THE EFFECT OF INDONESIAN VELVET BEAN EXTRACT ON THE FERTILITY OF ALBINO MALE MICE

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ABSTRACT

Objective: This research aims to determine the dose of Indonesian velvet bean (Mucuna pruriens) extract which has significant effect on the fertility of albino male mice [concentration, motility and morphology of mouse sperm]. The extract is expected to be an alternative infertility herbal medicine relatively more secure and affordable replacing synthetic hormonal drugs which tend to have negative effects.

Methods: The seed was taken from Yogyakarta Indonesia. Fertility test was done to fertile adult male albino mice 12 w old, weighing 25-35 grams. Fertility tests performed on seven groups of mice; i.e. negative control, positive control and treatment groups (five dose levels at 50, 100, 150, 200 and 250 mg/kg body weight). Subsequent fertility test results were statistically tested, including tests of normality (Kolmogorov-Smirnov) followed by T test (Independent-Samples T Test).

Results: The sperm concentration and motility increased as an increased dose of seed extract was admistered, as well as decreased abnormal morphology. The highest change in the quality and quantity of sperm occurred at the dose of 250 mg/kg body weight with increased sperm concentration of about 22 million, sperm motility increased by 18% and decreased abnormal sperm morphology by 12%. Statistical analysis showed a significant increase in sperm concentration and motility at doses of 100, 150, 200 and 250 mg/kg body weight, whereas a significant decrease of abnormal sperm morphology was found at doses of 150, 200 and 250 mg/kg body weight.

Conclusion: Based on this study it is concluded that Indonesian velvet beans seed extract can increase the fertility of albino male mice significantly at dose level 250 mg/kg body weight.

Keywords: Mucuna pruriens, L-dopa, Sperm Concentration, Sperm Motility, Sperm Morphology

INTRODUCTION

Modern lifestyle and certain environmental exposure may have caused male infertility [1]. Various factors can directly or indirectly lead to this sexual dysfunction [2-5]. These days, the number of male infertility problem keeps increasing across the world. Various types of modern medicine have been applied to solve this problem but many of which yield negative effects [6-7].

Infertility problem has been a global issue [8]. Various studies had been done to investigate whether there is a reduction of quality in men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9]. Many researches proved that there is a drop of density in human sperm. A study comparing the quality of men’s fertility in the last 40 y [9].

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Various researches have been conducted to restore male infertility [13-15]. Some endocrine substances such as cytomet thyroid, glucocorticoid, androgen, mesterolone, gonadotropin, and clomiphene citrate of low dose have been tested to restore male fertility. In addition to these endocrine substances, many types of herbal medicine have also been used to reinitate this fertility problem. Believed to have relatively lower negative side effects, herbal is more preferable than synthetic medicine [16-18].

One potential plant to solve this fertility problem is velvet beans (Mucuna pruriens) [19]. It has been identified that velvet bean can regulate steroidogenesis and improve the quality of infertile male sperm [20]. The extract of velvet beans is known to be able to increase the quantity and motility of male mice sperm and decrease its abnormal morphology. This extract can affect the morphology of male mice reproduction organ; testis and epididymis. The testis of a male mouse which has been given velvet bean extract is known to have increased tubular diameter and spermatoid quantity [21]. L-Dopa is an important component of velvet beans [22]. As previously studied, L-dopa and other substances found in velvet beans have been proven effective in improving the sexual performance of albino male mice [23]. It is also reported that L-dopa can increase libido and thus improve sexual behaviour [24].

The content of L-dopa in velvet beans may be varying depending on the origin and environmental conditions. The Indonesian velvet beans, especially originating from Bantul district, have L-dopa content of 7.56% [25]. Despite the numerous studies on the ability of L-dopa in velvet beans in increasing the fertility of men, however L-dopa in velvet beans from Indonesia has not been well investigated on their ability in increasing the fertility of men, as well as the dose which give the significant effect. The recent study is conducted to provide a more comprehensive understanding of the effect of Indonesia velvet beans extract on the fertility of male albino mice (Mus musculus), especially on the concentration, motility, and morphology of sperm.

MATERIALS AND METHODS

Material

Sample or material used in this research is the seed of velvet beans which originates from Yogyakarta Indonesia, whereas animals were obtained from the Biotechnology Laboratory of Institut Teknologi Bandung (ITB).

