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Identification and Characterization of a Gene Regulating Enzymatic Glycosylation which is Induced by Diabetes and Hyperglycemia Specifically in Rat Cardiac Tissue

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Abstract

Primary cardiac abnormalities have been frequently reported in patients with diabetes probably due to metabolic consequences of the disease. Approximately 2,000 mRNA species from the heart of streptozotocin-induced diabetic and control rats were compared by the mRNA differential display method, two of eight candidate clones thus isolated (DH1 and 13) were confirmed by Northern blot analysis. The expression of clone 13 was increased in the heart by 3.5-fold ($P < 0.05$) and decreased in the aorta by twofold ($P < 0.05$) in diabetes as compared to control. Sequence analysis showed that clone 13 is a rat mitochondrial gene. DH1 was predominantly expressed in the heart with an expression level 6.8-fold higher in the diabetic rats than in control ($P < 0.001$). Insulin treatment significantly ($P < 0.001$) normalized the expression of DH1 in the hearts of diabetic rats. DH1 expression was observed in cultured rat cardiomyocytes, but not in aortic smooth muscle cells or in cardiac derived fibroblasts. The expression in cardiomyocytes was regulated by insulin and glucose concentration of culture media. The full length cDNA of DH1 had a single open-reading frame with 85 and 92% amino acid identity to human and mouse UDP-GlcNAc:Galβ1-3GalNAcαβ1-6 N-acetylgalcosaminyltransferase (core 2 GlcNAc-T), respectively, a key enzyme determining the structure of O-linked glycosylation. Transect RNA expression of DH1 cDNA into Cos7 cells conferred core 2 GlcNAc-T enzyme activity. In vivo, core 2 GlcNAc-T activity was increased by 82% ($P < 0.05$) in diabetic hearts vs controls, while the enzymes GlcNAc-TI and GlcNAc-TV responsible for N-linked glycosylation were unchanged. These results suggest that core 2 GlcNAc-T is specifically induced in the heart by diabetes or hyperglycemia. The induction of this enzyme may be responsible for the increase in the deposition of glycoconjugates and the abnormal functions found in the hearts of diabetic rats. (J. Clin. Invest. 1995. 96:1759–1767.) Key words: glycosylation \cdot cardiomyopathy \cdot cardiomyocytes \cdot diabetic complications \cdot gene regulation

Introduction

Cardiovascular diseases are the major cause of morbidity and mortality in diabetic patients, involving cardiac tissues as well as large vessels in the brain, heart, and lower extremities (1). In the heart, the majority of the cardiac failure is probably due to atherosclerotic processes in the coronary vessels, but multiple studies have documented that a sizeable number of diabetic patients suffer from congestive heart failure without significant coronary disease (2, 3). In addition, type 1 diabetic patients with $< 5$ yr of disease have been reported to have abnormal cardiac function in the absence of significant coronary vessel disease (4). These clinical findings are supported further by animal studies documenting biochemical and functional changes in the cardiac tissue shortly after induction of diabetes (5–8). From these results, it has been postulated that diabetes mellitus and its metabolic sequelae can induce a specific form of cardiomyopathy (8, 9).

As with other chronic complications of diabetes, the cardiovascular changes once established are difficult to reverse, both in clinical and experimental settings (10–12). Most cardiovascular abnormalities are metabolically induced, with a great deal of interest directed towards identifying alterations in gene expression induced by diabetes or hyperglycemia in the vascular tree. Since thickening of basement membrane is a classical finding in diabetic microvasculature (10), many of the studies concerning glucose-regulated genes have primarily focused on changes in the basement matrix components using cultured vascular cells (13, 14). In contrast, we investigated the effect of diabetes on gene expression in cardiac tissue. We adapted mRNA differential display (developed by Liang and Pardee to identify novel oncogenes in tumor cells [15, 16]), to identify changes in gene expression induced by metabolic stimuli such as hyperglycemia or diabetes (17, 18). We have succeeded in characterizing several clones which are differentially expressed in the ventricles of streptozotocin (STZ)$^3$ diabetic rats. One clone (DH1) was sequenced, identified, and studied further since its genetic expression in cardiomyocytes was regulated by hyperglycemia and insulin.

