High Glucose Induces Lipid Accumulation via 25-Hydroxycholesterol DNA-CpG Methylation

HIGHLIGHTS
- High glucose induces lipids accumulation via epigenetic regulation
- High glucose induces lipids accumulation via DNA CpG methylation in promoter regions
- High glucose induces lipid accumulation by inhibiting multiple signaling pathways
- 25-Hydroxycholesterol is an endogenous agonist of DNMT1, an epigenetic regulator

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High Glucose Induces Lipid Accumulation via 25-Hydroxycholesterol DNA-CpG Methylation

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SUMMARY
This work investigates the relationship between high-glucose (HG) culture, CpG methylation of genes involved in cell signaling pathways, and the regulation of carbohydrate and lipid metabolism in hepatocytes. The results indicate that HG leads to an increase in nuclear 25-hydroxycholesterol (25HC), which specifically activates DNA methyltransferase-1 (DNMT1), and regulates gene expression involved in intracellular lipid metabolism. The results show significant increases in 5mCpG levels in at least 2,225 genes involved in 57 signaling pathways. The hypermethylated genes directly involved in carbohydrate and lipid metabolism are of PI3K, cAMP, insulin, insulin secretion, diabetic, and NAFLD signaling pathways. The studies indicate a close relationship between the increase in nuclear 25HC levels and activation of DNMT1, which may regulate lipid metabolism via DNA CpG methylation. Our results indicate an epigenetic regulation of hepatic cell metabolism that has relevance to some common diseases such as non-alcoholic fatty liver disease and metabolic syndrome.

INTRODUCTION
A high-glucose (HG) diet has been associated with metabolic syndrome (Moreno-Fernandez et al., 2018; Zhao et al., 2016), non-alcoholic fatty liver disease (NAFLD) (Liu et al., 2018), type II diabetes (Menegazzo et al., 2015; Vallon and Thomson, 2017), and cardiovascular disease (Che et al., 2018). A HG diet can induce fatty acid accumulation in liver, elevate insulin resistance index, cause oxidative stress, and increase body weight. This has led to the association of a HG diet with the development of the metabolic syndrome (Du et al., 2010). Hepatocytes play an important role in metabolizing carbohydrates and lipids in the body. Their function changes depending on the circulating levels of insulin and the sensitivity of the insulin receptor. The inability of insulin to suppress hepatic glucose production is a key defect found in metabolic syndrome (Balland et al., 2019; O-Sullivan et al., 2015). The role of insulin has been well documented. Hepatocytes respond to elevated glucose levels by packaging it into glycogen. When carbohydrates are in excess, hepatocytes can convert this excess energy into lipids. When glucose levels drop and insulin production falls, the shortage of insulin in the blood signals the hepatocytes to utilize their storage by hydrolyzing glycogen and lipids. Dysregulation of this process can result in lipid accumulation in hepatocytes, leading to the development of metabolic disorders, including NAFLD and type II diabetes (Saltiel, 2016). HG in culture media induces lipid accumulation in hepatocytes, which has been used as an in vitro model for the study of NAFLD (Swaminathan et al., 2013). The detailed biochemical mechanisms that allow for HG to produce lipid accumulation are not fully understood.

Epigenetic modification plays a major role in the interpretation of genetic information. Methylation at position 5 of cytosine (5-methylcytosine, 5mC) in DNA is an important epigenetic modification that regulates gene expression and other functions of the genome (Moore et al., 2013). Cytosine methylation in CpG is inversely correlated with transcriptional activity of associated genes as it causes chromatin condensation and thus gene silencing (Yu et al., 2014). Recently published literature has demonstrated that dysregulation of gene expression is important in metabolism and can affect tissue function and in turn the metabolic state (Castellano-Castillo and Moreno-Indias, 2019). It has been shown that HG levels affected gene expression through DNA methylation (Wang et al., 2018). However, the molecular mechanisms by which HG levels lead
to DNA methylation, the changes of gene expression patterns, and lipid accumulation in human hepatocytes are not well understood.

Oxysterols are oxidized products of cholesterol. They are short-lived but can be potent biologically active molecules involved in a plethora of functions including lipid metabolism and inflammatory responses (Cy- ster et al., 2014; McDonald and Russell, 2010). Oxysterols have long been known for their important role in cholesterol homeostasis, where they are involved in both transcriptional and posttranscriptional mechanisms in controlling cholesterol levels (Luu et al., 2016). Recent research with oxysterols has demonstrated their novel and sometimes surprising role associated with a wide variety of cellular functions (Vurusaner et al., 2016). They act as ligands for a growing list of receptors, including some that are of importance to the immune system and lipid metabolism. Oxysterols have also been implicated in several diseases, such as NAFLD and other metabolic syndromes. However, until now their specific binding sites and mechanism of action remain unclear. The objective of this study is to investigate the role of a specific oxysterol (25-hydroxycholesterol [25HC]) in the epigenetic regulation associated with HG induction of lipid accumulation in human hepatocytes.

RESULTS

HG Induces Intracellular Lipid Accumulation by Increasing Gene Expression Involved in Lipid Biosynthesis in Hepatocytes

To study the effect of HG on intracellular lipid accumulation, Huh-7 cells were cultured in media with HG for different time periods and total neutral lipids were estimated by oil red O staining. As shown in Figure 1A, lipid levels started to increase at 36 h and had significantly accumulated at 72 h. HG increases expression and activation of SREBP-1C. Biosynthesis of free fatty acids and triglycerides is regulated by SREBP-1C (Horton et al., 2003). To investigate how HG increases lipid biosynthesis, total RNA was isolated from the HG-cultured Huh-7 cells. The mRNA levels of SREBP-1C were determined by real time RT-PCR. HG increased RNA levels in a time-dependent manner (Figure 1B). To confirm the results, SREBP-1C proteins were analyzed by western blot. As expected, increases in SREBP-1C protein levels (p < 0.01) in nuclear fractions were time dependent (Figure 1C). Thus, HG increases intracellular lipid levels by increasing SREBP-1C expression, which is consistent with previous reports (Pang et al., 2018; Xu et al., 2013).

Quantitative analysis showed that HG culture increased intracellular total cholesterol by 1.5-fold (p < 0.05) (Figure 1D), cholesterol ester by 2.9-fold (p < 0.05) (Figure 1E), triglycerides by 2.1-fold (p < 0.01) (Figure 1F), and free fatty acids by 28% (p < 0.01) at 72 h (Figure 1G). It is noted that HG culture did not significantly change free cholesterol levels (total cholesterol minus cholesterol ester). To further study the increased lipids in nuclear fractions, their composition was quantified by lipidomic analysis as shown in Figures 1H–1L and Table S2. Compared with low glucose (LG), HG increased phosphatidylcholine, sphingomyelin, and triglyceride levels with saturated fatty acids in nuclear fractions of Huh-7 cells (Figure 1H and Table S3). Compared with normal human liver tissues, non-alcoholic steatohepatitis (NASH) liver nuclear fraction showed increases mainly in triglycerides with saturated fatty acids (Figure 1I and Table S4). Interestingly, HG increased nuclear 25HC levels, whereas no change was detectable in LG (Figure 1K and Table S3); NASH liver nuclear fraction showed increases in 25HC levels by 30%, whereas decreases in 27-hydroxycholesterol (27HC) by 90% (Figure 1L and Table S4). The results imply that 25HC might play a role in the lipid accumulation.

