Anti-diabetic activity of traditional Indian gold containing preparation: Shadguna Balijarita Makaradhwaja on streptozotocin induced diabetic rats

Sanjay Khedekar¹, Galib Rukkudin², Basavaiah Ravishankar³, Pradeepkumar Prajapati⁴

¹Department of Rasashastra and Bhaishajya Kalpana, Assistant Director for Post Graduate Studies, SSAM & H, Hirawadi, Maharashtra University of Health Sciences, Nashik, Maharashtra, India, ²Department of Rasashastra and Bhaishajya Kalpana, Institute of Post Graduate Teaching and Research in Ayurveda, Gujarat Ayurved University, Jamnagar, India, ³Department of Rasashastra and Bhaishajya Kalpana, SDM Centre for Research in Ayurveda & Allied Sciences, SDM College of Ayurveda, Udupi, Karnataka, India, ⁴Department of Rasashastra and Bhaishajya Kalpana, Institute of Post Graduate Teaching and Research in Ayurveda, Gujarat Ayurved University, Jamnagar, India

ABSTRACT

Background: Makaradhwaja a gold containing mercurial preparation used for diabetes mellitus in indigenous system of medicine. It is a popular aphrodisiac and rejuvenator traditional medicine. It is prepared by using processed gold, mercury and sulfur in different ratios by applying intermittent heating pattern in Valuka Yantra. Objectives: The aim of the study was to evaluate anti-diabetic effect of Shadguna Balijarita Makaradhwaja (SBM) on streptozotocin (STZ) induced diabetic rats. Materials and Methods: Diabetes was induced to normal rats by injecting STZ in dose 40 mg/kg. Powdered SBM and dried extract of Tinospora cordifolia were mixed with honey and administered orally for 20 days at dose 2.63 mg/kg and 42.34 mg/kg body weight, respectively. The effects of treatment on body weight changes and blood glucose levels were quantified on day 1, 5, 10, 15 and 21 of the experiments. On the 21st day, animals were sacrificed and gross histopathological changes in liver, kidney and pancreas were illustrated. Blood sugar level, glycated hemoglobin, blood urea, serum cholesterol, serum creatinine, serum triglyceride and serum protein were estimated with standard methods. The study was conducted in the year 2011. Results: Test drug observed significant decrease (P < 0.001) in glycated hemoglobin level compared to diabetic control rats. Blood sugar level of test drug group shown a significant decrease (279.11 ± 57.95) compared with diabetic rats. Conclusion: The present study demonstrates that SBM and dried extract of T. cordifolia with honey significantly reduces the blood glucose level and shows anti-diabetic effect.

KEY WORDS: Anti-diabetic activity, cinnabar gold, Kupipakwa Rasayana, mercury sulfide Shadguna Balijarita Makaradhwaja

INTRODUCTION

Diabetes mellitus is threatening for the 21st century. It will be leading cause of morbidity and mortality in near future due to increasing incidence worldwide. Herbo-mineral-metallic drugs of Ayurveda having potential of decreasing blood sugar levels and found efficient on experimental animal models. Makaradhwaja [1] is such herbo-mineral-metallic composition.

Medieval classical texts of Ayurveda quoted that Makaradhwaja is one of the best rejuvenator and aphrodisiac [2] agent. It is prepared by using processed gold, mercury and sulfur in the ratio of 1:8:24 or 1:8:48 by sublimation in the traditional system of heating device Valuka Yantra [1,2]. Nowadays, it is prepared in modified vertical electrical muffle furnace (modified Valuka Yantra) [3]. It gives synergistic action with different herbs in the various disorders. It is therapeutically efficient in disorders such as Madhumeha (diabetes mellitus), Jwara (remittent fever), Kushta (skin disorders), and Raktaja Vikara (blood disorders) [2]. Previous studies claimed its safe use without any untoward effect and toxicity [4]. Specific action on cell-mediated immunity is proved by its immune-modulator
action [5]. In experimental and clinical studies, it is found effective in diabetes [6].

Guduchi (Tinospora cordifolia Linn.) is well-known Madhumehahar (anti-hyperglycemic) herb described throughout ayurvedic classics [7]. Most commonly, its stem is used for medicinal purposes. It is well-known anti-oxidant, anti-hyperglycemic, immune-modulator, and rejuvenator [8]. Its different formulations are quoted in the ayurvedic classics like juice, decoction, powder and Ghana (dried extract) as per diseases.