Methods

Animals. Male Sprague-Dawley rats (Taconic Farms, Inc. German Town, NY) weighing 180–200 grams were injected by the intraperitoneal route with STZ (80 mg/kg of body weight). STZ was dissolved

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1. Abbreviations used in this paper: Core 2 GlcNAc-T, UDP-GlcNAc:Galβ1-3GalNAcαβ1-6 N-acetylgalcosaminyltransferase; NOD, nonobese diabetic; PAS, periodic acid Schiff; STZ, streptozotocin.
in 20 mM citrate buffer (pH 4.5) immediately before use. Cardiac tissue from spontaneous autoimmune-caused diabetic nonobese diabetic (NOD) mice and their control littermates were graciously provided by Dr. Masakazu Hattori of the Joslin Diabetes Center (Boston, MA) (19). Onset of diabetes was determined by the detection of urinary glucose and confirmed by blood glucose levels. Insulin pellets (Linshin, Scarborough, Canada) were implanted subcutaneously 1 wk after STZ injection in four STZ-diabetic rats for another week to normalize blood glucose levels. The mean plasma glucose level in insulin-treated STZ-diabetic rats was not significantly different from control animals (4.8±0.5 vs 5.6±0.7 mM), while, the mean body weight in the insulin-treated STZ-diabetic rats remained significantly less than control (270±17 vs 326±17 grams respectively, P < 0.01) but significantly greater than diabetic rats without insulin treatment (210±33 grams), (P < 0.01).

mRNA differential display. Rats were killed 2 wk after onset of diabetes. Cardiac ventricles and thoracic aorta were dissected and washed with ice-cold PBS, immediately frozen in liquid N₂, then crushed into frozen powder. Pieces of aorta from three rats were combined into one sample. Total RNA was extracted using Ultraspec RNA isolation system (Biotec Laboratories, Houston, TX). mRNA differential display was performed as previously reported (17, 18). Briefly, DNA-free RNA was obtained by treatment with DNase I (GIBCO BRL, Grand Island, NY) in the presence of placental RNase inhibitor (GIBCO BRL) for 30 min at 37°C. After phenol/chloroform extraction and ethanol precipitation, two reverse transcriptions were performed for each RNA sample using 0.2 μg DNA-free total RNA in 1× reverse transcription buffer (PCR buffer) containing 10 mM DTT, 20 μM each of dGTP, dATP, dTTP, and dCTP, and 1 μM of either T₂₀NG or T₂₀NC oligonucleotide (Midland Certified Reagent Co., Midland, TX) where N is threofold degenerate for G, A, and C. The solution was heated at 65°C for 5 min and cooled to 37°C, then superscript reverse transcriptase (20 U) (GIBCO BRL) was added for 1 h. PCR was performed in reaction mixtures containing 0.1 vol of reverse transcription reaction mixture, 1× PCR buffer, 2 μM each of dGTP, dATP, dTTP, and dCTP, 10 μCi [α-³²S]-dATP, 1 μM of the respective T₂₀NX oligonucleotide, 0.2 μM of 20 different specific arbitrary-10-mer oligonucleotides (OP-ERON Technologies Inc., Almeda, CA) and 10 U of AmpliTaq DNA polymerase (Perkin-Elmer Cetus Corp. Norwalk, CT). The PCR reactions were initiated at 95°C for 1 min, amplified 40 cycles at 94°C for 45 s, 40°C for 90 s, 72°C for 30 s, and finished at 72°C for 15 min. DNA sequencing stop buffer (U.S. Biochemical, Inc., Cleveland, OH) was added to samples which were heated at 80°C for 2 min before loading on a 6% polyacrylamide sequencing gel (National Diagnostics, Atlanta, GA). After electrophoresis, the gels were exposed to XAR-5 film (Eastman Kodak Co., Rochester NY) for 48 h. Bands evident under one gelicmy condition and absent in the other were identified and the PCR repeated to confirm the findings.

