Abstract. The dual expression system for the suppression and clearance of insulin has not been previously used to produce transgenic mice for diabetes-related disease. The aim of this study was to produce new transgenic mice coexpressing specific insulin small interfering RNA (siRNA) sequences and the human insulin degrading enzyme (hIDE) gene in order to examine the diabetes-like phenotype. To achieve this, a new lineage of transgenic mice was produced by the microinjection of the dual expression constructs (pH1/siRNA_insulin-CMV/hIDE) into mouse fertilized eggs. The results showed that overexpressing the insulin siRNA and hIDE genes resulted in the induction of the human enzyme, impaired glucose tolerance and lower serum insulin levels compared to the Non-Tg mice. Moreover, the Tg mice aged 20 weeks had a significantly activated ER stress signaling compared to their Non-Tg counterparts, which may be associated with the suppression of insulin production in the pancreas and the degradation of insulin in the liver, respectively. Therefore, insulin-suppressed transgenic mice can be used to examine diabetes as a new diabetes-like phenotype model, which results in a lower level of circulating insulin without the destruction of pancreatic islets.

Introduction

The suppression of insulin production and the degradation of insulin are very effective areas for the development of new model animals for type I diabetes. Insulin removal helps control the cellular response to hormones by decreasing the substrate availability. The degradation process may also be involved in mediating some aspects of insulin action. Most evidence suggests that an insulin-degrading enzyme is associated with the primary mechanism of insulin degradation, although other systems including PDI, lysosome, and other enzymes undoubtedly contribute to the metabolism of insulin (1). IDE (insulysin, insulinase, EC3.4.24.56) is a member of the M16 family of zinc-metalloendopeptidases containing a tetrad of conserved residues \[H-x-x-E-H-x (76)-E\] at its catalytic center (1). This protein is broadly expressed in both insulin-sensitive and insulin-insensitive tissues, and is localized mainly in the cytoplasm within the cells. However, different forms of this protein can be secreted into extracellular spaces or associated cell surfaces (2,3). Studies have reported various IDE high-affinity substrates, which were polypeptides 28-51 amino acids in length. A variety of physiological peptides have been shown to be substrates in vitro for IDE including insulin and glucagons (4), the atrial natriuretic factor (5), transforming growth factor \(\beta\) (6,7), the atrial natriuretic peptide (8), ß-endorphin and dynorphins (9), amylin (10), and amyloid ß peptides (11,12). IDE may also have other biological functions such as the clearance of the ß-amyloid precursor protein in the intracellular domain (AICD), which is suggested to regulate the transcriptional activity in the nucleus (13). Furthermore, it was demonstrated that IDE interacts directly and enhances the DNA binding of androgen and glucocorticoid receptor fragments in vitro (14).

Duckworth et al (15) reported that IDE exists in the cell as a complex with the 26S proteasome, which is regarded as the principal site of the ubiquitin-dependent intracellular protein degradation. However, no transgenic mice expressing a hIDE protein under the control of the human cytomegalovirus (CMV) immediate early promoter have been produced to study the IDE function in vivo.

Small interfering RNA or double-stranded RNA (dsRNA)-based gene silencing is a highly conserved mechanism for sequence-specific post-transcriptional gene silencing between various species (16,17). This technique has been used in the nematode Caenorhabditis elegans to suppress gene expression by an injection of a double-stranded mixture into adult
animals (18). Furthermore, Bernstein et al (19) identified the new enzyme, Dicer, which produces putative guide RNA. Dicer is a member of the RNase III family of nucleases that specifically cleave double-stranded RNA, and is evolutionarily conserved in worms, flies, plants, fungi and mammals. Until recently, siRNA has not been used to silence mammalian genes, since the introduction of dsRNA longer than 30 nt induces a sequence-non-specific viral response (20). However, Elbashir et al (21) suggested that the introduction of short duplexes of synthetic 21-23-nt RNAs into mammalian cells has gene specific silencing effects. Nevertheless, this method has a critical limitation in that the transfected synthetic siRNA works for only a few days in mammalian cells. This problem can be overcome by a vector-based technique for the synthesis of siRNAs, which was driven by the RNA polymerase III promoter (22,23,24). In addition, Hasuwa et al (25) demonstrated that transfected siRNA silenced the ubiquitously enhanced expression of green fluorescent protein in all tissues of mice and rats. These effects of gene suppression were transmitted into their progenies containing the expression vector. However, there are no reports as to whether the transgene-based siRNA system can suppress the expression of the insulin gene in adult mice.