Madhu (honey) is mentioned as Madhumehahar (anti-diabetic) agent within classical texts of Ayurveda [9]. Its anti-diabetic activity is reported by Erejuwa et al. [10]. It is used as a vehicle drug (logavahi) in the numerous herbo mineral formulations. It is mentioned in the reference to Makaradhwaja as a vehicle drug [11]. In the present study, it was used as a vehicle drug as prescribed in texts of Rasashastra.

The present study was planned to evaluate anti-diabetic activity of Shadguna Balijarita Makaradhwaja (SBM) and T. cordifolia Linn with honey in streptozotocin (STZ) induced diabetic rats.

MATERIALS AND METHODS

Preparation of SBM and Guduchi Ghana

Test drug SBM was prepared as per the classical text reference in the Department of Rasashastra and Bhaishajya Kalpana, Institute of Post Graduate Teaching and Research in Ayurveda (IPGT and RA), Gujarat Ayurved University, Jamnagar in 2011 [1]. Raw Material Swarna (gold) was purchased from the local market, and Hingula (cinnabar) and Gandhaka (sulfur) were collected from Pharmacy of Gujarat Ayurved University, Jamnagar. Gold was subjected to Shodhana and after Shodhana, its foils were prepared. Cinnabar was processed to prepared. Cinnabar was processed to

Makaradhwaja is consisted of red sulfide of mercury and having an empirical formula of HgS (mercury sulfide). Inductively coupled plasma optical emission spectrometry was observed that it contains 1.2% of gold with mercury and sulfur in finished product as a major element.

T. cordifolia’s stems were collected from the herbal garden of Gujarat Ayurved University, Jamnagar. Stems of T. cordifolia were cut into small pieces and crushed. These crushed pieces were cooked with 4 times of potable water ant reduced at 1/4th of the same to prepare decoction. The decoction was sieved, cooked to semisolid consistency. The semisolid mass was dried in hot air oven at 45°C to prepare dry extract [15]. Average 5.21% yield was obtained from the stem extract. The Guduchi Ghana (dried extract) was powdered and stored [16].

Honey was collected from the local forest of Jamnagar.

Experimental Animals

Albino rats (160-220 g) of either sex were selected for this study. Animals were procured from the Animal house of Pharmacology laboratory of IPGT and RA, Gujarat Ayurved University, Jamnagar. Permission for the experiment was granted by Animal Ethical Committee of the Same Institute (IAEC-06/09-11/02). The animals were fed pellet diet and water. As per the guidelines of National Institute of Health’s guide for the Care and Use of Laboratory Animals, the study was conducted [17].

Collection of Blood Samples

Tail veni-puncture method was applied for the collection of blood sample in the rats. For investigation Glucometer strip was used, and reading was noted down. By adopting this procedure blood glucose level of animals was estimated, prior injection of STZ, 5th, 10th, 15th, and 20th day during trials.

The sacrifice was carried out on the 21st day after completion of the study. Animals were anesthetized and stroked over tiles for sacrifice. The blood sample was collected by the dissection of jugular veins of animals. Biochemical parameters were investigated at pathology laboratory, IPGT, and RA, Jamnagar.

Experimental Induction of Diabetes

Diabetes was induced to rats by single intra peritoneal injection of STZ (40 mg/kg). STZ was weighed individually for each animal, according to its weight it was solubilized with 0.2 mL saline (154 mM NaCl) just prior to injection. After 72 h of STZ injection, rats with diabetic hyperglycemia (blood glucose more or equal to 250 mg/dL) were selected for experiment. Suspension of Makaradhwaja and T. cordifolia with honey was started fed to selected diabetic rats.

Anti-diabetic Activity

Anti-diabetic activity was evaluated by the effect of test drugs on the ponderal and biochemical parameters. Ponderal
parameters like gross body and different body organs weight were evaluated. Biochemical variables such as blood sugar, glycated hemoglobin (HbA1C), serum total cholesterol, serum high-density lipoprotein (HDL), serum triglyceride, serum creatinine, blood urea, serum glutamic-pyruvic transaminase (SGPT), serum glutamic oxaloacetic transaminase (SGOT), total protein, albumin, and globulin were estimated. Sacrificed animals gross and histological appearance of vital organs (liver, pancreas and kidney) were examined at the end of the study.

