Evaluation of in vivo Synergistic Hypoglycemic & Hypolipidemic Activity of	1
Ethanolic Extract of Calotropis gigantean Leaves in Combination to	2
Metformin	3
in Alloxan Induced Rats	4
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## 7 **ABSTRACT:**

Aim: The present study was designed to investigate the antidiabetic & hypolipidemic activity of *Calotropis gigantean* in alloxan-induced diabetic rat model.

Study Design: *in vivo* study was carried out by ethanolic leaf extract was administered in 250mg/kg body weight concentration and then subjected to different rats models to authenticate the antidiabetic and hyperlipidimic properties of the plant.

Place and Duration of Study: Department of Pharmacy, Southeast University, Banani, Dhaka-1213
 Bangladesh within a period of July 2018 to December, 2018.

Methodology: Diabetes was induced in rats by an intraperitoneal injection (i.p) of alloxan (100 mg/kg B.W). Ethanolic leaf extract of *C. gigantean* (250 mg/kg B.W) was administrated orally as a single dose per day to the diabetic rats for 7 days. The negative control group received 0.5 ml of sterile normal saline water & positive control group received metformin with 100 mg/kg B.W & 50 mg/kg B.W doses in combination to extract to evaluate synergistic effect. After 7 days study period, fasting blood glucose, total cholesterol, triglyceride, high-density lipoprotein cholesterol, liver weight & body weight were measured

21 **Results:** Individual plant extract & standard reduced FBG significantly by 52% (P<0.001) & 55.3% 22 (P<0.001) correspondingly. Metformin (100mg/kg B.W) potentiated reduction (68%) (P<0.001) when combined to plant extract (250 mg/Kg B.W). Significant dose dependent manner was followed when 23 24 metformin (50 mg/kg B.W) was combined to plant extract (250mg/Kg B.W). Our results clearly suggest 25 that C. gigantean exhibit hypoglycemic & hypolipidemic activity with an alteration in body-liver weight. The 26 present study also suggested to develop a combination therapy of extract along with metfromin in 27 different doses to minimize the intake of synthetic drug. Significant reduction of TG, TC were noted by 28 extract (250 mg/kg B.W) with 32.42% (P<0.001) & 41.32% (P<0.001) respectively where standard shown 29 the diminution 43.43% (P<0.05) & 47.21% (P<0.001) respectively as compare to Untreated diabetic rats. 30 50.21% (P<0.01) & 42.38% (P<0.001) reduction of TG & TC were estimated by C.gigantea extracts (250 31 mg/kg B.W) when combined with Metformin (100 mg/kg B.W). 34.53% (P<0.05) & 41.54% (P<0.001) 32 reduction of TG & TC by C.gigantea extracts (250 mg/kg B.W) were confirmed when combined to 33 Metformin (50 mg/kg B.W). Combination therapy also shown synergistic effect in elevation of plasma 34 HDL-cholesterol.

35 **Conclusion:** The results of the study concluded that *Tricosanthes tricuspidata and Clematis montana* leaf 36 and root extracts have potential antidiabetic and antioxidant properties.

37

38 *Keywords:* Calotropis gigantean, diabetes mellitus, hypolipidemic activity & antidiabetic activity.

39

40 1. BACKGROUND

41

42 As Diabetes mellitus is a public health challenge the complications are on a rise in day to day 42 life According to World Health Organization the diabetic population is likely to increase up to

43 life. According to World Health Organization the diabetic population is likely to increase up to

