Antifertility effect of aqueous and ethanol extracts of the leaves and roots of *Asparagus africanus* in rats

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Abstract

**Background:** Asparagus africanus is claimed to have use in reproductive related health problems in some areas of Ethiopia.

**Objective:** To study the potential antifertility effect of the aqueous and ethanol extracts of the leaves and roots of *Asparagus africanus* in rats.

**Methods:** Water and ethanol extracts were given by gavages to rats in the in vivo test at a dose of 300 mg/kg of body weight, and rat uterine tissue were used for the in vitro test at different concentrations.

**Results:** The aqueous extracts of the leaves and the roots showed an anti-implantation activity of 70% and 77%, respectively, while the ethanol extracts of the leaves and roots showed 48% and 61%, respectively. The antifertility activities of the aqueous and ethanol extracts were 40% (for leaves), 60% (for roots) and 20% (for leaves), 40% (for roots), respectively. All the extracts have resulted in significant (P< 0.05) reduction in the number of implants as compared with their respective controls. Each extract potentiated acetylcholine induced uterine contractions in a concentration dependent manner significantly (P< 0.05).

**Conclusion:** The results obtained in this study suggest that the leaves and roots of this plant may possess hormonal properties that can modulate the reproductive function of the experimental rats.

**Key words:** Antifertility, Anti-implantation, *Asparagus africanus*, aqueous extract, ethanol extract, leaves, roots, rats

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Introduction

*Asparagus africanus* (L) is one among the many plants with traditionally claimed antifertility properties in Ethiopia. This plant belongs to the Family Liliaceae. It is a perennial climbing or erect shrub that can grow between 700 and 3800 m above sea level. However, it is widely distributed and suitably grows higher up to 6m at the altitude range of 1450 and 2900 m. Its vernacular name is “Saritti”; this plant has been known for a long time to serve as a traditional cure for problems related to reproductive health. An infusion of its roots is employed as a remedy for venereal diseases. Moreover, at parturition the infusion of the root being mixed with water is taken by women to facilitate the process2, 3. In rural areas of Bale zone (southern parts of Ethiopia) the leaf or root of *Asparagus spp.* is traditionally chewed to facilitate childbirth4. Steroidal saponins, the most probable component of estrogen, were isolated from its roots5. The steroidal saponins, which were isolated from the roots of its related species of *A. officinalis*, were reported to have uterine contractile property6. The objective of this study is, therefore, to investigate the possible antifertility effect of the water and ethanol extracts of the leaves and roots of *A. africanaus* to substantiate its use as herbal medicine.

Materials and methods

**Plant collection and extract preparation**

The leaves and roots of *Asparagus africanus* were collected around Meki town about 130 km and the shore of Lake Langano about 210 km from Addis Ababa, the capital city of Ethiopia in May 2001. A botanist identified the plant and sample specimen was kept in the National Herbarium of Addis Ababa University under voucher number 1/2001. The leaves and roots were chopped with knife, and made dry in shade independently and then ground into powder with mortal and pestle. Measured amounts of the powder were macerated in distilled water (1 g / 6 ml, w/v) and in 90% ethanol (1 g / 4 ml, w/v). The macerated extracts were stirred with magnetic stirrer for 24 hrs at room temperature. Each extract was then filtered using cotton and Watman filter paper No 1. After filtration, the water extracts were lyophilized with a lyophilizer; while ethanol was evaporated from the ethanol extracts using Rota vapour at 40 °C. The resulting partially solid extracts were stored at -20 °C until used.
Experimental animals

Male and female albino rats were obtained from Faculty of Medicine, Addis Ababa University and let to breed continuously in the animal house of the Department of Biology, Addis Ababa University by allowing to be paired for any length of time as needed for mating. They were kept in cages in animal house with a 12:12 hr light-dark cycle. All the conditions were the same to all animals. They all made to feed on pellet diet and water ad libitum. The newly born female rats were separated from males at the age of four weeks in order to prevent uncontrolled mating. Virgin female rats 10 to 11 weeks of age weighing between 195 and 200 g were used for both in vivo and in vitro experiments.

