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Adaptogenic Activity of Morus Alba Extracts in Wistar Rats

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5 Abstract

⁶ Morus alba was broadly used with a long history of traditional medicinal remedy for the

7 regimen of various illnesses. The current investigation was designed to evaluate anti-stress

⁸ activity of aqueous and ethanol extracts of Morus alba. Anti-stress of aqueous and ethanol

 $_{9}\;$ extracts of Morus alba was estimated by inducing stress in rats through the forceful

¹⁰ swimming. Homovanillic acid (HVA), urinary vanillymandelic acid (VMA), 6-?-OH- cortisol,

¹¹ 5-hydroxyindoleacetic acid (5HIAA) and ascorbic acid were reckoned as non- invasive

¹² biomarkers to assess the adaptogenic activity. Daily oral administration of aqueous and

 $_{13}$ $\,$ ethanol extracts of Morus alba at dose of 200 and 400 mg/kg body weight one hour before the

¹⁴ induction of stress to retard stress-induced urinary biochemical changes in a dose dependent

¹⁵ manner. However, non-significant changes in the urinary excretion of Homovanillic acid

¹⁶ (HVA), urinary vanillymandelic acid (VMA), 6-?-OH-cortisol, 5- hydroxyindoleacetic acid

17 (5HIAA) and ascorbic acid was perceived when compared to basal levels in normal animals.

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19 Index terms— morus alba, stress, homovanillic acid, urinary vanillymandelic acid, 6-?-OH-cortisol, 5-20 hydroxyindo- leacetic acid, ascorbic acids.

²¹ **I.** Introducton

tress can be defined as the sum total of all the reaction of the body, which disorganise the normal physiological 22 condition and result in a state of threatened homeostasis. Stress is an internationally conceded phenomenon 23 fortified by advancement of industrialization in a demanding civilization. Thus every individual is likely to face 24 stressful situation in day to day life (Selye, 1998). Stress is a stimulus that activates the hypothalamic pituitary 25 adrenal (HPA) axis and Sympathetic Nervous System (SNS) and begets a physiological change. Physiological 26 responses to stressful stimuli, including the increases in blood pressure, heart rate, body temperature and plasma 27 concentration of adreno-corticotrophic hormone (ACTH), can be related to the stress induced activation of the 28 SNS. Stress prompts synthesis and release glucocorticoids (corticosterone and cortisol) and monoamines such 29 as epinephrine, dopamine, norepinephrine and serotonin which are characteristic stress hormones (Carrasco et 30 al., 2003). Adaptogens are the substances that help organisms to adapt to unfavourable stressful conditions, 31 which could be physical, chemical, biological or mental conditions (Rege et al., 1999). The prevalent objective 32 of adaptogenic therapy is due to diminish stress reactions during the alarm phase of the stress response, inhibit 33 or retard the state of exhaustion and consequently issue a certain level of protection against long-term stress 34 (Wagner et al., 1994). Morus alba belongs to family Moraceae commonly called as white mulberry. This plant 35 has been used traditionally as anti-asthma, antidiabetic (Singab et al., 2005), hypotensive (Fukai et al., 1985) and 36 neuroprotective (Kang et al., 2006). The current investigation was carried out to assess the antistress activity of 37 aqueous and ethanol extracts of Morus alba. 38

³⁹ 2 II. Materials and Methods

⁴⁰ **3** a) Plant material and Preparation of extracts

41 The fruits of Morus alba were collected from Chennai, Tamil Nadu, India and authenticated by Green Chem, 42 Bangalore, Karnataka, India, a voucher specimen (MAT-SIP-501) were preserved for future references. The fruits