44 300 million or more by the year 2025 [1]. Diabetes mellitus (DM) is a severe physiological problem being one of the major causes of death all over the world, and if not treated, it can lead to many complications 45 46 [2] such as long term damage, dysfunction, and failure of various organs [3]. This disease is caused by 47 the destruction or dysfunction of pancreatic of  $\beta$ -cell and insulin resistance which results in elevating 48 blood glucose level, known as hyperglycemia [4, 5]. Aldose reductases, a key enzyme in the polyol 49 pathway catalyze the glucose to be reduced to sorbitol. Accumulation of sorbitol in the body causes 50 various complications [6]. Over time, diabetic patients with poor glycemic control undergo various life 51 threatening difficulties which include nephropathy, retinopathy, neuropathy, and cardiovascular diseases [7]. Alongside with exercise, modern drugs such as pioglitazone, biguanides, meglitinides, 52 53 thiazolidinedione, alpha glucosidase inhibitors and sulphonylureas shows considerable benefits with side 54 effects like hypoglycemia, GIT disturbance, water intoxication, and hyponatremia, obesity with high cost for long term use [8]. Numerous agents that are currently used for the treatment of type 2 diabetes are 55 facing limited efficacy and tolerability [9]. For instance, sulforyly as induce  $\beta$ -cell death in isolated rodent 56 57 and human islets while glucagon-like peptide-1 receptor agonists and dipeptidyl peptidase-4 inhibitors 58 have potential risks for pancreatitis, pancreatic, and thyroid cancers [10]. So natural compounds could be 59 great substitute when taken with synthetic drugs.

60

Now a days medicinal plants show the proof to be used as hypoglycemic agent as most of plants contain glycosides, alkaloids, terpenoids, flavonoids, carotenoids **etc.**[11] that significantly posses antidiabetic effect. Antihyperglycemic activity of the plants is mainly due to their ability to renovate the function of pancreatic tissues by causing an elevation in insulin output or hindering the intestinal absorption of glucose, facilitating of metabolites in insulin dependent or amylase and glucosidase inhibitor as these enzymes are responsible for breaking  $\alpha$ - 1, 4 bonds in complex carbohydrate to elevate FBGL [12].

67

68 Calotropis gigantean (Family: Apocynaceae) is a common weed in open waste ground, roadsides, village surroundings and railway lines. It is native to continental Asia and South-East Asia and has been 69 70 introduced in the Pacific Islands, Australia, Central and northern South America and Africa [13]. Mainly known as crown flower, crown plant, giant milkweed or rubber bush [14]. Different parts of the plant 71 72 contains stigmasterol, ß-sitosterol [15], mudarine, glycosides (calotropin uscharin, calotoxin), lupeol, 73 calotropin, uscharin, calotoxin, calactin and uscharidin; gigantin, protease such as calotropin DI and DII 74 and calotropin FI and FII [16]. Calotropnaphthalene, calotropises juiterpenol, calotropisesterterpenol and 75 calotropbenzofuranone along with sucrose, have been isolated from Calotropis gigantean [17]. 76 Traditionally different parts of the plant are used such as in leprosy, eczema, syphilis, elephantiasis, 77 ulceration, and cough [18] purgative and gastrointestinal irritant and abortion inducer [19], paralysis, 78 swellings, intermittent fevers, asthma, anorexia, helmintic infections, inflammations, cutaneous infections, 79 intestinal worms, ascites, bronchitis and dyspepsia and promotes gastric secretions [20], in poisonous 80 snake or rat bites, periodic fever, vatha diseases, ulcers, cures dental problems, gonococcal arthritis and 81 other rheumatic complaints[21]. The plant proves to hold some pharmacological effects like antipyretic [22], proteolytic activity [23] antiamoebic [24] wound healing [25], hepatoprotective [26] and anti-oxidant 82 83 [27] properties. Other reported potentials are analgesic activity [28], antimicrobial [29] and cytotoxic 84 activity [30], anti-diarrhoeal activity anti-Candida activity [31], anti-pyretic activity [32], insecticidal activity 85 [33], CNS activity [34] and pregnancy interceptive properties [35], procoagulant activity [36]. C. gigantea is reported to possess major phytochemical groups as alkaloids, cyanogenic, glycosides, phenolics, 86 87 tannins [37], cardenolides, ester [38,39], flavonoids [40], terpenes [41] (antimosquito larvicidal activity) sterols (Campesterol, Stigmasterol, gamma-Sitosterol, Desmosterol anticervical cancer property), 88 89 Proteinases [42] and nonprotein amino acid [43]. Acetates and the benzoates,  $\alpha$ -and  $\beta$ -calotropeols and β-amyrin, tetracyclic triterpene compounds and traces of sterols, Giganteol acetate, Giganteol are also 90 91 reported by P. Bhaskara et al. [44].

