Preservation of Insulin mRNA Levels and Insulin Secretion in HIT Cells by Avoidance of Chronic Exposure to High Glucose Concentrations

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Abstract

Glucose toxicity of the pancreatic beta cell is considered to play a secondary role in the pathogenesis of type II diabetes mellitus. To gain insights into possible mechanisms of action of glucose toxicity, we designed studies to assess whether the loss of insulin secretion associated with serial passages of HIT-T15 cells might be caused by chronic exposure to high glucose levels since these cells are routinely cultured in media containing supramaximal stimulatory concentrations of glucose. We found that late passages of HIT cells serially cultured in media containing 11.1 mM glucose lost insulin responsivity and had greatly diminished levels of insulin content and insulin mRNA. In marked contrast, late passages of HIT cells cultured serially in media containing 0.8 mM glucose retained insulin mRNA, insulin content, and insulin responsivity to glucose in static incubations and during perfusion with glucose. No insulin gene mutation or alteration of levels of GLUT-2 were found in late passages of HIT cells cultured with media containing 11.1 mM glucose. These data uniquely indicate that loss of beta cell function in HIT cells passed serially under high glucose conditions is caused by loss of insulin mRNA, insulin content, and insulin secretion and is preventable by culturing HIT cells under low glucose conditions. This strongly suggests potential genetic mechanisms of action for glucose toxicity of beta cells. (J. Clin. Invest. 1992. 90:320–325.) Key words: glucose toxicity • insulin gene • diabetes mellitus

Introduction

The thesis that glucose toxicity plays a secondary role in the pathogenesis of type II diabetes mellitus maintains that after hyperglycemia is established, glucose itself in high concentrations has deleterious effects on pancreatic islet beta cell function (1–18). Type II diabetes is characterized by a specific lack of normal recognition of glucose signals, since all other insulin secretagogues (except glucose) elicit first phase insulin responses when injected intravenously (19–22). Type I diabetes is markedly different in this regard because it is caused by beta cell death and complete absence of insulin secretion. In support of the glucose toxicity concept, defective glucose-induced insulinen secretion in type II diabetic patients can be partially reversed by an infusion of exogenous insulin to lower circulating glucose levels into the normal range (1, 5). This suggests that hyperglycemia is not only an effect of type II diabetes, but may be a contributing cause of progressive deterioration of insulin secretion. However, as yet no biochemical mechanism for glucose toxicity has been identified (15).

The HIT-T15 cell is a clonal cell line of pancreatic islet beta cells derived from SV-40 transfection of Syrian hamster pancreatic islets (23). The HIT cell is a valuable research tool because it is one of only a few cell lines that secrete insulin in response to glucose and because it responds to physiologic inhibitors of insulin secretion (23–31). However, insulin content and insulin secretion by this cell line progressively diminish with serial passage (23, 26, 30). The explanation for this loss of secretion has not been forthcoming, although further transformation of the cell is often mentioned as a possibility by investigators who study this cell line. However, because loss of insulin secretion is highly predictable by passage number and not sporadic in nature, further cellular transformation as a sole explanation seems questionable.

HIT cells are routinely serially cultured by most investigators using media containing concentrations of 11.1 mM glucose (23–31), a concentration that is supramaximal for stimulating insulin secretion from these cells (30). Consequently, we designed studies to address (a) whether the loss of insulin secretion associated with serial passages of HIT cells might be caused by adverse effects of chronic culturing the cells under high glucose conditions; and, if so, (b) whether this phenomenon is associated with an insulin gene mutation, a deficiency in the pancreatic islet glucose transporter (GLUT-2), or decreased levels of insulin mRNA; and, if so, (c) whether such changes might be prevented by culturing the cells in media containing a lower glucose concentration that is more physiologic for HIT cells.

