The potential usefulness of taurine on diabetes mellitus and its complications

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Abstract Taurine (2-aminoethanesulfonic acid) is a free amino acid found ubiquitously in millimolar concentrations in all mammalian tissues. Taurine exerts a variety of biological actions, including antioxidation, modulation of ion movement, osmoregulation, modulation of neurotransmitters, and conjugation of bile acids, which may maintain physiological homeostasis. Recently, data is accumulating that show the effectiveness of taurine against diabetes mellitus, insulin resistance and its complications, including retinopathy, nephropathy, neuropathy, atherosclerosis and cardiomyopathy, independent of hypoglycemic effect in several animal models. The useful effects appear due to the multiple actions of taurine on cellular functions. This review summarizes the beneficial effects of taurine supplementation on diabetes mellitus and the molecular mechanisms underlying its effectiveness.

Keywords Taurine · Diabetic mellitus · Diabetic complications · Obesity · Insulin resistance

Introduction

Taurine (2-aminoethanesulfonic acid) is widely distributed and is found in millimolar concentration in mammalian tissues. Many evidences support that taurine is a cytoprotective agent in a variety of tissues. Taurine modulates a variety of cellular functions, including antioxidation, modulation of ion movement, osmoregulation, modulation of neurotransmitters and conjugation of bile acids etc. (Huxtable 1992; Satoh 1998; Schaffer et al. 2000; Sjovall 1959). The source of taurine in body is biosynthesis and dietary intake. Taurine is synthesized from methionine and cysteine mainly in the liver. It is well-known that biosynthetic capacity of taurine is very low in human and is absent in cats, while rodents have high synthetic capacity (Hansen 2001). On the other hand, dietary taurine is ingested from meat and sea food. Especially, sea food is rich in taurine. Yamori et al. (2001) demonstrated that urinary taurine excretion as a marker of taurine intake inversely correlated with mortality rate caused by ischemic heart diseases in world wide epidemiological study. Moreover, taurine depletion by taurine-deficient diet in cats causes various pathological conditions, including retinal degeneration, reproductive failure and dilated cardiomyopathy (Hayes et al. 1975; Pion et al. 1987; Sturman 1991). Therefore, taurine seems an essential nutrient and its deficiency may cause various tissue disorders in human. Moreover, treatment of taurine benefits many kinds of pathologies. The accumulating data show the effectiveness of taurine supplementation against both insulin dependent, non-insulin dependent diabetes mellitus and insulin resistance (Franconi et al. 2004, 2006; Hansen 2001; Schaffer et al. 2009). In addition, taurine supplementation is beneficial to diabetic complications, including retinopathy, nephropathy, neuropathy, atherosclerosis and cardiomyopathy. These useful effects appear due to the multiple actions of taurine on cellular functions. This review summarizes the beneficial effects of taurine supplementation on diabetes and the molecular mechanisms underlying the effectiveness.
The effect of taurine on hyperglycemia in diabetic animal models and its potential mechanisms

The effect of taurine on type 1 diabetic models

The effect of taurine administration for type 1 diabetes has been well investigated. Treatment of taurine before diabetic onset suppressed hyperglycemia and lowered plasma glycated hemoglobin, cholesterol and triglyceride in STZ-induced type 1 diabetic rats (Alvarado-Vasquez et al. 2003; Tokunaga et al. 1979, 1983). Moreover, taurine reduced plasma lipid peroxidation products induced in type 1 diabetes mellitus. The prevention of hyperglycemia by taurine was also reported in alloxan-induced type 1 diabetic rabbits (Tenner et al. 2003; Winiarska et al. 2009). Importantly, it has been reported that treatment of taurine started from the time-point of diabetic onset failed to improve hyperglycemia in type 1 diabetic animals (Goodman and Shihabi 1990), indicating that lowering effect of taurine on blood glucose level in type 1 models may be due to the protection of beta cells from STZ or alloxan (Chang and Kwon 2000; Gavrovskaya et al. 2008). Interestingly, taurine supplementation from 2 days later of STZ injection prolonged survival in diabetic rats (Di Leo et al. 2004). This observation indicates that taurine may confer the resistance against some stresses induced by hyperglycemia, which may associate with the beneficial role against the complications, as described below.

