C-peptide and Central Nervous System Complications in Diabetes

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Substantial evidence collected from clinical data and experimental studies has indicated that CNS is not spared from diabetes complications. Impairments in CNS function are well documented in both type 1 and type 2 diabetic patients as well as in various animal models of diabetes, in terms of alterations in cognition, neuropsychology, neurobehavior, electrophysiology, structure, neurochemistry and apoptotic activities. These data suggest that primary diabetic encephalopathy exists as a definable diabetic complication. The mechanisms underlying this CNS complication are not clear. Experimental studies have suggested that neuronal apoptosis may play an important role in neuronal loss and impaired cognitive function. In diabetes multiple factors are responsible for neuronal apoptosis, such as a perturbed IGF system, hyperglycemia and the aging process itself. Recent data suggest that insulin/C-peptide deficiency may exert an eminent role. Administration of C-peptide partially corrects the perturbed IGF system in the brain and prevents neuronal apoptosis in hippocampus of type 1 diabetes. In neuroblastoma SH-SY5Y cells C-peptide provides a dose-dependent stimulation on cell proliferation and an anti-apoptotic effect as well. These studies provide a basis for administration of C-peptide as a potentially effective therapy for type 1 diabetes.

Keywords C-peptide Deficiency; Diabetes; Hyperglycemia; Insulin Deficiency; Neuronal Apoptosis; Primary Encephalopathy

INTRODUCTION

Diabetes is a common metabolic disorder affecting many systems, including muscle, retina, kidney, blood vessels (small and large vessels), and the nervous system. With respect to the nervous system, the peripheral nervous system (PNS) has traditionally been considered as the sole nervous system complication in diabetes, whereas the central nervous system (CNS) was believed to be spared from diabetes. However, recent studies have suggested that diabetes causes primary as well as secondary impairments in CNS function. In this review, we will describe the CNS complications occurring in diabetic patients and experimental models of diabetes. Several possible mechanisms will be discussed, although we will focus on neuronal apoptosis and CNS dysfunction.

CNS COMPLICATIONS IN DIABETES

Clinical Studies

Diabetes-related cognitive dysfunction has been recognized since the early 1920s in the medical literature (Miles and Root, 1922). Subsequently, many studies of both type 1 and type 2 diabetic subjects have found that diabetes mellitus is associated with a series of neurobehavioral or neuropsychological changes. “Diabetic encephalopathy” could, therefore, be included as one of the complications associated with diabetes.

CNS Complications in Type 1 Diabetic Patients

Impairments in learning, memory, problem solving, and mental and motor speed are more common in type 1 diabetic patients than in the general population (McCarthy et al., 2002; Ryan, 1988; Ryan and Williams, 1993; Ryan et al., 1993). The cognitive impairments can be severe in rare cases (Deary et al., 1993; Gold et al., 1994), but is in most cases modest. The
variation in severity and pattern of impairment may be due to the subtle nature of the defects and variation in the diabetic populations selected for study, the patient number in each study, the psychological tests employed, and variations in glycemic controls among individual studies. A lower Wechsler Intelligence Scale for Children—Revised (WISC-R) performance IQ was found in type 1 diabetic children with disease onset at less than 7 years of age and disease duration greater than or equal to 5 years (Holmes and Richman, 1985). Diabetic patients suffered from significantly more introspective symptoms than their healthy counterparts, especially with respect to somatic symptoms, sleeping disturbances, compulsions, and depressive moods (Blanz et al., 1993; Kovacs et al., 1997; Lustman et al., 1988; Popkin et al., 1988). These studies thus indicate impaired CNS function in type 1 diabetes.

