**Epigenetic Modifications in the Biology of Nonalcoholic Fatty Liver Disease**

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**Abstract:** The 5-Hydroxymethylcytosine (5-hmC) is an epigenetic modification whose role in the pathogenesis of metabolic-related complex diseases remains unexplored; 5-hmC appears to be prevalent in the mitochondrial genome. The Ten-Eleven-Translocation (TET) family of proteins is responsible for catalyzing the conversion of 5-methylcyto- sine to 5-hmC. We hypothesized that epigenetic editing by 5-hmC might be a novel mechanism through which nonalcoholic fatty liver disease (NAFLD)-associated molecular traits could be explained.

Hence, we performed an observational study to explore global levels of 5-hmC in fresh liver samples of patients with NAFLD and controls (n = 90) using an enzyme-linked-immunosorbent serologic assay and immunohistochemistry. We also screened for genetic variation in TET genes.

**INTRODUCTION**

Nonalcoholic fatty liver disease (NAFLD) is a highly prevalent chronic liver disease that is clustered in the metabolic syndrome (MetS) along with other cardiometabolic comorbidities. 2-4 In fact, NAFLD is not only a component of the diagnostic criteria for MetS but is critically involved in the development of insulin resistance.5,6

The clinical picture of NAFLD might present in a relatively mild form, characterized by abnormal hepatic triglyceride accumulation, named simple or bland steatosis or nonalcoholic fatty liver (NAFL). However, it might progress to a more severe clinical condition, named nonalcoholic steatohepatitis (NASH), which is characterized by fat accumulation accompanied by liver cell injury, a mixed inflammatory lobular infiltrate, and...
variable fibrosis. Notably, NAFL is not necessarily associated with a benign prognosis. A recent meta-analysis showed that patients with NAFL could develop liver fibrosis over time. In addition, NAFL may seriously increase the probability of exacerbating associated co-morbidities, including cardiovascular disease.

Like many other complex diseases, NAFLD is under both genetic and environmental influences. Recent evidence from human studies indicates presence of tissue-specific epigenetic modifications in both nuclear and mitochondrial (mt) genome. These are involved not only in NAFLD pathogenesis and histological disease progression but are relevant to the pathophysiology of the MetS-associated phenotypes.

Furthermore, there is compelling evidence showing that mitochondrial function, and mitochondrial DNA (mtDNA) copy number are both diminished in NAFLD. Notably, the aforementioned molecular changes are not only determinants of the disease progression and severity but are also critical modifiers of disease-associated metabolic traits. In fact, we previously demonstrated that epigenetic modifications occurring in the liver of patients with NAFLD, such as methylation of target sites of the peroxisome proliferator-activated receptor gamma coactivator 1 (PPARGC1A), a master transcription factor involved in liver-related metabolic pathways, including mitochondrial biogenesis and fatty acid oxidation, gluconeogenesis, and lipogenesis adversely affect peripheral insulin sensitivity.

Finally, we and others have previously shown that liver epigenetic changes are not necessarily stable but subject to reversion. These are involved not only in NAFLD pathogenesis but are also critical modifiers of disease-associated metabolic traits. In fact, we previously demonstrated that liver epigenetic changes are not necessarily stable but subject to reversion.

The 5-hydroxymethylcytosine (5-hmC) is an old, but recently rediscovered and understood epigenetic modification that modulates gene transcription by influencing putative DNA demethylation and chromatin structure remodelling. Interestingly, 5-hmC profile seems to be tissue-specific and is highly dependent on the cellular state, resulting in changes in response to environmental stimuli and metabolic perturbations. Non-CG hydroxymethylation appears to be prevalent in the mitochondrial genome as well.

The Ten-Eleven-Translocation (TET1, TET2, and TET3) family of proteins, which share significant sequence homology at their C-terminal catalytic domains, were recently identified as 5mC hydrolases that catalyze the conversion of 5-mC to 5-hmC. These proteins, which frequently mutate in cancer, are involved in programmed cell death in neurons. However, whether this effect could be observed in hepatocytes remains unknown.

Because NAFLD is associated with altered DNA methylation profiles, changes in mtDNA content and PPARGC1A expression, we hypothesized that epigenetic editing by 5-hmC might be a novel mechanism through which the NAFLD-associated molecular traits could be explained. Hence, we explored the global levels of 5-hmC in the liver of patients with NAFLD at different stages of disease severity and controls.

In addition, we screened for genetic variation in TET1–3 loci by targeted NGS in a discovery group of 96 individuals (64 NAFLD patients and 32 controls); and independent replication of selected single nucleotide polymorphisms (SNPs) in a case–control association study (n = 380 participants).

Patients were included in the study if there was histopathological evidence of NAFLD, either NAFL or NASH, based on a liver biopsy (LB) conducted within the study period. Exclusion criteria were secondary causes of steatosis, including alcohol abuse (>30 g alcohol daily for men and >20 g for women), total parenteral nutrition, hepatitis B and hepatitis C virus infection, and the use of drugs known to precipitate steatosis. By using standard clinical and laboratory evaluation, as well as LB features when applicable, autoimmune liver disease, metabolic liver disease, Wilson’s disease, and α-1-antitrypsin deficiency were likewise ruled out in all patients.

Healthy subjects were selected for inclusion into the control group if their age and sex matched those of the NAFLD patients and in whom, in addition to the standard health assessment described below, a careful ultrasonographic (US) examination of the liver was performed to exclude fatty liver infiltration. In addition, controls were included at any study phase if they did not exhibit features of MetS and did not have fatty liver at liver US.