This study has developed novel Tg mice coexpressing siRNAinsulin and hIDE, which induce the suppression of insulin production and the activation of insulin clearance, respectively. In addition, these transgenic mice were shown to have diabetes-like phenotypes such as insulin suppression, impaired glucose tolerance, and an altered ER stress signaling pathway. These results suggest that siRNAinsulin-hIDE transgenic mice may be useful for developing an animal model for diabetes, which show a lower level of insulin in blood without the destruction of pancreatic islets.

Materials and methods

Plasmid construction. In an attempt to suppress insulin production in Tg mice, the siRNA sequences of 5 types were selected from the open reading frame of the rat insulin cDNA sequence using the siRNA target finder program (Ambion, USA). The transcription-based siRNA vector was constructed using the Silencer siRNA construction kit (cat. no. 5770, Ambion). Each siRNA duplex of 5 types was ligated into the pEGFP C-1 vector, and then ligated into the pCMV/hIDE, for extension. The 3,080-bp products of the amplified IDE gene, with 25 cycles of the following: 95°C for 30 sec for denaturation, 62°C for 30 sec for annealing and 72°C for 45 sec for extension. The 3,080-bp products of the amplified IDE sequence in length were cloned into the pGEM-T vector (Promega A3600). The IDE fragment (3,080-bp) was obtained by digesting pGEM-hIDE with AgeI/SacII enzymes, and cloning them into the AgeI/SacII site of pEGFP-C-1 (pCMV/hIDE) (Fig. 2).

Finally, the dual expression vector involving pH1/siRNAinsulin and pCMV/hIDE were constructed by cloning the H1/siRNAinsulin fragment into the AgeI site of pCMV/hIDE. The resulting plasmid, named pH1/siRNAinsulin-CMV/hIDE, was verified for the presence and orientation of the siRNA duplex by an automatic sequencer (ABI 373, Perkin Elmer).

Production of transgenic mice. The pH1/siRNAinsulin-CMV/hIDE plasmid was digested with BlnI in order to remove the prokaryotic sequence, and was then electrophoresized in an agarose gel. The linear gene fragment of the pH1/siRNAinsulin-CMV/hIDE fragment (6,214-kb) was excised and extracted from the agarose gel. The pH1/siRNAinsulin-CMV/hIDE fragment was purified by electrophoretisation, diluted to 4 ng/μl and microinjected into the male pronucleus of the fertilized embryos, which were obtained by crossing C57BL/6 (female) mice with DBA2 (male) mice. The injected eggs were then transferred into the oviducts of a pseudopregnant ICR recipient female.

Maintenance of mice. All of the mice were kept in an accredited Korean FDA animal facility in accordance with the AAALAC International Animal Care policies (Accredited Unit, Korean Food and Drug Administration, unit no. 000996). The mice were given a standard irradiated chow diet (Purina Mills Inc.) ad libitum, and were maintained in a specified pathogen-free state under a strict light cycle (light on at 06:00 h and off at 18:00 h). All pedigrees were hemizygous for their transgene.

