Effects of Camel’s Milk as a Vehicle for Insulin on Glycaemic Control and Lipid Profile in Type 1 Diabetics

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Abstract: Problem statement: To evaluate effects of oral insulin carried by camel’s milk on type 1 diabetes patient in comparison with both insulin injection requirements and camel’s milk alone.

Approach: Total of fifty randomly selected type 1 diabetic subjects were divided into three groups; A, B and C (n = 15). All subjects received a usual care (diet, exercise and insulin injection) but group B subjects in addition received 500 mL day$^{-1}$ of camel’s milk, while group C their subjects in addition received insulin mixed with 500 ml day$^{-1}$ of camel’s milk. Blood samples were taken at beginning and at end of treatment to measure lipid profile and HbA$_1c$ % while blood glucose measured weekly.

Results: Data shown significant improvements in FBG (205.3±2.16-47.26±1.89), PBG (339.86±3.18-161.26±2.89), HbA$_1c$ % (9.9±0.25±5.62±0.21) and significant reduction in insulin requirement (59.26±0.7-20±0.35) in group C receiving insulin mixed with camel’s milk in comparison with groups A and B.

Conclusion: Camel’s milk mixed with insulin was indicated to be an effective supplementation, as adjunctive therapy in management of type 1 diabetes than camel milk or insulin injection.

Key words: Camel’s milk, oral insulin, body mass index, diabetes type 1, anti-coagulum

INTRODUCTION

Type 1 diabetes mellitus is a common organ specific auto-immune and endocrine disorder, characterized by chronic hyperglycaemia and disturbances of carbohydrates, fat and protein metabolism associated with insulin deficiency. Prevention and early treatment is important because diabetes interrupts normal developments in children and carries the threat of severe complication in more active period of life (Singh and Palmer, 2005). Its primary treatment is insulin injection replacement, however at present neither entire physiological insulin replacement could be achieved in clinical practice nor could metabolic disturbances be normalized. Insulin therapy is still the best treatment, however, among certain social groups needle phobia, cost and routine treatment forces these patients to adopt alternative treatments in the context of traditional ethno-medical practices.

Oral insulin therapy has been known for many years but the important drawback is its coagulum formation in acidic environment such as the stomach, thereby neutralising its potency. The potential benefits of oral delivery of insulin include control of plasma glucose levels without peripheral hyper-insulinaemia and restoration of the physiological pathway of endogenous insulin, this result in preservation of the counter-regulatory responses to hypoglycaemia, with a concomitant reduction in hypoglycaemic events.

It has been observed that oral administration of insulin initiated at clinical onset of type 1 diabetes did not prevent the deterioration of beta cell function (Chaillous et al., 2000). Pozzilli et al. (2000) in recent-onset Type I diabetes (IMDIAB VII) study indicates that addition of 5 mg of oral insulin does not modify the course of the disease in the first year after diagnosis and probably does not statistically affect the humoral immune response against insulin (Pozzilli et al., 2000).
It is claimed that the value of camel’s milk is to be found in the high concentrations of volatile acids especially, linoleum acid and polyunsaturated acids, which are essential for human nutrition. El-Agamy et al. (1996) found good amount of lysozyme, lactoferrin, lactoperoxidase, immunoglobulin G and secretory immunoglobulin a in camel’s milk. The milk protein, lactoferrin, has some anti-viral and anti-bacterial properties. Camel’s milk was therapeutically used against dropsy, Jaundice, problems of the spleen, tuberculosis (Akhoundov et al., 1972; Urazakov and Bainazarov, 1974). Camel’s milk works as a laxative on people unaccustomed to drinking this milk (Rao et al., 1970), other lung ailments (Gast et al., 1969) and hepatitis (Sharmanov and Kadyrova, 1978). Several studies have been carried out at the Zayed Complex; UAE showed that camel milk has the IgA and IgG that have proved effective against several viral and bacterial pathogens (Amir, 2003). The milk also has slimming properties (Yasin and Wahid, 1957).