Band recovery and Northern blot analysis. Bands reproducibly exhibiting significant differences in expression were cut out and DNA was eluted by boiling in 10 mM Tris HCl and 1 mM EDTA solution for 30 min. After ethanol precipitation, the DNA was reamplified by PCR using appropriate primers and conditions described above except for dNTP concentrations of 20 μM and no radiolabel. Products were visualized on 2% agarose gels, eluted, and used as probes for Northern blot analysis or subcloned. Total RNA (20–25 μg) was fractionated by denaturing 1% formaldehyde agarose gel electrophoresis and transferred to Biotrans nylon membrane (ICN, Irvine, CA). 32P-labeled probes prepared by random priming using a commercially available kit (Amer sham Corp., Arlington Heights, IL) were hybridized to UV cross-linked blots in 0.1 M Pipes, 0.2 M NaPO₄, 0.1 M NaCl, 1 mM EDTA, 5% SDS, and 60 μg/ml salmon sperm DNA at 65°C and washed in 0.5 × SSC, 5% SDS at 65°C for over 1 h. mRNA expression was quantified using a phosphorImager and standardized volume integration with the accompanying ImageQuant Analyzing Software version 3.3 (Molecular Dynamics, Sunnyvale, CA) and loading differences were normalized using 36B4 as standard CDNA probe (17, 18).

DNA sequencing. Samples showing significant changes by Northern blot analysis were subcloned using the TA Cloning Kit (Invitrogen Corp., San Diego, CA). After the subcloned inserts were checked by Northern blot analysis, DNA sequencing was performed using commercially available Sequenase version 2.0 kit (U.S. Biochemical, Inc.,). Gene database searches were performed at the National Center for Biotechnology Information (NCBI) using the BLAST network service.

Construction of the diabetic heart cDNA library. Poly(A) + RNA was isolated from the total cellular RNA extracted from heart of diabetic rats using an oligo-dT cellulose column (Pharmacia LKB Biotechnology Piscataway, NJ) as previously described (17). CDNA was prepared and ligated into the EcoRI sites of Lambda gt10 (Stratagene, Inc., La Jolla, CA) by standard methods (20). After packaging the DNA, Escherichia coli (C600) was infected with the plaque and plated on a P150 plate yielding about 5 × 10⁴ independent plaque-forming units. Plaques were lifted onto nitrocellulose (Schleicher & Schuell, Inc., Keene, NH) and cross-linked to the membrane by ultraviolet light. 20 P150 plates were screened, which provided 1 × 10⁶ plaque-forming units for screening.

Screening the cDNA library. A 214-bp cDNA (DH1) probe obtained from differential display was hybridized to the cDNA library by standard methods at 44°C then washed at 56°C (21). After screening 1 × 10⁶ plaque-forming units, a positive cDNA insert was isolated and subcloned into pBluescript (Stratagene, Inc.). For sequencing, the inserts were restricted using BamHI and EcoRI and subcloned into pBluescript. cDNA cloning of mouse UDP-GlcNAc:Galβ1-3GalNAcR β1-6 N-acetylgalcosaminyltransferase (core 2 GlcNAc-T). Approximately 2 × 10⁷ colonies of a cDNA library prepared in pCDMB (Invitrogen Corp.) using poly A+ RNA from D33W25, a murine lymphoid tumor cell line (22), were screened by colony hybridization (23) to a 864-bp EcoRI–BamHI subcloned fragment of human core 2 GlcNAc-T isolated by PCR (gift of Dr. A. Datti, Perugia, Italy) corresponding to amino acids 85–372 of the human enzyme. Hybridization was performed overnight at 65°C in 500 mM sodium phosphate pH 7.2, 7% SDS, 1% BSA, 1 mM EDTA. After rinsing, filters were washed at 65°C in 100 mM sodium phosphate, 0.1% SDS. After three rounds of hybridization and purification, two clones, designated C2-251 and C2-352, were isolated and gave specific and strongly positive signals on Southern blots hybridized with the probe. The cDNA inserts were subcloned as Xhol fragments into SalI cut pGEM5zf (+) (Promega Corp., Madison, WI) and a series of exonuclease III-mung bean nuclease (GIBCO BRL)–nests deletions generated from each end. DNA sequencing was performed using the Autoread sequencing kit and the AFL DNA sequence primer according to the manufacturer’s instructions (Pharmacia LKB Biotechnol). Some sequences were also generated using internal primers. Data were analyzed and edited using the UWCGC suite.

Transient expression of DHI in Cos 7 cells. A cDNA insert containing the full open reading frame of DHI was isolated with Xhol and EcoRV and subcloned into pcDNA1/amp (Invitrogen Corp.). The plasmid was purified by double CsCl ultracentrifugation followed by phenol/chloroform extraction (21), then DNA (0.5 μg) was transfected into nonconfluent Cos 7 cells cultured on P100 dishes using 20 μg of Lipofectin for 16 h at 37°C (GIBCO BRL) (24). Cells were harvested 48 h later and used for measurement of core 2 GlcNAc transferase activity.