- materials (1kg) was dried, powdered and extracted with water and ethanol (60-80 o C) using soxhlet methods. 43
- The filtrate was evaporated at 70 o C in a vaccum dryer to give final yield 40.5g. 44

b) Chemicals 4 45

Homovanillic acid (CAS 306-08-1), urinary vanillymandelic acid (VMA), 6-?-OH-cortisol, 5hydroxyindoleacetic 46

acid (5HIAA) and ascorbic acid was purchased from Sigma, ST Louis, MO, USA. Acetonitrile and methanol 47 HPLC grade were supplied from Qualigens, Fischer Scientific, Mumbai. All other chemicals were analytical 48 grade and obtained from local store of Visveswarapura Institute of Pharmaceutical Sciences. 49

5 c) Animals 50

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Albino Wistar rats (150-200gm) of either sex obtained from the NIMHANS animal house, Bengaluru and were housed at room temperature in a wellventilated animal house under 12 hrs light / dark cycle in polypropylene 52 cages (29"x22"x14") with stainless steel grill top, bedded with paddy husk. The animals were maintained under 53 standard conditions in an animal house as per the guidelines of "Committee for the Purpose of Control and Supervision on Experiments on Animals" (CPCSEA) for at least one week prior to use. The rats had free access 55 to standard rat chow and water ad libitum. Simultaneous HPLC determination of HVA, VMA, 6-?-OH cortisol 56 and 5-HIAA in urine were determined to assess their standard value (Sreemantula et al., 2004). Different dilutions 25 ng/ml, 50 ng/ml, 100 ng/ml, 250 ng/ml, 500 ng/ml, 1000 ng/ml, 2500 ng/ml, 5000 ng/ml of the HVA, VMA, 58 6-?-OH cortisol and 5-HIAA were made with mobile phase from the working standards (1mg/ml) which consist 59 of 250 ng/ml internal standard (Ethyl-3-hydroxy-4-methoxy-mandelate). These dilutions were spiked into the 60 HPLC and calibration curves were plotted as peak area ratio vs. concentration. The peak area ratios of HVA, VMA, 6-?-OH cortisol and 5-HIAA to that of internal standard were calculated and substituted in the respective 62 regression equations to estimate the amount of the metabolite present in the sample. 63

e) Calibration curve for ascorbic acid 6 64

Calibration curve for ascorbic acid was done according method of (Roe and Kuether, 1943). Standard solution 65 of ascorbic acid (1mg/ml) was prepared by dissolving 10 mg of ascorbic acid in 10 ml of distilled water. By using 66 this different concentration of ascorbic acid 5, 10, 20, 30, 40 µg/ml were prepared with 4% Trichloro Acetic acid 67 (TCA). 10 ml of each of these were taken and mixed well with 0.375 gm of activated charcoal and filtered. 4% 68 TCA was used for the blank. From the filtrate 1 ml was taken in to the test tube and a drop of thiourea was 69 added to that. Then to those test tubes 1 ml of 2,4-Dinitro Phenyl Hydrazine (2,4-DNPH) was added and kept 70 in incubator for about 3 hrs, maintained at 37 0 C. The tubes were then placed in the beaker containing ice and 71 5 ml of 95% sulphuric acid was added drop by drop within 1 min interval with intermittent mixing. Finally they 72 were shaken and kept aside for 30 min. Then optical density was measured at 550 nm using spectrophotometer 73 74 [Shimadzu (UV-1601)]. Standard curve was plotted by taking concentration of ascorbic acid on X-axis and optical 75 density on Y-axis.