In vivo test

There were four experimental groups, i.e., aqueous extract of the leaf (LW), aqueous extract of the root (RW), ethanol extract of the leaf (LE) and ethanol extract of the root (RE). There was also one control group for the water and ethanol extracts. Each rat both in the experimental and control groups was kept singly in a cage to acclimatize for a week 9. After a week of acclimatization experimental groups were administered with 300mg/kg body weight of the extract by gavages, while the control groups received same volume of the vehicles. Each extract and the vehicle were dissolved in a 0.5 ml solvent. Distilled water was used to dissolve the aqueous extracts, while 70% ethanol and distilled water in the ratio of 2: 3 were used to dissolve the ethanol extracts. While dosing continued, males of proven breeding ability were introduced into each cage on the ninth day according to the method described by Williamson et al. 9. Vaginal lavages were used to determine the presence of sperm every morning. The animals were separated immediately after confirming mating. The oestrous cycle was monitored and the animals were still cycling after the nine days of dosing. By the method described by Williamson et al. 9 laparatomy was performed on day 20 of pregnancy. The animals were sacrificed, and the number of implantation sites was determined. Weight gained by each rat of all the groups was recorded.

Anti-implantation and antifertility activities were expressed as percentages using the following formula (9):

\[
\text{Antifertility activity} = \frac{\text{No. of animals showing no implantation}}{\text{Total No. of animals}} \times 100
\]

Anti-implantation activity =

\[
\frac{\text{No. implants in control} - \text{No of implants in test group}}{\text{No. of implants in control group}} \times 100
\]

In vitro test

A gentle blow on the head killed each rat. The abdomen was opened, and the uterine horns were cut at their junctions with the fallopian tubes, and placed in a dish containing De Jalon’s solution. For each experiment a uterine strip was set up in a thermostatically regulated organ bath that contains the solution, which was maintained at about 37°C and gassed with air 8,9.

The uterine strip was tied with a string to a transducer (Grass FT.03) that was connected to Grass Polygraph model 07 to record contractions. A tension of 1gm was applied to the tissue and was allowed to equilibrate for at least 30 min before starting the test. Acetylcholine (Ach) was used as a stimulant to record contractions. A dose response curve was plotted with acetylcholine at a final bath concentration each of 40ng/ml, 80ng/ml, 160ng/ml and 320ng/ml. Each time, the added acetylcholine was left in contact with the tissue for 30 seconds and then washed with De Jalon’s solution. It was then left to resume its normal contraction. Acetylcholine (80ng/ml) was selected as the control concentration that induced sub-maximal contraction of the uterine tissue.

A given concentration of an extract was added in the organ bath and left in contact with the tissue for 5 min. Acetylcholine (80ng/ml) was added at the end of the 5 min in the presence of the extract and then washed after 30 sec. After rhythmic contraction resumed, the same concentration of acetylcholine was added in order to establish the reversible contractile capacity of the tissue and test the extent the extract acted upon the uterine tissue. The same procedure was repeated whenever different extracts at different concentrations were tested. Each extract was tested at 80µg/ml, 160µg/ml, 240µg/ml and 320µg/ml, as final bath concentration.

Contraction peaks recorded by the polygraph were measured in cm. The contraction peak of the control acetylcholine (80ng/ml) was taken as a reference (100%). The peak produced by 80ng/ml of acetylcholine was compared with that produced by the extract plus acetylcholine. The effect of the extract at a given bath concentration was then recorded by measuring the length of the peak produced by the combination of the extract and acetylcholine. Each value was converted to a percentage contraction by considering a 100% acetylcholine contraction. Then the mean percentage contraction was taken to compare the result 10. Seven uterine strip preparations were used for each sample tested.
Statistical analysis
For the in vivo anti-implantation tests the mean ± SEM weight gain and the mean ± SEM number of implants in each test group were compared with the respective control groups. Independent Student’s t-test was used to analyze the result. In the in vivo test, the mean (± SEM) percentage tissue contraction due to acetylcholine was also compared with the mean (± SE) percentage tissue contraction due to acetylcholine with the extract at each bath concentration. One-way ANOVA test was used to analyze the results. In each test a 95% confident interval was used.

