Exposure to High or Low Glucose Levels Accelerates the Appearance of Markers of Endothelial Cell Senescence and Induces Dysregulation of Nitric Oxide Synthase

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To test the hypothesis that aging impairs endothelial cell response to glucose stress, we utilized a human umbilical vein endothelial cell in vitro model in which clinically relevant concentrations of normal (5.5 mM), high (25 mM), and low (1.5 mM) glucose were tested. With advancing population doubling, exposure to normal glucose gradually decreased endothelial nitric oxide synthase expression and activity, resulting in slow, progressive development of markers of senescence (by population doubling level [PDL] 44). High or low glucose treatment accelerated the appearance of markers of senescence (by ~PDL 35) along with declines in endothelial nitric oxide synthase expression and activity. Human umbilical vein endothelial cells exposed to alternating low and high glucose gave even more rapid acceleration in the appearance of markers of senescence (by ~PDL 18) and reduction in endothelial nitric oxide synthase levels. Thus, exposure to low and high glucose induces earlier appearance of markers of endothelial cell senescence and dysregulation of the nitric oxide synthase gene and protein expression and function. These findings will help to elucidate endothelial dysfunction associated with glucose intolerance and improve future therapy for diabetic seniors.

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Cardiovascular disease is widely recognized to increase in both prevalence and incidence with advancing age (1,2). Chronic hyperglycemia results in pathophysiology that resembles accelerated aging in a number of aspects and further increases the risk of cardiovascular disease, especially among the elderly patients. Diabetes has reached epidemic proportions in the United States, including 10.9 million people or 26.9% of those who are 65 years and older, and patients with hyperglycemia are at increased risk for microvascular and macrovascular complications (3,4). The benefit of glycemic control in reducing the risk for microvascular disease is well established. The possibility of hypoglycemia occurring in the tight glycemic management among the elderly patients is high and its role as a risk factor for cardiovascular events is a topic of increasing interest (5–8).

Homeostatic control is generally diminished with advancing age (1,2), including the regulation of glucose metabolism that is reflected by the declining glucose tolerance in seniors (9,10). Both aging and hyperglycemia are associated with endothelial dysfunction, vascular stiffening, and remodeling, as well as reduced nitric oxide function and impaired vasomotor control (11–14).

One process that has been increasingly linked with both aging and hyperglycemia and the development of vascular pathologies is the phenomenon of cellular senescence (15,16). More recently, it has been realized that the response of early replicative senescence can also be induced by a number of stress stimuli, especially by those causing intracellular oxidative stress. The process has been given a variety of names including “stress-induced premature senescence” and “stress or aberrant signaling-induced senescence” (17). Increased numbers of senescent endothelial cells are found in the vasculature of seniors, mature atherosclerotic plaques, vessels from younger adult diabetic patients, coronary vessels of patients with ischemic heart disease, and hypertensive patients. These findings suggest that the senescence of vascular cells likely contributes to a reduction of vasomotor control and age-related vascular diseases (18).

Endothelial cell function is essential for the homoeostasis of the vascular system. Due to its unique position in the vessel wall, the endothelium acts as a barrier and serves as the primary sensor and mediator of blood flow-mediated vasomotor control (19,20). Nitric oxide, produced by endothelial nitric oxide synthase (eNOS), is a key signaling molecule in vascular homeostasis and is an important regulator of vascular tone and arterial pressure. Loss of nitric oxide bioavailability is a cardinal feature of endothelial dysfunction and is an independent predictor of increased
cardiovascular disease risk (21–23). Furthermore, external stress stimuli such as hyperglycemia have been shown to raise the expression and activity of inducible NOS (iNOS) (24), which is associated with excessive production of nitric oxide and leads to increased cellular damage. Advanced age and hyperglycemia have been shown to be associated with impaired basal eNOS and enhance endothelial dysfunction (25–27). The disappointing results of recent clinical trials of tight glycemic control in seniors with type 2 diabetes highlight the potential cardiovascular risks of hypoglycemia and hyperglycemia (28–31).

In the present study, we tested the hypothesis that cellular aging reduces the endothelial cell’s ability to respond to glucose stress and that alternating levels of elevated and low glucose may further accelerate cellular senescence. We observed that exposure to low and/or high glucose induced the premature appearance of markers of cellular senescence and dysregulation of NOS.