HG Increases Nuclear Levels of Newly Synthesized Cholesterol, 25-Hydroxycholesterol, and 27-Hydroxycholesterol

Oxysterols play an important role in lipid metabolism (Figure 2A). To examine the nuclear oxysterol levels in hepatocytes, Huh-7 cells were cultured in DMEM media with HG for 72 h. [14C]-Acetate was added to the media, and the cells were cultured for another 3 and 9 h (Figures 2B and 2C). Total lipids were extracted and partially purified. Individual [14C]-acetate derivatives were analyzed by high-performance liquid chromatography (HPLC). HPLC analysis showed that the product has the same retention time as 25HC and 27HC in our HPLC system (Figures 2D–2F), suggesting that the nuclear oxysterol derivative is sulfonated 25HC and 27HC. Compared with cells cultured in LG, the cells cultured in HG exhibited significantly increased levels of [14C]-cholesterol and [14C]-oxysterols, including [14C]-25HC and [14C]-27HC in nuclear fractions. About 50% of the total [14C]-25HC was detected in the nuclear fraction (Figures 2G and 2H). We also studied the distribution of exogenous [3H]-25HC and found that similar percentilte of the [3H]-25HC will go to nuclei in 1 h (data not shown). As oxysterols have been reported to increase SREBP-1C
Figure 1. The Effects of HG on Lipid Accumulation in Hepatocytes

(A–L) Huh-7 cells were treated in HG media for 0, 12, 24, 48, and 72 h. The total intracellular neutral lipids were stained with 0.2% oil red O (A); inserts are shown at 400 x magnification of the boxed areas (scale bar, 10 µm); mRNA levels of SREBP-1C in HG-cultured hepatocytes were determined by real-time RT-PCR analysis (B). SREBP-1C mature form protein levels in the nuclear fraction were analyzed by western blot (C); total lipids were extracted, and individual lipids were biochemically analyzed. Hepatic total cholesterol (D), cholesterol ester (E), triglyceride (F), and free fatty acid (G). Each individual level was normalized by protein concentration. Nuclear lipid composition from hepatocytes and human liver tissues was determined by lipidomic analysis. The lipid profile in nuclear fraction from Huh-7 cells cultured in HG or LG media (HG minus LG) (H); between human normal and NASH liver tissues (NASH minus normal) (I); 25HC as standard (J); the different levels of nuclear 25HC in Huh-7 cells cultured in HG and LG (K); the different levels of nuclear 25HC in human normal and NASH liver tissue (L). The lipidomic profile represents results from one of two experiments. All values are expressed as the mean ± SD.
expression and 25HC is one of the most potent regulatory oxysterols (Spann and Glass, 2013; Xu et al., 2010), cholesterol, 25HC, and 27HC were selected for further study to determine which molecule serves as an epigenetic regulator and plays a key role in lipid accumulation.

25HC Serves as an Agonist of DNMT1

Epigenetic regulators are DNA methyltransferases/demethylases and acetyltransferases/deacetylases. To study the effects of nuclear sterols on epigenetic regulation, 12 recombinant epigenetic regulators, DNA methyltransferase-1 (DNMT1), DNMT3a, DNMT3b, GCN3 (Giant congenital nevi), p300 (histone acetyltransferase), Pcaf (KAT2B lysine acetyltransferase 2B), HDAC1 (histone deacetylase 1), HDAC2 (histone deacetylase 2), HDAC3 (histone deacetylase 3), HDAC6 (histone deacetylase 6), HDAC10 (histone deacetylase 10), and KDM6B-JMJD3 (lysine demethylase 6B), were used to determine whether 25HC or cholesterol
serves as their endogenous ligand(s). Interestingly, 25HC and 27HC, but not cholesterol, act on only DNMT1, and not on DNMT3a/3b (Figures 3A and 3B) and did not affect any demethylases, acetyltransferases, or deacetylases (data not shown). The kinetic study showed that 25HC increased DNMT1 by 8-fold at 3.7 μM, whereas 27HC reached the level at 100 μM, which is not physiologically significant, although both are in a dose-dependent manner. Then negative control S-adenosyl homocysteine (SAH) at 1 mM inhibited DNMT1 activity by 90% as shown in Figure 3C. Cholesterol did not have any effect on the DNMTs. The results indicate that 25HC is much more potent than 27HC (~30-fold). It is reasonable to believe that

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**Figure 3. Effects of HG and 25HC on DNMT1 Expression and Enzyme Kinetics**

(A–G) Effects of 25HC on DNMT1/3a/3b activities were determined by enzyme kinetics. The effects of different concentration on the enzymatic activities were tested from 0 to 0.001 M (A); effects of cholesterol, 25HC, 27HC, and S-adenosyl homocysteine (SAH, inhibitor as a negative control) on the DNMT1 activity (plots versus linear) (B); and plots versus logarithm (C). Effects of HG and 25HC on the expression of DNMT1 were determined by real-time RT-PCR. Huh-7 cells were cultured in HG media for 0, 12, 24, 48, and 72 h; mRNA levels of DNMT1 were determined by real-time RT-PCR analysis (D). Huh-7 cells were treated with 25 μM 25HC for 3, 6, and 12 h. The mRNA levels of DNMT1 were determined by real-time RT-PCR analysis (E). Protein levels of DNMT1 in the nuclei were analyzed by western blot (F). Effects of HG and 25HC on the global methylation (G). After Huh-7 cells were cultured in HG for 72 h, or treated with 25 μM of 25HC for 4 h, total DNA was purified as described in Methods. The levels of methylation were estimated by LINE-1 assay. Four CpG sites in promoter region of LINE-1 element were chosen as the target position. Data are represented as mean ± SD and statistic by t test. *Significant difference in expression between LG and HG (p < 0.05).
25HC, but not 27HC, is an endogenous ligand of DNMT1, which can methylate DNA CpG and subsequently regulate gene expression. In addition, HG also increased expression of DNMT1 in hepatocytes. The RT-PCR and western blot results showed that HG and 25HC led to a time-dependent increase of DNMT1 both in mRNA and protein levels as shown in Figures 3D–3F.

Recent studies have shown that global DNA methylation (long interspersed nucleotide element 1 [LINE-1] assay) and the methylation of specific genes are related to adipogenesis (Malodobra-Mazur et al., 2019), lipid metabolism (Chen et al., 2019; Li et al., 2020), and inflammation in visceral adipose tissues (Barajas-Olmos et al., 2018; Castellano-Castillo et al., 2018), which are in turn related to the etiology of metabolic syndrome. To study the effects of 25HC on DNA cytosine methylation in the promoter region, LINE-1 analysis was first performed to estimate CpG methylation in promoter regions. Methylation usually occurs in repetitive elements, such as LINE elements. There are 500,000 LINE elements and 40 trillion copies in total. Human LINE-1 is a retrotransposable region (promoter region) and has only 700,000 copies, which correlates to ~17% of the human genome (Lander et al., 2001). The specific sequence includes four CpG dinucleotides (Pos 1, 2, 3, and 4) as methylation targets in LINE-1 (Figure 3G). The results showed that HG significantly increases CpG methylation of LINE-1 elements in Huh-7 cells. Following culture in HG for 72 h, increases in methylation of 11% at Pos 3, and 8% at Pos 4, were observed. The results indicate that HG increases CpG methylation in promoter regions.

Profiles of Whole Genome-wide DNA Methylation in HG- and LG-Treated Huh-7 Cells

To understand the possible function of cytosine methylation in HG-treated Huh-7 cells, the cells were harvested for the construction of bisulfite-treated genomic DNA libraries. In total whole-genome bisulfite sequencing (WGBS) generated 401 million (LG) and 359 million (HG) raw reads from the two libraries by paired-end sequencing, respectively. Of the 394 million clean reads from LG library, 77% (305 million) were uniquely mapped to the reference genome of “human reference genome (hg38),” whereas of 354 million clean reads from HG library, 79% (279 million) were uniquely mapped to the reference genome, exhibiting an average read depth of 22 and 20, respectively. More than 82% cytosines were covered by at least 10 reads in the reference genome. The depth and density of the sequencing were enough for a high-quality genome-wide methylation analysis. Meanwhile, the bisulfite conversion efficiencies represented by the lambda DNA added into the libraries were over 99%, providing reliable and accurate results for the WGBS (Table S5).

DNA methylation levels in whole genome and differentially methylated regions (DMRs) are shown in Figure 4A. To present the global DNA methylation profiles of the two libraries, the uneven methylation levels throughout the chromosomes under CG, CHG (H represents adenosine or thymidine), and CHH contexts are shown in Figure 4B. A total of 6,012 differentially methylated genes (DMGs) were screened out among two libraries; 2,591 DMGs were identified under CG context, 862 under CHG context, and 2,559 under CHH context. In these DMGs, 277 were identified under CG and CHG contexts, 567 under CG and CHH contexts, 1,088 under CHG and CHH contexts, and only 516 under CG, CHG, and CHH contexts. Furthermore, 3,549 were identified as promoter region different, and 2,225 were identified under CG context, 468 under CHG context, and 856 under CHH context. For these DMGs, 169 were identified under CG and CHG contexts, 88 under CG and CHH contexts, 283 under CHG and CHH contexts, and only 89 identified under CG, CHG, and CHH contexts.

