PUFAs from Stem Bark of *Alstonia boonei* Synergistically Modulates Diabetic, Hepatic and Androgenic Damage by Low Expression of COX-2 and iNOS in Rats

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**Abstract**

PUFAs, from the stem bark extract of *Alstonia boonei* (SBEAB) was hypothesized to possess anti-inflammatory, antioxidant, anti-diabetic, pro-spermatogenic and hepatoprotective activities. The present study investigated the possible biochemical and molecular mechanisms underlying the hepatoprotective and testoprotective effects of SBEAB in diabetic rat. Biomarkers of hepatic and testicular damage, histological and immunohistochemical techniques were used. The expression of cyclooxygenase (COX-2) and inducible nitric oxide synthase (iNOS) were also estimated. SBEAB administered orally at dose of 100 mg/kg for 14 days significantly lowered the activities of serum transaminases and MDA levels induced by single intraperitoneal administration of streptozotoxin (STREP) (80 mg/kg) and preserved the integrity of both hepatocytes and spermatocytes. Also, SBEAB elevated the STREP-induced reduced activities of Δ5-17β-HSD and Δ5-17β-HSD with corresponding decrease in the activity of CAT. SBEAB inhibited the STREP induced expression of COX-2 and iNOS. The protective effect of SBEAB was compared to that of metaglomide (METAG), an established anti-diabetic drug. METAG treatment on hepatic damage was most efficacious in diabetic rats; followed by post and pre-treatment respectively while pre-and post-treatment were more efficacious on testicular damage than anti-diabetic drug. Furthermore, pre and post-treatment were more efficacious in preventing pro-inflammation and testicular cancer in diabetic rats than METAG-administration. We therefore concluded that the repression of genes encoding COX-2 and iNOS proteins by SBEAB validates the molecular basis of testicular protection and further suggests the links between the hepatocellular damage and male reproductive dysfunctions in diabetic individuals.

**Keywords:** SBEAB; PUFAs; Pro-inflammation; Synergy; Diabetic rats

**Introduction**

Diabetes mellitus (DM) is one of the well-known public health risks in contemporary societies and its occurrence is geometrically increasing. A recent report speculated that before 2030, diabetes would have affected approximately 40% increase of the world population [1]. Also, male reproductive challenges of modern societies reveal that the increased incidence of DM was closely linked with falling birth rates and infertility [2,3]. This could be attributed to the disturbing increase of diabetic men in reproductive age. Previous work reported that patients with type 1 diabetes (T1D) were identified before the age of 30 [4] and that the number of young persons with T1D and type 2 diabetes (T2D) were uncontrollably high [5]. Further investigations also reported that sexual disorders, such as erectile dysfunction [6] or retrograde ejaculation [7], were manifested in diabetic individuals and generally resulted in a reduced libido [8]. Although, the decline in reproductive health of male diabetics can be attenuated in patients but the molecular mechanism of metabolic pathways aside glucose transport to the cells, remain undiscovered and obscure [9,10]. Essentially, the links between male reproductive dysfunctions and hepatic damage in diabetic induced rats are scarcely elucidated.

Cyclooxygenase-2 (COX-2) and inducible nitric oxide synthase (iNOS) can be induced quickly by pro-inflammatory mediators, endotoxins and carcinogens [11]. Some recent studies have demonstrated that carcinogens were capable of provoking COX-2 expression via activation of transcription factors [12,13]. However, targeted inhibitions of COX-2 and iNOS were recently recognized as the molecular basis for cancer prevention [13]. Also, health-promoting plants origin are linked to the presence of essential phenolic phytochemicals including phenolic acids, PUFAs and flavonoids [14-16]. Phenolic compounds are secondary metabolites of plants which exhibit strong antioxidant activity and play crucial role in health promotion and disease aversion [17].

