Antioxidant and Hypoglycemic Effects of *Momordica cochinchinensis* Spreng. (Gac) Aril Extract on Reproductive Damages in Streptozotocin (STZ)-Induced Hyperglycemia Mice

Efectos Antioxidantes e Hipoglucémicos del Extracto de Aril de *Momordica cochinchinensis* Spreng. (Gac) sobre los Daños Reproductivos en Ratones con Hiperglucemia Inducida por Streptozotocin (STZ)

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**SUMMARY:** The aim of present study was to investigate the effect of *Momordica cochinchinensis* (Gag) aril (GA) aqueous extract on male reproductive system of streptozotocin (STZ)-induced hyperglycemia (HG) mice. GA were extracted with distilled water (DW) and analyzed for *in vitro* antioxidant capacities. ICR male mice were divided into 7 groups: 1) control, 2) DW, 3) GA 1000 mg/kg BW, 4) HG, 5) HG + glibenclamide, 6 and 7) HG + GA 500 and 1000 mg/kg BW respectively (7 mice/group). In HG groups, mice were induced by STZ at single dose (150 mg/kg BW). They were treated for consecutive 35 days. All groups were compared for blood glucose levels, weights and histopathologies of reproductive organs, sperm concentration including testicular tyrosine phosphorylation protein patterns by Immuno-Western blotting. The results showed that GA processed antioxidant activities and could significantly decrease blood glucose levels and increase sperm concentration in HG mice. Moreover, GA could change the density of a testicular 70 kDa protein in HG-GA groups. In conclusion, GA extract could improve hyperglycemia and male reproductive damages in STZ-induced HG mice.

**KEY WORDS:** *Momordica cochinchinensis* aril; Streptozotocin (STZ); Testicular damage; Testicular tyrosine phosphorylated protein; Hyperglycemia (HG) mice.

**INTRODUCTION**

Diabetes mellitus (DM), also called hyperglycemia, is a group of chronic disorders in glucose metabolism caused by insulin impaired secretion from pancreatic beta cells resulting in elevating glucose levels. DM is involved in many diseases such as cardiovascular disease (de Ferrant *et al.*, 2014), diabetic peripheral neuropathy, diabetic retinopathy (Wang *et al.*, 2014) and diabetic nephropathy (Jin *et al.*, 2014). Moreover, it has been reported that DM affects the male reproductive system of both diabetic patients and animals induced by chemicals (Baccetti *et al.*, 2002; Ballester *et al.*, 2004). In addition, the reductions of sperm motility, viability, concentration, normal morphology and sex hormones have been observed in DM condition. It seemed that testicular tyrosine phosphorylation patterns, in streptozotocin (STZ)-induced hyperglycemia (HG) rats were different from control (Ballester *et al*.). Interestingly, previous studies suggested that an increase in phosphorylation levels of testicular proteins caused by administration of antioxidant plant extracts could be ascribed to testosterone production and spermatogenesis (Iamsaard *et al.*, 2013; 2014). In recent studies, it has been shown that medicinal plants possessing...
antioxidants including *Momordica cochinchinensis* Spreng can significantly reduce blood glucose levels in diabetic patients and STZ-induced animals (Vajpeyi *et al*., 2007; Zhang *et al*., 2014).

*M. cochinchinensis* Spreng (gac) is a tropical plant found in many countries including Thailand. This plant was identified to have various phytochemical components (Ishida *et al*., 2004; Kubola & Siriamornpun, 2011). These fruits have been reported to have antioxidant activity (Kubola & Siriamornpun), anti-inflammatory effect (Jung *et al*., 2013), immunomodulatory activity (Tsoi *et al*., 2006), antitumor properties (Chuethong *et al*., 2007), anti-cancer activity (Zheng *et al*., 2014; Petchsak & Sirapanidkulchai, 2015). However, there was no report of the hypoglycemic effect of Gac aril (GA) on adverse male reproductive system in type- I DM animal model.

