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**ORIGINAL ARTICLE** 

# 2 Aqueous Extract of Nigella sativa Seeds Suppresses <sup>3</sup> Testicular Steroidogenesis in Mice Leydig Cells in <sub>4</sub>vitro

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### 10 ABSTRACT

11 Nigella sativa (black seed) is an important medicinal herb with folkloric use in wide range of diseases. It is 12 well studied for its biological activities. However, there is limited information regarding its effect on the 13 male reproductive system. This study describes the effect of the aqueous extract of N. sativa (NSE) on 14 testicular steroidogenesis from mice Leydig cells in vitro. Mice testicular cells were incubated in a media 15 containing either no treatment or NSE or LH alone or combination of LH and NSE. Incubations were 16 carried out for three hours in a shaking water bath at 34°C. Testosterone was measured by 17 radioimmunoassay. At all doses, NSE significantly (p < 0.05) inhibited both basal and LH-stimulated in 18 vitro testosterone secretion. At a dose of 1000 µg, NSE inhibited 52% of basal testosterone and 97% of 19 LH-stimulated testosterone, compared to control (0.32 ± 0.008 ng/ml) and LH alone (0.33 ± 0.01 ng/ml) 20 respectively. Thus, it is concluded that that both the basal and the LH-stimulated secretion of testosterone 21 from Leydig cells are suppressed significantly in the presence of different doses of NSE in vitro. However, 22 further studies are needed to explore the effect of chronic treatment with NSE in male and its potential to 23 be used as a contraceptive in male.

24 Keywords: Nigella sativa, Black seed, Male reproductive system, aqueous extract, Leydig cell, 25 testosterone

27 commonly, known as black seed or black cumin and 42 antioxidant, hepatotprotective [3], nephroprotective, 28 locally as Kalonji have been used in folk (herbal) 43 antihypertensive [4], muscle relaxant, bronchodilator 2º medicine for centuries for treatment of many acute as 44 [5,6], CNS depressant effects [7], antlipidemic [8], <sup>(30</sup> well as chronic conditions worldwide [1,2]. It has been 45 antidiabetic [9,10], anticancer [11,12], analgesic 3 used in the treatment of asthma, diarrhea, indigestion, 46[13,14], anti-inflammatory [13,15], antiulcer [16] and 32 dizziness, influenza, dyslipidemia, many dermatological 47 neuroprotective effects [17,18]. 33 conditions and as a diuretic and immune modulator. The 48 34 seeds contain 36%-38% fixed oils, proteins, alkaloids, 49 have been shown to be due to the presence of 35 saponins, 0.4%-2.5% essential oil, crude fiber, minerals, 50 thymoquinone, which is the major component of the 36 vitamins, aliphatic alcohols and ketones [1].

38 pharmacological action(s) of Nigella sativa seed extract 53 be very effective in inhibiting histamine release induced 39 or its active compound(s) on various body systems in 54 by the secretagogues: antigen in sensitized cells [19]. 40 vivo or in vitro. The herb has been extensively studied 55 However, the herb is not well studied for its effect on

The seeds of Nigella sativa Lin. (Ranunculaceace), 41 for its different biological activities which includes

Much of the biological activities of the black seeds 51 essential oil and fixed oil. Nigellone, is another Many studies have been conducted on the 52 compound of Nigella sativa, which has been shown to

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56 reproductive system. Moreover, the existing information 57 in this regard is quite scanty and rather contradictory. 58 Significant abortificient activity of N. sativa seed 59 powder, ethanolic and hexane extracts was 60 demonstrated in rats [20]. However, Prakash et al [21] 61 did not find any anti-fertility activity in aqueous, 62 ethanolic and petroleum ether extracts of the seeds of N. 63 sativa when tested at a dose of 150-200 mg/kg daily in 64 rats on the days 1-7 post-coitum schedule.