The herbs produced from stem barks have been used as healing mediators and as anti-hepatic damage for thousands of years [18]. Herbs have also been recognized to possess medicinal properties [19]. The herbal therapies with high phenolic content are preponderant in the West Africa especially the tropical Africa [20,21]. They are administered with a drink to treat colds and several assaults [18,20]. The present stem bark of this study belongs to the family of apocynaceae (*Alstonia boonei*); also, known as God’s tree. It is employed as an alternative medicine for many complicated ailments and used for medicinal purposes such as antirheumatic [22], pain–reliever, antipyretic, antimicrobial, and antibiotic properties [23]. Recent findings reported the hypoglycaemic properties of stem bark
The aim of the present study was to evaluate the effect of combined PUFAs of stem bark of *Alstonia boonei* (SBEAB) on activity of enzymes linked to testicular pro-inflammation, liver and androgenic damage in streptozotocin (STREP)-induced diabetic rats.

**Materials and Methods**

**Sample selection**

Fresh sample of stem bark of *Alstonia boonei* was purchased from the local market, in Ota metropolis, Ogun State, Nigeria. Authentication of the plant was carried out in the Department of Biology, University of Ibadan, Nigeria. Adult male wistar strain albino rat was purchased from the Biochemistry Department animal colony, University of Ibadan, Nigeria and maintained ad libitum on commercial diet and water.

**Chemicals and reagents**

STREP was purchased from Sigma Chemical Co. (St Louis, MO, USA). Rabbit polyclonal COX-2 and iNOS antibodies were products of Cayman Chemical Co. (Ann Arbor, MI, USA). Bovine Serum Albumin (BSA), hydrogen peroxide, trichloroacetic acid (TCA) and thiobarbituric acid (TBA) were purchased from Sigma (St Louis, MO, USA). All the kits used for the bioassay were sourced from Randox Laboratories Ltd. (Crumlin, Dublin, Northern Ireland, UK). All other reagents used were in the purest form available commercially.

**Extraction of stem bark *Alstonia boonei***

The bark of the stem was thoroughly washed in distilled water to remove any contaminant, chopped into small pieces before being milled. The ethanol extract of the stem was subsequently prepared by soaking the grinded sample (10 g) in ethanol (200 ml) for about 24 h at 37°C; the mixture was filtered and the filtrate was concentrated by rotator evaporator designated as SBEAB and stored in the refrigerator for subsequent analysis.

**Gas Chromatography - Mass Spectrum Analysis (GC-MS)**

The crude extract was subjected to column chromatography over silica-gel (100-200 mesh) and eluted with ethanol. The ethanol fraction of stem bark of *Alstonia boonei* was taken for GC-MS analysis. GC-MS analysis was carried out on a GC Clarus 500 Perlin Elma system comprising a AOC-20i auto-sampler and gas chromatography interfaced to a mass spectrometer (GC-MS) instrument employing the following conditions: column Elite-1 fused silica capillary column (30 x 0.25 mm x 1D 1 EM df, composed of 100% Dimethyl polysiloxane), operating in electron impact mode at 70 eV; helium (99.999%) was used as a carrier gas at a constant flow of 1 ml/min and an injection of volume of 0.5 EI was employed (split ratio of 10:1 injector temperature 250°C), ion-source temperature 280°C. The oven temperature was programmed from 110°C (isothermal for 2 min) with an increase of 10°C/min, to 200°C then 5°C/min to 280°C, ending with a 9°C/min isothermal at 280°C. Mass spectra were taken at 70 eV; a scan interval of 0.5s and fragments from 40 to 550 Da.

**Identification of components**

Interpretation on mass spectrum GC-MS was conducted using database of National Institute Standard and Technology (NIST) having more than 62,000 patterns. The spectrum of the unknown components was compared with the spectrum of the known components stored in the NIST library. The molecular name and their percentage composition of the components of the test materials were ascertained.

**Animal treatment**

Adult male Wistar rats (200–300 g) from the Central Animal House of the Bells’ University of Technology, Ota, Nigeria were used in this experiment. The animals were maintained at a constant temperature (22 ± 2°C) on a 12 h light/dark cycle with free access to food and water. Animal care and handling was done according to the institutional guidelines of Nigeria Academy. The study was approved by the Institutional Animal Ethical Committee.