f) Evaluation of anti-stress activity 7 76

Rats of either sex weighing between 180-220 gm were divided into five groups (I, II, III, IV & V) each containing 77 78 six animals. The 24 hr urine sample from each group were collected into two different beakers, one containing 79 5 ml of 10% oxalic acid for the spectrophotometric determination of ascorbic acid at 550 nm and the other containing 0.5 ml of 6N hydrochloric acid for the determination of stress metabolites. The experimental protocol 80 was divided into four phases: In the first phase of the experiment, 24 hr urine samples were collected in all the 81 groups and subjected to analysis for HVA, VMA, 6-?-OH cortisol, 5-HIAA and ascorbic acid and the normal 82 values were recorded for four consecutive days. In the second phase, after a recovery period of one week, the 83 animals in each group were subjected to fresh water swimming stress individually. In this method, rats are forced 84 to swim until exhausted in a cylindrical vessel of 60 cm height and 45 cm diameter containing water at room 85 temperature (28°C). Water depth was always maintained at 40 cm. The 24 hr urine samples were collected in all 86 the groups and subjected to analysis for HVA, VMA, 6-?-OH cortisol, 5-HIAA and ascorbic acid and the values 87 were recorded for four consecutive days. In the third phase of the experiment, after a recovery period of one week, 88 89 the experimental animals were administered as follows for four consecutive days. Group 1 rats served as normal 90 control and received 2ml/kg distilled water, group 2,3 rats were administrated orally with aqueous extracts of 91 Morus alba at dose of 200 mg/kg and 400 mg/kg respectively. Group 4,5 rats received ethanol extracts Morus 92 alba orally at dose of 200 mg/kg and 400 mg/kg respectively. In the final phase of the experiment, after a recovery period of one week, the administration of Morus alba extract were done as mentioned in the third phase, one 93 hour prior to the daily induction of stress for four consecutive days while group I serving as control. The 24 hr 94 urine samples were collected in all the groups and subjected to analysis for HVA, VMA, 6-?-OH cortisol, 5-HIAA 95 and ascorbic acid and the values were recorded for four consecutive days to study the influence of the aqueous 96

and ethanol extracts of Morus alba on the stress induced biochemical changes (Sreemantula et al., 2004). 97

g) Statistical analysis 8 98

The data were expressed as mean \pm S.E.M. Statistical analysis was performed by using student's paired t-test, 99 where the difference was considered significant if p < 0.05. 100

9 **III.** Results 101

a) Calibration curves of biomarkers for determination of 10 102 standard value 103

A typical chromatogram was manifested in Figure ??. The retention times of VMA, IS (Ethyl-3hydroxy-4-104 methoxy-mandelate), 5-HIAA, 6-?-OH cortisol and HVA were found to be 4.46, 5.210, 9.31, 10. There was 105 variation in each biomarkers from group to group in normal state. The amount of VMA (0.343 $?g \pm 0.019$), 106 5-HIAA (0.348 $g \pm 0.013$) and HVA (0.113 $g \pm 0.009$) were low in group I in normal state and high amount 107 of VMA (0.431? g \pm 0.017), 5-HIAA (0.463 ? g \pm 0.022) and HVA (0.176 ? g \pm 0.006) were found in group IV in 108 normal state. The level of 6-?-OH cortisol was low in group V (0.424 ?g \pm 0.032) and high in group I (0.557 ?g 109 \pm 0.010), also the amount of ascorbic acid was low in group IV (43.92 ?g \pm 2.33) and high in group I (52.64 ?g \pm 110 5.64) in normal condition. Significant increase (P<0.05) in urinary levels of VMA, 5-HIAA, 6-?-OH cortisol and 111 112 HVA was noted in group I to V. Significant decrease (P<0.05) in urinary levels of ascorbic acid was observed in 113 group I to V in stress condition. There were slight changes in VMA, 5-HIAA, 6?-OH cortisol, HVA and ascorbic acid levels in urine of animals treated with aqueous and ethanol extracts of Morus alba in the normal state. There 114 was variation from day to day and the variation is different from group to group. However observed the changes 115 in the levels of the urinary metabolites when compared to normal basal levels were found to be non-significant. 116 Aqueous and ethanol extracts of Morus alba significantly (P<0.05) diminished urinary levels of VMA, 5-HIAA, 117 6-?-OH cortisol and HVA in group II, III, IV and V, also significant increased (P<0.05) in urinary ascorbic acid 118 levels was perceived in group II, III, IV and V compared to their respective stress condition. 119