Southern blot analysis of the transgene inserted in genomic DNA. The presence of the inserted transgene was determined from the genomic DNA extracted from the tail of the 4-week-old founder mice, and the transgenes were detected by Southern blot analysis of the HindIII-digested tail DNA. After electrophoresis, the DNA was transferred overnight to nylon membranes in 10X SSC according to the manufacturer's instructions. The membranes were then immersed in 0.4 N NaOH, neutralized in 0.2 M Tris-HCl (pH 7.5) and 2X SSC, and then allowed to air dry. Each membrane was prehybridized at 65°C for 2 h in a hybridization buffer. The 32P-labeled siRNA-hIDE probe was added to the membranes, and hybridization was allowed to proceed at 65°C for 18 h. The membranes were then washed using the following procedure: 15 min in 2X SSC/0.2% SDS at room temperature (once); 15 min in 1x SSC/0.2% SDS at 65°C (three times) and 30 min in 0.5X SSC/0.2% SDS at 65°C (three times). The filters were then exposed to Kodak X-ray film at -70°C. The hybridization signal was quantified using a GS-690 imaging densitometer (Biorad, USA).
Real-time PCR analysis. The transgene was identified by DNA-PCR analysis of the genomic DNA that had been isolated from the tails of the 4-week-old founder mice. This transgene DNA was synthesized using the sense and antisense primers, 5'-GGT TTT CCC AGT CAC GAC-3 and 5'-GAG TTA GCT CAC YCA TAG GC-3, respectively, with a complementary siRNA oligonucleotide plasmid ranging from 359-376 and 663-683 nucleotides as the DNA template. After 25 cycles of amplification, the level of the siRNA-oligo-hIDE product (324-bp) was quantified using a Kodak electrophoresis documentation and analysis system 120 on 1% agarose gels. For RT-PCR analysis, each tissue sample was frozen in liquid nitrogen. The frozen tissues were chopped with scissors and homogenized in an RNAzol B solution (CS104, Tet-Test Inc.). The isolated RNA was then determined by UV-spectroscopy. The expression of the transgenes was examined using RT-PCR with 5 μg of the total RNA from each tissue. Oligo-dT primer (Gibco BRL (18418-012)) (500 ng) was annealed at 70°C for 10 min. The complementary DNA, which served as the template for further amplification, was synthesized by adding dATP, dCTP, dGTP and dTTP with 200 units of the reverse transcriptase. Ten pmole of the sense and antisense primers were then added, and the reaction mixture was subjected to 28 cycles of amplification. Amplification was carried out in a Perkin-Elmer thermal cycler using the TaqMan Universal PCR master mix (AB4034437) on 4-20% SDS-PAGE gel for 3 h and transferred to nitrocellulose membranes for 2 h at 40 V. Each membrane was incubated separately with the primary, anti-IDE (PC730, Calbiochem), anti-JNK (SantaCruz), anti-p-JNK (SantaCruz), anti-βF2 antibody (Cell Signaling), anti-p-βF2 antibody (Cell Signaling), and anti-actin (A5316, Sigma) antibodies overnight at 4°C. The membranes were washed with a washing buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na 2HPO 4, 2 mM KH 2PO 4 and 0.05% Tween-20) and incubated with horseradish peroxidase-conjugated goat anti-rabbit IgG (Zymed) at a 1:1,000 dilution at room temperature for 2 h. The membrane blots were developed using a chemiluminescent reagent plus kit (ECL, Pharmacia).

Glucose tolerance test. The glucose tolerance was determined from five mice/group using an intraperitoneal glucose tolerance test (1.5 mg glucose/g) (IPGTT) after a 24-h fast with blood glucose measurements performed at 0, 15, 30, 60, 90 and 120 min. An IPGTT study was carried out at 20 weeks of age. The glucose levels in the blood from the Tg and Non-Tg mice were detected with the sensitive strip method using a glucose monitoring system (I-sens Co., Korea).