Materials and Methods

Endothelial Cell Culture

Human umbilical vein endothelial cells (HUVECs) were obtained from Lonza (Clonetics, Walkersville, MD) as pooled primary cell lines and cultured as previously described (32–34). HUVECs were cultured in BD Falcon 75-cm² flasks, containing EGM-2 bullet kits supplemented with 5% heat irradiated fetal bovine serum (Lonza). Cell cultures were incubated at 37°C in 5% CO₂. Primary cultures received new medium 24 hrs after seeding and were subcultured upon reaching 80% confluence by the use of 0.25% trypsin-EDTA, inactivated by trypsin neutralizing solution (Lonza). Population doubling levels (PDLs) were estimated at each passage using the following equation: \( n = (\log_2 X - \log_2 Y) \), where \( n = \text{PDL} \), \( X = \text{number of cells at the end of one passage} \), \( Y = \text{number of cells that were seeded at the beginning of one passage} \); see Supplementary Material 1).

Glucose Stress

The following clinically relevant glucose concentrations were tested in varying exposures: normal (100 mg/dL or 5.5 mM), high (450 mg/dL or 25 mM), and low (30 mg/dL or 1.5 mM). Experimental groups (Figure 1) included early through late passage HUVECs treated with (i) continuous normal (5.5 mM), (ii) continuous high (25 mM) glucose, (iii) alternating glucose (normal and high), (iv) alternating low (1.5 mM)/normal glucose (low and normal), and (v) alternating low and high glucose (low and high). For simulating hyperglycemia and/or hypoglycemia exposure, early (PDL 6–8) through late (PDL 44±2) passage cells were incubated in 25 mM high glucose for 48 hrs and compared with that of normal glucose control cells at the respective PDLs (see Supplementary Material 1).

Experimental Design

1. Continuous normal glucose (NG)
2. Continuous high glucose (HG)
   48 h
   Normal glucose or high glucose
3. Alternating high/normal glucose (24 h HG/24 h NG)
   96 h
   Alternating glucose
4. Alternating low/normal glucose (6 h LG/6 h NG)
5. Alternating low/high glucose (6 h LG/6 h HG)
   96 h
   Alternating glucose

Figure 1. Glucose treatment protocol. This figure shown illustrates the duration and concentration levels used for all glucose experimental groups tested in this study. The following clinically relevant glucose concentrations were used: normal (100 mg/dL or 5.5 mM), high (450 mg/dL or 25 mM), and low (30 mg/dL or 1.5 mM). Human umbilical vein endothelial cells were incubated at 37°C in 5% CO₂ and grown in T-75 flasks. The experimental values for all tested groups were obtained from four repeated cultures (in parallel) of endothelial cells from early (population doubling level [PDL] 6) to late (PDL 44) passage.

Senescence-Associated Beta-Galactosidase Staining

To visualize a marker of cellular senescence, in vitro staining for senescence-associated beta-galactosidase (SA-β-gal) was performed using a commercial kit from Cell Signaling Technology (Boston, MA), designed to detect β-galactosidase activity at pH 6, a known characteristic of senescent cells that is usually not found in presenescent, quiescent, or immortal cells (35) (see Supplementary Material 1).

Western Blotting

Western blotting of the cell lysates was conducted as performed previously in our lab (36,37). In brief, the cells were lysed in a lysis buffer (10 mM Tris, pH 7.4, 100 mM NaCl, 1 mM EDTA, 1 mM ethylene glycol tetraacetic acid, 1% Triton X-100, 10% glycerol, and 0.1% sodium dodecyl sulfate) supplemented with 1x protease inhibitor cocktail (Sigma-Aldrich, St Louis, MO). Proteins were denatured in Laemmli sample buffer for 5 minutes at 95°C and resolved by Mini-PROTEAN TGX gels (Bio-Rad, Hercules, CA). Following electrophoresis, western protocol was adjusted based on Licor In-Gel detection instructions (Lincoln, NE). The gel was incubated with gentle shaking in 50% isopropanol + 5% acetic acid (prepared with ultrapure
GLUCOSE STRESS MAY ACCELERATE ENDOTHELIAL SENESCENCE