92 Therefore, the aim of this study was to find out the scientific basis of the use *C. gigantean* in the 93 management of diabetes & hyperlipidemia used by traditional practitioners using ethanol extracts on 94 alloxan-induced diabetic mice.

## 95 2. MATERIALS AND METHODS

### 96 2.1 Experimental Animals

97 30 Long Evan rats with (gender: male, wg: 80±10g) were obtained from ICDDR, B Mohakhali, Dhaka, 98 Bangladesh. Rats were housed under standard laboratory conditions (22-25<sup>0</sup>C, humidity 40-60%,12 hr 99 light:12 hr dark cycle) and housed in standard size metallic cages (5 rats/ cages) in properly ventilated 100 room. Through the experiments all rats were fed with standard laboratory diet. Prior to the beginning of 101 the study, animals were allowed for two weeks to acclimatize to laboratory conditions.

### **2.2 Collection of Plant Material and Preparation of Extracts**

C. gigantea plant was collected from the natural population growing in the Gazipur, Dhaka, Bagladesh & 103 authenticated by the expert taxonomist from Bangladesh National Herbarium, Mirpur, Dhaka, Bangladesh 104 105 (Accession number: 45130). Leaves were washed and shade dried for several days followed by grinding using mechanical grinder. About 200 gm dried powder were soaked in 800 ml ethanol and kept for a 106 107 period of about 7 days with occasional shaking and stirring. The whole mixture is then filtered through Whatman No.1 filters paper and concentrated by a rotary evaporate under reduced pressure at 50°C 108 109 temperature to afford crude extract with gummy or semisolid appearance. The concentrate was stored in an airtight container and kept in a cool, dark and dry place until the next course of action. 110

#### 111 2.3 Chemicals

- 112 Alloxan and metformin were purchased from Sigma-Aldrich and Merck company (Germany) respectively.
- All other used chemicals were of analytical grade and were obtained from standard commercial suppliers.

## 114 **2.4 Induction of Diabetes**

Diabetes was induced in overnight fasted Evan rats by single-dose intraperitoneal injection of freshly prepared alloxan at 140 mg/kg body weight dissolved in 0.5 ml of sterile normal saline water and drank 10% glucose solution to overcome drug induced hypoglycemia. After 72 hours blood glucose level was measured by using tail blood sample. Rats with fasting blood glucose level above 7.0 mmol/L were selected for further study.

#### 120 2.5 Experimental Design

Long Evan rats were randomly assigned into group I, II, III, IV, V, VI with 5 rats in each group for the respective one week treatment due to determination of blood glucose, lipid profile test studies.

- 123 Group I: Non Diabetic Normal Control (Only water & normal diet)
- 124 **Group II:** Diabetic Control (Only water & normal diet)
- 125 **Group III:** Diabetic Control+ Metformin (100 mg/kg B.W in 0.5 ml 99% DMSO)

- Group IV: Metformin(50 mg/kg B.W) + Ethanolic Extract of *C.gigantea* (250mg/kg B.W in 0.5 ml 99%
   DMSO)
- **Group V:** Metformin (100 mg/kg B.W) + Ethanolic Extract of *C.gigantea* (250 mg/kg B.W in 0.5 ml 99%
   DMSO)
- 130 **Grroup VI:** Diabetic Control+ Ethanolic Extract of *C.gigantea* (250 mg/kg B.W in 0.5 ml 99% DMSO)

## **2.6 Collection of blood and determination of Biochemical Parameters**

At 0<sup>th</sup>, 3<sup>th</sup>, 5<sup>th</sup> & 7<sup>th</sup> day, blood samples were collected from tail vein after the administration of metformin 132 & ethanolic extract of *C.gigantea* and blood glucose levels were determined by using by glucose meter. 133 After completing the one week treatment the rats were at first anesthetized with chloroform and 3 ml of 134 blood was directly collected from heart by syringe. Immediately after blood samples collection, serum was 135 isolated by centrifugation at 4000 rpm for 20 min and then analyzed for various biochemical parameters. 136 The serum samples were stored at -80 °C in a freezer until they were analyzed. The concentration of TC, 137 TG, HDL-Cholesterol were measured colorimetrically by blood analyzer using commercially available wet 138 139 reagent diagnostic kits (HUMAN GmbH, Germany).

