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CHICORIC ACID, A PHYTOCHEMICAL COMPOUND OF *SOLENOSTEMON MONOSTACHYUS*: POSSIBLE DRUG CANDIDATE FOR THE RELIEF OF ERECTILE DYSFUNCTION

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Abstract- *Solenostemon monostachyus* is commonly used in folkloric treatment of many ailments including erectile dysfunction. In this study, its aphrodisiac component and underlying mechanism were investigated. Behavioural evaluation was done, following the oral administration of three different doses (100, 200 and 400 mg/kg) of ethanolic extract of *S. monostachyus*, sildenafil citrate (Viagra) and corn oil were used as positive and negative controls respectively. Mounting, intromission and ejaculation frequencies, mating performance, and orientation activities towards females. Result showed a striking effect in male rats. Molecular docking was done and heat map generated to probe the likely mechanism of action as *in vivo* studies compared with sildenafil. A library containing 45 phytochemicals previously characterized from *S. monostachyus* was generated and docked (Autodock Vina) into homology models of Arginase II, Phosphodiesterase 5 and Aromatase. From the docking result above, PDE 5 was identified as a major target for two key compounds, *Chicoric acid* and *Hesperidine*. Although the latter violated more than two rules of the Lipinski's rule of oral drug.

Conclusion: The results of this study support that Chicoric acids from *S. monostachyus* may inhibit the activity of PDE5, leading to relaxation of erectile smooth muscles. This finding may underscore the aphrodisiac potency of *S. monostachyus*

Key words; *Solenostemon monostachyus*, Sildenafil citrate, Phosphodiesterase 5, *Chicoric acid*, erectile dysfunction.

I. INTRODUCTION

Sex is the most cherished, indispensable and an integral part of every individual and can be a cradle of pleasure and satisfaction [1]. It is essential for procreation and general wellbeing, as it bonds a relationship and also has

a calming effect. Most men enjoy sexual activity, which may include penetrative intercourse. In order to achieve this successfully, erection has to be sustained till the female partner reaches orgasm. When a male is repeatedly unable to get or keep an erection firm enough for sexual intercourse, Erectile dysfunction is said to have resulted.

Erectile dysfunction, also referred to as "impotence," is commonly managed with the help of aphrodisiacs. An aphrodisiac is any substance or agent (food, drug, scent or device) that stimulates the erotic instinct, induces venereal desire and surges pleasure and performance [2]. The usage of many common medicines such as; blood pressure drugs, suppressants and cimetidine (an ulcer drug) can produce ED as a side effect. Nevertheless, psychological factors such as stress, anxiety, guilt, depression, low self-esteem, and fear of sexual failure cause 10 to 20 percent of ED cases. Other possible causes are smoking, which affects blood flow in veins and arteries, as well as hormonal abnormalities, like inadequate testosterone, overactivity of some enzymes like aromatase and arginase II. The availability of Phosphodiesterase type 5 inhibitors, like tadalafil (Cialis) and sildenafil (Viagra) are the most commonly used aphrodisiac agents, but these drugs cause dizziness, headache, visual disturbance, pulse irregularities, dyspepsia, priapism, diarrhoea and flushing [3]. Any medicinal plant with aphrodisiac tendencies should produce statistically significant increase in mating frequencies [4]

Traditional preparations (herbs) have been reported to have contributed to revolutionary insight in the treatment of sexual inadequacies and have become acceptable all over the world as "instant" treatment. About 80% of the world's population still rely on local medicines and traditional treatments mainly from plant extracts. Interestingly, herbal medicine is becoming very popular in the developing countries.



S. monostachyus is an aromatic, medicinal plant that belongs to the family Lamiaceae. In English language, it is called Monkey potato. Its names among some ethnic groups in Nigeria include Ntorikwot (Ibibio) and Olojogbodu (Yoruba). The use of *S.monostachyus* plant in traditional medicine by the people of Nigeria has been documented [5,6,7]. The phytochemical screening of *S. monostachyus* leaf extract revealed the presence of saponins, tannins, cyanogenic glycosides, flavonoids and alkaloids [8]. Despite the acclaimed use of *S. monostachyus* as sex enhancer in some parts of Nigeria, there seems not to be any substantive information in the open scientific literature with respect to its clear mechanism of action. Therefore, this study was designed to investigate and establish the traditional aphrodisiac potential of the herb.

