Early neural and vascular dysfunctions in diabetic rats are largely sequelae of increased sorbitol oxidation

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Early Neural and Vascular Dysfunctions in Diabetic Rats Are Largely Sequelae of Increased Sorbitol Oxidation

Yasuo Ido,1 Jens R. Nyengaard,2 Kathy Chang,3 Ronald G. Tilton,4 Charles Kilo,3 Banavara L. Mylari,5 Peter J. Oates,6 and Joseph R. Williamson7

Abstract

These experiments were undertaken to assess the importance of cytoplasmic (c) sorbitol oxidation versus mitochondrial (m) pyruvate oxidation in mediating neural and vascular dysfunction attributable to hyperglycemia in diabetic rats. Increased oxidation of sorbitol is coupled to enzymatic reduction of free oxidized NAD$^+$ to reduced NADHc, manifested by an increased ratio of NADH to NAD$^+$. Likewise, increased oxidation of pyruvate is coupled to reduction of NAD$^+$m to NADHm, which increases the NADH/NAD$^+$m ratio. Specific inhibitors of sorbitol production or sorbitol oxidation normalized: increased diabetic nerve NADH$^{=}$$\text{NAD}^+$ ratio, impaired nerve-conduction velocity, and vascular dysfunction in sciatic nerve, retina, and aorta; however, they had little or no impact on increased NADH/NAD$^+$m. These observations provide, for the first time, strong in vivo evidence for the primacy of sorbitol oxidation versus pyruvate oxidation in mediating the metabolic imbalances, impaired nerve conduction, and vascular dysfunction evoked by diabetes. These findings are consistent with (a) the fact that oxidation of sorbitol produces “prooxidant” NADHc uncoupled from subsequent production of “antioxidant” pyruvate required for reoxidation of NADHc to NAD$^+$c by lactate dehydrogenase, and (b) the hypothesis that neural and vascular dysfunction in early diabetes are caused primarily by increased NADHc, which fuels superoxide production by NADH-driven oxidases. Antioxid. Redox Signal. 12, 39–51.

Introduction

Increasing evidence supports the importance of superoxide ($O_2^\cdot$) and related reactive oxygen species (ROS) in mediating diabetic complications attributed to hyperglycemia (4, 5, 8, 20, 29, 52); however, the primary source(s) of electrons that fuel superoxide production is controversial. Two distinctly different hypotheses have been proposed: (a) increased oxidation of pyruvate (produced by increased glycolysis) in mitochondria coupled to reduction of free NAD$^+$m to NADHm, which promotes superoxide production by the mitochondrial electron transport chain (5, 29); and (b) increased oxidation of sorbitol (produced by increased flux of glucose via the sorbitol pathway, which does not produce pyruvate) by sorbitol dehydrogenase (SDH) to fructose coupled to reduction of cytosolic NAD$^+$c to NADHc (equimolar to fructose) that drives superoxide production primarily by NADH-driven oxidases (20, 30, 55):

$$\text{Sorbitol} + \text{NAD}^+ \rightarrow \text{fructose} + \text{NADH} + \mathrm{H}^+;$$

$$\text{NADH} + \text{O}_2 \rightarrow \text{NAD}^+ + \text{O}_2^-.$$

The first hypothesis suggests that pyruvate supplementation might mimic or exacerbate metabolic imbalances and vascular and neural dysfunction evoked by hyperglycemia. However, pyruvate supplementation (a) normalizes/attenuates vascular dysfunction and metabolic imbalances evoked by hyperglycemia in several different paradigms of diabetes (16, 24, 44, 48, 50, 57), and (b) attenuates cataract formation in diabetic rats (58).
The second hypothesis suggests that sorbitol supplementation (at normal glucose levels) might cause oxidative stress and associated metabolic imbalances and vascular dysfunction comparable to hyperglycemia/diabetes. This prediction has been confirmed in many investigations in tissues and cells exposed to elevated sorbitol levels in vivo and in vitro [9, 12, 26, 30 (pages 9–10 in Online Appendix Section (OAS)-IV-A see Supplemental Appendix at www.liebertonline.com/ars], 46, 47, 49, 53]. These effects of sorbitol also are prevented or substantially attenuated by coadministration of pyruvate, SOD (superoxide dismutase), and/or by inhibitors of SDH (SDI), or both (12, 26, 46, 47, 53). These effects of pyruvate and sorbitol are consistent with a potentially important role for sorbitol oxidation in mediating oxidative stress and vascular and neural dysfunction evoked by diabetes.

Observations that SDI and SOD prevent sorbitol-induced vascular dysfunction and superoxide production are consistent with numerous observations in animal models of diabetes that inhibition of sorbitol production by aldose reductase (AR) inhibitors (ARI) also prevent/attenuate vascular and neural dysfunction, oxidative–nitrosative stress, and the predicted increases in free NADH/NAD⁺c (6, 10, 20, 30–37, 39, 48, 50, 55, and OAS I-D, I-E).

To the extent that metabolic imbalances and vascular and neural changes in early diabetes are largely sequelae of increased sorbitol oxidation rather than oxidation of NADPHc to NADP⁺c by AR, they should be prevented by ARI or SDI:

\[
\text{Glucose} + \text{NADPHc} + H^+ \xrightarrow{\text{ARI}} \text{NADPH} + \text{H}^+ \\
\text{Sorbitol} + \text{NAD}^+ c \xrightarrow{\text{SDI}} \text{fructose} + \text{NADH} + H^+ 
\]

However, Cameron et al. (6) reported that ARI-WAY-121,509, but not SDI-WAY-135,706, reversed impaired nerve conduction and sciatic nerve blood flow, and Obrososa et al. (34, 36) reported that SDI-157 failed to attenuate increases in retinal and sciatic nerve NADH/NAD⁺c or NADH/NAD⁺m in diabetic rats.

The current studies were therefore undertaken to clarify and further to investigate the (a) contributions of increases in NADH/NAD⁺c versus. NADH/NAD⁺m in mediating neural and vascular dysfunction evoked by diabetes, and (b) efficacy of SDI versus. ARI in preventing and reversing sorbitol pathway–linked increases in NADH/NAD⁺c and associated neural and vascular dysfunction.

Materials and Methods

Animal protocols

Male Sprague–Dawley rats were purchased from Sasco (O’Fallon, MO) and cared for in accordance with guidelines of the University Committee for the Humane Care of Laboratory Animals. Rats were housed one per cage and had free access to standard rat chow (Ralston Purina, Richmond, IN) and tap water. Diabetes was induced after an overnight fast by injection of 50 mg/kg body weight (bwt), zopolrestat was added to chow to provide 50 mg/kg bwt/day in the 5-week study and 100 mg/kg bwt/day in both reversal studies; sorbinil was added to chow to provide 50 mg/kg bwt/day in the 18-week study. Two structurally different ARIs were used to minimize the possibility that efficacy was an off-target effect. Dosages were based on dose–response effects of the SDI and ARIs (33). Body weights were monitored weekly, and consumption of drinking water and chow was checked every 2–3 days; quantities of inhibitors were adjusted to provide the appropriate dosage.

