The Motor Coordination Activity Of Aqueous Extract Of Withania Coagulans Fruits In Swiss Albino Mice By Rota Rod Test

Amit S. Kamdi, Devesh D. Gosavi, Suvarna M. Kalambe, and Pankaj N. Bohra

Abstract — Background: The various sedative and hypnotic medications used today have the central nervous system (CNS) depressant effects. A very little work has been done on the Withania coagulans – a vulnerable species as it is not found rampant in the world except in late seventies. Therefore, it was important to explore the CNS depressant activities of aqueous extract of Withania coagulans fruits in Swiss albino mice by using rota rod test.

Methods: Motor coordination was assessed by using the Rota Rod Test. The CNS depressant drugs decrease the endurance time of mice on the rota rod as they impair the motor coordination so that mice fall early on the rotating rod. This endurance time is statistically correlated among the control, standard and the test drugs.

Results: There was statistically highly significant (p-value <0.001) association observed between aqueous extract of Withania coagulans fruits with endurance time in Swiss albino mice on rota rod test.

Conclusion: The aqueous extract of Withania coagulans fruits demonstrated the CNS depressant activity in Swiss albino mice by rota rod test.

Index Terms — CNS depressant; rota rod test; Swiss albino mice (SAM); Withania coagulans.

I. INTRODUCTION
Insomnia is defined as the individual perception of trouble with sleep commencement, extent, consolidation, quality, which occurs in spite of adequate opportunity for sleep, thus results in daytime impairment [1]. Prevalence of insomnia in the world is 10-30% [2]–[4]. However, the prevalence of insomnia in India is 9% in the general population [5].

There are various drugs like benzodiazepines and barbiturates used for the treatment of insomnia. However, these drugs are not devoid of the side effects. The Rota rod test is widely used to evaluate the motor coordination in rodents [6]. When a mouse is repeatedly placed on a rod or cylinder which is rotating at a constant speed, the animal gradually learns to walk on it, adapting itself to the rotation speed. After ingestion of a central depressant, however, the animal easily falls from the rod. This test was first introduced by Dunham and Miya for assaying the drug effects on the motor activity [7]. Since then, the effects of various central depressants, investigated by this test have been reported [8]–[10].

Withania coagulans is a rare species, not found rampant in the territory. It is distributed in east of the Mediterranean region extending to South Asia i.e. Iran, Afghanistan, Pakistan (Sind and Baluchistan), Nepal and India. In India, it is found in the North-West region mainly Himachal Pradesh (Simla), Uttarakhund (Garhwal & Kumaon hills), Punjab, and Western Rajasthan (Barmer, Jaisalmer and Jodhpur districts) [11]. This plant is mainly used for the milk coagulation. Though lot of work has been done to explore its hypoglycemic activity, not much work is done to evaluate its action on the central nervous system [12]–[14]. In 1977 Budhiraja et.al. reported CNS depressant activity of this plant [15]. Thereafter this plant was not much explored for the CNS activity. Therefore, it was thought worthwhile to assess the CNS depressant action of this medicinal herb in Swiss albino mice by Rota rod test.

II. MATERIALS AND METHODS
A. Rota Rod Test Apparatus
The Rota rod consists of experimental compartments, with a common rotating rod of about 25 mm diameter with selectable speeds of approximately 5, 10, 15, 20 and 25 revolutions per minute. The interval counters are provided in each compartment. The apparatus works on 220/30 Volts, single phase, 50 Hz, AC. On the floor of each compartment there is a cantilever platform that is hinged at the rear end.

B. Rota Rod Test Procedure
The shaft’s angular velocity was adjusted by changing the position of the Drive Belt, from one pair corresponding grooves to another. The shaft rotated in anticlockwise direction and such the animal in general fell on the front side of the platform i.e. towards the free end of the platform. In our experiment the shaft’s angular velocity was adjusted to 25 revolutions per minute. On switching on, the counter showed some arbitrary reading as the counter was running i.e. counting. The counter was stopped by pressing the platform gently till the switch below it operated by making a click sound. The platform below it was raised gently making sure that the switch below it was released. The apparatus made ready to operate. Placing the mouse on the rotating shaft the RESET switch was simultaneously pressed. On pressing the RESET switch counter started counting the time in seconds.

Regardless of the previous reading on the counter, the counter started counting from “zero” only. When animal fell off the
rotating shaft on the platform, the platform was lowered down by
the falling impact-stopping the counter. This showed the
animal’s endurance time in seconds. Endurance time is
defined as the time period for which each mouse remained on
the rotating rod during test session [16]. To restart the counter
again for the next trial, the platform was lifted gently, animal
was placed on the rotating rod and the RESET switch was
pressed, so that the counter started. The Rota rod was
designed for the normal rat of around 150 gm in weight. Since
the mice weight was much lower than that of rats, the
difference of weight was added to the platform by putting
securing weight with the help of adhesive tape at the front end
of the platform. All the compartments were used
simultaneously.

C. Control, Standard and Test Drugs

Distilled water was given as vehicle for control. Diazepam
was used as the standard drug. The animals were treated 30
min before the experiment with the test drugs (WCFAqE of
200 mg/kg, 500 mg/kg and 1000 mg/kg doses p. o.).
However, the test drug was given every day for 30 days
throughout the period of experiment. The mice were observed
for 5 min. Recordings were done on Day 1, Day 8, Day 15,
Day 23 and Day 30 for all the groups. The recordings were
taken half an hour after drug administration to the respective
group. Drugs were given in the following manner:

Control: Vehicle (Distilled Water) 2 ml/kg p. o. once a day for
30 days.
Standard: Standard drug (Diazepam) 5 mg/kg i. p. half an hour before test.
AQ-200: WCFAqE 200 mg/kg body weight p. o. once a day for 30 days.
AQ-500: WCFAqE 500 mg/kg body weight p. o. once a day for 30 days.
AQ-1000: WCFAqE 1000 mg/kg body weight p. o. once a day for 30 days.