II. MATERIALS AND METHODS

Collection of Plant materials and identification

The root of *S. monostachyus* was obtained in April 2019 from the University of Ibadan. The plant was harvested around the forest near new stadium in Ibadan. The plant was botanically identified and authenticated by Mr. Donatus O. Esimekhuai of the Department of Botany and Microbiology, University of Ibadan, Oyo state, Nigeria. with voucher no UIH -22874 with a specimen deposited at University of Ibadan.

Plant Extract Preparation

The whole plant of *S. monostachyus* was air dried for 72 h after which it was oven dried at a temperature of 80°C for 72 h. Thereafter, it was blended into powder, using a grinding machine. About 634 g of the powder was macerated in n-Hexane, ethyl acetate and ethanol (2500 ml) for 72 h with intermittent stirring in the morning and evening. The mixture was filtered on the 3rd day and the filtrate was concentrated using a rotary evaporator. The extract yield weighed 12.13g for n-Hexane, 11.83 for ethyl acetate and 12g ethanolic extracts. They were stored in the refrigerator for further use.

Animals

The male and female albino rats were obtained from the Department of Zoology in the University of Benin, Edo state and kept in the animal house of the Department of Pharmacology and Toxicology, Madonna University, Elele Campus. They were allowed to acclimatize for over 1 week before the experiments commenced. They were given growers feed (Vital feed growers by grand feed cereals, Onitsha.) kept in clean cages with access to water and under normal conditions. The rats with the weight range of 130-150g were used for this study. All the animals received humane care according to the criteria outlined in the 'Guide for the Care and Use of Laboratory Animals' prepared by the National Academy of Science and published by the National Institute of Health.

Phytochemical Screening

Preliminary phytochemical screening of the root extract of *S. monostachyus* was carried out using standard methods described by [9] and [10]. The extract was screened for the presence of flavonoids, alkaloids, tannins, terpenoids, saponins and glycosides.

Pilot studies

Pilot studies was done using graded doses of the three solvent extracts. Ethanolic extract showed more promise and was used for the study.

Acute toxicity study: Healthy male albino rats were starved for 3-4 hr and subjected to acute toxicity studies as per (OECD) Organization of Economic Co-operation and Development guidelines No: 423 [11]. They were divided into 4 groups of 4 animals each and kept singly in separate cages during the experiment. Group 1 represented the control group, which received 10 ml/kg of distilled water orally. Groups 2- 4 received suspension of ethanolic extract of *S. monostachyus* orally at the doses of 1000, 2000 and 5000 mg/kg daily for 7 days respectively. The rats were observed continuously for 2 hours for behavioural, neurological and autonomic profile, and for next 24 and 72 hours for any lethality or death

Invivo aphrodisiac studies

Preparation of male rats: Sexual behaviour training was given to male rats for 10 days. The animal that did not show any sexual interest was replaced by another sexually active rat [12].

Preparation of female rats Oestrus (heat) was developed artificially in female rats by giving clomiphene orally 2-3 days before mating. Before conducting the experiment the receptivity of female rats was ensured [12].

Preparation of female rats

Oestrus (heat) was developed artificially in female rats by giving clomiphene citrate 50mg/kg body weight (orally), 2 days before conducting the experiment. the receptivity of female rats was ensured.

Grouping of Rats

The rats were divided into 5 groups each having 4 rats.
Group I: Control that was not given any extract but food and water and corn oil
Group II: Sildenafil group as our standard for comparison
Group III: 100 mg/kg body weight of extract dissolved in corn oil
Group IV: 200 mg/kg body weight of extract dissolved in corn oil
Group V: 400 mg/kg body weight of extract dissolved in corn oil

Observation of behavioural sexual parameters



Observation of rats Close observation was made once every week for 14 days for the sexual parameters of the rats which entails;

Mount latency: Time interval (seconds, minutes) from the introduction of the female into the cage and the first mount by the male.

Mount frequency: Mounting is defined as the climbing of one animal by another usually from the posterior end with the intention of introducing one organ into another. Mount may also be operationally defined as the male assuming the copulatory position but failing to achieve intromission. Mount Frequency (MF) is therefore defined as the number of mounts without intromission from the introduction of the female until ejaculation.

Intromission latency: Time interval from the introduction of the female to the first intromission (vaginal penetration) by the male.

Intromission frequency: Intromission is the introduction of one organ or parts into another e.g. the penis into the vagina. Intromission Frequency (IF) is defined as the number of intromissions from the time of introduction of the female until ejaculation.