Motor-nerve conduction

Motor-nerve conduction velocity (MNCV) was measured in rats anesthetized with intramuscular injections of ketamine (100 mg/kg body wt). Core body temperature was maintained at 37±5°C with the use of heating pads and lamps controlled by a rectal temperature probe. Nerves were stimulated with a Grass stimulator (model SDP; Quincy, MA) by generating a rectangular pulse 0.2 ms in duration at a voltage sufficient to give a maximal response. The compound muscle action potential was amplified and recorded on a digital-storage oscilloscope 1425 (Gould, Oxnard, CA).

MNCV in the distal tibial branch of the sciatic nerve was quantified by insertion of stimulating electrodes through the skin at the popliteal fossa and the ankle to stimulate the nerve and insertion of a concentric bipolar recording electrode into the plantar muscle. MNCV was calculated by dividing the distance between the two stimulating electrodes by the la-
tency difference in the onset of the electromyographic (EMG) potentials in the recording electrodes evoked by the two stimulating electrodes. MNCV in the distal peroneal branch of the sciatic nerve was quantified by inserting stimulating electrodes at the sciatic notch and at the popliteal fossa to stimulate the nerve and by insertion of the concentric bipolar recording electrode into the anterior tibial muscle. MNCV was calculated as described for the tibial nerve. MNCVs are reported in meters per second (m/s).

**Blood flow and vascular permeability measurements**

One week after the last assessment of MNCV, sciatic nerve and retinal blood flows and vascular permeation (VAP) were assessed, as previously described (39, 40), and reported as microliters per gram (μl/g) wet wt/min and μg plasma/g wet wt/min, respectively. Rats were anesthetized with thiobutabarbital (Inactin), rather than ketamine, because it has a longer duration of action needed for preparation and assessment of blood flow and VAP. Blood flow was assessed by injection of 11.3 μm ⁴⁶Sc-labeled microspheres. VAP was quantified by the injection of [¹²⁵I]-bovine serum albumin followed by [³⁵S]-labeled microspheres. Mean arterial blood pressure (MABP) was monitored by inserting a polyethylene cannula filled with heparinized saline (connected to a pressure transducer) into the right subclavian artery (10).

**Metabolite measurements**

One week after the final assessment of MNCV, rats were again anesthetized with thiobutabarbital, and the portion of sciatic nerve between the sciatic notch and the popliteal fossa was rapidly removed, plunged into liquid nitrogen, and kept at −80°C before measurement of metabolites. Nerve polyols, lactate and pyruvate levels, plasma glucose levels, and HbA1c were quantified as described previously (30, 39, 45, 50). Plasma β-hydroxybutyrate was determined with the standard fluorescence enzymatic method in PCA-deproteinized samples (3). Plasma NEFA levels were assessed by using the WAKO NEFA-C test kit (Wako Chemicals, Richmond, VA).

The ratio of sciatic nerve molar concentrations of free cytosolic NADHc and free NAD⁺c was estimated by the redox metabolite indicator method (54). This method is based on the near-equilibrium between the NAD⁺/NADHc ratio and the ratio of the molar concentrations of lactate and pyruvate established by lactate dehydrogenase (LDH), as depicted in Eqs. 1 and 2. The value of the unitless equilibrium constant K_{LDH} is 1.111×10⁻⁴ (at pH 7.0) (54):

\[
\text{NAD}^+c + \text{lactate} \rightleftharpoons \text{NADH}_c + \text{pyruvate} \quad (\text{Eq. 1}).
\]

\[
\text{NADH}_c/\text{NAD}^+c = (\text{lactate/pyruvate}) \times K_{LDH} \quad (\text{Eq. 2}).
\]

Sciatic nerve molar ratios of free NADPH/NAD⁺c and mitochondrial NADH/NAD⁺m were assessed based on the near-equilibria established between (a) NADPH/NAD⁺c and malate/pyruvate ratios by the malic enzyme (19), and (b) NADH/NAD⁺m and β-hydroxybutyrate/acetoacetate ratios by β-hydroxybutyrate-DH (54).

**Statistical analysis**

All results are reported as mean±SD. Multiple analysis of variance was performed on Blomm-transformed data with the SAS general linear models procedure; overall differences among experimental groups for each parameter were first assessed by the Van der Waerden test, and individual pairwise group comparisons were evaluated with least-square means, as previously described (45).

**Results**

**5 weeks of diabetes**

Diabetes increased sciatic nerve endoneurial blood flow by 57%, VAP, 83%, and NADH/NAD⁺c, ~80% (Table 1). Retinal blood flow and VAP were increased 14% and 83%, respectively, and aortic VAP was increased 90% (from 98±16 μg plasma/g wet wt/min in controls to 191±52; p<0.001). SDI and ARI prevented all of these changes.

Diabetes increased sciatic nerve sorbitol and fructose levels by 12-fold and sevenfold, respectively, but decreased myoinositol levels 30%. The SDI increased sorbitol levels another 5.4-fold, whereas the ARI normalized them; nevertheless, both inhibitors decreased fructose levels ~80%.

Nerve sorbitol levels in SDI-treated controls equaled those in untreated diabetic subjects, whereas in ARI-treated controls, sorbitol levels were decreased to 22% of controls; fructose levels in SDI- and ARI-treated controls were significantly decreased versus controls; blood flow, VAP, and myoinositol levels in SDI- and ARI-treated controls did not differ from controls.

Plasma glucose levels in untreated diabetic rats were 25.7±2.7 mM vs. 6.2±0.7 in controls (p<0.001); glucose levels in SDI- and ARI-treated diabetic rats were 26.0±4.0 and 27.0±3.6 mM, respectively, and p<0.001 vs. controls for both groups. HbA1c levels were 11.1±2.0% in untreated diabetic rats vs. 3.7±1.3% in controls (p<0.001); HbA1c levels in SDI- and ARI-treated diabetic rats were 10.6±1.4% and 10.8±1.1%, respectively; p<0.001 vs. controls. Plasma glucose and HbA1c levels in SDI- and ARI-treated diabetics did not differ (p>0.05) from those in untreated diabetics.

Mean arterial blood pressure (MABP) was 128±9 mm Hg in diabetics vs. 144±14 in controls (p<0.01). Initial body weights were 240±12 g in diabetics vs. 259±17 in controls (p<0.02). Body weights increased 9.0±14% in diabetics vs. 33±11% in controls (p<0.001). MABP and body weights in SDI- and ARI-treated diabetics did not differ from those in untreated diabetics (p>0.05).

**18 weeks of diabetes**

Diabetes decreased MNCV 11.5% in the distal tibial nerve (Fig. 1) and 10.4% in the peroneal nerve (from 65.3±4.2 m/s to 58.5±5.1; p<0.01). Both inhibitors prevented these MNCV deficits. Diabetes increased sciatic nerve NADH/NAD⁺c and NADH/NAD⁺m by 80% and 60%, respectively (Fig. 1); both inhibitors prevented the increase in NADH/NAD⁺c, but not the increase in NADH/NAD⁺m.

Diabetes also increased sciatic nerve NADPH/NAD⁺c ratios ~80%, which also were normalized by the SDI (Table 2). The ARI partially attenuated the increase in NADPH/NAD⁺c (which did not differ significantly from controls or diabetics) but did not affect malate/pyruvate ratios.

