Estrous Cycle Cessation Caused via Activating the Apoptotic Pathway by NNSS2 Isolated from Naja naja Shedded Skin

Keywords: Anestrus; Folk and traditional medicine; Follicular atresia; Phase arrest

Abstract

The snake shed skin has long been used in the Chinese and Indian traditional, folk medicine for various medicinal purposes. Previously it was reported that N. naja shed skin caused temporary cessation of the estrous cycle. Based on these reviews and reports this work was undertaken to purify, characterize bioactive compound from Naja naja shed skin and to assess its bioactivity in animal model.

The N. naja shed skin extract was subjected to thin layer chromatography, HPLC for purification to obtain active factor, and it was characterized (MALDI, CD spectral analysis, 1H NMR & XRD). Purified fraction was injected subcutaneously in female Swiss albino mice. Serum progesterone, estradiol, IL-1β, TNF-α levels were assessed; histopathology of uterus and ovary was done. Uterine and ovarian apoptotic marker expressions were studied.

NNSS2 was purified having molecular weight of 882.6 Dalton. It caused cessation of estrous cycle for 5 days, decreased serum progesterone, P/E ratio, increased IL-1β, caspase 3,9, Bax, decreased BCL2, HSP70, HSP90 expressions and cleaved PARP. Histological studies of ovary showed immature follicles; and uterus displayed constricted structure. NNSS2 influenced reproductive cycle of female albino mice by altering hormones, cytokine profiles, apoptotic markers, and by involvement of intrinsic apoptotic pathway. This study provided a novel outline of the probable bioactive compounds present in the N. naja shedded skin.

Introduction

The estrous cycle of female mice involved four phases: diestrous, metestrous, proestrous and estrous, it lasted for 4-5 days. It is regulated very intricately by endocrine and cytokine circuitry. The short length and less complexity of the estrous cycle in mice made it ideal animal experimental model for study on reproductive cycle, including the effect of anti fertility agents from natural sources.

The female anti fertility drugs generally prescribed by doctors are chemically steroids, available in a combination of estrogen and progestins, also known as oral contraceptive pills. The adverse side effects of the oral contraceptive pills included cardiovascular diseases, venous risk and metabolic disorder like lipoprotein changes, altered insulin response to glucose and prolonged use of those even might lead to endometriosis [1-3]. The nonsteroidal anti fertility drugs e.g. Gonadotropin-releasing hormone analogues (goserelin, leuprolide) lead to endometriosis [1-3]. The nonsteroidal anti fertility drugs e.g. insulin response to glucose and prolonged use of those even might lead to endometriosis [1-3]. The nonsteroidal anti fertility drugs e.g. insulin response to glucose and prolonged use of those even might lead to endometriosis [1-3].

The snake shed skin has long been used in the Chinese traditional medicine for various disorders including malignant sores, such as mammary abscess and tumor, boils, carbuncles, and furuncles. The shedded skins are usually roasted and then used both internally and topically. The snake shed skin considered useful in reducing clouding of the cornea. In Indian traditional, folk and Santhal medicinal system mentioned the use of snake shed skin for reproductive disorders, and previous studies had shown that Naja naja shedded skin influenced rat estrous cycle [7]. However no reports are available regarding the presence of any bioactive compounds in the N. naja shedded skin. The present study was designed to isolate bioactive molecule from N. naja shedded skin and the probable molecular action on female estrous cycle.

Materials and Methods

Chemical and reagent

Carbinol and HPLC water were purchased from Spectrochem (India), IL-1β, TNF-α ELISA kits were purchased from R&D system (Minneapolis, USA), Hormones ( Estradiol, Progestrone) were procured from Cusabio, China. Prestained molecular weight marker (Fermentas, USA), Primary antibody ( β-actin, Bax, BCL2, Caspase 3,8,9, PARP, HSP70, and HSP90) and alkaline phosphatase conjugated secondary antibodies were purchased from Santacruz CA (USA), Ripa Lysis Buffer (Sigma Aldrich, USA). All other chemicals purchased are of AR grade or otherwise mentioned.

