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Oleic acid differentially affects lipid droplet storage of *de novo* synthesized lipids in hepatocytes and adipocytes[†]

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Lipogenesis is a vital but often dysregulated metabolic pathway. Here we use optical photothermal infrared imaging to quantify lipogenesis rates of isotopically labelled oleic acid and glucose concomitantly in live cells. In hepatocytes, but not adipocytes, we find that oleic acid feeding at 60 μ M increases the number and size of lipid droplets (LDs) while simultaneously inhibiting storage of *de novo* synthesized lipids in LDs. Our results demonstrate alternate regulation of lipogenesis between cell types.

Liver diseases have become a significant health concern worldwide, especially with both obesity and alcohol consumption on the rise.¹ Non-alcoholic fatty liver disease (NAFLD) is the most common liver disease at a staggering 25% lifetime prevalence worldwide.² Dysregulation of *de novo* lipogenesis (DNL) is a significant factor in these diseases.³ Therefore, it is important to study the DNL pathway at the cellular level, including how responses to external stimuli, such as scavenging pathways, adjust overall rates of DNL and total lipogenesis. Thus, observing competing lipogenic pathways *in cellulo* is crucial to answering these pressing questions.

As a monounsaturated fatty acid, dietary oleic acid (OA) enters lipogenesis during fatty acid synthesis and undergoes esterification and storage in lipid droplets (LDs, Fig. 1A).⁴ While fatty acid overload is typically associated with hepatic steatosis and the onset of diseases such as obesity and NAFLD, counter-evidence suggests that unsaturated fatty acids act as a protectant from cell death.^{4–6} Oleic acid in particular has been shown to prevent cholesterol synthesis and liver lipotoxicity,⁴ while simultaneously affecting LD morphology and formation in ways characteristic of disease (increasing both their number and size).^{7,8} This suggests OA can specifically regulate deleterious

aspects of lipogenesis associated with increased general lipid content and, furthermore, that LDs play a key role in this regulation.⁶ Enzymatic activity assays and mRNA studies have indicated OA may regulate cholesterol generation by inhibiting the expression and activity of acetyl-CoA carboxylase (ACC, Fig. 1A), at least short term (< 4 h).^{5,9} Additionally, unsaturated fatty acids like OA have been shown to decrease expression of and inhibit sterol regulatory element binding protein 1c (SREBP-1c), a master regulator of DNL.^{10–12} Thus, a better understanding of morphologic and metabolic effects of OA on DNL and other lipid pathways is needed to elucidate its role in disease.

Measuring metabolic pathways directly *in vivo* would provide this vital information, however, techniques for doing so are limited. Fluorescently labelled lipogenesis precursors like glucose and fatty acids are non-metabolizable or significantly disruptive. Other glucose analogs present a similar issue and



Fig. 1 Schematic of experiments and data collection. (A) Isotope labelling of *de novo* lipogenesis and fatty-acid scavenging pathways where oleic acid is surmised to inhibit ACC. (B) Live cell data collection by OPTIR. Cells grown on a calcium fluoride cover slip are mounted in PBS using a spacer. (C) Multispectral imaging at single frequencies.

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are primarily used in uptake assays.¹³ The exception is isotope labelling of precursors, which is nonperturbative, nontoxic, and well metabolized.^{13–15} While isotopic labels are easy to incorporate and have been used extensively for mass spectrometry experiments,^{16,17} a non-destructive technique is necessary to monitor these probes in living cells. Furthermore, research has shown that metabolism is heterogeneous throughout the cell, with external stressors stimulating condensation of metabolic components and other regulating factors.¹⁸ This presents a need for techniques that can provide both spatially and temporally resolved metabolic information.

Vibrational microspectroscopy is a promising means of tracking metabolism throughout the cell using nonperturbative isotopic probes.^{13–15,19} Biomolecules have specific vibrational signatures that can provide information about the surrounding environment.²⁰ FTIR and Raman microspectroscopy, stimulated Raman scattering (SRS) in particular,^{13,21} have been successfully used in cellular metabolism studies, but each present its own challenges when working with live cells.^{22,23} Raman-based imaging is restricted by the small Raman scattering cross section, resulting in long imaging times, high powered lasers, and/or limited spectral coverage, making live cell work difficult.²³ FTIR is limited by the strong IR absorbance of water overlapping with the peaks of many other biomolecules and spatial resolution in the micron range, which is larger than most cellular organelles and therefore provides no sub-cellular information.^{14,24}