Non-obese diabetic (NOD) mice genetically develop autoimmune diabetes caused by infiltration of the pancreatic islets by mononuclear leucocytes. Taurine supplementation starting before birth (pregnant mice were received a diet supplemented with taurine) until weaning significantly increased in pancreatic islet mass in NOD mice (Arany et al. 2004), suggesting that taurine alters islet development. Moreover, taurine treatment delayed the onset time of diabetes, and 20% of treated female mice remained free of diabetes.

The effect of taurine on obese-induced diabetic models

The Otsuka Long-Evans Tokushima Fatty (OLETF) rat is a model of non-insulin dependent diabetes, which exhibits hyperglycemia and insulin resistance and has accumulated abdominal fat as compared to the normal rats. It has been demonstrated that taurine supplementation improved hyperglycemia and insulin resistance in OLETF rats (Harada et al. 2004; Nakaya et al. 2000). Taurine also suppressed the increase in serum triglyceride and cholesterol, but not body weight and abdominal fat mass. Whereas it was expected that taurine could enhance the calorie consumption and/or lipid oxidation, taurine supplementation did not increase but decrease the energy expenditure and did not alter the lipid oxidation in OLETF rats (Harada et al. 2004). Therefore, the other pathways may underlie. Concerning with body weight and fat mass, taurine supplementation prevented the high fat diet-induced increase in body weight as well as fat mass in high fat diet-induced obese mice (Tsuboyama-Kasaoka et al. 2006). Then, the effect of taurine on obesity may differ dependent on animal species and/or experimental procedures.

The effect of taurine on fructose-fed rat model

High fructose diet impairs glucose tolerance and insulin sensitivity, taurine supplementation suppressed hyperglycemia and insulin resistance in high fructose-fed rat model (Nandhini et al. 2004, 2005; El Mesallamy et al. 2010). Furthermore, taurine supplementation to high fructose-fed rats decreased in the biomarkers of oxidative stress, such as lipid peroxidation and conjugated dines. Additionally, while taurine improved urinary kallikrein activity in fructose-fed rats, the effect of taurine supplementation was prevented by co-administration of Hoe 140, a kinin B2 receptor antagonist (Nandhini and Anuradha 2002, 2004). Since kinins influence insulin release and insulin action, this action may be involved in the molecular mechanism of the effect of taurine against insulin resistance.

The effect of taurine on insulin secretion and insulin sensitivity in acute glucose or lipid infusion models

Prolonged elevation of glucose is associated with insulin resistance. While 6-h infusion of high glucose induced a decrease in insulin-stimulated peripheral glucose uptake, co-infusion of taurine prevented the defect of glucose uptake (Haber et al. 2003). Furthermore, co-infusion of taurine suppressed lipid peroxidation induced by high glucose infusion in soleus muscle, indicating that antioxidative role of taurine in skeletal muscle is involved in the pathway of effectiveness against peripheral insulin resistance. However, taurine failed to prevent a decrease in glucose-stimulated insulin secretion and an increase in reactive oxygen induced by 48-h infusion of high glucose (Tang et al. 2007). On the other hand, some reports demonstrated various protective actions of taurine, such as the modulation of mitochondrial calcium handling and the stabilization of protein folding, against high glucose exposure in the cultured beta cells (Han et al. 2004; Kaniuk et al. 2007).

Meanwhile, the beneficial effect of taurine on islet dysfunction induced by free fatty acid was observed (Oprescu et al. 2007). While 48-h intravenous infusion of oleate decreased the glucose-stimulated insulin secretion, co-infusion of taurine prevented defective insulin secretion in islets induced by oleate. Furthermore, taurine suppressed
oleate-induced ROS production in islets. Moreover, taurine also prevented hepatic insulin resistance induced by intravenous infusion of fatty acids (Wu et al. 2010). This beneficial effect was accompanied by inhibition of fatty acid-induced oxidative stress and JNK1 activation which impairs insulin signaling. These reports indicate the beneficial effect of taurine against lipotoxicity in islets and livers, which in turn may contribute to the prevention of diabetic onset in obesity.

