasites in their blood was determined after months 3 and 6 of prophylaxis. Parasitaemia was checked independently by two experienced laboratory technicians and where variation was observed a third laboratory technician examined the slides. Occurrence of serious adverse effects was monitored over 12 months through weekly self-report by the participants and monthly clinical examination by the study physician in addition to the biochemical and haematological tests. The clinical signs looked for include: skin rash, jaundice, changes in blood pressure and heart rate, and were documented for each participant. Female participants who missed their menses had pregnancy test done and those found positive were counseled and excluded from the study. Study participants
Figure 1: Participants’ recruitment, enrolment and follow up chart

Table 1: Study participants’ demographic/baseline data

| Descriptive statistics | Artemisia (n= 67) | Control (n=65) | P-value |
|------------------------|------------------|----------------|---------|
| Sex:                   |                  |                |         |
| Female                 | 38 (56.7%)       | 39 (58.3%)     | 0.702   |
| Male                   | 29 (43.3%)       | 26 (41.7%)     |         |
| Marital status:        |                  |                |         |
| Married                | 25 (37.3%)       | 28(43.1%)      | 0.561   |
| Single                 | 32 (47.8%)       | 31 (47.7%)     |         |
| Not stated             | 10 (14.9%)       | 6 (9.2%)       |         |
| Bed net use:           |                  |                |         |
| Yes                    | 24 (35.8%)       | 12 (18.5%)     | 0.025   |
| No                     | 43 (64.2%)       | 53 (81.5%)     |         |
| Mean age (years)       | 26.6 ± 6.4       | 25 ± 4.3       | 0.100   |
| Mean temperature (°C)  | 36.7± 0.5        | 36.7 ± 0.5     | 0.78    |
| Average weight (kgs)   | 56.3± 6.4        | 54.7± 6.1      | 0.15    |
| Haemoglobin (g/dl)     | 14.2±1.8         | 14.6±1.6       | 0.35    |
| White Blood cell x 1000/mm³ | 5.8±2.0 | 5.7±2.0       | 0.93    |
| Creatinin level (ummol/L) | 99.4±15.5 | 95.5±17.0       | 0.15    |
| Alanine aminotransferase (IU/L) | 23.02±12.5 | 25.0±13.8 | 0.26 |

With the exception of bednets, the randomisation ensured equal distribution of participants’ characteristics between the groups. The effect of bednets on the incidence of malaria was accounted for in the analysis.
Table 2: Distribution of clinical malaria between Artemisia and placebo groups over a 9-month period

| Episodes | Artemisia % (n) | 95% CI | Placebo % (n) | 95% CI | Z   | P    |
|----------|----------------|--------|---------------|--------|-----|------|
| 0        | 53.7 (36)      | 42.1, 65.9 | 35 (23)       | 23.4, 46.6 | 2.2 | 0.028*|
| 1        | 28.4 (19)      | 17.2, 38.8 | 25% (16)      | 14.5, 35.5 | 0.4 | 0.696|
| 2        | 13.4 (9)       | 5.0, 21.0 | 28 (18)       | 17.1, 38.9 | -2.14 | 0.032*|
| 3        | 3.0 (2)        | -1.1, 7.1 | 11 (7)        | 3.4, 18.6 | -1.81 | 0.071|
| 4        | 1.5 (1)        | -1.3, 5.3 | 1.5 (1)       | -1.4, 5.5 | 0.00 | 1.00 |
| >1       | 17.9 (12)      | 8.8, 27.2 | 40 (26)       | 28.1, 51.9 | -2.79 | 0.005*|

*Artemisia annua infusion had a significant protective effect against malaria especially in reducing multiple episodes of malaria.

A signed written consent was obtained from each study participant and filed appropriately. Written permission to conduct study in Wagagai Flower Farm and publish the findings was granted by the farm's management. The study was approved by the Makerere University Institutional Review Board and permission to conduct study was granted by the Uganda National Council for Science and Technology (UNCST) under protocol registration number HS 528.

