Modifications in Retinal Mitochondrial Respiration Precede Type 2 Diabetes and Protracted Microvascular Retinopathy

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PURPOSE. To characterize retinal mitochondrial respiration associated with type 2 diabetes (T2D) progression in a cone-rich diurnal rodent, the Nile rat (genus Arvicomus, species niloticus).

METHODS. Nile rats were fed a standard rodent diet that resulted in rising glucose levels from 6 months. Age-matched control animals were fed a high-fiber diet that prevented diabetes up to 18 months. The functional status of specific retinal mitochondrial components and mitochondrial outer membrane integrity were studied by using high-resolution respirometry. Ocular complications were documented with funduscopy, electrotretininograpy (ERG), and tryptin digestion of retinal vasculature.

RESULTS. Mitochondrial functional changes were detected during hyperinsulienemia with maintained normoglycemia (2 months), corresponding to stage 1 of human T2D. Our data showed increased contribution of mitochondrial respiration through the NADH pathway relative to maximal oxidative phosphorylation capacity, with simultaneous electron entry into NADH (Complex I and related dehydrogenases) and succinate (Complex II) pathways. These compensatory events coincided with compromised mitochondrial outer membrane integrity. The first clinical sign of retinopathy (pericycle loss) was only detected at 12 months (after 6 months of sustained hyperglycemia) alongside a common ocular complication of diabetes, cataractogenesis. Further prolongation of hyperglycemia (from 12 to 18 months) led to capillary degeneration and delayed photopic ERG oscillatory potentials.

CONCLUSIONS. Oxidative phosphorylation compensatory changes in the retina can be detected as early as 2 months, before development of hyperglycemia, and are associated with reduced mitochondrial outer membrane integrity.

Keywords: hyperinsulienemia, hyperglycemia, type 2 diabetes, animal model, oxidative phosphorylation

With a worldwide prevalence of 415 million and importantly, estimates that only one of every two adults has been diagnosed,1 the number of individuals living with diabetes may be closer to 1 billion. In adults, type 2 diabetes (T2D) accounts for most (90%-95%) diagnosed cases of diabetes.2 A well-recognized microvascular complication of diabetes is diabetic retinopathy (DR).

Clinically, DR is an inner retina pathology that begins with the nonproliferative form characterized by alterations in blood flow, death of retinal pericytes, and basement membrane thickening.3,4 Subsequently, the disease progresses to non-perfusion, microaneurysms, vascular loops, and hemorrhages evident on ophthalmic examination.5 Ultimately, DR evolves to a proliferative stage, typified by neovascularization, neurodegeneration, and retinal detachment, which cause irreversible vision loss.6 The cellular mechanisms underlying microvascular complications involve neurovascular cross talk, which is currently under intense scrutiny.7 There is accumulating evidence that the outer retina, although asymptomatic itself, contributes to these microvascular changes.8,9 Photoreceptor inner segments, in view of their extremely high metabolic activity,10 their constitutive production of high amounts of reactive oxygen species (ROS),11 and their high capacity for mitochondrial oxidative phosphorylation,12 are attracting increasing attention in the pathophysiology of DR.13 A major hurdle in identifying early pathologic events in DR has been the prevalence of animal models with early onset of diabetes.14–15 In this study, we relied on the combined advantages of the laboratory rodent Arvicomus niloticus: (1) it is a well-established model of T2D16–18 and of associated retinopathy19; (2) unlike mouse and rat models, retinopathy progresses in a cone-rich retina (30%-35%, as opposed to 1%-2% in mouse and rat),20,21 therefore more closely approximating the human macula (cone-dominated area); and (3) unlike mice and rats, Nile rats are diurnal, as humans are.22 The latter advantage is particularly pertinent in the context that development of T2D
has been associated with circadian rhythm disruption via the dysregulation of clock genes.\textsuperscript{25}

During development of T2D, Nile rats undergo an initial compensation for insulin resistance with increased insulin secretion at 2 months, followed by rising glucose levels from 6 months, increasing β-cell dysfunction and associated endoplasmic reticulum stress at 12 months, and finally, severe decompensation and progression to ketosis at 18 months.\textsuperscript{16–18} Of note, feeding of a high-fiber diet completely prevents T2D in Nile rats.\textsuperscript{18} Noda et al.\textsuperscript{19} have described the protracted accumulation of leukocytes in retinal arteries at high plasma insulin levels, which is correlated with plasma leptin (body fat) in this model, and furthermore, demonstrated microvascular biomarkers of human early DR,\textsuperscript{24} including tortuosity, acellular capillaries, and pericyte ghosts, all of which corroborate pericyte apoptosis. Hajmousa et al.\textsuperscript{25} have attributed pericyte apoptosis, in vitro, to hyperglycemia-induced accumulation of ROS. Long-term exposure of “pericyte-like” adipose-derived stromal cells to hyperglycemia is also associated with impaired glucose uptake and mitochondrial electron transport system capacity measured in the intact cells after uncoupling.\textsuperscript{26–28}