Experimental Design

Diabetic animals were divided into four groups. Food and water were provided to the animals.

- **Group 1**: Normal control (NC)
- **Group 2**: Diabetic control (DC)
- **Group 3**: DC + SBM and Guduchi Ghana with honey suspension (SBM)
- **Group 4**: DC + glibenclamide (reference standard [RS])

Dose Fixation

The dose of the drug was calculated by extrapolating the therapeutic dose to rat on the basis of body surface area ratio by referring to the table of Paget and Barnes (1964) [18].

Dose

Powdered SBM 2.63 mg/kg rat and dried extract of *T. cordifolia* 42.34 mg/kg rat were grinded with honey to prepare suspension. This suspension was administered orally to test drug group SBM diabetic animals. Glibenclamide, in the dose 0.45 mg/kg was administered to RS drug control group.

Statistical Analysis

All the values of were expressed as mean ± standard error mean. A statistical analysis was performed by using Student’s *t*-test. It was calculated by using Microsoft excel programmer.

RESULTS

In NC rats, during the course of 21 days, 13.90% weight was increased. Insignificant decrease in body weight was observed in SBM and RS group in comparison to DC rats [Table 1]. In DC rats, weight of kidney and liver was increased up to significant extent. Treatment with test drug did not affect the weight of these organs to significant extent in comparison to the DC group [Table 2].

Initial blood sugar level was decreased by 44.27% and 44.04% in SBM and RS treated groups respectively [Table 3]. Raised levels of HbA1C were significantly attenuated by test drug SBM and RS [Table 4]. Increased blood sugar levels were insignificantly decreased by SBM and RS test drugs.

Non-significant rise in serum cholesterol, triglyceride and HDL were moderately decreased by administration of SBM. Blood urea levels decreased significantly in group SBM and moderately decreased in RS group. Elevated serum creatinine levels non-significantly decreased by administration of SBM which was non-significantly increased in RS group in comparison to DC group. Decreased SGOT parameter in DC group was significantly attenuated by SBM and RS drugs. Increased levels of SGPT were decreased up to significant extent by SBM and RS drugs [Table 5].

The accumulation of fat in liver was observed in histopathological study. SBM and RS treated animals restored the histological changes [Figure 1a-d]. Inflammation in blood vessels, increase in the thickness of bowman capsules, fat deposition, and change in size of the glomerulus were found in the kidney of

**Table 1: Effect of test drug on body weight in STZ diabetic Wister strain albino rats**

| Group       | 0 day (g)       | 5th day (g)      | 10th day (g)     | 15th day (g)     | 21st day (g)     | % change in comparison to 0 day |
|-------------|-----------------|------------------|------------------|------------------|------------------|--------------------------------|
| NC          | 187.00±10.95    | 192.00±11.27     | 196.33±11.03     | 205.67±10.28     | 213.00±10.57     | 13.90↑                         |
| DC (STZ)    | 170.67±11.73    | 165.67±11.49     | 165.67±14.00     | 167.00±13.61     | 166.67±15.05     | 02.34↓                         |
| SBM         | 176.33±07.79    | 170.33±08.20     | 164.33±08.54     | 158.00±09.18     | 152.00±09.24     | 13.80↓                         |
| RS          | 172.67±04.15    | 162.00±09.62     | 158.33±14.04     | 161.17±14.38     | 161.33±17.15     | 06.57↓                         |

Mean±SEM, NC: Normal control, DC: Diabetic control, SBM: *Shadguna Balijarita Makaradhwaja* control, RS: Reference standard control, STZ: Streptozotocin, ↑: Increase ↓: Decrease, SEM: Standard error of mean
diabetic rats. The treatment with SBM showed the normal histopathology of the kidney without any inflammatory vessels and fat deposition [Figure 2a-d]. Decreased Islets of Langerhans, smashed size of β cells and extensive necrosis, fibrosis and atrophy were observed in the pancreas of diabetic rats. STZ induced diabetic rat treated with SBM and glibenclamide restored the necrotic and fibrotic changes and up to moderate levels, raised the number of β cells. [Figure 3a-d].

**DISCUSSION**

Excessive breakdown of tissue proteins decreases body weight in diabetes. It was observed in DC rats in comparison to normal rats [19-21]. Treatment with SBM improved body weight to a certain extent, indicating that control over muscle wasting resulted from glycemic control.