**MATERIAL AND METHOD**

**Plant collection and extraction: M. cochinchinensis** arils (GA) was cultured from January to May, 2014 at Kuchinarai district, Kalasin province, Thailand. The plants were authenticated for its actual species by Prof. Dr. Pranom Chantaranothai, Department of Biology, Faculty of Science, Khon Kaen University, Thailand and kept samples in the Herbarium (#Apichakan Sampannang 01 [KKU]), Khon Kaen University, Thailand. The 8000 g of fresh red GA arils were blended. Then GA juices were filtered by the nylon cloth. 200 ml of GA fresh weight was mixed with distilled water at the ratio of 1 to 1 and blended. Then GA juices were filtered by the nylon cloth. The GA filtrate was dried using Spray dryer. The percentage (% yield) of GA extraction was calculated by using below formula:

\[
\text{% yield of GA extraction} = \frac{839.88 \text{ g of GA dried weight} \times 100}{8,000 \text{ g of GA fresh weight}} = 10.50
\]

**Antioxidant assays**

**Determination of total phenolic compound (TPC):** The TPC was measured by Folin-Ciocalteu’s colorimetric method as modified by Singleton *et al*., (1999). The standard calibration was prepared using the gallic acid (Sigma-Aldrich Co. Ltd., USA). Briefly, 1.580 µl of distilled water and 100 µl of Folin–Ciocalteu’s reagents (Darmstadt, Germany) were added and mixed. After 8 minutes, 300 µl of sodium carbonate (Na₂CO₃) (E. Merck KG, Darmstadt, Germany) solution was added, mixed and then incubated for 2 hours. The absorbance of blue color was measured using UV-VIS spectrophotometer (Jasco V530, Japan) at 765 nm.

**Determination of total flavonoid content (TFC):** The TFC was measured using colorimetric method as described by Bakaret *et al*., (2009). The standard calibration was prepared by using the quercetin (Sigma-Aldrich Co. Ltd., USA). Twenty µl of diluted GA extract and quercetin solution were added into test tubes. Briefly, 0.5 ml of GA extract was mixed with distilled water 2.25 ml and added 5 % NaNO₂ solution 0.15 ml. After 6 minutes, 0.3 ml of 10 % AlCl₃·H₂O solution was added before 1 M NaOH 0.1 ml. The absorbance of pale yellow color was measured using UV-VIS spectrophotometer at 420 nm. The concentration of total flavonoid content in GA extract was measured as milligram quercetin equivalent (QE) per gram extract.

**Determination of DPPH free radical scavenging activity:** The scavenging effect was measured using 1,1-diphenyl-2-picryl-hydrazyl (DPPH) assay based on method of Brand-Williams *et al*., (1995). The GA extract solutions in methanol with a series of concentrations were prepared. A standard calibration was prepared using the ascorbic acid (E. Merck KG, Darmstadt, Germany). Then 500 µl of ascorbic acid or GA extract solutions were mixed with 500 µl of 1 mM DPPH (Fluka, Switzerland) solution and incubated for 30 minutes. The absorbance of the resulting solution was measured by using UV-VIS spectrophotometer at 517 nm. The radical scavenging capacity was calculated by using the following equation:

\[
\% \text{ DPPH radical scavenging} = \frac{(\text{Abs}_{control} - \text{Abs}_{sample})}{\text{Abs}_{control}} \times 100
\]

Abs<sub>control</sub> is the absorbance of the control and Abs<sub>sample</sub> is the absorbance in the presence of the sample. The different concentrations of standard and GA extract solutions were used to obtain percent radical scavenging curves for calculating the IC50 values that required to scavenging the 50 % DPPH free radicals.

**Determination of ferric reducing antioxidant power (FRAP):** The total antioxidant activity was measured using ferric tripyridyltriazine (FeIII-TPTZ) complex as reported by Benzie & Strain (1996). A standard calibration was prepared by using various concentrations of ascorbic acid (E. Merck KG, Darmstadt, Germany). Then 1.500 µl of freshly prepared FRAP reagent (E. Merck KG, Darmstadt, Germany) was mixed with 50 µl of ascorbic acid and GA extract solutions and warm at 37 ºC for 10 minutes. The absorbance was measured by using UV-VIS spectrophotometer at 593 nm. The FRAP value of GA extract was calculated from an equation from linear standard curve.
Animals and treatment regime: Forty-nine ICR male mice, aged 6-8 weeks were purchased from National Laboratory Animal Center (NLAC), Mahidol University, Salaya, Nakhon Pathom, Thailand. Mice were acclimatized at Northeast Laboratory Animal Center (NELAC), Khon Kaen University, Thailand. This study was approved by the Animal Ethics Committee of NELAC, Khon Kaen University, Thailand, based on the Ethics of Animal Experimentation of the National Research Council of Thailand (ref. No. 0514.1.12.2/35 with record No. AEKKU-NELAC 29/2557). All mice were grouped and treated for consecutive 35 days (shown in Table I). All animals were measured the glucose levels from tail prick blood using blood glucose oxidase reaction monitoring system (Johnson & Johnson Ltd.). Hyperglycemic (HG) mice were considered when the levels blood glucose was greater than 250 mg/dl.