While mitochondrial dysfunction has been extensively studied in T2D, with distinct links between obesity, inactivity, and energy homeostasis, a causal relationship has yet to be established for the development of DR.\textsuperscript{29} Deficiencies in mitochondrial biogenesis and/or activity, as well as chronic hyperglycemia from 6 months and decompensation at 18 months,\textsuperscript{16–18} known to induce hyperinsulinemia by 2 months, followed by rising glucose levels from 6 months and decompensation at 18 months,\textsuperscript{16–18} or (2) control animals fed high-fiber diet (4.1% fat, 15.0% fiber, Mazuri Chinchilla Diet, 5M01; Purina Mills, LLC, St. Louis, MO, USA), normoglycemic to 18 months. Food and water were provided ad libitum. Animals were maintained on a 14:10 light–dark cycle, room temperature 21°C ± 2°C, and relative humidity ~40%. This research was done under the approval of the Institutional Animal Care and Use Committee (University of Alberta). Experiments were carried out in accordance with National Institutes of Health (Bethesda, MD, USA) guidelines regarding the care and use of animals for experimental procedures and the ARVO Statement for the Use of Animals in Ophthalmic and Visual Research.

**Fasting Blood Glucose, Glycated Hemoglobin, and Plasma Insulin**

Fasting blood glucose (FBG) values were measured from tail blood samples after overnight (16–18 hours) fasting by using an Accu-check Compact Plus glucose monitoring system (Roche, Mississauga, ON, Canada). Hyperglycemia was determined as FBG levels > 5.6 mM (100 mg/dL) as previously described.\textsuperscript{18} Nile rats were euthanized with a lethal dose (480 mg/kg) of Euthanyl (Bimeda-MTC Animal Health, Inc., Cambridge, ON, Canada). When animals reached surgical plane, cardiac puncture was performed and blood was collected in K$_2$EDTA-coated BD Microtainer Tubes with Microgar Closures (Becton, Dickinson and Company, Mississauga, ON, Canada). After centrifugation at 1000g for 20 minutes at 4°C, plasma was collected and stored at −80°C until use.

Plasma insulin levels were assessed by using an insulin ELISA kit (Ultra Sensitive Mouse Insulin ELISA kit No. 90080; Crystal Chem, Inc., Downers Grove, IL, USA). Fasting insulin levels > 2 ng/mL were indicative of compensation as previously described.\textsuperscript{18} Glycated hemoglobin (HbA1c) was measured by using a GLYCO-Tek affinity column kit (No. 5351; Helena Laboratories, Beaumont, TX, USA). Absorbance was measured at 415 nm. Percentage of HbA1c in the sample was calculated as recommended by the manufacturer. Nile rats with HbA1c levels > 6.5% were considered diabetic, as previously defined.\textsuperscript{18,31}

**Funduscropy**

Nile rats were anesthetized by using ketamine:xyazine (75:30 mg/kg). Eyes were dilated with corneal application of 1% tropicamide and 2.5% phenylephrine and kept hydrated with Tear-gel (Novartis Pharmaceuticals Canada Inc., Mississauga, ON, Canada). Images from both eyes were captured with a Micron III retina imaging system for rodents (Phoenix Research Laboratories, Inc., Pleasanton, CA, USA) equipped with a Semrock FF01-554/211 filter for white light, FF01-469/35 exciters, and BLP01-488R barrier filters for blue excitation with green emission (Semrock, Inc., Rochester, NY, USA) and StreamPix 5 software (Norpix, Montreal, QC, Canada). Fluorescein angiography images were captured after intraperitoneal injection with fluorescein (1.5 μL/g body weight of 100 mg/mL fluorescein in PBS, F6377; Sigma, Oakville, ON, Canada).

**Trypsin Digestion of Retinal Vasculature**

After overnight fixation (16–18 hours) in 4% paraformaldehyde, retinas were removed and immersed in 3% trypsin in 0.1 M Tris, pH 7.8 (preheated to 37°C) for 130 minutes. Following digestion, the vitreous was carefully removed and the retina was subjected to gentle trituration with water to separate disintegrated neuronal tissue from the vasculature. Isolated vascular networks were mounted, treated with periodic acid

**Funding**

This study was performed on male Nile rats (Arvicanthis nilotica) aged 2 to 18 months; females were excluded on the basis of their reduced susceptibility to develop T2D\textsuperscript{17} and to control for potential sexual dimorphisms with respect to metabolic pathways. After weaning at 21 days, animals were divided into two diet groups: (1) prediabetic and diabetic animals fed standard rodent diet (9.6% fat, 3.2% fiber, ProLab RMH 2000, 5P06; LabDiet, Nutrition Intl., Richmond, IN, USA), known to induce hyperinsulinemia by 2 months, followed by hyperglycemia from 6 months and decompensation at 18 months,\textsuperscript{18} or (2) control animals fed high-fiber diet (4.1% fat, 15.0% fiber, Mazuri Chinchilla Diet, 5M01; Purina Mills, LLC, St.
solution, and stained with Schiff’s reagent and hematoxylin (Periodic Acid–Schiff kit 395B-1KT; Sigma-Aldrich Corp., St. Louis, MO, USA).