DNA methylation levels generally show a varied distribution across different functional regions of the genome. The methylation levels in the CGI (CG island), CGI-shore (up to 2 kbp away from the CGI), promoter (upstream 2-kb sequence from transcription starting site), 5’ UTR, exon, intron, 3’ UTR, and repeat were significantly different between HG- and LG-treated groups. It is interesting that HG has significantly higher hypermethylation levels than LG in CpG positions only (Figure 4C). In total 60,633 DMRs were identified; 4,711 (2,832 hypermethylated and 1,879 hypomethylated) were distributed in CGI, 4,986 (2,727 hypermethylated and 2,259 hypomethylated) in CGI-shore, 5,105 (2,724 hypermethylated and 2,381 hypomethylated) in exon, 15,291 (7,198 hypermethylated and 8,093 hypomethylated) in intron, 5,312 (2,754 hypermethylated and 2,558 hypomethylated) in promoter, 21,838 (13,023 hypermethylated and 8,815 hypomethylated) in repeat region, 210 (131 hypermethylated and 79 hypomethylated) in transcription start site elements, 811 (529 hypermethylated and 282 hypomethylated) in transcription start site (TSS) elements, 757 (350 hypermethylated and 407 hypomethylated) in 3’ UTRs, and 1,612 (916 hypermethylated and 696 hypomethylated) in 5’ UTRs. In almost all DMRs, CpGs are significantly more hypermethylated than...
hypomethylated. It has been reported that CG methylation in promoter regions plays a key role in silencing gene expression (Baylin, 2005).

Total DMGs, 6,012 genes, were significantly enriched in 288 Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways (189 hypermethylated and 129 hypomethylated). DMGs in promoter regions were significantly enriched in 169 KEGG pathways (83 hypermethylated and 51 hypomethylated) (Table S5), of which 79 (57 hypermethylated and 5 hypomethylated) were under CG context. Seventy-two DMGs (25

Figure 4. Whole-Genome Bisulfite Sequencing (WGBS) Analysis of the Effects of HG on the DNA Methylation in Hepatocytes
(A–D) Huh-7 cells were cultured in HG or LG media for 72 h. Detailed global methylation was measured by WGBS. Circos maps of DMR distribution in chromosomes (A): the first circle shows the distribution of hypermethylation DMRs, the second circle shows transposable element (TE) density, and the third circle shows the distribution of hypomethylation DMRs. Venn diagrams of DMR-associated genes (DMGs) among LG and HG libraries under CG, CHG, and CHH contexts of whole genome and promoter region (B). DNA methylation levels in different genomic functional regions of the whole genome (C), where the x axis represents the different genomic regions (CGI, CGI-shore, promoter, UTR 5, exon, intron, UTR 3 and repeat) and the y axis represents the methylation levels for LG and HG libraries under CG, CHG, and CHH contexts. Significant enrichment KEGG pathways of DMGs (D).
hypermethylated and 34 hypomethylated) were significantly enriched under CHG context and 18 (1 hypermethylated and 12 hypomethylated) under CHH context.

To analyze the function of DMGs in promoter regions under CG context, 79 KEGG pathways (p < 0.05) were identified. In these pathways 57 were identified as hypermethylation and 5 as hypomethylation. Thirteen of them are related to metabolic syndrome (Figure 4D). Six of these pathways including PI3K-Akt signaling pathway (Table 1), cAMP signaling pathway (Table 2), NAFLD (Table 3), type II diabetes mellitus (Table 4), and insulin secretion (Table 5) are directly related to carbohydrate, lipid, and energy metabolism. The most significant genes involved in these signaling pathways include insulin (INS), retinoid X receptor alpha (RXRa), insulin receptor substrate 1 (IRS1), insulin receptor (INSR), phosphoinositide-3-kinase regulatory subunit 5 (PIK3R5), mitogen-activated protein kinase 1 (MAPK1), cyclin-dependent kinase inhibitor 1B (CDKN1B), protein kinase N2 (PKN2), calcium/calmodulin-dependent protein kinase II beta (CAMK2B), peroxisome proliferator-activated receptor alpha (PPARα), mitogen-activated protein kinase 1 (MAPK1), potassium calcium-activated channel subfamily N member 3 (KCNN3), glucagon-like peptide 1 receptor (GLP1R), potassium calcium-activated channel subfamily U member 1 (KCNU1), potassium calcium-activated channel subfamily N member 4 (KCNN4), and calcium voltage-gated channel subunit alpha1D (CACNA1D). It has been well believed that insulin resistance is the major mechanism in the development and progression of NAFLD/NASH (Engin, 2017). Interestingly, DMRs in the promoter regions of key genes in these signaling pathways are hypermethylated, indicating that the gene expressions are down-regulated, which fits well with the mechanism of NAFLD mechanism.

**Target Next-Generation Bisulfite Sequencing to Confirm the WGBS Results**

To confirm the results from WGBS, the methylation levels of CpG sites in promoter regions of PPARα, INS, CAMK2B, and PIK3R5, key genes in NAFLD, type II diabetes mellitus, insulin secretion, and cAMP signaling pathways, were determined using target next-generation bisulfite sequencing (tNGBS) (Figure 5). The results showed that HG increases the methylation levels of every CpG site not only in promoter region but also in exon and intron regions. For CAMK2B gene (Figure 5A), 24 CpG sites were identified including 8 (Pos 1–8) in promoter region, 4 (Pos 9–12) in exon region, and 12 (Pos 13–24) in intron region. HG increases the methylation levels of Pos 1–8 by 22%, Pos 9–12 by 18%, and Pos 13–24 by 19%. For PIK3R5 gene (Figure 5B), 24 CpG sites were identified including 8 (Pos 1–8) in exon region, 13 (Pos 9–21) in promoter region, and 3 (Pos 22–24) in intron region. HG increases the methylation level of Pos 1–8 by 60% on average, Pos 9–21 by 15%, and Pos 22–24 by 19%. For PPARα gene (Figure 5C), HG increases the methylation levels of 14 CpG sites (Pos 1–14) in promoter region by 27%. For INS gene (Figure 5D), HG increases the methylation levels of 10 CpG sites (Pos 1–10) by 40% on average.

To explore whether DNA methylation in promoter region is involved in down-regulation of gene expression, four multiple function genes were selected for confirmation (Figure 5). Gene expressions were estimated by RT-PCR, and CpG methylations were confirmed by tNGBS. The relationship of gene expression with CpG methylation of CAMK2B is shown in Figures 5A; with PIK3R5, in 5B; with PPARα, in 5C; and with INS, in 5D. The results showed that all these gene expressions were significantly decreased when Huh-7 cells were cultured in HG, which is highly consistent with the results from the tNGBS analysis. The results confirm that the CpG methylation in the genes is responsible for the down-regulation of their expression.

**DISCUSSION**

This study proposes a regulatory pathway for HG to lead to induction of lipid accumulation through 25HC and epigenetic regulation through hypermethylation. HG induces lipid accumulation through increased nuclear 25HC levels. The potent regulatory molecule 25HC, but not 27HC, activates DNMT1, which methylates cytosine of CpG in promoter regions, suppresses their expression, and down-regulates the associated signaling pathways. This proposed mechanism sheds new light on understanding how consuming excess sugar can lead to an increase in lipid biosynthesis, lipid accumulation, and risk of weight gain. These results propose some of the epigenetic mechanisms by which high-sugar diet may lead to insulin resistance and inflammation, both of which are risk factors for metabolic syndrome, through epigenetic regulation.

