Potential Role of Mitochondrial Dysfunction in Diabetic Hypertriglyceridemia

Parihar Mordhwaj1, Dmitry Litvinov1, Sainath Babu2, Chandrakala Aluganti Narasimhulu1 and Sampath Parthasarathy1,*

1Burnett School of Biomedical Sciences, University of Central Florida, USA
2Ohio State University Medical Center, USA

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*Corresponding author: Sampath Parthasarathy, PhD, MBA, Burnett School of Biomedical Sciences, University of Central Florida, 6900 Lake Nona Blvd, Orlando, FL 32827, USA, Tel: +1-(407)-266-7121; Fax: +1-407-266-7002; Email: sparth@ucf.edu

Abstract

Type 2 diabetes patients have increased oxidative stress and hypertriglyceridemia. We tested the hypothesis that these two are related and the latter could be the result of peroxide-mediated mitochondrial dysfunction and increased acetate production. Treatment of isolated liver mitochondria or primary hepatocytes with oxidized linoleic acid (LOOH) or hydrogen peroxide (H2O2) resulted in a drastic decrease in activities of pyruvate dehydrogenase (PDHC), aconitase and α-ketoglutarate dehydrogenase (KDHC). In contrast, the incorporation of 14C-acetate into lipids was not affected by peroxides suggesting that fatty acid synthesis was not affected. The livers of Db/db diabetic mice showed reduced enzyme activities as compared to control non-diabetic mice. In vitro reaction of pyruvate in the presence of LOOH or H2O2 showed that pyruvate was non-enzymatically converted to acetate together with the release of carbon dioxide (CO2).

These results show that diabetic mice may convert more pyruvate non-enzymatically into acetate in the cytoplasm in the presence of peroxides. In addition, mitochondria in diabetic state may have poor capacity to utilize acetate by TCA cycle to generate energy. Combined with the findings that peroxides did not affect acetate incorporation of acetate into fatty acids, one could expect a net increase in fatty acid triacylglycerol (TG) production.

Abbreviations: EDTA: Ethylene Diamine Tetraacetic Acid; EGTA: Ethylene Glycol-Bis (β-aminoethyl Ether)-N,N,N',N'-Tetraacetic Acid; FCCP: Carbonyl Cyanide-P-Trimfluoromethoxy Phenylhydrazone; HBSS: Hanks Balanced Salt Solution; HNE: Hydroxy Non Enal; KDHC: Α-Ketoglutarate Dehydrogenase; LMB: Leukomethylene Blue Reaction; LOH: Lipid Hydroxide; LOOH: Lipid Peroxide; MSH: Mannitol Sucrose Hepes; NADH: Nicotinamide Adenine Dinucleotide Hydrogenase; NMR: Nuclear Magnetic Resonance; PBS: Phosphate Buffer Saline; PDHC: Pyruvate Dehydrogenase; RNS: Reactive Nitrogen Species; ROS: Reactive Oxygen Species; TBARS: Thiobarbituric Acid Reactive Substances; TCA: Tricarboxylic Acid Cycle

Introduction

Increased oxidative stress observed in both clinical and experimental diabetes mellitus has been implicated in the etiology of chronic diabetic complications [1-4]. Hyperglycemia leads to an increase in lipid peroxidation in diabetic patients and animals reflecting a rise in reactive oxygen species production [5-8]. Association of diabetic pathology to mitochondrial dysfunction and oxidative stress have been well documented [9-12]. Various studies point to generalized mitochondrial dysfunction in type 2 diabetes patients [13]. For example, mitochondria of type 2 diabetes patients have been shown to possess reduced electron transport chain capacities and reduced citrate synthase activity [14]. Type 2 diabetes patient’s show reduced fatty acid oxidative capacities [15]. Free fatty acid levels are increased together with decreases in fat oxidative capacity in obese, and diabetic patients, and in time this can result in accumulation of fatty acids and acyl glycerols in tissues [13,16,17]. Under both normal and pathological conditions, mitochondria are considered as the major endogenous source of ROS [18-20]. During normal metabolism 1-2% of the electrons that flow into the respiratory chain catalyze the incomplete reduction of O2 generating superoxide anion and hydrogen peroxide. However, under certain pathophysiological conditions the generation of these oxidants dramatically increases, leading to an imbalance between the pro-oxidant and the antioxidant systems. In addition, the high content of polyunsaturated fatty acids in mitochondrial membranes enhances mitochondrial susceptibility to lipid peroxidation, leading to alterations in major enzymes involved in energy production. Liver is heavily dependent on mitochondrial oxidative catabolism for the majority of their ATP requirements.
Elevated levels of ROS in liver cells are particularly dangerous because they mediate mitochondrial damage, which in turn can generate further oxidative stress in the cells [21]. Fatty acids are particularly sensitive to ROS/RNS oxidation, resulting in the formation of lipid peroxides, which are cytotoxic and lead to free-radical damage to other lipids, proteins and DNA [13].