Animals and experimental design

Female Swiss albino mice 10-12 weeks old were purchased from approved animal breeders of Maulana Azad College, Kolkata; they were housed in polypropylene cages at temperature (26±2 °C) and light controlled environment (12 h light/dark). All the animals were fed with standard laboratory diet, had free access to food and water ad libidum. Estrous cycle was followed by carrying out vaginal smears daily. Vaginal lavage was performed using approximately 30 μl of 0.9% NaCl, dispensed in the vagina through plastic tip applicator. Cells were collected and studied under microscope to confirm each stage of estrous cycle. Animals showing two consecutive cycles were used.
for the experimental study. Regulations of the Animal Experimental Ethics Committee were followed, and were in accordance with the guidelines of the Committee for the Purpose of Control and Supervision of Experiments on Animal (CPCSEA) [25/250/2012-AWD], Government of India. Vaginal lavage was observed every 24 h for 5 consecutive days.

Collection of N. naja shedded skin

Fresh Naja naja shedded skins of both sexes were collected from North 24-Parganas of West Bengal, India as per the permission granted by the Ministry of Forests & Wild Life, Govt. of West Bengal, India [2105/WL/4R-l (PI-IX)].

Isolation, purification and characterization of active fraction

Prior to TLC, 8 g N. naja shedded skin was dissolved in 2 ml HPLC water, kept overnight at 4 °C followed by centrifugation. The supernatant was separated and TLC was done on preactivated silica gel G coated glass plates using isopropanol: 0.1 N HCl (7:3, v/v). Spots were visualised in 0.1% ninhydrin and Rf was calculated, same protocol was followed for preparative TLC. Zones were scraped and eluted in methanol followed by centrifugation. The supernatant was kept at room temperature for 18-24 h to obtain zone 2. Prior to RP-HPLC zone 2 was washed in solvents (diethyl ether, ethyl acetate, chloroform, methanol), and was further purified by RP-HPLC (Jasco V 2080 HPLC system, C18 column 5 μm, 250 mmX 4.6 mm) using methanol: water (70: 30, v/v) and visualized at 254 nm for 90 min to obtain NNSS2. M.W of NNSS2 was confirmed through MALDI-TOF (4700 Linear Spectrophotometer). Secondary structure of the peptide was confirmed by CD spectral analysis. The probable constituents and chemical nature were analysed by FTIR spectroscopy (Perkin Elmer RX-1 FTIR express version 1.03.00 spectrophotometer), H1NMR (AV 300 MHz NMR system dual probe) and x-ray diffraction (PAN analytical PW.3040/60, Netherlands).

Effect of NNSS2 on serum hormones and cytokines level

On day 5 following NNSS2 exposure serum 17-β estradiol, progesterone, IL-1β and TNF-α levels were measured by ELISA.

Histopathological studies of NNSS2 treated mice uterus and ovary

On day 5 following NNSS2 exposure uterus and ovaries of mice were isolated and fixed in 10% buffered formalin followed by dehydration, clearing and paraffin embedding. 5 μ sections were stained with haematoxylin-eosin and observed under bright field microscope.

Western immunoblot analysis

Expressions of different proteins (Bax, BCI2, HSP 70, HSP 90, PARP, and β-actin) of uterus and ovary tissue homogenate were analysed by western blot. For preparation of tissue homogenate uterus and ovaries were isolated and placed in cold PBS (pH 7.2). 10% tissue homogenate was prepared in RIPA lysis buffer in the presence of protease and phosphatase inhibitors (Roche, Cat. No. 10760388001). Western blot analysis was performed using antibodies (Santa Cruz Biotechnology, Inc., CA, USA) and detected using an enhanced chemiluminescence method (Pierce™ ECL Plus Western Blotting Substrate). The protein levels were quantified using an optical density reader (SpectraMax M2e, Molecular Devices, CA, USA) and expressed as relative protein abundance. Table 1 shows the effect of NNSS2 on serum hormones and cytokines level and histopathological studies of NNSS2 treated mice uterus and ovary.

Table 1: FTIR and H1NMR analysis of NNSS2.

