Polyherbal Formulation (PDBT Capsules) Containing Extracts of Ayurvedic Plants for Treatment of Prediabetes

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Abstract: In Prediabetes blood sugar levels are found to be higher than normal levels. It is considered as pre stage of diabetes as people with prediabetes, if remain undiagnosed, have chances of developing type 2 diabetes. Studies show that if diabetes is treated in the prediabetic stage, it results in complete cure of diabetes. The objective of the present study was to investigate effect of PDBT capsules containing extracts of well-known Ayurvedic plants such as Zingiber officinale, Tinospora cordifolia, Pterocarpus marsupium, Gymnema sylvester, and Momordica charantia on prediabetic condition induced in Wistar rats. High fat fructose diet was fed to male Wistar rats to induce prediabetes. After induction on 11th week of study, the prediabetic rats were divided in different groups. Treatment groups received Pioglitazone (3 mg/kg) (standard control), test formulation, PDBT capsule (200mg/kg) (test group) along with high fat fructose diet. Normal diet control group received normal diet and high fat fructose diet control group received only high fat fructose diet. The different biochemical parameters, blood glucose levels and body weights were monitored. Animals were sacrificed at the end of study period. Histopathological studies on the major organs such as liver, kidney, pancreas was performed. High fat fructose diet induced of prediabetes in rats. Significant decrease in levels of fasting blood glucose and biochemical parameters in treatment groups was observed. The histopathological study showed no abnormalities. The present research show that PDBT capsules can be efficiently used in prediabetes treatment. Hence, PDBT capsule can be a potential agent for the treatment of prediabetes.

Keywords: Prediabetes, Zingiber officinale, Tinospora cordifolia, Pterocarpus marsupium, Gymnema sylvester, Momordica charantia

1. Introduction

Diabetes mellitus is a cluster of different disorders. It is diagnosed by hyperglycemia and glucose intolerance. In diabetes there may be lack of insulin or action of insulin might be defective or both [1].

Diabetes mellitus has affected more than 170 million individuals worldwide [2]. In India the diabetes is one of the major health problem. In the treatment of the diabetes, new area for the research has been developed, that is to treat diabetes in its primary stage [3, 4]. The condition is commonly known as prediabetes. As per Tabák A et al Prediabetes, typically defined as blood glucose concentrations higher than normal, but lower than diabetes thresholds, is a high-risk state for diabetes development [3, 5].

In prediabetes the fasting glucose levels are between 100 and 125 mg/dl and while in oral glucose tolerance test (OGTT) glucose levels are between 140 and 199 mg/dl [6, 7].

Prediabetes is becoming more common in the South Asian population (both in India and abroad). As per the International Diabetes Federation reports, 280 million individuals were pre-diabetic in year 2011. It has been proposed that 398 million individuals having prediabetes are
likely to develop type 2 diabetes if remain untreated. [8]. International Diabetes Federation indicates that approximately 471 million may have prediabetes by 2035.

According to WHO, if people have one of two distinct states: impaired fasting glucose (IFG), or impaired glucose tolerance (IGT), then they are at high risk of developing diabetes [9].

It is proved that type 2 diabetes can be cured when treated in its prediabetic stage [3]. Lifestyle modification can reduce 40–70% relative-risk and help in diabetes prevention. Pharmacological interventions are also found useful in clinical studies [10].

Oral hypoglycemic agents help to maintain the blood glucose level but they show several unwanted consequences like weight gain, hypoglycemia, insulin resistance, complications of cardiovascular system and gastrointestinal disorders [11]. Pathogenesis of diabetes mellitus involves multiple factors. Hence the drugs which can target at multiple receptors, and which are safe, cheap and easily available are the need of the time. The traditional Ayurvedic drug treatments offer good therapeutic results [12]. In Ayurvedic literature several medicinal herbs are suggested for the treatment of diabetes mellitus [13]. Traditional medicinal plants and traditional formulation have potent phytochemicals are helpful in developing new types of therapy. Potential phytochemicals from Medicinal plants show multiple beneficial effects in treatment of diabetes as well as diabetes-related complications [14, 15].

The scientific validation of pharmacological effects and phytochemicals in Indian medicinal plants with modern technique is very limited. A large number of plants used with enormous potential are yet to be validated scientifically and systematically for their pharmacological activity. The systematic evaluation and information on the evidence-base of these plants will enhance the global acceptance about Ayurveda and Indian herbs [14, 16].