**Experimental protocol**

The rats were acclimatized for two weeks and randomly divided into five groups of eight animals each (n=8). Group 1 (Control): serve as the normodiabetic control group; Group 2 (Induced): serve as the diabetics group induced with STREP plus nicotinamide adenine dinucleotide (NAD) (STREP+ NAD); Group 3 serve as the positive control induced with STREP plus metaglomide (METAG) plus NAD (STREP + METAG+ NAD); Group 4 (Pre-treatment): serve as diabetic group pre-treated with 200 mg/kg body weight of stem bark *Alstonia boonei* extract (SBEAB before+ NAD + STREP after) [24]; Group 5 (Post-treatment): serve as diabetic group post-treated with 200 mg/kg body weight of stem bark *Alstonia boonei* extract (STREP after + NAD + STREP before) 120 mg/kg body weight of nicotinamide (NAD), 80 mg/kg body weight of streptozotocin and 5 mg/kg body weight of metaglomide (METAG) were used. In the normodiabetic groups, the animals received water by gavage throughout the entire experiment to be subjected to the same stress (normodiabetic groups). The experiment lasted for two weeks (14 days).

**Diabetic determination**

The animals were subjected to overnight fast prior to the induction of diabetes. Streptozotocin (STREP) freshly prepared in citrate buffer (0.01 M, pH 4.5). Rats in groups 2, 3, 4 and 5 received a single intraperitoneal dose of STREP (80 mg/kg) for 72 hr [25]. Blood samples were taken by tail vein puncture and glucose levels was monitored using automatic auto-analyzer (Fine test Auto-coding TM). Animals with blood glucose ≥ 200 mg/dl after 72 h were considered diabetic and were used in the study. The non-diabetic animal received 1 ml of 0.1 M citrate buffer intraperitoneally. During the experiment, fasting blood glucose was monitored at 3 days interval. The blood was rapidly collected by direct heart puncture and the liver and testes were isolated, rinsed in cold saline (0.9%) and homogenized in phosphate buffer (pH 7.4). The protective effect of *Alstonia boonei* on STREP-induced diabetic was compared to that of metaglomide (5 mg/kg) which has been confirmed to have antidiabetic effects [26]. At the end of each experiment, rats were sacrificed by cervical dislocation. Blood was collected by heart puncture for serum isolation. Liver and testes were rapidly frozen in liquid nitrogen and kept at −80°C for later analysis.
were excised, rinsed in physiological saline and stored at 4°C until use for histopathology and immunohistochemistry.

Measurement of liver enzyme activities

The activity of Liver enzymes [(Alanine aminotransferase (ALT) and aspartate aminotransferase (AST)] were measured using commercially available kits (Randox Laboratories Kits, St Louis, MO, USA) [27].

Determination of lipid peroxidation and catalase activity

Lipid peroxidation was assessed in terms of malondialdehyde (MDA) formation in the rat liver 10,000 g supernatant fractions. The measurement of thiobarbituric acid reacting substances (TBARS) was performed as described previously [28]. MDA was quantified and measured as nmole/mg protein. The estimation of catalase [CAT] activity using hydrogen peroxide as substrate was measured according to the method of Clairborne [29]. Protein concentration was determined by the method of Lowry et al. [30].

Histological studies and immunohistochemical staining

Liver and testes specimen were fixed in 10% neutral buffered formalin, embedded in paraffin, and sectioned. After deparaffinization and dehydration, the paraffin blocks were stained with hematoxylin and eosin for microscopic examination. For immunohistochemical analysis, the enzymatic activity of endogenous peroxidases in the testes section was first blocked with 3% hydrogen peroxide, followed by incubation with rabbit polyclonal anti-rat COX-2 and iNOS antibody (Cayman Chemical CO., Ann Arbor, MI, USA) at room temperature for 40 min. The peroxidase binding sites were detected by staining with 3,3-diaminobenzidine tetrahydrochloride (Dako, Glostrup, Denmark). Finally, counterstaining was performed using Mayer’s hematoxylin.