Sugars, such as fructose and glucose, can be catabolized to acetyl-CoA, which is the major substrate for producing energy, ATP, in vivo. When ATP concentrations reach high levels, excess acetyl-CoA is shunted into synthesis of cholesterol, free fatty acids, and triglycerides as a means of energy storage. The correlation between cholesterol and triglyceride biosynthesis has not been elucidated. In the present report, we demonstrate evidence that...
| Gene Name | DMR Location in Promoter Region | DMR (Methylation %) |
|-----------|---------------------------------|---------------------|
|           | Chromosome | Start | End | HG | LG | HG-LG |
| COL11A1   | Chr1       | 1.03 x 10^8 | 1.03 x 10^8 | 92.12 | 51.66 | 40.46 |
| GNG5P2    | ChrX       | 1.1 x 10^8 | 1.1 x 10^8 | 47.26 | 6.76 | 40.5 |
| KRAS      | Chr12      | 25250819 | 25250967 | 17.42 | 5.25 | 12.18 |
| INS       | Chr11      | 2162594  | 2162810  | 79.13 | 23.5 | 55.62 |
| JAK1      | Chr1       | 64967244 | 64967529 | 67.24 | 15.6 | 51.65 |
| LAMA5     | Chr20      | 62368047 | 62368299 | 31.14 | 8.64 | 22.5 |
| BCL2      | Chr18      | 63320785 | 63321134 | 63.22 | 35.11 | 28.1 |
| HRAS      | Chr11      | 536242   | 537214   | 61.26 | 18.46 | 42.8 |
| FGFR3     | Chr4       | 1792656  | 1792887  | 60.41 | 18.3 | 42.11 |
| CD19      | Chr16      | 28935890 | 28936133 | 88.21 | 47.13 | 41.08 |
| LAMB1     | Chr7       | 1.08 x 10^8 | 1.08 x 10^8 | 71.14 | 24.24 | 46.9 |
| RAC1P2    | Chr4       | 46724260 | 46724573 | 73.99 | 47.16 | 26.82 |
| MAPK1     | Chr22      | 21867333 | 21867621 | 23.71 | 3.56 | 20.15 |
| CDKN1B    | Chr12      | 12714273 | 12715037 | 93.4 | 23.71 | 69.69 |
| PIK3R5    | Chr17      | 8888466  | 8888776  | 89.6 | 36.73 | 52.87 |
| BCL2L11   | Chr2       | 1.11 x 10^8 | 1.11 x 10^8 | 72.75 | 42.65 | 30.1 |
| PDGFB     | Chr22      | 3924292  | 3924277  | 83.66 | 54.37 | 29.29 |
| ANGPT1    | Chr8       | 1.07 x 10^8 | 1.07 x 10^8 | 46.11 | 5.16 | 40.96 |
| IL6       | Chr7       | 22726101 | 22726775 | 80.53 | 30.36 | 50.18 |
| MDM2      | Chr12      | 68806524 | 68806756 | 75.71 | 33.96 | 41.74 |
| INSR      | Chr19      | 7295041  | 7295182  | 57.84 | 14.92 | 42.92 |
| FGF11     | Chr17      | 7437822  | 7438331  | 50.31 | 19.51 | 30.81 |
| LPAR3     | Chr1       | 84894412 | 84894734 | 90.41 | 45.1 | 45.31 |
| FGF12     | Chr3       | 1.93 x 10^8 | 1.93 x 10^8 | 85.93 | 40.52 | 45.41 |
| FGFR2     | Chr10      | 1.22 x 10^8 | 1.22 x 10^8 | 24.32 | 8.01 | 16.3 |
| COL2A1    | Chr12      | 48004804 | 48005182 | 83.53 | 12.91 | 70.62 |
| FGF7P3    | Chr9       | 40106680 | 40106914 | 32.35 | 7.07 | 25.28 |
| CSF3      | Chr17      | 40014561 | 40014935 | 73.05 | 18.59 | 54.46 |
| FOXO3     | Chr6       | 1.09 x 10^8 | 1.09 x 10^8 | 25.28 | 4.11 | 21.17 |
| NR4A1     | Chr12      | 52051741 | 52052037 | 61.95 | 14.22 | 47.73 |
| F2R       | Chr5       | 76715992 | 76716157 | 27.58 | 4.89 | 22.69 |
| PKN2      | Chr1       | 88684561 | 88684789 | 37.22 | 4.97 | 32.25 |
| EIF4E1B   | Chr5       | 1.77 x 10^8 | 1.77 x 10^8 | 42.38 | 11.45 | 30.93 |
| RXRa      | Chr9       | 1.34 x 10^8 | 1.34 x 10^8 | 36.99 | 6.91 | 30.08 |

Table 1. Methylation of DMR in Gene Promoter Regions of PI3K-Akt Signaling Pathway (p = 0.000523)

(Continued on next page)
HG increases nuclear 25HC levels and subsequently increases triglyceride biosynthesis. We also may have explored for the first time that 25HC serves as an activator of DNMT1 and plays an important role in global regulation of metabolic pathways, cell survivals, and cell death by regulating critical signaling pathways. It has been

| Gene Name | DMR Location in Promoter Region | DMR (Methylation %) |
|-----------|---------------------------------|---------------------|
|           | Chromosome | Start | End     | HG | LG | HG-LG |
| AC113189.4| Chr17       | 7437822| 7438331 | 50.31 | 19.51 | 30.81 |
| CSF3R     | Chr1        | 36482185| 36482512 | 90.94 | 41.71 | 49.23 |
| IRS1      | Chr2        | 2.27 × 10^8| 2.27 × 10^8 | 34.01 | 5.95 | 28.06 |
| PDGFA     | Chr7        | 521434 | 521719 | 83.23 | 59.88 | 23.35 |

Table 1. Continued

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| Gene Name | DMR Location in Promoter Region | DMR (Methylation %) |
|-----------|---------------------------------|---------------------|
|           | Chromosome | Start | End     | HG | LG | HG-LG |
| PDE4C     | Chr19      | 18220846| 18221310 | 85.7 | 24.91 | 60.79 |
| PPARγ     | Chr22      | 46149487| 46149878 | 94.42 | 59.47 | 34.95 |
| HCAR2     | Chr12      | 1.23 × 10^8| 1.23 × 10^8 | 87.48 | 45.91 | 41.57 |
| GNAS      | Chr20      | 58891431| 58891578 | 35.11 | 5.71 | 29.4 |
| CACNA1D   | Chr3       | 53493469| 53494019 | 74.84 | 25.81 | 49.03 |
| CAMK2B    | Chr7       | 44221409| 44221698 | 86.07 | 45  | 41.07 |
| RAC1P2    | Chr4       | 46724260| 46724573 | 73.99 | 47.16 | 26.82 |
| PLD2      | Chr17      | 4806842 | 4806937 | 18.7 | 5.64 | 13.06 |
| MAPK1     | Chr22      | 21867333| 21867621 | 23.71 | 3.56 | 20.15 |
| PIK3R5    | Chr17      | 8888466 | 8888776 | 89.6 | 36.73 | 52.87 |
| GLP1R     | Chr6       | 39048762| 39048955 | 86.25 | 59.95 | 26.3 |
| ADCY1     | Chr7       | 45574683| 45574877 | 59.55 | 33.76 | 25.79 |
| PDE4B     | Chr1       | 66331798| 66332539 | 86.73 | 37.91 | 48.82 |
| NFATC1    | Chr18      | 79399054| 79399517 | 78.35 | 33.06 | 45.29 |
| PDE4D     | Chr5       | 59215742| 59215941 | 94.05 | 54.39 | 39.66 |
| VAV3      | Chr1       | 1.08 × 10^8| 1.08 × 10^8 | 88.49 | 24.15 | 64.34 |
| F2R       | Chr5       | 76715992| 76716157 | 27.58 | 4.89 | 22.69 |
| ADCY4     | Chr14      | 24334404| 24334719 | 86.49 | 53.89 | 32.6 |
| HCN2      | Chr19      | 587907  | 588099  | 56.37 | 20.45 | 35.92 |
| GIPR      | Chr19      | 45669075| 45669744 | 83.03 | 39.41 | 43.62 |
| HTR4      | Chr5       | 1.49 × 10^8| 1.49 × 10^8 | 95.66 | 78.7 | 16.96 |
| ALS9063S.1| Chr6       | 91816380| 91819294 | 93.85 | 28.92 | 64.92 |
| GRIN1     | Chr9       | 1.37 × 10^8| 1.37 × 10^8 | 37.91 | 13.55 | 24.36 |
| PPAR1B    | Chr17      | 39630003| 39630266 | 65.33 | 17.07 | 48.25 |
| NPR1      | Chr1       | 1.54 × 10^8| 1.54 × 10^8 | 76.11 | 53.96 | 22.15 |

Table 2. Methylation of DMR in Gene Promoter Regions of cAMP Signaling Pathway (p = 0.001699)
reported that 25HC as a ligand of liver X receptor (LXRs) plays an important role in the control of lipid metabolism
(Janowski et al., 1996). LXR ligands up-regulate expression of cholesterol reverse transporters such as ABCA1 and
ABCG1 via activation of LXR/RXR heterodimers. ABCA1/G1 is known to mediate efflux of cellular cholesterol and
phospholipids, which is the target for therapy of anti-atherosclerosis (Zhu et al., 2012). Unfortunately, administra-
tion of the synthetic ligands to mice triggers induction of the lipogenic pathway and elevates plasma triglyceride
levels (Schultz et al., 2000). Addition of 25HC to human hepatocytes increases gene expression of key enzymes,
ACC and FAS, in lipid biosynthesis and increases intracellular lipids levels via SREBP-1C pathway (Ma et al., 2008).
25HC has also been described to function in immune system, suppressing immunoglobulin production in B cells
(Bauman et al., 2009) and inducing expression of the inflammatory cytokine interleukin-8 (Wang et al., 2004).
Recently, 25HC has been shown to directly restrict target cell entry of enveloped viruses by inhibiting fusion of
virus and cell membranes (Doms et al., 2018; Ke et al., 2017; Li et al., 2017; Liu et al., 2013; Wang et al., 2017).
More interestingly, it has been reported that 25HC can be sulfated to 25-hydroxycholesterol 3-sulfate,
25HC3S, which has been shown as a distinct yet potent regulator of cellular functions (Li et al., 2007; Ren et al.,
2007). Both 25HC3S and 25HC play important roles in lipid metabolism, inflammatory responses, and cell survival
but act in a direction opposite to each other (Ren and Ning, 2014). The observation of global regulation is consis-
tent with the present conclusion that 25HC serves as an agonist of epigenetic regulator, DNMT1. Whether the
sulfation of 25HC to 25HC3S is the fundamental regulatory mechanism needs to be further investigated.

