ROLE OF WITHANIA SOMNIFERA AND IT’S ACTIVE PRINCIPLE WITHANOLIDE-A IN BALANCE AND MOTOR CO-ORDINATION BY FOOT PRINT TEST

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ABSTRACT

Background: The basal ganglia historically been considered as a part of the motor system because of the variety of motor deficits that occur when they are damaged. But now it is considered as “extrapyramidal” motor system, and the disorders of basal ganglia are called extrapyramidal disorders. One type of symptoms that result from basal ganglia disorders is called as Huntington’s Chorea. As this disorder involves with symptoms like dyskinesias - abnormal involuntary movements, we felt it is necessary to protect our nervous system from such a disorder if possible in a painless regular fashion by a herb.

Materials and Methods: We used adult male Sprague Dawly rats for this study. Animals were divided into 5 groups and were given either Withania somnifera extract or the active component Withanolide A in different concentrations 10 days prior to lesion surgery and continued 5 days post surgery. The neuroprotective role of the drug employed was analyzed on the 5th day post lesion by using foot print test in a run way.

Result: The gait and balance of the animals were taken as a measure to analyze the protective nature of the striatum and so the activity of the drug employed here. The gait and balance of the LC animals were poor stating the unprotective nature of striatum. But the balance and gait of both drug group animals were comparatively better than the LC animals. That clearly stated the neuroprotective capacity of both the drugs used for this study.

Conclusion: Based on the observations and results we came to a conclusion that both the ethanolic extract and the active component withanolide A have the capacity in protecting the striatum and so can be used as a food supplement on a daily basis to protect our striatum. If needed further research can be conducted to analyze deep into the therapeutic effects of these herbal drugs.

KEY WORDS: Basal ganglia, Extrapyramidal, Huntington’s Chorea, Dyskinesias, Withania somnifera, Withanolide A, Gait, Striatum.

INTRODUCTION

The basal ganglia (BG) are a highly organized network, where different parts are activated for specific functions like movement control, associative learning, planning, working memory, and emotions [1] by secreting some inhibitory neurotransmitters. In general the basal ganglia...
are associated with a variety of functions, including voluntary motor control, procedural learning related to routine behaviors or “habits” such as bruxism, eye movements, and cognitive, emotional functions [2] by inhibition.

Huntington’s disease (HD) is a fatal neurodegenerative disorder that causes defects in behavior, cognition, and uncontrolled rapid, jerky movements. Huntington’s disease is associated with basal ganglia degeneration [3]. This degeneration of striatal neurons leads to hyperkinetic movement disorders characterized by increased uncontrollable motor functions due to inability to stop unwanted movement.

Huntington’s disease is also known as Huntington’s chorea because it is characterized by a continuous, choreiform movements of the body (Greek word for “dance”) Helmes and Shulman 2005 [4]. There is no cure for HD, and full-time care is required in the later stages of the disease [5]. The neuronal degeneration and other complications such as pneumonia, heart disease, and physical injury from falls reduce the life expectancy and eventually cause death within 10 to 20 years.

According to World Health Organization (WHO), 80 percent of the world’s population presently uses herbal medicine for some aspect of primary health care [6]. Ashwagandha, also called as “Queen of Ayurveda” is a very important plant in Ayurveda, the Indian traditional medicine. This herb was used 4000 years plus in India.

Soman 2012 investigated the effect of Withania somnifera (WS) root extract and Withanolide A (WD) in restoring spatial memory deficit by oxidative stress induced alteration in the hippocampus of epileptic rats and concluded that treatment with WS and WA has ameliorated spatial memory deficits by enhancing antioxidant system.

All the above said conditions paved ways to conduct a study to find the neuroprotective role of Withania somnifera and Withanolide A in experimental Huntington’s chorea rats by analyzing the gait difference of animals using a runway.