Statistical analysis

The data in each group were expressed as mean ± standard deviation. A one way analysis of variance (ANOVA) was used to analyze the results and Duncan multiple test was used for the post hoc [31]. Statistical package for Social Science (SPSS) 17.0 for windows was used for the analysis and the least significance difference (LSD) was accepted at P<0.05.

Results

In other to establish the phyto-active compounds responsible for the biochemical activities of SBEAB, the ethanol extract was quantified for various phytochemicals. As shown in Table 1, GC-MS Chromatogram (peak not shown) provided 22 different poly unsaturated fatty acids (PUFAs). The compound names and their percentage compositions of the unsaturated fatty acids identified in SBEAB were given (Table 1). As observed, GC-MS spectrum showed the predominant presence of 1,2-epoxyoctocloctane trans-2-nonal, 1-(2-hydroxyl-1-nitroethyl)cyclohexanol and Spiro[1,3-dioxolane-2,2'-6(6,7]diazobicyclo [3.2.2 non-6-ene].

The protective effect of SBEAB on diabetic-induced liver injury was evaluated by determining the levels of AST and ALT. As shown in Table 2, SBEAB pre and post-treatment as well as metaglomide (anti-diabetic drug) significantly (P<0.05) lowered the activities of AST and ALT in diabetic rats which were released into serum as an end point of hepatic damage. In addition, there was a significant decrease in intracellular CAT activity of testes of diabetic-induced rats (Figures 1 and 2) with concomitant increases in MDA production of both liver and testes (Figure 1).

| Phytochemical                      | % composition |
|------------------------------------|---------------|
| Trans-3-hepten-1-ol                | 3.3           |
| 1,2-epoxycloctane                 | 9.0*          |
| Cyclohexylmethyl trifluorocetane   | 3             |
| Trans-6-nitro-2-hexene             | 3             |
| Trans-5-decene                    | 3.3           |
| Cis-2-nitro-2-hepten-1-ol          | 3             |
| Trans-2-nonenal                   | 11.1*         |
| 1,2-epoxydocane                   | 1.5           |
| 10-undecyn-1-ol                   | 3.5           |
| 4-chloro-3-hexyltetrahydro-2H-pyran| 1.5           |
| 1-(2-hydroxy-1-nitroethyl)cyclohexanol| 9.2*       |
| 13-tetradec-11-yn-1-ol             | 3.8           |
| Cis, cis-6,9-pentadecadien-1-ol    | 3.5           |
| 2-pentadecyn-1-ol                 | 1.7           |
| Cis-10-pentadecen-1-ol             | 1.5           |
| Palmitic acid                     | 1.5           |
| Oleic acid                        | 1.7           |
| α-linolenic acid                  | 3.8           |
| n- nonadecanoic acid              | 1.5           |
| Octadecanoic acid, 2-(2-hydroxyethoxy) ethyl ester| 1.6 |
| Spiro[1,3-dioxolane-2,2'-6(6,7]diazobicyclo[3.2.2 non-6-ene] | 9.4* |

*Predominantly higher

| Phytochemical | % composition |
|---------------|---------------|
| 1,2-epoxycloctane | 9.0*        |
| Cyclohexylmethyl trifluorocetane | 3 |
| Trans-6-nitro-2-hexene | 3 |
| Trans-5-decene | 3.3 |
| Cis-2-nitro-2-hepten-1-ol | 3 |
| Trans-2-nonenal | 11.1* |
| 1,2-epoxydocane | 1.5 |
| 10-undecyn-1-ol | 3.5 |
| 4-chloro-3-hexyltetrahydro-2H-pyran | 1.5 |
| 1-(2-hydroxy-1-nitroethyl)cyclohexanol | 9.2* |
| 13-tetradec-11-yn-1-ol | 3.8 |
| Cis, cis-6,9-pentadecadien-1-ol | 3.5 |
| 2-pentadecyn-1-ol | 1.7 |
| Cis-10-pentadecen-1-ol | 1.5 |
| Palmitic acid | 1.5 |
| Oleic acid | 1.7 |
| α-linolenic acid | 3.8 |
| n- nonadecanoic acid | 1.5 |
| Octadecanoic acid, 2-(2-hydroxyethoxy) ethyl ester | 1.6 |
| Spiro[1,3-dioxolane-2,2'-6(6,7]diazobicyclo[3.2.2 non-6-ene] | 9.4* |