Our approach, optical photothermal infrared microscopy (OPTIR), addresses the issues posed by traditional Raman and FTIR imaging techniques. OPTIR is a nonperturbative and nondestructive infrared imaging method with a spatial resolution below 500 nm. In this pump–probe system the sample is "pumped" with an IR beam, inducing a photothermal effect, which is monitored by a change in intensity of the probe, a visible laser. The diffraction limit is based on the probe laser, thus surpassing the spatial resolutions of traditional FTIR spectroscopy (Fig. S1, ESI†). The specific geometry also leads to significantly less signal contribution from water.^{15,25} OPTIR has previously been used on fixed samples to obtain information on metabolic rates and chemical compositions,^{26,27} but live cell imaging, especially in H₂O, has been more limited.^{15,25,28}

We previously used OPTIR to track rates of DNL by feeding ¹³C glucose to differentiated 3T3-L1 cells (adipocytes) and monitoring incorporation of the labelled carbon into the ester carbonyl of triglycerides in both live and fixed cells.¹⁵ Here, we extend this study to the hepatocyte-derived Huh-7 cancer cell line because hepatic DNL is especially relevant to NAFLD and other metabolic diseases. Further, we incorporate a multiplexed imaging approach to monitor DNL and fatty-acid scavenging concomitantly (Fig. 1). Multiplexed imaging is widely used in single-cell studies because it provides a more comprehensive view of intertwined pathways.^{22,29}

Multiplexed imaging (Fig. 1C) in both live hepatocytes (Fig. 2A) and adipocytes (Fig. 2B) can be achieved because the vibrational probes used exhibit strong IR bands easily distinguishable amongst the fingerprint of other biomolecules (Fig. 2C). Briefly, by feeding Huh-7 and differentiated 3T3-L1



Fig. 2 Live cell characterization of the cell lines (A) Huh-7 and (B) differentiated 3T3-L1 after feeding with ²H oleic acid (60 μ M) and ¹³C glucose (4.5 mg mL⁻¹). Scale bars are 20 μ m. (C) Full IR spectra of a point within a fixed Huh-7 cell LD 48 hours after feeding ²H OA. ²H-C from the ²H oleic acid (green, structure in inset) which has been esterified in LDs. 2214 cm⁻¹ is assigned to ²H-C stretching. ¹³C=O stretch at 1703 cm⁻¹ where the ¹³C comes from metabolized ¹³C glucose (blue, structure in inset). Pictured alongside the unlabelled ¹²C=O band at 1744 cm⁻¹.

cells both ²H oleic acid and ¹³C glucose, the regulatory effect of oleic acid and resulting competing scavenging pathways can be observed (Fig. 1A and Fig. S2, ESI†). Live cell imaging is achieved by growing cells on a CaF₂ coverslip and mounting it in buffer on a glass slide (Fig. 1B). Rates of LD storage of DNL and OA scavenging/esterification are determined by ratios of probe bands to the lipid bands. These images are collected sequentially as single wavenumber images set to frequencies associated with metabolism of the isotope-labelled probes as well as the protein amide-I and unlabelled lipid carbonyl stretch (Fig. 1C).

The full spectrum of ²H oleic acid (Fig. S3A, ESI[†]) shows strong peaks at 2200 cm⁻¹ and 2100 cm⁻¹ attributed to C⁻²H stretches in the "cell silent region".¹⁴ These peaks also appear in LDs of cells fed ²H OA (Fig. 2C and Fig. S2, S3B, ESI[†]). Shoulders at 2214 cm⁻¹ and 2246 cm⁻¹ are assigned to the asymmetric stretch of the singular C⁻²H₃ and C⁻⁻C⁻²H in OA, respectively.^{30,31} The 2214 cm⁻¹ shoulder was used to collect single wavenumber images to specifically measure uptake and esterification of ²H OA. Free OA is generally not visible outside of LDs because the concentrations are below the detection limit of the instrument (Fig. S4, ESI[†]). The OA is esterified into triglycerides before storage in LDs, but this does not shift the vibrational bands of the C⁻²H stretches.³²

In the case of ¹³C glucose, as the glucose is metabolized the ¹³C is incorporated into fatty acids (Fig. 1A) that are eventually incorporated into triglycerides and cholesteryl esters that are stored in LDs. An increase in reduced mass affects the vibrational frequency, resulting in a redshift of lipid vibrational bands.³³ The most significant is the redshift of the triglyceride and cholesteryl ester ¹²C=O ester carbonyl stretch (hereafter referred to as ¹²C lipid band) from 1744 cm⁻¹ to 1703 cm⁻¹, which we use to monitor ¹³C triglyceride and cholesteryl ester generation and storage in LDs (Fig. 2C).¹⁵