The effect of taurine against diabetic complications and its molecular mechanisms

Alteration of taurine level in diabetes

Plasma and tissue taurine level are known to alter in a variety of pathophysiological conditions. Plasma taurine concentration in patients with insulin-dependent diabetes mellitus (IDDM) was significantly lower than in control subjects (Franconi et al. 1995, 1996). Consistently, the reduction of plasma taurine level has been demonstrated in STZ- or alloxan-induced diabetic animals (Franconi et al. 1996; Trachtman et al. 1995). In case of type2 diabetes, plasma taurine level is lower in the patients than non-diabetic healthy subjects (De Luca et al. 2001; Merheb et al. 2007). Since taurine deficiency associates with dysfunction in various tissues (Hayes et al. 1975; Pion et al. 1987; Sturman 1991), a decrease in taurine level in diabetic subjects may be involved in the diabetic complications.

The role of taurine against the production of advanced glycation end-products and modified LDL

Hyperglycemia accelerates non-enzymatic glycation of protein and causes the accumulation of advanced glycation end products (AGEs). It is known that AGEs play a key role in the development of diabetic complications, such as nephropathy and microvascular diseases. Since taurine has a high reactivity with aldehyde as compared to the other amino acids (Ogasawara et al. 1993), the preventive actions of taurine on the productions of AGEs in diabetes are expected. Indeed, taurine inhibited the AGE formation in vitro (Nandhini et al. 2004; Nandhini and Anuradha 2003; Selvaraj et al. 2006). Consistently, taurine supplementation prevented an increase in the plasma glycated proteins, such as fructosamine and glycated hemoglobin, in high fructose-fed rats (Nandhini et al. 2004).

While the modified LDL also contributes to development of vascular complications, some actions of taurine is likely to associate with prevention of LDL modification. The reactivity of taurine with aldehyde is also likely to contribute to decrease in malondialdehyde-related LDL modification (Ogasawara et al. 1993). Moreover, taurine also has the scavenging action for hypochlorous acid (HClO), while HClO, produced by myeloperoxidase in neutrophils and macrophages, possesses the antimicrobial properties and is also involved in oxidation of LDL (Pennathur and Heinecke 2007). In type 2 diabetic models, high myeloperoxidase activity has been found in the vessels of diabetic obese rats (Zhang et al. 2004), indicating that HClO may contribute to increase in oxidized LDL in diabetes. Moreover, lowering effect of taurine on the production of LDL-cholesterol itself may contribute the reduction of oxidized LDL (Bellentani et al. 1987; Gandhi et al. 1992; Nakamura-Yamanaka et al. 1987; Yokogoshi et al. 1999). The increased serum level of LDL cholesterol in STZ-treated diabetic mice was normalized by the chronic administration of taurine (Mochizuki et al. 1999; Nanami et al. 1996).

Endothelial dysfunction

Most of diabetic complications are associated with vascular disorder. Microangiopathy causes retinopathy, nephropathy and neuropathy, whereas macroangiopathy causes cardiomyopathy and atherosclerosis. In STZ-treated diabetic mice, it has been demonstrated that the chronic taurine supplementation normalized the acetylcholine-induced relaxation of aortic rings, while the vasodilatation capacity is attenuated (Kamata et al. 1996; Wang et al. 2008). Furthermore, pre-incubation of the tissues with taurine for 2-h ex vivo improved both the enhanced response to nor-epinephrine and the attenuated response to acetylcholine in aorta ring from STZ-treated diabetic rats (Abebe 2008). These data illustrate the protective action of taurine on the impaired endothelium-dependent vasodilator response in hyperglycemia.