Epididymal sperm concentration: Sperm concentration was performed as described by Iamsaard et al. (2014). Sperm fluid was collected from epididymis and vas deferens. Its fluid was dipped and re-suspended in 1 ml of PBS. Then sperm suspension was centrifuged (3000 r/min, 2 min) to wash and separate the mature sperm pellet from its fluid. To analyze the sperm concentration, the sperm pellets were re-suspended with 1 ml PBS before dilution. The sperm dilutions (1:20 dilution) were counted for three times of each animal by using a Neubauer’s counting chamber.

Histopathological examinations of the testes and epididymes: Right testis and cauda epididymes were fixed with 10 % formalin and embedded with paraffin and sectioned by microtome about 5 µm thicknesses. The testicular sections were deparaffinized and stained by

Table I. Treatment on each group of mice.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Day 1 (via i.p.)</th>
<th>Days 4-39 (via gastric)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>non</td>
<td>non</td>
</tr>
<tr>
<td>Vehicle I+DW</td>
<td>0.1 M Citrate buffer</td>
<td>DW</td>
</tr>
<tr>
<td>Vehicle II+GA1000</td>
<td>0.1 M Citrate buffer</td>
<td>GA extract (1000 mg/kg BW)</td>
</tr>
<tr>
<td>HG+DW (Negative control)</td>
<td>STZ (150 mg/kg BW)</td>
<td>DW</td>
</tr>
<tr>
<td>HG+Gli (Positive control)</td>
<td>STZ (150 mg/kg BW)</td>
<td>Glibenclamide (Gli)</td>
</tr>
<tr>
<td>HG+GA500</td>
<td>STZ (150 mg/kg BW)</td>
<td>GA extract (500 mg/kg BW)</td>
</tr>
<tr>
<td>HG+GA1000</td>
<td>STZ (150 mg/kg BW)</td>
<td>GA extract (1000 mg/kg BW)</td>
</tr>
</tbody>
</table>

BW= body weight; DW= distilled water; GA= gac aril; HG= hyperglycemia; i.p.= intraperitoneal injection; M= molar; STZ= streptozotocin dissolved in 0.1 M citrate buffer. HG groups (negative control, positive control, HG+GA500 and HG+GA1000); designed based on Muralidhara (2007). Day 1 = STZ injection. Days 4-39 = the treatment days.
hematoxylin and eosin before observing testicular histology under light microscope.

Investigation of testicular phosphotyrosine pattern: The total protein lysate of left side testis was prepared by homogenization with 1X Radioimmunoprecipitation assay buffer added- cocktail protease inhibitors (Cell Signaling Technology Inc., USA). The homogenate was centrifuged and the supernatant was collected to measure the total protein concentration by NANOdrop (NanoDrop ND-1000 Spectrophotometer V3.5 User’s Manual, NanoDrop Technologies Inc., USA). One-hundred micrograms of total testicular proteins were separated by 10% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and blotted onto nitrocellulose membrane to detect the intensity of phosphorylated protein by using the anti-phosphotyrosine primary antibody (Santa Cruz Biotechnology INC., USA) overnight respectively, followed by incubation of anti-mouse secondary antibody (Santa Cruz Biotechnology INC., USA) respectively. Bovine serum albumin (AMRESCO®, USA) and epidermal growth factor (EGF)-like growth factor (Millipore Co., USA) were used as negative and positive controls.

Statistical analysis: All quantitative results were expressed as mean ± standard deviation (SD). One-way analysis of variance (ANOVA) and Duncan test were used SPSS program (version 16) software to examine the significance of differences among sets of data. The level of statistical significance was P<0.05.

