Antilactogenic Characteristics of Methanol and Ethyl acetate Crude Extract of Aframomum melegueta seeds in Wistar rats

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Abstract
The serious side effects associated with orthodox drugs used in controlling excessive milk secretion has necessitated the search for natural alternative with little or no side effects. This study was conducted to ascertain the antilactogenic characteristics of methanol and ethyla cetate crude extracts of Aframomum melegueta seeds in female Wistar rats. The extracts were obtained using soxhlet apparatus with the first divided half re-extracted in methanol and the second in ethyla cetate. A total of 48 female Wistar rats were assigned into two groups of 24 rats, for lactating and non-lactating groups. These were assigned into 4 treatment groups of 6 rats each for non-lactating and lactating rats. Treatments I and ii (controls) received control drug (bromocriptine) and placebo (1ml dimethylsulphoxide in 9ml of normal saline) at doses of 0.11mg/kg and 5ml/kg body weight per day respectively. Treatment iii and iv received methanol and ethylacetate extracts at doses of 100mg/kg and 94 mg/kg body weight respectively via an oral intubation. The administration was carried out for a period of 21 days with food and water given ad libitum. The rats were sacrificed and blood collected by cardiac puncture and serum concentrations of prolactin, estrogen and progesterone were analysed. The result obtained revealed that reference control drug caused significant (P<0.05) reduction in serum levels (12.56±0.03; 19.90±3.16; 3.05±0.06; 3.17 ± 0.09; 323.43± 0.20 and 320.60± 0.19 ng/ml). The methanol extract of Aframomum melegueta seeds like the standard control drug produced significant decrease (26.15±0.15; 14.90±3.16; 2.68±0.08; 2.40± 0.03; 315.72±7.72; and 318.00± 2.26 ng/ml in serum prolactin, estrogen but no significant changes in progesterone of both non-lactating and lactating wistar rats relative to normal control rats. Ethylacetate extract however produced significant reduction (21.57±1.82; 22.40± 1.60; 2.45±0.10; 2.17± 0.06; 323.03±0.25, 323.42±1.68 ng/ml) in serum prolactin levels of non-lactating rats, estrogen of both non-lactating and lactating rats but no significant (P>0.05) alterations in serum levels of prolactin in lactating and progesterone of non-lactating and lactating rats relative to normal control rats. The result of this study indicates that both extracts of Aframomum melegueta exhibited antilactogenic characteristics with methanol extract on the lead.

Keywords: Antilactogenic characteristics, methanol/ ethylacetate extract, Aframomum melegueta, wistar rats.

1. Introduction
The complex physiologic hormonal involvement in mammary gland development and milk secretion has opened a wider area of research (Wolfang et al, 2010) [44]. The hormones range from corticosteroids, growth hormone, insulin, prolactin, estrogen and progesterone. The selected hormones of interest in this study are the prolactin, estrogen and progesterone (Agarwal et al, 2014) [2]. Prolactin, estrogen and progesterone all play one role or the other in mammary gland development and milk production (Tripathi, 2001) [41]. The inhibitory effect of progesterone on lactogenesis during pregnancy is evident (Lopez-fontana, et al, 2012) [26]. With parturition and expulsion of placenta, progesterone level falls and full milk supply is initiated (Neville 2001) [31]. During lactation, several key hormones are involved in the regulation of mammary gland cell number, secretory activity and consequent milk production potential (Malgwi et al, 2017) [27]. Prolactin is mainly synthesized and secreted by the lactotrophic cells of the pituitary gland (Freeman et al, 2000) [18]. It is the major generation of...
Lactation competence during pregnancy and it functions both indirectly, through its regulation of ovarian progesterone production, and directly via its effect on mammary epithelial cells (Hector and Lindsay, 2012) [21]. The role of prolactin during mammary gland morphogenesis has been investigated using prolactin receptor knockout mice (Ormandy et al, 1997) [34]. This experiment revealed normal development in the transplanted tissues, suggesting that defect in knockout glands are due to lack of prolactin/prolactin receptor signaling in other tissues.

