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Quantification and mechanisms of oleic acid-induced steatosis in HepG2 cells

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Abstract: Developing a quantifiable in vitro model of steatosis is critical in understanding the pathogenesis of non-alcoholic fatty liver disease (NAFLD) and searching for effective therapies. Using an ORO-based colorimetric measurement, we developed a convenient assay to qualify the degree of OA-induced steatosis in HepG2 cells. We demonstrated that in the absence of exogenous inflammatory mediators, OA-induced steatosis was associated with increased production and secretion of tumor necrosis factor alpha and decreased expression of peroxisome proliferators-activated receptor α in HepG2 cells. OA-induced steatosis was also associated with increased lipid peroxidation, apoptosis, but decreased proliferation in these cells. The increased lipid peroxidation was related to decreased SOD-1, a free radical scavenger enzyme; while increased apoptosis was related to increased active caspase-9. The decreased proliferation mediated by OA-induced steatosis was associated with increased production of p27 with unchanged alanine transaminase (ALT) level in the culture medium, indicating OA-induced steatosis alters cell cycle progression without direct toxicity to these cells. In conclusion, the present study developed a colorimetric assay that accurately quantifies OA-induced steatosis in HepG2 cells. In the absence of exogenous inflammatory mediators, OA-induced steatosis results in a series of pathophysiological changes in HepG2 cells, indicating direct pathogenic roles of hepatocytes in NAFLD.

Keywords: Steatosis, tumor necrosis factor α, peroxisome proliferators-activated receptor α, apoptosis, lipid peroxidation, cell proliferation

Introduction
Non-alcoholic fatty liver disease (NAFLD) is one of the most common liver diseases in Western countries. NAFLD encompasses a spectrum of liver diseases, ranging from simple steatosis to steatosis combined with varying degrees of necroinflammation and fibrosis [1]. It is estimated that approximately 20% to 30% of adults in the United States and other Western countries have excess fat accumulation in the liver [2]. Hepatic steatosis can either be a benign, noninflammatory condition, or can be associated with non-alcoholic steatohepatitis (NASH), a condition that can result in end-stage liver disease [3]. A convenient and quantifiable in vitro model will be valuable in understanding the pathogenesis and evaluating the effects of future therapies of this common liver disease.

Hepatic steatosis in human beings is associated with accumulation of excess oleic acid (OA), a monosaturated omega-9 fatty acid and the end product of de novo fatty acid synthesis [4]. Treatment of HepG2 cells, a human hepato-

Abbreviations: NAFLD, non-alcoholic fatty liver disease; OA, oleic acid; ORO, oil red O; TNF-α, tumor necrosis factor α; PPARα, peroxisome proliferator-activated receptors α; ALT, alanine transaminase; NASH, nonalcoholic steatohepatitis; ALD, alcoholic liver disease; MTT, 3-(4, 5-dimethyl-thiazol-2yl)-2, 5-diphenyl-tetrasodium bromide; DMEM, Dulbecco’s modified eagle’s medium; FBS, fetal bovine serum; LPO, lipid peroxidation; FFA, free fatty acid; ROS, reactive oxygen species; CDK, cyclin-dependent kinase.
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toblastoma cell line, with OA induces morphological similarities to steatotic hepatocytes [5, 6], but quantification of OA-induced steatosis has not been well established. Oil red O (ORO) is a lysochrome (fat-soluble dye) diazo dye used for staining of neutral triglycerides and lipids on frozen sections. ORO stains protein bound lipids in paraffin sections. This biochemical technique has been predominantly used for triglyceride staining on tissue sections [7]. Quantification of ORO stained cell with light microscopy is inconvenient, and its accuracy is operator-dependent that makes reliable quantification difficult.

Although studies demonstrated profound pathogenic changes in cytokines and signal transduction, lipid metabolism, and hepatocytic apoptosis and injury in both alcoholic liver disease (ALD) and NAFLD [8-12], the underlying mechanisms remain to be determined. Developing a human cell model in which steatosis is qualifiable will hold a very special value in understanding mechanisms of NAFLD from liver cell level, because such human cell model exclude the interference from the matrix and other non-hepatocytic cells.

In the present study, we used OA and HepG2 cells and developed a cell model of steatosis. Using an ORO-based colorimetric measurement developed in the present study, we were able to rapidly and accurately quantify the degree of OA-induced steatosis. This model was also used to determine the underlying mechanisms of OA-induced steatosis in HepG2 cells.