Lipid abnormalities are commonly associated with diabetes, especially high risk for hyperlipidemia, most commonly in the form of elevated triacylglycerol levels and decreased high-density lipoprotein (HDL) levels. The most important pathogenic mechanisms, such as increased oxidative stress and increase free fatty acids and triacylglycerols followed by inactivation of mitochondrial enzymes have been identified in experimental studies [4,22-27]. Although multiple oxidant moieties may participate, there is in vitro and ex vivo evidence to support a role for superoxide anion and H$_2$O$_2$ in the pathogenesis of vascular dysfunction, lipid peroxidation, and formation of glyco-oxidation products in diabetes [23,26-30]. Various markers of oxidative stress such as increases in oxidized lipoproteins, red cell membrane lipid peroxidation, advanced glycation end products have been documented in blood and tissues of human and experimental diabetic subjects [31-34].

Recent studies shown defects in TCA cycle enzymes are associated with accumulation of very long chain fatty acids and may be accompanied by alterations in the intracellular pool of fatty acid and fatty acyl CoAs, which are known to alter mitochondrial function [35]. Most importantly, mitochondrial alterations in TCA cycle enzymes directly cause defects in electron transport chain (ETC). The pyruvate dehydrogenase complex (PDHC) is a mitochondrial matrix enzyme located exclusively in the mitochondrial matrix that catalyzes the oxidative decarboxylation of pyruvate and represents the sole bridge between anaerobic and aerobic cerebral energy metabolism. α-Ketoglutarate dehydrogenase complex (KGDH) located in the inner mitochondrial matrices is crucial in the cellular production of reducing equivalents (NADH) and in the maintenance of the mitochondrial redox state [36]. Both pyruvate dehydrogenase [37] and α-Ketoglutarate dehydrogenase [38] are highly susceptible to oxidants inactivation in vitro. Aconitase contains an (4Fe-4S) cluster and is present in two isoforms. The mitochondrial isoform catalyzes the conversion of citrate to isocitrate in the tricarboxylic acid (TCA) cycle, and the cytosolic isoform is involved in iron metabolism [39]. The (4Fe-4S) cluster confers a marked sensitivity to oxidative stress, and the enzyme is inactivated by reactive oxygen species (ROS) [40-42].

Because mitochondria are a site of free radical production and oxidative damage in diabetes [43,44], it is likely that free radical events contribute to declines in mitochondrial function in those subjects. Since mitochondria are the major cellular site involved in fatty acid metabolism, and the main source of reactive oxygen species (ROS), they could play a key role in fat storage and related complications. However, only limited data exist on the involvement of the mitochondrial compartment in this process.

The present study show increased oxidative stress in diabetic compared to age matched normal mice. Diabetic mice show an inactivation of mitochondrial energy producing enzymes. In addition, these mitochondrial enzymes are very susceptible to inactivation by LOOH (HPODE) or H$_2$O$_2$. There is little doubt that one of the most reactive products of fat oxidation is lipid hydroperoxide. Our in vitro studies show that under oxidative stress pyruvate is converted non-enzymatically into acetate that further converted into free fatty acids and triacylglycerols. The goals of this study were to

(i) Determine the mechanism of increased production of free fatty acids and triglycerides in diabetic subjects and
(ii) To identify mitochondrial enzymes inactivated in diabetes under oxidative influence of LOOH or H$_2$O$_2$.

