Cannabis Smoke Causes Up-Regulation of Akt and Bax Protein in Subfertile Patient’s Sperm Cells

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Abstract

Background: Emerging worldwide evidences in support of adverse effects of cannabis smoke indicate its significant role in declining male fertility. The aim of the present study was to compare the percentage of damaged sperm cells and the expression profiles of cell survival protein p-Akt and pro-apoptotic protein Bax in non-smoker, tobacco smoke addicted and cannabis smoke addicted subfertile subjects.

Method: Sperm samples were collected from 80 male subjects of reproductive age group in Southern Assam of North-East India. 46 (57.5%), 25 (31.25 %) and 9 (11.325%) of these subjects were found to be cigarette smokers, cannabis smokers and non-smokers respectively. ROS levels in semen samples were measured by chemiluminescence assay. Sperm DNA integrity were assessed by acridine orange test, toluidine blue staining and TUNEL assay. Expression profiles of p-Akt and Bax were observed by flow cytometry and western blot analysis.

Results: Among three groups, the cannabis smoke addicted subjects showed the highest level of seminal ROS production along with the highest percentage of sperm DNA damage, chromatin abnormalities and apoptotic cells. High expression of Bax and low expression of p-Akt was observed in non-smoker and tobacco smoke addicted subjects. Conversely, cannabis smoke addicted group showed the highest expression of both p-Akt and Bax proteins.

Conclusion: The present study indicates cannabis smoke addiction to be more detrimental for male reproductive health compared to the tobacco smoke. The over-expression of both Akt and Bax proteins among cannabis smokers suggest that the up-regulation of pro-survival protein Akt, during sperm meiotic division could have triggered the oxidative apoptosis of sperm cells via the up-regulation of pro-apoptotic protein Bax.

Keywords: Infertility; Cannabis addiction; ROS; DNA damage; Apoptosis; Akt; Bax

Introduction

Since last few decades, a sharp decline in male fecundity has been observed all over the world. In addition to congenital abnormalities, environmental and occupational exposures, changed lifestyle factors were also found to impact male reproductive health [1-3]. Several studies reported direct association of excessive tobacco and alcohol consumption with the declining male fertility [4,5]. Emerging studies have also correlated addiction to cannabis smoke with poor semen quality of men [6,7]. Contents of cannabis smoke reduce antioxidant defence mechanism and increase oxidative stress in seminal plasma [8,9].

It has been estimated that around 150 million people across the globe were addicted to cannabis in the beginning of this millennia [10]. The hallucinogenic effects caused by this recreational drug, entices people of different age groups and socio-economic classes in different countries all over the world [11,12]. ‘Cannabis’ is a generic term used for marijuana, hashish and hash oil and it is derived from the Cannabis sativa plant [13]. Δ9-Tetrahydrocannabinol (THC) is the unique compound of cannabis with major psychoactive effects and is said to act upon a specific cannabinoid receptor (CB1) in the brain [14,15]. In the 1990s, it was observed that cannabinoid compounds are naturally synthesized in human body from fatty acid derivatives termed as endogenous cannabinoids or endocannabinoids [16,17]. Endocannabinoids modulate several pathophysiological processes like neuropathic pain, movement disorders such as Parkinson disease, Huntington disease and many other conditions like atherosclerosis, obesity as well as reproductive health [18]. However, the association between cannabis smoking and cancer is highly disputed as different case-control studies had inferred different results [19,20]. Both cannabinoid [21] and nicotine [22] receptors are coupled to the protein kinase B (Akt) signalling pathway. Akt is a serine threonine kinase which induces anti-apoptotic signal and inhibits apoptosis. However, the role of Akt coupled with cannabinoid receptor, varies from one disease to another. In Alzheimer’s disease, activation of cannabinoid receptor [23] and subsequent activation of Akt pathway can prevent brain cell death caused by the production of beta-amyloid protein [24]. Conversely, the Akt signalling cascade inhibits apoptosis and promotes tumour progression in cancer. Downstream signalling cascade of Akt has Bax protein. Bax is a member of the Bcl-2 (pro-apoptotic) family of proteins with accelerates apoptosis induced by a variety

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of stimuli [25,26]. It promotes mitochondrial cytochrome c leakage by dimerization and insertion into mitochondrial membrane which eventually leads to the nuclear fragmentation of the cell [27].