Previous studies claimed that STZ destructs beta cells, which leads to cells less active that makes poor utilization by tissues [22,23]. This suggests that test drug may possess as insulin-like effect on peripheral tissues by inhibiting hepatic gluconeogenesis by promoting glucose uptake or metabolism [24]. Or it might be possible that test drug increases absorption of glucose into the muscles and adipose tissues [25] by stimulation of regeneration process and revitalization of remaining beta cells [26]. Hence, the hypoglycemic activity of SBM may be due to its protective action against damaged pancreatic beta cells and possibly because of increased insulin release or secretion or regeneration of damaged beta cell. Makaradhwaja is a well-known therapeutic medicine of rejuvenation in Indian system of medicine. Its immune-modulatory action was also established [5]. The observed effect may be attributed to the rejuvenation property of Makaradhwaja. The previous study supports its action too [27]. Moderate but insignificant decrease in blood sugar levels were also observed in test drug-treated animals.

Higher levels of HbA1C were observed in the diabetic rats compared with those in normal rats, it might be due to poor glycemic control. SBM treated diabetic rats significantly decreased the level of HbA1C, may be glucose metabolism was improved. This action represents that SBM has an ability to prevent the development of diabetes associated complications.

Elevated levels of SGPT indicating impaired liver function due to hepato-cellular necrosis. Due to elevated transaminase activities leads to diabetic complications like increased ketogenesis and glucogenesis [28], test drug significantly restored this parameter toward normal levels.

Table 2: The effect of test drugs on weight of liver and kidney in streptozotocin induced diabetic Wistar strain albino rats

| Organ      | Kidney (g/100 g) | Liver (g/100 g) | % change in comparison to NC | % change in comparison to DC | % change in comparison to RS |
|------------|------------------|-----------------|-----------------------------|-----------------------------|-----------------------------|
| NC         | 0.65±0.05        | 2.38±0.08       | 52.31↑                      | 46.64↑                      | 4.01↑                       |
| DC (STZ)   | 0.99±0.07        | 3.49±0.19       | 3.49↓                       | 2.38↓                       | 2.38↓                       |
| SBM        | 1.06±0.08        | 3.63±0.08       | 92.02↑                      | 0.02↑                       | 2.38↑                       |
| RS         | 0.92±0.08        | 3.36±0.17       | 2.38↑                       | 0.02↑                       | 2.38↑                       |

Mean±SEM, **P<0.01, ***P<0.001 (comparison to normal control group, unpaired t-test) NC: Normal control, DC: Diabetic control, SBM: Shadguna Balijarita Makaradhwaja Control, RS: Reference standard control, SEM: Standard error mean, ↑: Increase ↓: Decrease

Table 3: The effect of test drugs on blood sugar level in STZ induced diabetic rats at various intervals

| Group      | Blood sugar level (mg/dl) | 0 day | 5th day | 10th day | 15th day | 21st day | % change in comparison to 0 day |
|------------|---------------------------|-------|---------|----------|----------|----------|--------------------------------|
| DC (STZ)   | 383.67±51.44              | 380.33±28.36 | 355.33±28.36 | 372.83±34.89 | 328.17±30.67 | 14.67↓ |
| SBM        | 500.83±42.61              | 503.67±31.81 | 455.67±35.60 | 486.67±51.93 | 279.11±57.95 | 44.27↑ |
| RS         | 512.83±21.43              | 374.33±22.16 | 334.83±32.03 | 314.83±29.68 | 287.00±33.78 | 44.04↑ |

Mean±SEM, *P<0.05, **P<0.01, ***P<0.001 (comparison to diabetic control, unpaired t-test). NC: Normal control, DC: Diabetic control, SBM: Shadguna Balijarita Makaradhwaja control, RS: Reference standard control, SEM: Standard error mean, STZ: Streptozotocin, ↑: Increase ↓: Decrease
Table 5: The effect of test drugs on various serum biochemical parameters