Researchers have found that simply restoring progesterone levels in ovariectomized prolactin mice restored ductal side branching (Vomacka et al, 2000) [43] providing further affirmation that prolactin and progesterone work together to generate lobuloalveolar growth.

Estrogen, a membrane soluble ligand is released from the ovary and activates gene expression through intracellular receptors (Bandyopadhayay, 2013) [8]. Estrogen unequivocally stimulates mammary ductal growth in both ovariectomized and intact animals and can be blocked by estrogen receptor antagonist, tamoxifen (Silberstein et al, 1994) [38]. Estrogen acts in concert with insulin growth factor 1(IGFI) to generate the burst of proliferation required for ductal morphogenesis at puberty (Rauan and Kleinberg, 1999) [37]. This study focused attention on serum levels of three of these hormones which play essential role in lactogenesis i.e prolactin, estradiol and progesterone.

The attendant complications (galactorrhea, hyperprolactinemia, prolactinoma) associated with excessive breast feeding and the serious side effect (nausea, fatigue, postural hypotension, nasal congestion, exacerbation, psychosis, seizure) associated with orthodox drugs (bromocriptine, pergolide, cabergoline, quinagolide) have necessitated the search for natural alternative with little or no side effect which can be used in alleviating the situation. *Aframomum melegueta* (Alligator pepper) has been incriminated. *Aframomum melegueta* methanolic extract has been noted to inhibit 56.7±3.4% of estrogenic activity in a yeast assay when at the concentration of 100μg/ml which was decreased with naringinase pretreatment (EI-Halawany et al, 2011) [13]. *Aframomum melegueta* lowering effect on serum prolactin has been reported (Ebong et al, 2000) [12]. The use of *Aframomum melegueta* seed extract in reducing excessive lactation and post partum haemorrhage in Ghana has been reported (Mark, 2004) [29]. Dopaminergic agonist with D2 receptor activity suppressed prolactin secretion (Clive et al, 2002) [8]. *Aframomum melegueta* has been noted to in vitro inhibit cytochrome P3A4 enzymes at 10mg/ml concentration (Agbonon et al, 2010) [1]. This may affect the capacity of this enzyme in metabolizing xenobiotics. Progesterone levels decreased significantly when wistar rats were administered with 13.3mg/kg body weight of a saline extract of Alligator pepper (Ingbenebor and Ebomoyi, 2014) [23].

Materials and Methods

Preparation of plant material.

Fresh fruits of *Aframomum melegueta* (A. melegueta) weighing 3000g were purchased from Apiapum market in Obubra Local Government Council of Cross River State of Nigeria. The fruits were sundried for two weeks to facilitate removal of the seeds from the pods (capsule). The seeds were further sundried to constant weight within four days and stored in an air-tight flask for extraction.

Preparation of *Aframomum melegueta* seed extract.

The seeds were ground to powdered form by the use of motor powered milling machine at Root Crop Research Institute, Umudike in Abia State. A 1000g weight of *Aframomum melegueta* seed powder was first extracted with petroleum ether (60-80 °C) to defat the powder. The ether extract was carefully decanted leaving the residue for further extraction in methanol and ethylacetate solvents in two divided doses. The soxhlet apparatus was filled with methanol and 300g of the residue was wrapped in a thimble and placed in a soxhlet extractor fitted to a 500ml round bottom flask seated on a hot plate. The reflux condenser was attached to the extraction tube and power was supplied by switch linking the electric hot plate cable. The methanol vapour passes up the side tube of condenser and runs back on to the residue in the thimble. After siphoning over for 24 times, the experiment was stopped just before the next lot of methanol was at the point of siphoning over. The methanol soluble fraction was evaporated of its methanol content using a rotary evaporator and was preserved at 4 °C. The same process was carried out to produce ethylacetate extract of *Aframomum melegueta* seed.

