Antifertility Effect of Inhaled Cannabis Sativa on Male Wistar Rats

Udokang Nsikak Ephraim, Udom Utibe Godwin

Department of Physiology, University of Uyo, Uyo, Nigeria

Email address: nsikakudokang@yahoo.com (U. N. Ephraim)

To cite this article:

Received: November 30, 2018; Accepted: December 29, 2018; Published: January 29, 2019

Abstract: A total of forty (40) male wistar rats randomly divided into four groups of ten (10) rats per group were used for this work. Group 1 served as the control and was treated with distilled water, group 2 was the low dose group (exposed to cannabis smoke for 5 minutes daily), group 3 was the high dose group (exposed to cannabis smoke for 10 minutes daily) and group 4 was the high dose + ascorbic acid group (exposed to cannabis smoke for 10 minutes and orally gavaged with ascorbic acid at 2.8mg/kg body weight daily). The rats were weighed weekly and the administration lasted for 28 days. The animals were sacrificed on the 29th day and blood was collected by means of cardiac puncture. Serum was obtained for hormonal assay. The result showed a significant increase in the body weight (p>0.05) in the high dose group compared to low dose and high dose + ascorbic acid group, while the control group recorded the least body weight gained. There was significant decrease in testicular weight (p>0.05) in the experimental groups compared to the control group. Prolactin level was significantly increased (p>0.05) in the high dose group compared to low dose and high dose + ascorbic acid group, while the control group recorded the least body weight gained. There was significant decrease in testicular weight (p>0.05) in the experimental groups compared to the control group. Testosterone level was significantly increased (p>0.05) in the high dose + ascorbic acid group compared to the other groups. Estradiol level was significantly decreased (p>0.05) in the high dose group compared to the other groups. Follicle Stimulating Hormone (FSH) level was significantly increased (p>0.05) in the high dose and high dose + ascorbic acid group compared to the other two groups. Luteinizing Hormone (LH) level increased significantly (p>0.05) in the control and high dose groups compared to the other two groups. From this result, it can therefore be concluded that cannabis sativa decreases testicular weight, estradiol and testosterone level and increases body weight, FSH, LH and prolactin level in male wistar rats. Also that ascorbic acid (vitamin c) was able to ameliorate some of these effects.

Keywords: Cannabis Sativa, Ascorbic Acid (Vitamin C), Antioxidant, Oxidative Stress, Antifertility

1. Introduction

Cannabis specifically refers to the green, brown, or gray mixture of dried, shredded leaves, stems, seeds, and flowers of the cannabis plant [1]. It is also called Marijuana. It is the most widely used illicit drug worldwide and became the second most commonly smoked substance after tobacco, with an estimated 160 million users (3.8% of the world’s population of 15-64 year olds) [2].

Though Cannabis is consumed by various routes, it is mostly smoked as cigarette [3]. While some people use it as an ingredient in foods, it has also been made available as beer.

Cannabis contains many chemicals, with Δ9-tetrahydrocannabinol (Δ9-THC) and Cannabidiol (CBD) being the main ingredients of Cannabis sativa plant and having distinct symptomatic and behavioral effect [4].

Two cannabinoid receptor systems have been discovered, cannabinoid 1 receptor (CB1R) in the brain specific for Δ9-THC [5] and cannabinoid 2 receptor (CB2R) discovered by [6]. CB1Rs are mainly in the brain, particularly in the substantia nigra, the basal ganglia, limbic system, hippocampus and cerebellum, but are also expressed in the peripheral nervous system, liver, thyroid, uterus, bones and testicular tissue [7-9]. CB2Rs are mostly expressed in immune cells, spleen and the gastrointestinal system, and to some extent in the brain and peripheral nervous system [9, 10].

Studies have demonstrated that cannabis and its active component have widespread effects on multiple hormonal systems, including gonadal, adrenal, prolactin, growth hormone, and thyroid hormone regulation in experimental models [11].

