Antidiabetic Synergistic Effects of Medicinal Plant Extract Mixtures on db/db Mice

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This study investigates the effects of Psidium guajava L. leaf (Pg) extract, Lagerstroemia speciosa L. leaf (Ls) extract, and mixture A (Pg, Ls, Morus indica L. leaf, Pinus densiflora needles, Acanthopanax senticosus M. roots extract) on db/db mice. For four weeks, db/db mice were fed powdered extracts of Pg, Ls, and mixture A. Compared to the diabetic control, extracts of Pg, Ls and mixture A decreased body weight, glucose and insulin. The greatest decreases were caused by mixture A. These extracts decreased the levels of total cholesterol, triglyceride and free fatty acid compared to the diabetic control. The antihyperlipidemic effect of mixture A was the greatest. Mixture A also significantly decreased injuries of Langerhans’ islets compared to the diabetic control. Mixture A showed a beneficial synergistic effect due to the supplementary pharmacological actions of the ingredients in contains, indicating that it improved hyperglycemia without the side effect of weight gain.

Key words: db/db mice, hyperglycemia, synergistic effect

Introduction

Type 2 diabetes mellitus (DM) is characterized by obesity and insulin resistance [31]. Regulation of DM involves exercise, diet, and pharmacotherapy. However, drugs developed for the treatment of diabetes may fail to prevent long-term complications of diabetes and have side effects that may decrease patient tolerance [14,17]. New therapeutic modalities which provide better management of diabetes, and better tolerance by patients would be an important improvement. Currently, more than 400 plants are reported to have the potential to lower blood glucose levels [3,9]. These medicinal plants are known to be effective in treating diabetes and are expected to achieve a high level of antihyperglycemic effect without adverse reactions, unlike conventional antidiabetic drugs. Antidiabetic activities are caused by various polyphenol, flavonoids, glycosides and terpenoids in the plants, effectively contributing to the improvement of diabetic patients [6,40]. The plant extracts accomplish antidiabetic actions by way of pharmacological mechanisms including insulin-like activity [25,10], as well as increases in insulin sensitivity [15,22,26,28], insulin secretion [19,24,34,38,39], antioxidant activity [7,8,21,29], and restrain the activity of intestinal glycosidase [18-21,28]. The hypoglycemic effect of Lagerstroemia speciosa L. leaves was reported initially in early 1940 [11], and its antidiabetic effect has since been confirmed [16,34]. Effective antihyperglycemic activity has also been reported for Psidium guajava L leaves [1,4,25]. Furthermore, effective antidiabetic activity of Morus indica L. leaves, Pinus densiflora needle, and Acanthopanax senticosus M. roots, which grow wild in Korea, was confirmed [10,18,20,22,30,38]. The objective of this study was to compare the antidiabetic effects of individual L. speciosa and P. guajava leaf extracts on db/db mice, with effects due to the combined antidiabetic mechanisms provided by mixture A (L. speciosa, P. guajava M. indica leaf extract, P. densiflora needles extract, A. senticosus roots extract). It was thought that synergism might make the mixtures superior to extracts from single plant sources.

Materials and Methods

Preparation of the plant extracts

Psidium guajava L. leaves (Pg), Morus indica L. leaves (Mi), Pinus densiflora needles (Pd), and Acanthopanax senticosus M. roots (As) were purchased from the Daegu Yangyeongsi herbmarket (Daegu, Korea). 15 kg of well dried herb, were added to 150 L of distilled water for each of the four (Pg, Mi, Pd, and As). Each herb was boiled for 6 hours at 98 °C to get extracts. Each of the water soluble plant extracts was filtered (50 μm), concentrated under vacuum, then freeze-
dried to form a powder (Pg 0.9%, Mi 1%, Pd 0.8, As 1% in yield). *Lagerstroemia speciosa* L. leaf (Ls) extract, which contained 1% corosolic acid powder extracted using ethanol, was purchased from the Use Techno Corporation Co., Ltd (Kyoto, Japan). Mixture A contained the same amounts of freeze-dried Pg, Ls, Mi, Pd, and As powders.