Materials and methods

Reagents

Dulbecco’s modified eagle’s medium (DMEM), fetal bovine serum (FBS), trypsin-EDTA, and penicillin-streptomycin-fungizone were purchased from Cambrex Bio Science Walkersville, Inc. (Walkersville, MD). The cell proliferation assay was performed using a Cell Titer 96 AQueous One Solution Reagent purchased from Promega Corporation (Madison, WI). OA-conjugated BSA solution and ORO were purchased from Sigma Chemical Company (St. Louis, MO). Antibodies to p21waf1/cip1, p27kip1, Bcl-2, Bax, PPARα, cleaved caspase-9, TNF-α, SOD1, and β-actin were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Rabbit anti-alanine transaminase (ALT) peptide antibody was produced in our laboratory and its specificity (54 kDa) was confirmed by Western blot. Protein A/G PLUS-agarose was from Santa Cruz Biotechnology (Santa Cruz, CA). SuperSignal West Dura Extended Duration Substrate for Western blots was purchased from Pierce Biotechnology (Rockford, IL). The protein assay kit, gel electrophoresis buffers, 4%-15% gradient Tris-HCl gels and 10% Tris-HCl glycine gels, and the nitrocellulose transfer membrane were purchased from Bio-Rad Laboratories (Hercules, CA).

Cell culture

HepG2 cells were cultured in DMEM-based medium as described before. [7, 8]. The experiments were performed when cells reached about 80% confluence, and after cultured in FBS-free media for 24 hrs.

OA-induced steatosis

Once approximately 80% confluence and cultured in FBS-free medium for 24 hrs in 96-well culture plate, HepG2 cells were treated with 200 μl of different concentration (0.1-2.0 mM) of OA solution for 24 hours. After the medium was removed, 100 μl of fixative solution were added, and incubated at room temperature for 10 minutes. Control cells were treated with OA-free medium containing albumin.

ORO cell staining

ORO solution was made by mixing 2.4 ml of ORO stock solution with 1.6 ml of distilled water then undergoing filtering. After removing the fixative solution from each well of the 96-well culture plate, HepG2 cells were treated with 200 μl of different concentration (0.1-2.0 mM) of OA solution for 24 hours. After the medium was removed, 100 μl of fixative solution were added, and incubated at room temperature for 10 minutes. Control cells were treated with OA-free medium containing albumin.

ORO-based steatosis quantification

To develop a rapid and convenient quantification assay for OA-induced steatosis in HepG2 cells, ORO-based cell staining was combined with a colorimetric assay using 96-well plate
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Cell proliferation assay

Cell proliferation was determined using an MTT assay as previously reported [13, 14]. Briefly, 5x10^3 cells were plated into a 96-well plate containing 100 µl of cell culture medium in triplicate and were treated with different concentrations of OA. The effects of OA on HepG2 cell growth were then determined by optical density absorbance as previously reported [13, 14].

Assay of apoptosis

Apoptosis was determined with two different methods, as previously reported [14]. Briefly, after treatment of HepG2 cells with respective doses of OA for 24 hrs, cell death detection was performed in duplicate and measured by calculating the ratio of absorbance of treated cells relative to the control cells. The activated caspase-9, Bax, and Bcl-2 were measured by Western blots [13, 14].

Western blots

Protein expression was determined by Western blot [7, 8]. Briefly, HepG2 cells were cultured with OA for 24 hours, and then the cell pellets were lysed with 1 ml of the lysis buffer. The lysates were centrifuged at 12,000 rpm for 10 minutes at 4°C. The supernatants were used to detect TNF-α, PPARγ, ALT, SOD1, p27kip1, p21waf1/cip1, Bax, Bcl-2, and cleaved caspase-9. All the Western blots were repeated for three times. Western blot for β-actin was used as internal control. To quantify the results, the relative amount of each protein was determined by digital scanning of the hybridizing bands, as previously described [13, 14].

Assay for LPO in OA-induced HepG2 cells

Lipid peroxidation (LPO) levels were determined using a commercial lipid hydroperoxide assay kit from Cayman Chemical Co. (Ann Arbor, MI). Medium and cellular homogenates (500 µl) were deproteinated and extracted under acidic conditions with 1 ml ice-cold deoxygenated chloroform, and the chloroform extract was removed following centrifugation (1500 X g for 5 minutes at 0°C) for LPO determination. After mixing with chromogen reagent, 200 µl of the extracted samples were transferred to a 96-well plate. The absorbance was determined at 500 nM. The 13-hydroperoxy-octadecadienoic acid was used as a lipid hydroperoxide standard to construct a standard curve (linearity from 0.5 to 5 nM hydroperoxide).