We have heard of many folkloric stories which describe the use of camel’s milk in the treatment of type-1 diabetes mellitus. There is also an account in memories of Emperor Jahangir (1579-1627 AD) referring to the usefulness and acceptability of camel milk (Rogers, 1989). Camel’s milk is used traditionally to treat diabetes (Hamers-Casterman et al., 1993; Hull, 2004). Breitling (2002) and believed that camel’s milk had an anti-diabetic activity possibly because of insulin-like activity, regulatory and immuno modulators effect on beta cells. One of the camel milk proteins has been reported to have similar characteristics to insulin (Beg et al., 1986a; 1986b).

It is important to note that a certain level of scientific testing on camel’s milk has already been attempted and documented, particularly, insulin levels in camel milk and this scientific wisdom can be remarkable achievement for diabetic patients while there is no any attempting to assess anti-coagulum effect of camel milk on oral insulin in treatment of diabetes mellitus type 1. Depending on above literatures, we aim in the current study to evaluate the effects of raw camel’s milk mixed with insulin orally on glycaemia control, lipid profile parameters and insulin requirements in type 1 diabetic subjects in comparison to treatments with camel’s milk alone or insulin replacement.

MATERIALS AND METHODS

Materials:
Milk sample source, collection and preservation: Milk samples were obtained from multifarious lactating camels (Camelus dromedarius) of the Experimental Station of Al-Mahra in Yemen republic. Samples were collected from bulk milk of single Naga from early morning milking in dry sterile screw clean bottles (250 mL) after thoroughly mixing and immediately labelled, cooled to 4°C and transferred in an icebox to the laboratory within 24 h. To prevent microbial growth 10 mg of 2-bromo-2-nitro-1, 3 propanediol (Bronopol Tablets, preservative systems Ltd from Chemgo Organica, Basel, Switzerland) were added to each 50 mL of camel milk.

Insulin vials: Insulin (Human Mixtard(R) 30/70, 100 IU mL-1) vials were purchased from NOVO NORDISK Denmark. Mixtard(R) 30/70 is a pre-mixed neutral suspension of human insulin consisting of neutral soluble insulin injection (30%) and isophane insulin injection (70%). When shaken gently, the liquid appears white and cloudy. Together, the combination of insulins is called biphasic isophane insulin.

Reagents and chemicals: Kits of TG, Tc, LDLc and HDLc were obtained from Beijing Chengxinde Biochemistry Reagent Company (Beijing, China). 2-bromo-2-nitro-1, 3 propanediol (Bronopol Tablets, preservative systems Ltd) was purchased from Chemgo Organica, Basel, Switzerland.

Equipments: The HbA1c level was measured by HPLC (variant BIO-RAD Company, USA), Centrifuge (HERNIE Z 400, Wehingen), UV/Visible spectrophotometer (Shimadzu), micropipette of different size, deep freeze, Accu-Chek(R) (Roche) and plastic syringe were used.

Patients and treatment schedule: A fifty Yemeni subjects previously diagnosed with type 1 diabetes mellitus since two years were randomly recruited from the outpatient of Diabetes and Endocrine Center at Al-Thawra Hospital in Sana’a Capital of Yemen Republic. Subjects gave written consent before participation in the study and the study protocol approved by local ethical committee. The patients were advised to follow strict diet, exercise and insulin treatment for 1 month. During this period frequent monitoring of blood sugar was done to maintain euglycemia.

After one-month period these patients were randomly divided into three groups. Group A (Control; n = 15), comprise patients whom received usual care i.e., diet, exercise and insulin replacement for three months (age = 21.2 ± 2.4; 10 M/5 F). Group B (n = 15), comprise patients whom received received 500 mL day-1 of camel’s milk (in two equally divided doses) in addition to usual care for 3 months (age = 19.4±1.5; 9 M/6 F), Group C (n = 15), comprise patients whom received 500 mL day-1 (in two equally divided doses) of
camel milk (in which X unit of insulin dissolved) in addition to usual care for 3 months (age = 20.22 ±1.9; 11 M/ 4 F).