Methods

Cell culture. Stock cultures of HIT-T15 cells were routinely grown in 5% CO2/95% humidified air at 37°C, maintained in RPMI 1640 medium supplemented with 10% fetal calf serum, passed once weekly following detachment using trypsin-EDTA, and fed every 48 h by changing medium as previously described (30). The final concentration of insulin (from fetal calf serum) in the working media averaged 5–10 nU/ml. No antibiotics were used. Beginning with passage 70, cells were split weekly and continuously passed in media containing either 0.8 mM glucose, a slightly stimulatory concentration, or 11.1 mM glucose, a supramaximal stimulatory concentration (30). Cell population doubling levels and doubling times were determined by plating cells into 12-well plates (4 cm2/well) at 0.1 million or 0.5 million cells/well. These conditions reflect the densities at which the cells were routinely cultured. Daily thereafter for 6 d, cells were detached with trypsin-
EDTA and counted. Viable cells were determined by trypan blue exclusion, and both viable and nonviable cells were counted. The cumulative doubling levels and doubling times were calculated using the following equations:

\[
\text{Number of doubling on day } i = n_i = \frac{\log(\text{TC}/\text{AC}_{i-1})}{\log 2}
\]

where

\[
\text{TC}_i = \text{total number cells on day } i \text{ (alive and dead)}.
\]

\[
\text{AC}_{i-1} = \text{number of alive cells on the day previous to day } i.
\]

\[
\text{Cumulative doubling level} = \sum n_i
\]

\[
\text{doubling time (h)} = \frac{24 \text{ h/d } \times 6 \text{ d}}{\text{cumulative doubling level}}
\]

**Insulin secretion.** Static insulin secretion in response to progressively increasing concentrations of glucose (0-11.1 mM) was examined as previously described (30). Phasic insulin secretion in response to 11.1 mM glucose perfusion in the presence of 0.1 mM 3-isobutyl 1-methylxanthine (IBMX) was examined as previously described (32).

**Insulin mRNA.** Cells were rinsed twice with ice-cold Dulbecco's PBS and scraped on ice with denaturing solution (4 M guanidine thiocyanate; 25 mM sodium citrate, pH 7; 0.5% sarcosyl; 0.1 M 2-mercaptoethanol). The lysed cells were passed several times through a 22-gauge needle to shear chromosomal DNA. RNA was isolated by the procedure of Chomczynski et al. (33). Total RNA (5-15 μg) was fractionated on 1.5% agarose formaldehyde gels and transferred to a nylon hybridization membrane (0.22 μm, Micron Separations, Inc., Westboro, MA) by electroblotting. The membrane was prehybridized in 50% formamide; 5× standard saline citrate (SSC); 10× Denhardt's; 50 mM sodium phosphate, pH 6.5; 1 mg/ml salmon sperm DNA, and 0.1% SDS at 30°C overnight. The membrane was then hybridized overnight with 32P-labeled human insulin genomic DNA probe in 50% formamide; 5× SSC; 2× Denhardt's; 50 mM sodium phosphate, pH 6.5; 0.1 mg/ml salmon sperm DNA; 0.1 mg/ml yeast tRNA; and 0.1% SDS at 30°C. The membrane was washed twice for 45 min at 50°C in 0.2× SSC and 0.1% SDS, and exposed to x-ray film (Kodak X-Omat AR) for 16-24 h. The probe was human insulin genomic DNA (phins 214; Amer. Type Culture Collection, Rockville, MD) oligolabeled with [32P]dCTP (34). The probe hybridized with a 0.5-kb band on agarose gel fractionation of total HIT cell RNA consistent with HIT cell insulin mRNA (35). Under the hybridization conditions employed, the probe also labeled HIT cell 18S and 28S RNA, and the density of labeled 18S rRNA was proportional to total RNA over the range of total HIT cell RNA applied to gels. Insulin mRNA levels were quantitated by scanning densitometry of autoradiographs; advantage was taken of 18S rRNA labeling by normalizing insulin mRNA blots for the density of the corresponding 18S rRNA. The data, therefore, are expressed as the ratio of density of insulin mRNA to density of 18S rRNA. The validity of this approach has been verified by de Leeuw (36). Also, we periodically used a probe for β-actin mRNA to verify use of 18S rRNA for normalization.