Several plausible theories explain the probable mechanism of abnormal sperm formation [28,29]. Recent advances in the field of genetics and molecular biology provided a great impetus to explore sperm chromosomal abnormalities at molecular level [30]. These studies suggested that mutation during sperm meiotic division may trigger DNA fragmentation and subsequent apoptosis of the cells in humans and experimental animals [31-33]. On the other hand, several literatures revealed that DNA damage in sperm cells is associated with elevated levels of reactive oxygen species (ROS) production causing oxidative stress [34-36]. At lower level, ROS play an important role in sperm maturation and functions such as capacitation and acrosome reaction [37]. However, increased ROS production, beyond the antioxidant capacity limit in seminal plasma often resulted in cell and DNA damage. The polyunsaturated fatty acid (PUFA) content makes the sperm cells susceptible to the peroxidation in the presence of seminal ROS, resulting in the up-regulation of apoptotic pathways [38-42].

The aim of the present study was to compare the seminal oxidative stress, percentage of sperm DNA damage and expression profiles of Akt and pro-apoptotic protein Bax in sperm cells of three groups of non-smoker, tobacco smoke addicted and cannabis smoke addicted subfertile subjects in Southern Assam of North-East India.

Materials and methods collection of semen samples

Total (n=80) semen samples were collected for the study with 96% power from the subjects of reproductive age group (25–40 years) living in Southern Assam of North-East India. Each subject was asked to sign an informed consent form approved by Institutional Ethical Committee (IEC), Assam University, Silchar and were asked to fill in a questionnaire. The subjects were instructed 2-3 days of sexual abstinence prior to semen ejaculation by masturbation into a sterile, wide mouthed, labeled container. Semen samples were prepared and the air-dried and fixed in freshly prepared 96% ethanol-water. Under light microscopic evaluation, a total of 300 spermatozoa were counted in different areas of each slide using oil immersion with ×200 magnifications. Sperm cell heads with good chromatin integrity stained light blue and those of diminished integrity were deep blue in colour. 

Measurement of reactive oxygen species

ROS levels in seminal ejaculates were measured by chemiluminescence assay using luminol (5-amino-2, 3-dihydro-1, 4-phthalazinedione; Sigma, St Louis, MO) as the probe. 10 µL of 5 mmol/L luminol prepared in dimethyl sulfoxide (Sigma Chemical) were added to 400 µL of the washed sperm suspension. Negative controls were prepared by replacing the sperm suspension with 400 µL phosphate buffered saline. Positive control included 400 µL of PBS and 50 µL of hydrogen peroxide (30%; 8.8 M) in triplicates. Chemiluminescence was measured for 15 minutes using a luminometer (Promega Glomax 20/20 Luminometer). The results were expressed as relative light units (RLU)/sec/10⁶ sperm [43].

Tests for sperm DNA integrity acridine orange test (AOT):

Acridine orange test (AOT) is a simple microscopic procedure based on acidic conditions to denaturant DNA followed by staining with acridine orange. The AOT measures the metachromatic shift of AO fluorescence from green (native DNA) to red (denatured DNA). Sperm smears were fixed in Carnoy's solution (60% ethanol, 30% chloroform and 10% glacial acetic acid) and were subsequently stained with acridine orange solution (0.02% acridine orange in citratephosphate buffer, pH 2.5) according to the procedure of Tejada et al. [44]. After 5 minutes of staining, each smear was washed with distilled water, covered with a cover slip and sealed with a synthetic resin to prevent the smear from drying. Smears were examined within 1 day using a Leica fluorescence microscope (Leica DM 4000 B) with the following filter combination: 450 nm to 490 nm excitation and 520 nm barrier filters. All spermatozoa with fully compacted nuclei were examined. Nuclei of 300 spermatozoa were scored on the basis of their fluorescence (green/red). In recent studies percentage of DNA fragmentation has been represented as DFI (DNA fragmentation Index).

\[ \text{DFI} = \frac{\text{% of Red fluorescence}}{\text{% of Total (red+green) fluorescence}} \]

Toluidin blue staining for sperm chromatin abnormalities

Toluidin blue (TB) staining had been reported to be a sensitive test for incomplete DNA structure and packaging [45,46]. A thin smear was prepared and the air-dried and fixed in freshly prepared 96% ethanol-acetone (1:1) at 4°C for 1 hour. After that the samples were hydrolyzed in 0.1 N HCL at 4°C for 5 minutes. Thereafter, the slides were rinsed 3 times in distilled water for 2 minutes and finally stained with 0.05% TB (in 50% McIlvaine's citrate phosphate buffer, pH 3.5, Merck) for 5 minutes at room temperature. The slides were rinsed briefly in distilled water. Under light microscopic evaluation, a total of 300 spermatozoa were counted in different areas of each slide using oil immersion with ×200 magnifications. Sperm cell heads with good chromatin integrity stained light blue and those of diminished integrity were deep blue in colour.