Cannabis use is on the increase as many countries have
decriminalized it. As such, interest has been drawn to the
effects in which inhaled cannabis may have on the
reproductive system.

The delay in conception as observed in couples in recent
days compared to previous generation coincides with the
increase use of cannabis [12]. Its use tends to be more
prevalent among youths aged<35 years [13] and males
appear to be more likely to use cannabis [14].

Following the above facts, it is of essence to investigate the
harmful effects of cannabis as relates to fertility and to
give an account of such with possible remedy.

1.1. Purpose of the Study

With the existence of cannabinoid receptors in the brain and
gonads, it is also likely that cannabis can alter the sex hormone
levels because of its direct action on these organs which
synthesize and regulate these hormones and possibly affect
fertility. For this reason, this study soughts to investigate the;

i. Antifertility effect of inhaled cannabis on male wistar
rats using body weight, testicular weight and sex hormone
levels as the criteria.

ii. Potency of ascorbic acid in ameliorating the inhaled
cannabis effects.

1.2. Significance of the Study

With increasing number of countries decriminalizing the
use of cannabis, this study so to explore its antifertility
effects, if any and the potency of ascorbic acid in reducing
these effects.

2. Methodology

2.1. Drugs and Animal Collection

2.1.1. Cannabis Sativa

An approval and sample of cannabis sativa was gotten
from The State Command, National Drug and Law
Enforcement Agency (NDLEA), Akwa Ibom State
Command, Uyo, for the study.

2.1.2. Ascorbic Acid (Vitamin c)

A packet of Emzor 100mg vitamin c tablets was bought
from Leadsons pharmacy in Uyo, Akwa Ibom state.

2.1.3. Animals

Prepubertal rats of the Wistar strain (60-80g) were
obtained from the animal house of the faculty of Basic
Medical Sciences, University of Uyo and kept in wooden
cages of 50 x 30cm dimension in a well-ventilated section of
the same animal house for acclimatisation. The animals were
fed with rat chow (vital feed) and were allowed free access to
drinking water while the experiment lasted.

2.2. Experimental Design

A total of forty (40) prepubertal male albino rats were
randomly designed into four (4) groups of ten (10) animals
per group with group 1 as control group. Group 2, 3 and 4
served as experimental groups. Group 2 rats (low dose
cannabis) were exposed to cannabis smoke through
inhalation for 5minutes a day. Group 3 rats (high dose
cannabis) were exposed to cannabis smoke through
inhalation for 10minutes a day. Group 4 (high dose cannabis
+ ascorbic acid) was exposed to cannabis smoke through
inhalation for 10minutes a day and ascorbic acid. Group 1
(control group) were given distilled water and feed
throughout the experimental days.

2.3. Administration

2.3.1. Cannabis Sativa

Daily, animals from group 2, 3 and 4 were exposed to
cannabis smoke. This was done by putting 1g of the cannabis
rolled with rizla rolling paper in a red-hot charcoal in a
stainless plate and placing it in an airtight smoking chamber
with 10rats per smoking session for five (5) minutes. The
procedure was repeated for the group 3 and 4 for another five
(5) minutes each day, making the total time of exposure of
this two groups (3 and 4) to be ten (10) minutes each day.
The air-tight smoking chamber was made from polythene
plastic cage of 60cm×50cm×40cm dimension.

2.3.2. Ascorbic Acid

After the cannabis smoke exposure, the group 4 rats were
orally gavaged with 2.8mg/kg body weight of ascorbic acid.

2.4. Sample Collection and Analysis

2.4.1. Sample Collection

After 28 days of administration, the rats were
anaesthetized using chloroform and sacrificed. Blood was
collected by cardiac puncture into plain bottles for
hormonal analysis. The testis was also collected and weighed. Approval
was gotten from the Local Research Ethical Committee of the
University of Uyo, Uyo, Akwa Ibom State, Nigeria.