Ejaculation latency: Ejaculation is the act of ejecting semen brought about by a reflex action that occurs as the result of sexual stimulation. Ejaculation latency (EL) is therefore the duration from first intromission till ejaculation.

Post ejaculation interval: Time interval between ejaculation and next intromission.

All these parameters were observed for the various groups of rats.

Ligand Library Generation

Identified secondary metabolites of *S.monostachyus* employed for this study were determined from published literature and were used in the creation of the ligand library. The secondary metabolites were retrieved from NCBI PubChem database [13]. In Standard Database Format format (2D). The ligand library generated were imported to Maestro and prepared using the [14]. Utilizing Ligprep v4.5[14], Epik v4.3 [15]. With OPLS3 force field, for protonation, stereo-isomerization, tautomers generation, and to attain biological conformer.

Energy minimization was achieved for all tautomeric state at pH of 7±2.

Protein Preparation

Human Arginase II (1.8 Å), PDE-5 (2.8 Å), and Aromatase (2.75 Å) bound with ligands were retrieved from the PDB [16] With the PDB ID 4I06, 2H42, and 5JKV, respectively. Protein Preparation Wizard [17]. Module in maestro 11.5 was used to prepare each protein complex. Protein structure's missing hydrogen atoms, missing loop, and missing sidechains were fixed while the added hydrogen atoms were optimized at pH 7.0. Optimized structures were then minimized using the OPLS3 force field by converging heavy atoms to RMSD of 0.3Å.

The protein binding site was identified by utilizing the receptor grid generation tool in maestro 11.5. Receptor grid defines the region of interaction between the protein and the ligand. The co-crystal ligands of each protein were used to identify the binding cavity employing default parameters of van der Waals scaling factor 1.00 and charge cutoff of 0.25 around the binding site residues of the protein structures.

Docking Studies

Molecular docking studies were performed using Glide v7.8 [16]. Module on maestro 11.5. The ligand library compounds and co-crystal ligands were docked into the binding cavity of the targets using the standard precision algorithm (SP) applying a scaling factor of 0.8 and partial charge cutoff of 0.15, and the ligand was handled as flexible. Five (5) of the lowest energy poses were selected for post-docking minimization seething the threshold at 0.50 kcal/mol [16]. Lastly, the binding affinity of the receptor-ligand complex was ranked according to Glide score and ligand pose.

ADME/TOX Prediction

The absorption, distribution, metabolism, and excretion of the hit ligands were predicted using the Qikprop module in maestro 11.5. QikProp predicts physically significant descriptors and pharmaceutical features of organic molecules. It uses descriptors that are gotten from molecular structure and calculated molecular properties [18]

Results:

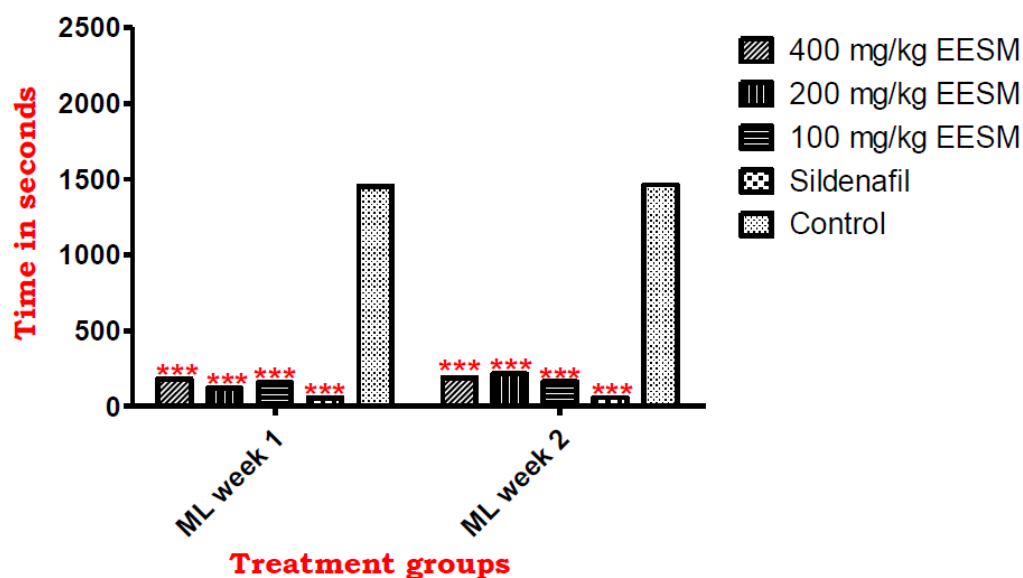


Fig 1. Effect of *S. monostachyus* on mount latency for 2 weeks in male rats.