Fig. 2. A. Showing effect of GA extract on absolute (1) and relative (2) weights of male reproductive organs. Each data point represented as mean ± SD (n=7). * P<0.05 was statistically significant. B. The blood glucose levels of all groups at day 4 and effect of GA on BGL at day 35 in STZ-induced hyperglycemic (HG) mice. DW: distilled water, Gli: glibenclamide. Each data point represented as mean ± SD (n=7). * P<0.05 was statistically significant. C. Effect of gac aril (GA) extract on sperm concentration in all experimental groups induced hyperglycemia. Each data point represented as mean ± SD (n=7). * P<0.05 was statistically significant. D. The testosterone levels of all experimental groups.
RESULTS

Total phenolic compound and antioxidant activity of GA extract.

Total phenolic compound: The absorbance of GA extract was represented by gallic acid standard linear equation \( y = 0.0004x + 0.0748 \) (and calculated as compared with gallic acid concentration. The GA extract of 47.15 ± 0.21 mg possesses equivalence to 1.00 g of gallic acid (mg/g gallic acid equivalence, GAE).

Total flavonoid content: The absorbance of GA extract was represented by gallic acid standard linear equation \( y = 0.0154x + 0.0484 \) and calculated as compared with quercetin concentration. The GA extract of 11.38 ± 0.02 mg possesses equivalence to 1.00 g of quercetin (mg/g quercetin equivalence, QE).

DPPH free radical scavenging activity: The various concentration of GA extract scavenged the free radical in concentration-dependent DPPH free radical scavenging \( y = 9.6929x + 10.7938 \) with the IC50 of 40.44 µg/ml and maximum response of 67.29 %. Ascorbic acid showed eminent scavenging activity \( y = 4.9724x + 6.6005 \) with the IC50 of 8.07 µg/ml and maximum response of 84.41 %.

Ferric reducing antioxidant power: The absorbance of GA extract was represented by gallic acid standard linear equation \( y = 0.098x + 0.1333 \) and calculated as compared with ascorbic acid concentration. GA extract of 481.12 ± 4.10 mg possesses reducing equivalence to 1.00 g of ascorbic acid mg/g GA extract.

Effect of GA extract on morphology and weights of reproductive organs: Gli/GA 500 or 1000 could obviously improve testicular size as compared to HG group (Fig. 1). Contrasting, all HG groups had small size of reproductive organs as compared to those of control and vehicle (Fig. 2A).

In Figure 2A, the absolute and relative weight of testis tended to be improved after treatment with GA extract but they were not significantly different (P>0.05). However, all
Fig. 4. Photographs showing the seminiferous histopathologies in HG-STZ mice. A) seminiferous atrophy with multinucleated giant cells (arrows) and B) atrophy with germ cells degeneration (asterisk).

HG groups had significantly decreased weight as compared to those of control and vehicle (P<0.05).

Effect of GA extracts on blood glucose levels STZ could induce mice to be actual HG condition with significantly elevated blood glucose levels (BGL) (average 388.21 mg/dl) as compared to control (average 71.29 mg/dl) (P<0.05) (Fig. 2B). After treatment, only GA extract 1000 and glibenclamide could significantly reduce BGL in HG mice as compared to HG-untreated group (P<0.05). In contrast, GA extract could not decrease BGL in GA-500 HG mice compared to HG-untreated group (Fig. 2B).

Effects of GA extract on sperm concentration and testosterone: In Figure 2C, Gli/GA 500 or 1000 can significantly improve sperm concentration compared to that of the HG group (P<0.05). However, sperm concentration in Gli-HG group was not different from GA treated groups (Fig. 2C). Interestingly, GA (1000 mg) tends to improve sperm concentration compared with HG-mice administered low dose of GA. It was noted that the sperm concentration of all HG groups were significantly reduced as compared to that of the non-HG groups (Fig. 2C).

The Gli/GA 500 or 1000 tends to slightly increase testosterone levels as compared to that of the HG group. However, the results showed that the testosterone levels were not significantly among treated groups (Fig. 2D).