There is a growing demand for men to share the 66 burden of responsibility and risks of contraception 67 because of growing population pressures and the 68 increasing dissatisfaction of women in assuming almost 69 all the risks of adequate contraception. A major 70 challenge in this field is that the most of the male 71 contraceptive agents currently in use offer little promise 72 and about 15% of the 200 most commonly prescribed 73 drugs can have adverse effects on male reproduction<sub>110</sub> dissected out immediately and de-capsulated. Leydig 74 either by influencing its hormonal profile or impairing 111 cells were isolated as described earlier [22]. Leydig 75 their sexual performance. The discovery of key<sub>112</sub> cells (80,000/tube) were pre-incubated for 1 h to remove 76 regulators of gonadal hormones and gametogenesis<sub>113</sub> the endogenous testosterone, the media were replaced 77 from black seed may provide opportunities to alter our114 with either fresh medium or medium containing graded 78 approaches towards management of contraception.

Since, no data on the effect of *N* sativa on testicular<sub>116</sub>  $\mu$ g/tube).  $_{\rm 80}$  steroidogenesis is available, we designed this in  $vitro_{\rm 117}$ 81 study to investigate the direct effect of crude aqueous 118 modulate stimulated testosterone secretion, samples  $^{82}$  extract on basal and LH-stimulated testicular<sub>119</sub> were challenged with LH (500  $\mu$ IU/tube) alone or with 83 steroidogenesis by mice Leydig cells.

### MATERIALS AND METHODS

### 85 Preparation of the crude extract

Dried black seeds of *Nigella sativa* were purchased 87 from the local market in Karachi. The plant seeds were 88 cleaned of any adulterant materials. NS seeds were 126 89 ground with an electric grinder into a coarse powder. A127 medium directly by a highly-sensitive RIA according to 90 measured quantity was soaked in 70% aqueous-128 WHO protocol, using <sup>3</sup>H-labeled testosterone, as tracer. 91 methanol (30:70) at room temperature by cold129 Highly specific antiserum for testosterone was acquired 92 maceration for a total of 3 days. Thereafter, the filtrate130 from Guildhay UK. RIA reagents were directly added to 93 was collected through Whatman's qualitative grade 1131 tubes containing incubation medium. After addition of 94 filter papers and the plant material was again subjected 132 all the reagents, tubes were incubated for 30 min. at 4 C. 95 to the same treatment as the first macerate. The 133 The bound and unbound fractions were separated by the 96 combined filtrate was concentrated using a rotary134 addition of 0.1% activated charcoal. Radioactivity was 97 evaporator at 40°C under reduced pressure. Extract was 135 measured in a scintillation counter. Testosterone 98 stored at -4°C until used for biological activity.

### 99 Leydig cells preparation

Three bulbce male mice (weight  $36 \pm 2$ ) were used 101 for each experiment. Animals were obtained from the 102 AKU animal facility, where they were maintained under 103 standard conditions of 14-hour light and 10-hour dark 104 cycle.

Direct effect of aqueous extract of Nigella sativa<sup>142</sup> Statistical analysis 106 seeds (NSE) on testosterone secretion was studied by143 107 the incubation of Leydig cells as described by Van-144 analyzed for statistical significance using an 108 Damme et al, 1974 [22], with minor modifications. Mice145 independent t test on SPSS. A p value < 0.05 was 109 were killed by cervical dislocation. Testes were 146 considered significant.

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Fig 1. Effect of aqueous extract of Nigeel sativa seeds (NSE) on basal testosterone by mice Leydig cells in vitro \*Significant difference between control and treated groups (p < 0.05)

115 doses of crude aqueous extract of NS seeds (1.0-1000

Moreover, to test the ability of the extract to 120 different doses of NSE (1.0-1000µg). After 3 h, the 21 incubation reaction was stopped by dipping the tubes in 122 water bath at 60 C for 10 min. Samples were kept frozen 123 until testosterone was measured by highly-specific 124 radioimmunoassay.

### 5 Radioimmunoassav

Testosterone was measured in the incubation 136 concentration calculated was by logit-log 137 transformation [23].

138 The sensitivity of T assay was 0.0125 ng and the 139 intra-assay coefficient of variation was less than 10%. 40 The levels of testosterone in the media are expressed as 41 ng/ml.

