Interaction of Aqueous Extract of Trigonella Foenum-Graecum Seeds with Glibenclamide in Streptozotocin Induced Diabetic Rats

Lal, V.K., P.P. Gupta, P. Tripathi and A. Pandey

Abstract: Problem statement: Interaction between herbs and drugs may increase or decrease the pharmacological or toxicological effects of either component. Herb traditionally used to decrease glucose concentrations in diabetes could theoretically precipitate hypoglycaemia if taken in combination with conventional drugs. Approach: The present study was undertaken to determine the interaction of Glibenclamide, a Sulphonylurea with Trigonella foenum-graecum seed extract, an herbal drug widely used as an antidiabetic agent. The pharmacodynamic interaction was evaluated in streptozotocin induced diabetic rats. Glibenclamide was given orally in two different doses of 0.25 and 0.50 mg kg$^{-1}$. Trigonella seed extract was administered at a dose of 20 mL kg$^{-1}$. The blood glucose estimation was carried out. Results: Both glibenclamide and Trigonella seed extract showed hypoglycemic effect. The hypoglycemic effect observed with combination of glibenclamide and Trigonella seed extract was significantly more than either of drug given alone. Conclusion: It is concluded that Trigonella seed extract shows synergistic effect with glibenclamide. This could be important in reducing the dose of glibenclamide to achieve enhanced therapeutic effect with minimum adverse effect.

Key words: Glibenclamide, trigonella foenum-graecum, hypoglycemic, synergistic effect

INTRODUCTION

Diabetes Mellitus (DM) is a major chronic life-threatening disorder, in which homeostasis of carbohydrate and lipid metabolism is improperly regulated by the pancreatic hormone, insulin; resulting in an increased blood glucose level. DM is a serious metabolic disease that has a significant impact on the health, quality of life and life expectancy of patients, as well as on the health care system Sancheti et al. (2009). The condition affects the metabolism of carbohydrates, fat, water and electrolytes leading to structural changes in a range of cells especially those of the vascular system, subsequently leading to long term complications of diabetes (King et al., 1998; Amos et al., 1997; Zimmet, 2000; Kumar and Clark, 2002; Wild et al., 2004). On the basis of laboratory findings, WHO has defined DM as a fasting venous plasma glucose concentration greater than 7.8 m mol/l (140 mg dL$^{-1}$) or greater than 11.1 m mol/l (200 mg dL$^{-1}$), two hours after carbohydrate meal or two hour after an oral ingestion of the equivalent of 75 g glucose, even if the fasting condition is normal (Grossman and Messerli, 1996). It is also one of the most prevalent (the estimated lifetime risk of developing diabetes for individuals born in 2000 is 32.8% for males and 38.5% for females) and costly chronic diseases, which significantly reduces life expectancy. In the year 2000, the total number of people with DM was 151 million and the number is projected to increase by 46% to reach 221 million by the year 2010 and 300 million in 2025 Al-Shamsi et al. (2007). Most of these cases will be type 2 diabetes, which is strongly associated with a sedentary lifestyle and high calorie-nutrition and obesity (Zimmet et al., 2001; 2003). This presents a moving therapeutic target that requires a range of different agents to address the different features of the disease at different stages of its natural history. Although biomedical science has unrevealed substantially the pathologically processes involved in causing/fostering diabetes and has designed therapeutic agents with a range of action to fight hyperglycemia, the efficacy of these therapeutic agents is compromised in several ways. Individual agents act only on part of the pathogenic process and only to a
partial extent. This may be the reason that even after so much advancement in understanding the disease process and availability of a wide range of therapeutic agents, the disease is still progressing. Although several drugs targeted for carbohydrate hydrolysing enzymes (pseudosaccharides), release of insulin from pancreatic b-cells (sulphonyl urea), glucose utilization (biguanides), insulin sensitizers, PPARγ agonists (glitazones) are in clinical practice, the growing diabetes market observes a number of changes. The glitazones are meant to target the problem of insulin resistance and enhance insulin action at the cellular level; however, some of these drugs are linked to liver toxicity (troglitazone), including a number of deaths from hepatic failure (Krische, 2000; Gale, 2001; Stern, 1999) and raising the symptoms and risk factors of heart disease leading to heart failure (rosiglitazone).