Western blot. The protein prepared from the tissues of the Tg mice and Non-Tg mice was separated by electrophoresis in a 4-20% SDS-PAGE gel for 3 h and transferred to nitrocellulose membranes for 2 h at 40 V. Each membrane was incubated separately with the primary, anti-IDE (PC730, Calbiochem), anti-JNK (SantaCruz), anti-p-JNK (SantaCruz), anti-βF2 antibody (Cell Signaling), anti-p-βF2 antibody (Cell Signaling), and anti-actin (A5316, Sigma) antibodies overnight at 4°C. The membranes were washed with a washing buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na 2HPO 4, 2 mM KH 2PO 4 and 0.05% Tween-20) and incubated with horseradish peroxidase-conjugated goat anti-rabbit IgG (Zymed) at a 1:1,000 dilution at room temperature for 2 h. The membrane blots were developed using a chemiluminescent reagent plus kit (ECL, Pharmacia).

The quantification of insulin by ELISA. The level of insulin in the serum from the Tg and Non-Tg mice was determined using the ultra-sensitive assay procedure and reagents included in the Mercodia rat insulin ELISA kit (cat. no. 10-1137-01; Mercodia, Sweden). The sera and standards were incubated in antibody-coated plates for 2 h at room temperature on a plate shaker at 500-600 rpm. The wells were then washed six times using an automatic plate washer (PV100; Hoefer, USA). The HRP conjugate was added to all plates and incubated for 30 min at room temperature on a shaker. The reaction was terminated with the addition of a 50-μl stop solution. The plates were read using a Molecular Devices Vmax plate reader (Sunnyvale, CA, USA) at 450 nm.

Immunohistochemistry. Immunohistochemical staining was performed as previously described (26). Briefly, the level of IDE protein synthesis was determined using optical microscopy after fixing the tissue samples in 5% formalin for 12 h, embedding them in paraffin, and slicing them into 4-μm thick sections. These sections were de-paraffinized with xylene, rehydrated, and pretreated for 30 min at room temperature with a PBS-blocking buffer containing 10% goat

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Real-time PCR. Real-time PCR analysis was performed using a TaqMan Universal PCR master mix (AB4034437) on an ABI PRISM sequence detection system (ABI-SDS) (Applied Biosystems). After reverse transcription, PCR was carried out in triplicate for each mixture with the synthesized cDNA added to the TaqMan Universal master mix and CYP Adb product (mixture of sense, antisense, and FAM primer). PCR for glyceraldehyde-3-phosphate dehydrogenase (GAPDH), as an endogenous control was performed simultaneously, with the exception of the rodent GAPDH control reagent (AB4308313). TaqMan probes have two fluorescent dyes, one at the 5′-terminal (Reporter, R) for displacing the strand, and the other at the 3′-terminal (Quencher, Q) for blocking the extension. The reaction mixture was then subjected to amplification with the following sequence: 1 cycle at 50°C for 2 min (annealing stage), 1 cycle at 95°C for 10 min (denaturation stage) and 40 cycles at 95°C for 15 sec and at 60°C for 2 min (extension stage). In all cases, an initial denaturation was carried out at 95°C for 5 min in order to decrease the formation of a primer-dimer complex under these PCR conditions. The ABI-SDS was programmed to take R and Q fluorescent dye readings after each cycle at a temperature several degrees (60°C) lower than the melting temperature of the target amplicon. This step at 60°C was taken to avoid, or at least minimize, any potential contribution to the overall fluorescent dye signals from primer-dimer formation. A calibration curve was constructed by plotting the R/Q ratio against the amounts of hIDE and siRNA cDNA synthesized based on the RNA isolated from the livers of the transgenic and non-transgenic mice, which were used as the control.
The samples were then incubated with the mouse anti-IDE antibody diluted 1:1,000 on a PBS-blocking buffer. These antigen-antibody complexes were visualized with the biotinylated secondary antibody (goat anti-rabbit)-conjugated HRP streptavidin (Histostain-plus kit, Zymed) diluted 1:1,500 in a PBS blocking buffer. The IDE proteins were detected using a stable DAB (ResGen, Invitrogen Corp.) and an imaging densitometer (Model GS-690, Bio-Rad).