2.4.2. Determination of Hormonal Assay

The collected blood was allowed to clot and centrifuged at
300rev per minutes for 20mins. The serum was collected
with the aid of a micropipette and hormone levels determined
by ELISA (Enzyme-linked immunosorbent assay) method
using ELISA hormonal kits as follows:

1. Prolactin
   i. All reagents, samples and calibrators were prepared;
   ii. 25 µl of each calibrator was pipetted into control and
      samples into the wells prepared.
   iii. 50 µl of Rat Prolactin Sample Buffer was added to
      every well.
   iv. The mixture was shaken for 2 hours at room
      temperature (18 - 28°C).
   v. The content of the wells was discarded and the wells
      were washed 4 times with 300 µl buffered wash solution. The
      microplate was beaten carefully to remove as much wash
      solution as possible.
   vi. 200 µl of Enzyme-Labeled Anti-Rat Prolactin Antibody
      was added to all wells.
   vii. The mixture was shaken again for 1 hour.
viii. The content of the wells were discarded and washed 4 times with 300 µl buffered wash solution.

ix. The microplate was beaten carefully to remove as much wash solution as possible.

x. 200 µl of liquid Substrate Solution was added to all wells.

xi. The solution was incubated without shaking for 30 minutes in the dark.

xii. 50 µl of Stop Solution was added to each well and mixed carefully.

xiii. The optical density at 450 nm was read. Bi-chromatic measurement with a reference at 600-690 nm was taken.

xiv. A 4-Parameter-Fit with log-log coordinates for optical density (linear scale) and concentration (logarithmic scale) was used to calculate the result.

ii. Testosterone and Estradiol

i. All reagents, samples and calibrators were prepared.

ii. 100 µL calibrator or sample was added to the appropriate well in the antibody pre-coated microtiter plate. 100 µL of PBS (pH 7.0-7.2) was added in the blank control well.

iii. And then conjugate was added immediately. It was shaken, mixed and incubated for 1 hour at 37°C.

iv. It was aspirated and washed 5 times.

v. 50 µL of substrate A and 50 µL substrate B was added to each well including blank control well and incubated for 10-20 minutes at 37°C.

vi. 50 µL Stop Solution was added to each well including blank control well and mixed well.

vii. The optical density was read at 450 nm using a microplate reader immediately.

vi. The content of the wells were discarded and washed 4 times with 300 µl buffered wash solution.

x. 200 µl of liquid Substrate Solution was added to all wells.

xi. The solution was incubated without shaking for 30 minutes in the dark.

xii. 50 µl of Stop Solution was added to each well and mixed carefully.

xiii. The optical density at 450 nm was read. Bi-chromatic measurement with a reference at 600-690 nm was taken.

xiv. A 4-Parameter-Fit with log-log coordinates for optical density (linear scale) and concentration (logarithmic scale) was used to calculate the result.

II. FOLLICLE STIMULATING HORMONE (FSH) AND LUTEINIZING HORMONE (LH)

i. All reagents, samples and calibrators were prepared.

ii. 50 µL calibrator or sample was added to each well. And then prepared Detection Reagent A was added immediately. It was shaken and mixed and incubated for 1 hour at 37°C.

iii. It was aspirated and washed 3 times.

iv. 100 µL prepared Detection Reagent B was added and incubated for 30 minutes at 37°C.

v. It was aspirated and washed 5 times.

vi. 90 µL Substrate Solution was added incubated for 10-20 minutes at 37°C.

vii. 50 µL Stop Solution was added and read using a microplate reader at 450 nm immediately.

viii. Regression analysis was used to determine the result.

2.4.3. Statistical Analysis

Data obtained were analyzed using Mean, Standard Error of Mean and Analysis of Variance followed by Duncan’s test which was used to determine the direction of significance. The level of weight and hormones were reported in the form mean± SEM and statistical significance was established at 0.05 level of significance with p<0.05 signifying significance. Data were analyzed using the Statistical Package for Social Sciences (SPSS version 22.0) and GraphPad Prism 5.0.