Preparation of stock solution of crude extract of *Aframomum melegueta* seeds.

One gram (1g) of methanol extract of *Aframomum melegueta* seed was dissolved in one (1ml) milliliter of Dimethyl sulfoxide (Dmso) and suspended in 9ml of normal saline (0.9% NaCl). The same 1g of ethylacetate of *Aframomum melegueta* seed was used for its stock solution preparation. The stock solutions were immediately administered to rats in treatment III and IV of this study respectively.

Animal and animal treatment.

A total of 24 wistar rats each for non-lactating and lactating groups were assigned on the basis of weight (150-200g) into 4 treatments of 6 rats each. The animals were housed in the Department of Biochemistry, University of Calabar under standard laboratory conditions of ambient temperature of 26°C and adequate ventilation with relative humidity of 50% and a 12 hour day-light cycle. Animal were fed ad libitum during the course of treatment. The administration of *Aframomum melegueta* seed extract commenced 3 days after parturition and lasted 3 weeks for both non-lactating and lactating rats. Rats in Treatment Group I: received bromocriptine (reference standard drug) at a dose of 0.11mg/kg/day according to manufacturer’s instruction. Treatment Group II rats received 100mg/kg/day of dimethyl sulfoxide (1ml in 9ml of normal saline) for normal control treatment. The rats in treatment III and IV received 100mg and 94mg/kg/day of methanol and ethylacetate seed extracts of *Aframomum melegueta* via oral intubation respectively. The treatment was terminated after a 21 day period of administration of crude extract. The rats were starved of food overnight in preparation for sample collection. The blood sample was aseptically collected via cardiac puncture and transferred into sample labeled bottles. The collected blood was allowed to stand for 2 hours to perfect clotting of blood and centrifuged at 1000rpm and serum removed with Pasteur pipette for assay of prolactin, estradiol and progesterone concentrations.
Estimation of serum prolactin concentration.
This was carried out by microwell enzyme immuno-assay technique using the method of Frantz, (1978) [17]. A total of 53 streptatavidin located micro-plate was removed from the zip-lock kit being-one for a substrate and two each for reference standard and control and 48 micro-plates for the test samples. The micro-plates were properly labeled and placed in a micro-plate holder. Twenty five (25 µl) micro-liter of standards, controls and samples were dispensed into each micro-well. A total of 100µl of conjugate reagent was dispensed into each well and thoroughly mixed for 30 seconds. These were incubated at room temperature for 60 minutes and the content of the well discarded by decantation and the micro-plate blotted dry with absorbent paper. A 300N1V wash was aspirated and repeated twice making a total of three washes. One hundred (100 µl) micro liter of 3, 3’ 5’ 5’ tetramethyl benzidine (TMB) was added into each well and gently mixed for 110 seconds. These were incubated at room temperature in the dark for 15 minutes. A 50 µl of stop solution was gently added to each well and mixed after the blue colour of the well has completely changed to yellow. The absorbance of each well was read at 450nm wave length using spectrophotometer. The average absorbance values were calculated from the duplicate standards, controls and sample and a standard curve was constructed by plotting the average absorbance obtained from each reference standard against its assigned concentration in ng/ml on linear graph paper with the absorbance on the vertical (Y) axis and the concentrations on the horizontal (x) axis. The corresponding concentration of prolactin in the sample was determined from the intersecting point of the curve.

Estimation of serum estrogen (estriadiol) concentration.
This was determined by microwell enzyme immunoassay according to the method of March et al, (1979), and progesterone serum concentration according to the method of Radwanska et al, (1978) [35].

Stastical Analysis
One way analysis of variance was used to test for significance of difference and significant means were separated using least significant difference of Fishers.