Animal groups and treatment

Six-week-old male C57BLKs/J-db/db mice (n=80) and C57BL/6CrSlc mice (n=20) were purchased from the Japan Shizuoka Laboratory Center (Shizuoka, Japan) and acclimatized to a controlled temperature of 20±2℃, humidity of 55±5%, and a 12h light and 12h dark cycle for 1 week before starting the experiments. They had free access to standard pellets (Shizuoka Laboratory Center Inc, Japan) and water. The mice were randomly divided into five groups with 20 animals in each group. The NC group served as normal controls (C57BL/6CrSlc mice) and received vehicle (normal saline) only. The DC group served as diabetic controls (C57BLKs/J-db/db mice) and also received vehicle only. The PG and L5 groups received the single plant extracts; Pg extract and Ls extract respectively. The MA group received mixture A. The vehicle or plant extract (100 mg/kg, w/w) was orally administered daily over the experimental period of 4 weeks.

Reagents and equipment

The levels of blood glucose, HDL-cholesterol (HDL-C), total cholesterol (TC), triglyceride (TG), and free fatty acid (FFA) were measured using an automatic analyzer (ADVIA 1650, Bayer, Deerfield, IL, USA). Blood insulin was measured by using the Sensitive Rat RIA kit (Linco Research, USA) and automatic analyzer (Gamma counter COBRA-II, Packard, Ramsey, MN, USA). The Periodic Acid-Schiff (PAS) stain for the liver was measured with an S8002 Kit (BioGenex, USA). The mouse anti-insulin and rabbit antiguacagons antibody (BioGenex, USA) was used as a primary antibody, and a Sensitive™ polymer-HRP detection kit (BioGenex, USA) as a secondary antibody for the immunohistochemical stain of the pancreas. The entire histological examination process was done using light microscopy (Axio Scope, Carl Zeiss, Göttingen, Germany).

Measurement of blood glucose, insulin, lipoprotein and lipids in serum

Five mice from each group were selected each week over 4 weeks, and blood from the heart was collected using a syringe by making an incision in the chest after using ethyl ether as anesthesia. The collected blood was centrifuged for 10 min at 4,000 rpm and 4℃, and the serum was stored at -70℃. After completion of the experiment, blood glucose, insulin, HDL-cholesterol (HDL-C), total cholesterol (TC), triglyceride (TG), and free fatty acid (FFA) levels were measured using an automatic blood chemical analyzer, and the HDL-cholesterol/total cholesterol ratio (HTR) calculated.

Histological examination

After the 4th week of the experiment, the liver and pancreas of each mouse were dissected and fixed in 10% neutral formalin for 1 day. Then, these were processed to produce 4 µm thick paraffin sections (m.p. 58℃). The PAS reaction was employed for liver, and the immunohistochemical avidin-biotin complex (ABC) method was used for insulin or glucagon-detects in pancreas tissue. All the histological examination results reported were the means of measures from 10 different fields on each slide. Histological damage was scored as follows, based on the ABC stain using insulin antibodies: 0 - normal; I - minor injury; II - moderate injury; III - obvious injury; IV - severe injury. Each sample was observed at 400× magnification [37]. While the insulin or glucagon occupied proportion in Langerhans' islet cells was magnified 200×, the picture of each slide was analyzed using the Cell Image Scanner program. The Cell Image Scanner program for determining the insulin or glucagon occupied proportion was developed by Professor Jong Hae Kim from the Division of Electronic Engineering, Sun Moon University. The proposed Cell Image Scanner program on the basis of HIS (Hue, Intensity, and Saturation) color coordinates has been developed by Visual Basic Net 2005 in order to allow detailed analyses of cell images. For the various analyses of cell images, the user can adjust the ranges of hue, intensity, and saturation.

Statistical analysis

All values were expressed as means±S.D. The results were compared using the Kruskal-Wallis test of SPSSWIN, ver. 12.0. The level of statistical significance in the study was either p<0.05 or p<0.01.

Results

Effect of plant extracts on body weight

After 4 weeks, the body weight gain of the diabetic con-
trol increased over 4 times compared to that of the normal control, and showed obesity which is characteristics of type 2 diabetes (Table 1). The body weight gain of the PG, LS, and MA groups decreased 54%, 37%, and 60% compared to that of the diabetic control (p<0.01) and, among these, the body weight gain of the LS group was significantly high compared to other groups (p<0.05).