Statistical analysis. The tests for significance were performed using a one-way ANOVA test of variance (SPSS for Windows, release 10.10, standard version, Chicago, IL). All of the values are represented as the mean ± standard deviation (SD). The mean difference between dietary treatments was evaluated at p<0.05.

Results

A pH1/siRNA_inso insulin-CMV/hIDE system in transgenic mice. Insulin production was suppressed and insulin degradation was activated in the mice by introducing the H1/siRNA_inso insulin/PolIII fusion gene into the CMV/hIDE/SV40 fusion gene to produce the pH1/siRNA_inso insulin-CMV/hIDE dual expression system. The samples were then incubated with the mouse anti-IDE antibody diluted 1:1,000 on a PBS-blocking buffer. These antigen-antibody complexes were visualized with the biotinylated secondary antibody (goat anti-rabbit)-conjugated HRP streptavidin (Histostain-plus kit, Zymed) diluted 1:1,500 in a PBS blocking buffer. The IDE proteins were detected using a stable DAB (ResGen, Invitrogen Corp.) and an imaging densitometer (Model GS-690, Bio-Rad).
system (Fig. 1). The H1/siRNA insulin-CMV/hIDE fragment (6.214-kb), in which the prokaryotic sequence had been eliminated through the digestion of pH1/siRNA insulin-CMV/hIDE (7.2-kb) with BlnI, was used for the microinjection. Of a total of 34 offspring (15 males and 19 females), 1 male (#9381) and 2 females (#9384, #9385) of the first lineage founder mice possessed the H1/siRNA insulin-CMV/hIDE gene, which was identified by DNA-PCR (Fig. 2A). Also, the intensity and restriction patterns of the DNA obtained from the Southern blotting were compared among Tg mice using a 32P-labeled siRNA insulin-hIDE sequence in the Southern blot analysis (Fig. 2B). The Southern blot analysis revealed that the copy numbers of the integrated transgenes varied from line to line, ranging from 3 to 12 copies. The first lineage founder mouse (#9381) containing 6 copies of the H1/siRNA insulin-CMV/hIDE gene was selected for further study.

The founder mice containing siRNA insulin and hIDE were then backcrossed to the C57BL/6 parental strain in order to establish independent lines. The introduced siRNA insulin and hIDE gene in their genomes was transmitted to all offspring with approximately 50% hemizygotes in a Mendelian fashion.

Tissue-specific regulations of siRNA insulin and hIDE expression in the Tg mice. In order to determine if the regulation of the introduced siRNA insulin and hIDE genes were expressed under the control of the H1 and CMV promoter in all tissues, the transcriptional levels of IDE and siRNA insulin were measured by RT-PCR and real-time PCR analysis (Fig. 3). The expression patterns of IDE and siRNA insulin in various tissues were analyzed by RT-PCR (Fig. 3A). The β-actin signal was used as the control, and the transcript (650-bp) indicates the RNA loading. In addition, the RT-PCR products for IDE (340-bp) are indicated. The IDE expression in the Non-Tg and the Tg groups is shown. The transcript was quantified using a Kodak electrophoresis documentation and analysis system 120. (B) Real-time PCR analysis of hIDE expression. Calibration was constructed by plotting the R/Q ratio against the amounts of hIDE cDNA synthesized based on the RNA isolated from the tissue of the transgenic mice and from the tissue of the non-transgenic mice used as the control. The data represents the mean ± SEM from three replicates. *p<0.05, significant difference between treatments.
from various tissues including the brain, heart, lung, liver, kidney, intestine, and muscle of H1/siRNA\textsubscript{insulin}-CMV/hIDE Tg mice were examined by RT-PCR and real-time PCR analysis. In the RT-PCR analysis, the highest hIDE and siRNA\textsubscript{insulin} expression level was observed in the muscles, followed by the lung, brain, heart, liver, kidney, and intestine (Fig. 3A). Real-time PCR analysis showed a significantly higher level of the hIDE and siRNA\textsubscript{insulin} transcripts (RQ level) in the muscle and liver of the H1/siRNA\textsubscript{insulin}-CMV/hIDE-Tg mice (Fig. 3B). Furthermore, total IDE protein expression in the liver of the Tg mice was examined using Western blotting and immunostaining in order to determine if the CMV-controlled hIDE proteins from the livers of the Tg mice stably expressed the IDE proteins. As shown in Fig. 4A, the total IDE proteins (110 kDa) were detected by Western blotting using the specific antibody for mouse/human IDE proteins in the liver tissue of the H1/siRNA\textsubscript{insulin}-CMV/hIDE Tg mice. The level of the IDE proteins was significantly higher in the liver and pancreas of the Tg mice than of the Non-Tg mice. In contrast, a very low level of the endogenous mouse IDE protein was detected in the Non-Tg mice (Fig. 4A). The IDE immunoreactivity in the liver was next analyzed using optical microscopy. The immunostaining intensity in the H1/siRNA\textsubscript{insulin}-CMV/hIDE Tg mice compared to the Non-Tg mice, at x200 magnification. The data represent the mean ± SEM from three replicates. *p<0.05, significant difference between treatments.