Data management and analysis

All data were captured into participants' case report forms and laboratory forms at the study sites. Participants’ files were then transported at the end of data collection to the central office for data entry and further management. Double data entries were done in Epidata (version 3.1), with pre-designed check programs for quality control, and in Microsoft Excel 2007 as backup. The data were then exported to Stata, version 9.0, for cleaning and analysis. The risk of first and multiple episodes of malaria were determined by intention-to-treat (ITT) approach using a two sample proportion analysis in Stata. Malaria incidence rate reduction (protection efficacy) was determined by per protocol using Poisson regression model for rates computed in person-months. Haematological and biochemical data was compared using paired Student’s t-test and only significant data are reported. Study participants’ demographic and baseline data were compared between the groups using Chi-square and paired Student’s t-tests. All data were analysed at 95 % confidence interval and a difference was considered statistically significant if $p < 0.05$.

RESULTS

Effect of A. annua tea on clinical and subclinical malaria

A total of 124 episodes of malaria were recorded over a period of nine months with 88.8 % being infections caused by *Plasmodium falciparum*. The effect of *A. annua* infusion on malaria episodes is shown in Table 2.

Malaria incidence rate, protection efficacy of *A. annua* ‘tea’ and effect of mosquito bednets

The average follow-up time for malaria incidence rate was 7.4 ± 2.4 months for the Artemisia group and 6.9±3.0 months for the placebo. Malaria incidence rate was 10 %, (95% CI 7.5, 13.5) person months and 16 % (95% CI 12.5, 19.5) person months for Artemisia and placebo groups, respectively giving protection efficacy of 37.5 % ($p = 0.015$). Comparison of malaria incidence rate reduction in participants with no mosquito bed nets showed a similar protective effect of 38.4 %, $p = 0.03$. bednets use had a non significant effect of 26.7 %, ($p = 0.41$) in participants who used both *A. annua* and bed nets. The malaria protective effect of the *A. annua* infusion showed increasing trend in Chi square test for trend ($p = 0.003$) with no cases of clinical malaria occurring in the
Figure 2: Monthly incidence of malaria in Artemisia (dashed line) and placebo (unbroken line) groups

Artemisia group by the Month 8 of prophylaxis (Figure 2).

Effect of A. annua tea on parasite carriage and white blood cells

Malaria parasites (sub-clinical malaria) in participants in A. annua group were reduced by more than 60% by months 3 and 6 of prophylaxis with the infusion. These participants also had higher white blood cell count than those on placebo (see Table 3).

Table 3: White blood cells levels in participants on A. annua and control groups

| Blood sampling time | Diagnostic marker | Artemisia group Mean ±SD (n) | Control group Mean ±SD (n) | Difference (t value) | p value |
|---------------------|-------------------|-----------------------------|----------------------------|----------------------|---------|
| At baseline         | T.WBC             | 5.76±1.94 (47)              | 5.72±2.02 (49)             | 0.0899               | 0.9286  |
|                     | Lymphocytes       | 2.45±0.96 (47)              | 1.95±0.83 (49)             | 1.65                 | 0.1023  |
|                     | Monocytes         | 0.67±6.8 (47)               | 0.70±5.0 (49)              | -0.20                | 0.844   |
|                     | Granulocytes      | 2.84±1.21(47)               | 3.29±1.43 (49)             | 1.63                 | 0.1306  |
| At 3 months         | T.WBC             | 5.80±1.77 (33)              | 6.05±2.36 (44)             | -0.52                | 0.606   |
|                     | Lymphocytes       | 2.37±0.74(33)               | 2.35±1.08(44)              | 0.084                | 0.933   |
|                     | Monocytes         | 0.54±0.47(33)               | 0.72±0.59(44)              | 1.47                 | 0.145   |
|                     | Granulocytes      | 2.89±1.54(33)               | 3.00±1.19(44)              | -0.34                | 0.737   |
| At 6 months         | T.WBC             | 5.26±1.51 (39)              | 4.71±1.47(43)              | 1.71                 | 0.046*  |
|                     | Lymphocytes       | 2.14±0.72(39)               | 1.98±0.58(43)              | 1.13                 | 0.135   |
|                     | Monocytes         | 0.49±0.29(39)               | 0.39±0.18(43)              | 1.91                 | 0.029*  |
|                     | Granulocytes      | 2.66±1.03 (39)              | 2.34±1.04(43)              | 1.39                 | 0.084   |
| At 12 months        | T.WBC             | 6.00±1.75 (36)              | 5.21±1.46(39)              | 2.12                 | 0.0187* |
|                     | Lymphocytes       | 2.67±0.89(36)               | 2.27±0.74(39)              | 2.14                 | 0.0179* |
|                     | Monocytes         | 0.93±0.42(36)               | 0.78±0.29(39)              | 1.85                 | 0.034*  |
|                     | Granulocytes      | 2.41±1.30(36)               | 2.17±0.92(39)              | 0.94                 | 0.174   |