A variety of molecular mechanisms underlie the beneficial role of taurine against endothelial dysfunction in diabetes mellitus (summarized in Fig. 1). As described above, lowering effect of taurine on AGE and modified LDL may be involved in the molecular pathway of beneficial effect of taurine. Additionally, HClO consumes the nitric oxide and in turn causes vasoconstriction and endothelial dysfunction (Pennathur and Heinecke 2007). Taurine may increase the bioavailability of NO through scavenging HClO. Furthermore, taurine suppressed the expressions of vascular cell adhesion molecule-1 (VCAM-1) and intercellular adhesion molecule-1 (ICAM-1) induced by high glucose in cultured endothelial cells (Ulrich-Merzenich et al. 2007). Consistently, taurine supplementation started after the diabetes onset in STZ-treated diabetic rats prevented the induction of ICAM-1 and lectin-like oxidized LDL receptor-1 (LOX-1), which is responsible for the incorporation of oxidized LDL into cells, in aortas (Wang et al. 2007).
et al. 2008). Furthermore, while acute hyperglycemia induced by intravenous infusion of glucose activated leucocyte adhesion and migration to endothelium and increased in endothelial ICAM-1 and apoptosis in rats, taurine supplementation for 5 days prior to the experiment prevented leucocyte actions and the elevation of ICAM-1 and apoptotic cell death (Casey et al. 2007). Therefore, taurine may prevent the leukocyte-endothelial cell interaction and endothelial apoptosis enhanced by hyperglycemia.

Diabetic nephropathy

It has been reported that taurine supplementation started concurrently with STZ injection reduced albuminuria and diminished glomerulosclerosis and tubulointerstitial fibrosis (Trachtman et al. 1995). More recently, it has been demonstrated that the taurine administration from fourth month later of the induction of diabetes significantly suppressed further increase in urinary protein excretion in diabetic rats, accompanied by the reduction of mesangial extracellular matrix expansion, TGF-β expression and oxidative stress in the renal glomerulus in rats (Higo et al. 2008). In vitro study also demonstrated that taurine treatment suppressed the increases in lipid peroxidation and TGF-β by high glucose in renal proximal tubule cells (Park et al. 2001). Moreover, taurine attenuated impairment of cellular growth and tubule cell hypertrophy induced by high glucose, associated with the suppression of high glucose-induced signal activations including MAPK cascade and STAT3 (Huang et al. 2007). Furthermore, taurine attenuated the induction of cytochrome P450 2E1 which metabolizes a variety of endogenous and exogenous compounds and is a potential source of ROS in kidney of STZ-treated diabetic rats (Yao et al. 2009). Taurine also attenuated cell hypertrophy and fibrosis induced by AGE exposure in renal tubular epithelial cells (Huang et al. 2008). Moreover, it has been reported that taurine suppressed the induction of fibrosis-related genes in AGE-treated renal tubular cells (Huang et al. 2009). Therefore, taurine may prevent renal injury and fibrosis in diabetic animals through suppression of ROS induced by glucose and AGE in kidney.

Diabetic retinopathy

In STZ-induced diabetic model, taurine supplementation after diabetic onset effectively improved the changes in ultrastructure and attenuated induction of glial fibrillary acid protein (GFAP), a marker of gliosis, and apoptosis in retinal glial cells of STZ-treated diabetic rats without the effect on plasma glucose concentration (Yu et al. 2008; Zeng et al. 2009, 2010a), indicating the beneficial role of taurine on diabetic retinopathy. Furthermore, the taurine supplementation in STZ-treated rats significantly decreased in retinal carbonyl dienes (Di Leo et al. 2002, 2003). Additionally, taurine supplementation attenuated the induction of retinal VEGF, which associates with vascularization, in STZ-diabetic rats, suggesting that taurine may normalize the retinal vascular function in diabetes (Obrosova et al. 2001b; Zeng et al. 2009). Furthermore, while elevation of glutamate in retina is associated with the development of diabetic retinopathy, taurine prevented the elevation of retinal glutamate content and reduction of the proteins involved in glutamate uptake and degradation in STZ-treated diabetic rats (Zeng et al. 2009). Consistently, taurine suppressed high glucose-induced defect of glutamate uptake and degradation in cultured Muller cells (Zeng et al. 2010b).