<table>
<thead>
<tr>
<th>Absorption spectra of NNSS2 (FTIR)</th>
<th>Composition</th>
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<tbody>
<tr>
<td>3400.98 N-H stretching, amines</td>
<td>1611.81 Glycine (NH₂ banding) B-sheet</td>
</tr>
<tr>
<td>1564.17 Aspartate (Asp), bending Glutamine (Glu)</td>
<td>1450.34 Phenyl alanine, benzene ring like structure</td>
</tr>
<tr>
<td>1028.24 -OH group</td>
<td>818.01 Out-of-plane NH bending</td>
</tr>
<tr>
<td>1053.69 Amine</td>
<td>1213.38 C-OH group</td>
</tr>
<tr>
<td>946.64 -CH bending</td>
<td>946.64 -CH bending</td>
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Table 2: FTIR and H1NMR analysis of NNSS2.

<table>
<thead>
<tr>
<th>Random chemical shift (H1NMR) of NNSS2</th>
<th>Residue</th>
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<tbody>
<tr>
<td>δH shift of NNSS2</td>
<td>Residue</td>
</tr>
<tr>
<td>8.387</td>
<td>aldehyde</td>
</tr>
<tr>
<td>7.809</td>
<td>Aromatic or RCH=CHR</td>
</tr>
<tr>
<td>4.6-4.9</td>
<td>Aspartate/ Asparagine/ Tryptophan</td>
</tr>
<tr>
<td>4.089</td>
<td>Ester or ether or alcohols</td>
</tr>
<tr>
<td>4.066</td>
<td>Ester or ether or alcohols</td>
</tr>
<tr>
<td>3.520</td>
<td>Alcohol or Ar-CH</td>
</tr>
<tr>
<td>2.971</td>
<td>RCO=CH or CH-RNH₂</td>
</tr>
<tr>
<td>2.763</td>
<td>R₂CO=CH or CH-RNH₂</td>
</tr>
<tr>
<td>2.102</td>
<td>R₂NH or R-CO-CH</td>
</tr>
<tr>
<td>1.637</td>
<td>R-H or ROH</td>
</tr>
<tr>
<td>1.281</td>
<td>R-H or ROH</td>
</tr>
<tr>
<td>0.984</td>
<td>R-H</td>
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of protease inhibitor cocktail. After incubation period of 1 h at 4 °C, it was centrifuged at 12,000 rpm for 30 min (4 °C). The supernatant was collected and stored in -20 °C for future use. Equal concentration and volume of cellular protein samples were run in 12.5% SDS PAGE and transferred to nitrocellulose membrane in an immunoblot apparatus using transfer buffer (193 mM Glycine, 25 mM Tris and 20% methanol) and were blocked using 5% BSA in TBST (Tris buffered saline-Tween 20) for 2 h. The membrane was next incubated with respective primary antibody diluted with TBST containing 0.3% BSA and kept for overnight. Next day, the membrane was washed with TBS (Tris buffered saline) to remove unbound primary antibodies and next incubated with respective secondary antibody (alkaline phosphatase conjugated). The membrane was washed with TBS and colour was developed using NBT-BCIP mix in dark. Finally membranes were scanned for future documentation. After western blot, each bands (except PARP, as it was cleaved after treatment) were analyzed through ImageJ software for quantitative measurement. In brief, a constant unit area was selected and average density of the total band area of the respective proteins in control and treated groups were measured and represented graphically.

Statistical Analysis

The results were expressed in terms of mean ± SEM (n=6). The data were subjected to one way analysis of variance (ANOVA) followed by Tukey’s test using GraphPad InStat 3 software to establish statistical significance (*p<0.05, **p< 0.01, ***p< 0.001).