Pericytes were identified as round, dark extracapillary protrusions when compared to endothelial cells, which were oval and lighter in color.² Images were captured on a Reichert Polyvar 2 microscope with Infinity Analyze imaging software (Lumenera Corporation, Ottawa, ON, Canada) using an ×40 oil objective. Counts of pericytes, endothelial cells, and degenerated capillaries were performed and normalized to capillary area by using ImageJ 1.48v software (http://imagej.nih.gov/ij; provided in the public domain by the National Institutes of Health, Bethesda, MD, USA). Pericytes were categorized into three subtypes by location; pericytes located on straight capillaries, forked capillaries, and bridged between capillaries were classified as longitudinal, forked, and bridging pericytes, respectively. Six diabetic and six control animals were assessed at each time point (6, 12, 18 months), and five randomly selected images per animal were analyzed. All variables met the criteria of normality and homogeneity of variance for ANOVA as tested with Kolmogorov-Smirnov (Lilliefors’ correction) and Spearman tests, respectively. Therefore, differences between diet groups could be analyzed by using 2-way ANOVA with Bonferroni post hoc comparisons. Significance was set to \( P < 0.05 \).

**Immunohistochemistry**

After enucleation, corneas were punctured, and eyes were fixed in 4% paraformaldehyde for 30 minutes at 4°C. Corneas and lenses were carefully removed, and posterior eyecups were fixed in 4% paraformaldehyde for 30 minutes at 4°C, cryoprotected in sucrose (10% and 20% for 1 hour each, 30% overnight), embedded in O.C.T. (Tissue-Tek; Sakura Finetek, Torrance, CA, USA), frozen in liquid nitrogen, and stored at –80°C.

For cross sections, 20-μm cryosections were cut along the nasotemporal axis. After blocking in 10% normal serum and 0.3% Triton X-100 in PBS for 2 hours, sections were reacted overnight with goat anti–choline acetyltransferase (ChAT, 1:200; AB144P; Chemicon, Etohoicke, ON, Canada). All incubations were performed at room temperature. The following day sections were washed (3 × 5 minutes) in PBS and reacted for 2 hours with donkey anti-goat Alexa594 antibody (1:500; Molecular Probes, Inc., Eugene, OR, USA). After washing in PBS, slides were coated with ProLong Gold antifade (1:50; Molecular Probes, Inc., Eugene, OR, USA), coverslipped. Images were captured as above.

**ERG Recordings**

ERGs were recorded at 18 months (\( n = 6 \) diabetic animals, \( n = 5 \) control animals). Nile rats were dark-adapted 1 hour before recording, and all subsequent preparations were made under dim red light. Animals were anesthetized with inhaled isofluorane (Somnosuite, Small Animal Anesthesia System; Kent Scientific, Torrington, CT, USA) and placed over a homeothermic blanket to maintain their body temperature at 38°C. Pupils were dilated with one drop of 1% tropicamide, and corneas were kept hydrated with methylcellulose. ERG recordings were performed as previously described.²⁰ Data were analyzed in a single eye per animal, selected on the basis of largest maximal a-wave amplitude in the scotopic intensity series. For oscillatory potential (OP) analysis, to ensure consistency between traces, amplitudes and implicit times were calculated by manually placing cursors at the base and apex of the first four OPs; cursors were sequentially labeled from 1 to 8 and values (amplitude or implicit time) were plotted for each cursor against stimulus strength.

**Scotopic ERG.** Intensity response series in dark-adapted animals were generated by presenting flashes (10-μs duration, 0.500°K, xenon bulb) ranging between –3.70 to 2.86 log cd s/m² in luminance over 16 consecutive incremental steps. Intensity levels were measured with a photometer at the cornea level. Interstimulus intervals ranging from 10 seconds to 2 minutes were used for lowest to highest intensity flashes, respectively, providing sufficient time for rods to recover from photobleaching. Responses were recorded three to six times and averaged to optimize signal-to-noise ratio. For all recordings, a-wave amplitude corresponded to the difference between baseline at time 0 second (flash delivery) and negative a-wave trough, while b-wave amplitude corresponded to the difference between negative a-wave trough and positive b-wave peak (excluding OPs). Criterion amplitudes were set to 10 μV.

**Mesopic ERG.** After completion of scotopic ERG recordings, animals were exposed to a background illumination of 0.01 cd/m² for 10 minutes before applying the same tests as described above.

**Photopic ERG.** Finally, after completion of mesopic ERG recordings, animals were light-adapted at 30 cd/m² background for 10 minutes. Intensity response series were generated by presenting white flashes ranging between –1.63 to 2.86 log cd s/m² in luminance over 11 consecutive incremental steps. Responses were recorded six times and averaged. Interstimulus intervals of 3 seconds were used. Flicker response series were then recorded to flashes of 1.36 log cd s/m² presented at the following 13 incremental frequencies: 3, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, and 60 Hz. The first 3 seconds were not harvested, avoiding averaging responses to single flashes. Amplitudes were measured from adjacent trough to peak and latencies from flash onset to peak.