Plasma glucose levels in diabetic rats were increased ~sixfold (39.7±3.6 mM vs. 6.9±0.5 in controls; p<0.001);
Neither the SDI nor the ARI significantly attenuated NEFA or treated diabetics.

Table 1. 5 Weeks of Diabetes: Effects of the SDI and the ARI (Zopolrestat) on Blood Flow and Vascular Albumin Permeation in Sciatic Nerve Endoneurium and Retina and on Sciatic Nerve NADH/NAD⁺c, Sorbitol, Fructose, and Myoinositol Levels

<table>
<thead>
<tr>
<th></th>
<th>Blood flow</th>
<th>VAP</th>
<th>NADH/NAD⁺c</th>
<th>Sorbitol</th>
<th>Fructose</th>
<th>Myoinositol</th>
<th>Blood flow</th>
<th>VAP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>65 ± 16</td>
<td>76 ± 13</td>
<td>1.1 ± 0.4</td>
<td>159 ± 55</td>
<td>814 ± 197</td>
<td>4,198 ± 1692</td>
<td>374 ± 15</td>
<td>75 ± 12</td>
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<tr>
<td>+SDI</td>
<td>64 ± 17</td>
<td>75 ± 13</td>
<td>-</td>
<td>2,050 ± 697*</td>
<td>524 ± 201*</td>
<td>4,351 ± 1290</td>
<td>360 ± 20</td>
<td>71 ± 8</td>
</tr>
<tr>
<td>+ARI</td>
<td>68 ± 30</td>
<td>76 ± 18</td>
<td>-</td>
<td>24 ± 11*</td>
<td>179 ± 60*</td>
<td>4,283 ± 938</td>
<td>376 ± 16</td>
<td>74 ± 19</td>
</tr>
<tr>
<td>Diabetic</td>
<td>102 ± 17*</td>
<td>139 ± 16*</td>
<td>1.9 ± 0.4</td>
<td>1,863 ± 623*</td>
<td>5,815 ± 1,711*</td>
<td>2,875 ± 737*</td>
<td>426 ± 21*</td>
<td>137 ± 16*</td>
</tr>
<tr>
<td>+SDI</td>
<td>67 ± 26*</td>
<td>89 ± 25*</td>
<td>1.3 ± 0.4*</td>
<td>9,975 ± 4397*</td>
<td>1,001 ± 519*</td>
<td>2,844 ± 793*</td>
<td>380 ± 45*</td>
<td>85 ± 24*</td>
</tr>
<tr>
<td>+ARI</td>
<td>75 ± 24*</td>
<td>94 ± 3*</td>
<td>1.2 ± 0.4*</td>
<td>170 ± 85*</td>
<td>1,217 ± 253*</td>
<td>3,521 ± 1,229</td>
<td>382 ± 33*</td>
<td>91 ± 38*</td>
</tr>
</tbody>
</table>

BF, blood flow; VAP, vascular albumin permeation. Data are expressed as mean ± SD for 10–12 animals. Blood flow is expressed as ml/g wet wt/min; vascular albumin permeation is expressed as µg plasma/g wet wt/min; sorbitol, fructose, and myoinositol levels are expressed as nmol/g wet wt; see Research Design and Methods for calculation of free NADH/NAD⁺ c × 10⁻³.

Different from Controls: *p < 0.001, †p < 0.005, ‡p < 0.04.
Different from Diabetics: ipsis p < 0.001, ††p < 0.01.

HbA1c levels were increased by ~3.3-fold (14.7 ± 0.6% vs. 4.5 ± 0.2% in controls; p < 0.001). Plasma glucose and HbA1c levels in ARI- and SDI-treated diabetic rats did not differ from those in untreated diabetics (Table 2).

Nerve sorbitol and fructose levels were increased by diabetes by 13.4- and 5.7-fold, respectively (Table 2). The SDI increased sorbitol levels another 3.9-fold and normalized fructose levels. The ARI reduced sorbitol and fructose levels to levels lower than those in controls (p < 0.01). Myoinositol levels were decreased ~37% by diabetes and were normalized by the ARI but unaffected by the SDI (p < 0.0001 vs. SDI-treated diabetics).

Plasma levels of NEFA were 69 ± 17 µEq/dl in diabetics versus 42 ± 8 in controls (p < 0.004); plasma β-HBA levels were 102 ± 51 µM in diabetics vs. 27 ± 3 in controls (p < 0.001). Neither the SDI nor the ARI significantly attenuated NEFA or β-HBA levels in diabetics. Initial body weights were similar in controls and diabetics; however, diabetics gained significantly less weight than did controls (p < 0.05).

Reversal of MNCV deficits and NADH/NAD⁺c by SDI and ARI

Reversal Study 1. After 6 weeks of untreated diabetes, the MNCV was decreased ~13–14% in all three groups of diabetic rats (Fig. 2, left panel). After 8 weeks of untreated diabetes, MNCV was decreased 17% (Fig. 2, middle panel). SDI and ARI administered during the last 2 weeks attenuated (but did not normalize) MNCV deficits to 11% and 4.6%, respectively. After 10 weeks of untreated diabetes, MNCV was decreased 14% and was statistically normalized by SDI and by ARI administered during the last 4 weeks (Fig. 2, right panel).

Diabetes increased sciatic nerve NADH/NAD⁺c and NADH/NAD⁺m about twofold (Table 3). The ARI normal-
failed to normalize NADH with the substantially larger variance in NADH in diabetics and ARI-treated diabetics and the unfavorable pharmacokinetics of SDI, as discussed earlier (31, 33, 45).

Diabetes increased plasma glucose levels 4.4-fold (35.1 ± 4.7 mM vs. 8.0 ± 1.8, p < 0.001) and increased HbA1c levels 3.8-fold (14.1 ± 1.0% vs. 3.7 ± 0.4%; p < 0.001). Plasma glucose levels and HbA1c levels in diabetic rats treated with SDI and ARI did not differ from those in untreated diabetics (Table 3).

Diabetes increased NEFA and β-HBA levels by about two- and fourfold; p ≤ 0.001. All three groups of diabetics failed to gain weight in contrast to controls. These parameters were unaffected by the SDI or ARI.

Reversal Study 2. After 6 weeks of untreated diabetes, MNCV deficits ranged from 9.3 and 14.3% before administration of SDI and ARI (Table 4). After 8 weeks of untreated diabetes, the MNCV deficit was 10.7% and was not attenuated by SDI or ARI treatment during the last 2 weeks. After 10 weeks of untreated diabetes, the MNCV deficit was 10.4% and was normalized by SDI and by ARI administered during the last 4 weeks. Four weeks of ARI treatment also normalized the associated 19% reduction in nerve blood flow; however, blood flow in SDI-treated diabetics was not significantly different from that in controls or untreated diabetics. Neither inhibitor attenuated the associated 2.5-fold increase in VAP. MABP was 132 ± 18 mm Hg in diabetics versus 148 ± 8 mm Hg in controls (p < 0.01) and was unaffected by the SDI or ARI.