Use of Complementary and Alternative Medicine (CAM) in the United States has been increasing in recent years (Eisenberg et al., 1998). Indeed, the dietary supplement industry is currently estimated to be a $20 billion industry and according to recent statistics from the Food and Drug Administration (FDA), there are at least 29,000 dietary supplement products on the market. Dietary supplements are defined by the Dietary Supplement and Health Education Act (DSHEA) of 1994 Anon, 1995 and include such products as herbal, vitamins, minerals, sports nutrition supplements, weight management products, specialty supplements and other oral dosage forms intended to supplement the diet.

Hence antidiabetic herbs are also utilized frequently and effectively. A multitude of plants have been used for the treatment of DM throughout the world. In fact in many parts of world especially in poor countries, this may be the only form of therapy available for treating diabetic patients. Complementary and alternative medicine involves the use of herbs and other dietary supplements as alternatives to mainstream Western medical treatment. A recent studies has estimated that up to 30% of patients with DM use complementary and alternative medicine (Ryan et al., 2001).

The interaction of herbs with drug is well known. Herbal drug interaction can be characterized as either Pharmacodynamic (PD) or Pharmacokinetic (PK) in nature. Pharmacodynamic interaction may occur when constituents of herbal product have either synergistic or antagonistic activity in relation to a conventional drug. Pharmacokinetic interaction result from alteration of absorption, distribution, metabolism or elimination of a conventional drug by an herbal product or other. Seeds of *Trigonella Foenum-graecum* L. (Leguminosae) are known to exhibit hypoglycemic activity when taken orally. The hypoglycemic effect of *Trigonella* seeds and their major alkaloids, trigonilnine was first described by Fournier and by Nadakarnis. The seeds are widely recommended for non insulin dependent diabetes mellitus patients. Ajabnoor and Tilmisany reported the hypoglycemic effect of *Trigonella foenum graecum* seeds on the serum glucose level (Abdel-Barry et al., 1997).

Now a day many people use the antidiabetic herb and antidiabetic drug along with and hence there may be chance of interaction between them. Thus the present study was undertaken to evaluate any possible pharmacodynamic interaction *Trigonella* seed extract with oral hypoglycemic agent named glibenclamide.

**MATERIALS AND METHODS**

**Plant material:** *Trigonella foenum-graecum* (Fenugreek) seeds were purchased from the local market and identified by the NBRI, Lucknow, India. A voucher specimen no is 97378.

**Extraction of aqueous plant material:** About 100 g of powdered *Trigonella foenum-graecum* seeds were added in 750 mL of boiling water and macerated well for 10 min. The 400 mL final filtrate was then cooled, filtered, stored as aqueous extract at 4°C and used for oral administration.

**Experimental animal:** Healthy adult rats of wistar strain weighing 110-160 mg were used in the present study. The animals were housed in clean polypropylene cages and maintained in a well ventilated temperature controlled animal house with constant 12h light:dark schedule. The animals were fed with standard rat pellet diet and clean drinking water was made available ad libitum.

**Experimental design:**

**Induction of diabetes:** Rats were fasted overnight before inducing diabetes with streptozotocin. The rats were given an intraperitoneal injection of streptozotocin (50 mg kg\(^{-1}\)) freshly prepared in 0.1M sodium citrate buffer. The diabetic state was confirmed 48 h after streptozotocin injection. Threshold value of fasting blood glucose was taken as > 200 mg DL\(^{-1}\).