Diabetic cataract

Taurine is known to be the most abundant free amino acid in eye lens, and lens taurine level is decreased in diabetic cataract group in STZ-treated diabetic rats (Anthrayose and Shashidhar 2004; Malone et al. 1990; Vilchis and Salceda...
Chronic taurine supplementation reduced malondialdehyde level in lens of STZ-induced diabetic rats (Obrosova and Stevens 1999). Furthermore, although taurine did not improve opacity of eye lens induced by the exposure to high glucose medium for 6 days in cultured lens, taurine inhibited protein carbonylation induced by high glucose (Son et al. 2007). These studies indicate that taurine protects the lens from oxidative stress induced by hyperglycemia, while the effect of taurine against cataract is arguable.

Diabetic neuropathy

It is well-known that sorbitol accumulation in the nerve is associated with diabetic peripheral neuropathy. When intracellular glucose is increased in hyperglycemic condition, excessive glucose is metabolized to sorbitol through aldose reductase. Organic osmolytes, including sorbitol, taurine and myo-inositol, are regulated in response to the change of extracellular osmolality to maintain the cell volume. Stevens demonstrated that taurine and myo-inositol are decreased in nerve of STZ-treated diabetic rats, whereas administration of aldose reductase inhibitor attenuated the depletion of taurine and myo-inositol (Stevens et al. 1993), suggesting that excessive accumulation of sorbitol led to the depletion of other organic osmolytes. Interestingly, the exposure of cells to high glucose reduces the expression of taurine transporter, whereas the treatment of an aldose reductase inhibitor or an antioxidant with the high glucose reversed the expression of taurine transporter (Askwith et al. 2009; Stevens et al. 1999), suggesting the crucial role of sorbitol in the regulation of intracellular taurine concentration. While diabetic neuropathy is observed in STZ-treated diabetic model, taurine supplementation improved the defective nerve functions, such as nerve conductance deficits and hyperalgesia, and ameliorated the deficit of nerve blood flow (Li et al. 2005b; Obrosova et al. 2001a, b; Pop-Busui et al. 2001; Stevens et al. 1993). Taurine supplementation reduced the oxidative stress in nerves and prevented the impairment of calcium handling in sensory neuron of STZ-treated diabetic rats (Li et al. 2005b). Additionally, a decrease in nerve growth factor (NGF) in STZ-treated rats was prevented by taurine supplementation (Obrosova et al. 2001a). Similarly, usefulness of taurine against diabetic peripheral neuropathy, including deficits of hind limb sciatic motor and digital sensory nerve conduction velocity, nerve blood flow, and sensory thresholds, was observed in Zucker diabetic fatty rats (Li et al. 2006).

Atherosclerosis and thrombosis

Diabetes is a major risk factor of atherosclerosis. While there are several reports on the beneficial effects of taurine against atherosclerosis (Kondo et al. 2000; Murakami et al. 1999a, 1999b, 2002), no evidence has been demonstrated in the diabetic models. Since endothelial dysfunction and accumulation of oxidized LDL in vessel are the critical features of atherosclerosis, various actions of taurine in endothelial cells mentioned above, such as lowering oxidized LDL and anti-apoptosis, can work effectively on atherosclerosis in diabetes.

Moreover, platelet activation and aggregation caused by atherosclerosis are the critical events in thrombosis. It is known that platelet function alters in diabetic subjects, which may sensitize the platelet for stresses to generate thrombi. It has been reported that platelet taurine level as well as plasma taurine level is decreased in both type 1 and type 2 diabetic patients (De Luca et al. 2001; Franconi et al. 1995). Because taurine depletion causes an increase in platelet sensitivity to aggregation (Hayes et al. 1989), it has been hypothesized that taurine supplementation can prevent platelet aggregation in diabetic patients. Franconi et al. (1995) reported that taurine supplementation at 1.5 g/day for 90 days in IDDM patients suppressed platelet aggregation induced by arachidonic acid in the isolated platelet. Additionally, they demonstrated the pre-incubation of taurine inhibits arachidonic acid-induced aggregation in platelet from diabetic patients, but not in platelet from control subjects. Spohr et al. (2005) demonstrated that taurine supplementation at 1.5 g/day for 8 weeks had no effect on ADP-stimulated platelet aggregation in high risk subjects with a positive family history of NIDDM.