In all studied groups, insulin injection doses gradually reduced according to glycaemia control and case improvements. In addition, insulin injection doses in group C reduced according to previously added doses to milk. In groups C insulin added to milk at beginning of each month in gradually increased mixed doses (5, 10 and 20 units). Patients with any acute metabolic complications like hypoglycaemia, ketoacidosis, cardiovascular event, renal or acute infections were excluded from the study. Baseline data of the experimental groups were similar in demographic and clinical variables.

Blood sugar was measured once weekly before- and 2 h after breakfast using one touch profile memory glucometer (Accu-Chek®) which were provided for all patients. At beginning and at end of treatment regimen, blood samples were collected from participants for measuring HbA1c % and serum separation, then serum samples preserved at-20°C until measurement (within 2 weeks) of total cholesterol (Tc), low density lipoprotein cholesterol (LDLc), High Density Lipoprotein cholesterol (HDLc) and triglycerides (TGc). Finally, BMI for all individual before and after treatments was recorded. Diagnosis of diabetes was made if fasting blood sugar was ≥126 mg dL−1 and 2 h blood glucose was ≥200 mg dL−1 or a personal detail of known diabetes with the history of treatment was available.

**Biochemical estimations:** Fasting and postprandial blood glucose level was measured by Accu-Chek Active device as described by Joslin (1971). Body Mass Index (BMI) was measured according to the equation, BMI = weight (kg)/length² (m²). HbA1c % was measured by High Performance Liquid Chromatography (HPLC) using a spherical cation exchange gel. Serum lipids were estimated by colorimetric methods. TGc were estimated using the phosphate oxidase method as described by Trinder (1969), total cholesterol was estimated using the Chod-Pap method as described by Zoppi and Fellini (1976), LDLc was measured as described by Bergmeyer (1983) and HDLc was estimated using the dextran-sulphate Mg(II) method as described by Wieland and Siedel (1981).

**Statistical analysis:** Statistical calculations were performed using SPSS (version 15.0) Computer software. Values before and after treatment within each group was analyzed by using paired student’s t test. Data were prepared as mean ± S.E.M. P value <0.05 was considered statistically significant.

**RESULTS**

Data of the present study were illustrated in Table 1-5 and Fig. 1-5 The present study showed that a significant decrease in FBG, PBG and HbA1c % after treatment in three studied groups after treatment except a non-significant change in HbA1c % of group A as compared with baseline data (Fig. 1; Table 1). Diabetes mellitus is usually complicated with hyperlipoproteinemia. The effects of these treatments on lipid profile in type 1 diabetic subjects were shown significant decrease in TGc, Tc and LDLc after treatment in three studied groups while a significant increase in HDLc of three groups after treatment except in group A as compared with baseline data (Fig. 2; Table 1).

| Table 1: Effects of camel’s milk mixed with insulin (C), camel’s milk alone (B) and insulin injection (A) on biochemical parameters within group before and after treatment of type 1 diabetic patients |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| Variable                        | Group A (control group) | Group B (camel milk group) | Group C (camel milk + insulin roup) |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| FBG (mg/dL)                     | 195.6±2.01      | 173.4±1.66***  | 199.46±4         | 155.13±3.5***   | 205.3±2.16      | 147.2±1.89***   |
| PBG (mg/dL)                     | 339.6±10        | 259.1±8.9***   | 345.6±6.3        | 239.2±5.5***    | 339.8±3.18      | 161.2±2.89***   |
| HbA1c (%)                       | 9.30±0.39       | 9.27±0.36      | 9.7 ± 0.39       | 7.28±0.23***    | 9.9±0.25        | 5.62±0.21***    |
| BMI (Kg/m²)                     | 18.04±0.34      | 17.79±0.27     | 16.78±0.25       | 17.38±0.18**    | 17.68±0.30      | 20.79±0.28***   |
| TG (mg/dL)                      | 193.1±1.7       | 175.7±3.0***   | 184±2.1          | 133.6±4.2***    | 182.8±2.15      | 100.8±2.15***   |
| Tc (mg/dL)                      | 271.8±3.35      | 248.63±3.7***  | 251.8±9.3        | 209.2±3.2***    | 203.6±2.56      | 153.3±1.60***   |
| LDLc (mg/dL)                    | 109.9±2.45      | 102.6±1.51**   | 110±2.9          | 92.4±2.6***     | 103.5±2.91      | 70.6±3.32***    |
| HDLc (mg/dL)                    | 43.1±1.53       | 43.7±1.26      | 44.3±2.0         | 49±1.3**        | 41±1.89         | 48.9±1.22**     |
| Dose of insulin (units/day)     | 50±0.64         | 45.46±0.9***   | 55.1±1.4         | 36.2±1.22***    | 59.2±0.7        | 20±0.35***      |