Control and diabetic rats were weighed matched for body weight and divided into following group consisting five animal each:

- Group I-Non diabetic control: Treated with single ip injection vehicle
- Group II-Diabetic control: Treated with single ip injection vehicle
- Group III-Chlorpropamide: Treated with chlorpropamide (15 mg kg\(^{-1}\)) ip injection vehicle
- Group IV-Trigonella foenum-graecum extract: Treated with extract (100 mg kg\(^{-1}\)) ip injection vehicle
- Group V-Chlorpropamide+Trigonella foenum-graecum extract: Treated with chlorpropamide (15 mg kg\(^{-1}\)) + extract (100 mg kg\(^{-1}\)) ip injection vehicle

- Group II-Diabetic Control: Administered orally with equal volume of vehicle alone
- Group III-Diabetic rats administered with extract of Trigonella foenum graecum seed at a dose of 2 mL 100 g⁻¹ body weight day
- Group IV-Diabetic rats administered with Glibenclamide at low dose of 0.25 mg kg⁻¹
- Group V-Diabetic rats administered with Glibenclamide at high dose of 0.5 mg kg⁻¹
- Group VI-Diabetic rats administered with Fenugreek extract and low dose of Glibenclamide
- Group VII-Diabetic rats administered with Fenugreek extract and high dose of Glibenclamide

**Blood glucose estimation:** Blood sample was obtained through puncture tail vein and glucose was estimated on 0, 7, 14, 21 and 28th day by Accu-Chek Glucometer.

**Statistical analysis:** Result were expressed as mean ± SEM. Statistical analysis was carried out by using one way analysis of variance followed by Dunnet test. A value of p<0.05, p<0.01 and p<0.001 were considered significant.

**RESULTS**

The mean blood glucose level of animals at various time intervals after the oral administration of drug, extract and drug extract combinations are shown in Table 1. The glucose levels were compared to values obtained for animals given only normal saline (Diabetic control).

Aqueous extract of Fenugreek seeds showed significant decrease (p<0.05) in blood sugar level in comparison to diabetic control group. Glibenclamide alone and the combination of extract and glibenclamide caused a steady and significant reduction in the glycemia throughout the duration of the monitoring period as shown in Fig. 1. Table 1 showed that aqueous extract of Fenugreek seeds significantly increase the hypoglycemic effect of half dose of glibenclamide and its nearby similar to the hypoglycemic effect obtained by glibenclamide high dose. The maximum hypoglycemic effect was seen by the high dose of glibenclamide with aqueous extract of Fenugreek seeds.

![Fig. 1: Effect of Trigonella seed extract, Glibenclamide and Glibenclamide + Trigonella seed extract on blood glucose](image)

**Table 1: Effect of Trigonella seed extract, Glibenclamide and Glibenclamide + Trigonella seed extract on blood glucose**

<table>
<thead>
<tr>
<th>Blood glucose</th>
<th>0 day</th>
<th>7 day</th>
<th>14 day</th>
<th>21 day</th>
<th>28 day</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non Diabetic</td>
<td>90±3.9</td>
<td>93±4.6</td>
<td>91±2.6</td>
<td>94±3.5</td>
<td>89±2.7</td>
</tr>
<tr>
<td>Diabetic</td>
<td>235±2.7</td>
<td>281±3.5</td>
<td>297±4.6</td>
<td>333±4.8</td>
<td>327±5.1</td>
</tr>
<tr>
<td>Diabetic + Trigonella seed extract</td>
<td>227±4.6</td>
<td>215±3.2*</td>
<td>168±2.6*</td>
<td>129±5.2*</td>
<td>120±3.7*</td>
</tr>
<tr>
<td>Diabetic + Glibenclamide low dose</td>
<td>229±3.2</td>
<td>203±3.3**</td>
<td>162±3.6*</td>
<td>121±3.8*</td>
<td>113±4.1*</td>
</tr>
<tr>
<td>Diabetic + Glibenclamide high dose</td>
<td>244±3.2</td>
<td>203±6.2***</td>
<td>159±4.2**</td>
<td>119±4.8**</td>
<td>99±2.3***</td>
</tr>
<tr>
<td>Diabetic + Trigonella seed extract + Glibenclamide low dose</td>
<td>233±4.6</td>
<td>185±5.8***</td>
<td>131±2.8***</td>
<td>110±2.6***</td>
<td>105±3.7***</td>
</tr>
<tr>
<td>Diabetic + Trigonella seed extract + Glibenclamide high dose</td>
<td>233±3.7</td>
<td>179±3.8***</td>
<td>123±3.7***</td>
<td>98±3.0***</td>
<td>71±2.9***</td>
</tr>
</tbody>
</table>