TUNEL assay of sperm cells

Apoptosis in sperm cells was determined by the TUNEL technique using TACS® 2 TdT DAB kit (Trevenag, Catalogue No.4810-30-K). The terminal deoxynucleotidyl transferase-mediated (TdT) deoxyuridine triphosphate (dUTP) nick end labeling assay (TUNEL) is a direct quantification of sperm DNA fragmentation. dUTP is incorporated at single stranded and double stranded DNA fragments in a reaction catalyzed by the enzyme TdT. The DNA breaks based on acidic conditions to denaturant DNA followed by staining, each smear were washed with distilled water, covered with a cover slip and sealed with a synthetic resin to prevent the smear from drying. Smears were examined within 1 day using a Leica fluorescence microscope (Leica DM 4000 B) with the following filter combination: 450 nm to 490 nm excitation and 520 nm barrier filters. All spermatozoa with fully compacted nuclei were examined. Nuclei of 300 spermatozoa were scored on the basis of their fluorescence (green/red). In recent studies percentage of DNA fragmentation has been represented as DFI (DNA fragmentation Index).

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Flow cytometric analysis

The sperm cell pellets were suspended in 900 µL in PBS and centrifuged at 4000 rpm for 5 minutes at room temperature. The pellet was resuspended in 900 µL PBS and this step was repeated twice. The final pellet was resuspended in 100 µL PBS and the number of cells were counted using haemocytometer. 5 x 10⁶ sperm cells were dissolved again in 100 µL PBS and the suspension incubated with 0.5% TritonX-100 for 20 minutes at room temperature followed by PBS wash. The


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Volume 6 • Issue 4 • 1000247
cells were next incubated with Alexa Fluor 488 tagged anti-p-Akt and Bax antibodies for 1 hour at room temperature. After two consecutive PBS washes, the cells were fixed using 100μl paraformaldehyde. Prior to acquiring, 200 μl sheath fluid was added. Expression of proteins were obtained using BD FACS Calibur machine.

Protein extraction and western blot analysis

To prepare a whole sperm cell extract aliquots of 0.5 ml samples were centrifuged at 7500 g for 5 minutes at room temperature and the supernatant was discarded. The resulting sperm pellet was resuspended in 500 μl extraction medium (2% SDS, 28% sucrose, 12.4 mM N,N,N9,N9-tetramethylethylenediamine and 185 mM Tris–HCl, pH 6.8) and immediately incubated for 5 minutes at 100°C. After centrifugation the concentration of proteins in the supernatant was measured using BSA kit (Thermo Scientific) as per the manufacturer’s protocol. Finally, extracts were stored at -20°C until used for western blot analysis [48]. The proteins were subjected to SDS-PAGE and electro-transferred to nitrocellulose membranes. The membrane was blocked with 5% non-fat dry milk in Tris-buffer saline (20 mM Tris HCl and 137 mM NaCl, pH-7.5) for 1 hour at room temperature. Immunogenicity was detected by incubation of the membrane overnight with appropriate primary antibodies p-Akt and Bax (1:200) and specific proteins were detected by the enhanced chemiluminescence system (Biovision ECL Western Blot substrate). The Signal intensity of the band was detected by densitometer (Bio Rad, GS 800).β-actin expressions were tested for confirming equal distributions of proteins.

Statistical analysis

Statistical analyses were performed using Graph Pad Prism 5.00 gold software package. t-test was used to compare the means. Median values were calculated as 25th and 75th percentile. Spearman rank correlation coefficient (r) was calculated to find the correlation between variables. P ≤ 0.05 was taken to be statistically significant.