Preparation of cardiomyocytes. Cardiomyocytes were prepared by collagenase digestion as described before (25). Briefly, hearts were excised and perfused through the aorta with Krebs-Henseleit bicarbonate buffer containing 5.5 mM glucose and 2.5 mM calcium. The perfused medium was switched to the same buffer without calcium to stop contraction, followed by a final perfusion with Krebs-Henseleit buffer containing 50 μM calcium, 0.1% BSA, 312 U/ml hyaluronidase (Worthington, Freehold, NJ) and 0.1% collagenase (Worthington). Ventricular tissue was dissociated by shaking in Krebs-Henseleit buffer containing 50 μM calcium, 0.2% BSA, 312 U/ml hyaluronidase, and 0.1% collagenase. The cells were allowed to settle under gravity and were washed twice in the Krebs-Henseleit buffer containing 100 μM calcium and 0.5% BSA before resuspension in minimal essential medium containing Earle’s salts, 26 mM sodium bicarbonate, 5 mM creatine, 20 mM Hepes, 100 U/ml penicillin G, 100 μg/ml streptomycin, and 1.8 mM calcium. The cells were seeded onto laminin-coated dishes and maintained in a
37°C humidified 95% air-5% CO₂ incubator. All cells were allowed to equilibrate for 2 h then washed and refed with the same media containing 0.24% glucose and 10^{-7} M insulin for 3 d with daily changes of media.

**Measurement of core 2 GlcNAc-T activity.** Transfected Cos-7 cells were washed in PBS, frozen, thawed, and lysed in 0.9% NaCl. 0.4% Triton X-100 at 0°C. PBS-rinsed, fresh frozen rat hearts were rinsed again in PBS and homogenized using a polytron in 0.9% NaCl, 0.4% Triton X-100, 0.1 mM PMSF, 0.1% Trasylol at 0°C. The core 2 GlcNAc transferase reactions contained 50 mM 2-[(N-morpholino)ethanesulfonic acid (MES) pH 7.0, 1 mM UDP-GlcNAc, 0.5 μCi UDP-6²H]-N-acetylglucosamine (16,000 dpm/nmol, New England Nuclear, Boston, MA), 0.1 M GlcNAc, 1 mM of Galβ1-3GalNAca-pNP (Toronto Research Chemicals, Toronto, Canada) as substrate, and 16 μl of cell lysate (8–12 mg/ml of protein) for a final volume of 32 μl (26, 27). The GlcNAc-TV reactions were the same except that Triton X-100 was added to a final concentration of 1%, and 1 mM of GlcNAcβ1-2Manα1-6Manβ1-O(CH₂)₅COOCH₃ (gift of Dr. O. Hindsdaul, University of Alberta, Edmonton, Canada), was used as acceptor (28). The GlcNAcTT reactions were the same as GlcNAc-TV but with the addition of 10 mM MnCl₂ and 1 mM Manα1-3-Manα1-6-Manβ1-O(CH₂)₅COOCH₃ as used as acceptor (29). Reactions were incubated for either 1 or 2 h at 37°C, then diluted to 5 ml in H₂O and applied to a C₈ Sep-Pak (Millipore Corp., Bedford, MA) in H₂O, washed with 20 ml H₂O. The products were eluted with 5 ml of methanol and radioactivity was counted in a liquid scintillation β-counter. Endogenous enzymatic activity was measured in the absence of acceptor and subtracted from values determined in the presence of added acceptor.

**Statistical analysis.** Differences in signal intensity between controls and diabetic animals were expressed as percentage of controls. Because percentages tend to deviate from normal distribution, mean and standard error were calculated after transformation of data to logarithmic values and data were expressed as mean (±SE range). Statistical analysis (Student’s t test) was performed using the logarithmic values.

**Results**

**Differential display.** The expression of mRNA species derived from the cardiac ventricles of diabetic and control rats was compared using mRNA differential display. Approximately 2,000, presumably different, mRNA species were screened in this study using 40 primer sets. As exemplified by Fig. 1 A, eight candidates appeared differentially expressed when ventricular tissue from control and diabetic rats was compared; five increased and three decreased their expression in the diabetic state. These changes were confirmed by repetition at least twice using different preparations of total RNA.