It is also interesting to notice that HG hyper-methylates CpG but hypomethylates CHG and CHH in whole
gene regions including CpG island, CpG island shore, promoter, and TSS region as shown in Figure 4C. It is
well-believed that CpG methylation in promoter regions is directly related with silencing gene expression

| Gene Name | DMR Location in Promoter Regions | DMR (Methylation %) |
|-----------|----------------------------------|---------------------|
| RAC1P2    | Chr4 46724260-46724573           | HG: 73.99, LG: 47.16, HG-LG: 26.82 |
| MTCYBP22  | Chr5 1 × 10⁶-1 × 10⁶             | HG: 90.01, LG: 52.51, HG-LG: 37.50 |
| RXRα      | Chr9 1.34 × 10⁶-1.34 × 10⁶        | HG: 36.99, LG: 6.91, HG-LG: 30.08 |
| MTCO3P2   | Chr5 1 × 10⁶-1 × 10⁶             | HG: 88.87, LG: 45.71, HG-LG: 43.16 |
| MTCO1P46  | Chr2 2.12 × 10⁶-2.12 × 10⁶        | HG: 83.83, LG: 26.28, HG-LG: 57.54 |
| MTCO2P2   | Chr5 1 × 10⁶-1 × 10⁶             | HG: 88.87, LG: 45.71, HG-LG: 43.16 |
| INSR      | Chr19 7295041-7295182            | HG: 57.84, LG: 14.92, HG-LG: 42.92 |
| AC234779.1| Chrx 1.41 × 10⁶-1.41 × 10⁶       | HG: 89.36, LG: 19.05, HG-LG: 70.31 |
| COX6CP2   | Chr20 5047826-50478554           | HG: 81.66, LG: 43.62, HG-LG: 38.04 |
| BID       | Chr22 17774417-17774536          | HG: 20.06, LG: 5.89, HG-LG: 14.17 |
| MAP3K5    | Chr6 1.37 × 10⁶-1.37 × 10⁶        | HG: 81.00, LG: 40.19, HG-LG: 40.81 |
| PIK3R5    | Chr17 8888466-8888776            | HG: 89.60, LG: 36.73, HG-LG: 52.87 |
| BCL2L11   | Chr2 1.11 × 10⁶-1.11 × 10⁶        | HG: 72.75, LG: 42.65, HG-LG: 30.10 |
| COX6B2    | Chr19 55354472-55354665          | HG: 85.45, LG: 65.5, HG-LG: 19.95 |
| PPARα     | Chr22 46149487-46149878          | HG: 94.42, LG: 59.47, HG-LG: 34.95 |
| INS       | Chr11 2162594-2162810            | HG: 79.13, LG: 23.5, HG-LG: 55.62 |
| IL6       | Chr7 22726101-22727675           | HG: 80.53, LG: 30.36, HG-LG: 50.18 |
| IRS1      | Chr2 2.27 × 10⁶-2.27 × 10⁶        | HG: 34.01, LG: 5.95, HG-LG: 28.06 |
| DDI1T3    | Chr12 57521695-57521932          | HG: 68.03, LG: 14.01, HG-LG: 54.02 |
| NDUFA4L2  | Chr12 57240882-57241348          | HG: 72.33, LG: 37.01, HG-LG: 35.32 |

Table 3. Methylation of DMR in Gene Promoter Regions of Non-alcoholic Fatty Liver Disease (NAFLD) (p = 0.003206)
However, the physiological significance of hypomethylation of CHG and CHH is unknown. DNA methylation is established and maintained by three essential DNA methyltransferases, DNMT1, DNMT3a, and DNMT3b, in mammals (Koerner and Barlow, 2010; Long et al., 2017). Methylated DNA is then recognized by methyl CpG-binding proteins along with associated co-repressors that lead to silencing of the associated promoter (Clouaire and Stancheva, 2008). The in vivo role of DNMT1 in carcinogenesis has been widely studied using hypomorphic mice instead of DNMT1−/− mice that die during embryogenesis (Subramaniam et al., 2014). DNMT1 hypomorphic mice exhibit resistance to alcohol-induced hepatic steatosis compared with the wild-type mice (Kutay et al., 2012). It has been reported that metabolic syndrome has an impact on the global methylation pattern in visceral adipose tissue (Castellano-Castillo and Moreno-Indias, 2019). The results indicate that the steatosis is most likely dependent on the DNA methylation in related genes, which silences gene expression. In this study, we report that HG incubation induces DNA CpG methylation in promoter regions of the key signaling pathways, which induces lipid accumulation in human hepatocytes. We found that HG incubation increases 25HC levels in hepatocyte nuclei. 25HC activates DNMT1 and increases 5mCpG in promoter regions of 2,225 genes.

Table 4. Methylation of DMR in Gene Promoter Regions of Type II Diabetes Mellitus (p = 0.032118)

| Gene Name | DMR Location in Promoter Regions | DMR (Methylation %) |
|-----------|----------------------------------|---------------------|
|           | Chromosome Start End HG LG HG-LG |
| INSR      | Chr19 7295041 7295182 57.84 14.92 42.92 |
| INS       | Chr11 2162594 2162810 79.13 23.5 55.62 |
| SOCS2     | Chr12 93572989 93573585 87.30 53.22 34.08 |
| MAPK1     | Chr22 21867333 21867621 23.71 3.56 20.15 |
| HK1       | Chr10 69318270 69318400 63.47 43.17 20.30 |
| PIK3R5    | Chr17 8888466 8888776 89.60 36.73 52.87 |
| CACNA1D   | Chr3 53493469 53494019 74.84 25.81 49.03 |
| IRS1      | Chr2 2.27 × 10^8 2.27 × 10^8 34.01 5.95 28.06 |

Furthermore, the HG-induced CpG methylation in promoter regions covers 79 KEGG pathways. Fifty-seven pathways were identified as significantly hypermethylated. To our knowledge, 13 of these pathways are directly related to lipid metabolism, including metabolic signaling pathway, Ras signaling pathway

Table 5. Methylation of DMR in Gene Promoter Regions of Insulin Secretion (p = 0.035573)

| Gene Name | DMR Location in Promoter Regions | DMR (Methylation %) |
|-----------|----------------------------------|---------------------|
|           | Chromosome Start End HG LG HG-LG |
| ADCY4     | Chr14 24334404 24334719 86.49 53.89 32.6 |
| INS       | Chr11 2162594 2162810 79.13 23.5 55.62 |
| KCNN3     | Chr1 1.55 × 10^8 1.55 × 10^8 83.22 46.68 36.54 |
| CHRM3     | Chr1 2.4 × 10^8 2.4 × 10^8 58.07 21.85 36.22 |
| GLP1R     | Chr6 39048762 39048955 86.25 59.95 26.3 |
| KCNU1     | Chr8 36928346 36930601 64.67 11.97 52.7 |
| KCNN4     | Chr19 43774123 43774347 31.44 10.69 20.76 |
| GNAS      | Chr20 58888560 58888756 41.48 18.28 23.2 |
| CACNA1D   | Chr3 53493469 53494019 74.84 25.81 49.03 |
| ADCY1     | Chr7 45574683 45574877 59.55 33.76 25.79 |
| CAMK2B    | Chr7 44221409 44221698 86.07 45 41.07 |
Figure 5. The Relationship between DNA CpG Methylation and Gene Expression

(A–D) Huh-7 cells were cultured in HG or LG media for 72 h, and the methylation levels of the CpG sites of key genes involved in signaling pathways were analyzed by tNGBS. Huh-7 cells were cultured in HG media for 12 h, and the mRNA levels of these genes were determined by real-time RT-PCR analysis. The CpG methylation of the CAMK2B gene is shown in (A), where the top image shows the mRNA levels, the middle shows the methylation levels of 5' upstream (Pos 1–14) region, and the bottom shows the sequence. CpG methylation of the PIK3R5 gene is shown in (B), where the top image shows the mRNA levels; the middle, the methylation levels of 5' upstream (Pos 1–13), intron (Pos 14–16), and exon (Pos 17–24) regions; and the bottom shows the gene sequence. CpG methylation of the PPARα gene is shown in (C), where the top shows the mRNA levels; the middle, the methylation levels of 5' upstream (Pos 1–8), exon (Pos 9–12), and intron (Pos 13–24) regions; and the bottom, the gene sequence. CpG methylation of the INS gene is shown in (D), where the top image shows the mRNA levels; the middle, the methylation levels of 5' UTR (Pos 1–2) and 5' upstream (Pos 3–10) regions; and the bottom, the gene sequence. Data are represented as mean ± SD and statistic by t test. *Significant difference in expression between LG and HG (p < 0.05).
As hypermethylation in promoter regions is believed to silence the gene expression, it is reasonable to conclude that HG induces metabolic disorders via methylation of their key genes in the pathways, which were confirmed by analysis of mRNA levels as shown in Figure 5. Only one (p < 0.05) tight junction pathway was identified as hypomethylation. However, its physiological significance is unknown.