Adverse effects of A. annua tea

Although bitter taste was a major side effect, none of the participants on A.annua experienced any serious adverse clinical or systemic effects associated with it. One participant on A. annua infusion, however, presented with severe jaundice after 4 weeks of prophylaxis. Laboratory investigation revealed hepatitis B as the cause of the jaundice. The participant was thereafter excluded from the study and treated; the participant recovered in six weeks.

*Once a week dose of A.annua significantly elevated the levels of monocytes by the 6th months of prophylaxis
DISCUSSION

In this study, by using a randomised trial design, we established that *A. annua* herbal tea (leaf infusion), equivalent to 5 g in 250 ml (20 mg/ml), taken once a week at Wagagai Flower Farm in Uganda has a protective effect against malaria. It is especially effective in reducing the risk of having multiple episodes of malaria in an individual living in a malaria-endemic setting. In Uganda, it is estimated that on the average, an individual suffers between 3 and 6 episodes of malaria in a year if he/she is not using insecticide-treated nets [8]. In this study, 17.9% of the participants using *A. annua* suffered more than one episode of malaria compared to 40% in the placebo group and also had statistically significant reduction in malaria incident rate when adjusted and not adjusted for bednet use.

The use of *A. annua* in this farm benefited not only the farm workers through reduced incidence of malaria, reduced hospital visits and expenditure on treatment but also the farm management of the farm through increased staff productivity. Although *A. annua* can grow well in most parts of Uganda and is effective in preventing the risk of multiple episodes of malaria, the infusion used in this study community however had a bitter taste and also contained artemisinin. These two elements are a major limitation to mass use in Uganda and other communities. The use of *A. annua* infusion containing artemisinin heightens the risk of development of malaria parasites resistant to artemisinin and related drugs.

The low levels of artemisinin in the infusion, however, indicate the unlikely role of artemisinin in the observed protective effect against malaria. It is probable that the antimalarial and immune-modulating group of flavonoids, previously reported to be present in *A. Annua*, may be responsible for the prophylactic effect of the infusion. This is buttressed by the fact that participants took the treatment once a week, and usually within 24 h of oral administration of artemisinin, almost all of it and its metabolite (dihydroartemisinin) are eliminated from the body [6].

Previous *in vitro* studies of *A. annua* flavonoids, especially casticin, cirsilloineol, chrysoplenol D, chrysopleninetin and artemetin, have shown that these compounds also have powerful immunomodulatory activities [13-14,15]. In this study, participants who took *A. annua* infusion demonstrated both lower risk of malaria attack and higher levels of monocyte blood cells than the placebo group. Monocytes, have been documented to phagocytize human erythrocytes infected by *Plasmodium falciparum* malaria parasites at a rate of 7.5 and 2.9 per monocyte for erythrocytes containing young and mature parasites, respectively, compared to 0.8 per monocyte for non-infected erythrocytes [16]. Monocytes have also been shown to play an important role in blood stage malaria in mice [17]. The increase in monocyte count was not transient as levels remained significant from the 6th month of prophylaxis even when no cases of malaria occurred in the *A. annua* group from the 8th month. The effect of *A. annua* flavonoids as both antimalarial and immune modulator may, therefore, explain the observed prophylactic effect of *A. annua* against malaria; however, this can only be proved in a study that uses *A. annua* tea that does not contain artemisinin. Since immune suppression and malnutrition have been documented as major risk factors for malaria, cancers and other diseases in sub-Saharan Africa, this study reveals the potential use of *A. annua* for the control of malaria and other burdensome diseases in Africa.[16,18-20]. Combining *A. annua* flavonoids with locally available nutritional products may provide hope for affordable and effective means of combating malaria.

CONCLUSION

*A. annua* infusion consumed once a week as a herbal tea is effective in preventing multiple
episodes of malaria attacks and its mode of action may involve stimulation of white blood cells production especially monocytes.

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