Result
The results presented in table 1, II, and III depict the outcome of various treatment administered to wistar rats in the course of the investigation. Table I shows the serum levels of selected lactogenic hormones (prolactin, estradiol and progesterone) resulting from the various extract of Aframomum meleguetia seed and reference control drug bromocriptine administered in the course of this study to non-lactating wistar rats. Bromocriptine significantly produced generalized reduction in all the hormones tested (10.56±0.89ng/ml prolactin, 2.81±0.12ng/ml estradiol and 12.96±0.06ng/ml progesterone amounting to 7.86 and 60.41% reduction respectively relative to (33.23±1.82ng prolactin, 3.05±0.06ng estradiol and 323.43±0.20ng/ml progesterone) control treatment accordingly.

The 100mg/kg body weight extract of Aframomum meleguetia seed caused a significant (P<0.05) reduction in serum levels of prolactin 26.15±0.15ng/ml or 21.31%, estrogen 2.68±0.08ng/ml or 12.13% and non-significant reduction in progesterone 31.72±7.72ng/ml or 2.0% relative to (33.23±1.82ng/ml, prolactin, 3.05±0.06ng/ml and 323.43±0.02ng/ml progesterone) relative to normal control treatment. Administration of 94mg/kg body weight of rats revealed significant decrease in (21.57±1.82ng/ml prolactin or 35.08%, 2.45±0.01ng/ml or 19.67% estrogen and non-significant 323.03±1.82ng/ml of prolactin, 3.05±0.06ng/ml estrogen and 323.43±0.02ng/ml progesterone) relative to control experiment in non-lactating rats. When methanolic extract was compared with the standard reference drug bromocriptine, the result showed a significant reduction (12.56±0.89ng/ml or 51.97 of prolactin, 127.96±0.06ng/ml or 60.39% of progesterone but 2.81±0.12ng/ml or 12.81% increase in estrogen) relative to ethylacetate extract (21.57±1.82ng/ml prolactin, 323.03±0.25ng/ml progesterone and 2.45±1.82ng/ml of estrogen for non-lactating rats. When compared within themselves, ethylacetate extract of Aframomum meleguetia seed caused a significant marginal decrease in serum levels of (21.57±1.82ng/ml of prolactin or 17.51%, 2.45±0.01mg/ml or 8.58% of estrogen but non-significant increased 323.03±0.25ng/ml of progesterone relative to 26.15±0.15ng/ml, 2.68±0.06ng/ml and 315.72±7.72ng/ml of prolactin, estrogen and progesterone respectively.

Table 1: Treatment effects on lactogenic hormones of non-lactating rats

<table>
<thead>
<tr>
<th>Lactogenic Hormones</th>
<th>Bromocriptine 0.11mg/kgb,w</th>
<th>Normal control 5ml/kgb,w</th>
<th>Methanol extract 100mg/kgb,w</th>
<th>Ethylacetate extract 94mg/kgb,w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactin</td>
<td>12.56±0.89b</td>
<td>33.23±1.82b</td>
<td>26.15±0.15b</td>
<td>12.57±1.82b</td>
</tr>
<tr>
<td>Estradiol</td>
<td>2.88±0.12b</td>
<td>3.05±0.06b</td>
<td>2.68±0.08b</td>
<td>2.45±1.10b</td>
</tr>
<tr>
<td>Progesterone</td>
<td>127.96±0.06b</td>
<td>323.43±0.20b</td>
<td>315.72±7.72b</td>
<td>323.03±1.68b</td>
</tr>
</tbody>
</table>

Mean is ± SD. Mean values n, b or m superscript are significantly (P<0.05) different from normal control, bromocriptine and methanol extract of Aframomum meleguetia seed.