Animals per group (n) = 4. The values are mean $\pm$ SEM.; * p < 0.05; ** p < 0.01; *** p < 0.001 when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test).

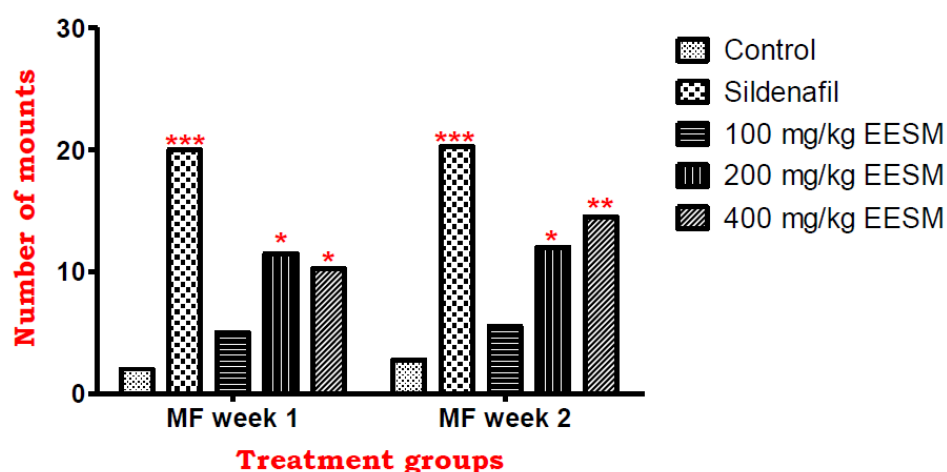


Fig 2. Effect of *S. monostachyus* on mount frequency in male rats for 2 weeks.

Animals per group (n) = 4. The values are mean $\pm$ S.E.M.; * p < 0.05; ** p < 0.01; *** p < 0.001 when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test).

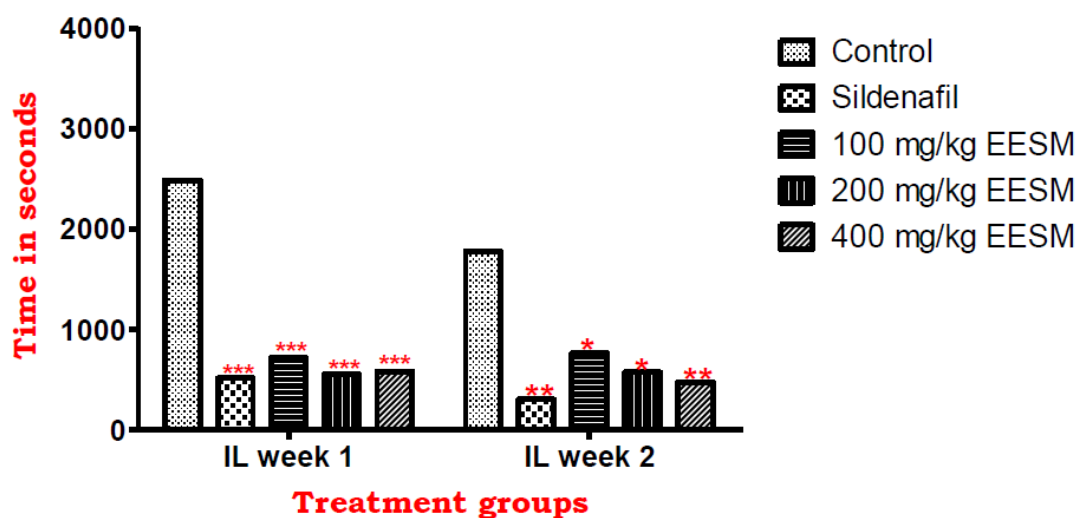


Fig 3. Effect of *S. monostachyus* on intromission latency in male rats for 2 weeks.

Animals per group (n) = 4. The values are mean $\pm$ S.E.M.; * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$ when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test).