Figure 4. The level of IDE protein expression in the liver and pancreas. (A) Expression of the total IDE protein according to Western blot analysis. The membranes were incubated with the antibodies for IDE and the β-actin protein from the liver cytosol fraction. These antibodies were recognized by the IDE protein originating from the mouse and the human. Three mice/group were assayed by Western blotting. The values are reported as a mean SD, p<0.05 vs control. (B) Immunostaining analysis of IDE expression. The expression profile of IDE in the liver and the pancreas at 20 weeks of age by immunostaining analysis. A high intensity was observed in the liver (B-c and B-d) and pancreas (B-a, and B-b) of the H1/siRNA\textsubscript{insulin}-CMV/hIDE Tg mice compared to the Non-Tg mice, at x200 magnification. The data represent the mean ± SEM from three replicates. *p<0.05, significant difference between treatments.
Post-prandial intraperitoneal glucose tolerance response in the Tg mice. The effect of the siRNA<sub>insulin</sub> and hIDE overexpression in all tissues of the Tg mice on the glucose tolerance response was determined by measuring the serum glucose level during the time course of an IPGTT in the H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice aged 20 weeks using a post-prandial intraperitoneal glucose tolerance test. The H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice showed significant impairment in glucose tolerance during the course of an IPGTT (Fig. 5). In particular, the Tg mice aged 20 weeks showed a greater impairment of glucose tolerance than their Non-Tg counterparts with a higher difference being observed in the male Tg mice (Fig. 5A). These results suggest that H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice at this age have a lower insulin activity.

Suppression of serum insulin in the Tg mice. In order to verify the suppression of insulin production by the overexpressed siRNA<sub>insulin</sub> and hIDE protein, the serum insulin level was assayed in the H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice using ELISA with the anti-insulin antibody. The serum insulin level was significantly lower in the H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice than in the Non-Tg mice (Fig. 6). Therefore, the overexpressed siRNA<sub>insulin</sub> and hIDE protein can effectively suppressed insulin production in the pancreas and stimulated insulin degradation in the liver of the Tg mice.