**Restriction enzyme analysis of insulin gene DNA.** DNA was isolated from HIT cells by Maniatis' procedure (37). Genomic DNA was divided into four samples, which were each digested with one of four restriction endonucleases (EcoRI, PstI, HindIII, or BamHI). The restriction fragments were separated by electrophoresis in a 0.8% agarose gel. BstEII digested lambda phage DNA was used as a size marker. The separated DNA fragments were transferred from the agarose gel to a nylon membrane as described by Southern (38). DNA was retained on the nylon membrane by ultraviolet crosslinking and then hybridized with 32P-labeled human insulin genomic DNA probe as described above. The restriction fragments containing sequences complementary to the insulin DNA probe were detected by exposing x-ray film to the nylon membrane.

**Glucose transporter.** The amount of glucose transporter (GLUT-2) was assessed by immunoblot. Antisera (W-19) against GLUT-2 was a generous gift from Dr. Mike Mueckler of Washington University, St. Louis, MO. HIT cell extracts obtained from various passages cultured under various conditions were prepared as described previously and were electrophoresed in 11% polyacrylamide gels (39). After electro-transfer to Immobilon-P, the immunoblot procedure of Harris et al. (40) was employed using W-19 antisera at a dilution of 1/250.

**Statistics.** Intergroup comparisons for densitometric data and insulin secretion data were performed by Student's t tests. Data are presented as mean±SE.

**Results**

**Insulin secreted by HIT cells during cell growth.** After splitting passage 71 into flasks with media containing either 0.8 mM glucose or 11.1 mM glucose, cells were serially passed for 40 wk by weekly splitting and continued culturing in media containing the initial glucose concentration of either 0.8 or 11.1 mM. To compensate for the empirical observation of lesser numbers of cells attached to the walls of flasks when 0.8 mM glucose was used in media, approximately four to five times as many cells were initially plated at the beginning of each week for earlier passages serially cultured in media containing 0.8 mM glucose. In this manner, approximately equal numbers of cells were present by the end of each week when comparing low and high glucose conditions. However, it was ascertained by daily total cell counts in duplicate wells performed in duplicate (the sum of attached cells plus cells floating in media) for seven consecutive days for passages 75, 85, and 95 that the population doubling times and cumulative doubling levels were not significantly different (Fig. 1). Rather, a greater proportion of cells reproducibly detached into the media when they were cultured under conditions of low glucose, which accounted for the lesser numbers of cells attached to the walls of the flasks at the time of subculturing. Initially and through passage 79, HIT cells exposed to the higher glucose concentration secreted insulin into the media that reached levels of ~3,500-4,500 μU/million cells per 48 h (Fig. 2). Thereafter, insulin levels in the media diminished progressively through passage 90 to ~100-250 μU/million cells. Media insulin levels secreted by cells cultured in 0.8 mM glucose were initially 1,000-2,500 μU/million up to

![Figure 1. Comparison of HIT cell population doubling times and cumulative doubling levels over 6 d at passages 75, 85, and 95, when serially cultured in media containing either 11.1 mM or 0.8 mM glucose.](image-url)
passage 79, and then declined gradually over time, remaining stable at ~400–1,000 μU/million cells from passage 81 to passage 95.

Insulin secretion during static incubation and during perifusion. Glucose concentration–insulin response curves were determined using selected passages serially cultured in media containing either 0.8 or 11.1 mM glucose. Cells cultured in 0.8 mM glucose were subcultured in 0.8 mM glucose for 2 or 3 days or in 11.1 mM glucose for 2 or for 9 days before secretion experiments were performed to stimulate insulin gene transcription and insulin synthesis (35). HIT cells from passages 96–97 serially passed in media containing 11.1 mM glucose failed to secrete insulin in response to glucose during 1-h static incubations. In striking contrast, late passages of HIT cells serially passed in media containing 0.8 mM glucose demonstrated two- to threefold insulin responses to glucose (Fig. 3), a magnitude usually found in passage 70 cultured in media containing 11.1 mM glucose (30).