| Parameters               | NC         | DC (STZ)    | SBM       | RS         |
|--------------------------|------------|-------------|-----------|------------|
| Blood sugar (mg/dl)      | 117.50±2.50| 321.00±27.80 | 261.22±60.01 | 286.33±29.52 |
| Cholesterol (mg/dl)      | 57.17±4.72 | 62.00±4.03  | 45.67±4.46* | 58.83±5.87  |
| Triglyceride (mg/dl)     | 64.33±7.28 | 80.83±4.10  | 67.50±6.86  | 70.83±6.02  |
| HDL (mg/dl)              | 25.00±3.22 | 34.17±2.60  | 29.50±3.77  | 28.33±3.26  |
| Blood urea (mg/dl)       | 90.33±4.92 | 137.17±11.66 | 57.00±5.49***| 100.50±10.43*|
| Creatinine (mg/dl)       | 0.58±0.03  | 0.72±0.05*  | 0.65±0.03   | 0.75±0.07   |
| SGPT (IU/L)              | 75.50±8.60 | 329.00±21.29 | 88.83±10.37***| 90.33±3.10***|
| SGOT (IU/L)              | 223.00±12.84| 120.83±11.67 | 173.50±9.16**| 304.67±20.66***|
| Total protein (g/dl)     | 7.35±0.11  | 6.70±0.23*  | 6.70±0.28   | 6.08±0.31   |
| Albumin (g/dl)           | 3.38±0.11  | 3.08±0.09   | 3.45±0.07** | 3.00±0.18   |
| Globulin (g/dl)          | 3.93±0.16  | 3.67±0.25   | 3.25±0.25   | 3.13±0.27   |

Data: Mean±SEM, *P<0.05, **P<0.01, ***P<0.001, (comparison to normal control group, unpaired t-test) *P<0.05, **P<0.01, ***P<0.001 (comparison to DC group, unpaired t-test). NC: Normal control, DC: Diabetic control, SBM: Shadguna Balijarita Makaradhwaja control, RS: Reference standard control, SEM: Standard error of mean, HDL: High-density lipoprotein

Figure 3: (a) Normal cytoarchitecture in normal control group, (b) Photomicrographs of representative section of Pancreas. Marked degeneration of Islets of Langerhans and degranulation of streptozotocin control treated diabetic rats (1×400 magnification), (c) SBM treated rats shows comparatively less degeneration of Islets of Langerhans with intact granules in comparison with streptozotocin control group diabetic rats (1×400 magnification), and (d) Glibenclamide treated rat showed normal cyto-architecture with intact granules in comparison with streptozotocin control group diabetic rats (1×400 magnification)

Elevation of urea and creatinine levels results due to renal dysfunction caused by free radical generation mediated stress in diabetes, persistent hyperglycemia, and hemo-dynamic changes within the kidney tissue [29-31]. Administration of SBM and glibenclamide to the diabetic rats significantly reduced the creatinine and urea levels, which represent the preventive action of SBM on kidney damages in diabetic condition perhaps due to the anti-oxidant properties. Blood urea level was significantly increased in diabetic rats compared to normal rats due to excessive breakdown of protein. This elevation was significantly decreased by test drug.

STZ induced diabetic rat’s increases the level of lipid peroxidation, as an indirect evidence of production of free radical [32]. Hyperlipidemia commonly associated with diabetes [33]. Test drug significantly attenuated serum lipid profiles in diabetic rats. STZ induced diabetic groups treated with glibenclamide and SBM brought back the increased level of total cholesterol and triglyceride near to the normal levels.

Histopathological changes in liver, kidney and pancreas were well restored by the test drug in comparison with DC group. The observed results show anti-diabetic potential of SBM. Observed results may be due to synergistic action of Makaradhwaja with adjuvant T. cordifolia and honey.

CONCLUSION

The present study demonstrates that SBM and dried extract of T. cordifolia with honey significantly reduces the blood glucose level and shows anti-diabetic effect. Restoration histopathological changes in different organs support safe and effective anti-diabetic action of test drug. A significant decrease in glycedated hemoglobin shows effect of Makaradhwaja on diabetes-related complications.