**MATERIALS AND METHODS**

**Animals:** We used adult male Sprague Dawly rats weighing (200–240 gm) for this study and maximum effort was taken to minimize the unwanted stress to the animals and to reduce the number of animal to be used for this study. Animals were divided into 5 groups with 6 animals in each group. They are control group (CO), lesion control (LC), sham control (SC), withania somnifera ethanolic extract 25mg/kg body weight (WS 125) and withanolide A 100mg/kg body weight (WD 100). The drugs were dissolved in normal saline and the volume was adjusted to 1ml. The administration of drug was started 10 days prior to lesion surgery so as to access the protective role of the drug in striate neurons. Dosages of drugs were given IP. around 10 O’clock every day. (Table-1)

**Preparation and administration of the drug:**

**Ethanol Extract of Withania Somnifera (WS)**

**Preparation:** The ethanolic extract was prepared by soxhlation method following Elayaraja et al., 2010(7). He proved that the ethanolic extract expressed more antioxidant activity than other extracts of withania somnifera. The I.P . dosage of this drug was prepared by dissolving the ethanolic extract of the drug in normal saline (Mohan et al., 2008(8)), the volume was adjusted to 1ml for each animal

**Withanolide A:** Withanolide A was purchased from Sigma Fluka-USA. The I.P . Oral dosage was avoided to reduce the wastage of drug and to minimize the stress condition of rats. The drug was started 10 days prior to lesion and 5 days post lesion so as to access the protective role and the therapeutic power of it in striatum. Drug administrating time was maintained around 10’ clock every day.

**FOOT PRINT ANALYSIS FOR GAIT (Carter et al, 1999 [9])**

Paw Print is a simple assay to measure gait analysis and balance.

**Apparatus (Figure-1)**

The runway apparatus was a elongated box with a walkable 50-cm-long, 10-cm-wide runway with 10-cm-high walls.

**Procedure**

**Trial:** One run per day per animal for one week was given as trial.

**Test:** The rats were acclimated in the testing room for 15 min

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The rats were held in supine position and the limbs were painted with a nontoxic watercolor paint, one color for fore paw and another for hind paw.

The animals were then placed in a clean runway lined with white paper and allowed to explore and run way. The white sheet with the paw print was taken, the first and the last paw prints were omitted and the remaining prints were analyzed for coordination of gait.

**Parameters**

- **Print length (PL):** Distance from the heel to the third toe
- **Toe spread (TS):** Distance from the first to the fourth toe
- **DOF:** Distance to opposite foot.
- **Gait difference:** Distance from left or right front footprint /hind footprint overlap was used to ensure the uniformity of step alternation. When the center of the hind footprint fell on top of the center of the preceding front footprint, a value of zero was recorded. When the footprints did not overlap, the distance between the center of the footprints were recorded.

**Scores:**

- **short print length:** Poor balance and gait coordination
- **low toe spread:** Good balance and gait coordination
- **Less distance to opposite foot:** Poor gait coordination
- **Gait difference:** Poor balance and gait coordination

**Hippocampal lesion surgery:** The animals were maintained in empty stomach 10h before the procedure and were anaesthetized using pentathol sodium. The hair in the head region was shaved using diluted savlon as sanitizer. The animals were fixed in the stereotaxic frame with the help of the tooth and nose bar.

A 2cm long incision was made along the scalp of the rats and the fascia was cleaned to point out the bregma [10]. Necessary steps were taken to avoid infections at all levels. The striata were marked in the scalp region by moving the manipulator from the bregma 2.2mm anteriorly, 3mm bilaterally and a small hole with 1mm diameter was made in the marked region.

0.5µl of kainic acid [11] was taken in a Hamilton syringe and was fixed in the frame. With the help of the manipulator the syringe was moved 5mm inferiorly from the dura to the striatum. The same chemical was injected in the rate of 1 µl per 1 minute following Mcginty et al., 1983 [12]. The syringe was withdrawn and the scalp was sutured with proper care.

**Post operative care:** Proper antibiotic care was given post lesion with 2mg/kg/day gentamysin for 3 days

**RESULTS AND DISCUSSION**

**Foot Print Analysis (figure-1, figure-2):** This study was performed to analyse the gait coordination and balance of the animals. For this study the fore limb of the animals were painted with fluorescent green colour and the hind limbs were painted with fluorescent pink and made to run in the apparatus. Three parameters were studied to analyse the animal’s behaviour.