Changes in the proteins in the ER stress signal pathway. ER stress is significantly stimulated by glucose or nutrient deprivation, viral infection, lipids, increased synthesis of secretory proteins and expression of mutant or misfolded protein (27,28). To examine whether ER stress was increased in H1/siRNA<sub>insulin</sub>-CMV/hIDE Tg mice, the expression patterns of several molecular indicators of ER stress were detected using specific antibodies in the liver tissues from Tg mice. The level of p-eIF2<sup>α</sup> protein was significantly increased in the Tg mice compare to the Non-Tg mice, indicating that the Tg mice were involved in the phosphorylation of the p-eIF2<sup>α</sup> signaling system (Fig. 7). In addition, the Tg mice showed a significantly higher phosphorylation of JNK protein in the liver than Non-Tg mice. Therefore, these results suggest that decreased insulin and glucose impairment in the Tg mice can effectively induce the activation of JNK and eIF2<sup>α</sup> proteins in the ER stress signaling pathway.

Discussion

Type 1 diabetes mellitus, accounting for the early onset of diabetes in children, is characterized by the down-regulation of insulin secretion in the pancreas, lower circulating insulin levels, and subsequently reduced insulin action on glucose uptake in the peripheral tissues and the liver. With an overall aim of developing model animals for type 1 diabetes, this study examined whether or not the suppression by the transgene-based siRNA<sub>insulin</sub> sequence and the degradation as a result of the overexpression of the hIDE protein in Tg mice could contribute to a decrease in the level of circulating insulin and insulin action in vivo. Hasuwa et al (25) reported that transgenically supplied short duplex siRNA (21-23 nt) could silenced the ubiquitously overexpressed green fluorescent protein and simultaneously enhanced the expression of the red fluorescent protein in every tissue of Tg mice and rats. This suggests that this dual expression system may be an alternative method for the simultaneous gene disruption and overexpression in the same animals. In the present study, two
insulin related genes, insulin siRNA and hIDE, were introduced into this dual expression system with the aim of developing a new diabetes-related model. Most studies have demonstrated that IDE is the primary enzyme for initiating cellular insulin degradation. This protein has a major role in insulin-degradation in cell homogenates (29). The features of cellular insulin degradation including the effects of various modifiers (e.g. Ca++) and inhibitors (e.g. bacitracin and N-ethylmaleimide) are consistent with the properties of IDE. However, the successful expression of the human IDE gene in a transgenic animal has not been reported. In this study, the hIDE gene and insulin siRNA sequence were applied to this dual expression system for overexpression in transgenic mice. Furthermore, the 5 types of siRNA sequences with specific binding to the insulin sequence were synthesized using the siRNA target finder program (Ambion) and inserted into the pSilencer 3.1 vector containing the H1 promoter, which successfully regulated the siRNA sequence both in vitro and in vivo (30). The level of insulin in the cultured transfectants was significantly lower in the 1-1 and 2-2 clones of the pH1/siRNAinsulin-CMV/hIDE transfectants than in the pEGFPC-1 control transfectants (31).

The dual expression system containing siRNAinsulin under the control of the H1 promoter and the hIDE cDNA under the control of the CMV promoter was expressed successfully in all tissues of the H1/siRNAinsulin-CMV/hIDE Tg mice at the RNA level. In particular, the highest hIDE and insulin siRNA expression level was observed in the muscle, followed by the lung, brain, heart, liver, kidney, and intestine. There was also a significantly higher level of the total hIDE proteins in the liver in the Tg mice than in the Non-Tg mice used as the control, as indicated by Western blotting and immunostaining. Therefore, this dual expression system coexpressing the insulin siRNA and hIDE genes was successfully introduced into the genomic DNA of the Tg mice, which expressed the RNA and protein for down-regulating the insulin level in the Tg mice for the first time.

An impaired glucose tolerance and β-cell function are widely recognized as features of type 1 diabetes (32,33). The animal models used for diabetic study were the Non-obese diabetic (NOD) mouse and the bio-breeding rat (34). The NOD mice discovered in Japan in the late 1970's (35) were inbred and distributed worldwide to establish numerous animal models. However, in order to understand type I diabetes mellitus, the physiological characteristics of the Tg mice need to be determined because these Tg mice may be used to examine the entire insulin-related metabolism in comparison with chemically-induced diabetic model animals.

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