In the present study therefore attempts were made for formulation and evaluation of poly herbal capsules (PDBT capsules) against the prediabetic conditions induced by high fat fructose diet in Wistar strain rats.

Poly-herbal formulations have diversity of active principles and hence they have potent effect. These actives may give synergistic effect. PDBT capsules contain aqueous extracts of Zingiber officinale, Tinospora cordifolia, Pterocarpus marsupium, Gymnema sylvestre and Momordica charantia in equal quantity. Previous studies of these plants were carried out by different authors for proving the actions of individual drug for its safety and antidiabetic efficacy studies [15, 17-20]. Here in this study we have evaluated the combination of these medicinal plants against prediabetic condition in rats.

2. Materials and Methods

2.1. Chemicals

PDBT capsules were procured from Podar Ayurved College Mumabi. Pioglitazone (5mg) by Aventis Pharma were purchased from local market.

Ready to use biochemical kits for different parameters were purchased ARK diagnostics. The different blood parameters such as total cholesterol (TC), triglyceride (TG), high-density lipoprotein (HDL), alanine aminotransferase (ALT), alkaline phosphatase (ALP) for liver tests serum creatinine and blood urea nitrogen were evaluated using these kits. For estimation of all biochemical parameters automated bioanalyzer (ChemWell) was used.

2.2. Experimental Animals

Current study was conducted as per prior approval from the Institutional Animal Ethics Committee of C. U. Shah College of pharmacy SNDT University (IAEC approval number is CUSCP/IAEC/31/2012-2013).

Wistar rats of both sexes, weighing 180-210 g were procured from Bharat Serum and Vaccine Pvt. Ltd., Mumbai, India. Animals were provided balanced laboratory diet and given clean water ad libitum. They were kept at 20±2°C, 65–70% humidity, and day/night cycle (12 h/12 h).

2.3. Experimental Design

2.3.1. Induction of Prediabetes

Prediabetes was induced in animals by orally feeding High fat fructose Diet to animals for 15 weeks. The high-fat diet comprises approximately 21% protein, 66.5% fat, 12.5% carbohydrates. The fasting blood glucose level and oral glucose tolerance test (OGTT) were estimated. [21] The rats showing fasting glucose level between 100 and 125 mg/dl and glucose level between 140 and 199 mg/dl in OGTT test were considered as prediabetic [6, 7, 22] and selected for the study. The animals were then randomized into four groups (n=6/group). The drug treatment was given for 4 weeks. During the treatment the animals were divided in following groups.

Group 1: Normal diet control group, this group received normal diet throughout the study and received vehicle 0.1% Sodium Carboxy methyl cellulose (Na CMC) during the intervention.

Group 2: High fat fructose diet control group (HFD group) received High fat fructose diet only and no treatment.

Group 3: Standard control group received treatment of Pioglitazone (3 mg/kg) with high fat fructose diet during intervention.

Group 4: PDBT treatment group received treatment of PDBT capsules (200 mg/kg) during intervention and high fat fructose diet throughout the study.

Fasting blood glucose levels were estimated weekly using one touch electronic glucometer and glucose test strips. To evaluate the other biochemical parameters, blood samples were withdrawn from the retro orbital plexus of the animals after overnight fasting. Serum was separated and used for estimating various biochemical parameters. All biochemical parameters were estimated at different intervals, before induction to find basal values while every month during induction as well as twice during drug intervention. The
animals were sacrificed at the end of 4 weeks of intervention and vital organs viz. kidney, liver, pancreas were isolated for histopathological study.

2.3.2. Statistical Analysis
All the values of in this study were expressed as mean±standard error of mean (S. E. M). The results were analysed for statistics using InStat software. All results were analysed by one way ANOVA and Dunnet’s t-test.

3. Results

3.1 Effect of PDBT Capsules on Different Biochemical Parameters

3.1.1. Effect of PDBT Capsules on Fasting Blood Glucose Levels
Levels of fasting blood glucose in all groups are shown in Figure 1. There was an age-related progressive increase in fasting blood glucose levels in the vehicle control group. Feeding of high fat fructose diet in the HFD group caused increase in fasting blood levels. Compared to vehicle control, there was a significant increase in the fasting blood glucose values at the 24th and 30th day. On 30th day 25% increase in fasting glucose levels were found as compared to normal control group. In the treatment groups, there was a significant decrease in fasting glucose levels on 16th day, 24th day, 30th day of drug treatment. Treatment of PDBT formulation and pioglitazone reduced fasting glucose levels of prediabetic rats upto 84±0.72 mg/dL and 87±0.8 mg/dL respectively on 30th day of intervention. However, there is no statistically significant difference in glucose levels between the pioglitazone and PDBT capsule treated groups.