Results

The results of the present study showed no significant deviation (p>0.05) in the mean age (in years) and average BMI (body mass index) (Kg/m²) of non-smoker, tobacco smoke addicted and cannabis smoke addicted subjects [Mean age, non-smokers-29.28 ± 1.97, tobacco smokers-29.63 ± 2.95, cannabis smokers - 29.08 ± 2.43; BMI, non-smokers-24.14 ± 1.28, tobacco smokers-24.08 ± 1.44, cannabis smokers-24.22 ± 1.41] which could be the contributing factors of their fertility status.

ROS production in semen samples

The oxidative stress was measured by assessing the production of ROS level in seminal ejaculates of three groups of subjects. The highest and the lowest ROS production was observed in subjects addicted to cannabis smoke [161.5 (154; 163.25) (RLU/sec/10⁶ sperm)] and in non-smoker group respectively [123.25 (132; 136.8) (RLU/sec/10⁶ sperm)] whereas, the tobacco smoke addicted subjects showed the intermediate level of ROS production [144.5 (142; 153.5) (RLU/sec/10⁶ sperm)] (Table 1).

Acridine orange fluorescence study of sperm nuclei

Table 1 summarizes significant differences (p<0.001) in the percentages of sperm cell DNA damage (red fluorescence; single stranded/denatured DNA) among three groups of subjects [cannabis smokers- 51.77 ± 4.61, non-smokers-36.33 ± 3.97, tobacco smokers-41.82 ± 8.14 respectively] (Figure 1A-1D).

Assessment of sperm chromatin abnormalities among different subjects

The percentage of abnormal sperm chromatin structure and its

<table>
<thead>
<tr>
<th>Variables</th>
<th>Non-smoker</th>
<th>Tobacco smoker</th>
<th>Cannabis smoker</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n=9)</td>
<td>(n=46)</td>
<td>(n=25)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ROS production [(RLU/sec/106 sperm)]</td>
<td>123.25 (132; 136.8)**</td>
<td>144.5 (142; 153.5)**</td>
<td>161.5 (155; 163.6)**</td>
<td></td>
</tr>
<tr>
<td>% of Sperm DNA damage (DFI)</td>
<td>36.33 ± 3.97 (47.53)</td>
<td>41.82 ± 8.14a (53.32)</td>
<td>51.77 ± 4.61a,b (62.17)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% of Sperm chromatin abnormalities</td>
<td>36.81 ± 4.46</td>
<td>45.06 ± 2.80a</td>
<td>52.83 ± 4.67a,b</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% of Apoptotic Sperm Cells</td>
<td>37.93 ± 3.84</td>
<td>42.92 ± 1.98a</td>
<td>52.44 ± 2.87a,b</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

** Values are represented as median (25th; 75th percentile)
Percentages are calculated as Mean ± SD
DFI: DNA Fragmentation Index
ap<0.01 statistically significant compared to non-smoker group.
bp<0.01 statistically significant compared to tobacco addicted group.

Table 1: Relative ROS production and percentage of Sperm cell DNA integrity among three groups of patients.
condensation was compared between the three groups by Toluidine Blue staining (Table 1). The results showed highest percentage (52.83 ± 4.67) of sperm chromatin abnormalities among cannabis smokers. Tobacco smokers (45.06 ± 2.80) and non-smokers (36.81 ± 4.46) also showed moderately high percentage of sperm chromatin abnormalities (Figure 2A-2D).

**Variation in the percentage of apoptotic sperm cells among three groups**

The TUNEL assay results were again consistent with the DNA fragmentation results. The overall incidence of sperm cell apoptosis was significantly different (p<0.001) among the three groups of subjects (Table 1). The highest percentage (52.44 ± 2.87) of apoptotic sperm cells were observed again in subjects addicted to cannabis smoke. Tobacco smokers (42.92 ± 1.98) and non-smokers (37.93 ± 3.84) also showed moderately high percentage of apoptotic sperm cells (Figure 2A-2D).

**Correlation between increasing seminal ROS production and percentage of DNA damage, chromatin abnormalities and apoptosis in sperm cells**

The Spearman correlation analysis showed significant (p<0.0001) positive correlations between increasing seminal ROS production and percentage of sperm DNA damage, chromatin abnormalities and the percentage of apoptotic sperm cells in cannabis smoke addicted subjects. However, no significant association between these parameters was observed among non-smokers and tobacco smokers (Table 2).