3. Results

Table 1. Weight.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group 1 (Control)</th>
<th>Group 2 (Low dose cannabis)</th>
<th>Group 3 (High dose cannabis)</th>
<th>Group 4 (High dose cannabis + Ascorbic acid)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight gained (g)</td>
<td>35.50±1.89</td>
<td>46.53±4.44</td>
<td>53.17±3.67</td>
<td>46.35±3.8</td>
</tr>
<tr>
<td>Testicular weight (g)</td>
<td>3.17±0.19</td>
<td>2.54±0.09</td>
<td>2.06±0.12</td>
<td>1.90±0.35</td>
</tr>
</tbody>
</table>

Values reported in the form Mean ± SEM. Similar superscript letters mean not significantly different (p>0.05) while different superscript letters mean significantly different (p<0.05).

Result reveals that exposure to high dose of Cannabis caused a significant increase in weight than in groups exposed to low dose, high dose of Cannabis with ascorbic acid and the control group(p<0.05). There was a significant decrease in testicular weight in the group treated with high dose of Cannabis (group 2) and those treated with high of dose Cannabis combined with Ascorbic acid but between the control and the group exposed to low dose of Cannabis, there were no significant difference in their testicular weight(p>0.05) (Table 1).

Table 2. Hormonal Assay.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group 1 (Control)</th>
<th>Group 2 (Low dose cannabis)</th>
<th>Group 3 (High dose cannabis)</th>
<th>Group 4 (High dose cannabis + Ascorbic acid)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactin(ng/ml)</td>
<td>0.06±0.002</td>
<td>0.06±0.003</td>
<td>0.07±0.001</td>
<td>0.05±0.002</td>
</tr>
<tr>
<td>Estradiol(pg/ml)</td>
<td>2.15±0.08</td>
<td>2.15±0.09</td>
<td>1.85±0.03</td>
<td>2.23±0.05</td>
</tr>
<tr>
<td>Testosterone(ng/ml)</td>
<td>0.87±0.15</td>
<td>0.23±0.07</td>
<td>0.30±0.04</td>
<td>1.13±0.26</td>
</tr>
<tr>
<td>FSH (miu/ml)</td>
<td>0.04±0.01</td>
<td>0.42±0.07</td>
<td>0.47±0.16</td>
<td>0.19±0.05</td>
</tr>
<tr>
<td>LH(miu/ml)</td>
<td>2.50±0.41</td>
<td>0.47±0.16</td>
<td>2.88±0.24</td>
<td>1.08±0.07</td>
</tr>
</tbody>
</table>

Values reported in the form Mean ± SEM. Similar superscript letters mean not significantly different (p>0.05) while different superscript letters mean significantly different (p<0.05).
Result presented in Table 2 reveals that exposure to Cannabis caused a significant increase in Prolactin level, FSH and LH (p<0.05) but a significant decrease in Estradiol and Testosterone (p<0.05) in the group treated with high dose of Cannabis. In Group treated with combination of high dose of Cannabis and Ascorbic acid, there was a significant increase in testosterone only (p<0.05) while for other hormones, an insignificant effect was established (p>0.05). Result also shows that exposure to low dose of Cannabis caused a significant increase in FSH and LH and a significant decrease in testosterone (p<0.05) but no effect on Prolactin and Estradiol (p>0.05) (Table 2).
4. Discussion

Inhaled cannabis interferes with normal body function as observed in the deviation of the tested parameters in the experimental groups from the control group. This deviation may be caused probably by the effects of intoxication by cannabis compounds; it can also be due to oxidative stress-induced damage since cannabis smoke generates free radicals (reactive oxygen species, ROS) which cause oxidative stress, [15]; or by the combined action of these two factors.

Reactive oxygen species (ROS) are reported to be tumorigenic [16] and can also induce cellular senescence and cell death [17]. According to [18] whether ROS causes tumor or apoptosis depends on the cell and tissues, the location of ROS production, and the concentration of individual ROS.