Data are expressed as mean  $\pm$  S.E.M. Results were

### Nigella sativa and testicular steroidogenesis



stimulated testosterone production by mice Leydig cells in vitro \*Significant difference between control and treated groups (p<0.05)

### RESULTS

149 significantly (p < 0.05) both basal and LH-stimulated<sup>206</sup> serves two crucial functions: synthesis and secretion of 150 testicular testosterone secretion in vitro. Moreover, the 207 testosterone by Leydig cells and production of a 151 inhibitory effect of NS seed extract was more 200 sufficient number of competent spermatozoa supported 152 pronounced at the higher doses.

### 153 Effect on basal testicular steroidogenesis

155 in the cells treated with NS seed extract (1.0-1000213 Follicle stimulating hormone (FSH), and testosterone  $156 \,\mu\text{g/tube}$ ) was significantly (p < 0.05) reduced compared 214 during fetal and postnatal life [29]. A deficiency of 157 with the control in a dose-dependent manner. The215 these hormones leads to hypogonadism and sterility, a 158 inhibition was more pronounced at the higher doses.216 condition that can be treated with specific replacement 159 Inhibitory effect of NSE was able to inhibit 52% of the217 therapies [30]. Testosterone biosynthesis in the Leydig 160 basal testosterone production and this inhibition was<sup>218</sup> cells is primarily regulated by LH [31]. Deficiency of 161 still present at the lowest NS dose of  $1.0 \,\mu g$ .

164 of NS seed extract (1.0-1000 µg) caused a significant<sup>223</sup> used as a contraceptive. Oral administration of crude 165 (p < 0.05) and a dose-dependent inhibition of LH-224 ethanol extracts showed significant contraceptive effect 166 stimulated (500 μIU) testosterone production. The<sup>225</sup> in male rats [20]. Since, no data about the effect of NS 167 inhibition was more pronounced at higher doses of NSE226 seed extract on testicular steroidogenesis have yet been 168 with maximum effect (97% inhibition) obtained at 1000227 available; these results open new fronts in the 169 µg dose of NSE. LH (500 µIU) was used for maximal228 exploration of possible effects of Nigella sativa on the dose 170 stimulation. This was selected 171 LH/testosterone dose-response curve to variable doses230 potential contraceptive effects of Nigella sativa on the 172 of LH (16-500µIU) (data not shown). Treatment with231 hormonal regulation of male reproductive axis. This 173 NSE caused dose-dependent inhibition of the LH-232 study has provided us with important insight towards 174 stimulated testosterone production when compared to233 formulation of a new contraceptive pill that would 175 LH 500 μI response (Fig 2, with maximum effect (97%234 temporarily stop spermatogenesis, thus producing 176 inhibition) obtained at 1000 µg dose).

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### DISCUSSION

This study provides the first evidence for a strong 78 79 effect of N. sativa seed extract on testicular 80 steroidogenesis indicating a potential contraceptive role. 81 Our data suggest that N sativa extract inhibits both basal 82 and LH-stimulated testosterone biosynthesis signaling 83 pathways. The mechanism behind its effect is not clear 84 and further studies are needed to elucidate its further 85 role and mechanism of action. The crude extract of N. 86 sativa seeds has been reported to possess calcium 87 channel blocking activity [5] and there is evidence that 88 calcium may be involved in the signaling mechanism 89 [24]. Significant abortificient activity of N. sativa seed 90 powder, ethanolic and hexane extract, is demonstrated 91 in women [20,25] and rats [21]. However, Prakash et al. 92 [26] did not find any anti-fertility activity in aqueous, 93 ethanolic and petroleum ether extracts of the seeds of 94 Nigella sativa when tested at a dose of 150-200 mg/kg 95 daily in rats on the days 1-7 post-coitum schedule. The Fig 2. Effect of aqueous extract of Nigeel sativa seeds (NSE) on LH196 volatile oil of Nigella seeds inhibits the spontaneous 197 movements of rat and guinea pig uterine smooth muscle +Significant difference between LH alone and treated groups (p<0.05)<sup>98</sup> and also the oxytocin-induced contractions [27]. A 99 single report in male rats has suggested that seed extract 200 treatment not only causes a general reduction in the size 201 of reproductive organs but also suppresses 202 spermatogenesis at the spermatocyte stage. However, 203 similar changes in the reproductive hormones of the 204 treated animals was not observed [28].