**Toe Spread (figure-3, chart-1):** In this parameter the distance from the first to the fourth toe was measured in cm’s and analysed. The animals with low toe spread were considered as having good balance and gait coordination according to literatures.

The toe spread of the LC animals were significantly high when compared with the CO(figure-2) animals. The SC group of animals were not showing any significance in their toe spread and was equal with the CO animals. The animal group WS125 shown a slight increase in the toe spread in comparison with the CO group and significantly low toe spread in comparison with the LC group of animals. The WD100 group also was not showing significance in the toe spread and can be comparable with the CO group. This is because this dosage of drug was highly effective in the striatum in protection and so the animals were retaining their gait and co-ordination as like the normal animals.

**Distance to Opposite Foot (DOF) (figure-4, chart-2):** The distance between the centre of the one forelimb and the centre of the next forelimb print was measured to analyze the gait
co-ordination of the animals. Less distance to opposite foot was considered as poor gait coordination as per literature. The LC animals had significantly low DOF when compared with the CO group of animals. The SC group of animals had an equal DOF with the CO animals. The drug group animals WS125 and WD100 were showing slight high DOF. That shows the gait co-ordination was perfect with the WS125 and WD100 animals as their striatum was protected by the drug given.

Table 1: Showing the animal groups used for footprint test.

<table>
<thead>
<tr>
<th>S.No</th>
<th>Animal groups</th>
<th>Lesion surgery</th>
<th>Withania somnifera ethanolic extract treatment before and after surgery</th>
<th>Withanolide A treatment before and after surgery</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Control (CO)</td>
<td>NO</td>
<td>NO</td>
<td>NO</td>
</tr>
<tr>
<td>2</td>
<td>Sham control (SC)</td>
<td>YES (DUMMY)</td>
<td>NO</td>
<td>NO</td>
</tr>
<tr>
<td>3</td>
<td>Lesion control (LC)</td>
<td>YES</td>
<td>NO</td>
<td>NO</td>
</tr>
<tr>
<td>4</td>
<td>WS 125mg (WS 125)</td>
<td>YES</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>5</td>
<td>WA100µg (WD100)</td>
<td>YES</td>
<td>NO</td>
<td>YES</td>
</tr>
</tbody>
</table>

Chart 1: Bar diagram showing the toe spread of the animals in footprint test.

Chart 2: Bar diagram showing the distance to the opposite foot in footprint test.

Chart 3: Bar diagram showing the gait difference of the animals in footprint test.

* Indicates significance with CO group  
# Indicates significance with LC group
The gait difference was recorded as gait difference. High gait difference explains poor balance and gait incoordination.

The LC animals presented maximum gait difference by showing significant increase in the gait difference in comparison with the CO group of animals. The other animals including the SC and WS125 were not showing any significant gait difference with the CO animals. The WD100 animals shown a little gait difference which was not so significant with the CO group. These parameters proved the drugs were acting on the striatum and protecting it from free radicals produced by the lesion.

CONCLUSION

Vandeputte et al., 2010 [13], investigated the gait parameters of rat models for Parkinson’s disease (PD), Huntington’s disease (HD) and stroke using the Catwalk method by a novel automated gait analysis test and confirmed that Catwalk analysis proved more sensitive than rotarod test. In this present study the animals belonging to WS125 and WD100 were analyzed for three parameters. Toe spread, distance to the opposite foot and gait difference. The toe spread of the drug group animals were minimum as like the CO animals in that, WD100 showed minimal toe spread. The gait of the animals belongs to WS125 were normal as like the CO group of animals and the WD100 animals were showing a slight difference in their gait but was negligible. In WS125 group the distance to the opposite foot was more than the LC animals and was more equal to the CO animals but for in WD100 group the foot distance were more than the WS125 group animals. This clearly states both the drugs were effective in neuroprotection and the drug WD100 was bit more effective than the other drug.

As the herbal drugs are free of side effects we can learn the habit of taking these herbal products as food supplements in a very low dose along with our daily drinks or food. This will regularly check our nervous system protect it and will keep it active for a few more years.

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Conflicts of Interests: None

REFERENCES


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