**Northern blot analysis.** Signals from all the candidates were detectable by Northern blot analysis using total RNA preparations. As shown in Fig. 1 B, two of the eight candidate species showed significant and reproducible changes in diabetic rats compared to controls. Fig. 2 demonstrates that in rats diabetic for 2 wk, the level of DH1 (5.0 kb) increased to 680% (580–790%, n = 8, P < 0.001) of control in the heart but was not detectable in the aorta. Furthermore, in rats with diabetes for just 1 wk, the mRNA level of DH1 in the heart had already significantly increased to 410% of control levels. A significant increase of DH1 in the heart of diabetic rats was observed even after 4 wk of disease. The expression of clone 13 mRNA increased to 350% (260–470%, n = 3, P < 0.05) of control in the heart and decreased to 43% (38–49%, n = 3, P < 0.05) of control in the aorta.

**Sequence analysis of DH1 and clone 13.** The nucleotide sequences of cDNA fragments of DH1 and clone 13 derived from differential display were determined. Both had flanking primer sequences identical to those used in the differential display. Searching the national gene databases (GenBank/EMBL) revealed that clone 13 had 99% identity to the Wistar rat mitochondrial 16S ribosomal RNA gene (30) while DH1, which was 214 bp in size, did not reveal any homology to previously reported sequences.

**Cloning of full-length DH1 cDNA.** To facilitate identification, a cDNA library derived from diabetic rat heart mRNA was screened using the 214-bp-clone DH1 PCR fragment as a probe. Five overlapping recombinants were identified and the composite of the full cDNA sequence was determined (Fig. 3 A). It contained 5,010 bp inclusive of poly A tail and corresponded to the size detected by the original Northern blot analysis. Open reading frame analysis showed that the longest possible coding region which was from position 802 to 2085 and encoded 428 amino acids. The GXXATGC pattern was observed flanking the region of the presumptive start codon (31).
### Figure 3. Nucleotide and deduced amino acid sequences of DH1. (A), The full-length nucleotide sequence of DH1 is shown. The sequence is numbered relative to presumed start codon. The longest open reading frame (underlined) and putative polyadenylation signal (dashed) are noted. (B), The deduced amino acid sequence of DH1 is compared with mouse and human (32) core 2 GlcNAc-T. The sequences are numbered from the presumed initiation methionine. Dashes represent identical sequences. The putative signal/membrane-anchoring domain is underlined.
and a polyadenylation signal, AATAAAA, was found 15 bp upstream from the polyA tail. Searches for homologous sequences in Genbank/EMBL revealed that this cDNA shared 80% identity at the nucleotide level and 85% identity at the amino acid level with human core 2 (GlcNAc-T) (32). We also cloned and sequenced the mouse core 2 GlcNAc-T and found that DH1 shared 92% identity with the amino acid sequence of mouse core 2 GlcNAc-T (Fig 3 B). These findings strongly suggested that DH1 was rat core 2 GlcNAc-T.

Characterization of DH1 expression in the NOD mouse. To check that increased expression of DH1 was diabetes specific and not due to other effects of streptozotocin, DH1 expression in the hearts of spontaneous autoimmune-caused diabetic NOD mice was measured. As shown in Fig. 4, DH1 hybridizing signals were detected by Northern blot analysis at 6.0, 4.6, and 1.9 kb from animals which had experienced 2–3 wk of hyperglycemia and diabetes. The 4.6 and 6.0 kb bands in hearts from diabetic NOD mice increased to 560% of control animals.

Effects of insulin on DH1 expression in diabetic rats. After 1 wk of diabetes, four rats were treated with insulin for an additional week. Blood glucose level was normalized from 24.7 to 4.8 mM (P < 0.01). The Northern blot analysis shown in Fig. 5, demonstrated that cardiac expression of DH1 in rats diabetic for 2 wk increased to 680% of control levels, consistent with earlier data, whereas insulin treatment normalized the expression of the DH1 to 169% (134–214% P < 0.001 vs 2-wk diabetic rats) of control levels.