Effect of plant extracts on blood glucose
The glucose level of the diabetic control increased as the experiment went on, and showed hyperglycemia of over 600 mg/dl in week 4 (Table 2). After 4 weeks, the glucose level of the PG and LS groups were decreased 11% and 6% respectively compared to that of the diabetic control (p<0.05). The glucose level of the MA group was decreased 17% respectively compared to that of the diabetic control (p<0.01). The glucose level of the MA group was lower compared to that of the other groups (p<0.05), the mixture A extract had the highest glucose-lowering effect.

Effect of plant extracts on serum insulin
The insulin level of the diabetic control increased over 6 times compared to that of the normal control since week 1 and showed hyperinsulinemia (Table 3). After 4 weeks, the insulin level of the PG and LS groups were decreased 15% and 10% respectively compared to that of the diabetic

Table 1. Effect of plant extracts on body weight gain in db/db mice (n=5)

<table>
<thead>
<tr>
<th>Group</th>
<th>Initial (g)</th>
<th>Final (g)</th>
<th>Gain (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>23.3±0.7</td>
<td>26.9±0.6</td>
<td>3.6±0.5</td>
</tr>
<tr>
<td>DC</td>
<td>33.4±1.8</td>
<td>49.9±3.4</td>
<td>16.5±2.5</td>
</tr>
<tr>
<td>MA</td>
<td>32.1±0.6</td>
<td>38.6±2.7</td>
<td>6.5±2.2</td>
</tr>
<tr>
<td>PG</td>
<td>34.3±0.7</td>
<td>41.8±2.3</td>
<td>7.5±2.1</td>
</tr>
<tr>
<td>LS</td>
<td>33.5±2.1</td>
<td>43.8±2.8</td>
<td>10.3±1.4</td>
</tr>
</tbody>
</table>

Probability less than 0.01 indicated by "**". Letter superscripts refer to comparison groups.
NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with mixture A, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.

Table 2. Effect of plant extracts on blood glucose level in db/db mice (n=5)

<table>
<thead>
<tr>
<th>Group</th>
<th>1 week (mg/dl)</th>
<th>2 week (mg/dl)</th>
<th>3 week (mg/dl)</th>
<th>4 week (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>151.1±18.7</td>
<td>154.3±13.7</td>
<td>152.5±14.9</td>
<td>155.4±15.5</td>
</tr>
<tr>
<td>DC</td>
<td>559.2±22.7**DC</td>
<td>569.9±24.7**DC</td>
<td>571.6±30.1**DC</td>
<td>620.4±27.7**DC</td>
</tr>
<tr>
<td>MA</td>
<td>551.6±19.0**DC</td>
<td>561.1±20.0**DC</td>
<td>548.2±15.9**DC</td>
<td>513.8±18.9**DC</td>
</tr>
<tr>
<td>PG</td>
<td>555.2±17.7**MA</td>
<td>568.0±21.2**MA</td>
<td>554.8±18.5**MA</td>
<td>551.8±17.1**MA</td>
</tr>
<tr>
<td>LS</td>
<td>564.0±16.3**MA</td>
<td>562.4±20.9**MA</td>
<td>568.2±19.1**MA</td>
<td>580.8±19.2**MA</td>
</tr>
</tbody>
</table>

Probability less than 0.05 indicated by "*" , and less than 0.01 indicated by "**". Letter superscripts refer to comparison groups.
NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with Mixture A, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.

Table 3. Effect of plant extracts on serum insulin level in db/db mice (n=5)

<table>
<thead>
<tr>
<th>Group</th>
<th>1 week (ng/ml)</th>
<th>2 week (ng/ml)</th>
<th>3 week (ng/ml)</th>
<th>4 week (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>0.71±0.47</td>
<td>0.74±0.49</td>
<td>0.73±0.44</td>
<td>0.74±0.44</td>
</tr>
<tr>
<td>DC</td>
<td>4.35±0.85**NC</td>
<td>4.67±0.54**NC</td>
<td>5.26±0.49**NC</td>
<td>5.71±0.38**NC</td>
</tr>
<tr>
<td>MA</td>
<td>4.30±0.55**NC</td>
<td>4.58±0.45**NC</td>
<td>4.61±0.44**NC</td>
<td>4.05±0.37**NC</td>
</tr>
<tr>
<td>PG</td>
<td>4.37±0.46**NC</td>
<td>4.65±0.52**NC</td>
<td>5.02±0.49**NC</td>
<td>4.81±0.32**NC</td>
</tr>
<tr>
<td>LS</td>
<td>4.38±0.60**NC</td>
<td>4.68±0.44**NC</td>
<td>5.13±0.53**NC</td>
<td>5.09±0.42**NC</td>
</tr>
</tbody>
</table>