To ensure that the technique used previously on adipocytes¹⁵ can be applied to hepatocytes, a similar experiment was performed in which Huh-7 cells were incubated over the course of 72 hours with ¹³C glucose as sole source of glucose and imaged live. As anticipated, the resulting images depict smaller LDs



Fig. 3 Visualization of rates of glucose-derived *de novo* lipid storage in live Huh-7 cells fed with both ²H OA (60 μ M) and ¹³C glucose (4.5 mg mL⁻¹). (A) Brightfield images of live cells. (B) ²H OA ratio images of ²H lipid to the ¹²C lipid band, after correction. (C) ¹³C glucose ratio images of the ¹³C lipid band to the ¹²C lipid band, after correction. (D) Representative spectra of the live Huh-7 cell at the 48-hour time point both inside a LD (pink) and outside the droplet (blue) (E) Average ratios of ²H lipid band (grey) and ¹³C lipid band (red) to ¹²C lipid band across many cells (*N* = 14). Scale bars are 10 μ m.

compared to those of adipocytes, but similar ${}^{13}C = O/{}^{12}C = O$ lipid ester carbonyl ratios and therefore rates of glucose-derived *de novo* lipid storage in LDs (Fig. S5, ESI†). IR spectra collected with 500 nm spatial resolution through a lipid droplet of a live Huh-7 cell demonstrate the spatial resolution of the instrument (Fig. S6, ESI†). In this work, cell data was collected using multispectral single wavenumber imaging as it is approximately 25 times faster than full hyperspectral imaging.¹⁵ This enables increased statistics with similar, though not as detailed, spectral information.

Live cell multispectral imaging was then performed on Huh-7 (Fig. 3) and differentiated 3T3-L1 cells (Fig. S7, ESI†) fed both ²H OA and ¹³C glucose. Trends were confirmed with fixed cell multispectral imaging (Fig. S8 and S9, ESI†). We found that both cell lines readily took up ²H OA and sequestered it in LDs by 24 hours. Ratio images of the ²H lipid band of ²H OA to the ¹²C lipid band reveal that ²H OA concentrations remain constant in LDs over the 72 hour time period (Fig. 3B and Table S1, ESI†). Ratio images of the ¹³C lipid band to the ¹²C lipid band specifically track DNL resulting in LD lipid storage because the ¹³C in the red-shifted lipid band comes only from the ¹³C glucose. Fig. 3C shows the increasing ratio of ¹³C labelled lipid, allowing us to estimate the rate of glucose-derived *de novo* lipid storage in LDs (Fig. 3 and Tables S1–S4, ESI†). Notably, ²H OA and ¹³C glucose signal was primarily confined to LDs in living cells (Fig. 3A–D and Fig. S7B–D, ESI†). As LDs are the primary endpoint of DNL and critically implicated in disease we focused our analysis on the storage rates within LDs. 3T3-L1 cells had a much lower ratio of ²H lipid to total lipid in both live and fixed cells, but similarly showed a constant ratio from 24 to 72 hours (Fig. S7 and Table S5–S8, ESI†). This may be due to higher lipid content and larger LDs in adipocytes as compared to hepatocytes, leading to a lower ratio of OA to total lipid. In agreement with prior work,¹⁵ there was significant cell to cell variability in both cell lines as well as limited heterogeneity of ratios within cells.

Huh-7 cells showed a marked increase in both number and size of LDs after ²H OA feeding (Fig. S10, ESI[†]), agreeing with previous literature.^{8,34} At the 48 hr time point, there were on average 9.2 \pm 4.7 LDs in OA fed cells compared with 5.6 \pm 2.4 LDs in vehicle fed (BSA only) conditions (Table S9, ESI[†]). Similarly, LD diameter was 2.5 \pm 0.8 µm in ²H OA fed conditions and 2.1 \pm 0.5 µm in vehicle fed conditions (Table S9, ESI[†]). The differences in both number and diameter of LDs were significant with *p* values below 0.0005.

Interestingly, ²H OA fed 3T3-L1 cells had much higher levels of glucose-derived *de novo* lipids stored in LDs compared to ²H OA fed Huh-7 cells with a final ¹³C/¹²C lipid ratio of ~0.7 as compared to ~0.3 (Fig. S7E, ESI† and Fig. 3E red), respectively. To further investigate this, we compared the average ratios of ²H OA fed cells to cells fed only the BSA vehicle without OA (Fig. 4). Both sets contained reduced FBS to limit other sources of fatty acid scavenging. In Huh-7 hepatocytes, at every timepoint the ratio of ¹³C lipid to ¹²C lipid was dramatically and significantly (p < 0.05) reduced in ²H OA fed conditions (Fig. 4A). Conversely, the ratio remained largely unaffected in 3T3-L1 adipocytes (Fig. 4B). The same relationship was observed in fixed Huh-7 and 3T3-L1 cells (Fig. S11 and Tables S1–S8, ESI†). Therefore, DNL resulting in lipid storage in LDs is downregulated in ²H OA fed Huh-7 cells and unchanged in ²H OA fed 3T3-L1 cells.