The neuropsychological changes are often accompanied with objective electrophysiological, structural, and neurochemical abnormalities. In electrophysiological studies, measurements of latencies of evoked potentials have been widely used to examine the functional integrities of the CNS in diabetic patients (Di Mario et al., 1995). Delayed conduction velocity and data-processing time in the CNS were demonstrated in type 1 diabetic patients, including brainstem auditory-evoked potential, somato-sensory-evoked potentials, visual-evoked potentials, and motor-evoked potentials (Comi, 1997; Cracco et al., 1984; Harkins et al., 1985; Pietravalle et al., 1993; Uberall et al., 1996; Verrotti et al., 2000). The latency of the P100 wave of the visual-evoked potential, which is believed to be generated in the visual cortex, is increased in diabetic patients (Paris and Ucciolli, 2001; Ziegler et al., 1994). The event-related (P300) potential is an objective measure of cognitive function (Madl et al., 1994; Picton, 1992). Several studies have shown that P300 latency, a neurophysiological correlate of information processing, such as stimulus evaluation, alertness, and memory updating (Kramer et al., 1998), is increased in diabetic patients (Kramer et al., 1998; Pozzessere et al., 1991; Uberall et al., 1996), indicating an impairment of higher brain functions.

In terms of structural changes, several abnormalities have been described in type 1 diabetic patients, such as diffuse and local degenerative changes of cerebral cortex (Reske-Nielsen and Lundbaek, 1964; Reske-Nielsen et al., 1965), neuronal loss, demyelination and gliosis, and infarction secondary to microangiopathy (De Jong, 1977). Magnetic resonance imaging and computed tomography have shown that widened sulci and/or enlarged lateral ventricles (Araki et al., 1994; Lunetta et al., 1994; Perros et al., 1997) and increased occurrence of white matter hyperintensities (Araki et al., 1994; Perros et al., 1997) are more pronounced in diabetic patients than in age-matched control subjects. These studies indicate structural changes in CNS of diabetes.

Because hypoglycemia itself can cause brain damages (de Courten-Myers et al., 2000; Vannucci and Vannucci, 2001; Yager, 2002), it remains controversial as to whether the above CNS abnormalities are caused by diabetes per se (primary) or are the result of hypoglycemic episodes (secondary), which occur frequently in type 1 diabetic patients receiving intensive insulin treatment. Recent studies have shown that a duration-dependent decline in cognitive function occurs in type 1 diabetic patients without hypoglycemic episodes (Kramer et al., 1998), and that impaired intellectual and cognitive development occur in type 1 diabetic children, who have not experienced hypoglycemic episodes (Schoenle et al., 2002). These latter impairments correlate with diagnosis at young age, male sex, and metabolic status at time of diagnosis. Therefore, CNS complications seem to be impacted by diabetes itself without implementing hypoglycemic episodes as a causative mechanism and hence being primary.

In summary, evidence obtained from pathologic, physiologic, psychologic, and behavioral studies point toward a “primary diabetic encephalopathy” in type 1 diabetes.

CNS Complications in Type 2 Diabetic Patients

Neuropsychologic studies in type 2 diabetes patients have shown more consistent results compared to type 1 diabetes patients. Cognitive deficits (Gradman et al., 1993; Perlmuter et al., 1984; Ryan and Geckle, 2000a; Strachan et al., 1997; Tun et al., 1990; Worrall et al., 1993) and poor performance in abstract reasoning and complex psychomotor functioning (Reaven et al., 1990) occur in type 2 diabetes. Complex cognitive functions, as demonstrated by complex cognitive tasks requiring storage and retrieval of new information, are affected, whereas performances of less demanding tasks, such as immediate memory and simple reaction time, are not significantly altered (Tun et al., 1990). Electrophysiological studies have demonstrated delayed CNS conduction velocity (Donald et al., 1984; Varsik et al., 2001; Verrotti et al., 2000) and increased latency of P100 (Ziegler et al., 1994). Several studies demonstrate that P300 latency is increased in type 2 diabetic patients (Dey et al., 1997; Kurita et al., 1995; Mochizuki et al., 1998; Mooradian et al., 1988), indicating an impairment of higher brain function. Recently, increased blood-brain barrier (BBB) permeability in type 2 diabetes was demonstrated by magnetic resonance imaging when intravenous administration of gadolinium diethylenetriamine pentaacetic acid (Gd-DTPA) was used to identify BBB permeability, suggesting that loss of BBB integrity may play a role in CNS dysfunction in diabetes (Starr et al., 2003). In the All Wales Research into Elderly (Aware) Diabetes Study, elderly type 2 diabetic patients displayed significant excess of cognitive dysfunction associated with a poorer ability in diabetes self-care (Sinclair et al., 2000). However,
no significant impairments in learning, memory and problem-solving skills were found in middle-aged type 2 diabetic patients compared to control subjects (Ryan and Geckle, 2000b). In fact, few studies have been conducted regarding cognitive deficits in middle-aged type 2 diabetic patients, among which the largest patient number was 80 type 2 diabetic patients versus 81 nondiabetic controls (Lowe et al., 1994). Therefore, more studies with large number of patients are needed in determining cognitive function in middle-aged type 2 diabetes. The current thought is that learning and memory dysfunction are more prominent in elderly type 2 diabetic subjects (Ryan and Geckle, 2000a). As to whether this is due to a potentiation of the normal aging process by superimposed diabetes or a function of diabetes duration alone is not settled.