Probability less than 0.05 indicated by "*", and less than 0.01 indicated by "**". Letter superscripts refer to comparison groups.
NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with Mixture A, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.
Effect of plant extracts on serum TC, HDL-C and HTR

The level of serum total cholesterol (TC), HDL-cholesterol (HDL-C), and HDL-cholesterol/total cholesterol ratio (HTR) in each group is shown in Fig. 1. In Fig. 1A, the TC level of the diabetic control constantly increased until week 4 and showed hypercholesterolemia by over 3 times compared to that of the normal control (p<0.01). After 4 weeks, the TC level of the PG and LS groups decreased 52% and 47% respectively compared to that of the diabetic control (p<0.01). The TC level of the MA group decreased 62% compared to that of the diabetic control (p<0.01). Also, the TC level of the MA group was lower compared to that of the other groups (p<0.01). The HDL-C levels of the normal and diabetic control increased during 4 weeks, but there was no significant difference between the two groups (Fig. 1B). After 4 weeks, the HDL-C level of the PG and MA groups were higher than that of the diabetic control (p<0.01), but in LS group no significant difference was found compared to the diabetic control (p<0.01).

**Fig. 1.** Effect of plant extracts on serum total cholesterol (A), HDL-cholesterol (B) and HTR (C) level in db/db mice (n=5; mean±S.D.). Probability less than 0.01 indicated by "**". Letter superscripts refer to comparison groups. HTR: HDL-cholesterol/total cholesterol ratio, NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with Mixture A, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.
to diabetic control. After 4 weeks, the HTR of the diabetic control decreased to 31% of the normal control (p<0.01), showing no complementary increase of HDL-C corresponding with the increased TC level (Fig. 1C). The HTR of the PG, LS, and MA groups were higher than that of the diabetic control (p<0.01), and in particular, the HTR of the MA showed no significant difference compared to that of the normal control.

Effect of plant extracts on serum TG and FFA

The levels of serum triglyceride (TG) and free fatty acid (FFA) in each group is shown in Fig. 2. In Fig. 2A, the TG level of the diabetic control constantly increased until week 4 and showed an abnormal lipid profile of 6 times higher than that of the normal control (p<0.01). After 4 weeks, the TG level of the PG and LS groups decreased 30% and 8% respectively compared to that of the diabetic control (p<0.01), and the TG level of the PG group was lower than that of LS group (p<0.01). Also, the TG level of the MA group decreased 48% respectively compared to that of the diabetic control (p<0.01), and mixture A extract showing to have the highest effectiveness compared to other extracts (p<0.01). In Fig. 2B, The FFA level of the diabetic control increased until week 4 and showed an abnormal lipid profile of over 2 times higher than that of normal control (p<0.01). After 4 weeks, the FFA level of the PG and LS groups decreased 12% and 7% respectively compared to that of the diabetic control (p<0.01), and the FFA level of the PG group was lower than that of the LS group (p<0.01). The FFA level of the MA group decreased 24% respectively compared to that of the diabetic control (p<0.01). Also, the FFA level of the MA group was the lowest compared to other groups (p<0.01).

Effect of plant extracts on lipid deposits in liver

Changes to lipid accumulation are displayed in Fig. 3. Based on PAS-stained tissue secretions, the oval portions not stained show lipid deposits. After 4 weeks, the normal control showed few lipid deposits in the liver (Fig. 3A) but the diabetic control showed an abnormally large amount of lipid deposits in the liver (Fig. 3B). The MA group showed a reduced lipid deposits compared to that of the

Fig. 2 Effect of plant extracts on serum triglyceride (A) and free fatty acid (B) level in db/db mice (n=5; mean±S.D.). Probability less than 0.05 indicated by *, and less than 0.01 indicated by **. Letter superscripts refer to comparison groups. NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with Mixture A, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.
diabetic control (Fig. 3C). The PG and LS groups also showed a reduced amount of stored lipids compared to that of the diabetic control (Fig. 3D and 3F), but still higher than that of MA group.