Statistical analysis

The student t test was used to compare the difference of the means. Linear regression was used to calculate correlation coefficient and determine the association between OA-induced steatosis and OD values measured by ORO colorimetric assay developed in the present study.

Results

OA-induced steatosis in HepG2 cells

The histological definition of steatosis is the visible accumulation of lipid droplets in more than 5% of hepatocytes [15]. To determine the optimal concentration of OA to induce steatosis, HepG2 cells were cultured at OA concentrations of 0.1 mM, 0.5 mM, 1 mM, and 2 mM for 24 hours. In untreated control of HepG2 cells, ORO staining revealed almost absence of intracellular lipid (Figure 1A). After treatment with OA, lipid droplets were accumulated in the cytoplasm of HepG2 cells (Figure 1B). OA at concentrations between 0.1 mM to 2 mM, reliably induced steatosis in HepG2 cells in a dose-dependent pattern as determined by ORO staining.
Steatosis staining with ORO and its quantification under microscopy can be inconvenient, and its accuracy is operator-dependent. To objectively measure the degree of OA-induced steatosis in HepG2 cells, we develop a colorimetric assay to quantify steatosis, based on the unique feature of high organic solubility of ORO that can be colorimetrically quantifiable after its release by an extract solution from stained steatosis in the cells.

After confirming the feasibility and optimizing the conditions of the ORO-based colorimetric assay, we then tested its sensitivity and reproducibility. As shown in Figure 1C, our ORO-based colorimetric quantitative assay revealed an excellent correlation of the measured optical density with OA-induced dose-dependent steatosis in HepG2 cells with a correlation coefficient ($r^2$) as high as 0.97 ($p=0.001$). The tests were repeated three times with the same results. This method easily and reliably quantifies the difference of steatosis induced by OA at the dose ranged from 0.1 to 2 mM, as observed under microscopy. Thus, ORO-based colorimetric quantitative assay developed in the present study is not only convenient, quantifiable, but also highly sensitive and reproducible.

**Effects of OA-induced steatosis on production of TNF-α and PPARα in HepG2 cells**

It is well known that elevated TNF-α plays a key role in the pathogenesis and disease progression of NAFLD [16-18]. Overexpression of TNF-α mRNA in both liver and adipose tissue has been reported in severely obese patients with NASH [19]. It was also reported that treatment of HepG2 cells with free fatty acid (FFA) resulted in increased production of TNF-α mRNA [20]. To ascertain if this effect also occurs at the translational level, we tested if OA treatment of HepG2 cells alters production and secretion of TNF-α. We found that OA treatment significantly induced TNF-α expression in HepG2 cells and its secretion to the culture medium (Figure 2A and 2B).
A growing body of literature implicates that peroxisome proliferators activated receptors (PPARs) play important roles in the pathogenesis of NAFLD [10]. For instance, PPARα may augment fatty acid oxidation and protects against steatosis [21-23]. To determine whether steatotic hepatocytes are a significant source of PPARα expression in HepG2 cells before and after OA treatment. We demonstrated that OA-induced steatosis was significantly associated with reduced expression of PPARα in HepG2 cells (Figure 2C).

**Effects of OA-induced steatosis on lipid peroxidation in HepG2 cells**

Many studies have indicated the important pathogenic role of oxidative stress in NAFLD. For instance, oxidative stress can initiate membrane lipid peroxidation and loss of cell viability in cultured hepatocytes [24-26]. The OA-induced steatosis in HepG2 cells provides a unique model to study the changes of lipid peroxidation in the hepatocytic level. To determine how OA-induced steatosis alters oxidative stress, we determined lipid peroxide formation by measuring lipid hydroperoxide in HepG2 cells after 24 hour exposure to OA. We found a significant rise in lipid peroxides in the OA-treated HepG2 cells (Figure 3A). Superox-
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SOD-1 (SOD-1) is a free radical scavenger enzyme that protects against cellular membrane injury mediated by lipid peroxidation. We observed that SOD1 expression was significantly reduced in HepG2 cells with OA-induced steatosis (Figure 4A).