Each group represents the mean ± S.E.M. (n = 15). Control = insulin (X units/day, S.C. injection); Group B = camel milk (500 mL day−1) + insulin injection (X units/day, S.C. injection); Group C = camel milk (500 mL day−1) + insulin injection (X units day−1, S.C. injection). FBG = fasting blood glucose; PBG = postprandial blood glucose; HbA1c % = glycated haemoglobin; BMI = body mass index; TGs = triglycerides; Tc = total cholesterol; LDLc = low density lipoprotein cholesterol; HDLc = high density lipoprotein cholesterol. **P<0.01, ***p<0.001 when compared with data before treatment.

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Fig. 1: The effects of S.C. insulin injection [X units/day] (a), camel milk [500 mL day$^{-1}$] (b) and camel milk [500 mL day$^{-1}$] in which X units of insulin mixed (c) on FBG, PBG and HbA$_1c$% in type 1 diabetic subjects. Data were shown as mean ± S.E.M. (n = 15), *** p<0.001 when compared with data before treatment. B = before treatment; A = after treatment; FBG = fasting blood glucose; PBG = postprandial blood glucose; HbA$_1c$% = glycoselated haemoglobin. Data had shown a significant decrease in FBG, PBG and HbA$_1c$% after treatment in three studied groups except a non-significant change in HbA$_1c$% of group A when compared with baseline data.

Table 2: Percentage reduction in insulin requirements in control (A) and experiment groups (B and C) with respect to baseline values.

| Treatment | FBG (mg/dL) | PBG (mg/dL) | HbA$_1c$ (%) | Doses of Insulin (units/day) |
|-----------|-------------|-------------|--------------|-----------------------------|
|           | Before T    | After T     | R (%)        | Before T     | After T     | R (%)        | Before T     | After T     | R (%)        | Before T     | After T     | R (%)        |
| Group A   | 195.60      | 173.40      | 11.34        | 339.60       | 299.10      | 11.92        | 9.3          | 9            | 9.27         | 1.27         | 50.00       | 45.46        |
| Group B   | 199.46      | 155.13      | 22.22        | 345.60       | 239.20      | 30.78        | 9.7          | 0            | 7.28         | 24.94        | 55.10       | 36.20        |
| Group C   | 205.30      | 147.27      | 28.27        | 339.86       | 161.26      | 52.55        | 9.90         | 5.62         | 43.23        | 59.26        | 20.00       | 66.25        |

Each group represents the mean ± S.E.M. (n = 15). Group A = insulin (X units day$^{-1}$, S.C. injection); Group B = camel milk (500 mL day$^{-1}$) + insulin injection (X units day$^{-1}$, S.C. injection); Group C = camel milk (500 mL day$^{-1}$) in which X units of insulin mixed + insulin injection (X units/day, S.C. injection). FBG = Fasting Blood Glucose; PBG = Postprandial Blood Glucose; HbA$_1c$% = glycedated haemoglobin; R = reduction.
Fig. 2: The effects of S.C. insulin injection [X units/day] (a), camel milk [500 mL day$^{-1}$] (b) and camel milk [500 mL day$^{-1}$] in which X units of insulin mixed (c) on lipid profile in type 1 diabetic subjects. Data were shown as mean ± S.E.M. (n = 15). **p<0.01, ***p<0.001 when compared with data before treatment. B = before treatment; A = after treatment; TG = triglycerides; TC = total cholesterol; LDLc = low density lipoprotein cholesterol; HDLc = high density lipoprotein cholesterol. Data had shown a significant decrease in TG, TC and LDLc after treatment in three studied groups while a significant increase in HDLc of three groups after treatment except group A when compared with baseline data.