The current results most likely represent the etiology for the development of high sugar diet-induced fatty liver diseases. Based on the results, we summarize the regulatory pathway, which may play an important role in the occurrence of NAFLD and metabolic syndrome. When cells are incubated in HG media, high sugar consumption will increase intracellular 25HC, which in turn will activate DNMT1, as shown in Figure 6. Limitations of the Study

We are investigating the effect of knockout and knockdown of 25HC syntheses on the activity of DNA methyltransferases and considering this as a limitation of this study. The enzyme kinetic results provide strong advance of this question. All the materials are available to all interested scientists per request. Subsection lead contact to Yaping Wang.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101102.
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AUTHOR CONTRIBUTIONS

Y.W. designed and conducted the experiments and wrote the paper; S.R. designed experiments, wrote the paper, and provided response to the project; and L.C., W.M.P., D.H., and P.B.H gave a suggestion and edited the manuscript.

DECLARATION OF INTERESTS

S.R. and Virginia Commonwealth University obtain license-related payments from DURECT Corporation, Cupertino, CA.

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Supplemental Information

High Glucose Induces Lipid Accumulation via 25-Hydroxycholesterol DNA-CpG Methylation

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### Table S1. Primer Sequence for Real-time Polymerase Chain Reaction, Related to Figure 1, Figure 3 and Figure 5.

| Gene name  | Forward primer         | Reverse primer         |
|------------|------------------------|------------------------|
| SREBP1     | CAGCCCCACTTCATCAAGG    | ACTGTTGCAAGATCAAGG    |
| DNMT1      | TACCTGGACGACCTGACCTC   | CGTTGGCATCAAAGATGGACA |
| INS        | CATATCCAGCTACTCCCAAAG  | ACAAGTCTGAATGCTCCACT  |
| CAMK2B     | GTCCACCCGGCCCTC        | TTTTGGTGCATTTCTGCTGGG |
| PPARG      | TCGACTCAAGCGTGGTG      | TTCTGAGAGGATGACCC     |
| PIK3R5     | TCGCAAGGAAGAAGGAGATCCCTC | GTCTTCAATATTTGGTGCTGGATG |
| β-actin    | CATGTACGTTGCAATCCAGGC  | TCTCTAATGTCACGCAAGC   |
| GAPDH      | CAATGACCCCTTCATTGACC   | TTGATTTTGAGGGATCTCG   |

### Table S3. Fold Change of Nuclear Lipid Levels Between HG and LG Cultured Huh-7 Cells, Related to Figure 1.

| Lipid name       | Folds  | Lipid name       | Folds  |
|------------------|--------|------------------|--------|
| Cer (d26:1)      | 53.44  | PE (18:1e_22:4)  | 4.75   |
| Cer (d18:2_18:0) | 7      | PE (18:1_22:6)   | 5.08   |
| Cer (m18:1_23:0) | 4.95   | PG (38:1e)       | 6.24   |
| ChE ()           | 55.57  | PI (16:0_18:1)   | 199.62 |
| DG (14:0_18:2)   | 63.45  | PI (18:0_18:2)   | 40.53  |
| DG (16:0_18:2)   | 5.01   | PIP (42:9e)      | 10.35  |
| DG (18:3_18:2)   | 6.05   | PS (16:0_18:2)   | 20.32  |
| Hex1Cer (d18:1_16:0) | 21.52 | PS (16:0_20:4) | 10.97 |
| Hex1Cer (d18:1_24:1) | 250.23 | PS (18:0_18:2) | 29.64 |
| LPE (18:1)       | 6.59   | SM (d34:2)       | 19.37  |
| LPE (22:6)       | 5660.37| TG (16:0_9:0_18:2)| 10.26 |
| PC (30:0)        | 17.06  | TG (16:1_13:0_14:0)| 10.93 |
| PC (16:1_16:1)   | 12.05  | TG (16:1_10:0_18:1)| 31.37 |
| PC (37:3e)       | 6.23   | TG (10:0_18:2_18:2)| 57.11 |
| PC (18:1_20:4)   | 5.33   | TG (16:0_14:4_18:3)| 41.59 |
| PC (38:5)        | 7.67   | TG (16:1_14:1_18:2)| 15.98 |
| PC (18:0_20:2)   | 22.1   | TG (16:0_18:1_18:1)| 19.06 |
| PC (38:1)        | 12.73  | TG (18:1_18:1_18:1)| 49.69 |
| PC (40:5)        | 4.51   | TG (16:0_20:2_20:4)| 12.23 |
| PE (16:0p_18:1)  | 18.93  | TG (18:0_18:1_20:5)| 49.7   |
| PE (18:0_16:0)   | 5.41   | SM (d41:2)       | 5.54   |
| PE (18:1_18:2)   | 5.58   | SPH (d18:1)      | 2.15   |
| PE (16:0_20:3)   | 5.31   | ZyE ()           | 87.39  |
| PE (18:1_18:1)   | 81.3   | 25HC             | 0.37   |
| PE (18:1e_20:4)  | 8.49   | 27HC             | -0.91  |
| PE (18:1e_22:5)  | 94.3   |                  |        |
Table S 4. Fold Change of Nuclear Lipid Levels Between Human NASH and Normal Liver Tissues, Related to Figure 1

| Lipid name                  | Folds | Lipid name                  | Folds |
|-----------------------------|-------|-----------------------------|-------|
| AEA (15:1)                  | 3.55  | PG (18:1_18:2)              | 5.8   |
| CL (18:1_16:1_16:1_18:1)    | 2.75  | PG (28:1_16:0)              | 102.71|
| CL (20:4_16:0_18:0_18:1)    | 9.46  | PG (38:1e)                  | 80.39 |
| Cer (t20:0_23:3)             | 2.11  | PG (18:1_18:1)              | 3.94  |
| Cer (d18:1_23:3)             | 2.54  | PG (18:1_18:2)              | 2.26  |
| Cer (m18:0_24:0)             | 17.35 | PS (18:0_22:5)              | 2.09  |
| ChE ()                      | 7.66  | SM (d40:1)                  | 2.73  |
| ChE ()                      | 42.3  | SM (d41:1)                  | 2.29  |
| CerE (30:6)                 | 2.8   | SM (d42:3)                  | 5.61  |
| CerE (20:4)                 | 2.03  | TG (16:0_18:1_18:1)         | 6.16  |
| CerE (10:0)                 | 15.94 | TG (16:0_18:1_18:2)         | 8.15  |
| Co (Q10)                    | 2.42  | TG (16:0_18:1_18:3)         | 32.11 |
| Co (Q10)                    | 9.25  | TG (18:1_18:1_18:1)         | 10.35 |
| Co (Q9)                     | 2.34  | TG (16:1_16:1_18:1)         | 29.59 |
| DG (16:0_18:1)              | 6.9   | TG (16:1_18:1_18:2)         | 14.48 |
| DG (36:4e)                  | 6.15  | TG (16:0_16:1_18:1)         | 4.22  |
| DG (18:1_18:1)              | 6.94  | TG (16:0_16:0_18:1)         | 2.68  |
| DG (18:1_18:2)              | 12.46 | TG (16:0_14:0_18:2)         | 38.08 |
| DG (16:0_18:2)              | 9.1   | TG (18:1_18:2_18:2)         | 40.05 |
| DG (32:1e)                  | 10.05 | TG (16:0_18:2_18:3)         | 72.33 |
| DG (34:2e)                  | 12.14 | TG (16:0_18:1_18:2)         | 216.21|
| DG (16:0_16:0)              | 2.61  | TG (18:1_14:0_18:3)         | 85.11 |
| DG (16:1_18:2)              | 15.2  | StE (4:0)                   | 2.45  |
| DG (34:3e)                  | 15.38 | StE (30:5)                  | 5.05  |
| Hex1Cer (d18:1_24:1)        | 2.65  | ZYE (13:1)                  | 2.08  |
| LPE (18:1)                  | 3.81  | ZYE ()                      | 4.23  |
| MLCL (18:2_18:2_18:2)       | 2.69  | **25HC**                    | **0.31**|
| PG (28:0_16:0)              | 50.44 | **27HC**                    | **0.004**|
| PG (28:0_16:0)              | 480.71|                             |       |
TRANSPARENT METHODS