In the lactating, rats the observations and findings are not much different. Administration of 0.11mg/kg bromocriptine per body weight of rats produced significant reduction (10.56±26ng/kg or 46.93% of prolactin, 2.41±0.80ng/ml or 23.97% of estrogen and 110.21±15ng/ml or 65.63% of progesterone) in selected lactogenic hormones relative to (19.90±3.16ng/ml, of 3.17±0.09ng/ml and 320.60±0.019ng/ml of prolactin, estrogen and progesterone respectively) of control treatments. In this group of rats methanolic extract caused significant decreased (14.90±3.16ng/ml or 25.12% of prolactin 2.40±0.03ng/ml or 24.29% of estrogen and progesterone) relative to 19.90±3.16ng/ml or 3.17±0.09ng/ml and 320.60±3.19ng/ml of prolactin, estrogen and progesterone respectively relative to control lactating rats. When compared with methanol extract of Aframomum meleguetia, bromocriptine showed significant decrease in serum levels of prolactin by (29.13% and 65.34% for progesterone but 0.00% changes for estrogen) relative to control rats. Ethylacetate extract in relation to bromocriptine, the result revealed 22.40±1.60ng/ml or 52.86% increase in
serum prolactin 2.17±0.06ng/ml or 10.00% decrease in estrogen and 323.42±1.68ng/ml or 0.00% progesterone relative to (19.90±3.16ng/ml prolactin, 3.17±0.09ng/ml of estrogen and 320.60±0.19ng/ml progesterone) control rats. When compared within themselves ethylacetate extract produced 22.40±1.60ng/ml or 33.48% increase in prolactin 2.17± 0.06ng/ml or 9.58% decrease in estrogen and 323.42±1.68ng/ml or 160% increase in progesterone relative to methanol extract treated rats of 14.90±3.16ng/ml or prolactin, 2.40±0.03ng/ml estrogen and 318.00±2.26ng/ml progesterone.

Table II: Treatment effects on lactogenic hormones of lactating rats

<table>
<thead>
<tr>
<th>Lactogenic hormones</th>
<th>Treatment schedule</th>
<th>Bromocriptine 0.11mg/kg b.w</th>
<th>Normal control 5ml/kg b.w</th>
<th>Methanol extract 100mg/kg b.w</th>
<th>Ethylacetate extract 94mg/kg b.w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactin NLF</td>
<td>2.68±0.06</td>
<td>19.90±3.16</td>
<td>3.17±0.09</td>
<td>14.90±3.16</td>
<td>2.40±0.03</td>
</tr>
<tr>
<td>Estradiol NLF</td>
<td>2.41±0.08</td>
<td>320.60±0.19</td>
<td>318.00±2.26</td>
<td>2.17±0.06</td>
<td>2.45±0.10</td>
</tr>
<tr>
<td>Progesterone NLF</td>
<td>110.21±1.15</td>
<td>110.21±1.15</td>
<td>110.21±1.15</td>
<td>110.21±1.15</td>
<td>110.21±1.15</td>
</tr>
</tbody>
</table>

Mean is ± SD. Mean values with n, b, or m superscripts are significantly different from normal control, bromocriptine and methanol extract of Aframomum melegueta seed.

Presented in Table III below is the serum concentration of lactogenic hormones (prolactin, estrogen and progesterone) in non-lactating and lactating rats. Bromocriptine produced significant but marginal reduction in serum levels of prolactin, estradiol and progesterone of lactating rats relative to non-lactating rats. The normal control treatment revealed a significant reduction in serum levels of prolactin in lactating rats relative to non-lactating rats but no significant changes were observed in serum levels of estradiol and progesterone of lactating rats relative to non-lactating rats. Methanol extract of Aframomum melegueta seed caused a significant reduction in serum levels of prolactin and estradiol but no significant changes in serum levels of progesterone of lactating relative to non-lactating rats. Ethylacetate extract of Aframomum melegueta seed however produced a significant reduction only in serum concentration of estradiol but none in prolactin and progesterone of lactating rats relative to non-lactating rats.

Table III: serum concentration of lactogenic hormones in non-lactating and lactating rats.