Sample preparation

The chosen velvet beans were cleaned, peeled, dried up, and then mashed using a grilling machine. The velvet bean powder was then...
extracted using the maceration method (3 x 24 h) with water and ethanol (1:1) as the solvent and citric acid added until the pH reached 3 in room temperature. Every 24 h, the samples were filtered and re-macerated. The addition of citric acid was done to increase the solubility of the main compound in velvet bean, L-dopa. The solvent in the velvet bean macerate was then evaporated using the rotary vacuum evaporator. The evaporated macerate was then dried up using the freeze dryer, resulting brown-black dry velvet bean extract in 1.58%.

**Determining the level of L-dopa**

The percentage of L-dopa identified uses high performance liquid chromatography (HPLC) analysis [26]. This step started with making a standard solution of L-dopa with concentrations 25, 50, 75, 100 and 125 ppm. The velvet bean extract was tested using HPLC with testing parameter of $\lambda = 280$ nm, speed flow 1 ml/minute and solvent ratio of $H_2O$: Methanol: H$_3$PO$_4$, that is 97:20:1 [27].

**Animal treatments**

In this study, 45 male mice (12 w old) weighing about 25-30 g were used for the investigation. The mice were acclimatized to an average temperature of 23-29 °C for seven days so that these tested animals could adapt themselves to their new environment during the treatment. They were grouped into cages of 30x20x12 cm$^3$ based on treatment with the five mice per cage. During the acclimatization period, these animals were fed following PC551 standard and water ad libitum.

All the animals were divided into seven groups of five each and treated as follows: Group I: negative control; group II: positive control (L-dopa treatments, 50 mg/kg); group III-VI: administration of velvet beans extract (50, 100, 150, 200, 250 mg/kg). Velvet beans extract was administered for 30 d, through oral intake, once per morning. Each mouse of treatment group received certain doses of velvet beans extract whereas that in positive control group was given pure L-dopa concentration of 50 mg/kg body weight. The amount of velvet beans extract given was 0.3 ml/day for each concentration.

**Assessments the sperm concentration**

Neck-dislocated died mice were located on a tray for surgery. Their cauda epididymis was isolated using phosphate buffered saline (PBS). Sperm liquid was emptied from cauda epididymis by a syringe before it was well shaken. The calculation of sperm concentration was done before it was well shaken. The calculation was conducted for five boxes of counting chamber (a total of 25 boxes) for each sample, prior to average calculation. The result of calculation is the sperm concentration in $10^4$ sperm count average /mm$^3$.

**Assessments the motility of the sperm**

Sperm motility was observable from sperm suspension dropped on neubauer counting chamber observed by a microscope of 1000x magnifications. Sperm motility is valued on the basis of percentage of good sperm motility, that is, sperm which moves fast, straight forward and active [28].

**Assessment the morphology of the sperm**

Sperm morphology is observable with colouring eosin Y 1%. Sperm morphology testing was conducted by differentiating the shape of normal and abnormal sperm of 100 sperms observed before it was made into percentage [28]. Abnormal sperm includes abnormality such as broken, detached and thin head; broken, crooked and droplet cytoplasm middle part; or broken, crooked and coil tail. The observation used a microscope of 400x magnifications.

**Analysis of the data**

Results are expressed as means±SD for 5 animals per group. The normality of the data acquired was then tested using the 'T' test (Independent-Samples T Test). The Software SPSS 20 was used for the analysis.

**RESULTS**

**The level of L-dopa**

The chromatogram of HPLC indicated that the peak standard L-dopa appears at the retention time (RT) of 3.44. The seed extract chromatogram showed seven peaks at different retention period. The obtained data show that percentage of L-dopa in the Indonesian velvet bean extract was 13.9%.

**The effect of the velvet bean extract on sperm concentration, motility, and morphology**

Mice sperm concentration was observed on the 31st day after the application of velvet beans extract for 30 d. The result of this observation can be seen in table 1.

### Table 1: The experimental results of mice sperm concentration

<table>
<thead>
<tr>
<th>Mice groups (n=5)</th>
<th>Treatments</th>
<th>Mice sperm concentration (x10^4/ml)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Negative control</td>
<td>348±1.91*</td>
</tr>
<tr>
<td>II</td>
<td>Positive control</td>
<td>372±3.25</td>
</tr>
<tr>
<td>III</td>
<td>Treatment group (50 mg/kg body weight)</td>
<td>394±4.00</td>
</tr>
<tr>
<td>IV</td>
<td>Treatment group (100 mg/kg body weight)</td>
<td>419±1.43**</td>
</tr>
<tr>
<td>V</td>
<td>Treatment group (150 mg/kg body weight)</td>
<td>436±1.91**</td>
</tr>
<tr>
<td>VI</td>
<td>Treatment group (200 mg/kg body weight)</td>
<td>495±1.22**</td>
</tr>
<tr>
<td>VII</td>
<td>Treatment group (250 mg/kg body weight)</td>
<td>567±3.13**</td>
</tr>
</tbody>
</table>

*) Data given in mean±SD, **) Significantly different from control (p<0.05).