Materials

Cell culture reagents and supplies were purchased from GIBCO BRL (Grand Island, NY); 25HC was purchased from New England Nuclear (Boston, MA). Huh-7 cells were obtained from American Type Culture Collection (Rockville, MD). Human liver tissues were purchased from Sekisui XenoTech, LLC (Kansas City, KS). The reagents for real time RT-PCR were from AB Applied Biosystems (Warrington, UK). The chemicals used in this research were obtained from Sigma Chemical Co. (St. Louis, MO) or Bio-Rad Laboratories (Hercules, CA). Polyclonal mouse antibodies against SREBP-1C, were purchased from Santa Cruz Biotechnology (Santa Cruz, CA), and DNMT1 was purchased from Abcam Inc. (Burlingame, CA). All solvents were obtained from Fisher (Fair Lawn, NJ) unless otherwise indicated. Biochemical assay reagents were purchased from Wako Chemicals USA, Inc. (Richmond, VA). For lipidomic analysis, internal standards were purchased from Avanti Polar Lipids (Alabaster, AL) as their premixed splash lipidomixmass spec standard. Internal standards were added to samples in 10 µL aliquots. Standards included 15:0-18:1(d7) PC, 15:0-18:1(d7) PE, 15:0-18:1(d7) PS, 15:0-18:1(d7) PG, 15:0-18:1(d7) PI, 15:0-18:1(d7) PA, 18:1(d7) LPC, 18:1(d7) LPE, 18:1(d7) Cholesterol Ester, 18:1(d7) MAG, 15:0-18:1(d7) DAG, 15:0-18:1(d7)-15:0 TAG, 18:1(d9) SM, and Cholesterol (d7). For LC-MS/MS analyses, a Thermo Scientific Q Exactive HF Hybrid Quadrupole-Orbitrap Mass Spectrometer was used. Samples were separated via a Thermo Scientific Vanquish Horizons UHPLC System functioning in binary mode.

Cell Culture

Huh-7 cells were cultured in DMEM medium supplemented with 10% heat-inactivated fetal bovine serum (FBS), with either low glucose (LG, 1.5 g/L) or high glucose (HG, 4.5 g/L) at 37 °C in a humidified atmosphere of 5% CO₂.

Qualification and Quantification of Intracellular Neutral Lipids in Hepatocytes
Oil Red O Staining: Huh-7 cells were seeded on 22 × 22 mm glass coverslips in six-well plates. Cells were cultured in DMEM media with LG or HG for 0, 36 and 72 hrs, fixed with 3.7% formaldehyde in PBS (phosphate-buffered saline) for 30 min, followed by three washes with PBS. The cells were stained with 0.2% Oil Red O in 60% isopropanol for 10 min and washed three times with PBS. Cell nuclei were stained with Harris hematoxylin for 1 minute, washed with PBS three times. The stained lipids were examined by microscopy (Ma et al., 2008).

Total levels of cellular triglyceride (TG), total cholesterol, cholesterol ester, and free fatty acids were determined by biochemical kits according to the manufacturer’s instructions. Lipids contents were normalized to protein concentrations tested with protein quantitative assay kit (Bio-Rad) (Xu et al., 2013). Briefly, Huh-7 cells were homogenized, and then lipids were extracted with a mixture of chloroform and methanol (2:1, vol/vol). The filtered extracts, 0.2 ml, were evaporated to dryness and dissolved in 100 µl of isopropanol containing 10% of triton X-100 for total cholesterol and cholesterol esters assay (Wako Chemicals USA, Richmond, VA); NEFA solution (0.5 g of EDTA-Na₂, 2 g of Triton X-100, 0.76 ml of 1N NaOH, and 0.5 g of sodium azide/l, pH 6.5), for free fatty acid assay (Wako Chemicals USA, Richmond, VA); and isopropanol only, for triglyceride assay (Fisher Scientific, Pittsburgh, PA), respectively.

**Extraction and Determination of DNA and mRNA Levels**

Genomic DNA from 5,000 Huh-7 cells cultured in DMEM medium with LG or HG for 72 hours were extracted using QIAamp DNA Mini Kit (QIAGEN, Hilden, Germany). Each sample, 2 µg, was sent to EpigenDx, Inc. (Hopkinton, MA) for analysis of global methylation and target next generation bisulfite sequencing. The same samples, 6 µg /each, were sent to Novogene Co., Ltd (Tianjin, China) for analysis of whole genome bisulfite sequencing (WGBS). Total RNA was isolated with an SV total RNA isolation system (Promega, Madison, WI) with DNase treatment. Each sample, 2 µg, was used for the first-strand cDNA synthesis as recommended by the
manufacturer (Invitrogen, Carlsbad, CA). Real-time RT-PCR was performed using SYBR Green as the indicator on ABI 7500 Fast Real-Time PCR System (Applied Biosystems, Foster City, CA). Amplifications of β-actin or GAPDH were used as internal controls. Relative messenger RNA (mRNA) expression was quantified with the comparative cycle threshold (Ct) method using the primer set shown in Table S1 and was expressed as $2^{\Delta\Delta Ct}$ (Livak and Schmittgen, 2001).

**Western blot Analysis**

Specific proteins in nuclei were analyzed by western blot. The nuclear proteins were extracted using NE-PER Nuclear and Cytoplasmic Extraction Reagents (Fisher Scientific). For each sample, 10 µg of nuclear proteins were separated on 8%-12% SDS polyacrylamide gel electrophoresis (SDS-PAGE) gels. Electrophoresis was performed at 100V for 15 min and 200V for another 25 min in a Bio-Rad mini-gel system. After electrophoresis, samples were transferred onto a polyvinylidene difluoride (PVDF) membrane at 30V for 50 min. The specific proteins on the membrane were detected by incubation with various primary antibodies including anti-SREBP-1C (Santa Cruz Biotechnologies) or anti-DNMT1 (Abcam) at 4°C overnight, followed by further incubation with an appropriate horseradish peroxidase (HRP)-conjugated secondary antibody (Bio-Rad) at room temperature for 1 hr. Lamin B1 protein levels were used as controls for equal nuclear protein loading. Each positive band was quantified by Advanced Image Data Analyzer (Aida, Straubenhardt, Germany) (Zhang et al., 2012).

**Lipidomic Analysis of Nuclear Neutral Lipids**

**Extraction of Nuclear Lipids**

The human liver tissue (80 mg) from normal people or NASH patients were homogenized in 1 ml PBS and centrifuged. Huh-7 cells were cultured in 100 mm dishes with 8 ml of media with LG or HG for 72 hrs. The cells were washed twice with cold phosphate-buffered saline (PBS). Both liver and Huh-7 cellular pellets were resuspended in 0.5 ml cytoplasmic extraction reagent I and incubated on ice for 10 min. Then 27.5 µl of
cytoplasmic extraction reagent II were added and the mixture was incubated on ice for 1 min, centrifuged for 5 min at 16,000 × g, the supernatant was completely removed, and the pellet was washed twice with cold PBS. Fifty µl of nuclear extraction reagent, 450 µl of PBS and 50 µl of proteinase K (1mg/ml) were added to the tube and the mixture was incubated for 3 hrs at 50 °C. Total lipids were extracted and partitioned by adding 3.3 volumes of chloroform: methanol (1:1) as previously described (Ren et al., 2006). Cholesterol and hydroxycholesterols were distributed into the chloroform phase.

The extracted lipids in chloroform phases from both human liver tissues and Huh-7 cells were collected into 13 x 100 mm borosilicate tubes with a Teflon-lined cap (West Chester, PA), respectively. Then 2 mL of CH₃OH and 1 mL of CHCl₃ were added along with the internal standard cocktail (10 µL). The contents were dispersed using an ultra sonicator at room temperature for 30 s. This single phase mixture was incubated at 48 °C overnight. Debris was then pelleted in a centrifuge for 5 minutes at 5000 × g, and the supernatant was transferred to a clean tube. The extract was reduced to dryness using a Speed Vac. The dried residue was reconstituted in 0.2 ml of the starting mobile phase solvent for untargeted analysis, sonicated for 15 sec, then centrifuged for 5 minutes in a tabletop centrifuge before transfer of the clear supernatant to the auto-injector vial for analysis.