Where WCFAqE=Withania coagulans fruits aqueous extract.

D. Conflict of Interest

There is no conflict of interest among authors.

E. Source of Support

Mahatma Gandhi Institute of Medical Sciences (MGIMS),
Sevagram.

F. Ethical Approval

Study was approved by Institutional as well as the Animal
Ethics Committee of MGIMS, Sevagram.

III. RESULTS

As observed in the Table I, on Day 1 to Day 8 there was no
statistically significant difference in endurance time by mice.
However, on days 15, 23 and 30 endurance time by mice on
rota rod apparatus decreased highly significantly (p<0.001)
for all the three doses of 200 mg/kg, 500 mg/kg and 1000
mg/kg of WCFAqE compared to control. Furthermore, dose
response relationship was observed for these doses. To
conclude, this decrease in endurance time by test drug was
comparable to that of standard diazepam.

<table>
<thead>
<tr>
<th>Day</th>
<th>Control</th>
<th>Standard</th>
<th>AQ-200</th>
<th>AQ-500</th>
<th>AQ-1000</th>
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<tr>
<td>Day 1</td>
<td>18.66</td>
<td>17.16</td>
<td>20.16</td>
<td>19.33</td>
<td>18.83</td>
</tr>
<tr>
<td></td>
<td>±9.11</td>
<td>±8.08</td>
<td>±11.58</td>
<td>±8.35</td>
<td>±10.94</td>
</tr>
<tr>
<td>Day 8</td>
<td>17.83</td>
<td>12.50</td>
<td>16.33</td>
<td>15.16</td>
<td>14.66</td>
</tr>
<tr>
<td></td>
<td>±12.49</td>
<td>±5.82</td>
<td>±11.10</td>
<td>±11.23</td>
<td>±11.23</td>
</tr>
<tr>
<td>Day 15</td>
<td>23.16</td>
<td>8.66</td>
<td>10.33</td>
<td>9.66</td>
<td>9.00</td>
</tr>
<tr>
<td></td>
<td>±3.86</td>
<td>±2.58***</td>
<td>±1.75***</td>
<td>±4.13***</td>
<td>±1.41***</td>
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<tr>
<td>Day 23</td>
<td>21.33</td>
<td>5.83</td>
<td>8.16</td>
<td>6.83</td>
<td>5.50</td>
</tr>
<tr>
<td></td>
<td>±2.42</td>
<td>±1.94***</td>
<td>±2.56***</td>
<td>±1.94***</td>
<td>±1.37***</td>
</tr>
<tr>
<td>Day 30</td>
<td>19.50</td>
<td>3.50</td>
<td>6.00</td>
<td>5.00</td>
<td>4.50</td>
</tr>
<tr>
<td></td>
<td>±3.72</td>
<td>±1.04***</td>
<td>±1.78***</td>
<td>±1.78***</td>
<td>±1.51***</td>
</tr>
</tbody>
</table>

* p<0.05, ** p<0.01 and *** p<0.001 compared to control group.

IV. DISCUSSION

In our study as observed in Table I, on days 15, 23 and 30
endurance time by mice on rota rod apparatus “decreased”
highly significantly (p<0.001) for all the three doses of
200 mg/kg, 500 mg/kg and 1000 mg/kg of WCFAqE
compared to control. Furthermore, dose response relationship
was observed for this ‘decrease’.

Our study is exceptional one as there are no other studies
reported which have tested the effect of any of the extract of
Withania coagulans on mice using Rota rod apparatus. Yet,
Withania somnifera (WS) which is a similar species as that of
Withania coagulans (WC) had been tested for the motor
coordination using Rota rod apparatus in middle cerebral
artery occluded (MCAO) rodents. It was found that WS had
significantly (p<0.05) “increased” the performance of MCAO
rodents treated with WS root extract on Rota rod
apparatus [17]. Hence it can be concluded that the different
withanolides in both the species might be responsible for
this activity.

There are two types of Rota rod test. One is the constant
speed Rota rod, and another is the accelerating Rota rod [18].
In our study we used the constant speed Rota rod apparatus.
But it has few drawbacks. Rod’s diameter can influence the
performance of mice [19], therefore we used Rota rod of 25
mm diameter. The drug to be tested can affect the memory of
mice rather than the motor coordination [20], to eradicate this
bias we trained the mice vigorously before test. The constant
speed Rota rod are time consuming because the time limit
needs to be set which decreases the sensitivity of this test [20].
To eliminate this bias, we used the maximum velocity of
25 rpm on the Rota rod test which had a range of 5 to 25 rpm.
The maximum endurance time of mouse on this apparatus
was 38 seconds. This test can evaluate the muscle relaxant
property of any compound but does not really differentiate
between anxiolytics and neuroleptics. Usually, the central
depressant drugs have muscle relaxant activity. Therefore,
this test can be useful to assess the central depressive effect
of any drug [7].

It has been proved that the stimulants increase the
performance of mice on the Rota rod test so that mice take
longer time to fall from the rotating rod [10]. On the other
hand CNS depressants decrease the endurance time of mice
REFERENCES


V. CONCLUSION

In conclusion the test drugs WCFaqEs if administered in chronic settings might have CNS depressant activity on mice.

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