<table>
<thead>
<tr>
<th>Lactogenic hormones NLF</th>
<th>Treatment schedule</th>
<th>Bromocriptine 0.11mg/kg b.w</th>
<th>Normal control 5ml/kg b.w</th>
<th>Methanol extract 100mg/kg b.w</th>
<th>Ethylacetate extract 94mg/kg b.w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactin NLF</td>
<td>12.56±0.89b</td>
<td>323.42±0.25</td>
<td>318.00±2.26</td>
<td>312.42±0.25</td>
<td>318.00±2.26</td>
</tr>
<tr>
<td>Estradiol NLF</td>
<td>12.56±0.89b</td>
<td>323.42±0.25</td>
<td>318.00±2.26</td>
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<td>318.00±2.26</td>
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<td>318.00±2.26</td>
</tr>
</tbody>
</table>

Mean is ± SD. Mean values with n, b, or m superscripts are significantly different from Normal control, bromocriptine and methanol extract of Aframomum melegueta seed.

Discussion

This study investigated antilactogenic hormones characteristics of methanol and ethylacetic extract of Aframomum melegueta seeds in Wister rats. The hormones of interest in this study are those of prolactin, estrogen and progesterone due to their involvement in imitating and maintenance of lactogenic activity of bromocriptine an ergot alkaloid which is known to lower down serum concentration of prolactin is used as standard or reference control. The result presented in table I revealed that bromocriptine caused a generalized reduction in the three hormones (prolactin, estrogen and progesterone) in this study. Bromocriptine as dopaminergic agonist acts via a specific receptor in the anterior pituitary tonically inhibits prolactin and lactotroph proliferation (Foord et al, 1983; Trouillas et al. 1994; Friedman et al. 1984) [16, 42, 19]. Bromocriptine produced approximately 62% reduction in serum concentration of prolactin relative to normal control treatment but only 8% marginal reduction in serum estradiol concentration relative to control treatment. This observation may not be unconnected with the fact that estradiol appears to be a potent anti-dopaminergic agent in vivo (Ferland et al, 1979) [14]. The decreased serum concentration of progesterone in rats administered with bromocriptine may not be attributed to decreased luteal phase progesterone concentration following prolactin suppression by bromocriptine as reported by Kauplia et al, 1988 signifying the involvement of other factors in this case. This observation is however in agreement with the report of Bohnet et al 1977; Kauplia et al, 1987 [6, 24], on over suppression of prolactin and isolated prolactin deficiency in a woman with puerperal alactogenesis. The mechanism by which prolactin participate in the regulation of ovarian functions in man are only partly understood. Progesterone prevents centrally the release of prolactin and casein synthesis in the presence of prolactin and the absence of progesterone at the level of mammary gland receptor (Deis et al, 1989) [11]. As long as both extracts have no significant reduction in serum concentration of progesterone, it may therefore be that the reduced level of prolactin is in part contributed by progesterone. The plausible explanation for the decreased serum level of progesterone can be backed up by the report of Robertson and King, 1974; Ash and Heap, 1975 [56, 4] who reported that a decline in progesterone concentration caused by the regression of copra lutea seems to play an essential role in parturition may extend into lactation in gilts. Suppression of serum prolactin by bromocriptine prevented the onset of lactation completely but has no obvious influence on changes of other hormone (Progesterone, relaxin etc.) concentration and course of parturition (Taverne et al, 1982) [19]. This goes to support the fact that the decreased level of progesterone in this study may not be solely on suppressed prolactin secretion by...
bromocriptine meaning other factors may have contributed. The significant decrease in serum prolactin caused by methanol extract of *Aframomum melegueta* (A.m) and ethylacetate extract of the same seed may be associated with presence of alkaloids in their phytochemistry as reported by Okwu 2005 [33], Okoli et al, 2007 [30]. Like bromocriptine (standard control drug) which is ergot alkaloid, methanol and ethylacetate extract of *Aframomum melegueta* produced in a similar manner reduction in a serum prolactin which may be attributed to the presence of alkaloid in the phytochemistry of *Aframomum melegueta* seeds. This finding is at variance with the report of Adefegha and Oboh 2012 [1] who failed to detect presence of alkaloid in their phytochemical screening of *Aframomum melegueta*. The solvent properties of methanol and ethyl acetate may have contributed to the differences in serum prolactin witnessed in this study. Polar protic property of methanol ought to have advantage in extracting more metabolites of A.M since these metabolites e.g [6]-paradol [6]-gingerol and shogoal contain (OH) bond in their structure when compared with polar aprotic property of ethylacetate. This however is not the case in this experiment as ethyl acetate extract reduce serum prolactin to 35.08% in non-lactating rats when compared with methanol extract that cause 21.3% reduction of the same hormone relative to normal control experiment this is not completely out of expectation especially as crude, extract was used in this research which contained both (OH) and (C-O) bonds. The anti estrogenic property of seed extract of A.M was observed in this study as both extracts caused significant decrease in serum concentration of estradiol with ethylacetate extract taking the lead. This observation is in consonance with the report of EL-Halawany et al. 2011 [13]; on screening for estrogenic and antiestrogenic activities of plant growing in Egypt and Thailand where A.M is the most antiestrogenic followed by Dalbergia candidatensis, then Dracaena loureiri and Mansonia gagei. There is a good evidence that consumption of plants’ secondary metabolites (poly-phenols, isoflavones, flavonoids) exhibits endocrine disrupting substance (EDS) properties (Grain et al. 2008) [20]. Most EDS with adverse effect on reproduction bind either to estrogenic receptors or androgen receptors and inhibit transcriptional mechanism (Wolfgang et al 2010) [44]. Most of them interfere with reproduction as either agonist or antagonist of the steroidal sex hormones which may be estrogen or androgen.