Effects of High and Low Glucose on SA-β-gal
Positive HUVECs
When staining for the presence of SA-β-gal in HUVECs treated with varying glucose exposures (Figure 1), we observed a general downward shift in the appearance of markers of cell senescence to an earlier population doubling point at which replicative senescence approaches 100%. The magnitude of this shift was found to be dependent upon the type and duration of glucose level exposure, with no major differences found among any treatment groups at the earliest and latest HUVEC passages. In early passage (PDL 6) HUVECs, appearance of markers of cellular senescence activity showed no significant difference among all normal (5 mM), high (25 mM), and low (1.5 mM) glucose exposed treatment groups: normal glucose 4% ± 2, high glucose 8% ± 4, alternating high and normal glucose 9% ± 4, high glucose and normal glucose 8% ± 3, alternating low and normal glucose 8% ± 2, and alternating low and high glucose 10% ± 5. Elevated glucose significantly increased SA-β-gal activity, 88% ± 6, in PDL 35 HUVECs compared with the 9% ± 2 normal glucose PDL 35 control group. Alternating high and normal glucose was found to accelerate senescence to a significantly earlier PDL, from 44 to 25, with a percent SA-β-gal activity of 86% ± 6 compared with 7% ± 3 in normal glucose PDL 25 controls. Alternating low and normal glucose exposure induced earlier appearance of markers of senescence to 84% ± 5 in HUVECs at PDL 30 compared with 12% ± 6 in the normal glucose PDL 30 control group. Lastly, HUVECs exposed to alternating low and high glucose were shown to manifest accelerated appearance of senescence markers most dramatically, with 81% ± 4 SA-β-gal activity at PDL 18, compared with only 7% ± 3 in the normal glucose PDL 18 control group (Figure 2a–e).

Real-Time Reverse Transcription PCR Quantification of mRNA
To detect each mature mRNA of interest, the following primers were used: eNOS, forward primer: 5'-ccacaatctctggtctgc-t-3' and reverse primer: 5'-gcctttttccagtttcca-3'; iNOS, forward primer: 5'-accaggtttgcaaatggaga-3' and reverse primer: 5'-tgcccacagcttactgaga-3'; and AKT, forward primer: 5'-tgcccacagcttactgaga-3 and reverse primer: 5'-tacctcagcaccagcatcacc-3'. The GAPDH gene was used as an internal control: GAPDH forward: 5'-actctacccacggcaaa-t-3' and GAPDH reverse: 5'-tgcccacagcttactgaga-3'. The expression of GAPDH mRNA was tested in all glucose treatments and across all PDL time points, and it was confirmed that GAPDH mRNA was unchanged throughout all the groups (see Supplementary Material 1).

Influence of Glucose Stress on Cell Proliferation Rates Over Time
The HUVEC proliferation rates also showed a similar general downward shift in the population doubling point toward earlier PDLs in which cell proliferation progressively slowed toward complete growth arrest. Again, the profile of this shift was found to be dependent upon the type and duration of glucose level exposure, with no major differences found among all treatment groups at the earliest and last passage HUVECs. Early passage endothelial cells (PDL 6) showed no statistical difference in proliferation rates (hours per population doubling) among all normal, high, and low glucose treatment groups: normal glucose 22 hrs ± 2, high glucose 22 hrs ± 4, alternating high glucose 26 hrs ± 4, alternating low and normal glucose 22 hrs ± 2, and alternating low and high glucose 25 hrs ± 5. At PDL 35, both normal and high glucose exposed HUVECs showed a significant decrease in proliferation rate compared with that of the early passage normal glucose control group. Normal glucose HUVECs at PDL 35 gave a rate of 36h/PDL, whereas high glucose exposed PDL 35 endothelial cells showed a significantly lower rate of 90h/PDL (p < .01). Before complete replicative senescence (growth arrest), late passage HUVECs at PDL 44 were found to have significantly lower proliferation rates in both normal and high glucose treatment groups compared with that of the early passage control. Normal glucose PDL 44 group was shown to have a proliferation rate of 92 hrs and high glucose HUVECs at this same PDL were significantly slower, with a rate of 103 h/PDL (Figure 2a–e).

p16INK-4a Protein Expression in Early and Late Passage HUVECs Exposed to Glucose Stress
To further examine changes in markers of cellular senescence in response to glucose stress (both high and low concentrations), we analyzed the protein expression of p16INK-4a in early through late passage HUVECs. Figure 3a and b shows the change in expression levels for water) for 15 minutes (Sigma-Aldrich). Fixative solution was removed and gel was washed in ultrapure water for 15 minutes with gentle shaking. Primary antibodies (p16INK-4a, AKT, p-AKT, iNOS, eNOS, and p-eNOS) were diluted in Licor Odyssey Buffer with 0.1% Tween 20 (see Supplementary Material 1).