The etiology of cognitive dysfunction in type 2 diabetes is not fully understood. It is likely that cognitive impairments are caused by an interaction between metabolic abnormalities intrinsic to diabetes and diabetes-specific complications. After all, the higher frequency of cerebral stroke in type 2 diabetes may be associated with cognitive deficits. Several studies have suggested that the cerebrovascular mortality rate is higher in patients with type 2 diabetes (Barrett-Connor and Khaw, 1988; Lehto et al., 1996). The risk for developing stroke is increased 2- to 5-fold compared to nondiabetic control subjects (Manson et al., 1991; Stamler et al., 1993). Recently, 5102 patients with type 2 diabetes in the UK Prospective Diabetes Study (UKPDS) were analyzed using a newly developed mathematical model. The results confirmed the higher risk of stroke in type 2 diabetes (Kothari et al., 2002). Although cerebrovascular disease is barely mentioned in patients with type 1 diabetes, recent analyses of mortality from 23,000 patients with type 1 diabetes has shown for the first time that cerebrovascular mortality is raised at all ages in this patient group (Laing et al., 2003). Therefore, type 1 diabetes is at least as great a risk factor for cerebrovascular mortality as is type 2 diabetes. Based on these reports, the continuing study of cerebral stroke in diabetes will provide useful insight into possible underlying mechanisms.

**Experimental Studies**

**Structural Changes**

A variety of structural changes has been described in the CNS of streptozotocin (STZ)-induced diabetes in rats and Chinese hamsters (Bestetti and Rossi, 1980, 1982; Garris et al., 1982; Jakobsen et al., 1987; Luse, 1970; Mukai et al., 1980). Structural alterations of the ventromedial hypothalamus, including accumulation of glycogen, degeneration of neurons, and atrophy of tanycytes, have been described. Ultrastructural abnormalities, such as dilated and fragmented endoplasmic reticulum, degranulated ergastoplasm, increased number of microtubuli, myelin figures, irregularities in the form of nuclei, and appearance of chromatin, were described. A significant loss of neocortical neurons occur in STZ-diabetic rats compared to nondiabetic control rats (Jakobsen et al., 1987). In addition, significant structural alterations in the BBB were shown in experimental diabetic animals (Mooradian, 1997), consistent with the magnetic resonance imaging findings in diabetic patients (Starr et al., 2003).