Statistical comparison for each marker (1 to 8) was assessed with nonparametric Kruskal-Wallis 1-way analysis of variance. Significance was set to \( P < 0.05 \).
Retinal Mitochondrial Changes Precede T2D

High-Resolution Respirometry

High-resolution respirometry (Oxygraph 2k; OROBOROS Instruments, Innsbruck, Austria) was performed by using freshly isolated retinas from individual Nile rats from 2 to 18 months (n = 5–12 per group, replicate measurements from a single animal were pooled). The oxygraph was calibrated at 37°C as per manufacturer’s instructions with each chamber filled with 2 mL Mitochondrial Respiration Medium 05 (MiR05; 0.5 mM EGTA, 3 mM MgCl2·6H2O, 60 mM K-lactobionate, 20 mM taurine, 10 mM KH2PO4, 20 mM HEPES, 110 mM sucrose, and 1 g/L BSA essentially fatty-acid free, pH 7.1). Datlab software (OROBOROS Instruments) was used for data acquisition and analysis.

Wet weight was measured and retinas were immediately transferred into 1 mL ice-cold MiR05. Retinas were homogenized on ice with a Potter-Elvehjem attached to an overhead stirrer (Wheaton Instruments, Millville, NJ, USA). After five passes at intensity level two, 200 μL homogenate was immediately placed in each oxygraph chamber containing 1.8 mL MiR05. The remaining homogenate was frozen at −80°C for determination of total protein (Pierce BCA protein assay kit, Cat. No. PI-23227; Thermo Fisher Scientific, Rockford, IL, USA) and citrate synthase (CS) activity (according to Kuznetsov et al.34). CS (EC 4.1.3.7) was measured at 37°C with a UV/Vis spectrophotometer (Ultrospec 2100 Pro; Biochrom, Cambridge, MA, USA) equipped with a thermostated cell holder and a circulating water bath. After thawing the sample on ice, an additional cycle of homogenization with a conical glass homogenizer for 30 seconds on ice was performed to ensure complete homogeneity of the sample. The absorbance was measured at 412 nm following the reduction of 0.1 mM 5,5′-dithiobis-2-nitrobenzoic acid (c: 13.6 mL cm−1 μmol−1) in the presence of 0.31 mM acetyl-CoA, 0.5 mM oxaloacetic acid, 0.25% Triton X-100, 100 mM Tris-HCl buffer (pH 8.0), and 25 μL retinal homogenate in a spectrophotometer cuvette containing a total volume of 1 mL.

The protocol used for evaluating mitochondrial function is presented in Figure 1. Two different states are measured: LEAK respiration represents the nonphosphorylated state in the absence of ADP; and OXPHOS represents oxygen consumption coupled to phosphorylation of ADP to ATP in the presence of saturating ADP. The following substrates and inhibitors were added (final chamber concentration): pyruvate (5 mM), malate (5 mM), ADP (2.5 mM), cytochrome c (10 μM), succinate (10 mM), rotenone (1 μM), antimycin A (5 μM), ascorbate (2 mM), tetramethylphenylenediamine (TMPD; 0.5 mM), and sodium azide (100 mM). Pyruvate and malate were added before the retina homogenate and the chamber was closed immediately after homogenate addition. An uncoupler titration (dinitrophosphonol; 5–10 M steps) was performed after succinate in order to determine the maximal capacity of the electron transport system (ETS). Furthermore, addition of digitonin after ADP did not show any increase in respiration, confirming that none of the plasma membranes were left intact after homogenization (results not shown). Mitochondrial respiration was corrected for oxygen flux due to instrumental background, and for residual oxygen consumption (ROX) after inhibition of Complexes I and III, with rotenone and antimycin A, respectively. This protocol provides an indirect assessment of Complex III activity. Although electrons are initially fed into Complexes I, II, or I and II, defects in steps downstream of the Q-cycle, requiring electron transfer by Complexes III and IV, impact the capacity of Complexes I and II. For Complex IV respiration, the chemical background measured in presence of sodium azide was subtracted.

For the detection of qualitative changes in the OXPHOS system, respiration was expressed as flux control ratio (FCR), normalized for maximal OXPHOS capacity in presence of substrates simultaneously feeding electrons into both NADH and succinate pathways (Complex I and II, respectively).

Statistical analyses were performed with SigmaStat 4 (Aspire Software International, Ashburn, VA, USA). Criteria of normality and homogeneity of variance for ANOVA were tested for each variable with Kolmogorov-Smirnov (Lilliefors’ correction) and Spearman tests, respectively. For variables meeting the above criteria (FCR, for NADH and succinate pathways; and CS

![Figure 1](http://arvojournals.org/)
activity), differences between age-matched diet groups were tested with 2-way ANOVA, followed by pairwise Tukey comparison. For variables not meeting the criteria (coupling control ratio; FCR for Complex IV; and cytochrome c control factor), age-matched differences were analyzed with Mann-Whitney U test. Unless otherwise stated, values reported in this study represent mean ± standard error of the mean (SEM). Significance was set to $P < 0.05$.

## Results

### Changes in Retinal Mitochondrial Respiration Precede T2D

FCR (when normalized for maximal OXPHOS capacity) allows detection of qualitative changes, and this approach revealed a greater NADH pathway contribution to maximal OXPHOS capacity at 2 months (Fig. 2A). At hyperglycemia onset (6 months), no differences were observed; however, statistical comparisons segregating hyperglycemic and hyperinsulinemic status confirmed that this increase was attributed to increases in FBG (18 months).