Total RNA Isolation
Total cellular RNA was extracted from cells using the RNeasy kit (Qiagen, Valencia, CA) in accordance with the manufacturer’s recommendations for mammalian cells. RNA with a high RNA integrity number and an A260 to A280 absorbance ratio ranging from 1.8 to 2.1 was utilized for cDNA synthesis.

Results
PDL 12, PDL 25, and PDL 44 HUVECs exposed to glucose stress. Normal glucose control cells at PDL 12 showed very little expression of p16INK-4a, whereas HUVECs of the same PDL, exposed to high and/or low glucose, displayed a small but significant increase in expression levels. HUVECs exposed to normal glucose at PDL 25 showed a slight increase in p16INK-4a compared with that of earlier passage control cells. Furthermore, endothelial cells exposed to all tested glucose treatments showed significant increases in p16INK-4a protein expression. Late passage
HUVECs showed significant increases in protein expression of p16INK-4a compared with that of PDL 12 and PDL 25 endothelial cells.

**Analysis of Protein Expression in Early and Late Passage HUVECs Exposed to Glucose Stress**

The initial Western blotting experiments were conducted to examine whether early and/or late passage HUVECs exposed to temporary (48 hrs) high glucose would demonstrate changes in regulation of AKT, p-AKT (threonine 308), iNOS, eNOS, and the active form of eNOS, p-eNOS (serine 1177) with that in endothelial cells exposed to normal glucose. Additional studies were performed to determine whether this regulatory change in protein expression would persist following glucose normalization and/or alternating high and low glucose exposure.

Protein expression of AKT was found to be statistically unchanged among all the tested glucose treatments in both early and late passage HUVECs as seen in Figure 4a and b. However, in PDL 25 endothelial cells, AKT expression was markedly increased in response to alternating low and high glucose exposure. AKT protein expression in PDL 25 HUVECs was also slightly increased following treatment with alternating high and normal and alternating low and normal glucose.

We also analyzed protein expression of threonine 308 phosphorylated AKT induced by in vitro cellular aging and glucose stress. Early passage HUVECs (PDL 12) show no significant difference in p-AKT protein expression among all glucose treatments and the control group. Similarly, no statistical difference was observed for p-AKT in PDL 25 and PDL 44 endothelial cells among all tested glucose treatment groups and the control. A slight overall decline in p-AKT protein expression was seen in PDL 25 and PDL 44 HUVECs compared with that of early passage (PDL 12) endothelial cells (Figure 5a and b).

Total protein expression for eNOS was also measured in early through late passage endothelial cells (Figure 6a and b). In early passage (PDL 12) HUVECs, virtually all tested glucose treatments showed no significant difference in total eNOS protein expression, with alternating low and high glucose exposure producing a slight decline in eNOS levels.
HUVECs exposed to normal glucose at PDL 25 showed no significant difference in eNOS levels compared with that of early passage PDL 12 normal glucose controls. However, endothelial cells treated with high and/or low glucose at PDL 25 displayed a significant decline in overall eNOS protein, evident in all glucose treatments as shown in Figure 6a and b. Interestingly, HUVEC exposure to alternating low and high glucose produced the greatest reduction in total eNOS levels compared with normal glucose controls. Lastly, late passage PDL 44 showed a similar trend as observed in PDL 25, with all glucose treatments resulting in greater significant reductions in eNOS protein expression compared with that of the early passage normal glucose controls.

Next, we tested the differences in AKT-dependent phosphorylation of the serine 1177 eNOS site induced by in vitro cellular aging and elevated and reduced glucose levels. For early passage (PDL 12) HUVECs, we found a slight reduction of eNOS phosphorylation in high, alternating high and normal, and alternating low and normal glucose conditions compared with normal glucose PDL 12 controls (Figure 7a and b). HUVECs exposed to alternating low and high glucose were found to have an even greater reduction in p-eNOS levels at PDL 12. Endothelial cells exposed to normal glucose at PDL 25 showed no significant change in the level of p-eNOS expression compared with the early passage PDL 12 normal glucose control. However, all tested high and/or low glucose treatment groups were found to have significantly lower p-eNOS expression compared with the normal glucose control. In late passage HUVECs (PDL 44), all high and/or low glucose treatments resulted in significant decreases in phosphorylated (serine 1177) eNOS compared with that of early passage normal glucose control. Normal glucose exposure also produced lower p-eNOS levels in late compared with early PDL cells, though the difference was not as significant as the previously described high and/or low glucose treatments compared with that of the early passage normal glucose control group.