Results

Isolation, purification and characterization of NNSS2

TLC of N. naja shedded skin aqueous extract showed four zones (Zone 1-Zone 4) having Rf 0.65, 0.77, 0.827 and 0.98 (Figure 1a). RP-HPLC produced a single peak with retention time of 5.8 min and nomenclature as NNSS2 (Naja naja shedded skin zone 2) (Figure 1c). Spectral analysis of NNSS2 showed λ max at 287.2 nm (Figure 1d). MALDI-TOF analysis confirmed M.W of NNSS2 of 882.6 Dalton (Figure 1e). CD-spectral analysis of NNSS2 showed presence of negative bands between 217-229 nm, 236-331 nm, positive region between 230-235 nm, 196-215 nm. CD-spectral analysis showed 4% α-helix, 48% β sheet, 0.4% random coils and β-turn (Figure 2a). For FTIR and H1NMR see Table 1 and Figures 2b and 2c. X-ray crystallographic structural analysis showed presence of sharp peak which revealed that NNSS2 is a highly crystalline compound (Figure 2d).

Effect of NNSS2 on estrous cycle

NNSS2 (1 mg kg⁻¹ body weight, s.c) produced cessation of the estrous cycle at the diestrous phase for 5±1 day when vaginal lavage was observed every 24 h for 5 days (Data not shown).

Effect of NNSS2 on serum hormones and cytokines

NNSS2 (1 mg kg⁻¹ body weight, s.c) significantly decreased (p<0.001) progesterone on Day 5 and no significant change was observed in estradiol (p>0.05) when compared to diestrous control adult female mice but it significantly decreased serum P/E2 ratio (Table 2). NNSS2 (1 mg kg⁻¹ body weight, s.c) caused a significant increase in the serum IL-1β (p<0.001) and did not produce significant change in serum TNF-α (p>0.05) when compared with diestrous control adult female mice (Table 2).

Histopathological studies of NNSS2 treated mice uterus and ovary

NNSS2 (1 mg kg⁻¹ body weight, s.c) treatment produced loss of normal uterine architecture, thinning of endometrial, myometrial, perimetrial lining, constriction of luminal structure, formation of vacuolated structure and loss of uterine glands (Figure 3a). NNSS2 (1 mg kg⁻¹ body weight, s.c) treatment produced phase arrest in the follicle, showed distorted follicular structure, absence of matured follicle, ovum, granulosa and thecal cell lining as compared with control diestrous ovary (Figure 3b).

Figure 2: Characterization of NNSS2. (a) CD spectral analysis of NNSS2; (b) FTIR analysis of NNSS2; (c) H1NMR of NNSS2; (d) XRD analysis of NNSS2.

Figure 3: Histopathological studies of NNSS2 treated ovary and uterus. Histological structure of control diestrous uterus (A) NNSS2 treated mice uterus (B) Histological structure of control diestrous ovary (C) NNSS2 treated ovary (D) Control diestrous uterus. a,b) presence of perimetrium, myometrium, endometrium, uterine glands c) Star shaped lumen. Control diestrous ovary d) follicles at different stage of development. NNSS2 treated uterus x) absence of uterine glands y) presence of vacuolated structure, NNSS2 treated ovary e) distorted follicle f) immature constricted follicle g) phase arrest of follicles. All the sections were stained with haematoxylin and eosin and magnification was 65X.
Western immunoblot analysis

NNSS2 (1 mg kg⁻¹ body weight, s.c) treated uterus and ovary showed up regulation of Caspase 3,9, Bax, and down regulation of BCL2 (Figure 4a). NNSS2 (1 mg kg⁻¹ body weight, s.c) treated uterus and ovary showed PARP cleavage. NNSS2 (1 mg kg⁻¹ body weight, s.c) treated uterus and ovary caused down regulation of heat shock proteins (HSP70 and HSP90) as compared with uterus and ovary from diestrous control mice (Figure 4a). Quantitative analysis through ImageJ software also confirmed significant up regulation of caspase 3, caspase 9, Bax and down regulation of BCL2, HSP70 and HSP90 expression significantly (Figure 4b).

Discussion

This present study was designed to isolate a bioactive factor from Naja naja shedded skin. In the present study, a small molecular weight crystalline peptide NNSS2 of 882.6 Dalton was isolated from Naja naja shedded skin. CD spectral analysis indicated NNSS2 is crystalline and is composed of β sheet random coils and α helix. NNSS2 is crystalline and is composed of β-sheeted structure, N-H bonding, scissoring, bending and double bonds. Six amino acids had been identified so far namely asparagine, aspartate, glutamine, glycine, phenylalanine and tryptophan from.