Materials and Methods

Animals

Male BKS.Cg-m +/- Lepr$^a$/+ (Db/db) mice (Jackson Laboratories) of 7-9 months of age were housed in a temperature controlled room with 12:12hr light cycle and maintained with access to food and water ad libitum. Db/db mice have a spontaneous Leprdb mutation and develop hyperinsulinemia, hyperglycemia and obesity by 1 to 2 months of age. Aged matched, C57BKS/J mice were used as controls (BKS). All procedures were conducted with the approval of the Institutional Animal Care and Use Committee of the Ohio State University, Columbus, OH, USA and in accordance with National Institutes of Health Guidelines for the Care and Use of Laboratory Animals. Blood glucose was checked via tail stick with a freestyle glucose monitor in order to verify the presence of hyperglycemia in the Db/db mice.

Isolation of mitochondria

Liver mitochondria were isolated from male BKS.Cg-m +/- Leprdb/J (Db/db) or C57BKS/J mice and Sprague-Dawley rats by differential centrifugation and purified by Percoll centrifugation as described [45]. In brief, liver was removed, washed, and homogenized in MSH EDTA (mannotol, 220mM; sucrose, 70mM; HEPES, 5mM; EDTA, 1mM; pH 7.40). The homogenate was centrifuged at 1,000 x g for 10min, and the supernatant was re-centrifuged at 10,000 x g for 10min to obtain crude mitochondrial fraction. The enriched mitochondrial pellet was layered on a Percoll solution (25%) and centrifuged for 30min at 100,000 xg. The middle layer was extracted and washed twice in MSH (mannotol, 220mM; sucrose, 70mM; HEPES, 5mM; pH 7.40) and centrifuged for 10 min at 10,000 xg. The purified mitochondria pellet was resuspended in 0.5ml MSH and the purity of the mitochondrial preparation was determined. All steps were carried out at 4°C. Purity of the isolated mitochondria was assessed by measuring cytochrome a using $F_{560-650nm}$ 12mM$^{-1}$ cm$^{-1}$ and by measuring transmembrane potential that is rapidly
reversed by the uncoupler FCCP. Only mitochondria with less than 5% impurity were used.

**Human HepG2 culture and treatments**

Human HepG2 cells (ATCC) were cultured in DMEM containing 10% fetal bovine serum, 2mM glutamine, 100 units/ml penicillin and 100µg/ml streptomycin and 4.5mg/L D-glucose. The cells were cultured in 75cm² flask s for 5 days at 37 °C under a humidified atmosphere of 95% air and 5% CO₂ to about 80% confluence. Cells were treated with 50µM LOOH and 1mM H₂O₂ for a period of 4 hours. After treatment, cells were washed and supplemented with fresh DMEM and incubated with (¹⁴C) Na-Acetate (1mM) for 2 hours. At the end of incubation, the medium was aspirated, cells were washed twice with PBS. For the determination of de novo fatty acid synthesis, cells were scrapped from flasks, homogenized with glass homogenizer and protein content was assayed by Bradford method [1976] [46]. The cells were saponified with aqueous KOH at 37°C for 120min. The total fatty acids were extracted with chloroform. Fatty acid synthesis was determined by LC/MS.

**Liver perfusion and culture of primary hepatocytes**

Sprague-Dawley rats were anesthetized by intraperitoneal injection of pentobarbital sodium (50mg/kg b.w.). Initially inferior vena cava was cannulated and the liver was perfused in situ with an oxygenated Hank’s buffer salt solution (HBSS; pH 7.4) containing penicillin/streptomycin (100U/ml) at the rate of 98ml/min for 10min 37 °C. Liver was further perfused with oxygenated HBSS containing penicillin/streptomycin (100U/ml), and insulin (1 x 10⁻⁷ M) (pH 7.4). Livers were perfused for 4 hours. After treatment, cells were re-homogenized with fresh DMEM and incubated with (¹⁴C) Na-Acetate (1mM) for 2 hours. At the end of incubation, the medium was aspirated, cells were washed twice with PBS. For the determination of de novo fatty acid synthesis, cells were scrapped from flasks, homogenized with glass homogenizer and protein content was assayed by Bradford method [1976] [46]. The cells were saponified with aqueous KOH at 37°C for 120min. The total fatty acids were extracted with chloroform. Fatty acid synthesis was determined by LC/MS.