Results

In vivo test
The weight gained with all extracts was significantly less than that of the controls (p < 0.005) as shown in table 1. The weight gained by the aqueous extracts of the leaves and roots of the plant was not significantly different from that gained by their ethanol counterparts. The root extracts, however, appeared to show less weight gain than the leaf extracts. But the difference is not significant. All rats in both control groups have shown implantation, while some rats among those, which received aqueous and ethanol extracts of both parts of the plant have shown no implantation (Table 2). The average numbers of implants in the pregnant rats with both the aqueous and ethanol extracts of the leaves and the roots of the plant were significantly less than that of the pregnant rats which received the vehicle (p < 0.005). Though it appeared that the root showed less number of implants than the leaf, and the aqueous extract showed less number of implants than the ethanol extract, the differences were not significant. The anti-fertility and anti-implantation activities produced by the aqueous extract of the leaves were 40% and 70%, respectively, while those of ethanol extract were 20% and 48%, respectively (Figure 1). The same figure shows that the anti-fertility and anti-implantation activities produced by the aqueous extract of the roots were 60% and 77%, respectively, while those of ethanol extract were 40% and 61%, respectively. Figure 1 also shows that the root showed more antifertility and anti-implantation activities than the leaf, and the water extracts were more potent than the ethanol extracts.

In vitro test
As shown in figure 2, all extracts except the ethanol extract of the leaves increased the uterine contractile activity of acetylcholine significantly (p < 0.05). Furthermore, the capacity of the extracts to increase uterine contractile activity of acetylcholine was concentration dependent (Figure 2). The aqueous extracts of both the leaves and roots showed significant increase (p < 0.05) in the contractile activity of acetylcholine as compared to the ethanol extract (Figure 2). The extracts alone did not effect contraction. The root extracts were significantly more potent (p < 0.05) than the leaf extracts as depicted in the same figure.

Table 1: Average weight gained by rats treated with 300-mg/kg of aqueous and ethanol extracts of the leaves and roots of Asparagus africanus by gavage for 19 days as compared to that of the controls.

<table>
<thead>
<tr>
<th>Extract</th>
<th>Weight gain in g</th>
<th>P-value</th>
<th>95% CI of the mean (Mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LW</td>
<td>42.7 ± 7.2</td>
<td>0.001</td>
<td>-54.85 -21.23</td>
</tr>
<tr>
<td>RW</td>
<td>37.4 ± 6.9</td>
<td>0.000</td>
<td>-59.56 -27.08</td>
</tr>
<tr>
<td>CONa</td>
<td>80.7 ± 1.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LE</td>
<td>43.2 ± 5.7</td>
<td>0.001</td>
<td>-52.12 -20.00</td>
</tr>
<tr>
<td>RE</td>
<td>39.1 ± 7.9</td>
<td>0.002</td>
<td>-60.82 -19.46</td>
</tr>
<tr>
<td>CONb</td>
<td>79.2 ± 4.1</td>
<td></td>
<td></td>
</tr>
</tbody>
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<td>CONb</td>
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<td></td>
<td></td>
</tr>
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</table>

Table 2: The number of rats showing implantation and the average number of implants counted during laparotomy on day 20 of pregnancy after treatment with the aqueous and ethanol extracts of the leaves and roots of Asparagus africanus as compared to the controls.