**Cognitive Functional Changes**

Impaired spatial learning and memory have been demonstrated using the Morris water maze system in various diabetic animals including BB/W rats, STZ-diabetic rats, STZ-diabetic mice, and Otsuka Long Evans Tokushima Fatty (OLETF) rats (Biessels et al., 1998; Flood et al., 1990; Kamal et al., 2000; Li et al., 2002a; Luesse et al., 2001). Multiple cognitive components are involved in this task, such as problem solving, enhanced selective attention, formation of internal
representation of the external environments, and storage and retrieval of relevant information (Bannerman et al., 1995). The hippocampal LTP and LTD are two forms of neuronal synaptic plasticity. In young adult STZ-diabetic rats, hippocampal LTP expression is impaired, whereas LTD expression is enhanced (Biessels et al., 1996; Kamal et al., 1999). The alterations in hippocampal synaptic plasticity is associated with defects in spatial learning and memory as detected by the Morris water maze (Biessels et al., 1996, 1998). The impaired performance in the Morris water maze started 10 weeks after the injection of STZ (Biessels et al., 1998; Kamal et al., 2000) and the degree of LTP impairment correlated with the duration of diabetes (Kamal et al., 1999). In diabetic BB/W rats, we failed to show impaired spatial learning and memory in acutely diabetic rats (2-month), whereas significant deficits in the Morris water maze were demonstrated in 8-month diabetic BB/W rats (Li et al., 2002b). These results suggest that progressive learning and memory deficiencies in type 1 diabetic rats occur in a duration-related fashion.

**Neuronal Loss/Apoptosis in Type 1 Diabetes**

A significant loss of neocortical neurons occurs in STZ-diabetic rats (Jakobsen et al., 1987). In the diabetic Chinese hamster, neuronal death occurs in the arcuate and ventromedial nuclei (Garris, 1984). The nature of cell death was not defined in these early studies, nor was it determined as to whether neuronal loss was associated with changes in size. In the spontaneously diabetic BB/Wor rat, neuronal counts of hippocampal pyramidal cells were performed in the hippocampal CA1–CA4 regions. No significant differences were demonstrated at 2 months of diabetes. In 8-month diabetic animals, there were a 37% and 24% loss of pyramidal cells in CA1 and CA3 regions, respectively, whereas other hippocampal regions showed non-significant decreases in neuronal densities (Li et al., 2002b). To exclude the effect of body weight on neuronal density, we compared 8-month diabetic rats with weight-matched control rats, which did not hippocampal neuronal loss. These data suggest that decreased neuronal density is not weight related but diabetes duration related. Comparisons with hyperglycemia- and duration-matched spontaneously type 2 diabetic BBDRZ/Wor rats revealed a much milder but still significant neuronal loss of the hippocampal CA4 region (Li and Sima, unpublished data).

The above studies indicate that neuronal loss occurs in type 1 diabetic animals. As to the nature of this neuronal loss, we have proposed that apoptosis might play an important role. Several in vitro and in vivo studies appear to support this notion (Li et al., 2002b, 2003). No apoptotic activity of hippocampal pyramidal neurons was shown in 2-month diabetic BB/Wor rats, nor was any cognitive deficits detected by the Morris water maze procedure, nor was any neuronal loss detectable. On the other hand, in 8-month diabetic rats, TUNEL-positive neurons, positive DNA laddering, and increased Bax expression and caspase-3 activity were evident and associated with decreased neuronal density and impaired Morris water maze performances (Li et al., 2002b). Hence, duration-related apoptosis is likely to account for the neuronal loss and the concomitant emergence of cognitive impairments in the type 1 diabetic BB/Wor rat.