Tissue distribution of DH1. Fig. 6 A shows a representative Northern blot analysis of DH1 expression using total cellular RNA isolated from tissues of control and diabetic rats. Relative signal intensity was calculated using the 36B4 signal for normalization and is shown in Fig. 6 B. In normal rats, DH1 transcripts were relatively high in the brain, kidney and liver and low in the heart, aorta, lungs, and skeletal (soleus) muscle. A significant and cardiac specific increase (6.8-fold) in the expression of DH1 mRNA was observed in diabetic animals.

The expression of DH1 in cultured cells. Cultured cardiomyocytes were measured to determine whether they could be the source of the increased expression of DH1. As shown in Fig. 7 A, DH1 hybridizing signals were detected by Northern blot analysis of cultured cardiomyocytes at the same mobility as the mRNA from heart tissue (5.0 kb). However, DH1 expression was not detectable in fibroblasts cultured from rat heart or rat aortic smooth muscle cells even when using 25 μg of total cellular RNA. Furthermore, in cardiomyocytes, cultures elevating media glucose concentrations from 5.5 to 22 mM increased the expression of the DH1 by 78% (54–106%, P < 0.05) while insulin (10−7M) decreased the expression by 53% (40–62%, P < 0.05) of control levels as shown in Fig. 7 B.

Core 2 GlcNAc-T activity in cells transiently transfected with DH1. Although Cos 7 cells lipofected with pcDNAI/amp showed significant endogenous core 2 GlcNAc-T activity of 1.71±0.27 nmol/mg per h (n = 3), cells transfected with the expression vector containing a full-length cDNA for DH1 in correct orientation had 3.85±1.6 nmol/mg per h (P < 0.05, n = 3). The assay is specific for core 2 GlcNAc-T, as confirmed by analysis of the reaction product Galβ1-[3(GlcNAcβ1-6)-GalNAcα-pNp] by 1 H-NMR (Nuclear Magnetic Resonance) and HPLC which had been reported in previous studies (28). Therefore, DH-1 encodes an enzymatically active core-2 GlcNAc-T.

GlcNAc-T activity in heart. With the identification of DH1 being an enzyme involved in mediating the biosynthesis of O-linked sugar chains, we tested the specificity of the diabetes effect by measuring the activities of core 2 GlcNAc-T (which branches maturing O-linked sugar chains) and two other GlcNAc transferases (which are specific for branching N-linked sugar chains) in the hearts of control and diabetic rats (Fig. 8). Core 2 GlcNAc-T activity increased significantly and specifically in diabetic hearts by 82% of control (0.39±0.03 vs 0.71±0.10 nmol/h per mg protein n = 3, P < 0.05). In contrast, GlcNAc-TI and GlcNAc-TV activities were not significantly different between control and diabetes (GlcNAc-TI:1.05±0.11 vs 0.79±0.09 nmol/h per mg protein, GlcNAc-TV: 0.078±0.024 vs 0.077±0.023 nmol/h per mg protein). The changes thus seem restricted to O-glycosylation.

Discussion

Multiple studies have reported metabolic and functional alterations in the heart of animals with a short duration of diabetes (6, 7). With chronic duration of diabetes, structural changes in the vasculature and increased carbohydrate content of the heart, denoted by an increase in periodic acid-Schiff (PAS) staining...
are indicative of less reversible alterations (5, 8, 9, 33). Changes in the expression of glucose transporter (GLUT-4) (34), Ca²⁺ ATPase (35), and alpha- and beta-adrenergic receptors (7) have been reported in the diabetic state and have been suggested as possible candidates causing acute cardiac dysfunctions. However, it is unlikely that these changes are directly responsible for the chronic and structural changes since their biological actions are transient and their relationship to the in-

Figure 6. Tissue distribution of gene expression of DH1 is shown. (A), Northern blot analysis was performed using 20 μg total cellular RNA isolated from aorta, brain, heart, kidney, liver, soleus muscle, and lung of control (C) and 2-wk diabetic (D) rats. DH1 signals are indicated by an arrow. The same blot was reprobed with 36B4 cDNA as a loading control. (B) Quantitation of changes induced by diabetes mRNA level for DH1 in aorta, brain, heart, kidney, liver, lung, and skeletal muscle. Data are expressed relative to the message level of control rat heart. Closed bars represent control (C) and hatched bars represent 2-wk diabetes (DM). Results are shown as mean±SE of three experiments using different animals. * P < 0.05 as compared with control.