Table 1: Characterisation of phyto-active components (PUFAs) in ethanol extracts of stem bark Alstonia boonei (SBEAB).

Histological examination of liver samples also agreed with the above observations. Thus, liver specimens from diabetic rats revealed hepatocyte necrosis towards the centrilobular region (Figure 3B) when compared with the control liver (Figure 3A). In metaglomide, SBEAB pre and post-treated rats, the integrity of the hepatocytes was moderately well preserved (Figures 3C–3E). Thus, the defensive effect of metaglomide on hepatic damage was most efficacious in diabetic rats when compared to SBEAB post and pre-treatment respectively.

Table 2: Comparison of biochemical activities of SBEAB post and pre-treatment with metaglomide (SBEAB).

| Animal Group | ALT (U/l) | AST (U/l) |
|--------------|-----------|-----------|
| Control      | 0.01490 ± 0.000083a | 0.02334 ± 0.00025a |

*Predominantly higher
Table 2: Effect of stem bark extract *Alstonia boonei* (SBEAB) on the activities of ALT and AST in STREP-induced diabetic rats.

| Condition   | ALT (nmol/L/min/mg protein) | AST (nmol/L/min/mg protein) |
|-------------|----------------------------|----------------------------|
| Induced     | 0.0152 ± 0.00011b          | 0.04524 ± 0.00275b         |
| Metaglomide | 0.01482 ± 0.00018a         | 0.02332 ± 0.000642a        |
| Pretreated  | 0.01482 ± 0.00011a         | 0.02366 ± 0.000321a        |
| Posttreated | 0.01470 ± 0.000212a        | 0.02488 ± 0.002696a        |

Data are presented as mean ± SD (n=8). Values with different letters are statistically different.

Figure 1: Malondialdehyde (MDA) level from germinal epithelial cells and hepatic cells in diabetic-induced rats treated with stem bark extract *Alstonia boonei*. Data are presented as mean ± SD (n=8). Bars with different letters are statistically different (p<0.05).

Figure 2: Catalase activity from germinal epithelial cells in diabetic-induced rats treated with stem bark extract *Alstonia boonei*. Data are presented as mean ± SD (n=8). Bars with different letters are statistically different.

(a) Rats showed no visible lesions to the hepatocytes.
(b) Diabetic rats showed mild single-cell necrosis (MSCN) especially hepatocytes towards left of the centre. Also, closely-packed hepatic cells with a few foci of random necrosis were seen.
(c) Rats treated with metaglomide (anti-diabetic drug) depicted closely packed hepatic cells with moderate congestion of hepatic sinusoids (CHS).
(d) Pre-treated group of animals with the SBEAB demonstrated closely-packed hepatic cells with marked widespread congestion of hepatic sinusoids (CHS). Also, mild Kupffer cell hyperplasia was observed (e) Post-treated group of animals with SBEAB showed closely-packed hepatic cells with no significant lesions (NSL).

The activity of Δ5-3β-HSD, indicator linked to testosterone production was significantly (p<0.05) inhibited in diabetic rats when compared with the control group (Figure 4). The substantial low activity of Δ5-3β-HSD was remarkably raised by metaglomide, SBEAB pre and post-treatment in diabetic-induced animals (Figure 4).
Similarly, the activity of Δ5-17β-HSD (Figure 5) was significantly (p < 0.05) depleted in diabetic rats relative to control animals. Whereas the depleted activity of Δ5-17β-HSD was significantly (p < 0.05) reversed after metaglomide, pre and post- treatment with SBEAB (Figure 5).