## 140 2.7 Statistical Analysis

141 The results were expressed as mean  $\pm$  SD. Data analysis was performed by the SPSS version 24 142 (SPSS/IBM, Chicago, IL) using one-way analysis of variance (ANOVA) and Dunnett's test. To assess the 143 individual variations between the control and treatment groups, *P* < 0.05 was considered significance 144 level.

## 145 **3. RESULTS**

## 146 **3.1 Antidiabetic Activity:**

147 **Table 1**: Effect of *C. gigentea* on fasting blood glucose levels in alloxan induced diabetic rats.

	Day of treatment			
Groups	Day 1	Day 3	Day 5	Day 7
Control	4.3±0.29	3.9±0.37	3.7±0.20	4.02±0.26
Untreated diabetic	8.02±0.53	9.05±1.02	13.6±1.02	$15.00 \pm 3.15$
		(12.84)	(69.58)	(87.03)
Diabetic+Metformin	9.6±0.98***	6.07±0.37***	5.62±0.07***	4.30±0.07***
(100 mg/kg B.W)		(36.77)	(41.46)	(55.21)
Metformin (100 mg/kg	8.6±0.37***	6.2±0.12***	5.44±0.17***	2.75±0.35***
B.W)+Extract (250		(27.91)	(36.74)	(68.02)
mg/kg B.W)				
Metformin	8.62±0.28***	7.2±0.12***	5.92±0.09***	3.48±0.37***
(50 mg/kg)+Extract (250		(16.47)	(31.32)	(59.63)

mg/kg B.W)				
Extract (250 mg/kg	8.74±0.46***	6.98±0.24***	5.76±0.29***	4.18±0.24***
<b>B.W</b> )		(20.14)	(34.1)	(52.17)

Values are expressed as mean  $\pm$  SD (*n* = 5 rats). Significance level among different groups at *P* < 0.05. (*P*<0.05; *P*<0.01, *P*<0.001); Diabetic rats were compared with normal rats. **Metformin** and *C.gigantea* 

150 treated diabetic rats were compared with diabetic rats.

At all-time points, blood glucose concentration remain unchanged (p<0.001) in normal rats treated with 151 distilled water. BGL were gradually decreased for each group at 1<sup>st</sup>, 3<sup>rd</sup>, 5<sup>th</sup> & 7<sup>th</sup> day. The FBGL of all 152 groups were compared to untreated diabetic group (Table: 01, Figure: 01). At 7<sup>th</sup> day, oral administration 153 of C.gigantea extracts (250 mg/kg B.W) significantly decreased the blood glucose level 52.17% 154 155 (P<0.001). Combination therapy was performed to establish synergistic effect with two doses. Metformin (100 mg/kg B.W) has shown significant FBGL reduction by 55.21% (P<0.001) individually but potentiated 156 157 reduction (68.02%) (P<0.001) when combined to plant extract (250 mg/Kg B.W). Dose dependent manner 158 (59.63%) (P<0.001) was followed when metformin (50 mg/kg B.W) was combined to plant extract (250 159 mg/Kg B.W) with reduced dose. The possible mechanism by which C.gigantea brings about its 160 hypoglycemic action may be stimulating the insulin effect of serum by increasing either the pancreatic 161 secretion of insulin from the beta - cells of islets of langerhans or its release from bound insulin. Thus, the 162 significant antidiabetic effect of the extracts of A.remota could be due to the presence of the flavonoids, 163 tannin and alkaloid in the extracts, which could act synergistically and/or independently to enhance the 164 activity of glycolytic enzymes.