Figure 7. Northern blot analysis of DH1 mRNA. (A) Total cellular RNA from cultured cardiomyocytes (10 μg) for lanes 1–3, fibroblasts (25 μg) for lane 4, and rat aortic smooth muscle cells (25 μg) for lane 5 were used. Cardiomyocytes were cultured as described in Methods with 5.5 mM glucose without insulin (C), 22 mM glucose without insulin (HG), or 5.5 mM glucose with 100 nM insulin (I) for 3 d. DH1 signals are indicated by an arrow. The same blot was reprobed with 36B4 cDNA as a loading control. (B) Quantitation of changes induced by 22 mM glucose or 100 nM insulin for DH1 expression in cultured cardiomyocytes. Data are expressed as percentage of control. Closed bar represents the results of cells cultured with 22 mM glucose and hatched bar represents the result of cells cultured with 100 nM insulin. Results are shown as mean±SE of four independent experiments. * P < 0.05 as compared with control.
increase in PAS staining is not obvious. To discover new candidate genes which may be responsible for causing diabetic cardiac dysfunction, we have used mRNA differential display to screen for alteration of gene expression in the cardiac ventricles of diabetic rats.

In the present study, eight possible candidate genes were suggested by the initial screening of ~2,000 mRNA species and two of the eight showed reproducible differences in expression by Northern blot analysis, a rate of confirmed positives similar to that reported in previous studies (17, 18). Genes previously identified as altered in the diabetic state such as GLUT-4 (34) and Ca\textsuperscript{2+} ATPase (35) mRNA were not observed in this study, however, this may be due to several potential limitations of the differential display method such as the insensitivity of this method to quantitate small changes in gene expression as compared with Northern blot analysis and differences in primer efficiencies. Alternatively, it is also possible that not all the potential mRNA species have been compared since only ~2,000 were analyzed. Previous studies using this method to identify glucose regulated genes in the vascular cells showed that genes with as little as twofold change in mRNA and consequent protein levels can be detected (18).

The sequence of clone 13 was identified as 99% identity to mitochondria 16S ribosomal RNA (30). At present, the significance of a change in mitochondrial ribosomal RNA to the development of cardiovascular abnormalities in diabetes is not clear. However, mitochondrial swelling has been observed in early stage of diabetic rats and preceded the abnormalities in cardiac function and histology which occurred after 4–8 wk and 12–24 wk of diabetes, respectively (36–38). Altered gene expression in the mitochondria could be due to changes in glucose metabolism and rates of oxidative phosphorylation in the cardiomyocytes of diabetic rats (39). Since all mitochondrial genes are regulated in a similar manner with a common promoter region (40), this finding would suggest that diabetes should affect them all equally and may have significant effect on mitochondrial functions, which could possibly decrease cardiac functions.

Clone DH1 encoded an open reading frame and analysis of its complete cDNA sequence inferred polypeptide showed 85% identity with human core 2 GlcNAc-T and 92% identity to the murine enzyme. The high degree of homology between DH1 and mouse core 2 GlcNAc-T is similar to that observed for other glycosyltransferase homologues of rat and mouse (41). Furthermore, the identity of the DH1 cDNA was confirmed by the transient transfection in Cos-7 cells which produced core 2 GlcNAc-T activity. Northern blot analysis showed three species of messages (5, 4.7, and 2 kb) similar to that previously reported for core 2 GlcNAc-T in human lymphoid cells (32). Since preliminary data from our laboratory has shown that the coding region of the gene at the genomic level is uninterrupted, suggesting that alternatingly spliced 5' or 3' untranslatable regions account for the multiple messages. Three polyadenylation sequences were identified and correspond to the message sizes observed by Northern blot analysis. The 100 nucleotides preceding the most distal poly A adenylation signal were conserved in mouse and rat sequences. The unusually long 3' non-coding region may affect message half life as observed in other systems (42). The finding of multiple sizes of messages in the mouse heart is consistent with the finding of multiple possible termination sites.