Effect of plant extracts on the histological micrograph in Langerhans' islet

Histopathological changes of Langerhans' islets are displayed in Table 4, Fig. 4 and Fig. 5. In Table 4, the pathological grading score of the MA group was significantly higher compared to that of the diabetic control (p<0.05). But the scores of the PG and LS groups were not significantly different compared to that of the diabetic control. The insulin occupied proportion of the Langerhans' islets of the diabetic control, decreased to 2 times less than that of the normal control (p<0.01) and the insulin occupied proportion of the Langerhans' islets of the PG, LS, and MA groups were significantly higher compared to that of the diabetic control (p<0.01). As for the MA group, the insulin occupied proportion of the Langerhans' islets and pathological grading of pancreas tissue did not show any significant difference compared to that of the normal control. In sum, mixture A was most effective in preventing beta cell failure among the plant extracts used for the present work. The glucagon occupied proportion of the Langerhans' islets of the diabetic control, increased to 5.5 times much than that of the normal control (p<0.01). The glucagon occupied proportion of the Langerhans' islets of the PG, LS, and MA groups were lower than that of the diabetic control (p<0.01). In particular, the glucagon occupied proportion of the Langerhans' islets of the MA group was lower than those of the PG and LS groups (p<0.01). In Fig. 4, based

Table 4. Effect of plant extracts on the histopathological damage of Langerhans' islet in db/db mice (n=10)

<table>
<thead>
<tr>
<th>Group</th>
<th>Pathological grading of pancreas</th>
<th>p</th>
<th>Insulin</th>
<th>Glucagon</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>I</td>
<td>II</td>
<td>III</td>
</tr>
<tr>
<td>NC</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>DC</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>MA</td>
<td>4</td>
<td>5</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>PG</td>
<td>1</td>
<td>4</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>LS</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Pathological grading of pancreas was scored using the light microscopy. (ABC stain using insulin antibody, 400×). Occupying proportion was calculated using the Cell Image Scanner program. (ABC stain, 200×)

Probability less than 0.05 indicated by '*', and less than 0.01 indicated by '**'. Letter superscripts refer to comparison groups. NC: Normal control, DC: Diabetic control, MA: Diabetic mice fed with Mixture A, MB: Diabetic mice fed with Mixture B, PG: Diabetic mice fed with Pg, LS: Diabetic mice fed with Ls extract.
Fig. 4. Effect of plant extracts on the histological micrograph of db/db mice Langerhans' islets cell and insulin secretion in 4 week (ABC stain, 200×). A: Normal control, B: Diabetic control, C: Diabetic mice fed with Mixture A, D: Diabetic mice fed with Pg, E: Diabetic mice fed with Ls extract, scale bars represent 100 μm.

Fig. 5. Effect of plant extracts on the histological micrograph of db/db mice Langerhans' islets cell and glucagon secretion in 4 week (ABC stain, 200×). A: Normal control, B: Diabetic control, C: Diabetic mice fed with Mixture A, D: Diabetic mice fed with Pg, E: Diabetic mice fed with Ls extract, scale bars represent 100 μm.

on ABC-stained tissue secretions using insulin antibodies, the portions stained yellowish brown color show insulin secretion. After 4 weeks, the normal control showed a normal shape of Langerhans' islets cell and insulin secretion (Fig. 4A) but the diabetic control have injured Langerhans' islets cell and few insulin secretion (Fig. 4B). The MA, PG, and LS groups showed recovery of Langerhans' islets cell and insulin secretion (Fig. 4C, 4D and 4E). In Fig. 5, based on ABC-stained tissue secretions using glucagon antibodies, the portions stained yellowish brown color show glucagon secretion. After 4 weeks, tissues of the normal control showed a normal shape of Langerhans' islets cell and glucagon secretion (Fig. 5A), but the diabetic control showed an injured Langerhans' islets cells. The majority of beta cells were damaged as glucagon secretion is present over the entire tissue (Fig. 5B). The MA and PG groups showed that the damage to the beta cells of Langerhans' islets was prolonged compared to the diabetic control (Fig. 5C and 5D). Damage to the beta cells of Langerhans' islets of the LS group was prolonged compared to the diabetic control, but
the damage level was higher than that of the PG and MA groups (Fig. 5B).