<table>
<thead>
<tr>
<th>Group</th>
<th>No of rats showing implantation (Mean ± SEM)</th>
<th>P-value</th>
<th>95% CI of the mean (Mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LW</td>
<td>3</td>
<td>2.8 ± 1.2</td>
<td>0.001</td>
</tr>
<tr>
<td>RW</td>
<td>2</td>
<td>2.2 ± 1.4</td>
<td>0.001</td>
</tr>
<tr>
<td>CONa</td>
<td>5</td>
<td>9.6 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>LE</td>
<td>4</td>
<td>4.8 ± 1.8</td>
<td>0.011</td>
</tr>
<tr>
<td>RE</td>
<td>3</td>
<td>3.6 ± 1.5</td>
<td>0.008</td>
</tr>
<tr>
<td>CONb</td>
<td>5</td>
<td>9.0 ± 0.3</td>
<td></td>
</tr>
</tbody>
</table>

a = control for water group  b = control for ethanol group, SEM is standard error of the mean, n= 5; LE is ethanol extract of the leaf, RE is ethanol extract of the root, LW is aqueous extract of the leaf, RW is aqueous extract of the root
The ability of all the extracts to potentiate acetylcholine effect on uterine contraction in the present study suggests the possibility of interaction of the plant material with endogenous acetylcholine to induce abortifacient effect. The results obtained in the present study correlate with that reported earlier by Makonnen et al (14) for Ricinus communis and Jatropha curcas seeds on guinea pig uterus and by Mekonnen (10) for Moringa stenopetala on mouse uterus. The fact that the root was found to be more potent than the leaf in potentiating the acetylcholine effect suggests that more amount and/or number of active principles and/or active principles with better efficacy might be present in the root than in the leaf.

The leaf and root of this plant may possess hormonal properties that can modulate the reproductive function of the experimental rats. Different mechanisms might have been involved to produce the antifertility activity that needs further elucidation.

The present study demonstrated the possible therapeutic application of the leaf and root of A. afric anus in fertility control. However, further study on the pharmacokinetic profile, safety and active principles of the plant have to be carried out.

References
Is there need for a desensitization program for patients who shared the same ward with a late colleague?

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It is Saturday morning, and though still battling a sleep deficit from last night’s ordeals in the Casualty Emergency Room, I have to head to my new residency posting at the National Tuberculosis and leprosy Program (NTLP) in-patient wards. The “aha” insight central to the theme of this filler is birthed in two occurrences on my first day here. While patients in most Western and developed country settings enjoy a relatively better nursing privacy, the picture in most developing and especially African hospitals is that of an open aura-except of course in a few private settings.

On this day, a long staying male in-patient of TB pleura (recurrent bilateral effusions) is seen by his colleagues heading to the “loo” never to return (reason! He is found dead several hours later after collapsing and hitting his head on the door). Considering his tender age, I rule out the possibility of a Vaso-Vago manœuvre—most probably it was respiratory distress leading to the syncope then head injury. That same evening, on the female ward—another seemingly stable two-times retreatment ISS patient develops DIB due to PCP—passing away despite all my “conventional interventions.” The following day, two male and three female in-patients request undue discharge. As I struggled to empathize with these patients—three facts became clear to me:

➢ The dying process is quite a traumatizing one, especially to patients (relatives and attendants aside) who shared the same ward/Cubicle as the late colleague—regardless of the diagnosis and prognosis. The open aura scenario doesn’t allow for privacy at this time, and the consequence is that witnessing non-medics are traumatized.

➢ Often, we in the medical profession take for granted ‘a’ death, having perhaps been desensitized by our training and past experiences—yet to the lay person, the picture is that of his saviours failing their mission—hopeless.

➢ Death apart—what about those bedside and corridor procedures we undertake using our seemingly ‘conventional’ yet scarcely manoeuvres—the tools: saws, blades, name it! While these may seem routine to a medic, Oh!, what a horror they are to the stranger—more so when they fail to yield good results.

While still caught up in trying to explain to those 5 patients that their stay on the ward did not mean they will be the next—it hit me hard how privacy during the dying process is a much needed thing in this setting! What do you think? May be a desensitization program for patients (&attendants) who witnessed a death could serve the purpose—given the high morbidity (or should I say poverty and poor governance) rates here that wouldn’t permit the Western picture. Regardless, such a program should serve to explain the reason for the occurrence of death, reduce fear and ultimately offer hope and trust in the system to the “survivor”.