REFERENCES

1. Sen G. In: Mishra S, editor. Bhaishajya Ratnavali. Varanasi: Chaukhamba Surabharti Prakashan; 2007. p. 194.
2. Sen G. In: Mishra S, editor. Bhaishiyaratnavali. Varanasi: Chaukhamba Surabharti Prakashan; 2007. p. 1135.
3. Khedekar S, Prajapati PK. Standard manufacturing process of Shadguna Balijarita Makaradhwaja. Ayu 2014;35:428-32.
4. Prajapati PK, Joshi D, Dubey GP, Kumar M, Prakash B. Pharmaceutical and Experimental Study on Makaradhwaja. MD Dissertation. Varanasi: BHU; 1994.
5. Patgiri BJ, Prajapati PK. A Pharmaceutical and Toxicity Study of Makaradhwaja Prepared by Astasmsakarita Parada. PhD Thesis. IPGT & RA, Jamnagar: Gujarat Ayurved University; 2005.
6. Khedekar S, Patgiri BJ, Ravishankar B, Prajapati PK. A pharmaceutico-pharmacoclinical study of Makaradhwaja Prepared by Swarna Patra-Varkha and Bhasma war to Madhumeha (Diabetes Mellitus). MD Dissertation. IPGT & RA. Jamnagar: Gujarat Ayurved University; 2009.
7. Bhavaprakash. In: Vaidya L, editor. Bhavaprakash Samhita. Part I. New Delhi: Motilal Banarasidas; 1986. p. 1135.
8. Saha S, Ghosh S. Tinospora cordifolia : One plant, many roles. Anc Sci Life 2012;31:151-9.
9. Bhavaprakash. In: Vaidya L, editor. Bhavaprakash Samhita. Part I. New Delhi: Motilal Banarasidas; 1986. p. 329.
10. Erejuwa OO, Sulaiman SA, Ab Wahab MS. Honey-A novel anti-diabetic agent. Int J Biol Sci 2012;8:913-34.
11. Prabhakar C, In: Rasachikitsa. Varanasi: Chowkhamba Vidya Bhawan; 1956. p. 356.
Khedekar, et al.: Anti-diabetic activity of Shadguna Balijarita Maka

12. Bhatta K. In: Bhatta RK, editor. Siddhabheshajamanimala. Varanasi: Chaukhambha Krishnadas Academy; 2008. p. 355.
13. Vagbhata. In: Prof. Kulkarni DA, editor. Vidhnyanobodhini Hindi Commentary. Rasaratnasamuccaya. New Delhi: Meherchanda Lachamandasa Publications; 2006. p. 45.
14. Khedekar SB, Prajapati PK. Standard manufacturing procedure of Shadguna Balijarita Makaradhwaja. Ayu 2014;35:428-32.
15. Sharangdhara. In: Parashar R, editor. Sharangadhara Samhita. Nagpur: Baidyanath Ayurveda Bhavan Ltd; 1994. p. 189.
16. Khedekar S, Ravishankar B, Prajapati PK. Role of Gandhaka Jarana in the Preparation of Makaradhwaja. PhD Thesis. IPGT & RA. Jamnagar: Gujarat Ayurved University; 2012.
17. Guide for the Care and Use of Laboratory Animals. National Institute of Health, Offices of Science and Health Reports. DRR/NIH. Bethesda, MD, USA: DHEW Publication No (NIH) 86-23. 1985.
18. Paget GE, Barnes JM. Evaluation of drug activities. In: Lawrence DR, Bacharach AL, editors. Pharmacometrics. Vol. 1. New York: Academic Press New York; 1964. p. 161.
19. Mishra SB, Verma A, Mukerjee A, Vijayakumar M. Anti-hyperglycemic activity of leaves extract of Hyptis suaveolens L. Poit in streptozotocin induced diabetic rats. Asian Pac J Trop Med 2011;4:889-99.
20. Sajeev T, Arunachalam K, Parimalazhahan T. Antioxidant and antipyretic studies on Pothos scandens L. Asian Pac J Trop Med 2011;4:889-99.
21. Salahuddin M, Jalalpure SS. Antidiabetic activity of aqueous fruit extract of Cucumis trigonus Roxb. in streptozotocin-induced-diabetic rats. J Ethnopharmacol 2010;127:565-7.
22. Junod A, Lambert AE, Stafffacher W, Renold AE. Diabetogenic action of streptozotocin: Relationship of dose to metabolic response. J Clin Invest 1969;48:2129-39.
23. Netchiporouk LI, Shram NF, Jaffrezic-Renault N, Martelet C, Cespuglio R. In vivo brain glucose measurements: Differential normal pulse voltammetry with enzyme-Modified carbon fiber microelectrodes. Anal Chem 1996;68:4358-64.
24. Gray AM, Wahab A, Flatt PR. The traditional plant treatment, Sambucus nigra (Elder), exhibits insulin-like and insulin releasing actions in vitro. J Nutr 2000;130:15-20.