However, as evidenced by the increased number and size of LDs (Fig. S10 and Table S9, ESI \dagger), the total ¹²C lipid content



Fig. 4 Effect of OA feeding on rates of *de novo* lipid storage. Ratio of ${}^{13}C$ lipid to ${}^{12}C$ lipid in live (A) Huh-7 and (B) 3T3-L1 cells fed with both ${}^{2}H$ oleic acid and ${}^{13}C$ glucose (red) or BSA vehicle and ${}^{13}C$ glucose (purple) at 24, 48, and 72 h. *N* = 8–16 cells per time point **p*-value < 0.05 ****p*-value < 0.005 N.S. *p*-value > 0.05.

Communication

also increases in ²H OA fed Huh-7 cells. This increase in LDs could artificially decrease the ratio used to approximate DNL. However, the rate is measured by the rate of change of the $^{13}C/^{12}C$ lipid and, therefore, should not be affected by a change in total lipid content. Thus, we fit the average ratio data of the Huh-7 cells and found very different rates with a linear slope of approximately 0.01 $^{13}C/^{12}C$ lipid ratio per hr for the BSA vehicle fed cells and 0.005 $^{13}C/^{12}C$ lipid ratio per hr for the ²H OA fed cells for both live and fixed cells (Fig. S12, ESI†). Rates of storage of glucose-derived *de novo* lipids in LDs is slowed by half by OA feeding. In 3T3-L1 cells no difference is seen between the ratio in the BSA vehicle and ²H OA fed conditions at any time point (Fig. 4B).

We found that ²H OA feeding in Huh-7, but not differentiated 3T3-L1 cells, is correlated with both an increase in total lipogenesis and therefore LD size and number and a decrease in LD storage of *de novo* lipids. This reduction after OA feeding has previously been reported in glioma cells and was associated with OA inhibition of ACC activity and expression.^{5,35} In line with our data, OA is also known to inhibit de novo fatty acid and cholesterol synthesis but increase triglyceride and general lipid content.^{5,9} OA has also been shown to be protective against cell death in hepatocytes, even when inducing steatosis.^{34,36} This may be due to its differential effects on DNL and total lipogenesis as DNL primarily generates palmitate, a fatty acid associated with cell death.^{3,36} Interestingly, OA feeding had little impact on rates of LD storage of de novo lipids in adipocytes, despite being readily taken up and stored in LDs (Fig. S7, ESI⁺). The difference between the two cell lines could simply be a concentration effect as cultured adipocytes contain more lipid, perhaps making them less sensitive to small changes in the synthesis of new lipid. However, this seems unlikely as rates are similar between adipocytes and hepatocytes in the BSA vehicle fed conditions (Fig. 4). OA feeding concentrations (60 μ M) were significantly below fatty acid levels known to cause fatty acid overload and cell death.³⁴ At the concentrations of OA (60 µM) and BSA vehicle (30 µM) used, OA uptake should depend primarily on BSA concentration and be similar between the cell lines.³⁷ Alternatively, OA is known to reduce the activity of SREBP-1c,^{10,11} a regulator of DNL enzymes, including ACC, that is sufficient and necessary for DNL in hepatocytes.^{38,39} In adipocytes, however, SREBP-1c knockout does not significantly decrease expression of lipogenic enzymes⁴⁰ and overall SREBP-1c seems to be a much smaller player for adipocyte DNL.^{39,41} We propose that OA may specifically regulate SREBP-1c and therefore hepatocyte ACC. This explains its dramatic effect on hepatocyte LDs, but not adipocyte. Further research is needed to fully disentangle the relationship between fatty acid scavenging and DNL, especially across cell lines.

This work uniquely highlights the ability of OPTIR not only to observe multiplexed vibrational probes in living cells, but also provide insight on competing metabolic pathways. We find that OA feeding at 60 μ M decreases the rate of glucose-derived *de novo* synthesized lipid storage in lipid droplets in hepatocytes but not adipocytes. We propose this is due to differences in regulation of DNL between the cell lines.

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Conflicts of interest

There are no conflicts to declare.

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