Diabetes increased sciatic nerve sorbitol and fructose levels by 12-fold and sevenfold, respectively (Table 4). The SDI increased sorbitol levels an additional fivefold (61 vs. 34.4 fold), while normalizing fructose levels. The ARI decreased sorbitol and fructose levels 35% lower than those in SDI diabetics and were unaffected by the SDI; however, the ARI normalized myoinositol levels, which were 1.7-fold higher than those in untreated diabetics (p < 0.02). Myoinositol levels were decreased 57% by diabetes and were unaffected by the SDI; however, the ARI normalized myoinositol levels, which were 1.7-fold higher than those in untreated diabetics (p < 0.01) and ~1.9-fold higher than those in SDI diabetics (p < 0.001).

Diabetes increased plasma glucose levels 4.5-fold (36.2 mM ± 3.8 vs. 8.0 ± 1.0 in controls; p ≤ 0.001); plasma glucose levels in SDI- and ARI-treated diabetics were 34.4 ± 3.5 and 35.0 ± 3.7 mM, respectively (p ≤ 0.001 vs. controls) and did not differ from those in untreated diabetics (p > 0.05).

Diabetes also increased HbA1c levels by 3.9-fold (14.0 ± 1.4% vs. 3.6 ± 0.2% in controls; p ≤ 0.001); HbA1c levels in SDI- and ARI-treated diabetics were 13.4 ± 1.6% and 13.4 ± 1.1%, respectively (p ≤ 0.001 vs. controls) and did not differ from those in untreated diabetics (p > 0.05).

**FIG. 2.** Reversal Study 1. Time course of reversal of impaired distal tibial MNCV by treatment with an SDI (200 mg/kg bwt/day) vs. an ARI (zopolrestat, 100 mg/kg bwt/day) initiated after 6 weeks of untreated diabetes. Mean ± SD, n = 9–10 rats for all parameters. Different from controls: *p < 0.001, †p < 0.05. Different from diabetics: #p < 0.001, §p < 0.04. Different from diabetics + SDI: ‡p = 0.013.
Controls 1.4 ± 0.5 32 ± 14 8.0 ± 1.8 3.7 ± 0.4 40 ± 12 27 ± 6 382 ± 7 +48 ± 8
Diabetics 2.9 ± 0.9 64 ± 19 35.1 ± 4.7 14.1 ± 1.0 73 ± 24 107 ± 99 380 ± 8 −17 ± 14
+SDI 2.5 ± 1.4 49 ± 30 36.6 ± 9.4 13.5 ± 0.9 65 ± 26 130 ± 103 397 ± 18 +0.9 ± 0.9
+ARI 1.8 ± 0.7 48 ± 18 33.4 ± 11.9 13.0 ± 0.9 75 ± 31 99 ± 57 377 ± 12 +3.8 ± 10

<table>
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<tr>
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Sciatic nerve NADH/NAD⁺c and NADH/NAD⁺m, HbA₁c, and plasma glucose (PG), nonesterified fatty acids (NEFA), and β-hydroxybutyrate (β-HBA) levels, initial body weights (BWIs) and body-weight change (BWC) for rats in which MNCV data are depicted in Fig. 2.

Data are expressed as mean ± SD for seven to 10 animals. NADH/NAD⁺c and NADH/NAD⁺m are calculated as described in the Research Design and Methods section and expressed as NADH/NAD⁺c×10⁻³ and NADH/NAD⁺m×10⁻³; plasma glucose levels are expressed as mM; HbA₁c, as gEq/dl; β-HBA as (μM); BWI in grams; and BWC as a percentage change from BWI.

Different from Diabetics: *p < 0.001; †p < 0.002; ‡p < 0.01; §p < 0.02; ¶p < 0.04.
Different from Controls: *p = 0.013.

Diabetes tended to increase NEFA (58 ± 21 μEq/dl vs. 42 ± 14 μEq/dl in controls); however, the difference was not significant (p > 0.05). NEFA levels in SDI- and ARI-treated diabetics were slightly higher than those in untreated diabetics (71 ± 39 and 77 ± 26 μEq/dl, respectively; p < 0.01 vs. controls for both groups) but did not differ from those in untreated diabetics (p > 0.05). Diabetes increased β-HBA levels by ~5.4-fold to 113 ± 84 μM vs. 21 ± 4 in controls (p < 0.001); β-HBA levels in SDI- and ARI-treated diabetics were 71 ± 39 and 77 ± 26, respectively (p < 0.01 vs. controls and p > 0.05 vs. untreated diabetics). Initial body weights were 417 ± 11 g in controls and diabetics; final body weights increased 35 ± 7% in controls vs. a weight loss of −9 ± 9% in untreated diabetics (p < 0.001). Weight losses in SDI- and ARI-treated diabetics were −6 ± 4% and −6 ± 10%, respectively (p < 0.05 vs. untreated diabetics).

**Effects of diabetes, SDI, and ARI on sciatic nerve:**
(a) malate levels and malate/pyruvate ratios and (b) lactate/pyruvate ratios manifested by changes in lactate and/or pyruvate levels Table 5

**Effects of untreated diabetes.** Lactate/pyruvate ratios were increased by 1.5- to twofold in all three studies and were attributable largely to increased lactate levels, except in the 18-week study in which lactate levels were identical to controls, whereas pyruvate levels were decreased 43% (Table 5). In the 10-week study, the increased lactate/pyruvate ratio also was contributed to by a 19% decrease in pyruvate levels that approached statistical significance (p = 0.05–0.1). Malate levels did not differ from controls, whereas malate/pyruvate ratios were increased ~80% (p < 0.001 vs. controls).

**Effects of SDI treatment.** The SDI normalized lactate/pyruvate ratios in the 5- and 18-week studies, but not in the 10-week reversal study. In the 5-week study, 87% normalization of lactate/pyruvate ratios was due largely to increased (33%) pyruvate levels, together with decreased (8%) lactate levels. In the 18-week study, 83% normalization was due entirely to increased pyruvate levels (77%). Malate levels did not differ from those in controls or diabetics, whereas malate/pyruvate ratios were normalized.

**Effects of ARI treatment.** The ARI normalized lactate/pyruvate ratios in all three studies. In the 5-week study, normalization was manifested by the combination of a marginally significant increase in pyruvate levels (28%; p = 0.05–0.01) and a 17% (p = 0.26) reduction in mean lactate levels. In the 10-week study, normalization was due to significantly increased pyruvate levels. In the 18-week study, normalization was due to significantly increased pyruvate levels and a statistically insignificant reduction in lactate levels. Malate levels and malate/pyruvate ratios were not significantly different from those in the other groups.

Table 3. Reversal Study 1

<table>
<thead>
<tr>
<th>NADH/ NAD⁺c</th>
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<td>32 ± 14</td>
<td>8.0 ± 1.8</td>
<td>3.7 ± 0.4</td>
<td>40 ± 12</td>
<td>27 ± 6</td>
<td>382 ± 7</td>
</tr>
<tr>
<td>Diabetics</td>
<td>2.9 ± 0.9</td>
<td>64 ± 19</td>
<td>35.1 ± 4.7</td>
<td>14.1 ± 1.0</td>
<td>73 ± 24</td>
<td>107 ± 99</td>
<td>380 ± 8</td>
</tr>
<tr>
<td>+SDI</td>
<td>2.5 ± 1.4</td>
<td>49 ± 30</td>
<td>36.6 ± 9.4</td>
<td>13.5 ± 0.9</td>
<td>65 ± 26</td>
<td>130 ± 103</td>
<td>397 ± 18</td>
</tr>
<tr>
<td>+ARI</td>
<td>1.8 ± 0.7</td>
<td>48 ± 18</td>
<td>33.4 ± 11.9</td>
<td>13.0 ± 0.9</td>
<td>75 ± 31</td>
<td>99 ± 57</td>
<td>377 ± 12</td>
</tr>
</tbody>
</table>

Also shown are sciatic nerve endoneurial blood flow (BF) and VAP (μg plasma/g/min) and sciatic nerve sorbitol, fructose, and myoinositol levels 4 weeks of treatment. Data are expressed as mean ± SD for nine to 10 animals.