Regardless of FBG levels, FCR for Complex IV did not vary at any of the time points examined (Fig. 2C). The phosphorylation system did not exert a limitation on electron transport in Nile rat retinas; ratios of maximal OXPHOS/ETS were $0.97 ± 0.03$ and $0.96 ± 0.01$ in diabetic versus control animals, respectively. Mitochondrial respiratory capacity expressed as a function of tissue mass did not vary between diets (Table 1) and mitochondrial content, as reflected by CS activity, was comparable between age-matched diet groups (Fig. 2D).

### Figure 2

Mitochondrial oxidative phosphorylation (OXPHOS) capacity and mitochondrial content in retina homogenates. OXPHOS capacity is expressed as FCR normalized for physiological OXPHOS capacity with substrates feeding electrons into NADH and succinate pathways simultaneously. FCRs are provided for (A) NADH pathway (pyruvate+malate+ADP), (B) succinate pathway (succinate+rotenone+ADP), and (C) Complex IV single step (ascorbate+TMPD+ADP). Finally, (D) CS activity (normalized against total protein content in homogenates) is used as an indicator of mitochondrial content. Asterisks indicate statistical significance.

### Table 1. Flux Per Mass Values

<table>
<thead>
<tr>
<th></th>
<th>2 mo</th>
<th>6 mo</th>
<th>18 mo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prediabetic</td>
<td>Control</td>
<td>Prediabetic</td>
</tr>
<tr>
<td>LEAK</td>
<td>$2.2 ± 0.7$</td>
<td>$2.8 ± 0.5$</td>
<td>$4.1 ± 0.6^*$</td>
</tr>
<tr>
<td>NADH pathway</td>
<td>$29 ± 5$</td>
<td>$32 ± 3$</td>
<td>$36 ± 3$</td>
</tr>
<tr>
<td>Succinate pathway</td>
<td>$22 ± 4$</td>
<td>$27 ± 3$</td>
<td>$26 ± 2$</td>
</tr>
<tr>
<td>NADH-succinate pathway</td>
<td>$46 ± 7$</td>
<td>$54 ± 5$</td>
<td>$55 ± 4$</td>
</tr>
<tr>
<td>Maximal COX capacity</td>
<td>$96 ± 17$</td>
<td>$111 ± 11$</td>
<td>$102 ± 8$</td>
</tr>
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COX, cytochrome c oxidase.

$^* P < 0.05$. 

IOVS | August 2017 | Vol. 58 | No. 10 | 3830

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Body weight, g 66
BMI, kg/m² 5.4

dimensions, regardless of insulin or FBG levels. The opposite effect was observed and was still attributed to hyperinsulinemia alone. No differences were detected at 18 months. The decline in this cell population progressed to 26% ± 4% at 18 months. In contrast, forked pericytes and bridged pericyte cell populations were unaffected. The total pericyte-to-endothelial cell (TP:EC) ratio at 6 months is approximately 1:1 (Fig. 4C) and declines thereafter owing to pericyte loss.

The number of degenerated capillaries was unchanged at 12 months; however, at 18 months there was a >3-fold increase in the number of degenerated capillaries (156 ± 36 in diabetic versus 47 ± 36 in control animals; Fig. 4D).

A second clinical sign of DR demonstrated in diabetic Nile rats is delayed ERG OP kinetics (Fig. 5). OP implicit times were delayed, exclusively under photopic adaptation. Cursor positions on OP base and apex were shifted to longer times, with the exception of two cursors (5 and 6) corresponding to the largest amplitudes in the OP bursts. OP amplitudes were unaffected.

No differences were observed in a- and b-wave amplitudes or implicit times between diabetic and control animals at 18 months, regardless of the adaptation background studied (scotopic, mesopic, or photopic). Finally, b-wave amplitudes were quantified at time 0 versus 10 minutes after switching background adaptation from scotopic to mesopic, as well as from mesopic to photopic, and again no changes were observed.

**Early Diabetic Retinopathy**

The first clinical sign of retinopathy (pericyte loss) is observed in T2D Nile rats from 12 months (Fig. 4A). These animals exhibited a preferential loss of longitudinal pericytes from 12 months onward (12 months: 1481 ± 111 vs. 1746 ± 183, 18 months: 1165 ± 131 vs. 1565 ± 84, in diabetic versus control animals, respectively; Fig. 4B). This corresponds to a 15% ± 3% loss in the total number of longitudinal pericytes at 12 months. The decline in this cell population progressed to 26% ± 4% at 18 months. In contrast, forked pericytes and bridged pericyte cell populations were unaffected. The total pericyte-to-endothelial cell (TP:EC) ratio at 6 months is

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**Table 2. Metabolic Phenotype**

<table>
<thead>
<tr>
<th></th>
<th>2 mo</th>
<th>6 mo</th>
<th>12 mo</th>
<th>18 mo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control (n = 11)</td>
<td>Prediabetic (n = 7)</td>
<td>Control (n = 10)</td>
<td>Prediabetic (n = 12)</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>5.4 ± 0.2</td>
<td>5.8 ± 0.2</td>
<td>6.9 ± 0.2</td>
<td>7.3 ± 0.3</td>
</tr>
<tr>
<td>FOG, mM</td>
<td>2.9 ± 0.3</td>
<td>2.7 ± 1.4</td>
<td>5.2 ± 1.6</td>
<td>5.4 ± 1.9*</td>
</tr>
<tr>
<td>HbA1c†</td>
<td>N/A</td>
<td>N/A</td>
<td>4.5 ± 0.1</td>
<td>4.1 ± 0.1</td>
</tr>
<tr>
<td>Insulin‡</td>
<td>0.9 ± 0.3</td>
<td>4.3 ± 1.5*</td>
<td>2.9 ± 1.5</td>
<td>5.6 ± 0.9*</td>
</tr>
</tbody>
</table>

Data represented as mean ± SEM, Mann-Whitney U test. N/A, not applicable.