Discussion

Insulin resistance is one of the major features of type 2 DM. Insulin resistance impairs glucose utilization by insulin-sensitive tissues and increases hepatic glucose output. Both effects contribute to hyperglycemia [31]. Thus, both hyperinsulinemia and hyperglycemia must be improved at the same time to effectively treat type 2 DM. In our previous study, it was confirmed that mixture A showed antidiabetic effect on type 1 DM [32]. The study, using in vitro pancreas beta cells and in vivo streptozotocin-diabetes rats, proved that mixture A ensured a more beneficial antidiabetic effect compared to the effect of either the Psidium guajava L. leaf (PG) or the Lagerstroemia speciosa L. leaf (LS) extract used alone. So, this study verified the antidiabetic effect of mixture A on type 2 DM. According to the result of this study, PG and LS extracts showed antihyperinsulinemic and anti-hyperglycemic activities, and there was no significant effect between the two. However, this result is insufficient to conclude that the antihyperinsulinemic and anti-hyperglycemic effect between these extracts is same as the level of glucose of LS extract increased gradually since Week 3 compared to that of PG extract. The PG extract contains various polyphenolic compounds such as polyphenol [29], terpenoids [4], flavonoids [23], and tannins [36]. These polyphenolic compounds showed various antidiabetic activities and PG extract actually increased insulin secretion [24,34], insulin sensitivity by inhibiting the protein tyrosine phosphatase 1B (PTP1B) [15,28], and blocked the uptake of intestinal glucose by inhibiting the activity of the carbohydrate hydrolyzing enzyme [28]. The LS extract used in this study was an ethanol extract including 1% corosolic acid, and polyphenolic compounds were removed during the extraction process and LS extract contained less polyphenolic compounds than the PG extract. Miura and co-workers [26] claimed that corosolic acid increases glucose transporter 4 (GLUT4) translocation and reduces insulin resistance. But after a study using water-soluble LS extract, Hayashi and co-workers [13] claimed that lagerstroemin, which is a polyphenol, but not the corosolic acid, is responsible for the antidiabetic effect of LS extract. Thus, the antidiabetic effect of corosolic acid is still in question. In sum, the glucose level of LS extract has increased gradually since Week 3 compared to that of PG extract for that the majority of hydrophilic polyphenol compounds contained in Lagerstroemia speciosa L. leaves were not extracted during the extraction process with ethanol, and that the antidiabetic activity of the extracted corosolic acid is lower than what has been widely known so far. Mixture A showed a higher level of antihyperinsulinemic and glucose-lowering activity compared to that of the PG and LS extracts. Mixture A is composed of a mixture of PG, LS, Morus indica L. leaf (Mi), Pinus densiflora needles (Pd), and Acanthopanax senticosus M. roots (As) extracts, and the Mi extract’s main components, fagomine and 1-deoxynojirimycin, stimulated insulin release [38] and blocked intestinal glucose uptake by inhibiting the carbohydrate hydrolyzing enzyme in the small intestine [20]. Liu and workers [22] proved As extract’s insulin-sensitivity increase effects. Also, the Pd extract has inhibition activity on the carbohydrate hydrolyzing enzyme [18], and insulin-like activity [10]. Then, mixture A extract can accomplish antidiabetic effects through antioxidant pharmacological mechanisms such as insulin-secretion increase activity by PG, LS, Mi, Pd, and As extracts; intestinal glucose-uptake blocking activity by PG, Mi, and Pd extracts; increase activity of glucose transportation to cells by the PG, LS, Pd, and As extracts.