3.1.2. Effect of PDBT Capsules on Body Weight of Rats
Changes in body weight in all groups are shown in Figure 2. All groups showed age-related increase in body weight. The prediabetic control rats showed statistically significant increase in body weight with prediabetes symptoms like polyuria, polydipsia, polyphagia indicating that prediabetic condition was induced.

The body weights of animals treated with formulations was significantly decreased as compared with HFD control group. At the 30th day of intervention pioglitazone and PDBT formulation treatment reduced weight to 191.91±6.26 gram and 201±6.37 gram respectively. The results indicate that the PDBT capsules were effective in controlling the obesity associated with prediabetic condition.

Figure 1. Effect of PDBT capsules on fasting blood glucose levels of rats.
Values are expressed as mean±SEM., n=6; n=Number of animals.
* Significant decrease when compared with high fat fructose diet control group, p<0.01
$ Significant increase when compared with control group, p<0.05
Control: vehicle control, HFD: High fat fructose diet control

Figure 2. Effect on body weight of rats treated with PDBT capsules
Values are expressed as mean±SEM., n=6; n=Number of animals.
* Significant decrease when compared with high fat fructose diet group, p<0.05
$ Significant increase when compared with control group, p<0.05
Control: vehicle control, HFD: High fat fructose diet control

Figure 3. Effect on Cholesterol levels in prediabetic rats treated with PDBT capsule.
Values are expressed as mean±SEM., n=6; n=Number of animals.
* Significant decrease when compared with high fat fructose diet group, p<0.05
$ Significant increase when compared with control group, p<0.05
Control: vehicle control, HFD: High fat fructose diet control
3.1.3. Effect of PDBT Capsules on Lipid Profile of Rats

i. Effect on Serum Total Cholesterol Levels

Serum total cholesterol levels in all groups are shown in Figure 3. The mean total cholesterol levels were significantly increased in prediabetic control animals when compared with vehicle control rats. Formulations treatment showed statistically significant decrease in the total cholesterol levels. Decrease in elevated levels with the standard drug (Pioglitazone) treatment was found to be more (80.16±0.76mg/dL) as compared to PDBT formulation (86.27±0.74 mg/dL). Results indicate that the formulations are effective in the elevated total cholesterol levels.

ii. Effect on Serum Triglycerides Levels

Figure 4 shows the serum triglycerides levels in all groups. There was a significant increase in the triglyceride levels on the 15th and 30th day in prediabetes control rats as compared to normal rats. The triglyceride levels in rats treated with formulation significantly decreased on 15th and 30th day of drug treatment when compared with prediabetes control animals. Standard drug Pioglitazone reduced elevated triglyceride levels to 78.29±0.61mg/dL while PDBT formulation treatment reduced elevated triglyceride levels to 80.36±0.70 mg/dL. However the decrease is not statistically significant and not different from each other groups.

![Figure 4. Effect on triglycerides levels in prediabetic rats treated with PDBT capsules.](image)

* Significant increase when compared with high fat fructose diet group, p<0.05
* * Significantly decrease when compared with high fat fructose diet group, p<0.01
* * * Significantly decrease when compared with high fat fructose diet group, p<0.005
Control: vehicle control, HFD: High fat fructose diet control

3.1.4. Effect of PDBT Capsules on Liver Biomarker of Prediabetic Rats

i. Effect on AST Levels

Levels of AST are given in figure 6. Prediabetes caused increase in AST levels in HFD control as well as treatment groups. On 30th day of drug intervention decrease in AST level with PDBT capsule treatment (36.07±0.12U/L) as well as pioglitazone treatment (30.31±0.14 U/L) was found. However the decrease is not statistically significant and not different from each other groups.

![Figure 6. Effect on AST levels in prediabetic rats treated with PDBT capsule.](image)

Control: vehicle control, HFD: High fat fructose diet control

ii. Effect on ALT Levels

The ALT levels in all groups are shown in figure 7. Increase in ALT levels were found with administration of high fat fructose diet. The prediabetic animals treated PDBT capsules and standard drug showed decrease in ALT levels to 38.15±0.15 U/L and 36.51±0.15 U/L respectively. However the decrease is not statistically significant.