*Nigella sativa* seeds extract was able to inhibit<sup>205</sup>. The testis is a complex male reproductive organ that 209 by Sertoli cells, to attain fertility. It is well known that 210 the essential prerequisite for normal testicular 211 development and maintenance of spermatogenesis is the As shown in the Fig 1, basal testosterone production<sup>212</sup> controlled secretion of Luteinizing hormone (LH), 219 these hormones leads to hypogonadism and sterility, a 220 condition that can be treated with specific replacement 162 Effect on LH-stimulated testicular steroidogenesis 221 therapies [30]. Reversible inhibition of these hormones As shown in Fig 2, administration of different doses<sup>222</sup> by any external measure may be beneficial as it can be from<sup>229</sup> reproductive functions. These data offer insights into 235 reversible infertility.

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### 242 REFERENCES

- 2431. Ali BH, Blunden G. Pharmacological and toxicological307
- 244 properties of Nigella sativa. Phytother Res 2003; 17:299-305. 308 20.
- 245 **2**. Farah IO, Begum RA. Effect of Nigella sativa (N. sativa L.) and 309
- oxidative stress on the survival pattern of MCF-7 breast cancer310 cells. Biomed Sci Instrum 2003: 39:359-64.

311 **21**. 2483.

- Iddamaldeniya SS, Thabrew MI, Wickramasinghe SM,312 Thammitiyagodage MGA. long-term<sub>31322</sub>. Ratnatunge N, investigation of the anti-hepatocarcinogenic potential of an314 indigenous medicine comprised of Nigella sativa, Hemidesmus315 indicus and Smilax glabra. J Carcinog 2006; 5:11.
- Zaoui A, Cherrah Y, Lacaille-Dubois MA, Settaf A, Amarouch<sub>317 23</sub>. 2534.
- H, Hassar M. Diuretic and hypotensive effects of Nigella sativa318 254 in the spontaneously hypertensive rat. Therapie 2000; 55:379-319 82
- Gilani AH, Aziz N, Khurram IM, Chaudhary KS, Iqbal A.321 24. 257 5. Bronchodilator, spasmolytic and calcium antagonist activities of 322 Nigella sativa seeds (Kalonji): a traditional herbal product with 323 multiple medicinal uses. J Pak Med Assoc 2001 51:115-20. 324 25.
- 261 6. Boskabady MH, Javan H, Sajady M, Rakhshandeh H. The<sub>325</sub> possible prophylactic effect of Nigella sativa seed extract in<sub>326</sub> asthmatic patients. Fundam Clin Pharmacol 2007; 21:559-66.
- 2647 Abdel-Fattah AM, Matsumoto K, and Watanabe H. Antinociceptive effects of Nigella sativa oil and its major component, thymoquinone, in mice. Eur J Pharmacol 2000; 400:89-97.
- 268 8. Dahri AH, Chandiol AM, Rahoo AA, Memon RA. Effect of 32.28 Nigella sativa (kalonji) on serum cholesterol of albino rats. J Ayub Med Coll Abbottabad 2005; 17:72-4.
- 2719. Meral I., Yener Z, Kahraman T, Mert N. Effect of Nigella sativa 5 29 on glucose concentration, lipid peroxidation, anti-oxidant defence system and liver damage in experimentally-induced 274 diabetic rabbits. J Vet Med A Physiol Pathol Clin Med 2001; 48:593-9.
- 39 30 El-Dakhakhny M, Mady N, Lembert N, Ammon HP. The 276 10. hypoglycemic effect of Nigella sativa oil is mediated by  $\frac{340}{341}$ extrapancreatic actions. Planta Med 2002; 68:465-6.
- Kaseb AO, Chinnakannu K, Chen D, Sivanandam A, Tejwani S, 342 31. 279 11. Menon M, Dou QP, Reddy GP. Androgen receptor and E2F-1 targeted thymoquinone therapy for hormone-refractory prostate cancer. Cancer Res 2007; 67:7782-8.
- 283 12. Randhawa MA, Alghamdi MS. Anticancer activity of Nigella sativa (black seed) - a review. Am J Chin Med 2011; 39:1075-345 Sheikh A. Saeed, Department of Basic Medical Sciences, College of 91
- 286 13. Al-Ghamdi MS. The anti-inflammatory, analgesic and antipyretic activity of Nigella sativa. *J Ethnopharmacol* 2001;<sup>348</sup><sub>349</sub> 76:45-8.
- 289 14. extract on experimentally induced pain in albino mice. J Coll<sup>351</sup> Physicians Surg Pak 2010; 20:464-7.
- Tekeoglu I, Dogan A, Demiralp L. Effects of thymoquinone<sup>353</sup> 29215
- models. Phytother Res 2006; 20:869-71.