Different from Controls: *p < 0.001; †p < 0.002; ‡p < 0.01.  
Different from Diabetics: *p < 0.001; †p < 0.002; ‡p < 0.01.  
Different from ARI-Rx Diabetics: *p < 0.001; †p < 0.01, ‡p < 0.02.
Table 5. Effects of Diabetes, SDI, and ARI on Sciatic Nerve: Lactate and Pyruvate Levels, and Lactate/Pyruvate (L/P) Ratios in Rats with Diabetes of 5-, 10-, and 18-Weeks Duration, and Malate and Malate/Pyruvate (M/P) Ratios in Rats with Diabetes of 18-Weeks Duration

<table>
<thead>
<tr>
<th>5 wk</th>
<th>10 wk</th>
<th>18 wk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td>Lactate</td>
<td>Pyruvate</td>
</tr>
<tr>
<td>3.52 ± 1.19</td>
<td>0.32 ± 0.08</td>
<td>20.3 ± 0.68</td>
</tr>
<tr>
<td>11.56 ± 4.32</td>
<td>3.72 ± 1.36</td>
<td>12.70 ± 4.39</td>
</tr>
<tr>
<td>Diabetics</td>
<td>Lactate</td>
<td>Pyruvate</td>
</tr>
<tr>
<td>6.36 ± 2.37</td>
<td>0.36 ± 0.13</td>
<td>3.13 ± 0.89</td>
</tr>
<tr>
<td>18.17 ± 4.41</td>
<td>0.48 ± 0.20</td>
<td>27.6 ± 0.88</td>
</tr>
<tr>
<td>D+SDI</td>
<td>Lactate</td>
<td>Pyruvate</td>
</tr>
<tr>
<td>5.88 ± 2.93</td>
<td>0.48 ± 0.20</td>
<td>3.76 ± 0.88</td>
</tr>
<tr>
<td>12.43 ± 3.19</td>
<td>0.48 ± 0.20</td>
<td>3.87 ± 1.57</td>
</tr>
<tr>
<td>D+ARI</td>
<td>Lactate</td>
<td>Pyruvate</td>
</tr>
<tr>
<td>5.28 ± 2.06</td>
<td>0.46 ± 0.16</td>
<td>16.13 ± 655</td>
</tr>
<tr>
<td>11.61 ± 3.24</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data are expressed as mean ± SD for lactate and pyruvate levels and mean L/P ratios for animals in each group. Numbers of animals for each duration of diabetes are shown in corresponding Tables 1 and 2 and Fig. 1.

Lactate, pyruvate, and malate levels are expressed as nmol/µg DNA.

Different from Controls: *p < 0.001, †p < 0.05, ‡p < 0.01.
Different from Diabetics: *p < 0.004, †p < 0.01, ‡p < 0.05.
Different from Diabetics + SDI: §p = 0.02.

Discussion

The observations in these experiments support two fundamental conclusions and concepts: (a) increased sorbitol oxidation plays a critical role in mediating diabetes-induced early neural and vascular dysfunction that precede pathologic structural changes, and (b) neural and vascular dysfunction evoked by sorbitol oxidation are primarily sequelae of increased cytoplasmic NADH/NAD+ c rather than elevated mitochondrial NADH/NAD+ m. Notably, these are the first studies in which neural or vascular dysfunction or both, NADH/NAD+ c or NADH/NAD+ m, or both have been assessed in the same animal. Previous investigators have assessed neural and vascular dysfunction but not NADH/NAD+ c or NADH/NAD+ m or both have been assessed in the same animal. Previous investigators have assessed neural and vascular dysfunction but not NADH/NAD+ c or NADH/NAD+ m; but not neural or vascular dysfunction (34).

The first conclusion is supported by the comparable efficacy of SDI and ARI in preventing and reversing MNCV deficits (Figs. 1 and 2, Table 4) and in preventing vascular dysfunction in sciatic nerve, retina, and aorta in rats with diabetes of 5 weeks' duration (Table 1).

In two independent studies, the SDI and the ARI reversed MNCV deficits (Fig. 2, Table 4) after 4 weeks of treatment, after 6 weeks of untreated diabetes. We confirm the previous observation of Cameron et al. (6) that SDI treatment for 2 weeks failed to reverse preexisting MNCV deficits. We observed in addition, however, that SDI treatment for 4 weeks was as efficacious as ARI in reversing MNCV deficits (Fig. 2). The biochemical explanation for clear SDI efficacy after 4 (but not after 2) weeks of treatment is unclear and will require future studies. The same SDI (albeit named differently) was used by Cameron et al. and by us. [Chemical structures of ARI and SDI used by Cameron et al. (6), Obrosova et al. (34), and by us are shown in OAS V.]

Failure of the ARI zopolrestat to reverse MNCV deficits after 2 weeks of treatment in our studies is likely attributable to its lower in vivo potency versus the ARI WAY-121,509 used by Cameron et al. Consistent with this notion, sciatic nerve fructose levels in WAY-121,509-treated diabetics were markedly reduced to only 16% of fructose levels in nondiabetic controls; in contrast, fructose levels in zopolrestat-treated diabetics were 76% of fructose levels in nondiabetic controls. The suggestion by Cameron et al. (6) that MNCV deficits in untreated diabetics are attributable to decreased blood flow (i.e., ischemia), even in diabetes of short duration (6, 7), is discordant with the current observations and with the findings of Chang et al. (10), Ido et al. (21), and others addressed in OAS II-A.

Treatment of diabetic rats with SDI-157 (also named S-0773) in an earlier study by Obrosova et al. (34) failed to prevent increases in neural NADH/NAD+ c. That finding differs from observations in the current experiments in which treatment with CP-166,572 (also named SDI-158) normalized sciatic nerve NADH/NAD+ c (Fig. 1, Table 1) and was as efficacious as ARI in preventing MNCV deficits in all three experiments in which MNCV was assessed (Figs. 1 and 2, Table 4). (Chemical structures of SDIs shown in OAS V.)

SDI-157 is a produg that is activated in vivo in the liver to the active chemical species SDI-158 (13). SDI-157 was administered at a dose of 100 mg/kg bwt/day, whereas the SDI-158 (CP-166,572) was administered at a twofold higher dose of 200 mg/kg bwt/day. Because the half-life of SDI-157 and SDI-158 in serum and nerve of rats is very short (31-page 359), i.e., <30 min, the duration of its pharmacologic effect is expected to be highly sensitive to the plasma level achieved. Moreover, the efficacy of SDI also is likely to be sensitive to variability in tissue drug levels because SDH blockade markedly elevates nerve levels of sorbitol substrate for SDH (e.g., about four- to fivefold vs. untreated diabetics (Tables 1, 2, and 4); as SDH blockade wanes, rates of sorbitol oxidation can exceed rates in untreated diabetics (45).