Figure 5: Δ5 17β-hydroxy steroid dehydrogenase (Δ5 17β-HSD) activity from germinal epithelial cells in diabetic-induced rats treated with stem bark extract Alstonia boonei. Data are presented as mean± SD (n=8). Bars with different letters are statistically different (p<0.05).

(a) Rat testis showing normal developmental stages of spermatogenesis i.e. No Visible Lesion (NVL).

(b) Diabetic rat testis showed depleted spermatogenic epithelial cells. Also, there were multiple foci of multinucleated cell formation (MCF) as well as cytoplasmic vacuolations (CV), and detachment of spermatogenic cells (DSC) from the basement membrane with numerous seminiferous tubules and fair regular outlines.

(c) Rats treated with metaglomide (anti-diabetic drug) depicted numerous variably-sized seminiferous tubules (NVS) with regular outlines. Most of the seminiferous tubules contain fewer amounts of spermatogenic epithelial cells. Also, the spermatogenic cells show various degrees of degeneration such as marked cytoplasmic vacuolations.

(d) Pre-treated group of animals with the SBEAB demonstrated numerous seminiferous tubules (NSTs) with regular outlines. Many of these seminiferous tubules contain moderate amounts of spermatogenic cells while a few contain sparse amounts of spermatogenic cells and only few of the spermatogenic cells were degenerated and vacuolated.
Post-treated group of animals with SBEAB showed numerous variably-sized seminiferous tubules with irregular outlines. The seminiferous tubules contain moderately depleted amounts of spermatogenic cells while the spermatocytes were few and elongated spermatids were frequent. Histological assessment of testicular samples in addition supported this finding. Hence, testes from diabetic rats showed depleted spermatogenic epithelial cells with cytoplasmic vacuolations and detachment of spermatogenic cells from the basement membrane (Figure 6B) when compared with the control liver (Figure 6A). For metaglomide, SBEAB pre and post-treated rats, the viability of the spermatogenic cells was reasonably conserved (Figures 6C–6E). Thus, the preservative effect of SBEAB pre and post-treatment on testicular damage via androgenic/steroidogenic pathway was more efficacious in diabetic rats when compared to metaglomide administration. The protein expression of iNOS and COX-2 were evaluated in testes of diabetic rats by immunohistochemistry. As shown in Fig. 7B and 8B respectively, both proteins were highly expressed in diabetic rats when compared to the control. However, metaglomide, SBEAB pre and post-treatment decreased the expression of testicular COX-2 and iNOS proteins in diabetic induced rats. The inhibition of COX-2 and iNOS expression in testes of diabetic rats by SBEAB pre and post-treatment were better than the reference anti diabetic drug-metaglomide (Figure 7 and Table 3).

| Treatment       | iNOS | COX-2 |
|-----------------|------|-------|
| Control         | 0    | 0     |
| Induced         | 2    | 2     |
| Metaglomide     | 1    | 1     |
| Pre-treatment   | 0    | 0     |
| Post-treatment  | 0    | 0     |

iNOS, inducible nitric oxide synthase; COX-2, cyclooxygenase-2.

Key to scores: 0= No testicular cancer formation, 1=mild testicular cancer formation, 2= severe testicular cancer formation. Pre and post-treatment are more efficacious in preventing testicular cancer in diabetic rats than METAG-treatment (pre=post>METG).

Table 3: Scoring of differential protective effect of SBEAB and metaglomide on iNOS and COX-2 expression in testes of diabetic induced rats.

(a) No visible expression of COX-2 (NVEC) i.e. there was intact and normal architectural structures of the membrane and the spermatids were not affected.

(b) Scattered or eroded spermatids with broken membrane architectures in diabetic induced rats were antecedent by the expression of iNOS.

(c) There was little expression of COX-2 (LEC) which resulted into mild distortion in the lumen of seminiferous tubules but with intact membrane architectures.