**Ischemic Injury in Diabetic Animals**

It is well known that diabetes aggravates brain damage in experimental and clinical stroke subjects. Diabetes accelerates maturation of neuronal damage, increases infarct volume, and induces postischemic seizures (Muranyi et al., 2003). A brief period of 30 minutes’ focal ischemia in normoglycemic rats leads to brain damage in a delayed fashion: infarction develops after 3 days and full maturation is reached by 2 weeks after recirculation (Du et al., 1996). If preischemic hyperglycemia exists or in STZ-diabetic rats, the damage is more severe than that in nondiabetic animals (Muranyi et al., 2003) and the damage evolves faster: infarction develops after 2 hours and is matured after 4 to 6 hours after the ischemic insult (Gisselsson et al., 1999; Li et al., 1998). The mechanisms underlying diabetes-related aggravation of ischemic brain damage are unclear. Apoptosis, or programmed cell death (PCD), is believed to play an important role in the pathogenesis of various human disorders including diabetes (Barr and Tomei, 1994; Fadeel et al., 1999; Orrenius, 1995; Thompson, 1995) and a vast body of studies has shown that apoptosis is associated with cerebral ischemia and ischemia-reperfusion injury. It has been demonstrated that a large number of neurons bordering the maturing infarct exhibit prominent TUNEL staining, and DNA prepared from the penumbral area of ischemic cortex show internucleosomal fragmentation (Du et al., 1996). Increased levels of either expression or activities of caspase-3 (Chen et al., 1998; Schulz et al., 1999) and induction of caspase-activated deoxyribonuclease activity (Luo et al., 2002) are detected in ischemic neurons. DNA fragmentation in ischemic brains has been demonstrated by either gel electrophoresis or TUNEL staining (Charriaut-Marlangue et al., 1998; Endres et al., 1998; Li et al., 1995). A strong correlation has been found among caspase-3 activity, TUNEL staining, and the appearance of apoptotic neurons (Cao et al., 2001; Charriaut-Marlangue et al., 1998; Chen et al., 1998; Namura et al., 1998). Furthermore, cytosolic cytochrome c release is markedly enhanced in both the ischemic focus and the penumbra in STZ-diabetic rats, together with increased activation of caspase-3 and accelerated cleavage of poly-ADP ribose polymerase (PARP) (Du et al., 1996), all of which suggest that apoptosis plays a role in the exaggerating effect of diabetes on cerebral ischemic lesions.
Recently, the effects of local ischemia on type 1 diabetic rats versus normal control rats were reported (Britton et al., 2003). A significant increase in the number of TUNEL-positive and caspase-3–positive cell was demonstrated in selected brain regions (hypothalamic preoptic area, piriform cortex, and parietal cortex) of STZ-diabetic rats subjected to middle cerebral artery occlusion for 24 to 48 hours, compared with nonoccluded diabetic rats (Britton et al., 2003). Consistent with these findings, preliminary data show that the ladder pattern of nucleosomal DNA fragmentation (ligation mediated–polymerase chain reaction [LM-PCR]) cannot be detected in unlesioned normal or STZ-diabetic animals, but is evident in infarcted cortex of diabetic rats (Li and Sima, unpublished data). Bax expression is increased in normal infarcted cortex and is further increased in infarcted diabetic cortex. These data suggest that diabetes confers an expansion of stroke via increased apoptotic activity.

In summary, data obtained from experimental type 1 diabetic animals show definite alterations in structure, neurotransmitters, electrophysiology, cognitive function, neuronal density, and apoptotic activity. These results clearly indicate that “diabetic encephalopathy” is an identifiable diabetic complication.

POSSIBLE MECHANISMS

The mechanisms underlying neuronal apoptosis and CNS dysfunction are not clear. We believe that contributing factors are insulin deficiency with concomitant C-peptide deficiency, as well as hyperglycemia and possibly the aging process itself (Figure 1).

**Insulin/C-peptide Deficiency**

Insulin plays an important role in regulation of brain metabolism. Recent studies show that insulin exerts additional modulatory roles on brain functions such as feeding (Gerozissis et al., 2001; Schwartz et al., 1992), learning, and memory (Gerozissis et al., 2001; Zhao and Alkon, 2001). In cerebral stroke, administration of insulin prevents neuronal death (Voll and Auer, 1991a; Voll and Auer, 1991b) and reduces neurological disability (LeMay et al., 1988). These neuroprotective effects of insulin may be associated with restoration of protein synthesis via increased dephosphorylation of eukaryotic initiation factor 2α (Sullivan et al., 1999). In STZ-diabetic rats, cognitive deficits is partially corrected by insulin (Biessels et al., 1998; Gradman et al., 1993). In vitro studies also show that insulin has an ant apoptotic effects (Bertrand et al., 1999; Lee-Kwon et al., 1998; Li et al., 2003). We have shown that the insulin receptor is downregulated in the brain of type 1 diabetic BB/Wor rats (Li et al., 2002b), suggesting that impaired insulin action caused not only by insulinopenia but also by impaired insulin receptor expression may play a role in CNS apoptosis in diabetes.