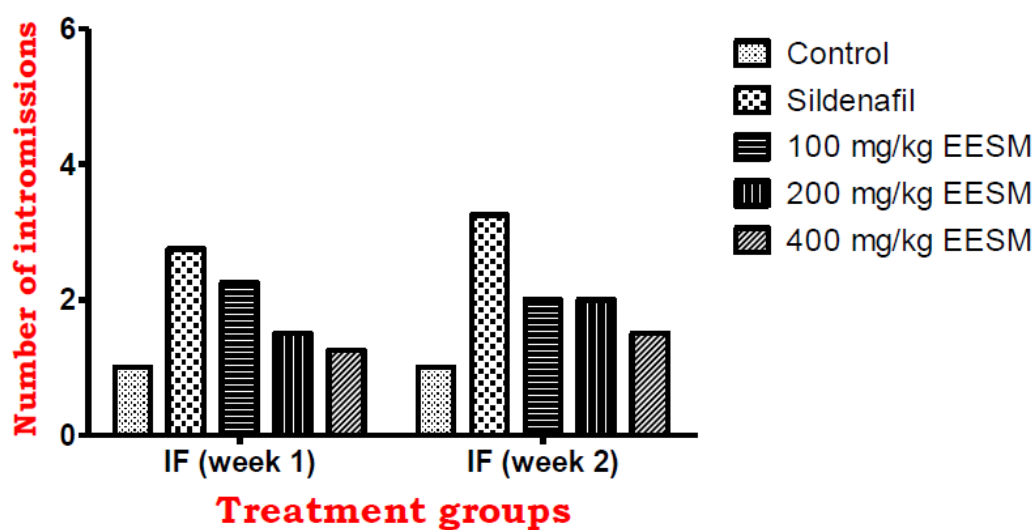


Fig 4. Effect of *S. monostachyus* on intromission frequency in male rats for 2 weeks.

Animals per group (n) = 4. The values are mean $\pm$ S.E.M.; * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$ when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test).

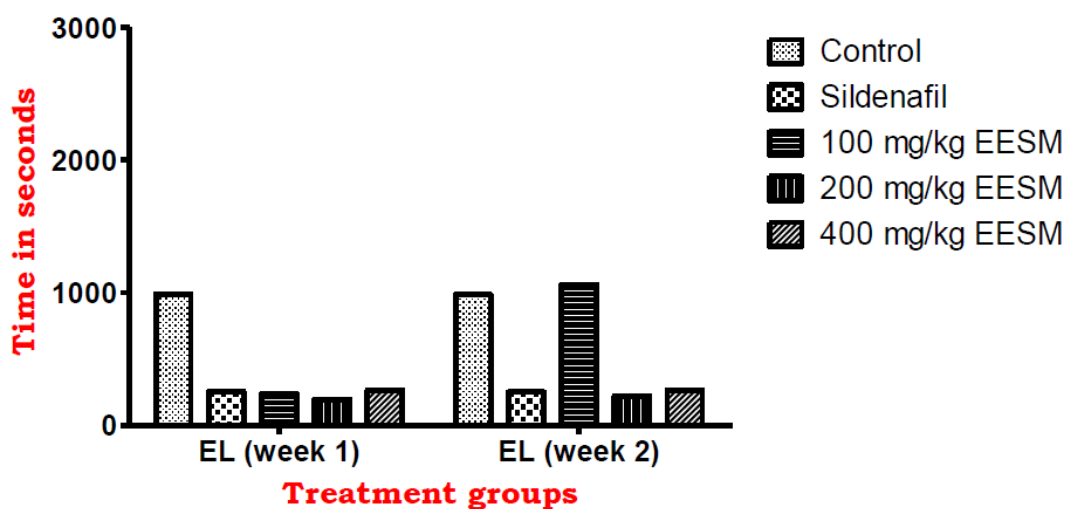


Fig 5. Effect of *S. monostachyus* on Ejaculatory latency in male rats for 2 weeks.