Since the antibody to rat core 2 GlcNAc-T is not available at the present time, it is not possible to determine whether the increase in the gene expression of this enzyme is translated into protein level directly. However, the finding that core 2 GlcNAc-T activity was doubled in hearts from diabetic rats compared to control animals supports this suggestion. Core 2 GlcNAc-T is a key enzyme in the O-linked glycosylation pathway which utilizes UDP-GlcNAc and substitutes Galβ1-3GalNAc-R at the 6 position of GalNAc. This results in Galβ1–3[Galβ1–6] GalNAc-R, a branched oligosaccharide which is commonly extended with polylactosamine chains and Lewis antigens. The latter sequences on O-linked oligosaccharides participate in leukocyte adhesion and homing, and are deficient in pathologic conditions, where core 2-GlcNAc-transferase expression is suppressed, such as leukemia (43), AIDS (44), and Wiskott-Aldrich syndrome (45). Furthermore, Datti et al. reported that this enzyme activity was induced by butyrate differentiation of Chinese hamster ovary cells, following an increase in intracellular cAMP level (27). Although it is presently unclear whether increased core 2 GlcNAc-T activity causes pathology in the heart of diabetic rats, it is possible that the additional branches and their sialylation results in the accumulation of PAS-positive material commonly observed in diabetic hearts. In fact, Reagan et al. and other investigators have emphasized that the accumulation of PAS-positive material in the interstitium opposing the surface of the cardiac fibers may compromise cardiac function (5). Further study with a longer duration of diabetes will be needed to determine the relationship between the increase of PAS-positive material and induction of core 2 GlcNAc-T in the cardiac tissue.

The increase in the expression of core 2 GlcNAc-T (DH1) by diabetes was very tissue specific involving only the heart among the various tissues studied. Similarly, core 2 GlcNAc-T message was observed in cultured cardiomyocytes but not in fibroblasts derived from cardiac tissue or aortic smooth muscle cells. The cardiac specificity of core 2 GlcNAc-T induction by diabetes and glucose contrasts with previous reports on the induction of extracellular matrix genes such as type I or IV collagen, where the changes were similar in all tissues examined including retina (46), kidney (47), and many types of cultured cells (13, 14). Since the clinical patterns of cardiomyopathy appear to be distinct from other diabetic vascularopathies, our
findings of tissue-specific changes in the gene expression suggest that the promoter of DH1 may contain cardiac-specific core 2 GlcNAc-T promoter elements which can be modulated by metabolic factors such as hyperglycemia and insulin.

Several observations from our study strongly suggest a role for metabolic regulation of core 2 GlcNAc-T in diabetic conditions. First, the up-regulation of core 2 GlcNAc-T is not due to streptozotocin or limited to rats since the same finding was also observed in the NOD mouse. Second, hyperglycemia regulates this enzyme directly. These findings suggest that core 2 GlcNAc-T is involved in diabetic cardiovascular complication and also shows that enzymes involved in glycosylation can be regulated by hyperglycemia. Further studies are needed to delineate the mechanism by which hyperglycemia can regulate core 2 GlcNAc-T expression. One possible mechanism is the activation by the diacylglycerol-protein kinase C (DAG-PKC) pathway since previous studies have reported that hyperglycemia and diabetes increased DAG levels and activated PKC in the heart of diabetic animals including the rat (13), and phorbol ester, a PKC agonist, has been reported to regulate core 2 transerase activity (27). Cloning and sequencing of the core 2 GlcNAc-T promoter regions is in progress to clarify this particular question.

In addition to changes directly related to hyperglycemia, insulin also decreased core 2 GlcNAc-T gene expression both in vivo and in cultured cardiomyocytes, suggesting that insulin may modulate its expression both directly and by improving glycemic control. It is not surprising that insulin was shown to affect gene expression in the heart since multiple studies have documented that myocardium is responsive to insulin at biochemical and physiological levels (34, 48). Clinically, insulin treatment has been reported to improve cardiac function in diabetic patients (49). At the molecular level, a great deal of information is available on insulin’s action on gene regulation, not specifically on cardiomyocytes (50). Therefore, further studies in the sequence of the 5’ region may provide information whether it contains the various insulin responsive elements which have already been reported (50). Similarly, other hormones and cytokines such as IGFs (51) and thyroxine (52), which are known to affect cardiac function, may also affect core 2 GlcNAc-T expression and will need to be characterized.

In summary, we have identified two novel molecular alterations in the myocardia of diabetic rats at the level of gene expression. One of these genes has been identified as core 2 GlcNAc-T, and provides strong evidence that hyperglycemia and insulin can regulate enzymes involved in the glycosylation of proteins, specifically in the heart. Thus, we have identified a possible mechanism for the pathology of increased carbonylates in the myocardium of diabetic rats and patients, which in turn could affect cardiac function.

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