The 44% higher coefficient of variation of NADH/NAD+ c ratios in SDI-treated diabetics versus ARI-treated diabetics (Table 3) suggests that inhibitory levels of the short-half-life SDI were not maintained in some of the animals in this group, despite the high dose of SDI administered (200 mg/kg bwt/day) and the water-dosing paradigm. This probability is consistent with evidence that a very high degree of SDH inhibition (i.e., >91% normalization of nerve fructose levels) must be achieved to demonstrate SDI efficacy, as
demonstrated for ARI (6). Thus, we hypothesize that the failure of SDI to normalize neural NADH/NAD$^+$c in the earlier experiment (34) may be explained by the combined effects of (a) the 50% lower dose administered of the prodrug SDI-157 versus the active drug SDI-158 in the present experiments, (b) the rapid build-up of sorbitol substrate for SDI caused by the SDI, and (c) the very short half-life of the active SDI.

Therefore, the suggestion that hypoxia-like metabolic changes [i.e., increases in NADH/NAD$^+$c in tissues of diabetic rats (55)] are sequelae of increased AR activity, independent of increased sorbitol oxidation (36) remains discordant with the present observations, as well as with previous findings demonstrating that exposure of normoxic normoglycemic cells and tissues to elevated sorbitol levels in vivo and in vitro evokes metabolic imbalances or vascular dysfunction or both, like those evoked by hyperglycemia (9, 12, 26, pages 9–10 in OAS IV-A of 30, 46, 47, 49, 53).

Likewise, the suggestion (4) that the increase in free NADH/NAD$^+$c evoked by diabetes is attributable to a marked decrease in total NAD$^+$t because of its consumption by poly(ADP-ribose) polymerase (PARP) is inconsistent with (a) increased levels of total NADHt with no change in NADt (NADH plus NAD$^+$t) (14) and increased levels of NADHt and NAD$^+$t (18) in retinas of diabetic rats, (b) the insignificant decrease in free NAD$^+$c associated with a twofold increase in NADH/NAD$^+$c discussed later, (c) the demonstration that hyperglycemia increases free NADH/NAD$^+$c = lactate/pyruvate $\times K_{LDH}$ ratios in human erythrocytes (48) that lack nuclei and mitochondria, and (d) observations of other investigators addressed in OAS II-E. Interestingly, recent observations suggest that activated PARP may further contribute to oxidative–nitrosative stress (OAS II-E, 9).

The second conclusion (see earlier; i.e., the primacy of increased NADH/NAD$^+$c vs. increased NADH/NAD$^+$m in mediating diabetes-induced neural and vascular dysfunction) is supported by observations that (a) SDH and AR are known to be primarily cytoplasmic enzymes (31), and (b) inhibition of either enzyme normalized/attenuated increased sciatic nerve free NADH/NAD$^+$c (Fig. 1; Table 1), impaired MNVC (Figs. 1 and 2; Table 4), and vascular dysfunction (Table 1), while having little or no impact on associated increases in free NADH/NAD$^+$m (Fig. 1; Table 3). Whatever putative mitochondrial changes may have contributed to neural and vascular changes, independent of the increase in NADH/NAD$^+$m, the observations that ARI and SDI prevented vascular and neural dysfunction indicates that those putative changes also were prevented. The most likely explanation for the increases in NADH/NAD$^+$m in these experiments is increased oxidation of fatty acids and/or ketones that were significantly increased in Reversal Study 1 (Table 3) and in the 18-week study (see Results) and were not attenuated by ARI or SDI.

Limitations of the current studies are that they do not directly address (a) potentially important increased superoxide production caused by hypoxic/ischemic vascular disease in chronic diabetes (OAS I-C2) or (b) mitochondrial superoxide production. The effects of the two types of inhibitors of cytoplasmic enzymes (AR and SDH) in the present studies do not exclude potentially important downstream enzymatic reactions/sequelae in the same or other cellular compartments. A positive control to normalize NADH/NAD$^+$m, but not NADH/NAD$^+$c, would be of interest for future studies.

The possible relationship between increased mitochondrial leakage, i.e., opening of MPTPs (1) and changes in NADH/NAD$^+$m also need further study. Because the effects of diabetes on NADH/NAD$^+$c and NADH/NAD$^+$m are complex and their roles in initiating complications of diabetes remain controversial (4–8, 10, 11, 21, 29, 34, 36, 56), they are discussed critically in more detail in the current OAS II and in the OAS in (30) for interested readers. Notably, observations and speculations thought to be inconsistent with the importance of sorbitol oxidation do not withstand scrutiny.

An additional limitation of the current studies is the question of the absolute specificity of the inhibitors used. Off-target effects are always possible for low-molecular-weight inhibitors. This is the reason that two structurally distinct inhibitors of AR were used (i.e., sorbinil and zopolrestat). That similar results were obtained with either ARI attests to the salutary effects being a result of inhibition of the intended target, AR. The result is further supported and refined by using the SDI, an inhibitor in the second step of the pathway. However, only one structural class of SDIs is active in vivo, the one used. SDI normalized impaired nerve conduction in diabetic rats, but paradoxically evoked ultrastructural changes in autonomic nerves (iliac mesenteric nerves) and in prevertebral sympathetic superior mesenteric ganglia in the same animals (OAS V-A in ref.30). The functional significance and mechanism of those ultrastructural changes remains unclear.

The fundamental issue raised by the current observations is the basic mechanism by which NADHc generated by oxidation of sorbitol causes (a) metabolic imbalances and associated neural and vascular dysfunction observed in early diabetes, and (b) late complications of diabetes that are now generally agreed to be linked to excess production of ROS (4, 5, 20, 29, 53). Equations 1 and 2 (see Methods) are the keys to understanding this mechanism and the pathophysiologic importance of sorbitol oxidation in generating relatively small molar increases in free NADHc, uncoupled from subsequent downstream production of equimolar pyruvate.