Animals per group (n) = 4. The values are mean $\pm$ S.E.M.; * p < 0.05; ** p < 0.01; *** p < 0.001 when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test)

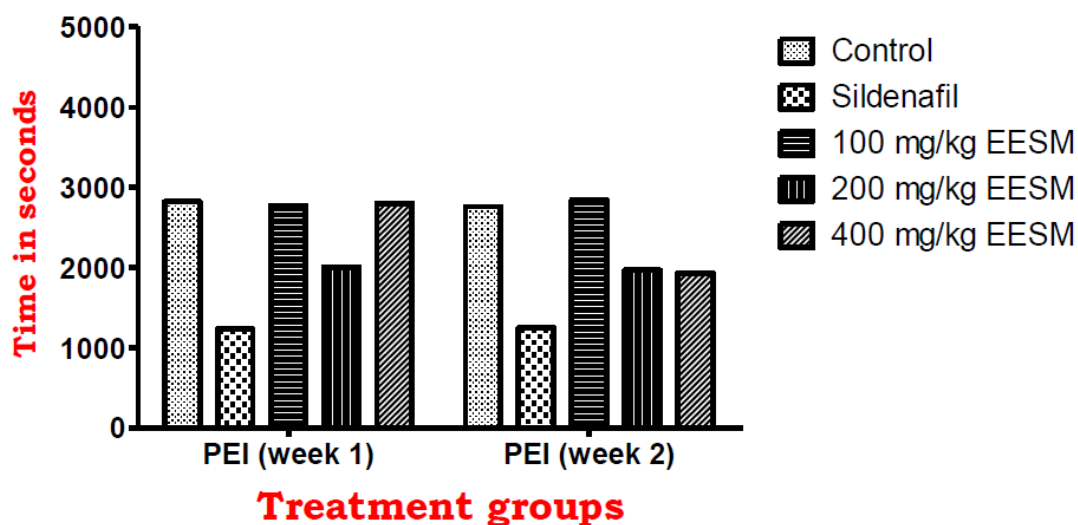


Fig 6. Effect of *S. monostachyus* on Post Ejaculatory Interval in male rats for 2 weeks.

Animals per group (n) = 4. The values are mean $\pm$ S.E.M.; * p < 0.05; ** p < 0.01; *** p < 0.001 when compared with control group. (Two-way ANOVA followed by Bonferonni post hoc test).

Table 2. Showing results of docked scores of some relevant mined compounds of *S.monostachyus* against erectile dysfunction targets.

Docking Result for <i>S. monostachyus</i>				
S/N	Compounds	ARGINASE II	PDE5	AROMATASE
1	CoLigands	-8.01	-11.127	-7.502
2	Chicoric acid	-5.393	-9.973	-7.221
3	Hesperidin	-5.465	-9.654	-8.048
4	Myricetin	-6.886	-9.129	-6.027
5	Quercetin	-6.624	-8.819	-5.776