NNSS2 caused cessation of the estrous cycle at diestrous phase for five days when observed microscopically, with corresponding change in ovarian and uterine histology. NNSS2 caused loss of three distinct layers, uterine glands and luminal structure with formation of vacuolated structure (Figure 3B). NNSS2 produced constricted ovarian structure, follicular atresia, phase arrests, absence of granulosa cell lining (Figure 3D), indicating uterine and damage and may be the cause behind phase arrest. The histological findings of both ovary and uterus supported tissue damage suggesting anovulation and phase arrest at diestrous. Tissue damage that had taken place in the uterus and ovary might have occurred either by apoptosis or by necrosis. Hence, to further elucidate the death pathway, expression of intracellular apoptogenic proteins of ovarian and uterine tissues were studied. In both the ovary and uterus increased the expression of caspase 3 and caspase 9 and cleaved PARP with no change in caspase 8 activities indicated involvement of intrinsic apoptotic pathway. PARP cleavage by caspase 3 is a considerable marker for apoptosis [8]. In the present study NNSS2 caused PARP cleavage indicating apoptosis through caspase 3 dependent pathway. No significant change in TNF-α level further confirmed absence of extrinsic pathway as TNF-α receptor is responsible for the activation of caspase 8 through the death receptor complex [9]. The intrinsic pathway of apoptosis gets activated by negative signal given by decreased level of progesterone and a positive signal from increasing expression of IL-1β, eventually activated the caspase dependent pathway [10]. This mitochondrial dependent pathway is regulated by BCL2 family of proteins which included proapoptotic protein like Bax and antiapoptotic protein like BCL2 by modifying mitochondrial membrane permeability. NNSS2 increased expression of proapoptotic protein Bax and decreased the expression of antiapoptotic protein BCL2, ultimately leading to the activation of effector molecule caspase 3. Activation of caspase 3 leads to the proteolytic cleavage of PARP responsible for morphological and biochemical changes in apoptosis [11,12].

HSP70 and HSP90 are molecular chaperone proteins associated with intracellular stress and play a critical role in cytoprotection [13]. HSP70 blocks apoptosis by binding to Apaf-1, prevented the
formation of apoptosomal complex further inhibiting caspase 3 induced proteolytic cleavage of PARP [14] HSP90 another chaperone protein directly binds to Apaf-1 inhibited apoptosome complex formation [15]. In the present study NNSS2 down regulated the expression of both HSP70 and HSP90 indicating apoptosis induction, suggesting follicular atresia and estrous cycle arrest at diestrous phase.

In the serum NNSS2 significantly decreased progesterone (Table 1); progesterone secreted from the corpus luteum is associated with anti apoptotic activity and exerts direct effect on luteal action [16]. Other than antiapoptotic activity progesterone is also associated with granulosa cell development [17], steroido bio synthesis and number of cell signalling activities and is also intricately associated with cytokine regulatory pathway [18,19]. Present study revealed that NNSS2 significantly increased serum IL-1β expression. It is a very well documented fact that increased expression of IL-1β is associated with induction of apoptosis by the involvement of ICE which converts the pro enzyme in to its active form [20]. From the present study it was revealed that NNSS2 influenced the estrous cycle in female albino mice by causing cessation of the estrous cycle at diestrous phase for 5 days.

Conclusion

NNSS2 caused anovulation hence it can lengthen a normal cycle by modulating the hormones, cytokines and intracellular apoptotic proteins. From the above findings it could be concluded that this study gave a novel insight into some of probable bioactive factors present in snake (N. naja) shed skin which remained unexplored until now.

References


Acknowledgements

We thankfully acknowledge University Grant Commission, New Delhi, India for partial financial assistance (Ref No: F.38-130 (SR), dated 19.12.2009 and we are thankful to the Department of Forests & Wild Life, Government of West Bengal, India (2105/WL/4R-1(PI- IX)) for kind permission of collection of the shed snake skin.