Phasic insulin secretion in response to glucose was examined during perifusion with 11.1 mM glucose and 0.1 mM IBMX. Passages 95–98 of HIT cells serially passed in media containing 11.1 mM glucose had no phasic insulin responses to glucose. However, the same late passages of HIT cells that had been serially passed in media containing 0.8 mM glucose and then subcultured in 11.1 mM glucose demonstrated first and second phase insulin responsiveness during glucose perifusion (Fig. 4) that was similar in magnitude to that observed in early passages cultured in media containing 11.1 mM glucose (32). Comparable insulin responsiveness was observed when either 2 or 9 days were used for subculturing in 11.1 mM glucose after cells had been serially passed under low glucose conditions.

Insulin content and insulin mRNA. HIT cell passages 70, 80, 90, 100, 110, and 130 cultured in medium containing 11.1 mM glucose showed a progressive diminution of insulin content (insulin content = 2468, 1730, 629, 32, 379, and 68 μU/mg protein, respectively). Insulin content was greater in passages 87, 95, and 96 when HIT cells were serially cultured in media containing 0.8 mM rather than 11.1 mM glucose (Fig. 5). Correspondingly, insulin mRNA in later passages of HIT cells cultured in media containing 11.1 mM glucose was markedly diminished (Fig. 6), whereas insulin mRNA of HIT cells serially cultured in media containing 0.8 mM glucose was greater compared to cells cultured in media containing 11.1 mM glucose (Figs. 5 and 7). The experiments in which insulin content and mRNA were observed did not involve subculturing under high glucose conditions but rather were carried out entirely under 0.8 mM glucose conditions.

Southern analysis of insulin gene and Western analysis of GLUT-2. Examination of HIT cells from early and late passages cultured in media containing either 11.1 mM or 0.8 mM glucose.
glucose failed to reveal differences in insulin gene fragments (Fig. 8) or levels of GLUT-2 (Fig. 9).

**Discussion**

The HIT cell has proven to be a valuable experimental model of beta cell function. Insulin secretion from HIT cells is glucose concentration-related and insulin release during perfusion is biphasic (23-31). However, disappointing loss of insulin secretion from HIT cells with serial passage has been a major shortcoming (23, 26, 30). We considered the possibility that an 11.1 mM concentration of glucose, which has been routinely used in media for culturing these cells (23-31) and is supramaximal for HIT cell insulin secretion (30), might be responsible for this effect. We found that late passages of HIT cells cultured in media containing 0.8 mM glucose demonstrated normal insulin responsivity to glucose in static incubations and normal first and second phase glucose-induced insulin secretion during perfusion. In contrast, late passages cultured with 11.1 mM glucose had no insulin responses to glucose stimulation as has been previously reported (23, 26, 30, 32). HIT cells as late as passage 96 cultured in media containing 0.8 mM glucose had levels of insulin and insulin mRNA that were greater than those found in late passages of cells cultured in 11.1 mM glucose. Because culturing under conditions of low glucose concentration caused greater numbers of cells to detach from the walls of the culture flasks, it was necessary to seed greater numbers of cells in earlier passages in order to harvest comparable numbers of attached cells by the end of each passage. Nonetheless, cell population doubling times and doubling levels were similar at a given glucose concentration at passages 75, 85, and 95, although with increasing passage doubling times decreased comparably for both glucose concentrations. No insulin gene mutation or decreased levels of GLUT-2, the functional glucose transporter of pancreatic beta cells, were found in late passages of HIT cells conventionally cultured with media containing 11.1 mM glucose. Thus, our data uniquely indicate that loss of beta cell function in HIT cells passed serially in media containing a high glucose concentration is caused by preventable effects of chronically culturing under conditions of high glucose concentrations that are associated with loss of insulin mRNA and insulin content and, consequently, loss of insulin secretion.