Effects of OA-induced steatosis on cell proliferation and cell cycle progression in HepG2 cells

Since steatosis has been reported to alter cell proliferation [27], we determined whether OA-induced steatosis affects cell proliferation by measuring effects of OA on HepG2 cell viability using MTT assay [13, 14]. As shown in Figure 3B, we found OA-induced steatosis was associated with significant inhibition of HepG2 cell proliferation in an OA-dose dependent manner at the concentration from 0.2 mM to 1.4 mM. For instance, OA inhibited HepG2 cell proliferation by 21% at 1 mM, and 37.5% at 1.4 mM of the concentration.

ALT release from hepatocytes has been a traditional marker of liver cell injury. To determine if OA-induced steatosis could result in direct liver cell injury that might count for decreased cell proliferation, we measured ALT level in the cell lysates and the culture medium without treatment (as controls) and after treatment with various OA concentrations. As shown in Figure 3C, compared to the untreated HepG2 cells, treatment with OA at the concentration from 0.5 mM to 2 mM for 72 hours did not increase the concentration of ALT in these cells and the culture medium. Thus, OA-reduced proliferation of HepG2 cells is not secondary to its direct toxic effects.

To further determine the effects of OA-induced steatosis on HepG2 cell proliferation, we assessed if OA-induced steatosis alters expression of several modulators involved in the progression of the cell cycle and cellular proliferation pathways. P21waf1/cip1 and p27kip1 are two cyclin-dependent kinase (CDK) inhibitors involved in suppression of the cell cycle at the G1-S checkpoint. We found that OA-induced steatosis significantly increased the expression of p27kip1, but not p21waf1/cip1 expression in HepG2 cells, as shown in Figures 4B and 4C.

Effects of OA-induced steatosis on apoptosis in HepG2 cells

Clinical studies have shown that NASH is associated with altered apoptosis activity [28, 29]. To evaluate the effects of OA-induced steatosis on apoptosis activity in HepG2 cell, we tested the cell death by measuring cytoplasmic histone-associated DNA fragments. We found that OA-induced steatosis significantly increased apoptosis activity in HepG2 cells in a dose-dependent manner at OA dose ranging from 0.1 mM to 1 mM (Figure 3D). At OA doses greater than 1 mM, the apoptosis activity measured in HepG2 cells appeared to reach a maximum level and began to plateau.

To determine the mechanisms involved in OA-induced apoptosis, we measured the expression of several apoptosis-related proteins. In OA-treated HepG2 cells, we found a significant increase in the expression of cleaved caspase-9, a marker of apoptosis signaling in the caspase activation pathway (Figure 4D). Although Bax, a pro-apoptotic marker, and Bcl-2 an anti-apoptosis marker, have been associated with apoptosis activity, we could not detect
changes in their expression after OA induction (Figures 4E and 4F).

Discussion

OA-induced steatosis in HepG2 cells may serve as an in vitro model for studying fatty liver disease. Consistent with previous reports [30], we revealed OA induces steatosis in HepG2 cells in a dose-dependent manner that can be assessed by ORO biochemical staining. However, this method is neither convenient, nor accurate in quantification, therefore, cannot be used as a routine steatosis assay, especially for the study of disease mechanisms and development of new therapy to NAFLD.

In the present study, we utilized the feature that after biochemical staining of steatosis, ORO can be released from the stained cells that could be further quantitatively measurable. Using this concept, we were able to develop an ORO-based colorimetric quantification assay to measure OA-induced steatosis in HepG2 cells. Our further analysis indicated that this novel method is not only convenient, and highly reproducible, but also accurate with a very high correlation between the dose of OA and the degree of steatosis as expressed as absorbance of optical density and histological ORO digital pixel measurement. Thus, the cell model of OA-induced steatosis, together with this novel ORO-based colorimetric quantitative assay, will provide valuable tools to study the pathogenesis and develop new treatment for NAFLD.

It is well known that TNF-α plays important pathogenic roles in both ALD and NAFLD [16-18]. For instance, TNF-α has been associated with insulin resistance and induce inflammatory cytokines formation. The mean plasma level of TNF-α was significantly higher in NAFLD patients with abnormal ALT than controls [31-33]. A key question is whether increased TNF-α in NAFLD is from hepatocytes and/or from other inflammatory cells. A recent study demonstrated that FFA treatment induces TNF-α mRNA in HepG2 cells [19], but it remains unknown if FFA-induced steatosis also promotes translation of TNF-α mRNA. In the present study, we demonstrated that OA-induced steatosis significantly increased TNF-α production and secretion from HepG2 cells. These in vitro results indicate that OA-induced steatosis promotes translation of TNF-α mRNA. Our data further support the pathogenic role of hepatocyte-derived TNF-α in NAFLD. Studies reported that TNF-α also stimulates ROS generation and induces lipid peroxidation [34]. We demonstrated that OA-induced steatosis is associated with increased lipid peroxides in HepG2 cells. However, the pathogenic role of increased TNF-α in up regulation of lipid peroxides in OA-induced steatosis remains to be determined.