We also measured the levels of iNOS protein expression among HUVECs treated with varying glucose exposures through early to late passage. As seen in Figure 8a and b, normal glucose control cells at PDL 12 had minimal iNOS expression. Early passage PDL 12 endothelial cells
exposed to high, alternating high and normal, or alternating low and normal showed significantly increased levels of iNOS protein compared with the PDL 12 normal glucose control. HUVECs exposed to alternating low and high produced even greater increases in iNOS expression compared with that of all other treatment groups at PDL 12 (Figure 8a and b). Endothelial cells treated with normal glucose at PDL 25 also showed virtually no iNOS expression and no significant difference compared with that of the early passage (PDL 12) control. However, HUVECs exposed to glucose stress (all the high and/or low glucose treatments) resulted in the most significant increases in iNOS protein expression at PDL 25. Endothelial cells exposed to these high and/or low glucose treatments at PDL 44 showed an overall slight reduction in the increase in iNOS protein expression compared with that seen in PDL 25. However, even normal glucose exposure resulted in a significant increase in iNOS presence compared with that of the early passage normal glucose control, suggesting that there is a glucose-independent increase of iNOS in senescent cells.

Changes in HUVEC Gene Expression

Real-time quantitative reverse transcription PCR experiments were performed to determine whether various glucose conditions might mediate alterations of transcriptional regulation in AKT and its downstream signaling target, eNOS, and iNOS. Expression of AKT mRNA was found to be statistically unchanged among all tested glucose treatments in both early and late passage HUVECs (Figure 9c). However, in PDL 25 endothelial cells, AKT expression was significantly increased (2.1 ± 0.047-fold) in response to alternating low and high glucose exposure compared with early passage normal glucose control HUVECs. AKT mRNA expression in PDL 25 HUVECs was also slightly increased following treatment with alternating high and normal and alternating low and normal glucose, 1.39 ± 0.035- and 1.54 ± 0.041-fold increases, respectively.

As shown in Figure 9a, there was no significant difference in mRNA expression of eNOS at PDL 12 among normal (1 ± 0.039), temporary high (0.94 ± 0.054-fold decrease), and high (0.98 ± 0.066-fold decrease). A slight decrease in eNOS expression at PDL 12 was observed...
among alternating high and normal (0.88 ± 0.071-fold decrease), alternating low and normal (0.89 ± 0.061-fold decrease), and alternating low and high (0.85 ± 0.052-fold decrease) glucose levels compared with that of normal glucose exposure. For middle passage (PDL 25) HUVECs, eNOS mRNA expression was reduced to a greater extent with all glucose treatments compared with that of the early passage normal glucose control. As shown in Figure 9a, normal and high glucose exposure at PDL 25 resulted in a 0.88 ± 0.039- and 0.84 ± 0.067-fold decreases in mRNA eNOS expression, respectively. HUVECs at PDL 25 exposed to alternating high and normal (0.67 ± 0.041-fold decrease), alternating low and normal (0.64 ± 0.043-fold decrease), and alternating low and high (0.52 ± 0.062-fold decrease) glucose levels showed significantly decreased eNOS mRNA levels compared with that of the normal glucose control. In late passage PDL 44 HUVECs, mRNA expression of eNOS was significantly downregulated for all glucose treatments. Figure 9a shows that exposure to normal glucose resulted in a 0.59 ± 0.047-fold decrease in eNOS mRNA expression at PDL 44 compared with that of early passage normal glucose control. Exposure to high and/or low glucose in HUVECs at PDL 44 resulted in even greater decreases in eNOS mRNA levels: high glucose (0.56 ± 0.034-fold decrease), alternating high and normal glucose (0.58 ± 0.061-fold decrease), alternating low and normal (0.58 ± 0.044-fold decrease), and alternating low and high (0.43 ± 0.057-fold decrease).