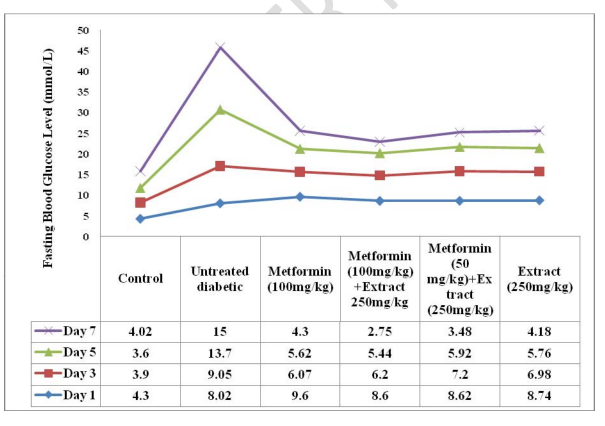


Figure 01: Effect of *C.gigentea* extracts & metformin on fasting blood glucose level in alloxan induced diabetic rats.

## 168 **3.2 Hypolipidemic Activity:**

Groups	Organ weight (mg/g)	Lipid profile (mmol/l)		
	Liver	TG	TC	HDL-C
Control	4.73	3.75±0.95	8.86±0.95*	8.12±0.41
Untreated diabetic	4.15***	4.72±0.64***	9.51±0.76***	5.43±0.20***
	(12.26)	25.87	7.34	33.13
Diabetic+ Metformin	4.49*	2.67±0.29***	5.02±0.12***	6.78±0.415**
(100 mg/kg B.W)	(8.19)	(43.43)	(47.21)	(24.86)
Metformin (100	4.61**	2.35±0.37**	5.48±0.46***	6.81±0.26**
mg/kg)+Extract (250 mg/kg	(11.08)	(50.21)	(42.38)	(25.23)
<b>B.</b> W)				
Metformin (50 mg/kg	4.31**	3.09±0.49	5.56±0.62***	6.64±0.98**
B.W)+Extract (250 mg/kg	(3.86)	(34.53)*	(41.54)	(22.28)
<b>B.W</b> )				
Extract (250 mg/kg B.W)	4.25***	3.19±0.40	5.58±0.35***	6.60±0.415**
	(2.41)	(32.42)*	(41.32)	(21.55)

### 169 **Table 02: Effect of** *C.gigentea* **on lipid profile in alloxan induced diabetic rats.**

170 Values are expressed as mean  $\pm$  SD (n = 5 rats). Significance level among different groups at P < 0.05. 171 (P < 0.05; P < 0.01, P < 0.001); Diabetic rats were compared with normal rats. **Metformin** and *C.gigantea* 172 treated diabetic rats were compared with diabetic rats

treated diabetic rats were compared with diabetic rats.

## 173 Table 03: Effect of *C.gigentea* on mean weight of liver in alloxan induced diabetic rats.

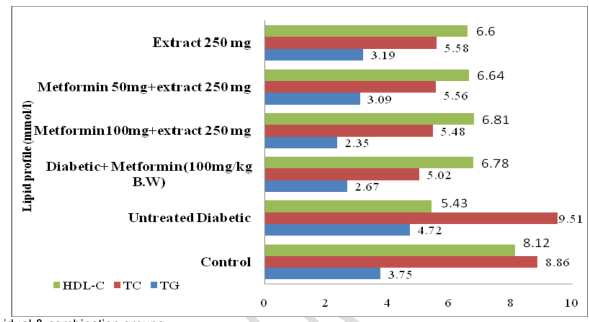
Groups	Mean weight of animals at sacrificial time (Grams)	Mean weight of Liver at sacrificial time (Grams)	Weight of liver in grams/kg body weight
Control	110	4.73	43 gm/kg
Untreated diabetic	95	4.15	43.68 gm/kg

#### 174

**Table 02 & Figure 02** showed that the effect of the *C.gigantea* extract on TG, TC, HDL in alloxanized diabetic rats. After alloxan induced, the result showed that TG, TC increased while HDL decreased compare to Untreated diabetic rats (Table: 02). Highest reduction of TG, TC were shown by Metformin (100 mg/kg B.W) like 43.43% (*P*<0.05) & 47.21% (*P*<0.001) respectively where extracts shown significant diminution by 32.42% (*P*<0.001) & 41.32% (*P*<0.001) respectively. Combination study of Extract to Meformin was performed to develop the synergistic effect with different doses in dose dependent activity.