All values are expressed as mean ± S.E.M (n=5). *p<0.05, **p<0.01, ***p<0.001 as compared to diabetic control. One-way ANOVA followed by Dunnet test.
DISCUSSION

Many medicinal plants and pharmaceutical drugs are therapeutic at one dose and toxic at another. Interaction between herbs and drugs may increase or decrease the pharmacological or toxicological effects of either component. Synergistic therapeutic effects may complicate the dosing of long term medications e.g., herb traditionally used to decrease glucose concentrations in diabetes could theoretically precipitate hypoglycaemia if taken in combination with conventional drugs.

The present study was undertaken to evaluate the effect of aqueous extract of Fenugreek seeds on hypoglycemic activity on hypoglycemic action of glibenclamide. The result observed suggest that aqueous extract of Fenugreek seeds when combined with glibenclamide enhances the hypoglycemic activity of latter.

The combination of high dose of glibenclamide (0.5 mg kg$^{-1}$) with aqueous extract of Fenugreek seeds shows maximum hypoglycemic activity and the effect produced by the combination of glibenclamide (0.25 mg kg$^{-1}$) with aqueous extract of Fenugreek seeds was similar to hypoglycemic effect shown by glibenclamide alone (0.5 mg kg$^{-1}$).

It is likely that the beneficial effect of TE is due to some of the bioactive compounds present in it, including 4-hydroxyisoleucine, a novel amino acid known to facilitate insulin secretion. In addition, the soluble dietary fibers present in TE could inhibit absorption of glucose in the gastrointestinal tract (Xue et al., 2007).

Glibenclamide is used to treat DM. Sulphonylurea enhance cell insulin release by blocking the ATP-dependent $K^+$ channel and are widely used in the treatment of type 2 diabetes-mellitus. In chronic therapy the mechanism of action of Sulphonylurea is less clear. Studies have shown that the long-term use of these oral hypoglycaemic agents does not increase basal insulin release or enhance insulin secretion in response to metabolic stimuli in patients with type 2 diabetes, the drug has been described as a classical inhibitor of the $K^+_{ATP}$ channels in pancreatic $\beta$ cells whose target is the SUR receptor, a protein belonging to the ABC transporter family (Anon, 1995).

Administration of an antidiabetic herb with a hypoglycemic drug for the treatment of diabetes may pose for potential drug-herb interaction that may have beneficial or adverse effects. It is generally believed that the use of herbs with medicine produces enhanced effect and reduces the adverse effect of drugs. The results of the present study indicate that combining aqueous extract of Fenugreek seeds with glibenclamide could provide an opportunity to reduce the dose of glibenclamide, which may help in minimizing the adverse effect of glibenclamide as well as achieve enhanced therapeutic effect. At the same time proper precaution and care should be taken to avoid severe hypoglycemia that may occur due to combination of these agents.

CONCLUSION

Furthermore this observed synergistic effect has a great clinical implication. A diabetic patients on be placed on a reduced dose of glibenclamide (Which also implied lower adverse effect) while being encouraged to consume Trigonella seed extract. The issue of interaction should be seriously considered while a patient is combining a potent antidiabetic agent and herbal remedies. So caution should be taken if any patient is taking antidiabetic drug and herb as this might cause severe hypoglycemia which could lead to coma and ultimately death.

REFERENCES