Diabetic cardiomyopathy

Unlike the other tissues, cardiac taurine level increases in STZ-treated diabetic models and does not alter in obese-induced diabetic models (Militante et al. 2000). Elevation of cardiac taurine level seems the feature of cardiomyopathy, since its elevation has also been reported in the animal models or patients with congestive heart failure (Huxtable and Bressler 1974). Although the biological significance of the elevation of taurine level in the failing heart is still unveiled, the elevated taurine is expected to modulate glycolytic capacity, such as pyruvate dehydrogenase phosphorylation, in diabetic cardiomyopathy (Militante et al. 2000). Recently, Li et al. (2005a) reported that taurine supplementation started after the development of cardiomyopathy prevented an increase in heart weight and improved the impaired $-dp/dt$ max but not $+dp/dt$ max. Furthermore, taurine supplementation suppressed the reduction of Bcl-2 expression in STZ-treated diabetic cardiomyopathy rats, indicating that anti-apoptotic action of taurine may be involved in the protective effect of taurine against diabetic cardiomyopathy.
Clinical studies

The clinical usefulness of taurine on diabetes has been evaluated in some clinical studies (listed in Table 1). In clinical trial for type 1 diabetic patients, Elizarova and Nedosugova (1996) investigated the effect of taurine on hyperglycemia in type 1 diabetic patients (n = 10) who had been already medicated with insulin. Additive supplementation of taurine (0.5 g twice a day) improved the carbohydrate metabolism and decreased in triglyceride. In the other study, however, Franconi et al. (1995) reported that the administration of taurine (1.5 g per day) for 90 day did not modify glucose metabolism in IDDM patients (n = 39). Clinical usefulness on type 1 diabetes is still arguable.

There are some clinical trials of taurine supplementation on the patients with type 2 diabetes mellitus. Chauncey et al. (2003) tested the hypoglycemic effect of taurine on type2 diabetes patients (n = 32), and they showed that taurine supplementation (3 g per day) for 4 month increased in plasma taurine level but did not change HbA1c level and the plasma lipid peroxide level compared to placebo group. Brons et al. (2004) also tested the effect of taurine in overweight non-diabetic men (n = 20) in crossover study. Although plasma taurine level was elevated after taurine administration (1.5 g per day) for 8 weeks, taurine had no effects on insulin secretion or sensitivity, and on plasma lipid level. These studies concluded that taurine does not influence hyperglycemia and insulin resistance in type 2 diabetic patients, inconsistent with animal studies. However, it should be noted that these clinical studies have some limitations such as other medications, given dose of taurine, duration of trial etc. On the other hand, usefulness of taurine supplementation against the impairment of insulin sensitivity was reported in the crossover clinical study by Xiao et al. (2008). They demonstrated the effect of taurine against chronic elevation of plasma fatty acids induced by the intravenous infusion of Intralipid (20% soybean oil, 1.2% egg phospholipids, 2.25% glycerin in water, heparin) on non-diabetic men who were either overweight or obese (n = 6). While 48-h infusion of intravenous lipid induced insulin resistance, a 2-week pretreatment of taurine (3 g per day) before lipid infusion improved the impaired insulin sensitivity and prevented the rise in lipid peroxidation products in plasma, indicating that oral taurine supplementation ameliorates fatty acid-induced insulin resistance in humans, possibly due to reducing oxidative stress.