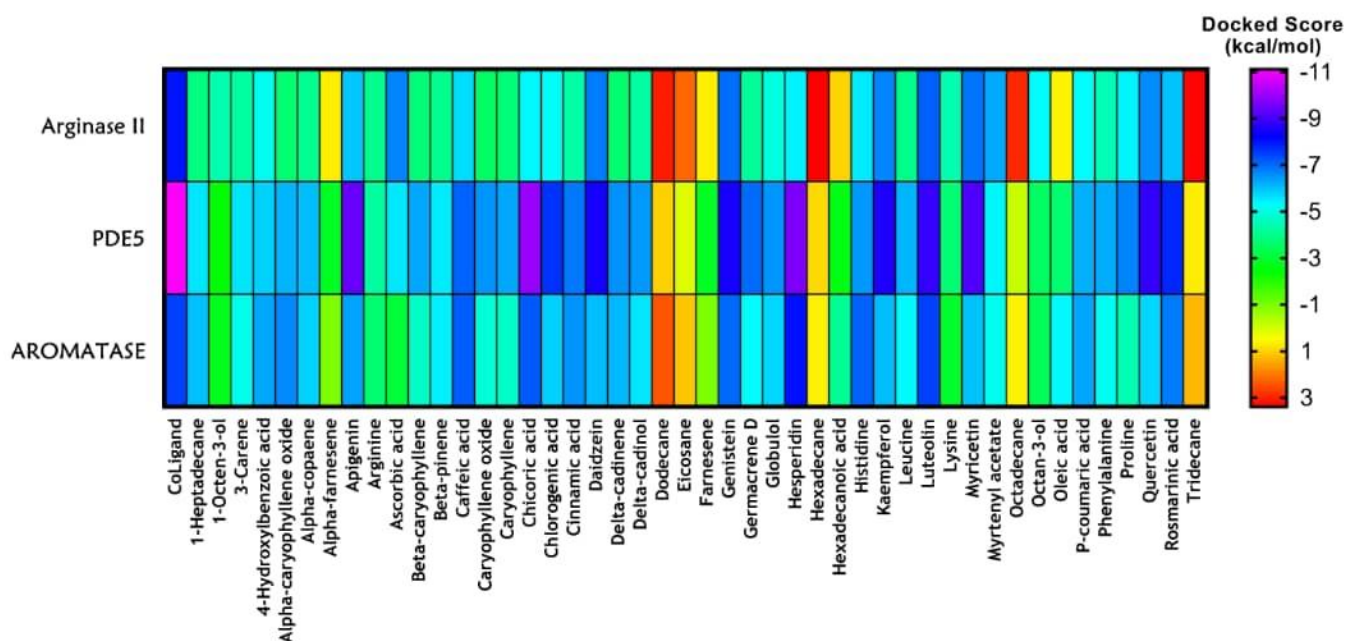


Fig 7. Heat map representation of docking result. The free energy of binding of phytochemicals of *W. indica* docked into the substrate binding sites of Aromatase, Arginase II and PDE 5, are represented as heat map. (The scale is a spectrum from red (3 kcal/mol) to pink (– 11 kcal/mol)).

Table 3. Result showing Adme/tox values of the compounds that compared well with the co crystallized PDE 5 inhibitor.

S/n	Molecule	Molecular weight	donor Hb	Accept Hb	QplogPo/w
1	PDE5 Co-Ligand	475.577	1	11.75	1.766
2	Chicoric acid	474.37	6	11	0.677
3	Hesperidin	610.568	7	20.05	-1.48

MW, molecular weight: R.V.: 130–725; donorHB, estimated number of hydrogen bonds

that would be donated by the solute to water molecules in an aqueous solution: R.V.: 0.0–

MW, molecular weight: R.V.: 130–725; donorHB, estimated number of hydrogen bonds that would be donated by the solute to water molecules in an aqueous solution: R.V.: 0.0–6.0; accptHB, estimated number of hydrogen bonds that would be accepted by the solute from water molecules in an aqueous solution: R.V. = 2.0–20.0; QPlogPo/w, predicted octanol/water partition coefficient: R.V. = -2.0 to 6.5; QPlogHERG, predicted IC50 value for blockage of human Ether-a`-go-go

related gene K⁺ channels, <-5 = concern; QPPCaco, predicted apparent gut-blood barrier permeability, <25 = poor, >500 = great; QPlogBB, predicted brain/blood partition coefficient, -3.0 to 1.2; QPPMDCK, predicted apparent Madin-Darby canine kidney cell permeability, <25 = poor, >500 = great; QPlogKhsa, prediction of binding to human serum albumin, -1.5 to 1.5; HOA, human oral absorption level, 1, 2, 3: 1 = low, 2 = medium; ROF, the number of violations of Lipinski's rule of five; R.V., recommended value.

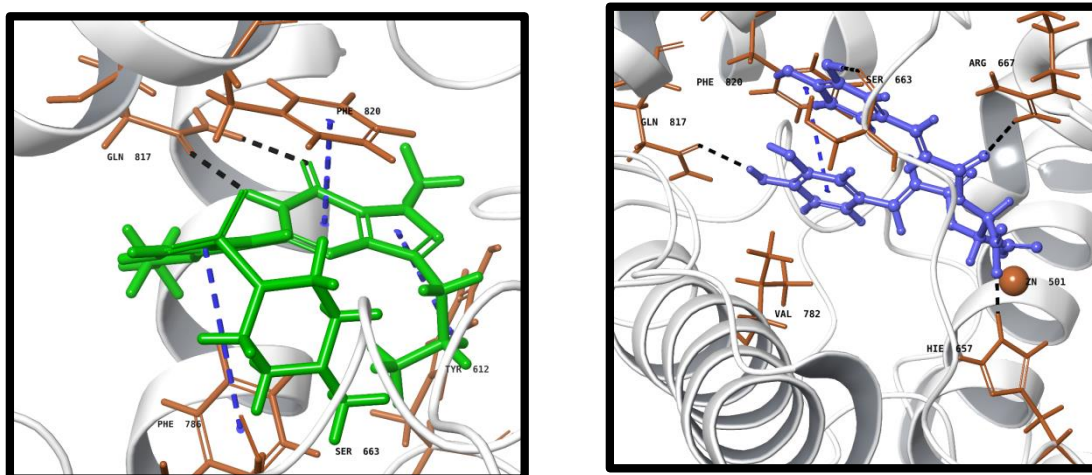


Fig 8. (A) Binding poses of Coligand; 7-(6-methoxypyridin-3-yl)-4-{{2-(propan-2 yloxy)ethyl}amino}-1-(2-propoxyethyl)pyrido[4,3-d]pyrimidin-2(1H)-one- (OH3) in the active site of PDE 5 **(B)** Binding poses of Chicoric acid in the active site of PDE 5.