(d) No visible expression of COX-2 (NVEC). There were intact and normal architectural structures of the membrane and the spermatids were not affected (Figure 8).
antioxidant and anti-diabetic properties of stem bark reported phytochemicals as chemoprevention of diverse diseases and non-6-ene. Antioxidant and models. However, the underlying biochemical and especially molecular elucidated. In the present study, we report that stem bark boonei mechanisms of hepato- and testo-protection by SBEAB remain to be recognized and acted as prophylactics of reacting compounds were recognized and acted as prophylactics of reacting oxygen species (ROS) [20]. This is in support of previous studies which reported phytochemicals as chemoprevention of diverse diseases and ailments [13]. Recent experimental studies have established the antioxidant and anti-diabetic properties of stem bark Alstonia boonei (SBAB) in experimental animal [26,32] and Escherichia coli [22] models. However, the underlying biochemical and especially molecular mechanisms of hepato- and testo-protection by SBEAB remain to be elucidated. In the present study, we report that stem bark Alstonia boonei and anti-diabetic drug (METAG) inhibit the activities of ALT, AST and MDA contents in rat liver of diabetic rats. The observed reduction in the activities of liver enzymes and lipid peroxidation by SBEAB in the present study is in agreement with previous observations on the hepatoprotective potential of several natural compounds on a number of hepatotoxic agents [18,23]. Similarly, as observed, there was a significant rise in hepatic and testicular MDA contents in diabetic induced rat. This result is in agreement with previous investigations where animals suffering from insulin dependent diabetics (type 1) showed increased lipid peroxidation [21,33]. It was also reported that diabetic patients are more susceptible to membrane damage [32,34]. However, mixture of PUFAs from stem bark Alstonia boonei and anti-diabetic drug (METAG) caused a significant reduction in MDA content of diabetic-induced rats. This is in agreement with Harlev et al. [35], where they reported little or no cell membrane lesions in group of diabetic animals treated with natural polyphenols. Hence, we report that people with degenerative diseases especially type 1 diabetes are prone to have hepatic necrosis and reproductive disorders. Also, antioxidant enzyme- catalase (CAT) was significantly increased in the testes of diabetic rats. Both pre and post-treatment with the extract and METAG caused a significant depletion in diabetic-induced rats. Post-treatment is most efficacious in inhibiting CAT activity in type 1 diabetic rat; followed by pre- and METAG-treatment respectively. Reduction on enzymatic antioxidant- CAT was complementary with the results of previous studies [32,36]. Also, protective effects of SBEAB on pre and post- treated group were due to free radical scavenging activity. The effect was connected to the phenolic compounds from stem bark Alstonia boonei. This was in agreement with the prior records which indicated polyphenols as scavengers of free radicals [14].

Histopathologically, livers from diabetic rats showed hepatocyte necrosis towards the centrilobular region. This discovery is line with recent development which reported that necrotic damage of the hepatic cells was responsible for the pathogenesis of the liver or its failure [37-40]. Also, another work reported that considerable damage to the hepatocytes was more pronounced in diabetic patients than physiological individuals [41]. This also suggests that the liver functions had been compromised. The group of animal pre and post treated with SBEAB showed moderately preserved hepatocytes. This may be attributed to the presence of essential PUFAs which act as promoter of liver functionality [41]. Totally, anti-diabetic drug (METAG) is most efficacious in potentiating hepatocytes function in diabetic rat followed by post- and pre-treatment respectively. This observation may be linked to the fact that the drug is most systemic to the survival of the liver cells.