Type 2 DM is characterized by obesity and abnormal fat metabolism. The increased adipocyte mass leads to increased levels of circulating free fatty acids and other fat cell products. The increased production of free fatty acids and some adipokines may cause insulin resistance in skeletal muscle and liver. This is also responsible for the dyslipidemia found in type 2 DM [31]. The activities of nutritional antioxidants can reduce cholesterol absorption, cholesterologenesis, and fatty acid synthesis [12,33]. In this study, PG and LS extracts improved dyslipidemia, and PG extract showed a higher effect due to its anioxidant activity. Apart from the activity increasing insulin sensitivity, PG extract also acts as an antioxidant activity [7]. Thus, the PG extract was more effective in improving dyslipidemia compared to the LS extract, which only increases insulin sensitivity by the remaining polyphenolic compound. Mixture A improved hyperlipidemia, and showed more effectiveness than PG and LS extracts. This is because mixture A extract’s antioxidant activity, antihyperinsulinemic and glucose lowering effect was higher than that of other extracts. The number of beta cell decrease in individuals with long-standing type 2 diabetes [31]. In the present study, beta cell failure was caused
by type 2 DM due to the decreased insulin occupied proportion of Langerhans’ islets of the diabetic control and increased glucagon occupied proportion compared to that of the normal control. The Pg extract’s insulin occupied proportion was significantly higher than that of the Ls extract, indicating the damage on the beta cells of Langerhans’ islets was delayed. The pathological grading of pancreas tissue, as well as the insulin and glucagon occupied proportion of mixture A was similar to that of the normal control. This suggests that mixture A contributed to the improvement of antihyperinsulinemia and antihyperglycemia, and to prevention of the functional damage of beta cells usually caused by type 2 DM. For mixture A, this appears to be the result of the extracts’ effectiveness in increasing insulin sensitivity, lowering glucose level, and its insulin-like and antioxidant activity. The obesity accompanying type 2 DM is thought to be part of the pathogenic process. As a result of insulin resistance in adipose tissue and resulting obesity, free fatty acid flux from adipocytes is increased, leading to increased lipid synthesis in hepatocytes. This lipid storage or seastosis in the liver may lead to nonalcoholic fatty liver disease and abnormal liver function [31]. Compared to the diabetic control, the Pg, Ls, and MA extracts contributed to improvement by decreasing glycogen storage and lipid accumulation in the liver as well as reducing obesity. Among treatments, the improvements due to mixture A were the greatest. It is well known that the majority of the current antidiabetic drugs have side effects such as gaining weight [27,41]. These drugs enhance hyperglycemia in a short time of period, and this can accelerate obesity, one of the main causes for type 2 DM. It is highly important to treat type 2 DM by ensuring the antihyperglycemic effect without side effects including weight gains. In this study, mixture A was effective not only in treating hyperglycemia, hypoinsulinemia and hyperlipidemia, but also in recovering from abnormal lipid accumulation in the liver. Furthermore, mixture A proved to be effective in restraining gaining weight gains while ensuring the antihyperglycemic effect.

In conclusion, it was proved that mixture A, which is the combination of functions such as intestinal glucose uptake blocking activity, increase activity of glucose transportation to cells, and antioxidant activity of each plant extracts, ensures a higher antidiabetic synergy effect compared to that of the other extracts. However, the components of each plant extracts proved of antidiabetic synergy effects through this study should be chemically analyzed and their chemical structure should be understood in order to provide a basis for future development of diabetes therapeutic agents.

References

초록: *db/db mice*에 대한 약용 식물추출 혼합물의 항당뇨 상승효과

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본 연구는 *db/db mice*에 대한 구아바 잎 (Pg), 바나바 잎 (Ls) 추출물 그리고 혼합물 A (바나바, 구아바, 뱃, 속 잎 그리고 가시오가피 뿌리 추출물)의 효과를 조사하였다. 4주간 *db/db mice*에 이들 추출물을 섭취시킨 결과 Pg, Ls 그리고 혼합물 A의 섭취는 당뇨 대조군에 비해 체중, 혈당, 인슐린을 감소시켰으며 이들 중에서 혼합물 A의 감소효과가 가장 능았다. 이들 추출물은 당뇨 대조군에 비해 총 콜레스테롤, 중성지방, 유리지방산을 감소시켰으며 이들 중에서 혼합물 A의 항고지혈증 효과가 가장 뛰어났다. 또한 혼합물 A는 당뇨 대조군에 비해 *Langerhans' islets*의 손상을 유의하게 감소시켰다. 따라서 혼합물 A는 포함된 구성물질의 상호보완적인 약리작용에 의해 체중증가의 부작용이 없이 고혈당을 개선시키는 유익한 상승효과를 발휘한 것으로 나타났다.