H-Bond - - - - -
pi-pi stacking - - - - -
Salt bridge - - - - -
pi-cation - - - - -



III. DISCUSSION

The higher access to herbs over expensive pharmaceutical drugs to treat diseases among non-industrialized societies is fast becoming revolutionized. In some countries, it has been integrated into the health scheme despite advances in orthodox medicine. It is thought that the natural products if used properly are less harmful than synthetic products, which most often elicit some side effects [19].

In vivo studies of ethanol extract of *S. monostachyus* somewhat justified the claim that it has some aphrodisiac effect as suggested in folklore medicine. The mount latency per time in seconds observed in various extract doses were seen to be significantly shorter compared to the untreated subjects and it wasn't dose dependent. Similar result was obtained in number of mounts as shown. There was significant difference in sperm counts treated rats when compared to untreated rats. Intromission latency sort of showed similar result, Indicating that the agent may well be very effective in terms of arousal.

Other behavioural parameters of intromission frequency, Ejaculatory latency and post ejaculatory interval didn't show such effect.

In-silico studies was done to predict the compounds if any that may be responsible for the effect hypothesized. It was therefore important to do molecular docking studies. Molecular docking procedure is based on the ability of the compounds to interact with amino acid residues, which is a function of the protein conformation, and the assumed pose of the ligand [20]. Previously identified and characterized compounds of *S.monostachyus* were retrieved from PubChem using ChemAxon suite (<https://www.chemaxon.com>). The 44 Library of phytochemicals generated was docked against the three target proteins which may be therapeutic in erectile dysfunction viz: Phosphodiesterase 5, (PDE-5), aromatase and arginase II. Sex enhancing potentials, may include the inhibition of the hydrolyzing action of PDE-5 with the result that active cGMP can accumulate. 'Undisturbed' and prolong the erection through increased blood flow [21]. Aromatase, is an enzyme involved in the production of estrogen that acts by catalyzing the conversion of testosterone (an androgen) to estradiol (an estrogen). Low testosterone and elevated estrogen have been implicated in the increase of the incidence of erectile dysfunction independently of one another [22]. Arginase II is a metalloenzyme, which catalyzes the hydrolysis of L-arginine to produce L-ornithine and urea. It is proposed that it competes for L-arginine and reduces Nitric oxide synthase activity in genital tissues (Nitric oxide synthase (NOS) utilizes L-arginine and oxygen as substrates to produce nitric oxide (NO) and citrulline.) thus modulating sexual function [23].

Results obtained from docking, indicated that chicoric acid and hesperidin may well inhibit PDE- 5 significantly though not as much as the co crystallized

ligand 7-(6-methoxypyridin-3-yl)-4-{{2-(propan-2-yloxy)ethyl}amino}-1-(2 propoxyethyl)pyrido[4,3-d]pyrimidin-2(1H)-one- (Sildenafil) at -11.127 Kcal/mol, but was able to substantively inhibit at -9.973Kcal/mol. While hesperidin also inhibited enzyme at -9.654 Kcal/mol. Both compounds have been identified to be components of *S. monostachyus* by [24].

The Admetox predictions for the compounds of interest, showed that Chicoric acid may well pass for a drug candidate as it only violated one of the Lipinski's rule of five. Which makes it rather eligible as a potential oral drug candidate [25].

IV. CONCLUSION

This result confirmed the earlier wet experiment, that Ethanolic extract of *S. monostachyus* has aphrodisiac properties as it looks to have inhibitory potential on PDE-5 and as such Chicoric acid should be considered for clinical trials as an aphrodisiac drug.

Acknowledgment

We are grateful to all the researchers at the Pharmacology and Pharmacognosy Department of Madonna University, Centre for Bio-computing and Drug Development, Adekunle Ajasin University, Ondo state, for their support and team spirit, without whom efforts this work would not have been possible.

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