Table 3: Levels of FBG (mg/dL) during 12 week in studied groups with respect to baseline values

| Treatment | 0       | 1       | 2       | 3       | 4       | 5       | 6       | 7       | 8       | 9       | 10      | 11      | 12      |
|-----------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| Group A   | 195.8±2.0 | 191.1±0.3 | 190.4±2.4 | 188.2±2.7 | 187.3±1.0 | 185.2±1.5 | 185.5±2.4 | 186.9±3.7 | 176.6±2.5 | 174.6±1.4 | 173.3±1.6 | 173.4±1.6 |
| Group B   | 199.4±6.0 | 196.2±5.6 | 195.6±2.3 | 190.4±3.3 | 186.2±1.2 | 182.6±0.8 | 177.6±0.7 | 170.6±2.2 | 164.4±6.7 | 159.3±3.4 | 156±2.90 | 156.1±3.50 |
| Group C   | 205.1±7.0 | 200±2.20 | 196±1.50 | 190.2±2.5 | 184±1.50 | 179.4±0.8 | 162±1.10 | 157±2.40 | 153±0.50 | 151±2.10 | 149±0.10 | 147.5±0.30 | 147.2±1.89 |

Each group represents the mean ± S.E.M. (n = 15). Group A = insulin (X units day$^{-1}$, S.C. injection); Group B = camel milk (500 mL day$^{-1}$) + insulin injection (X units day$^{-1}$, S.C. injection); Group C = camel milk (500 mL day$^{-1}$) in which X units of insulin mixed + insulin injection (X units day$^{-1}$, S.C. injection).

Table 4: Levels of PBG (mg/dL) during 12 week in studied groups with respect to baseline values

| Treatment | 0       | 1       | 2       | 3       | 4       | 5       | 6       | 7       | 8       | 9       | 10      | 11      | 12      |
|-----------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| Group A   | 339.6±10.0 | 340±2.2 | 335±7.6 | 330±3.40 | 325±3.8 | 321±1.9 | 327±8.8 | 318±5.5 | 311±3.3 | 309±3.4 | 304±0.7 | 301±0.67 | 299±1.68 |
| Group B   | 345.6±6.30 | 334±2.2 | 324±6.7 | 315±5.90 | 307±1.8 | 292±2.4 | 280±1.93 | 273±3.3 | 264±2.10 | 256±0.6 | 249±2.60 | 244±1.10 | 239±2.50 |
| Group C   | 339.86±3.18 | 332±1.2 | 306±3.5 | 292±2.2 | 291±0.4 | 276±4.5 | 258±1.1 | 233±2.9 | 205±0.30 | 183±2.90 | 173±0.37 | 166±2.10 | 161.26±2.89 |

Each group represents the mean ± S.E.M. (n = 15). Group A = insulin (X units day$^{-1}$, S.C. injection); Group B = camel milk (500 mL day$^{-1}$) + insulin injection (X units day$^{-1}$, S.C. injection); Group C = camel milk (500 mL day$^{-1}$) in which X units of insulin mixed + insulin injection (X units day$^{-1}$, S.C. injection).
Fig. 3: The effects of S.C. insulin injection [X units/day] (a), camel milk [500 mL day$^{-1}$] (b) and camel milk [500 mL day$^{-1}$] in which X units of insulin mixed (c) on BMI and insulin doses in type 1 diabetic subjects. Data were shown as mean ± S.E.M. (n = 15), ** p<0.01, *** p<0.001 when compared with data before treatment. B = before treatment; A = after treatment; BMI = body mass index; Ins D = insulin doses. Data had shown significant increase in BMI after treatment in both B and C studied groups while non significant change in group A when compared with baseline data. In addition, data were shown significant reduction in daily insulin requirements in three studied groups after treatment when compared with baseline values but with different percentages among studied groups.