### Table 2: The experimental results of mice sperm motility

<table>
<thead>
<tr>
<th>Mice groups (n=5)</th>
<th>Treatments</th>
<th>Mice sperm motility (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Negative control</td>
<td>73.8±0.83</td>
</tr>
<tr>
<td>II</td>
<td>Positive control</td>
<td>76.0±1.58**</td>
</tr>
<tr>
<td>III</td>
<td>Treatment group (50 mg/kg body weight)</td>
<td>77.2±1.92**</td>
</tr>
<tr>
<td>IV</td>
<td>Treatment group (100 mg/kg body weight)</td>
<td>78.4±2.07**</td>
</tr>
<tr>
<td>V</td>
<td>Treatment group (150 mg/kg body weight)</td>
<td>81.2±2.38**</td>
</tr>
<tr>
<td>VI</td>
<td>Treatment group (200 mg/kg body weight)</td>
<td>86.8±1.92**</td>
</tr>
<tr>
<td>VII</td>
<td>Treatment group (250 mg/kg body weight)</td>
<td>91.0±1.58**</td>
</tr>
</tbody>
</table>

*) Data given in mean±SD, **) Significantly different from control (p<0.05).
The sperm morphology was presented in table 3.

<table>
<thead>
<tr>
<th>Mice groups (n=5)</th>
<th>Treatments</th>
<th>Abnormal sperm morphology (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Negative control</td>
<td>16.6±1.81</td>
</tr>
<tr>
<td>II</td>
<td>Positive control</td>
<td>16.2±1.64</td>
</tr>
<tr>
<td>III</td>
<td>Treatment Group (50 mg/kg body weight)</td>
<td>15.4±0.89</td>
</tr>
<tr>
<td>IV</td>
<td>Treatment Group (100 mg/kg body weight)</td>
<td>13.0±3.31</td>
</tr>
<tr>
<td>V</td>
<td>Treatment Group (150 mg/kg body weight)</td>
<td>11.0±1.22**</td>
</tr>
<tr>
<td>VI</td>
<td>Treatment Group (200 mg/kg body weight)</td>
<td>8.6±0.99**</td>
</tr>
<tr>
<td>VII</td>
<td>Treatment Group (250 mg/kg body weight)</td>
<td>4.6±1.34**</td>
</tr>
</tbody>
</table>

*) Data given in mean±SD, **) Significantly different from control (p<0.05)

**DISCUSSION**

The experimental results showed the velvet bean extract apparently succeeded in increasing the sperm concentration of the mice, in all dosage used in this research (50, 100, 150, 200 and 250 mg/kg). Another interesting phenomenon observed is that L-dopa was able to increase the sperm concentration of the mice, but a higher increase was observed in mice that were given the velvet bean extract. The concentration of L-dopa which was given to the positive control group was 50 mg/kg, equivalent to 1.75 mg, while based on the HPLC result showed that the concentration of L-dopa in the velvet bean extract in each dosage was equivalent to 0.22, 0.44, 0.88, 1.77 and 3.55 mg. This fact proved that the L-dopa contained in the velvet beans was not the only one that affected the sperm concentration of the mice.

The result of the normality test showed that the sperm concentration of the mice in all seven treatments has a larger significance value than the degree of freedom 0.05. Thus, H0 was accepted, which means that the data was normally distributed. Meanwhile, the result of the T test (Independent-Samples T Test) showed that the average of sperm concentration of the positive control group and the treatment group with the 50 mg/kg dosage did not have a significant difference to the control group. However, the treatment groups that were given the 100, 150, 200, or 250 mg/kg dosage showed a significant difference from the control group.

The increase in sperm concentration indicated the success in spermatogenesis process. This process is heavily affected by gonadotrophine hormones, luteinizing hormone (LH) and follicle stimulating hormone (FSH). The hormones stimulates testis, sex organ in men. Both hormones are produced by the cells in anterior pituitary, known as gonadotrophs. The secretion of these hormones is affected by GnRH (gonadotropin releasing hormone) that is secreted by the hypothalamus. Both hormones work on different part of the testis. LH works on Leydig cells to control the secretion of testosterone, while FSH works on tubulus seminiferosa, especially on sertoli cells to improve spermatogenesis.