As shown in Eq. 2, free NADH/NAD$^+$c $\times K_{LDH}$ ($K_{LDH} = 1.11 \times 10^{-4}$ at $pH = 7.0$; i.e., ~1/10,000). Molar lactate/pyruvate ratios in a wide spectrum of normal cells and tissues range from ~10/1 to 20/1 [e.g., liver (54) and nerve (Table 5)]. Therefore, based on Eq. 2, a lactate/pyruvate ratio of 10:1 corresponds to a free NADH/NAD$^+$c ratio of ~10/10,000 or ~1/1000 or ~0.001 (i.e., ~1 mole of NADHc per 1,000 moles of NAD$^+$c). In light of this large difference in concentrations of NADHc and NAD$^+$c, molar changes in NADHc have a far greater impact on NADH/NAD$^+$c and lactate/pyruvate ratios than do equimolar changes in pyruvate, lactate, or NAD$^+$c, in decreasing order of impact. For example, accumulation of just 1 additional mole of NADHc per 1,000 moles of NAD$^+$c will increase NADHc levels by about twofold (from 1 to 2 moles of NADHc per 1,000 moles of NAD$^+$c). This will be manifested by a corresponding approximately twofold increase in NADH/NAD$^+$c, from ~0.001 to ~0.002, and in lactate/pyruvate ratios from ~10 to ~20, as observed (a) in the current studies in the 5-week experiment (Tables 1 and 5), the 10-week Reversal Study 1 (Tables 3 and 5), and the 18-week experiment (Fig. 1; Table 5); (b) in previous reports in retina, sciatic nerve, and liver of diabetic animals (17, 34–37, 41, 54), and (c) normal rat retinas and sciatic nerve endoneuria incubated in 30 mM glucose that are prevented by ARI (30, and current OAS IV).
If the 1 additional mole of NADHc per 1,000 moles of NAD\(^+\)c were generated by reduction of 1 mole of NAD\(^+\)c to NADHc \textit{via} oxidation of sorbitol, then SDI or ARI should prevent the increase in NADH/NAD\(^+\)c, as observed in these experiments (Fig. 1; Table 1, and by the ARI in Table 3). In contrast, if the increases in NADH/NAD\(^+\)c were due to increased NADHc formation independent of sorbitol oxidation (e.g., by hypoxia), ARI and SDI would not be expected to affect it. This prediction has been tested by Nyengaard \textit{et al.} (30) on the additive effects of hyperglycemia and hypoxia on free NADH/NAD\(^+\)c (manifested by increases in lactate/pyruvate ratios) in incubated normal rat retinas and sciatic nerve.
endoneuropathy (OAS IV). Hyperglycemia and hypoxia additively increased lactate/pyruvate ratios, triose phosphates, and G3P; the increases evoked by hyperglycemia were prevented by ARI, which did not affect the associated increases evoked by hypoxia.

The evidence that accumulation of just 1 mole of free NADH per 1,000 moles of free NAD⁺ c doubles the concentration of free neural NADHc and is associated with increased neural blood flow suggests that (a) the level of free NADHc is an exquisitely sensitive blood flow sensor due to "hyperglycemic pseudohypoxia" (55), as well as true hypoxia, manifested by increases in NADH/NAD⁺ c; and (b) increased levels of NADHc fuel signaling pathways that augment blood flow, (e.g., Table 1) (22, 23, 27, 51). This interpretation is consistent with observations (46) that exposure of granulation tissue to elevated levels of sorbitol (at normal glucose levels) or hyperglycemia increased blood flow and VAP (identical to those in nerve and retina in Table 1) that were prevented by SDI, SOD, or VEGF antibodies.

The near-equilibrium between free NADH/NAD⁺ c and lactate/pyruvate ratios maintained by LDH also plays a pivotal role in modulating energy metabolism, blood flow, and superoxide production in a spectrum of physiologic and pathologic conditions, including hypoxia and increased physiologic work in humans and animals addressed in OAS-I-C. Thus, infusion of lactate (which produces "prooxidant" NADHc when oxidized by LDH) (a) increases blood flow in sciatic nerve, retina, and other tissues in normal rats that are prevented by co-infusion of "antioxidant" pyruvate (23); and (b) augments the already increased blood flows evoked by increased physiologic work in normal humans and animals that also are prevented by co-infusion of pyruvate (22, 23, 27, 51, and OAS 1-C).

Observations in the 18-week study that diabetes increased sciatic nerve NADPH/NAD⁺ c (Table 2) and NADH/ NAD⁺ c by ~1.8-fold (Fig. 1), both of which were prevented by the SDI, are consistent with corresponding observations in an earlier report that were prevented by an ARI (35). Although the explanation for the increase in NADPH/ NAD⁺ c is not entirely clear, evidence that increased levels of NADHc generated by oxidation of sorbitol fuel superoxide production (46) and diacylglycerol synthesis (57) suggests that increased levels of NADHc might also fuel reduction of oxidized NAD⁺ c to NADPHc via the pyruvate/malate cycle (42), discussed in OAS I-B. This possibility is consistent with evidence that 40% of the NADPH+H⁺ used for fatty acid synthesis from glucose is provided by the pyruvate/malate cycle. These observations suggest that increased NADHc generated by oxidation of sorbitol may augment superoxide production by three mechanisms: (a) as substrate for NADH-driven oxidases, (b) as substrate for the pyruvate/malate cycle to increase NADPHc as fuel for NADPH-driven oxidases, and (c) by fueling DAG synthesis, which activates the PKC that activates some isoforms of NADPH oxidase (Fig. 3). The third mechanism is supported by observations (57) that exposure of skin-chamber granulation tissue vessels to elevated glucose levels in nondiabetic rats (circumventing other metabolic and hormonal imbalances associated with the diabetic milieu) increases DAG levels and VAP, both of which were prevented by addition of pyruvate (which compensates for the uncoupling of pyruvate production when NADHc is produced by sorbitol oxidation rather than by glycolysis).

Taken together, observations in these experiments support an important role for increased "prooxidant" NADHc generated by sorbitol oxidation in fueling superoxide production, metabolic imbalances, and neural and vascular dysfunction evoked by hyperglycemia. Uncoupling of prooxidant NADHc formed by sorbitol oxidation from equimolar formation of "antioxidant" pyruvate constrains reoxidation of NADHc to NAD⁺ c by LDH coupled to the reduction of pyruvate.

**FIG. 3** NADHc generated by sorbitol oxidation is reoxidized to NAD⁺ c largely by enzymes that promote the development of diabetic complications. (A) Under steady-state normoglycemic conditions, glucose is metabolized via glycolysis to synthesize ATP [in the cytoplasm via substrate phosphorylation (SP) and in mitochondria via oxidative phosphorylation (OP)] only after electrons and protons are transferred from G3P to oxidized NAD⁺ c (by G3P-DH), reducing it to NADHc followed by downstream formation of pyruvate equimolar to NADHc. Electrons and protons carried by NADHc can be transferred into mitochondria via the malate-aspartate (Mal-Asp) and glycerol phosphate (G3P) electron shuttles (Θ); pyruvate can be transferred into mitochondria via a mitochondrial pyruvate carrier. All cells generate NADHc and pyruvate via glycolysis faster than they are used for ATP synthesis in mitochondria. Excess NADHc is reoxidized to NAD⁺ c largely by lactate dehydrogenase (LDH), coupled to reduction of excess pyruvate to lactate (Θ) that diffuses out of the cell (e.g., for retina: 30, 50; for nerve: OAS IV). Much smaller amounts of glucose are metabolized by the sorbitol pathway through which sorbitol dehydrogenase (SDH) catalyzes oxidation of sorbitol to fructose coupled to reduction of equimolar NAD⁺ c to NADHc. However, SDH does not generate pyruvate required for reoxidation of NADHc to NAD⁺ c by LDH. This NADHc is reoxidized to NAD⁺ c largely via alternative pathways (Θ): (1) NADH-driven oxidases that generate basal levels of superoxide (O₂⁻) and (2) glucose 3-phosphate-DH (G3P-DHc), which fuels synthesis of basal levels of diacylglycerol (DAG) and protein kinase C (PKC) activity. Both of these mechanisms of reoxidation of NADHc to NAD⁺ c maintain normal steady-state ratios of NADH/NAD⁺ c = (lactate/pyruvate) × K₅LDH (Eq. 2 in Methods). (B) Hyperglycemia augments glucose metabolism via glycolysis and via the sorbitol pathway. In normal rat retina and sciatic nerve endoneuropathy incubated in 30 vs. 5 mM glucose, glycolysis was increased ~1.5-fold [manifested by increased lactate production (30, 50, current OAS-IV). Oxidation of sorbitol in the same endoneuropathy (manifested by production of fructose equimolar to NADHc) was increased ~11-fold. Because oxidation of sorbitol does not form pyruvate, NADHc levels increase by 1.5- to twofold (manifested by increased ratios of NADH/NAD⁺ c in Fig. 1, Tables 1 and 3; refs. 17, 34–37, 41), and much more NADHc is reoxidized to NAD⁺ c via alternative pathways (Θ) (i.e., by NADH-driven oxidases that increase superoxide production and by G3P-DHc, which increases synthesis of diacylglycerol (DAG), activation of protein kinase C (PKC), and activation of NADH-driven oxidases to augment superoxide production further. The 1.5- to twofold increases in NADHc also may contribute to product inhibition of G3P-DH, which increases levels of G3P and DHAP (i.e., triose phosphates-TP (30, 48, 50, OAS IV) that are in equilibrium. Nonenzymatic degradation of TP (primarily DHAP) forms methylglyoxal (MG), a potent nonenzymatic glycation agent. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article at www.liebertonline.com/ars).
ruvate to lactate that diffuses out of cells (Fig. 3). Thus, NADHc levels increase (~1.5- to twofold) until their mass-action effect drives reoxidation of NADHc to NAD⁺c by (a) NADH-driven oxidases that produce superoxide, and (b) G3P-DHc that promotes \( \text{de novo} \) synthesis of DAG and activation of PKC (Fig. 3) that activates some isoforms of NADPH-driven oxidases.