Expression of iNOS mRNA was found to undergo the most dramatic alterations in response to glucose treatments and also with advancing PDL (Figure 9b). In early passage PDL 12 HUVECs, iNOS mRNA expression was significantly increased following exposure to high (1.47 ± 0.055-fold increase), alternating high and normal (1.84 ± 0.046-fold increase), alternating low and normal (1.81 ± 0.033-fold increase), and alternating low and high (2.11 ± 0.051-fold increase) glucose. Endothelial cells at PDL 25 were observed to have greater increases compared with that of PDL 12 in iNOS expression among all glucose treatments. Expression of iNOS in normal glucose HUVECs at PDL 25 was found to be 1.51 ± 0.049-fold increase compared with that of the early passage control. Additionally, HUVEC
exposure at PDL 25 to high (2.17±0.041-fold increase), alternating high and normal (2.55±0.039-fold increase), alternating low and normal (2.7±0.049-fold increase), and alternating low and high (4.24±0.052-fold increase) glucose showed significantly increased iNOS mRNA levels compared with that of the PDL 25 normal glucose control. Finally, iNOS mRNA expression was most significantly elevated in late passage PDL 44 HUVECs in all glucose treatment groups: normal (2.61±0.049-fold increase), high (3.81±0.054-fold increase), alternating high and normal (4.19±0.061-fold increase), alternating low and normal (4.44±0.038-fold increase), and alternating low and high (5.12±0.042-fold increase).

**Discussion**

The present investigation has identified several novel findings. Our data indicate that exposure to glucose stress (both elevated and reduced glucose levels) significantly accelerated the appearance of markers of endothelial cell senescence. The results also strongly suggest that this accelerated appearance of markers of senescence is, in part, due to regulatory changes in the transcription and expression of AKT and its downstream signaling targets, including NOS. Additionally, we observed that exposure to alternating glucose levels rapidly impaired eNOS activity, resulted in overexpression of iNOS and decreased overall nitric oxide bioavailability and impaired endothelial function in the human endothelial cell. Taken together, these findings suggest that alternating between high and low glucose levels resulted in the acceleration of stress-induced senescence phenotype. Elevated and reduced glucose levels alter the late passage cell’s nitric oxide antioxidant function from low, basal, protective levels to higher, cytotoxic levels, and consequently produce cellular dysfunction, damage, and a senescent phenotype.

**Effects of Glucose Stress on Appearance of Markers of Senescence and Reduction of Proliferation in Early and Aged HUVECs**

Elderly patients are at higher risk of cardiovascular complications associated with the metabolic syndrome and they are more prone to hypoglycemia in response to pharmacological treatment strategies aimed at tight control of high blood glucose levels (38–41). In the Action in Diabetes and Vascular Disease: Preterax and Diamicron Modified Release Controlled Evaluation study, severe hypoglycemia induced by tight glucose control in elderly diabetics was clearly associated with increased risk of macrovascular events, microvascular events, and even death from both cardiovascular and noncardiovascular causes (42).

Our findings indicate that the inherent inability of the endothelial cell to regulate glucose flux in the face of glucose stress (high or low concentrations) can cause the accelerated appearance of markers of cell senescence. Under normal conditions, HUVECs usually maintain a relatively constant cell proliferation rate, and the progressive decline in proliferation rate and appearance of cellular senescence tend to be gradual until very late passage numbers (~PDL 40). Increased expression of p16INK-4a, an additional marker of senescence, was also observed in late passage (PDL 40) HUVECs. Elevated glucose levels significantly accelerated the replicative senescence and decreased the cell proliferation rates. Interestingly, when we exposed early passage HUVECs to temporarily elevated glucose levels, followed by a return to normal glucose levels, the senescence acceleration was found to be reduced compared with that of chronically elevated glucose. However, this was not the case in late passage cells, where the senescence acceleration after temporary elevation of glucose followed by normal glucose did not differ from that after chronically elevated glucose exposure. Exposure to alternating low glucose followed by normal glucose also resulted in reduced HUVEC proliferation. Moreover, the most deleterious effects were observed when HUVECs were exposed to alternating low followed by high glucose levels, which resulted in a dramatic acceleration of the appearance of markers of cellular senescence, with early passage endothelial cells (PDL 12) showing significant stress-induced growth arrest.

These results strongly suggest that later passage endothelial cells have a dramatically reduced ability to respond to glucose stress, resulting in an accelerated appearance of markers of the senescence phenotype and reduced replicative rates. As the rest of our data suggest, this diminished capability to tolerate glucose stress is, in part, due to regulatory changes in antioxidant mechanisms associated with NOS expression and activity. Taken together, these data indicate a novel mechanism of inducing early appearance of markers of endothelial cell senescence and endothelial dysfunction, namely, that of alternating high and/or low glucose levels, which may resemble the typical clinical scenario involving tight glycemic regulation among elderly diabetic patients.