For serum progesterone, both extracts produce non-significant (p>0.05) alteration in non-lactating rats relative to normal control rat. This observation runs contrary to the report of Inegbenebor and Ebomoyi 2014 [23] on anti-progesterone effect of *Aframomum melegueta* on Sprague dawley rats that received 6.7mg/kg and13.3mg/kg body weight significantly decreased serum progesterone only on day 7 of gestation for the rat that received 6.7mg/kg but those administered with 13.3mg/kg body weight of saline crude extract of Alligator pepper decreased serum progesterone on both day7 and 21th of gestation.

The significant (P<0.05) reduction observed in serum prolactin level with lactating rats treated with bromocriptine in table II goes a long way to support the agonistic action of bromocriptine on dopaminergic D2 receptor which is anti-prolactinemic (Clive et al, 2002) [8]. The reduction in serum prolactin concentration by methanol extract of *Aframomum melegueta* seed in lactating rat relative to normal control equally gives credence to that of Okoli et al, 2007 [32] on the presence of alkaloid which is present in A.M that mimicks that of ergot alkaloid in bromocriptine. Okwu 2005 [33] also reported presence of alkaloid in *Aframomum melegueta* (A.M) both qualitatively and quantitatively. Bromocriptine (standard reference control) has a higher degree of reduction in serum prolactin when compared with methanol and ethylacetate extract. This finding is in consonance with the report of Ebong et al, 2000 [32] on lowering effect of Alligator pepper on serum prolactin in non-lactating wistar rats. There was no significant change in serum prolactin of rats that received ethylacetate extract of A.M relative to control rats. The antiestrogenic activity observed in this study in all the treatment relative to control is in consonance with the report of EL-Halawany et al. 2011 [33] on screening for estrogenic and antiestrogenic activities of plants growing in Egypt and Thailand and Ibekwe et al; 2018 [22] on antilactogenic effect of methanol extract of A.M on wistar rats. For serum progesterone,only the bromocriptine treated rats (0.11mg/kg) produced significant decrease in serum concentrations when compared with the rats that received methanol, ethylacetate extract and even the normal control rats. Both methanol and ethylacetate extracts of *Aframomum melegueta* produced no significant (p>0.05) changes in serum concentration of progesterone. This observation is not in agreement with the report of Inegbenebor and Ebomoyi 2014 [23] on effect of saline extract of Alligator pepper (Zingiberaeace A.M) on serum progesterone in pregnant Sprague Dawley rats.