Loss of insulin responses to glucose in HIT cells serially passed in media containing high glucose concentrations is reminiscent of abnormal insulin secretion in type II diabetic patients and in rodents that have been made permanently hyperglycemic by a brief exposure to exogenously induced high circulating glucose levels. Patients with type II diabetes mellitus lose first phase insulin responses to glucose if their fasting glucose levels are greater than 115 mg/dl (41). However, this secretory defect is glucose-specific, since first phase insulin responses to...
non-glucose agonists such as isoproterenol (7), secretin (8), arginine (9), and glucagon (10) remain intact in hyperglycemic diabetic patients even though they are somewhat diminished. An interesting parallelism exists between these observations and our previously published report that HIT cells serially passed in media containing high glucose concentrations lose insulin responses to glucose before they lose insulin responses to arginine (30). It is important to note that lowering of circulating glucose levels into the normal range has been shown to partially restore first phase insulin responses to glucose in diabetic patients (1−5). Likewise, brief intravenous infusions with agents such as phentolamine (42), sodium salicylate (43), and naloxone (44) have also been shown to partially restore first phase insulin responses to glucose despite continued hyperglycemia. The ability of these drugs to restore responses in the face of continued hyperglycemia suggests the possibility that endogenous α-adrenergic tone, prostaglandins, or endorphins may mediate some of the adverse effects of chronic hyperglycemia on beta cell function.

The role of established hyperglycemia in the development of sustained abnormalities in insulin secretion has also been examined in animal models. For example, Rossetti et al. induced hyperglycemia and defective insulin secretion in rats by 90% pancreatectomy and reported that normalization of glycemia by phlorizin treatment of the pancreatectomized animals restored normal insulin secretion (10). Leathy et al. reported that rats undergoing 60% pancreatectomy developed persistent hyperglycemia and abnormal insulin secretion only if they were temporarily fed high concentrations of sucrose (12). Imamura et al. made similar observations in 80% pancreatectomized dogs (11). However, although studies in animals have made the point that artificially induced hyperglycemia can cause defective beta cell function, such studies are intrinsically limited by the short duration over which they can be conducted. Since type II diabetes mellitus in humans evolves over a much longer period of time than several weeks, a laboratory model that can be studied over many months or years has clear advantages in examining the dimension of time and precisely chosen glucose concentrations as major variables to assess the concept of glucose toxicity on the islet. We therefore propose the HIT cell as a valuable laboratory model for examining glucose toxicity of beta cells because putative biochemical mechanisms responsible for this phenomenon and maneuvers designed to subvert its mechanisms can easily be examined over many months and potentially years using this cell line. In this context, our finding of undetectable levels of insulin mRNA in late HIT cell passages that can be prevented by culturing cells serially in a low glucose concentration points specifically toward potential genetic mechanisms of action for glucose toxicity on beta cells, which in turn leads to defective insulin secretion. One potential proposal would involve an inhibitory effect of glucose on insulin gene transcription. This would represent a paradoxical effect because under normal conditions, glucose is a physiologic stimulator of insulin gene transcription (35, 45). Should it be found that prolonged exposure to excessively high glucose levels can cause progressively diminished rates of insulin gene transcription, this would provide a possible mechanism for the decreased insulin mRNA levels and loss of insulin secretion we have observed in HIT cells. Alternatively, deleterious effects on insulin mRNA stability and degradation rates need to be considered, and although our restriction fragment analysis studies revealed no abnormalities in the insulin gene in late passages serially cultured in high glucose concentrations, this analytical approach to possible gene mutations cannot be considered definitive. Nonetheless, our finding of preventable decrements in insulin mRNA associated with culturing under high glucose conditions points to preexocytotic events for the search for mechanisms of deleterious effects of chronic hyperglycemia on beta cell function in type II diabetic patients. Such effects may contribute to phenomena that are commonly referred to as glucose toxicity, glucose desensitization, or beta cell exhaustion.

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References

Glucose Toxicity in HIT Cells