The decrease in liver marker levels AST and ALT in the prediabetic rats treated with PDBT capsule indicating that the formulation is hepatoprotective.
3.1.5. **Effect on Kidney Parameters**

**i. Effect on Serum Creatinine Levels**

The serum creatinine levels are shown in figure 8. Prediabetic animals treated with PDBT capsules and standard drug (Pioglitazone) showed decrease in the serum creatinine levels 0.8±0.09 mg/dL and 0.8±0.07 mg/dL respectively, when compared with prediabetic control animals. PDBT capsules treatment decreased serum creatinine level same as that of standard drug treatment group.

Decrease in the Creatinine levels of prediabetic animals indicates that the capsules are nephroprotective.

**ii. Effect on Blood Urea Nitrogen Levels**

Blood urea nitrogen levels are shown in figure 9. There was rise in blood urea nitrogen level found in prediabetic control rats when compared with normal rats. PDBT capsule treatment showed decrease in the blood urea nitrogen 18.47±0.09 mg/dL when compared with prediabetic treated rats, the same action showed in standard drug (18.38±0.07 mg/dL). The results indicate that the PDBT capsules have therapeutic activity same as that of standard drug (Pioglitazone).

3.2. **Results of Histopathology**

At the end of intervention the animals were sacrificed. The vital organs were removed and fixed in 10% formalin, and subjected to histopathological study.

**3.2.1. Pancreas**

In normal control rats many islets were evenly distributed throughout the cytoplasm of the pancreas (Figure 10-a). The prediabetic control rats (Figure 10-b) and prediabetic rats treated with pioglitazone (Figure 10-c), pancreas showed no any degeneration and necrosis of islet of Langerhans. The prediabetic rats treated with PDBT capsules (Figure 10-d) do not showed abnormal structure of islets.

**3.2.2. Liver**

The liver of normal rats (Figure 11-a), prediabetic control rats (Figure 11-b) and standard drug (Pioglitazone) treated rats (Figure 11-c) appeared normal in histopathological studies. In prediabetic rats treated with PDBT capsules (Figure 11-d) does not showed any granular degeneration.

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**Figure 7.** Effect on ALT levels in prediabetic rats treated with PDBT capsules.

| Control | HFD | HFD + pine | HFD + PDBT |
|---------|-----|------------|------------|
| 31      | 37  | 38         | 35         |

Control: vehicle control, HFD: High fat fructose diet control

**Figure 8.** Effect on Creatinine levels in prediabetic rats treated with PDBT capsules.

| Control | HFD | HFD + pine | HFD + PDBT |
|---------|-----|------------|------------|
| 0.6     | 0.8 | 0.9        | 0.7        |

Control: vehicle control, HFD: High fat fructose diet control

**Figure 9.** Effect on blood urea nitrogen levels in prediabetic rats treated with PDBT capsules.

| Control | HFD | HFD + STD | HFD + PDBT |
|---------|-----|-----------|------------|
| 20      | 22  | 23        | 21         |

Control: vehicle control, HFD: High fat fructose diet control

**Figure 10.** Histopathology study of pancreas of prediabetic rats treated with PDBT capsules.

10a normal control rats, 10b prediabetic control rats, 10c prediabetic rats treated with pioglitazone, 10d prediabetic rats treated with PDBT capsule

**Figure 11.** Histopathology study of liver of prediabetic rats treated with PDBT capsules.

11a normal control rats, 11b prediabetic control rats, 11c prediabetic rats treated with pioglitazone, 11d prediabetic rats treated with PDBT capsule
3.2.3. Kidney

Normal glomeruli and normal cellularity was found in normal rats (Figure 12-a). The prediabetic control rats (Figure 12-b) and prediabetic rats treated with standard drug (Pioglitazone) (Figure 12-c) also showed normal glomeruli and normal cellularity structure. Prediabetic rats treated with PDBT capsule (Figure 12-d) does not showed any cellular degeneration.

Figure 12. Histopathology study of kidney of prediabetic rats treated with PDBT capsules.

12a normal control rats, 12b prediabetic control rats, 12c prediabetic rats treated with pioglitazone, 12 d prediabetic rats treated with PDBT capsule

4. Discussion

Knowledge of ethnopharmacology, its holistic and systems approach supported by an experiential basis from traditional Indian Medicine – Ayurveda, can serve as an innovative and powerful knowledge system for discovery of newer, safer and more affordable drugs [23].