The table III results shows that bromocriptine is a powerful antilactogenic agent seeing it can reduce serum prolactin even while the pups are suckling. In addition, the serum levels of estradiol and progesterone were equally reduced by bromocriptine in lactating rats’ relative to non-lactating rats. This in no doubt has given a strong support to its antilactogenicity. Lactation alone also brought a significant (P<0.05) reduction in serum prolactin as can be seen in table III. Lactogenesis or onset of milk synthesis is a 2-stage process in rats that occur during late pregnancy (Fleet et al, 1975) [13]. Lactogenesis I and II have been reported (Bussmann et al, 1996) [7]. Both stages of lactogenesis are controlled primarily in the rat by ovarian and adrenal steroid with lactogenic hormones from the placenta (Topper and Freeman, 1984). Some of these hormones have not been assayed in this investigation and may probably be part the discrepancies observed in the current study. There were however no significant changes in serum estradiol and progesterone in normal control rats with respect to non-lactating relative to lactating rats.

For the extract treated rats, methanol significantly reduced serum concentrations of prolactin and estradiol in lactating relative to non-lactating rats but no significant changes in serum levels of progesterone. This concurs with the reports of Mark, 2004 [29] on the use of A.M in reducing excessive lactation and postpartum haemorrhage in Ghana. Ethylacetate extract of A.M seed produced no significant reduction in serum levels of prolactin, estradiol and progesterone in lactating rats.

Here it may be likely that methanol extract wielded more influence in lactating rats when compared with ethylacetate extract bearing in mind its polarity index.. For the hormones investigated, the three hormones play crucial role in lactogenesis. Prolactin is the major generator of lactation competence during pregnancy and it functions directly via its effect on mammary gland epithelial cells and indirectly through its regulation of ovarian progesterone production, (Ibekwe et al, 2018) [22]. Prolactin in conjunction with
progesterone generate lobuloalveolar growth in mammary gland. Estrogen on the other hand stimulate mammary ductal growth which may be blocked by estrogen receptor antagonist tamoxifen (Daniel et al 1997; Silberstein et al 1994) [38]. Progesterone is responsible for extensive side-branching and alveologenesis required to create a lactation competent gland which occur in lactogenesis I.

In lactogenesis II, prolactin causes enlargement of mammary gland in pregnancy and increase milk secretion (Melmel and Kleinberg, 2008) [39]. During pregnancy, high circulating concentration of estrogen promote prolactin production (Daryl, 1996) [10]. Progesterone prevents centrally the release of prolactin and casein synthesis is initiated in the presence of prolactin and in the absence of progesterone (Deis et al 1989) [11]. Therefore progesterone withdrawal is the trigger hormone that initiates lactation.

**Conclusion**

Bromocriptine (standard reference drug) manifested clearly antilactogenicity by its ability to reduce significantly three selected hormones involve in lactogenesis I and II in lactating rats. Methanol crude extract of *Aframomum melegueta* seeds followed closely by bringing down the serum level of prolactin and estrogen in lactating rats. Its non-significant reduction in serum level of progesterone is perhaps a positive development because progesterone must be withdrawn in lactogenesis II for commencement of full lactation. The presence of progesterone is therefore antilactogenic. Ethylacetate extract of A.M seed at the dosage administered would not significantly influence any of the selected hormone involve in lactogenesis in lactating Wistar rats. While reduction in serum levels of prolactin and estrogen by the crude extract is antilactogenic, both extract at their specific dosage of administration portrayed antilactogenic characteristic with methanol extract taking the lead.

**References**

7. Busmann LE, Bussmann IM, Charreau EH. Role of receptor for epidermal growth factor and insulin like growth factor I and II in the differentiation of rat mammary gland from lactogenesis I to lactogenesis II.