* P < 0.05.
† A subset of animals was used, n = 4, 6 (6 months); 4, 8 (12 months), and 4, 10 (18 months); control and diabetic, respectively.
‡ A subset of animals was used, n = 4, 5 (2 months); n = 5, 9 (6 months); 13, 20 (12 months), and 6, 6 (18 months); control and prediabetic/diabetic, respectively.
FIGURE 4. Retinal digest preparations of diabetic and control Nile rat retinas. Representative images (A) of isolated retinal vasculature, 12 (lower row) and 18 months (upper row). Quantification of pericyte numbers (B), total TP:EC ratio (C), and number of degenerated capillaries (D). Scale bar: 50 μm in upper panel; 20 μm, enlarged images in lower panel. Black arrows: pericytes; arrowheads: endothelial cells; white arrows: tortuous capillaries; white asterisk: early capillary degeneration; black asterisk: pericyte ghost. Asterisks on graphs indicate statistical significance. LP, longitudinal pericyte; FP, forked pericyte; BP, bridged pericyte.
No evidence of either retinal edema or thinning was observed in diabetic animals from 6 to 18 months at any eccentricity (Fig. 6). ONL and INL measures taken from the central and peripheral retina did not differ between groups (Table 3). Cholinergic amacrine (ChAT-immunoreactive) total cell numbers, both in the INL as well as those displaced in the GCL, were unchanged (Table 3).

**Figure 5.** Oscillatory potential timing at 18 months, expressed as a function of stimulus (flash) strength. The upper panel shows traces from a diabetic (red) and a control (blue) in response to the highest strength stimulus (2.9 log cd s/m²). Timing versus strength graphs are provided separately for each of the eight markers (with the numbers indicated in upper boxes). The x-axis for the first marker is shown as an inset with strength values for each of the five stimuli presented at incremental strengths (strengths values could not be presented for all eight markers owing to space limitation). Asterisks indicate statistical significance.

**Figure 6.** Cross sections of central retina, immunostained for cholinergic amacrine cells (ChAT) in red with DAPI-stained nuclei in blue. Scale bar: 50 μm. ONL, outer nuclear layer.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Diabetic</th>
</tr>
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<tbody>
<tr>
<td><strong>6m</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Center</td>
<td>OLN thickness, μm</td>
<td>37.6 ± 3.8, n = 4</td>
</tr>
<tr>
<td></td>
<td>INL thickness, μm</td>
<td>24 ± 4, n = 4</td>
</tr>
<tr>
<td></td>
<td>ChAT+ cells</td>
<td>21.5 ± 38, n = 4</td>
</tr>
<tr>
<td>Periphery</td>
<td>OLN thickness, μm</td>
<td>25.0 ± 6.8, n = 4</td>
</tr>
<tr>
<td></td>
<td>INL thickness, μm</td>
<td>16 ± 4, n = 4</td>
</tr>
<tr>
<td></td>
<td>ChAT+ cells</td>
<td>18 ± 4, n = 4</td>
</tr>
</tbody>
</table>

**Table 3.** Central and Peripheral Retina Outer and Inner Nuclear Layer Thickness Measures and ChAT-Immunoreactive Amacrine Cell Counts

Values indicate average ± standard deviation.
Likewise, no differences were detected in retinal flatmounts stained with specific oxidative/nitrative stress markers (Fig. 7). Very low levels of extravascular lipid oxidative modifications were demonstrated with 4-hydroxynonenal–specific antibodies (12 months only); the retinal vasculature was devoid of any detectable immunoreactivity. The nitrative stress marker nitrotyrosine was detected at low levels in blood vessels of both groups at 2 months. This vascular staining was exacerbated at 12 months in both central and peripheral retina but did not differ between groups.