Table 5: Insulin requirements (units/day) during 12 week in studied groups with respect to baseline values

| Week | B | C | A |
|------|---|---|---|
| 0    | 50±0.640 | 59.26±0.7 | 50±0.640 |
| 1    | 55.1±1.40 | 50.1±0.99 | 55.1±1.40 |
| 2    | 57.2±1.11 | 47.2±1.12 | 57.2±1.11 |
| 3    | 45.1±1.95 | 37.2±0.13 | 45.1±1.95 |
| 4    | 34.7±0.35 | 33±1.40 | 34.7±0.35 |
| 5    | 25.3±0.9 | 25.3±0.9 | 25.3±0.9 |
| 6    | 22.5±0.35 | 22.5±0.35 | 22.5±0.35 |

Each group represents the mean ± S.E.M. (n = 15). Group A = insulin (X units day$^{-1}$), S.C. injection; Group B = camel milk (500 mL day$^{-1}$) + insulin injection (X units day$^{-1}$), S.C. injection; Group C = camel milk (500 mL day$^{-1}$) in which X units of insulin mixed + insulin injection (X units day$^{-1}$), S.C. injection.
Fig. 4: The effects of S.C. insulin injection [X units/day] (a), camel milk [500 mL day$^{-1}$] (b) and camel milk [500 mL day$^{-1}$] in which X units of insulin mixed (c) on FBG during 12 week of treatment of type 1 diabetic subjects. Data were shown as mean ± S.E.M. (n = 15). w = week; FBG = fasting blood glucose. Data had shown significant decrease in FBG after treatment in three studied groups but in both B and C studied groups than in group A and in group C than B when compared with baseline data.

Fig. 5: The effects of S.C. insulin injection [X units/day] (a), camel milk [500 mL day$^{-1}$] (b) and camel milk [500 mL day$^{-1}$] in which X units of insulin mixed (c) on insulin requirements during 12 week of treatment in type 1 diabetic subjects. Data were shown as mean ± S.E.M. (n = 15). w = week. Data had shown significant decrease in insulin doses after treatment in three studied groups but in both B and C studied groups than in group A and in group C than B when compared with baseline data.
In addition, data had shown significant increase in BMI after treatment in both B and C studied groups while non significant change in group A as compared with baseline data. In contrast, data were shown a significant reduction in daily insulin requirements in three studied groups after treatment as compared with baseline values but with different mode among studied groups (Fig. 3; Table 1). Data had shown significant decrease in FBG, PBG and insulin doses after treatment in three studied groups but in both B and C studied groups than in group A and in group C than B during 12 week of treatment as compared with baseline data (Fig. 4 and 5; Table 2-5). The acceptability of camel’s milk was fairly good and only two patients complained of flatulence which disappeared after 3 days. Nausea, flatulence and mild diarrhoea (2-4 semisolid) was reported by four patients who also subsided spontaneously. Five patients were excluded from the current study due to three of them have chronic diseases (renal failure and heart disease) and two of them hate the odour of mixed insulin with milk. No severe hypoglycaemic event or diabetic ketoacidosis were reported in all groups. Out of 45 patients, only ten patients in group A, five in group B and one patient in group C required the same doses of insulin and the other 29 patients had lower requirement.

**DISCUSSION**

The present study intended to explore the grounds on which camel’s milk is based as traditional use in achieving glycaemia control in type-1 diabetes patients, in addition; to assess the role of camel milk as a natural vehicle for carrying and protecting insulin hormone from acidity and digestion in stomach.

In the current study, significant improvements were observed in all studied biochemical parameter after 3 month of treatment in all studied groups. However, these improvements were clear and significant in group C than other studied groups; particularly in reduction of blood glucose and insulin requirements to obtain glycaemia control along with significant improvement in HbA1c, % level at the end of three months.