The velvet bean extract treatment also succeeded in increasing the motility of the sperm in all dosage given (50, 100, 150, 200 and 250 mg/kg). The lightest increase happened on the 50 mg/kg dosage, while the largest increase happened on the 250 mg/kg dosage, with the increase of about 18%. In general, it is observed that in the dosage range of 50-250 mg/kg, the motility of the sperm and the dosage given correlates positively.

A living spermatozoa correlates highly to the motility of the sperm as being alive is an absolute requirement for a spermatozoa to be able to produce energy and move. Semen of a mammal that has high fertility is characterized with a high level of living spermatozoa with normal morphology. Good motility depends on many things, including the morphology of the spermatozoa.

The velvet bean extract treatment succeeded in reducing morphologically abnormal sperm in all dosage given (50, 100, 150, 200 and 250 mg/kg). The lightest reduction happened on the dosage 50 mg/kg, while the largest reduction happened on the dosage 250 mg/kg with a decrease of about 12%. In general, it is observed that in the dosage range of 50-250 mg/kg, the percentage of morphologically abnormal sperms and the dosage given correlates negatively. The sperms suffered different kinds of abnormality, such as coiled tail, curved tail, and loose head (fig. 1).

Sperm morphology depends on testosterone hormone secreted by Leydig cells. Disturbance in testosterone supply leads to dysfunction of epididymis, in which spermatozoa grows [29]. Given the significant role of testosterone in spermatogenesis process, its limited supply will disturb this process and cause primary or spermatogenesis abnormalities such as over-or under-sized head, coiled tail, double tail and others [30].

The study shows that Indonesian velvet beans extract can increase sperm concentration and motility as well as reduce its abnormal morphology. This increase in sperm quality and quantity is inseparable from L-dopa component found in Indonesian velvet beans. This compound not only increase sexual activity but also hormones regulating spermatogenesis process such as FSH and LH. L-dopa in human body is changed into dopamine by aromatic L- amino acid decarboxylase enzyme as a catalyst. As previously explored, psychological stress reduces dopamine and testosterone level. Increased dopamine in body can directly reduce psychological stress and stimulate brain to secrete hormone LH [31].

Increased sperm concentration and motility as well as decreased abnormal sperm morphology of mice occur because of L-dopa and other components contained in velvet beans extract which influence the secretion of testosterone. Secreted by Leydig cells, this hormone plays significantly in spermatogenesis process. The hormone belongs to steroid which is synthesized from cholesterol. The synthesis process of steroid and its derivatives is called steroidogenesis [32].

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**Fig. 1: Normal and abnormal sperm (Magnified 400x)**

Normal sperm, Coiled tail, Curved tail
Sperm motility is an important aspect in fertilization/insemination process. A large amount of sperm is not sufficient for the insemination to take place successfully without being supported by good sperm motility. Having consumed L-dopa substance contained in velvet beans extract, the mice have increased sperm motility and level of dopamine in brain. This dopamine stimulates hypothalamus to secrete GnRH. Dopamine-stimulated hypothalamus increases the secretion of hormone LH and FSH [33]. Hormone FSH regulates spermatogenesis and nutrition needed in spermatogenesis process. Increased FSH hormone secretion improves the quality of spermatogenesis because this hormone provides all required nutrition. One benefit of increased FSH secretion is the provision of energy required by sperm to move/motile. Sperm abnormality is caused by lack of FSH and LH in testis. Gonadotropin serum (FSH and LH) correlates with the growth of sperm morphology [34]. L-dopa contained in velvet bean is known to increase dopamine level. The dopamine can increase the secretion of FSH and LH so that they can increase the quality of sperm morphology in spermatogenesis process [35-36].

CONCLUSION

The result of the fertility test showed that the Indonesian velvet bean cotyledon extract can increase the concentration and motility of sperm and reduce abnormal morphology of the sperm in mice. The change of quality and quantity in the sperm happened at the highest dosage given, 250 mg/kg. The increase in the sperm concentration was about 22 million sperms, while the increase of sperm motility was about 18%, and the decrease in abnormal sperm morphology was about 12%.

ACKNOWLEDGEMENT

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CONFLICT OF INTERESTS

We declare that we have no conflict of interest.

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