In addition to insulin deficiency, C-peptide deficiency has been considered as a pathogenic factor in type 1 diabetic
C-peptide is a 31–amino acid peptide that is cleaved from proinsulin during biosynthesis of insulin (Li et al., 2001; Steiner and Rubenstein, 1997). Recent studies have shown that C-peptide possesses physiological functions other than providing structural support for proinsulin cleavage (Wahren and Johansson, 1998; Wahren et al., 1994, 1996; Sima, 2003a, 2003b). In patients with type 1 diabetes, C-peptide improves renal function, reduces urinary albumin excretion and glomerular filtration, and decreases blood retinal barrier leakage (Fernqvist-Forbes et al., 2001; Forst et al., 1998; Johansson et al., 1992, 1996; Zierath et al., 1991, 1996). Chronic C-peptide replacement prevents functional and structural peripheral nerve changes in type 1 diabetic rat models (Huang et al., 2002; Ido et al., 1997; Li et al., 1999; Samnegard et al., 2001; Wu et al., 1996; Zhang et al., 2001; Sima et al., 2001; Pierson et al., 2003), suggesting that C-peptide deficiency is a participating factor in the causation of type 1 diabetic complications. We recently showed that administration of C-peptide partially corrects perturbed insulin-like growth factor (IGF) activity and prevents the neuronal apoptosis in hippocampus of type 1 diabetic BB/W rats (Li et al., 2002c), indicating a relationship between C-peptide deficiency, IGF perturbation, and neuronal apoptosis. In neuroblastoma SH-SY5Y cells, C-peptide provides a dose-dependent stimulatory effect on cell proliferation, whereas no effects were evident with the same concentration of scrambled C-peptide (Figure 2). In the same in vitro cell cultures, apoptosis is induced by high concentrations of glucose, mimicking hyperglycemia. Apoptosis of SH-SY5Y cells is inhibited by addition of insulin alone, whereas the combination of insulin and C-peptide show an enhanced antiapoptotic effect, as assessed by prevention of nuclear condensation and shrinkage, reduction in the number of apoptotic cells, and stimulation of Bcl2 expression and nuclear factor (NF)-κB (Li et al., 2003). The inactivated form NF-κB exists in the cytoplasm as a complex consisting of p50, p65, subunits of inhibitor-κB (IκB). On stimulation, IκB is phosphorylated and degraded, resulting in activation and translocation of NF-κB to the nucleus, where it activates target gene transcription. It is known that NF-κB and the genes regulated by this transcription factor, such as those coding for tumor necrosis factor (TNF) receptor–associated factor 1 (TRAF1), TRAF2, inhibitor-of-apoptosis (IAP) proteins c-IAP1 and c-IAP2, manganese superoxide dismutase (MnSOD), Bcl2, and BclxL, play important roles in the regulation of apoptosis (Aggarwal, 2000; Bours et al., 2000). In the SH-SY5Y cells grown under elevated glucose concentrations, C-peptide plus insulin gives rise to a significant increase in nuclear NF-κB staining (Figure 3), suggesting that insulin/C-peptide play effective antiapoptotic roles via activation and translocation of NF-κB. In summary, the data obtained from in vitro system are consistent with the results obtained from experimental diabetes complications.
in animals and support the notion that insulin/C-peptide deficiency plays an important role in type 1 diabetes-induced neuronal apoptosis.