**Differential expression of P-Akt and Bax protein in sperm cells of three groups**

The flow cytometric analysis showed high positive expression of Bax protein in all three groups of subjects, whereas high expression of p-Akt was observed among cannabis smokers only (Figure 4A-4D). This observation was further confirmed by the western blot analysis results (Figure 3). The quantitative densitometric analysis showed highest percentage (1.27 ± 0.15) of Bax protein expression in cannabis smoke addicted subjects compared to the non-smokers and tobacco smokers (0.845 ± 0.052 and 1.01 ± 0.02 respectively). The highest percentage (1.08 ± 0.090) of p-Akt was also observed in cannabis patients' sperm cells, whereas the other two groups [non-smokers (0.645 ± 0.01), tobacco smokers (0.86 ± 0.060)] showed comparatively low percentage of p-Akt expressions (Figure 5A and 5B).

**Discussion**

Worldwide evaluation of different infertility cases across the world suggests that the largest percentage of patients experience idiopathic infertility [49]. A proper understanding of the underlying mechanism causing male factor infertility requires insightful analysis of molecular and cellular events involved in human spermatogenesis [50]. Spermatogenesis is a complex process which produces mature sperm cells from undifferentiated germ cells via mitosis and meiosis. Previous work by Zhuang et al. [33] showed that chromosomal aberration during spermatogenesis causes apoptosis of haploid sperm cells and decline in the fertilizing potential [33]. An important review work by Diemer and his co-worker discussed about different...
genetic disorders during spermatogenesis [51]. A study by Jamieson et al. [52] showed that chromosomal aberration has profound effect on spermatogenesis and in some cases few mature spermatozoa are produced which eventually undergo apoptosis. On the other hand, increasing evidences have strongly correlated declining semen parameters with addiction to tobacco and cannabis smoke [4,6,7]. Some of the significant studies by Rossato et al. and Kolodny et al. have linked rampant cannabis consumption with declining fertility rate in
men [53,54]. In particular, it has been revealed by the works of Schuel H et al. and Berdyshnev EV et al. [55,56] that cannabinoids influence human sperm functions, leading to a reduction of their fertilizing ability in both invertebrates and vertebrates. In the present study, we observed the highest percentages of seminal ROS production, damaged sperm DNA, chromatin abnormalities, and apoptotic sperm cells in cannabis addicted individuals compared to the non-smokers and tobacco smokers. The percentage of apoptotic sperm cells was also high in the tobacco smokers which could be one of the predominant causes of their declining fertility status. The comparative analysis of our study suggested that cannabis consumption could be more deleterious for the fertility status of an addicted individual compared to those addicted to tobacco smoke. Previous works by Yamaguchi and his co-worker [57] as well as Gardai et al. [58] showed that over-expression of Akt suppresses the localization of Bax to mitochondria and subsequent apoptosis of the cell. Interestingly, in this study we observed high expression of both Akt and pro-apoptotic protein Bax in sperm cells of cannabis smokers but comparatively lower expression of Akt in the other two groups of subjects. Although the precise molecular mechanism behind this phenomenon is yet to be explored, one plausible hypothesis would be up-regulation of Akt during sperm meiotic division. Mutation during sperm meiotic division might trigger the up-regulation of Akt protein or the activation of Akt may occur naturally. A work by Andersen et al. showed that activation of Akt and the subsequent phosphoinositide 3-kinase signaling pathway promotes cell growth and differentiation in oocytes [59]. Furthermore, another work by Veronique Nogueira et al. suggested that Akt induces oxidative senescence and sensitizes cells to undergo apoptosis [60]. It can be hypothesized that the up-regulation of Akt and Bax could not induce the survival of haploid sperm cells as they do not divide. Alternatively, Akt triggered the oxidative apoptosis of the sperm cells via pro-apoptotic Bax protein. This could be the possible reason behind the synchronous expression of Akt and Bax in the sperm cells of cannabis smokers. In conclusion, the present study clearly shows that cannabis smoke is far more detrimental for the male reproductive health. Additionally, it causes up-regulation of Akt protein which in turn triggers the oxidative apoptosis of sperm cells via Bax protein. Although in-depth molecular studies need to be executed for a better understanding of these phenomena.

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Reference


Page 6 of 7 Volume 6 • Issue 4 • 1000247J Addict Res Ther ISSN:2155-6105 JART an open access journal