There are claims that cannabis and its chemical components appear to regulate feeding behavior. [19] state that the endocannabinoids are important mediators and metabolic regulators in mammalian physiology, with diverse and ubiquitous modulating actions, including the regulation of body weight. Agonism of CB1 receptors is known to stimulate feeding behavior [20]. According to [21], stimulation of the CB1 receptors in the mammalian cannabinoid system specifically increases food cravings and promotes the deposition of energy as fat into adipose tissues. In this work, the increase in weight gained (figure 1) also support those claims, as such, we can say that cannabis smoke increases appetite and causes increase in body weight.

A number of animal studies have reported direct effects on various reproductive organs and this is observed in the reduction in testicular weight (figure 2) in this study. Hypogonadism was also reported by [22]. The decrease in testicular weight is an indication of the direct action of cannabis smoke on the testis inducing apoptosis through oxidative stress.

The hormonal assay of this work changes greatly among groups depending on the hormone (table 2). These changes have been observed to show a pattern, as pituitary gland hormone (prolactin, FSH and LH) levels increase while organ hormone (testosterone and estradiol) levels decrease.

The decrease in estradiol and testosterone levels (figure 4 and 5 respectively) is possibly supporting that fact that cannabis has direct effect on the testis as seen in the reduction in its weight. This effect has also been observed by many including [23], who proposed that chronic cannabis use reduces levels of circulating testosterone. The significant increase in the level of testosterone in the ascorbic acid group (figure 5) supports the finding by [24] that ascorbic acid increases testosterone levels, but the mechanism of its action is not well understood.

The effect of cannabis on prolactin secretion varies a great deal in different researches, though there are well documented evidences of cannabis causing increase in prolactin. One of the studies was by [25], who concluded that ∆9-THC causes increase in prolactin in male rats.

Although most studies show that cannabis reduces the level of FSH, the result obtained in this work shows an increase in the levels of FSH (figure 7) and this is consistent with the other pituitary hormones (prolactin and LH) (figure 3 and 6). This may be due to a stimulating effect of inhaled cannabis smoke on the pituitary gland. Secondly, it may also be as a result of ROS-induced tumor in the pituitary gland caused by oxidative stress resulting in hypersecretion. Thirdly, it may be due to a feed-back effect of testosterone on...
the pituitary gland; this is specific in all the groups, as seen that the groups with low testosterone levels have increase LH levels and vice versa. The increased pituitary hormones secretion was unable to increase testosterone level since the leydig cells of the testis must have been severely destroyed by oxidative stress-induced apoptosis.

From the result obtained, ascorbic acid (an antioxidant) was observed to reduce the effect of cannabis in most of the measured parameters.

5. Conclusion

On the basis of this study, it can be postulated that inhalation of Cannabis sativa smoke can be anti-fertile probably due to its direct action on the testis and the pituitary gland. It causes decrease in testicular weight, testosterone and estradiol levels. It increases pituitary hormones possibly by the action on the pituitary gland in either of the following ways; direct stimulation, feedback mechanism from testosterone or by causing tumor growth resulting in hypersecretion.

It can also be assumed that cannabis smoke may be tumorigenic in the brain and apoptotic in other body organs.

Ascorbic acid (vitamin c) increases testosterone level and remedies the antifertility effects in male rats except in testicular weight in which the reduced weight was not ameliorated.

The antifertility effects of cannabis pose serious threat to the society. Considering the delay in conceptions in couples in recent days and the fact that many countries have decriminalized the use of cannabis, it can be suggested that in decades, the human race may be facing a very serious threat of extinction.

Therefore, it is necessary for the cultivation and use of this cannabis to be regulated by an authorised agency and researches on its effects be an area of interest to scientists.

Acknowledgements

We would like to appreciate the efforts of The State Commander, National Drug and Law Enforcement Agency (NDLEA), Akwa Ibom State Command for giving us the approval and sample of Cannabis sativa for this work.

References