**Assays of Mitochondrial Enzymes**

**Pyruvate dehydrogenase complex (PGDHC)**

PGDHC activity was measured spectrophotometrically using a modified method of Himin and Blass [50] by following the formation of NADH at 340nm at 37 °C. Isolated mitochondria were resuspended in the assay buffer (50mM phosphate buffer, pH 7.4). The reaction mixture contained the assay buffer, 0.2mM thiamine pyrophosphate, 1mM MgCl₂, 2mM NAD⁺, 0.2mM EGTA, 2.6mM L-cysteine, 0.5mM CaCl₂, 0.3mM dithiotreitol, 2mM pyruvate, and mitochondria (60µg/ml). The reaction was initiated by the addition of 0.2mM coenzyme A. Blank samples containing no pyruvate were included in all assays. The activity of the pyruvate dehydrogenase complex was expressed as nanomoles of NADH produced per minute per milligram of mitochondrial protein.

**Aconitase**

Aconitase activity was assessed by the method of Drapier JC and Hibbs (1976) [51]. The mitochondria (60µg/ml) was resuspended in 100mM Tris-HCl buffer, pH 7.4, containing 1mM MgCl₂, 1mM NAD⁺, and 1mM potassium citrate. The reaction was started by adding isocitrate dehydrogenase (2U/ml), carried out at 37°C. Blank samples containing no isocitrate dehydrogenase were included in all assays. The activity of aconitase was expressed as nanomoles of NADPH formed per minute per milligram of protein.

**α-Ketoglutarate dehydrogenase complex (KGDHC)**

KGDHC activity was measured according to the method of Trettier and Adam-Vizi, (2000) [52] by following the formation of NADH at 340nm at 37 °C. Mitochondrial aliquots (60µg/ml) were...
added to a 50mM phosphate buffer (pH 7.4) containing 0.2mM thiamine pyrophosphate, 1mM MgCl₂, 2mM NAD⁺, 0.2mM EGTA, 2.6mM L-cysteine, 0.5mM CaCl₂, 0.3mM dithiotreitol, and 2mM α-ketoglutarate. The reaction was initiated by the addition of 0.2mM coenzyme A. Blank samples containing no α-ketoglutarate were included in all assays. The activity of α-ketoglutarate dehydrogenase complex was expressed as nanomoles of NADH produced per minute per milligram of mitochondrial protein.

**Statistical analysis**

Unless otherwise indicated, data are shown as mean±SD. Data were compared by Student’s t-test. For all experiments, p<0.05 denoted statistical significance.

**Results**

**Body and weight in diabetic mice.**

Db/db mice displayed significantly increased body weight as compared to BKS mice (in g: 56.2±1.2 vs. 27.5±0.4, respectively, p<0.05).

**Lipid peroxidation**

To assess oxidative damage in liver mitochondria, TBARS was tested. Our results showed an increase in TBARS levels in liver mitochondria of diabetic mice compared to normal mice (data not shown).

**Decreased activities of PDHC, aconitase and KGDHC in diabetic mice**

Next we tested the possibility that increased lipid peroxides and consequently increased oxidative stress may cause alterations in mitochondrial energy producing enzymes, making it potentially difficult to maintain energy production in mitochondria. We measured activities of liver PDHC, aconitase and KGDHC in diabetic and normal mice. Our study show decrease in activities of PDHC, aconitase and KGDHC in liver of diabetic mice compared to normal (Figure 1).

**Figure 1: Decreased activities of mitochondrial enzymes upon H₂O₂ and HPODE treatment.**

**Figure 2: Reduced mitochondrial enzyme activities in diabetic mice.**

In order to verify potential effects of lipid peroxides on mitochondrial energy producing enzymes, we treated isolated rat liver mitochondria with LOOH or H₂O₂ for 1hr at room temperature followed by 1hr in ice. Treated mitochondria were
disrupted by freeze and thaw followed by sonication and clear homogenous liquid of mitoplasts was obtained for assay. As shown in Figure 2 treatment of mitochondria with both LOOH and H$_2$O$_2$ caused drastic inactivation of PDHC, aconitase and KGDHC. We further verified inactivation of these mitochondrial enzymes using cultured primary hepatocytes treated with LOOH or H$_2$O$_2$. After treatment mitochondria were isolated from cells, disrupted by freeze and thaw followed by sonication for determination of enzymes activities. As shown in Figure 2, our results show a similar decrease in PDHC, aconitase and KGDHC activities in mitochondria obtained from hepatocytes.