- 295 16. Al-Mofleh IA, Alhaider AA, Mossa JS, Al-Sohaibani MO, Al-Yahya MA, Rafatullah S, Shaik SA. Gastroprotective effect of an aqueous suspension of black cumin Nigella sativa on necrotizing agents-induced gastric injury in experimental animals. Saudi J Gastroenterol 2008; 14:128-34.
  - Kanter M. Coskun O, Kalayci M, Buyukbas S, Cagavi F. Neuroprotective effects of Nigella sativa on experimental spinal cord injury in rats. Hum Exp Toxicol 2006; 25:127-33.
- 303 18. Al-Naggar TB, Gómez-Serranillos MP, Carretero ME, Villar AM. Neuropharmacological activity of Nigella sativa L. extracts. J Ethnopharmacol 2003; 88:63-8.
  - Chakravarty N. Inhibition of histamine release from mast cells by nigellone. Ann Allergy 1993; 70:237-42.
    - Keshri G, Singh MM, Lakshmi V, Kamboj VP. Post-coital contraceptive efficacy of the seeds of Nigella sativa in rats. Indian J Physiol Pharmacol 1995; 39:59-62.
  - Prakash AO, Mathur R. Screening of Indian plants for antifertility activity. Indian J Exp Biol 1976; 14:623-6.
  - Van Damme MP, Robertson DM, Diczfalusy E. An improved in vitro bioassay method for measuring luteinizing hormone (LH) activity using mouse Leydig cell preparations. Acta Endocrinol (Copenh) 1974; 77:655-71,
  - Midgley AR Jr, Niswender GD, Rebar RW. Principles for the assessment of the reliability of radioimmunoassay methods (precision, accuracy, sensitivity, specificity). Acta Endocrinol Suppl (Copenh) 1969; 142:163-84.
  - Janszen FH, Cooke BA, Van Driel MJ, Van Der Molen HJ. The effect of calcium ions on testosterone production in Leydig cells from rat testis. Biochem J 1976; 160:433-7.
  - Siddiqui MB, Alam MM, Husain W, Sharma GK. Ethnomedical study of plants used for terminating prenancy. Fititerapia 1988; 59:250-252.
- 27 26. Prakash AO, Mathur R. Screening of Indian plants for antifertility activity. Indian J Exp Biol 1976; 14:623-6.
  - Aqel M, Shaheen R. Effects of the volatile oil of Nigella sativa 27. seeds on the uterine smooth muscle of rat and guinea pig. J Ethnopharmacol 1996; 52:23-6.
    - Agarwal C. Effects of seeds og 'Kalaunji' (Nigella Sativa L.) on the fertility and sialic acid content ogf the reproductive organs of the male rat. Geobios 1990; 17:269-72.
    - Gnessi L, Fabbri A, Spera G. Gonadal Peptides as Mediators of Development and Functional Control of the Testis: An Integrated System with Hormones and Local Environment. Endocrine Reviews 1997; 18:541-608.
    - WC H. Hypogonadotropic hypogonadism: gonadotropin therapy. In: CW Bardin (ed) Current Therapy. Endocrinology and Metabolism, 1991. BC Decker, Philadelphia, PA: p. 267-72.
    - Dufau ML. Endocrine regulation and communicating functions of the Leydig cell. Annu Rev Physiol 1988; 50:483-508.

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  - 352 Qaiser Jabeen, Dept. of Biological & Biomedical Sciences Aga Khan University, Stadium Road, Karachi.
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