**Cataracts of Increasing Severity**

Fluorescein angiography (FA) was performed to assess inner retinal vasculature. Imaging of diabetic animals was limited owing to impedance by cataract formation. Clearly evident in the eyes of diabetic animals (Fig. 8A; Fig. 8B, lens only), cataracts exhibited a range of severity (Figs. 8C–I), illustrating the barrier to examining retinal blood vessels in vivo (Figs. 8D–J). Incidence increased from 13% at 12 months, to 50% at 18 months. Cataracts were never detected in controls at any of the
The present study established that high-resolution respirometry allows the normalization of results as FCR values, which is crucial to accurately detect modest changes, and also allows many titration protocols to be performed in a single experiment. We showed evidence of well-coupled mitochondia in tissue homogenates, which allowed us to identify changes in oxidative phosphorylation and breaches in outer membrane integrity during insulin resistance (2 months), before the manifestation of hyperglycemia (6 months), in *Arvicanthis niloticus* (Nile rat). In addition, we provided evidence of delayed OPs (specific to the cone pathway) and increased incidence of cataractogenesis after prolonged hyperglycemia in retinas of this cone-rich diurnal rodent model, which recapitulates the five stages of human T2D progression 18 (Fig. 9). A key target in the development of DR is the synergistic interaction between neurons (amacrine and retinal ganglion cells), glial cells ( Müller cells and astrocytes), and blood vessels (endothelial cells and pericytes) involved in the autoregulation of vascular flow and metabolic activity. Dysregulation of this "neurovascular unit" has been implicated in diabetes, in part due to oxidative stress originating from the outer retina, and well before onset of clinically detectable DR. 38 According to Kooragayala et al., 36 photoreceptors have limited mitochondrial reserve capacity for ATP production from their electron transport system. Therefore, photoreceptors are particularly vulnerable to changes in homeostasis. Fort et al. 39 have observed that during the initial stages of type 1 diabetes, the retina compensates by lowering its constitutively high metabolic activity. Reductions in retinal electrical activity, 40 biosynthetic activity, 41 increases in autophagic flux in the outer plexiform layer, 42 and even apoptosis of marginal populations of neural and vascular cells 43 are all contributors. However, these mechanisms eventually fail, leading to the first clinically detectable signs of DR, specific to inner retina microvasculature: capillary basement membrane thickening and pericyte loss. Our results support that changes in mitochondrial function and integrity occur early and likely contribute to transient compensation in an attempt to maintain energy homeostasis.

Increased respiration at 2 months, as a result of exogenous cytochrome c entering the intermembrane space, indicates changes in mitochondrial membrane properties (such as composition and/or organization). Surprisingly, this transient increase in outer membrane permeability was associated with an increase in NADH pathway (Complex I and related dehydrogenases) contribution to maximal OXPHOS capacity, whereas it is generally associated with defects in mitochondrial respiratory complexes. 20, 44 At the onset of hyperglycemia (6 months), we observed increases in LEAK and decreases in cytochrome c access to mitochondrial respiratory complexes without paradoxical increases in NADH pathway activity. One possible explanation would be that early changes in mitochondrial function and integrity might entice transient compensation for the maintenance of energy homeostasis. Changes in mitochondrial membrane composition take several months to develop 15–17 and may explain the improved membrane integrity observed in 6-month diabetic animals, with no further changes at 18 months.

Mitochondria were well coupled in prediabetic animals, without decreased respiratory capacity, emphasizing the specificity of this change in membrane sensitivity in retinal mitochondria. These early changes did not correlate with fasting plasma insulin levels. Increased mitochondrial membrane permeability was documented in isolated mitochondria from retinas of Zucker diabetic fatty (ZDF) rats during hyperinsulinemia (before hyperglycemia), but alongside a decrease (rather than an increase) in Complex III activity. 46 Decreases in Complex III activity (without changes in Complex
I) were also reported in STZ mice. Respiration in the presence of substrates feeding electrons into Complex I (NADH pathway), II (succinate pathway), or I and II also involves Complexes III and IV, and so a decrease in Complex III would cause a decrease in maximal OXPHOS capacity, which was not observed in our study. The above differences between animal models epitomizes the debate about whether changes in mitochondrial capacity are causal or consequential to insulin resistance. Such discrepancies between models and studies might in fact provide pertinent insight into the intricate relationship between mitochondria and insulin action, and subsequently effective preventative targets.

Enhanced NADH pathway contribution to maximal OXPHOS capacity in retinas is reminiscent of data on cardiac mitochondria showing an increase in mitochondrial oxidation of palmityl carnitine, glutamate, and succinate in insulin-resistant mice and an increase in oxidation of palmitylCoA and octanoylcarnitine, as well as in Complex I activity, at early stages of diabetes in fructose-fed rats. Complexes I, II, and III have the highest capacity for superoxide production; therefore, decreases in their activity (albeit associated with maintenance of respiratory complex flux) would lead to excessive free radical production. In fact, most mitochondria produce superoxide high rates after addition of the Complex I inhibitor rotenone and furthermore, several pathologies involving defects in Complex I are associated with elevated production of superoxide. Therefore, the initial increase in relative contribution of electron flow through Complex I observed during prediabetes (2 months) could represent a compensatory mechanism serving to reduce the production of ROS. In contrast, at 18 months of age, after 1 year of hyperglycemia, reduced NADH pathway capacity suggests pathology as a result of oxidative stress. However, the age-related changes previously reported in other tissues (reduced CS activity and decrease in the activity of mitochondrial complexes) were not observed in Nile rat retinas at 18 months, regardless of their diabetic status. We failed to detect upregulation of targeted oxidative/nitrative stress markers in retinal vasculature of either prediabetic (2 months) or diabetic animals (up to 12 months) relative to controls. Our high-resolution respirometry results support these findings. We cannot rule out that examination of additional markers (and/or later time points) may have demonstrated oxidative stress in the vasculature. Assuming that oxidative stress would have occurred in the retina, then the expected outcome would have been a decrease (and not an increase as observed here) in NADH pathway activity. In STZ rats, at early stages (1–3 weeks post induction) retinal mitochondria have been shown to undergo adaptive changes associated with maintained energetic requirements and prevention of oxidative stress. However, at later stages (after 8–12 weeks post induction), upregulation of oxidative stress markers was reported. STZ rats are characterized by hyperglycemia due to insufficient insulin secretion (T1D model), whereas Nile rats undergo hyperinsulinemia without initial hyperglycemia (T2D model). While STZ studies show adaptive mitochondrial changes at early stages of hyperglycemia (without elevated insulin levels), the mitochondrial changes reported here in Nile rats predate manifestations of DR.