Observations that ARI and SDI increased sciatic nerve pyruvate levels that were statistically significant or approached significance in five of six interventions (Table 5) were associated with (a) normalization of MCNC deficits, and (b) normalization/attenuation of associated increases in NADH/NAD⁺c and vascular dysfunction that are consistent with well-known antioxidant effects of pyruvate (OAS I). Pyruvate (a) stoichiometrically degrades \( \text{H}_2\text{O}_2 \) coupled to its own nonenzymatic oxidative decarboxylation, and (b) prevents superoxide production by NADH-driven oxidases by driving reoxidation of “prooxidant” NADHc to NAD⁺c by LDH, coupled to reduction of “antioxidant” pyruvate to lactate that diffuses out of cells (Fig. 3).

We suggest that the second mechanism may be especially important, because it also will attenuate product inhibition of GA3P-DH (by increased levels of NADHc) and associated metabolic imbalances (i.e., increased triose phosphate levels that (a) fuel synthesis of DAG that activates PKC that activates some isoforms of NADPH-driven oxidases, etc. (30-OAS III p. 7, 57, and current OAS I); and (b) undergo concentration-dependent degradation to methylglyoxal, a toxic and potent glycating agent (Fig. 3). GA3P-DH activity also can be inhibited by oxidative damage caused by superoxide; the present observations support the importance of increased sorbitol oxidation as the primary source of superoxide rather than increased mitochondrial oxidation of pyruvate (4, 5, 29).

Although we conclude that our current observations (together with evidence published by other investigators) are consistent with a key role for increased sorbitol oxidation in mediating diabetic complications, it is important that we also identify credible alternative interpretations of observations and caveats to various speculations in the literature. For interested readers, these are discussed in detail in OAS II.

Therapeutic implications of these experiments include the need to minimize metabolism of glucose via the sorbitol pathway by (a) maintaining glucose levels as close to normal as feasible, and (b) developing more-potent ARIs to prevent sorbitol oxidation and associated production of superoxide and metabolic imbalances that mediate complications of diabetes (31, 32). Recent observations in animal models of diabetes suggest that the disappointing efficacy of ARIs in normalizing neural and vascular dysfunction in past clinical trials may be largely attributable to inadequate inhibition of the sorbitol pathway (31, 32). Pyruvate supplementation in some form also merits further exploration in view of its extraordinary antioxidant effects in animal models of diabetes and hypoxic injury (OAS I-C).

Conclusions

Observations in these experiments support the importance of increased “prooxidant” NADHc formed by sorbitol oxidation from equimolar formation of “antioxidant” pyruvate constrains reoxidation of NADHc to NAD⁺c by LDH, which is coupled to the reduction of pyruvate to lactate that diffuses out of cells (Fig. 3). Increasing NADHc levels drive reoxidation of NADHc to NAD⁺c by (a) NADH-driven oxidases that produce superoxide, and (b) G3P-DHc, which promotes synthesis of DAG, activation of PKC, and activation of NADPH-driven oxidases that further augment superoxide production (Fig. 3).

Attenuation of diabetes-induced vascular and neural dysfunction by ARI, SDI, and pyruvate supplementation is explained by (a) prevention of sorbitol formation by ARI, (b) inhibition of sorbitol oxidation by SDI, and (c) reoxidation of excess NADHc to NAD⁺c coupled to reduction of pyruvate to lactate by LDH rather than NADH- and NADPH-driven oxidases and G3P-DHc.

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Author Disclosure Statement

No competing financial interests exist. At the time the work in this manuscript was performed, BLM and FJO were employees of Pfizer Inc., Pfizer Global Research and Development, Groton, CT.

References

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**Abbreviations Used**

AR = aldose reductase  
ARI = aldose reductase inhibitor  
BREC = bovine retinal endothelial cell  
DAG = diacylglycerol  
DHAP = dihydroxyacetone phosphate (a triose phosphate-TP)  
G3P = glyceraldehyde phosphate (a triose phosphate-TP)  
G3P-DH = G3P-dehydrogenase  
G3P = glyceraldehyde 3-phosphate (another triose phosphate-TP)  
G3P-DHc = glyceraldehyde 3-phosphate dehydrogenase (cytosolic)  
G3P-DHm = glyceraldehyde 3-phosphate dehydrogenase (mitochondrial)  
GSH = reduced glutathione  
GSSG = oxidized glutathione  
H2O2 = hydrogen peroxide  
HVEC = human umbilical cord endothelial cell  
L/DH = lactate dehydrogenase  
L/P = lactate/pyruvate ratio  
MABP = mean arterial blood pressure  
MDA = malondialdehyde  
MG = methylglyoxal  
MNVC = motor nerve conduction velocity  
MPTP = mitochondrial permeability transition pore  
NEFA = nonesterified fatty acid  
NOS = nitric oxide synthase  
O2*− = superoxide  
OAS = Online Appendix Section  
OP = oxidative phosphorylation  
PARP = poly(ADP-ribose) polymerase  
PCK = protein kinase C  
ROS = reactive oxygen species  
SDH = sorbitol dehydrogenase  
SDI = sorbitol dehydrogenase inhibitor  
SIRT1 = silent information regulator 2 homolog 1 (an NAD-dependent deacetylase)  
SOD = superoxide dismutase  
SP = substrate phosphorylation  
TCA = tricarboxylic acid cycle (Krebs cycle, citric acid cycle)  
TP = triose phosphate  
VEGF = vascular endothelial growth factor