Increased fatty acid after treatment of LOOH

We anticipate that peroxides, in contrast to mitochondrial enzymes, would not adversely affect fatty acid synthesis in the liver cells. Cells were incubated with $^{13}$C-acetate in the presence of hydrogen peroxide for two hours, lipids were extracted and saponified. Tandem MS conditions were optimized with pure palmitic acid (PA). The results presented in Figure 3, shows that $^{13}$C-acetate was incorporated very efficiently. The number of $^{13}$C-atoms increased up to 4 acetate units in the 2hr incubation and then slowly decreased, indicating that the acetate was efficiently utilized for FA synthesis even in the presence of peroxides.

Loss of pyruvate and LOOH

First, we tested loss of pyruvate in the presence of H$_2$O$_2$. Incubation of 50mM pyruvate with 50mM H$_2$O$_2$ causes disappearance of pyruvate with the lapse of time. We noted a decrease of 37% to 22% with increase in incubation time from 30min to 90min (Figure 4A). The reaction was stoichiometric with 1:1 mole equivalents of reactants. We employed mill molar concentrations of reactants in order to follow spectrophotometric detection.

LOOH contains both peroxide component as well as conjugated diene structure (Figure 4B). When LOOH is reduced to LOH, the peroxide component (as measured by LMB reaction) will be lost while the conjugated diene structure will be unaffected. Incubation of LOOH alone did not result in the loss of either peroxide or conjugated diene content As shown in Figure 4, both the open and closed bars remained at 100 % of the initial levels at the end of incubation. Similar incubations of LOOH with pyridine for 60min at 37 °C resulted in a complete loss of LMB assay activity (P<0.05). The levels of conjugated diene however, remained at original levels. We used acetate instead of pyruvate as control in these studies and there was no reduction in either LMB reactivity or conjugated diene in these incubations (data not shown). We used several other α-keto acids (phenyl pyruvic acid, α-ketoglutarate and dehydroascorbate) with result similar to that obtained using pyridine. Thus α-ketoacids are readily decarboxylated by LOOH or H$_2$O$_2$ to yield acetate. As increased cytoplasmic glucose has been reported to generate increased levels of pyruvate, the results can be interpreted to suggest that increased levels of acetate might be generated when there is a oxidative stress.
Non-enzymatic conversion of pyruvate into acetate

Next we tested non-enzymatic conversion of pyruvate into acetate [53].

Release of CO₂ by decarboxylation of α-keto acids

Carbon dioxide evolved in H₂O₂ and pyruvate reaction was measured using radioactive calcium carbonate trapping method. Pyruvate was incubated alone, with H₂O₂ or H₂O₂ alone in a vial with a cap lined with ⁴⁵CaCO₃ (Figure 5A). Carbon dioxide released in experiment where pyruvate was treated with hydrogen peroxide was overwhelming (100%) compared to pyruvate (8%; p<0.001) or H2O2 alone (16%; p<0.002). Similarly formation of carbon dioxide evolution was higher when pyruvate was treated with LOOH (Figure 5B) (100%) when compared to pyruvate (10%; p<0.005) or LOOH alone (52%). The increased production of CO₂ from LOOH alone was perplexing but interesting and reproducible. This suggest a direct peroxide mediated decarboxylation reaction. Such reactions are not unprecedented as it might suggest the formation of a per-acid step.

![Image](image.png)

Figure 5: Release of CO₂ by decarboxylation of α-keto acids.

a) CO₂ trapping with Ca(OH)₂ using H₂O₂ b) CO₂ trapping with Ca(OH)₂ using LOOH.

Discussion

In this study, we establish for the first time that a mouse model of type II diabetes, Db/db mice exhibits not only oxidative stress in liver but also inactivation of three major mitochondrial enzymes, PDHC, aconitase and KGDHC activities.