Well-established clinical indicators of early DR, oscillatory potentials, were affected at 18 months in diabetic Nile rats; implicit times were prolonged while amplitudes were preserved. Of interest, Morlet wavelet transform failed to detect implicit time delays in this study. Morlet is limited to time domain analysis of the largest amplitude oscillatory components and these OPs were not delayed. Similar findings were reported in human DR and in animal models where delays in OPs also preceded changes in their amplitudes. Using a mouse model of T2D (high-fat diet), Rajagopal et al. have reported delayed OP timing as the first functional retinal phenotype, coinciding with the peak of insulin resistance (6 months). Once hyperglycemia develops (by 12 months), amplitudes become reduced, alongside microvascular changes. However, a- and b-waves (amplitudes and latencies) are unaltered as was the case in diabetic Nile rats. Our findings support previous evidence that changes in OPs precede those in b-waves. One major difference with mouse and rat models is that alterations in OPs in these nocturnal rodents were detected under scotopic adaptation. Interestingly, we only observed alterations in OPs when recorded under photopic adaptation, which could be related to the higher proportion of cones in Nile rats (35%) compared to rats and mice (1%–3%). Our observation of preserved OP amplitudes is in agreement with the preserved total number of cholinergic amacrine cells, which likely contribute to OP generation together with other inner retina neurons. Studies in induced and spontaneous diabetes type 1 diabetic rodent models (STZ rats and Ins2Akita mice, respectively) report the loss of both cholinergic and dopaminergic amacrine cells; however, ERG recordings have not been performed. Other studies on STZ models have reported defects in a-waves, but whether these reflect impaired photoreceptor response to light due to hyperglycemia or a direct toxic effect of STZ remains unclear. Our ERG results therefore further support that DR at its earliest stage of manifestation in Nile rats. The lack of changes in a-wave amplitudes and implicit times in our study implies that changes in photoreceptor metabolism likely
contribute to DR initiation,8 without early concomitant changes in photoreceptor response to light flashes.

The 50% incidence of cataracts in diabetic Nile rats, at this time point, is similar to the high incidence reported in T2D patients.7,5

Pericyte loss was preferential to the subtype lining straight capillaries as previously reported in the Ins2Akita mouse model of type 1 diabetes,76 as opposed to pericytes lining forked capillaries or bridging adjacent capillaries. While the underlying mechanisms elude us, one avenue warranting scrutiny is the potential for hyperglycemia to alter the types of integrins expressed on specific pericytes, and hence modulate their response to pro-apoptotic ligands such as angiopoietin-2.77 Recent in vitro studies by Zhang et al.78 have demonstrated increased levels of autoantibodies against pericyte cell surface antigens in the serum of DR patients. Of interest in terms of the Nile rat’s pertinence as a DR model, the ratio of total pericytes over endothelial cells (at 6 months, before any cell loss) was close to 1:1, as observed in humans,79 which contrasts with the 1:3 ratio reported in rats.80 Degenerated capillaries were only observed at the latest time point (18 months), consistent with DR progression.3–6

Control animals did not become hyperglycemic up to 18 months, but they did demonstrate hyperinsulinemia from 6 months onward, which alone is not associated with DR. Harris and colleagues81 have estimated that T2D likely develops over 12 years (and retinopathy over 7 years) before a clinical diagnosis of T2D. Our results from high-resolution respirometry support that early mitochondrial function changes might be part of these insidious changes that precede clinical detection of T2D and therefore, DR.

One major limitation to characterizing retinal mitochondrial change during the development of diabetes (and other insults) is the limited amount of tissue. Examples of specific changes that could not be directly assessed in our broad-scope study include (1) complementary assays to measure oxidative stress directly in isolated mitochondria; (2) specific content analysis of mitochondrial external membranes (such as phospholipid types and cardiolipin); and (3) direct measures of Complex III activity, using spectrophotometric assay for this protein. Although indirect, our overall data did not support a potential defect in Complex III.

CONCLUSIONS

There is overwhelming experimental evidence linking hyperglycemia with pericyte loss and development of microvascular complications in early DR.43 Application of high-resolution respirometry, with a multiple substrate-inhibitor-titration protocol, allowed detection of early-onset retinal changes in the Nile rat T2D model. Hyperinsulinemia is associated with increased contribution of NADH pathway to oxidative phosphorylation and with compromised outer membrane integrity. The diet-based difference between groups is of particular relevance, as it offers the potential to mimic clinical interventions aimed at reversing hyperglycemia itself, such as with diet or metformin. A key challenge is to understand why in some patients, even with hyperglycemia reversal, progression to retinopathy continues. This model would be a valuable tool to examine whether hyperinsulinemia alone might “program” retinopathy onset.

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References