**Protein Expression of NOS and AKT Is Differentially Regulated in Early and Late Passage HUVECs in Response to Glucose Stress**

In higher concentrations, nitric oxide can act as a reactive nitrogen species, the effects of which can contribute to pathological mechanisms that are involved in cardiovascular aging (43). All three isoforms of NOS have been found in cardiovascular tissues (44). eNOS is constitutively expressed in the endothelium under basal conditions and is responsible for the low level of nitric oxide production involved in endothelium-dependent vasodilatation. Our results show that under normal glucose levels, the HUVEC expression of eNOS gradually decreases over the passage of time. Additionally, early
passage protein expression of eNOS is virtually unchanged among all different glucose level treatment groups. However, in middle and late passage endothelial cells, high and/or low glucose exposure induces much greater declines in the expression of eNOS, most noticeably with alternating low and high glucose treatment. These results suggest that high and/or low glucose levels reduce the aged HUVEC’s nitric oxide signaling ability via depletion of basal eNOS and reduce its cytoprotective effects. The shift in iNOS:eNOS ratio (Figure 9d) toward a predominant iNOS presence likely results in much higher levels of free-radical (reactive nitrogen species) production and consequent injury, resulting in increased endothelial cell dysfunction and the accelerated appearance of markers of senescence.

iNOS has not been found to be constitutively present at detectable levels in the normal, younger adult heart or vasculature, but it can be induced by proinflammatory substances, such as cytokines, lipopolysaccharides, or proinflammatory events, including hypertension, atherosclerosis, ischemic stress, stroke, trauma, and infection (45). Recent studies have suggested that the aging process induces increased expression of iNOS in the brain (46) and the heart (47). For example, Ciszar and colleagues (48) demonstrated that aging is associated with a shift in coronary arteriole phenotype, including increased expression of iNOS and increased activity of NAD(P)H oxidases. Our data strongly suggest that glucose stress (both high and low glucose levels) is a potent inducer of iNOS protein upregulation and that these effects are further enhanced in the older (later passage) endothelial cells. Even at relatively early PDL numbers, iNOS protein levels were significantly increased in response to both high and low glucose exposure. iNOS is a more potent generator of nitric oxide and this shift to the predominant presence of iNOS protein results in an unfavorable intracellular free-radical environment due to the loss of low, basal level of nitric oxide production via eNOS. Therefore, alternating levels of low and high glucose appear to synergistically impair the endothelial cells’ nitric oxide-dependent antioxidant function partly by overcompensating NOS expression and activity. Consequently, we see a concomitant shift in the acceleration of appearance of markers of senescence and HUVEC growth arrest in response to both high or low glucose stress.

**Phosphorylation of eNOS**

AKT is an important mediator of many physiological pathways and is directly involved in metabolic regulation, cell growth, and proliferation. Recently, studies have shown its crucial role in p53-mediated senescence and proposed a potential affiliation with the “senescence-associated secretory phenotype.” These findings, coupled with AKT’s known involvement in inflammation and oxidative stress, suggest a potential mechanism in the alteration of NOS expression and development of senescence in response to glucose stress. AKT’s role in the activation of eNOS via phosphorylation of serine residue 1177 is well described (49,50). Perturbations of eNOS phosphorylation have been reported in a number of pathologies, thereby emphasizing the importance of the expression and regulation of eNOS activity in vascular health (51). Our data show some interesting findings related to expression of AKT in response to elevated and reduced glucose and endothelial cell aging. There was little change in early and late passage AKT protein expression among all treatment groups. However, in middle passage PDL 25 HUVECs, we observed an acute upregulation in AKT protein levels in response to alternating low and high glucose exposure. We propose that this increase is a compensatory response of the endothelial cells to upregulated NOS activity (antioxidant function) in the face of high and/or low glucose stress. The AKT protein phosphorylates a wide range of proteins, including eNOS. Therefore, small changes in its total protein expression may not have a direct effect on any one particular phosphorylation target. AKT is also directly activated and inhibited via mammalian target of rapamycin, which is known to be involved in nutrient sensing and in the progression of cellular senescence. Additionally, there could be other factors involved, including phosphorylation of eNOS via PKA (S1179) as well as other posttranslational events such as myrotylation, nitrosylation, and/or glycosylation of eNOS. Furthermore, eNOS is also known to be dysfunctional in glucose stress due to BH_{2}/BH_{4} alterations. Our data also show that protein expression levels of AKT phosphorylation at threonine 308 were, for the most part, unchanged among glucose treatment groups for early, middle, and late passage HUVECs. We did observe an overall decrease in p-AKT levels in middle and late passage cells compared with that of early passage control and treatment groups. The literature also supports that AKT phosphorylation may be biphase. Kinoshita and colleagues (52) showed that p-AKT levels were increased in human arteries in response to high glucose exposure for 1 h. Yet another study reported that long-term glucose exposure, 7 days, resulted in reduced expression of p-AKT in human aortic smooth muscle cells (53). Our results suggest that although phosphorylation of AKT at threonine 308 is slightly reduced in middle and late passage HUVECs, regulation of eNOS activity in response to high and low glucose in senescent cells may also be mediated through additional partners of the PI3K/AKT pathway, such as the nutrient-sensing mammalian target of rapamycin or AMPK. More work to delineate the role of these factors and the regulatory mechanisms involved is needed. These findings lend support to the notion that endothelial cell oxidative stress and accelerated cell senescence are likely the result of dysfunctional intracellular antioxidant response mechanisms.
Glucose Stress Induces Transcriptional Dysregulation of NOS