ROS production is indicated as one of the potential causes leading to insulin resistance and hepatic disease in diabetic animals [54]. For these reasons, we also assessed the oxidative status of liver mitochondria by measuring TBARS reactive substance. Our results show higher TBARS in diabetic mice that indicate an increased oxidative damage compared to normal mice. Increased ROS production in fatty liver has already been documented [55,56], and has been indicated as one of the causes rendering the liver cell more vulnerable to further injury. The main cellular site of fatty acid oxidation is the mitochondria. It is possible that excess fat deposition in the liver is partly due to alteration in mitochondrial function. Using the diabetic mice, in the present paper, we determined whether mitochondrial function is altered in diabetic mice compared to age-matched normal mice. Increase in oxidative stress can inactivate mitochondrial enzymes. In an attempt to determine whether diabetic mice show inactivation of key mitochondrial enzymes we measured PDH, aconitase and KGDHC. These dehydrogenases were chosen because the activities of these enzymes are required for the synthesis of NADH and because these enzymes have been shown to be highly susceptible to free radical-mediated inactivation. We found that all three mitochondrial enzymes are inactivated in mitochondria isolated from diabetic mice (Figure 2). A direct consequence of inactivation of key mitochondrial enzymes and increased oxidative stress as has been obtained in the present study is that all NADH producing steps in the TCA cycle are inhibited that may be reflected by dramatically reduced TCA cycle flux. These observations show that diabetic mice have lower capacity of mitochondrial NADH production. A low level of mitochondrial NADH may stimulate β-oxidation and TCA cycle flux and decrease triacylglyceride formation [57]. However, a high level of pyruvate that accumulate because of inactivation of PDH, aconitase and KGDHC could form acetate non-enzymatically that may convert into fatty acids and triacylglycerol. Unutilized pyruvate and inactivation of mitochondrial enzymes may be related to an increase in triacylglycerols in two possible ways. First, pyruvate non-enzymatically converted to acetate and triacylglycerol and secondly, enzymes that lead to formation of free fatty acids and triacylglycerol are stable in oxidative environments in contrast to PDH, aconitase and KGDHC that are highly susceptible to oxidative environments. Thus inactivation of mitochondrial enzymes and consequently over accumulation of pyruvate may be an alternative mechanism of triacylglycerol deposition in diabetic mice.

These diabetic mice have been reported to have higher triacylglycerols level. Under the conditions that found in those diabetic mice liver, we anticipate that treating isolated mitochondria or primary hepatocytes with LOOH or H₂O₂ reflect a combination of oxidative stress and increased lipid peroxides. Our study clearly show that PDH, aconitase and KGDHC are exquisitely sensitive to LOOH or H₂O₂.

Oxidative stress has been implicated in liver mitochondrial dysfunction in diabetic subjects [58-61]. Oxidative stress
is associated with increases in lipid hydroperoxides, lipid peroxidation, and production of lipid peroxidation products such as 4-hydroxy-2-nonenal (HNE) [62-64], and modification to mitochondrial protein by oxidized lipids [65,66]. HNE is a major product of lipid peroxidation that readily reacts with and inactivates protein [66-70]. Lipid peroxides and H$_2$O$_2$ have been viewed primarily from the perspective of the damage they may impart. It is becoming increasingly apparent, however, that by virtue of the ability to alter protein function [71] they can modify mitochondrial functions. To determine whether observed declines in mitochondrial enzymes could, at least in part, due to increased reactive oxygen species or oxidized lipids, mitochondria were isolated from perfused rat liver and treated with LOOH or H$_2$O$_2$. The inactivation of enzyme was then determined after two hours of incubation. As shown in Figure 2, PDH, aconitase and KGDHC activities are drastically inhibited with LOOH and H$_2$O$_2$. We further verified these results in primary hepatocytes treated with LOOH or H$_2$O$_2$. Our results show similar results in mitochondria isolated with hepatocytes treated with LOOH or H$_2$O$_2$. Based on results presented in Figure 2, we conclude that inactivation of PDH, aconitase and KGDHC could affect supply of NADH to the electron transport chain that controls the rate of NADH-linked mitochondrial respiration. The results of this study indicate that, LOOH and H$_2$O$_2$ interact with PDH, aconitase and KGDHC resulting in enzyme inactivation.