Concerning diabetic nephropathy, Nakamura et al. (1999) tested the taurine supplementation (3 g per day) on the patients with microalbuminemia of type 2 diabetes and treatment was continued for 12 months in intergroup trial (n = 10 each group). They demonstrated that taurine supplementation had no benefit against microalbuminemia and the biomarkers for fibrosis, such as serum collagen IV and plasma Matrix metalloproteinase-9.

Meanwhile, Moloney et al. (2010) recently reported the beneficial effect of taurine on endothelial dysfunction in type 1 diabetic patients in crossover study. While arterial stiffness and flow-mediated dilatation of brachial artery,

Table 1 Clinical studies with diabetes and complications

| Article                | Subjects                          | Duration | Dose (/day) | Endpoint                                  | Result                    |
|-----------------------|-----------------------------------|----------|-------------|-------------------------------------------|---------------------------|
| Diabetes              |                                    |          |             |                                           |                           |
| Franconi et al. (1995)| IDDM patients (n = 39)             | 90 days  | 1.5 g       | Glucose metabolism                        | NC                        |
| Elizarova and Nedosugova (1996) | IDDM patients (n = 10) | 30 days  | 1 g         | Glucose metabolism                        | Improved                 |
|                       |                                    |          |             | Triglyceride                              | Decreased                |
| Chauncey et al. (2003)| NIDDM patients (n = 32)            | 4 months | 3 g         | HbA1c                                     | NC                        |
|                       |                                    |          |             | Plasma lipid peroxide level               | NC                        |
| Brons et al. (2004)  | Overweight non-diabetic men (n = 20) | 8 weeks | 1.5 g       | Insulin secretion                         | NC                        |
|                       |                                    |          |             | Plasma lipid level                        | NC                        |
| Xiao et al. (2008)   | Overweight non-diabetic men (n = 6) | 2 weeks | 3 g         | Insulin sensitivity impaired by 48-h infusion of intravenous lipid | Improved                 |
|                       |                                    |          |             | Lipid peroxidation products               | Decreased                |
| Complications         | NIDDM patients with microalbuminemia (n = 10 each group) | 12 months | 3 g         | Microalbuminemia                          | NC                        |
|                       |                                    |          |             | Biomarkers for fibrosis                  | NC                        |
|                       | IDDM patients (n = 9)              | 2 weeks  | 1.5 g       | Hyperglycemia                             | NC                        |
|                       |                                    |          |             | Endothelium-dependent reaction            | Improved                 |

NC Not changed
which are endothelium-dependent reactions, were low in diabetic patients as compared to control subjects, taurine supplementation (1.5 g per day) for 2 weeks returned these parameters to control level without hypoglycemic effect, indicating the protective role of taurine on endothelium. Interestingly, same group previously reported that taurine supplementation (1.5 g per day) for 2 weeks also attenuated the impairment of flow-mediated dilatation in young cigarette smokers (Fennessy et al. 2003). Furthermore, while the culture medium conditioned with monocytes taken from smokers impaired the release of nitric oxide and increased in endothelin-1 in HUVECs, the levels of nitric oxide and endothelin-1 returned to control levels in HUVECs cultured with the monocyte-conditioned medium taken from smokers who had been treated with taurine. Therefore, the suppression of monocyte-endothelium interaction is likely to be a key action of protective role of taurine on endothelial function. Since the involvement of monocyte–endothelium interaction in endothelial dysfunction of diabetic animal models is also demonstrated as described above, it is possible that same pathway underlies the action of taurine against endothelial dysfunction in diabetic patients.