GROUP-I GROUP-II GROUP-III GROUP-IV 11 120

IV. Discussion 12121

Stress represents reactions of the body to a stimulus that tends to modify homestasis (Selye, 1998). Stress hor-122 123 mones are synthesised during stress condition for example the catecholamines (epinephrine and norepinephrine) produced by the SNS, and corticosteroids, produced by the ACTH stimulated adrenal cortex and glucocorticoid 124 stimulated increase in serotonin are the major stress hormones (Uresin et al., 2004). In the current investigation, 125 VMA as the peripheral metabolite of NA, 5-HIAA as the main metabolite of serotonin, 6-?-OH cortisol as 126 metabolite of cortisol, HVA as the predominant metabolite of dopamine and ascorbic acid as a metabolite of 127 glucose (in rats) were taken as an non-invasive biomarkers to display the increase in peripheral sympathetic 128 activity during stress to assess the anti-stress activity of aqueous and ethanol extracts of Morus alba. The 129 130 data indicated that VMA, 5-HIAA, 6-?-OH cortisol, HVA and ascorbic acid were excreted in urine daily at certain levels (basal values) as metabolites of NA, 5-HT, cortisol, DA and glucose respectively. The stress 131 affected on the neurotransmitter levels and increased VMA, 5-HIAA, 6-?-OH cortisol, HVA and diminished 132 ascorbic acid excretion. When aqueous and ethanol extracts of Morus alba administered to normal animals did 133 not change VMA, 5-HIAA, 6-?-OH cortisol, HVA and ascorbic acid in comparison with basal values but prior 134 administration of aqueous and ethanol extracts of Morus alba to stress induced rats exhibited the reduction in 135 urinary VMA, 5-HIAA, 6-?-OH cortisol, HVA and increased the ascorbic acid levels in dose dependent manner. 136 The previous phytochemical evaluation of Morus alba divulged the presence of phenolic compounds such as 137 flavonoids (Quercetin, rutin), tannin which could be expected to be responsible for anti-stress activity (Ayoola et 138 al., 2011). As VMA is a metabolite of norepinephrine (NE) and NE is synthesized by dopamine. Previous reports 139 exhibited that these phytochemicals can bind to the GABA A -BZDS complex, consequently, enhance GABA 140 level and decline dopamine and decrease plasma corticosterone level that lead to reduce level of VMA and 6-?-OH 141 142 cortisol repectively (Patil et al., 2006). Phenolic compounds such as flavonoids showed the affinity towards D 2 143 receptor; hence they can block the dopamine receptor and decrease the serotonin which causes to decrease level of HVA (Samson et al., 2006). These active compounds can prevent activity of tryptophan hydroxylase enzyme 144 which is involved in the biosynthesis of 5-HT, thus, they can reduce level of 5HIAA ultimately (Singh et al., 145 1990). Ascorbic acid is also utilized as a co-factor in the biosynthesis of NE from DA may also attribute to low 146 concentration of ascorbic acid in urine. Hence, effect of these bioactive compounds on reduction of dopamine 147 synthesis that can influence on the increase of urinary level of ascorbic acid. 148

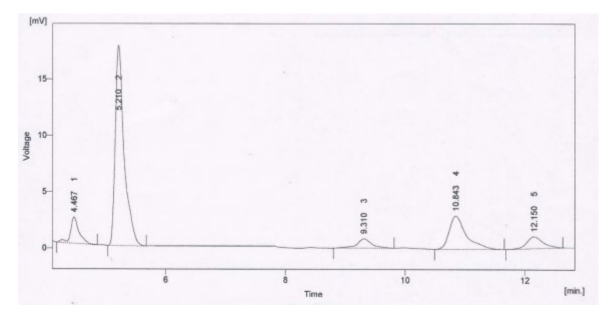


Figure 1:

¹⁴⁹ 13 V. Conclusion

The present investigation manifest that aqueous and ethanol extract of Morus alba have anti-stress activity due to normalization of monoamines and glucocorticoids homeostasis by acting on HPA axis and SNS.^{1 2 3 4}

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