As HepG2 cells accumulate intracellular lipids, there is a significant rise in lipid peroxides. In previous studies, unsaturated fatty acids have been shown to induce the cytochrome P450 2E1 (CYP2E1) enzyme pathway and stimulate lipid peroxidation, which subsequently promotes apoptosis and cell toxicity [35]. Consistent with these findings, we found that OA-induced steatosis was associated with a significant rise in lipid peroxide formation in the OA-treated HepG2 cells. The pathogenic role of increased lipid peroxides in the cell injury of OA-induced steatosis in HepG2 cells is further supported by our findings that OA-induced steatosis in these cells was associated with a significantly decreased SOD-1, a free radical scavenger enzyme that protects against cellular membrane injury mediated by lipid peroxidation. These finding also indicated a potential therapeutic role of anti-peroxidation agents for NAFLD.

PPARα activates expression of a series of target genes involved in the uptake, transportation, and β-oxidation of fatty acids [36, 37]. Several studies have shown that PPARα increases fatty acid catabolism, and therefore, may prevent hepatic fat deposition [38-42]. Our findings that OA-induced steatosis resulted in reduced PPARα expression in HepG2 cells indicated its possible pathogenic role in NAFLD. Taken together, our results suggested that the association of OA-induced steatosis with increased lipid peroxidation may be mediated by decreased PPARα expression in these cells. This speculation is supported by the clinical and experimental evidence that PPARα antagonists may improve steatosis in patients and animal model [43, 44]. Further determining this association will be help in developing novel therapeutic approach to this disease.

Hepatocyte apoptosis is a feature of fatty liver disease [45], which may reduce cell regeneration or proliferation. We found that OA, at 0.1-1mM dose range, leads to increased apoptosis
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of HepG2 cells with OA-induced steatosis. This was associated with increased production of activated caspase-9 of the apoptosis cascade. Although apoptosis may be associated with decreased proliferation of the liver cells with steatosis, it may also contribute to the pathogenesis of the liver injury in non-alcoholic fatty liver disease. We could not demonstrate any significant effects of OA-treatment on Bcl-2 and Bax expression, the two other apoptotic modulators, indicating that they are unlikely regulating apoptosis in OA-induced steatosis in HepG2 cells.

The effects of steatosis on hepatocyte proliferation are unknown. Using MTT assay, we demonstrated that OA-induced steatosis in HepG2 cells was associated with inhibition of cell proliferation. These results are supported by a recent report of microarray analysis [27]. Since OA-induced apoptosis was saturated at the dose of 1 mM, the decreased HepG2 proliferation could not be solely explained by apoptosis. On the other hand, we demonstrated its association with increased p27 expression. Since p27 functions as a cyclin-dependent kinase (CDK) inhibitor involved in suppression of the cell cycle at the G1-S checkpoint, our results indicated that OA-induced steatosis decreases cell proliferation by inhibiting p27 expression, therefore, G1-S progression.

The clinical presentation of NAFLD could be very variable from persistently normal to significantly increased ALT. It is not known if steatosis results in direct liver injury and elevated ALT. Using HepG2 in vitro model, we demonstrated that baseline level of ALT was comparable in the culture medium of both untreated and OA-treated HepG2 cells. At a wide range of concentration, OA treatment does not alter the ALT level in the culture medium. These findings suggest that OA-induced steatosis itself does not result in liver cell injury. Instead, it is likely that hepatocytes’ direct response to steatosis results in generation of a series of inflammatory mediators that may cause liver cell injury. Further studies will be needed to detail theses underlying mechanisms.

In conclusion, the present study developed an ORO-based colorimetric assay to quantify lipid accumulation in OA-induced HepG2 cells, which provides a convenient tool for studying the pathogenesis and therapy for NAFLD. We also assessed the effects of OA-induced steatosis on HepG2 cell proliferation, apoptosis, and lipid peroxidation through a complicated signaling. These data are very valuable in understanding the pathogenesis of NAFLD.

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