Plants as well as herbal formulations have been used for the treatment of diabetes for several years. Here in this study the combination of extracts of well-known Ayurvedic plants was formulated as PDBT capsule. The poly herbal formulation PDBT capsule was investigated for the activity against prediabetes. Our findings indicate that treatment of 200 mg/kg of a PDBT capsule for 4-weeks in prediabetic rats led to significant improvements in several features of the prediabetes. Because of the relationship between the metabolic syndrome and prediabetes, these results have important implications for public health.

The prediabetes was induced by orally feeding high fat fructose-fructose rich diet. Several reported studies are available on induction of prediabetic condition and insulin resistance with different combinations and for different time of induction [24-26]. The selected diet mimics an unhealthy diet containing high-fat products combined with high-sugar drinks [24].

In the present study it is found that the diet combination given to the Wistar strain albino rats caused increase in body weight, fasting blood glucose levels, serum TC, TG and HDL levels and liver and kidney markers. The diet caused the increase in fasting glucose level not very high but up to prediabetes level [27]. Insulin resistance and prediabetes was confirmed by oral glucose tolerance test. The diet was enriched with fructose and fats. This resulted in obesity and insulin resistance. High fat and obesity also induce oxidative stress, this might be reason for increase in liver and kidney markers [24, 28].

The herbal extracts used formulating of PDBT capsules exert cumulative action on hyperglycemia and insulin resistance. The treatment demonstrated statistically significant decrease from 2nd week onwards of drug treatment. At the end of 4th week the decrease in fasting glucose was more in PDBT capsule group that of pioglitazone treatment group.

The previous study reported suggests that increase in dietary fructose could be one of the factors for obesity and the insulin resistance development [29]. Consumption of high fat fructose diet resulted in increase in body weight in all groups of animals. Obesity is one of the underlying cause of insulin resistance. Pancreas unable to produce sufficient insulin due to obesity and which results into insulin resistance and finally type 2 diabetes [30]. Prediabetic groups when treated with PDBT capsule body weights significantly decreased as compared with HFD control group. This indicates that the PDBT capsules were effective in controlling the weight gain due to high fat fructose diet.

Consumption of cholesterol and saturated fats in diet was reflected in significant increase in cholesterol level. High amount of fructose in diet could be additional impact on lipid profile. High fructose in diet may cause increased lipogenesis and overproduction of triglycerides as well as cholesterol in hepatocytes [31]. The treatment with capsule demonstrated statistically significant decrease in elevated lipid profile. These results can be due to decrease in body weight as well as fasting glucose levels in treatment groups.

However high fat fructose diet consumption resulted in increased HDL cholesterol levels which was decreasing with with treatment. This may be due to the fact that lipid metabolism of rats is different than human [24]. Rats have strong capability to maintain their plasma cholesterol [29] additionally they do not have cholesterol-ester transfer protein (CETP) and HDL serves as major carrier of serum cholesterol [32]. The lipid metabolism in rats is primarily based on HDL rather than in human [33].

Oral feeding high fat fructose diet caused increase in the liver function markers AST and ALT, treatment of drugs decreased the elevated levels. Although the results are not statistically different this can be correlated with histopathological analysis of liver. The results show no impairment of liver cells no granular degeneration and all the samples are normal.

The consumption of large quantity of fructose caused increased blood urea nitrogen levels. The fructose diet has excessive consumption of ATP and degradation of nucleotides which results in increase blood urea nitrogen levels. The high-fructose diet also impacts on serum creatinine concentration. Phosphocreatine degradation produce creatinine is produced and excreted in urine. Increase in levels of plasma creatinine if kidneys are unable to eliminate such metabolic products [34].

In the present study the kidney function markers were increased with the consumption of high fat fructose diet but not very high. In histopathological study all the samples showed no abnormality in glomeruli and normal cellularity structure. The results from histopath study and kidney function markers levels in clearly indicates that drug treatment is not disturbing the kidney functions.
Similarly the histopathological study of pancreas also showed no inflammation and normal pancreas.

5. Conclusion

The results of this study indicate that PDBT capsules exhibited significant decrease in blood glucose levels on high fructose diet induced prediabetic rats. The effect of PDBT capsules on prediabetes was comparable with pioglitazone, a standard treatment for insulin resistance. The capsules resulted in improvement in hyperglycemia, dyslipidemia, obesity conditions in prediabetic rat model. The treatment did not cause any hepatotoxic, nephrotoxic and pancreatic toxic effects in rats.

Our findings suggest that, PDBT capsules may provide useful treatment for prediabetes, diabetes and related symptoms.

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