As described in the Results section, in vitro cellular aging and alternating reduced and elevated glucose levels induced differential protein expression of NOS and AKT in HUVECs. Therefore, we sought to determine if these alterations in protein levels correlated with transcriptional differences among the respective genes of interest. It is known that eNOS activity undergoes regulation at the posttranslational level, mainly via phosphorylation and/or dephosphorylation mechanisms that are partly mediated by AKT. Our results, however, also show an overall gradual decline in eNOS mRNA over passage of time in all of the glucose groups. This decrease was most significant in HUVECs exposed to alternating low and high glucose levels and are in agreement with that of other studies, which found that the aging process alone was associated with decreased eNOS gene expression in heart tissue and arterial vascular cells (54). Our results indicate that exposure to alternating low and high glucose levels enhances this documented progressive decline in eNOS transcription among endothelial cells. Normal endothelial production of nitric oxide plays an important role in preventing vascular disease. Endothelium-dependent vasodilation has been reported to be impaired in both the microcirculation and macrocirculation during acute hyperglycemia in normal (nondiabetic and euglycemic) (55,56) participants and in diabetic patients (57), suggesting that NOS activity may be chronically impaired in patients with hyperglycemia, insulin resistance, metabolic syndrome, and/or diabetes.

Lastly, in the present study, we found that iNOS gene expression was significantly higher among late versus early passage HUVECs, and that the difference was significantly enhanced in response to high and/or low glucose levels. We also found that although basal iNOS mRNA levels increased from PDL 25 to PDL 44, protein expression of iNOS decreased or remained the same. One explanation for this finding may be due to an overall reduction in protein translation among the late passage, senescent cells, or dysfunctional translational “machinery.” Recent studies have shown that global protein translation is reduced among senescent cells, and Dicer expression and activity is likewise dysfunctional (58–60). Glucose stress may accelerate or enhance this age-associated protein translation dysfunction.

The observed upregulation of iNOS mRNA levels together with the reduction in eNOS gene expression suggest that HUVECs, in part, shift or replace nitric oxide production source from that of the basal, protective eNOS to that of the more potent, deleterious iNOS enzyme. We propose that this shift in NOS regulation, including the decreased expression of eNOS, together with the increased expression of iNOS and AKT may underlie the acceleration of appearance of markers of endothelial cell senescence and vascular dysfunction. This alteration in NOS expression seems to be a potent proinflammatory response and a phenotypic consequence of senescent endothelial cells, exacerbated by glucose stress.

Conclusions

The results from the present study suggest that both high and low glucose levels may result in earlier appearance of markers of endothelial cell senescence. In addition, glucose stress may cause altered gene and protein expression among members in the AKT–NOS pathway. These changes may result in deleterious effects on the endothelium and the development of vascular dysfunction. Managing diabetes in the elderly population is difficult because of complex comorbid medical issues and the generally limited functional status of many senior diabetic patients. Nationally published guidelines for treating adults are often not easily applicable to geriatric care, and practitioners’ individualized approaches to therapy are highly variable. Understanding the special needs of geriatric patients and the mechanisms underlying their impaired physiological responses to glycemic stress, especially hypoglycemia, will likely aid in improving the overall care of elderly diabetic patients in the future.

Supplementary Material

Supplementary material can be found at: http://biomedgerontology.oxfordjournals.org/

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References