Discussion

As described in this article, numerous studies revealed that taurine supplementation is beneficial to diabetes and its complications in several animal models. Moreover, multiple actions of taurine coordinate to protect from diabetes and complications (Table 2). Especially, suppressive effect of taurine against oxidative stress is associated with various pathways in diabetic condition. First, reactivity of taurine against aldehyde can contribute to the reduction of AGE and modified LDL. Second, scavenging action against HClO can reduce the LDL modification and increase in bioavailability of the NO. Finally, taurine is likely to inhibit the ROS production via regulation of mitochondria (reviewed in Schaffer et al. 2009). While very high taurine concentration is found in mitochondria, several roles of taurine in mitochondria have been proposed. Taurine-containing modified uridine has recently been discovered at wobble position in mitochondrial transfer RNA (tRNA) (Suzuki et al. 2002). Taurine-modified tRNA may play a crucial role in the translation of proteins responsible for electron transport (Kirino et al. 2004), suggesting that taurine depletion might cause a decrease in taurine-modified tRNA and impairs electron transport capacity. Moreover, buffering property of taurine in mitochondrial matrix has been reported (Hansen et al. 2010). Therefore, taurine depletion in diabetes may contribute to mitochondrial dysfunction and it is possible that restoration of taurine contributes to normalize mitochondrial function, which may associate with inhibition of the ROS production from mitochondria. To elucidate the role of taurine depletion in mitochondrial function and in the development of diabetic complications, further studies, such as investigations using taurine transporter knock-out animals (Ito et al. 2008), will be required.

Nevertheless, most of clinical studies failed to prove the beneficial role of taurine on insulin resistance and diabetic

| Diabetic complications | Effects of taurine |
|------------------------|--------------------|
| Endothelial dysfunction| • Prevention of AGE production |
|                        | • Scavenging aldehydes → oxidized LDL ↓ |
|                        | • Scavenging HClO → oxidized LDL ↓, NO ↑ |
|                        | • LDL cholesterol ↓ |
|                        | • Inhibition of apoptosis in endothelial cells |
|                        | • Prevention of VCAM-1, ICAM-1 ↑ → leukocyte-endothelium interaction ↓ |
| Diabetic nephropathy   | • TGF-β↓ → prevention of fibrosis |
|                        | • Suppression of MAPK cascade, STAT3 → cell growth |
|                        | • Cytochrome P450 2E1↓ → oxidative stress ↓ |
| Diabetic retinopathy   | • Oxidative stress ↓ → prevention of Na+/K+/ATPase activity ↓ |
|                        | • Suppression of VEGF ↑ → retinal vascular function |
| Diabetic cataract      | • Oxidative stress ↓ |
|                        | • Prevention of protein carbonylation |
| Diabetic neuropathy    | • Oxidative stress ↓ in nerves |
|                        | • Prevention of the impaired calcium handling in sensory neuron |
|                        | • Suppression of NGF ↓ |
| Diabetic cardiomyopathy| • Suppression of Bcl-2↓ → apoptosis ↓ |
complications, whereas the others revealed the effectiveness. The discrepancies between animal experiments and clinical trials might be due to some limitations of clinical studies, such as a severity of the disease, other medications, given dose, duration of trial etc. Especially, the given dose of taurine per body weight is more than 10 times higher in animal experiments (e.g. diet containing 5% taurine) than in clinical trials (1.5–3 g taurine per day). Intake of taurine is thought to be quite safe as well as the amino acids found in food. While several reports strongly support that taurine is safe at levels up to 3 g/day, several clinical trials tested higher taurine dosages without adverse effects (Azuma et al. 1983, 1985; Shao and Hathcock 2008). Furthermore, since the pharmacological effect of taurine seems mild but not powerful, simultaneous therapy by using some medicines is also a problem. At present none of clinical studies have a sufficient numbers of patients. Therefore, long-term surveillance with large numbers of patients may be necessary to elucidate the effectiveness of taurine against diabetes or its complications in clinical study. Moreover, lifestyle, such as diet, and genetic factors, such as genomic polymorphisms which relate to individual differences, can affect to the result of trials. It is known that urinary taurine concentration in human varies widely among individuals (Yamori et al. 2001). Brons et al. (2004) reported a wide variation in the increasing rate of plasma taurine concentration after taurine administration among individuals. These variations of taurine movement among individuals must differ dependent not only on lifestyle but also genomic polymorphisms in taurine-related genes associated with the kinetics of taurine, such as taurine transporter. Therefore, we believe that the discovery of the genetic factors which determine the variation of taurine movement will help to elucidate the effectiveness of taurine against diabetes and its complications in humans.

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