**IGF System Impairments**

Perturbed IGF system has been shown in the CNS of STZ-diabetic rats. After 2 weeks of diabetes, IGF-II mRNA content is significantly decreased in brain and spinal cord. Insulin replacement partially restores IGF-II mRNA levels in cerebral, cortex, medulla, and spinal cord (Wuarin et al., 1996). We have systematically examined the IGF system (IGF-1, IGF-II, IGF-IR, and insulin receptor [IR]) in the BB/Wor model of type 1 diabetes. Using Western blotting, Northern hybridization, and in situ hybridization, we found significant reductions in the expression of IGF-I, IGF-II, IGF-IR, and IR already in 2-month diabetic BB/Wor rats, which persisted in 8-month diabetic rats, indicating that these abnormalities precede the functional cognitive impairments and the apoptotic neuronal loss in hippocampus (Li et al., 2002b).

**Hyperglycemia**

Hyperglycemia causes increased glucose levels in the brain. It is well established that hyperglycemia causes oxidative stress via the polyol pathway, enhanced advanced glycation end products (AGEs), and increased lipid peroxidation by-products and imbalances in the generation of reactive oxygen species and their scavengers (Ceriello et al., 1993; Lipinski, 2001; Mercuri et al., 2000; Opara, 2002). Oxidative stress is also associated with cerebral ischemic injury (Kaminski et al., 2002; Love, 1999) and neuronal apoptosis (Gorman et al., 1996; Nicotera et al., 1997; Sastre et al., 2000). Extreme hyperglycemia also causes nonketotic hyperosmolar coma, characterized by volume depletion, altered consciousness, confusion, and, less frequently, neurological deficit in the absence of ketonemia or acidosis. Hyperosmolar coma is a severe complication of diabetes with a high overall mortality rate (Amundson et al., 1996; Braaten, 1987; Rolfe et al., 1995; Ting, 2001). It is therefore likely that hyperglycemia per se may induce apoptotic activities, preferably via oxidative stress, as indicated by hippocampal apoptosis with neuronal loss in the type 2 hyperinsulinemic BBDRZ/Wor rat (Li and Sima, unpublished data.

**Aging**

Aging in developed countries is associated with an increasing prevalence of hypertension, atherosclerotic vascular diseases, reduced insulin sensitivity, and type 2 diabetes
(Barbagallo et al., 1997). In the Rotterdam Study, a community-based prospective cohort study, chronic disorders of the elderly were investigated. The relative risk for developing dementia was doubled in diabetic patients (Ott et al., 1999), suggesting that diabetes increases the risk of dementia, particularly in the elderly (Gardoni et al., 2002). It was recently demonstrated (Terry et al., 2001) that the expression of IGF-I, IGF-IR, and IR is reduced in susceptible hippocampal areas in Alzheimer’s disease. We have shown that impaired IGF and insulin activities precede the development of hippocampal apoptosis in type 1 diabetic BB/W rats (Li et al., 2002b). The similarity in perturbations of the IGF system suggests that possible common etiological factors may be present in Alzheimer’s disease and diabetes, two disorders in which apoptosis has been invoked as a mechanism. However, other studies have failed to find an association between diabetes and incident Alzheimer’s disease (Hassing et al., 2002; MacKnight et al., 2002). More work is needed to elucidate the effect of aging on impaired CNS function in diabetes.

CONCLUSION

Diabetes causes impairments in CNS functions. Primary diabetic encephalopathy is characterized by neurophysiological impairments, structural changes, and cognitive deficits. The mechanisms underlying this CNS dysfunction of diabetes are not fully elucidated. We propose that insulin/C-peptide deficiency in type 1 diabetes may be one important factor in the pathogenesis of CNS dysfunctions. It appears that apoptotic phenomena are contributing factors. However, much work is still needed to elucidate possible apoptotic pathways and likely alterations in contraregulatory mechanisms. The increased infarct size seen both in diabetic patients and animals following cerebral ischemia is likely to be caused by apoptosis. As to whether these apoptotic mechanisms are the same as those involved in primary diabetic encephalopathy remain to be examined.

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