Requirements of mean doses of insulin day\(^{-1}\) before treatment in patients of group C was 59.26±0.7. It was rapidly declined during the early experimental stages then gradually reached a mean value of 20±0.35, (p<0.001). There appears to be a connection between camel’s milk and diabetes; it is common practice in Africa, Asia and the Middle East to drink camel’s milk for treating diabetes mellitus, without defining which type there is (Yagil et al., 1994). The results of the present study in the same line with repeated trials of Agrawal et al. (2002) whom observed that there was 30-35% reduction in daily doses of insulin in patients of type 1 diabetes receiving raw camel’s milk. They stated that camel milk had shown to possess insulin like activities, which decrease the requirement of exogenous insulin in type 1 diabetes patients (Agrawal et al., 2002; 2003; 2005).

Camel’s milk dyes have special properties not found in milk of other mammals (Yagil, 1987), which favours it to treats diabetics. It contains large concentrations (52 units/liter) of insulin and used traditionally to treat diabetes (Hamers-Casterman et al., 1993; Yagil et al., 1994; Singh, 2001) thereby it may be the reason for lesser requirement of insulin in groups B and C than group A. In addition, Beg et al. (1986a; 1986b) found that amino acid sequence of some of the camel’s milk protein is rich in half cystine, which has superficial similarity with insulin family of peptides. Moreover, Breitling (2002) suggested that camel milk is having anti diabetic activity possibly because of: insulin like activity, regulatory and immuno modulatory functions on β cells. El-Agamy et al. (1996) found good amount of lactoferrin, in camel milk. The milk protein, lactoferrin, which known to enhance the immune response and is present in large quantities in camel milk (ten times higher than in cow milk) does have some anti-viral and anti-bacterial properties. Insulin is a metallic hormone contains Zn, in this point; Saitmuratova et al. (2001) stated that the proteins of camel’s milk and shubat were studied by electrophoresis. It was found that Fe and Zn occur in greater quantities in both camel’s milk and shubat than in cow’s milk. Then camel’s milk contains insulin like protein with its Zn.

There is strong evidence that oral insulin products would provide insulin in a more physiological manner, resulting in a decrease in peripheral insulin concentrations this “insulining” the live (Gwinup et al., 1991; Hoffam and Ziv, 1997). Scientists have developed hexylinulin monoconjugate 2(HIM2), in which a single amphiphilic oligmer is covalently linked to the free amino group on the lys-β 29 residue of recombinant human insulin via an amide bond (Still, 2002). HIM2 alterations in physio-chemical are a characteristic which resists the enzymatic degradation and facilitates absorption. A similar trial in France, using two doses of oral insulin (2.5 and 7.5 mg day\(^{-1}\)) also did not show any positive effect (Chaillous et al., 1999).

Using milk for treatment is itself a point of contention as insulin, being a protein, is normally destroyed in the stomach (Arthur et al., 2001). However, the concentration of insulin in human milk is also significantly higher (60.23±41.05 micro unit/mL).
whereas it is low in cow milk (16.32±5.98 micro unit/mL) (Shehadeh et al., 2001) but probably because of coagulation in stomach it is not available for absorption in the intestine. Intestinal absorption of large proteins and, therefore, insulin is only possible in the first few days of life (Lawrence and Lawrence, 2010). After this period proteins are destroyed in the stomach by acid and pepsin, especially in milk, which forms a coagulum in the stomach. However, among the pertinent properties is a high acid content ascorbic acid (Farah et al., 1992) and does not react to the acid and not form coagulum in acidic environment (Abu-Lehia, 1989; Wangoh, 1993) and so will pass into the intestines ready for absorption of insulin, so the therapeutic efficacy of camel’s milk may be due to lack of coagulum formation of camel’s milk in acidic media and antibacterial and antiviral properties (El-Agamy et al., 1992).