Previous studies showed the susceptibility of aconitase to free radical inactivation [72-76]. Aconitase contains an active site iron-sulfur [4Fe-4S]$^{2-}$ complex. Using electronic spin resonance, it was determined that when purified mitochondrial aconitase is treated with superoxide the (4Fe-4S)$^{2-}$ cluster is oxidized to (3Fe-4S)$^{3+}$, resulting in the release of Fe(II) and H$_2$O$_2$. Similarly KGDHC is also very susceptible to oxidative species. In vitro studies show that lipid peroxidation and lipid peroxidation products induced inactivation of KGDHC involves modification of essential lipoic acid residues covalently linked to E2 subunits of the enzyme [78]. One consequence of inactivation of KGDHC may be oxidation of glutamate by glutamate dehydrogenase that result in the accumulation of α-ketoglutarate. The conversion of glutamate to α-ketoglutarate by glutamate dehydrogenase is thermodynamically unfavorable ($K_{eq} = 1.8 \times 10^{-15}$), and a buildup of α-ketoglutarate would be expected to reduce the rate of glutamate utilisation. In addition, pyruvate dehydrogenase, an enzyme that shares structural and functional similarities with KGDH and contains covalently bound lipoic acid residues [79], may exhibit a similar response to LOOH or H$_2$O$_2$ inactivation. Together inactivation of these mitochondrial enzymes in diabetic mice as well as in vitro experiments in isolated mitochondria and hepatocytes treated with LOOH or H$_2$O$_2$ suggest impairment of NADH production.

During the course of present study, we noted an increase of $^{13}$C acetate as detected by $^{13}$C LC/MS of the hepatocytes treated with LOOH or H$_2$O$_2$. The impact of LOOH or H$_2$O$_2$ on increase in the fatty acid concentration is a direct reflection of its importance as a catastrophic pathway that lead to accumulation of triacylglycerols in the tissue. Under normal conditions, fatty acid synthesis and fatty acid oxidation must always balanced. However, with the inactivation of PDH, aconitase and KGDHC, pyruvate and TCA cycle intermediates may accumulate. Increase in these intermediates may cause allosteric inhibition of TCA cycle activity and fatty acid oxidation [80].

Thus, inactivation of PDHC, aconitase and KGDHC appears to be the primary mechanism by which LOOH or H$_2$O$_2$ may cause an increase in free fatty acids and triacylglycerols in tissues. Because the diabetic mice has more oxidative stress as shown by increased TBARS substances, therefore these mice would be expected to be more sensitive to mitochondrial enzyme inactivation.

It is noteworthy that, under the conditions of our experiments, the magnitude to which PDH, aconitase and KGDHC were inactivated was similar, regardless of the concentration of LOOH or H$_2$O$_2$ utilized (data not shown).

**Conclusion**

Maintenace of mitochondrial function depends on the ability of mitochondria to sense changes in redox status and respond in a manner commensurate with metabolic requirements. In this study, we have provided clear evidence that overall mitochondrial enzymes are inactivated by the addition of LOOH and H$_2$O$_2$ that promotes formation of fatty acids and triacylglycerols in a switch over mechanism from catabolic pathway to anabolic pathway. Whereas these observations suggest a role of oxidative stress in the liver mitochondrial impairments, this study points out a mechanisms whereby mitochondrial enzyme inactivation leads to formation of free fatty acids and triacylglycerols in diabetic subjects. In addition, experiments designed to test the effects of exogenously added LOOH and H$_2$O$_2$ will further enhance our understanding of the process and its physiological significance. In conclusion, our results indicate that alterations in the mitochondrial enzymes induced by LOOH or H$_2$O$_2$ are associated with the ectopic fat storage in the liver. Although this association cannot distinguish between causes and effects, it is interesting that our results fit with the emerging idea that mitochondrial dysfunction can lead to the development of metabolic diseases, such as obesity, type 2 diabetes mellitus. These results suggest that LOOH or H$_2$O$_2$ production may serve to regulate mitochondrial function.

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