Untargeted analysis

The lipids were separated by reverse phase LC using a Thermo Scientific Accucore Vanquish C18+ 2.1 (i.d.) x 150 mm column with 1.5 µm particles. The HPLC used a binary solvent system at a flow rate of 0.26 mL/min with a column oven set to 55 °C. Prior to injection of the sample, the column was equilibrated for 2 min with a solvent mixture of 99% Mobile phase A1 (CH₃CN/H₂O, 50/50, v/v, with 5 mM ammonium format and 0.1% formic acid) and 1% Mobile phase B1 (CH₃CHOHCH₂/CH₃CN/H₂O, 88/10/2, v/v/v, with 5 mM ammonium format and 0.1% formic acid) and after sample injection (typically 10 µL), the A1/B1 ratio was maintained at 99/1 for 1.0 min, followed by a linear gradient to 35% B1 over 2.0 min, then a linear gradient to 60% B1 over 6 min,
followed by a linear gradient to 100% B1 over 11 min., which held at 100% B1 for 5 min, followed by a 2.0 min
gradient return to 99/1 A1/B1. The column was re-equilibrated with 99:1 A1/B1 for 2.0 min before the next run.

Each sample was injected two times for analysis in both positive and negative modes. For initial full scan MS
(range 300 to 200 m/z) the resolution was set to 120,000 with a data-dependent MS\(^2\) triggered for any analyte
reaching 3E6 or above signal. Data-dependent MS\(^2\) were collected at 30,000 resolution. Data was analysed using
Thermo Scientific’s Lipid Search 4.2 software.

**Analysis of Cholesterol and Oxysterol Biosynthesis**

After incubation of Huh-7 cells in 100 mm dishes with 8 ml of media with LG or HG for 72 hrs, 5 µCi of
\([1-\text{\textsuperscript{14}}\text{C}]\)-acetate was added and culturing was continued for another 3 or 9 hrs. After 9 hrs of incubation at 37°C, the
media was removed and the cells were washed twice with cold PBS, harvested in 1ml PBS, and collected in
microcentrifuge tubes. The cells were sedimented by centrifugation, and the pellets were washed three times by
resuspension and sedimentation. The nuclear lipids were extracted as described above. \([\text{\textsuperscript{14}}\text{C}]\)-Acetate derivatives in
the chloroform phase were analyzed by HPLC on a Waters Spherisorb Analytical C18 column (5 µm, 4.0 mm x 50
mm). The mobile phase consisted of (A), 5 mM ammonia acetate in 95% of water, 5% acetonitrile, and (B), 5 mM
ammonia acetate in 95% of methanol, 5% of acetonitrile. The 45 min gradient was as follows: 0-25.0 min,
60%-100% B linear; 25.0-40.0 min, 100% B; 40.0-45.0 min, 100%-60% B linear. The elution stream from the
HPLC column was monitored by OD\(_{202\text{nm}}\) using Agilent Series 1100 solvent delivery system (Hewlett Packard) at
0.5 ml/min flow rate. The eluted fractions were collected every 0.5 min (0.25 ml per fraction) except as indicated.
The counts of \([\text{\textsuperscript{14}}\text{C}]\)-acetate derivatives in each fraction were determined by Scintillation Counting. The column
was calibrated with cholesterol, 25-hydroxycholesterol, and 27-hydroxycholesterol.

**Enzyme Kinetic Study of 25-Hydroxycholesterol as Epigenetic Regulator**

The activities of DNMT1, DNMT3a, DNMT3b, GCN3 (Giant congenital nevi), p300 (histone acetyl
transferase), Pcaf (KAT2B lysine acetyltransferase 2B), HDAC1 (histone deacetylase 1), HDAC2 (histone deacetylase 2), HDAC3 (histone deacetylase 3), HDAC6 (histone deacetylase 6), HDAC10 (histone deacetylase 10), and KDM6B-JMJD3 (lysine demethylase 6B) were measured as instructed by reaction biology company (Reaction Biology Corp. PA, USA).

For the DNMT1 activity assay, the substrate solution, 0.001 mg/ml Poly (dI-dC):Poly (dI-dC) in 50 mM Tris-HCl, pH 7.5, 50 mM NaCl, 5 mM EDTA, 5 mM DTT, 1 mM PMSF, 5% glycerol, 0.01% Brij35, 1% DMSO, was used. For the DNMT3a/3b activity assay, 0.0075 mg/ml Lambda DNA in 50 mM Tris-HCl, pH 7.5, 50 mM NaCl, 5 mM EDTA, 5 mM DTT, 1 mM PMSF, 5% glycerol, 1% DMSO, was used. The indicated DNMT1, DNMT3a and DNMT3b was added to the appropriate substrate solution and gently mixed, amounts of cholesterol, 25HC or 27HC ranging from 5.08E-09 to 0.0001 M in DMSO were added to the reaction mixture by using Acoustic Technology (Echo 550, LabCyte Inc. Sunnyvale, CA). The mixtures were incubated for 15 min, ³H-SAM was added to the reaction mixture to initiate the reaction, and the mixture was incubated for 60 min at 30°C. Following incubation, the reaction mixture was transferred to filter-paper for detection of radioactivity counts.

Analysis of Global Methylation, Long Interspersed Nucleotide Element 1 (LINE-1) Assay

For global DNA methylation analysis, 500 ng of extracted genomic DNA was bisulfite-treated using the EZ DNA Methylation kit (Zymo Research, Inc., CA). PCR reaction and product purification were performed as per the manufacturer’s protocol (GE Healthcare Life Sciences). Ten μL of the PCR products were sequenced by Pyrosequencing on the PSQ96 HS System following the manufacturer’s instructions (Pyrosequencing, Qiagen). The methylation status of each CpG site was determined individually as an artificial C/T SNP using QCpG software (Pyrosequencing, Qiagen). The methylation level at each CpG site was calculated as the percentage of the methylated alleles divided by the sum of all methylated and unmethylated alleles. The mean methylation level was calculated using methylation levels of all measured CpG sites within the targeted region of each gene. Each
experiment included non-CpG cytosines as internal controls to detect incomplete bisulfite conversion of the input DNA. In addition, a series of unmethylated and methylated DNA were included as controls in each PCR. Furthermore, PCR bias testing was performed by mixing unmethylated control DNA with in vitro methylated DNA at different ratios (0%, 5%, 10%, 25%, 50%, 75%, and 100%), followed by bisulfite modification, PCR, and Pyrosequencing analysis.

**Analysis of Whole Human Genome Bisulfite Sequencing (WGBS)**

Each sample, 5.2 µg, of genomic DNA spiked with 26 ng lambda DNA was fragmented by sonication to 200-300 bp with Covaris S220, followed by end repair and adenylation. Cytosine-methylated barcodes were ligated to sonicated DNA per manufacturer’s instructions. These DNA fragments were treated twice with bisulfite using EZ DNA Methylation-Gold TM Kit (Zymo Research), before the resulting single-strand DNA fragments were PCR amplified using KAPA HiFi Hot Start Uracil and Ready Mix (2X). Library concentration was quantified by Qubit® 2.0 Flurometer (Life Technologies, CA, USA) and quantitative PCR, and the insert size was assayed on an Agilent Bioanalyzer 2100 system.

The library preparations were sequenced on an Illumina Hiseq 2500/4000 or Novaseq platform and 125 bp/150 bp paired-end reads were generated. Image analysis and base calling were performed with Illumina CASAVA pipeline, and finally 125 bp/150 bp paired-end reads were generated. Trimmomatic (Trimmomatic-0.36) software was used for quality control. Bismark software (version 0.16.3; Krueger F, 2011) was used to perform alignments of bisulfite-treated reads to a reference genome (-X 700 --dovetail). DSS software (Park and Wu, 2016) was used to identify differentially methylated regions (DMRs). KOBAS software (Mao et al., 2005) was used to test the statistical enrichment of DMR related genes in the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways.

**Target Next Generation Bisulfite Sequencing**
The samples were prepared as described above for the LINE-1 assay. Libraries were prepared using a custom Library Preparation method created by EpigenDx. Next, library molecules were purified using AgencourtAMPure XP beads (Beckman Coulter) and quantified using the Qiagen QIAxcel Advanced System. Barcoded samples were then pooled in an equimolar fashion before template preparation and enrichment were performed on the Ion Chef™ system (Thermo Fisher) using Ion 520™ & Ion 530™ ExT Chef reagents. Following this, enriched, template-positive library molecules were then sequenced on the Ion S5™ sequencer using an Ion 530™ sequencing chip (Thermo Fisher). FASTQ files from the Ion Torrent S5 server were aligned to the local reference database using open-source Bismark Bisulfite Read Mapper with the Bowtie 2 alignment algorithm. Methylation levels were calculated in Bismark by dividing the number of methylated reads by the total number of reads.

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