Even if camel milk passes into the small intestine without forming cheese it is not known how insulin, a protein, is absorbed by humans with diabetes mellitus. The validity of this hypothesis was proven in the current study by decline in blood glucose after treatment in groups C and B than A. This decline validates the observation that camel milk acts on regulating cellular sugar in the absence of endogenous insulin and in presence of exogenous oral insulin.

Fat dispersed as small micelles in the camel’s milk instead of a layer of fat in other types of milk so non-reaction to acid (Shamsia, 2009). Hence we predict that micelles in camel’s milk could be take the insulin present in the milk within it and protect it from coagulation by acidity or digestion by pepsin in the stomach to be absorbed as it in the micelles or release it again in small intestine for absorption.

Diabetes mellitus is usually complicated with hyperlipoproteinemia in particular hypercholesterolemia. The effects of treatments in the current study on lipid profile in type 1 diabetic subjects were shown significant decrease in TG, Tc and LDLc after treatment in three studied groups but with different percentage of reduction. In contrast, a significant increase in HDLc of three groups after treatment was observed except in group a when compared with baseline data.

These results in agreement with Saitmuratova et al. (2001) and Hamers-Casterman et al. (1993) whom found that the camel’s milk, lower in fat, lactose and contains no cholesterol and more nutritious. It has been reported that camel’s milk contain high levels of vitamins C (three times that of cow milk and one-and-a-half times that of human milk), A, B2, E and very rich in potassium, iron, magnesium and other trace elements (Pozzilli et al., 2000; Gast et al., 1969; Al-Hashem, 2009). In addition, Gast et al. (1969) claim that the value of camel’s milk is to be found in the high concentrations of volatile acids, especially, linoleum acid and the polyunsaturated. These vitamins and polyunsaturated are antioxidants thereby removing free radicals and useful in preventing the tissues injury including β-cells of pancreas. High concentrations of antioxidants and removal of fat from the body may make the insulin receptors more responsive to available insulin. Moreover, these free radicals have a role in intensifying oxidized-LDLc and exhausting HDLc.

We observed a significant increase in mean BMI of group B and C (16.78±0.25-17.38±0.18 and 17.68±0.30-20.79±0.28 respectively) after three months of camel milk mixed with insulin suspension or camel milk alone treatment, while non-significant decrease in BMI in group A. This positive effect in weight gain and stability in lipid profile of the experimental group (B and C) may be attributed to the good nutritional value and relatively lower fat percent and extremely low cholesterol or anabolic effect of camel milk. However, camel milk is most beneficial for human nutrition (Agrawal et al., 2007; Konuspayeva et al., 2007; El-Fakharany et al., 2008); in particular the hypoallergenic properties of its protein fraction suggest that this product is more similar to human than cows’ milk (El-Agamy et al., 2009). Nutrition plays a vital role in the health of a community especially in malnutrition. Malnourished populations are more susceptible to most infections and severity of illness worsening the mortality rates (WHO, 2001).

CONCLUSION

We conclude that, the data of the current study shows significant hypoglycaemic effect of camel’s milk alone or camel’s milk carrying oral insulin when given as an adjunctive therapy. The hypoglycaemic effect of camel milk may be due to: (a) presence of insulin/insulin like protein with Zn where insulin like growth factor system in the bovine mammary gland has insulin receptors and binding proteins, (b) camel’s milk lactoferrin and immunoglobulins, of relatively small size and weight, might offer an interplay with host cell protein leading to an induction of regulatory cells and finally leading to a downward regulation of immune system and β-cell salvage and (c) fat dispersed as small micelles in the camel’s milk instead of a layer of fat in others milk, anti-coagulum (non-reaction to acid) effect and possible anti-pepsin effect of camel’s milk make it a suitable carrier, vehicle and protectant to insulin/exogenous insulin/or insulin like proteins to pass into small intestine and then absorbed.
However, more research is necessary and further studies are warranted to fractionate the active principle of camel's milk and to find out its exact mode of action as a suitable vehicle for oral insulin before a final conclusion can be made.

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