Metformin (100 mg/kg B.W) with *C.gigantea* extracts (250mg/kg) reduced TG & TC by 50.21% (*P*<0.01) & 42.38% (*P*<0.001) respectively & metformin (50 mg/kg B.W) with *C.gigantea* extracts (250mg/kg) lessened TC & TC level by 34.53% (*P*<0.05) & 41.54% (*P*<0.001) The administration of the extract of

184 C.gigantea produced a significant increase in the level of High-density lipoprotein-cholesterol (HDL-C) in



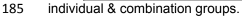
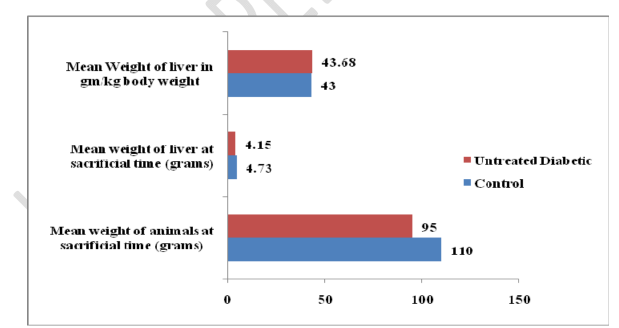




Figure 02: Effect of *C.gigentea* on lipid profile in alloxan induced diabetic rats.



#### 187

Figure 03: Effect of *C.gigentea* on mean weight of liver in alloxan induced diabetic rats.

Significant decrease of liver weight was revealed in diabetic rat (12.26%) (*P*<0.001) as compared to control. The tested compound confirmed slightly significant increase (2.41%) (*P*<0.001) of liver weight in

- comparison to untreated diabetic. Metformin (100 mg/kg B.W) with *C.gigantea* extracts (250 mg/kg B.W)
  & Metformin (50 mg/kg B.W) with *C.gigantea* extracts (250 mg/kg) increase liver weight 11.08% & 3.86%
  (*P*<0.01). During sacrificial time mean liver weight & body weight of alloxanized group also compared to</li>
  that of control group (Table: 03, Figure: 03). Liver weight was slightly increased in diabetic rats (43.00
- 194 gm/kg B.W & 43.68 gm/kg B.W) when compared with non-diabetic rats.

195 The observed significant reduction in serum total lipids, total cholesterol and LDL cholesterol by the 196 extract which can be attributed to the phytochemical constituents that propose the use of the plant to 197 prevent cardiovascular complications arising from hyperlipidemia [45]

198

### 199 Discussion

New antidiabetic drugs from natural plants are already in search that contain phytochemical compounds with high efficacy with minimum toxicity. As most of plants contain glycosides, alkaloids, terpenoids, flavonoids, carotenoids, *etc.*, that are significantly posses antidiabetic effect [12] Plant extracts are evaluated to balance the liberation and absorption of glucose is becoming a striking therapeutic choice in the treatment of diabetes mellitus.

205

206 Alloxan, a beta cytotoxic agent, rapidly and selectively accumulates in pancreatic beta cells] and causes 207 beta cell death and apoptosis by generation of reactive oxygen species (ROS), super oxide radicals and 208 hydrogen peroxide [46]. Sequential injection of alloxan caused a significant increase (p<0.05) in blood 209 glucose concentration for 7 days in all group of rats compared with their respective baseline blood glucose and to control values. Single & combination therapy was performed to establish synergistic effect 210 with two doses of metformin for 7 days. The estimated results were taken after 7<sup>th</sup> days. Individual plant 211 extract & standard reduced FBG significantly by 52% (P<0.001) & 55.3% (P<0.001) correspondingly. 212 Metformin (100mg/kg B.W) potentiated reduction (68%) (P<0.001) when combined to plant extract (250 213 214 mg/Kg B.W). Significant dose dependent manner was followed when metformin (50 mg/kg B.W) was 215 combined to plant extract (250mg/Kg B.W) with reduced dose. This results can led to a development of 216 new drug design with reduced dose of standard when taken with leaf extract of C.gigantea. The studied 217 plant may reduce absorption of glucose from the small intestine as glucose liberation from disaccharides 218 is reduced. In our study, it is found that extract have also hypoglycemic effect in glucose induced 219 hyperglycemic rats. Diabetes induction mainly alters morphological changes & level of enzymatic 220 metabolism. In the study liver size were measured for extract along with standard. Significant & dose 221 dependant increase of liver size were found in rats in comparison to untreated diabetic.