Furthermore, our present study showed marked decrease in the activities of steroidogenic enzymes- Δ5 3β-hydroxy steroid dehydrogenase (Δ5 3β-HSD) and Δ5 17β-hydroxy steroid dehydrogenase (Δ5 17β-HSD) in diabetic rats. These enzymatic activities advocate possible biochemical mechanisms of toxicity in diabetic men coupled with spermatogenic disorders. This observation was similar to previous study which reported that diabetic men have low levels of testosterone, which is a marker of spermatogenesis [42]. Also, further studies had reported diminution on both Δ5 3β-HSD and Δ5 17β-HSD activities among males distressed with reproductive dysfunctions [43,44]. However, pre, post and METAG treatments prevented the reduction of androgenic enzymes activities in diabetic rats. Pre and post treatment with SBEAB was more efficacious than anti-diabetic drug (METAG). The observed elevation could be attributed to synergistic, additive, mutual or joint interactions of the several phenolic compounds. Hence, the increased activity caused by the SBEAB substantiates the link between diabetes and male infertility.
Moreover, as portrayed by the study, diabetic animal had depleted spermatogenic epithelial cells and cytoplasmic vaculations with detachment of spermatogenic cells from the basement membrane. These abnormalities were related to the necropsinia and astenosperma [44,45]. This finding is line with recent study which reported that the necrosis of spermatogenic epithelium cells was connected to the low sperm count, reduced sperm motility and deleterious daily sperm production [46]. This observation further proposes that the abnormal production of primordial germ cells was as a result of the elevated COX-2 and iNOS expression, markers of pro-inflammatory and testicular cancer [13]. The group of animals pre and post treated with SBEAB showed that the possibility of the spermatogenic cells was realistically conserved. Generally, the differential protection (pre and post) of SBEAB is more efficacious in abrogating testicular damage in diabetic rat than anti-diabetic drug (METAG). This is not unconnected to the presence of essential fatty acids in stem bark of *Alstonia boonei*.

COX-2 and iNOS are well known pro-inflammatory genes and their expressions have been shown to be regulated by transcription factors [13,47]. Recently, attention was focused on molecules mediating inflammation and cancer [48]. This is because there is link between inflammation and cancer [46,47]. The inhibition of intracellular inflammatory response is now a sure recipe for developing molecular target-based chemopreventive agents [13]. The present study revealed that pre and post treatment with *Alstonia boonei* abolished the expression of these genes (COX-2 and iNOS) in testicular of diabetic–induced rat. Taken together, pre and post-treatment are more efficacious in preventing pro-inflammatory response and testicular cancer in diabetic rats than METAG-administration. The results therefore suggest that SBEAB may be significant not only in assuaging testes inflammation but also for the prevention of testicular or prostate cancer. It’s also suggests that physiological task of SBEAB was directly connected with its ability to inhibit the activation of transcription factors such as NF-kB, AP-1 and MAPK [47,48] thereby down-regulating the expression of COX-2 in rats’ testes stimulated with toxicant. This study corroborated the recent finding which reported that cyclooxygenase-2 is an enzyme engaged in inflammatory processes and a committed step in prostaglandin biosynthesis from arachidonic acid [49]. Further reports also discovered that inappropriate up-regulation of COX-2 was commonly observed in various premalignant and malignant tissues [50]. In addition, over expression of COX-2 was further noted in transgenic mice with tumor [16] and experimentally-induced carcinogenesis [13,51]. However, the repression of COX-2 and iNOS proteins underscores the molecular basis for the testicular cancer protection by SBEAB in diabetic-induced rats.

**Conclusion**

Essential fatty acids prevented the leakage of AST and ALT into the blood circulation with corresponding decreased in MDA levels of both liver and testes. The activities of Δ5-17β-HSD and Δ5-17β-HSD were significantly elevated in animal group subjected to SBEAB with corresponding decrease in the activity of CAT. Animals treated with SBEAB showed moderately and pragmatic preserved hepatocytes and normal spermatogenic cells. Anti-diabetic drug (METAG) treatment on hepatic damage was most efficacious in diabetic rats; followed by post and pre-treatment respectively while pre and post treatment were more efficacious on testicular damage than anti-diabetic drug (METAG). Furthermore, pre and post-treatment are more efficacious in preventing pro-inflammatory response and testicular cancer in diabetic rats than METAG-administration. We therefore concluded that the repression of genes encoding COX-2 and iNOS proteins validates the molecular basis of the testes protection by SBEAB and further suggests the links between the hepatocellular damage and male reproductive dysfunctions in diabetic individuals.

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