Effect of GA extract on histology of testis and epididymis: The Gli/GA 500 or 1000 could improve testicular damages compared to that
of the untreated group (Fig. 3). In addition, GA 1000 alone did not affect testicular tissue. Indeed, STZ decreased spermatogenetic cells and increased atrophic tubule (Fig. 3D). Interestingly, GA extracts could improve testicular damage (Fig. 3F-G) similar to that of glibenclamide treatment (Fig. 3E).

The testicular histopathologies observed in HG mice were classified into two types (atrophy with multinucleated giant cells (Fig. 4A), and with germ cell degeneration (Fig. 4B).

The histologies of epididymis in control and experimental groups were shown in Figure 5. In general, the epithelial cells of all groups were not different. However, the density of sperm mass in GA 500/GA 1000-HG groups (Fig. 5E-F) was greater than that of HG group (Fig. 5D). Such density was similar to that of glibenclamide and control groups. The GA 1000 extract alone did not change the sperm mass (Fig. 5C). In addition, STZ-induced groups were found to have various round cells within epididymal lumen (Fig. 5E).

Relative intensity of the testicular 70 kDa phosphorylated proteins: The intensity of the 70 kDa phosphorylated protein was divided by that of beta-actin (Fig. 6A). Its relative intensity in HG group was significantly increased compared with other groups (P<0.05) while that intensity of GA 1000 and control groups was not significantly different. It was found that relative intensity in GA/Gli-HG treated groups was significantly decreased as compared with the HG group (P<0.05) (Fig. 6B).

DISCUSSION

Kubola & Siriamompun reported that GA extracts were rich in total phenolic content. In comparisons, the antioxidant capacities determined in this study were closed to their results. A medicinal plant possessing antioxidants especially Momordica genus including GA can significantly reduce BGL in DM animals (Vajpeyi et al., 2006; Scarano et al., 2006). The treatments with GA extract in present study could also decrease BGL in HG mice. Although the doses of GA extracts treated in this study were higher than others, the LD50 of this extract documented in acute toxicity study was 2000 mg/kg BW (OECD, 2000). All experimental results indicated that GA extract has antioxidant capacity to reduce BGL in HG animals. In consistent with previous studies, the weights of male reproductive organs in DM animals were significantly decreased (Amaral et al., 2006; Scarano et al., 2006). In this study, the reduction of epididymal weight was related to reduction of sperm concentration and epididymis histology. In contrast to other investigator (Ballester et al.), the serum testosterone in this study was not significantly decreased in HG mice compared to the control. A previous study described that treatment with plants possessing antioxidants improved glycemic imbalance (Amaral et al., 2006; Scarano et al., 2006). In this study, the reduction of epididymal weight was related to reduction of sperm concentration and epididymis histology. In contrast to other investigator (Ballester et al.), the serum testosterone in this study was not significantly decreased in HG mice compared to the control. A previous study described that treatment with plants possessing antioxidants improved glycemic imbalance (Amaral et al., 2006; Scarano et al., 2006). Similarly, GA extract could improve testicular weight and sperm concentration. In present study, GA extract improved testicular and epididymal damages compared to HG mice. It is possible that antioxidants in GA extract have protective capacity against free radical from STZ-HG induction. Interestingly, GA (1000 mg) was not toxic to male reproductive system.

It is known that phosphotyrosine proteins are important in the regulation cell proliferations, and differentiations in both normal and cancer cells (Ulrich & Schlessinger, 1990). In post-translational modifications in male reproductive system, phosphotyrosine proteins are demonstrated to involve in sperm production, capacitation and acrosome reaction (Bailey, 2010; Iamsaard et al., 2014). Previously, Ballester et al. demonstrated that the presence of testicular 66 and 50 kDAs phosphorylated proteins were similar in the
healthy and DM rats with no different intensity. In the previous studies, plant extract processing antioxidant capacities could change the levels of testicular tyrosine protein phosphorylation in experimental animals (Iamsaard et al., 2013; 2014). In this study, only a 70 kDa of testicular phosphorylated protein seemed to interrupt spermatogenesis increased in HG group and improved in GA groups. However, actual roles of the proteins need to be elucidated. In conclusion, GA could improve hyperglycemia and male reproductive damages in STZ-induced hyperglycemia mice.

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REFERENCES


Petchsak, P. & Sripanidkulchai, B. Momordica cochinchinensis aril extract induced apoptosis in human MCF-7 breast cancer cells. Asian

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