222

223 Hyperlipidemia is a recognized outcome of Diabetes mellitus [47]. Abnormal high concentration of serum 224 lipids result from increase in the mobilization of free fatty acids from the peripheral storehouse. The 225 marked hyperlipidaemia that characterizes the diabetic state is the consequence of the dysfunction of 226 lipolytic hormones on the fat depots [48]. Hyperlipidemia associated with diabetes mellitus is reduced by 227 limited absorption of free fatty acids and free cholesterol following inhibition of pancreatic lipase and pancreatic cholesterol esterase [49]. Significant reduction of TG, TC were noted by extract (250 mg/kg 228 B.W) with 32.42% (P<0.001) & 41.32% (P<0.001) respectively where Standard shown the diminution 229 230 43.43% (P<0.05) & 47.21% (P<0.001) respectively as compare to Untreated diabetic rats. 50.21% (P<0.01) & 42.38% (P<0.001) reduction of TG & TC were studied by C.gigantea extracts (250 mg/kg 231 232 B.W) with Metformin (100 mg/kg B.W). TG (34.53%) (P<0.05) & TC (41.54%) (P<0.001) reduction by 233 C.gigantea extracts (250 mg/kg B.W) were expressed when combined to Metformin (50 mg/kg B.W). 234 Combination therapy alsoshown synergistic effect in elevation of plasma HDL-cholesterol that prevent risk 235 of developing cardiovascular disease. The administration of the extract of C.gigantea produced a 236 significant increase in the level of High-density lipoprotein-cholesterol (HDL-C). The plant demonstrated a

237 cardioprotective effect via an increase in HDL-cholesterol levels. The extracts of C.gigantea prove to 238 have a hypolipidemic potential. Alteration of liver weight is also related to diabetic. The present study has 239 shown related reduction of liver weight according to the dose of studied sample & standard in individual & 240 combination design. The liver is an insulin-sensitive organ that undergoes functional abnormalities in 241 individuals with untreated diabetes [50]. In this study, the liver of diabetic animals & control animals were 242 compared. An increase (hypertrophy) in the weight of liver in proportion to the body weight was observed 243 despite the reduction of the mean weight of all the animals in Alloxan induced group. It could be ascribed to increased triglyceride accumulation that can lead to liver enlargement by reason of increased entry of 244 245 fatty acids into the liver induced by hypoinsulinemia [51] and the less elimination of lipoprotein from liver. 246 Previous research articles also present the same agreement with the present findings [52]. 247

## 248 4. Conclusion

It had been concluded that in our study, decrease in the concentration of glucose, total triglyceride, total cholesterol, and increase in HDL cholesterol were observed for ethanolic extract of *C.gigantea* leaves along with metformin. The antidiabetic and hypolipidemic activity of the plant source is due to the phyto chemical constituents present in the plant. This justifies its use in ethnomedicine and can be exploited in the management of diabetes induced hyperlipidemia. Further studies are in progress for isolation and identification of lead compound to design a combination therapy in conjunction with synthetic drug.

255

## 256 **Ethical Considerations**

This was carried out in strict compliance with the National Research council guidelines on the care and use of laboratory animals to minimize research animal pain and suffering [53].

## 259 **Conflict of Interests**

- 260 The authors declare that they have no conflicts of interest.
- 261

## 262 Data Availability

263

The data used to support the findings of this study are included within the article.

265 266

# 267 **Reference**

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