For obvious ethical reasons, the control subjects were not exposed to the potential risk of an LB; hence, in explorations that involved molecular studies requiring fresh liver tissue, patients with near-normal liver histology (NNLH) were included. In the NNLH subjects, LB was justified owing to the presence of persistently mildly elevated serum activity of liver enzymes. In all NNLH subjects, all causes of common liver disease were ruled out, and these patients were included in the study if they did not present either histological evidence of fatty change or necro-inflammatory activity, and had either minimal changes or mild cholestasis as histological diagnosis.

The case participants and the controls were selected during the same study period from the same population of patients attending the liver unit, and all shared the same demographic characteristics.

Ethics, Consent, and Permissions

Human serum and DNA samples from healthy individuals and those diagnosed with NAFLD, as well as liver biopsies from all the patients, were obtained with written informed consent following Institutional Review Board-approved protocols (protocol number: 104/HGAZ/09, 89/100, and 1204/2012, Comité de Bioética Hospital Abel Zubizarreta). All the investigations performed in this study were conducted in accordance with the guidelines of the 1975 Declaration of Helsinki.

Physical, Anthropometric, and Biochemical Evaluation

Health examinations included anthropometric measurements, a questionnaire on health-related behaviors, and biochemical determinations.

Caspase-generated CK-18 fragment (CK-18)—a noninvasive quantification of hepatocellular apoptosis—concentration was measured by the 1-step in vitro immunonassay M30-aptosense enzyme-linked-immunosorbent serologic assay (ELISA) kit (PEVIVA BA; DiaPharma, OH) that selectively recognizes the caspase-generated fragments of keratin 18 containing the K18Asp396 neoepitope (CK-18).
Anthropometric measurements and blood samples were obtained from each patient at the time of LB or DNA sample collection, and before any intervention. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400.

Liver Biopsy and Histopathological Evaluation
LB was performed before any intervention with ultrasound guidance and a modified 1.4 mm-diameter Menghini needle (Hepafix, Braun, Germany) under local anesthesia, in the outpatient setting. A portion of each LB specimen was routinely fixed in 40 g/L formaldehyde (pH 7.4) embedded in paraffin before being stained with hematoxylin and eosin, Masson trichrome, and silver impregnation for reticular fibers. All the biopsies were at least 3 cm in length and contained a minimum of 8 portal tracts. The liver biopsies were read by a single experienced hepatopathologist, who was blinded to all the clinical and laboratory data. The degree of steatosis was assessed according to the system developed by Kleiner et al. NASH was defined as steatosis, accompanied by mixed inflammatory-cell infiltration, hepatocyte ballooning and necrosis, glycogen nuclei, Mallory’s hyaline, and any stage of fibrosis, including absent fibrosis. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400.

Liver Immunohistochemistry
Immunostaining for TET1 protein and 5-hmC was performed on liver specimens previously obtained from NAFLD patients and controls and included in paraffin before use. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400.

Electron Microscopy of the Liver Tissue
Electron microscopy (EM) studies were performed on liver specimens by immersion of a small sample (1 mm) in 2.5% glutaraldehyde and 0.1 M phosphate buffer (pH 7.4). Sections were mounted in copper grids, counterstained with lead citrate, and observed in a transmission electronic microscope (Carl Zeiss, Carl-Zeiss-Straße 56, 73447 Oberkochen, Germany). Further details are provided in extant literature.

RNA Preparation and Real-Time RT-PCR for Quantitative Assessment of mRNA Expression and Bisulfite Treatment of DNA for Methylation-Specific PCR
Total RNA was prepared from liver tissue using phenol extraction step method, with an additional DNase digestion. Genomic DNA was isolated from liver biopsies at the same time as RNA by standard methods, and DNA bisulfite modification was performed as previously reported. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400, which includes details on reverse transcription polymerase chain reaction (RT-PCR), data normalization, and the primer sequences.

Global Liver 5-hmC Quantification
We measured global 5-hmC levels in the DNA isolated from fresh liver specimens of patients and controls using Qubit 5-hmC DNA ELISA Kit (Zymo Research, Irvine, CA) according to the manufacturer’s instructions. This system is highly sensitive to 5-hmC DNA and has a detection threshold of 0.02% per 100 ng input DNA. In brief, 5-hmC was detected using a sandwich-based ELISA kit that includes a control DNA set, which was calibrated to accurately quantify the percent 5-hmC in sample DNA by a standard curve. All samples were measured in triplicate, allowing us to estimate the mean % 5-hmC level. The % 5-hmC levels were not normally distributed and were thus log-transformed for statistical analyses.

Characterization of Genetic Variation in TET 1–3 by NGS
NGS technology was employed to search for the presence of genetic variation in TET1–3 loci covering all protein-coding exons (TET1: 9.7 kb, 12 exons, TET2: 19 kb, 13 exons, and TET3: 11 kb, 9 exons), in addition to exon–intron boundaries and 5’ and 3’ UTR regions, which were examined by semiconductor technology offered in the Ion Torrent PGM system (Life Technologies, Carlsbad, CA) using a 316 chip. All reagents were obtained from the same provider. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400.

Replication Study: Genotype and Association Analysis, Power and Sample Size Calculation, and Population Stratification
The genetic analyses were performed on genomic DNA extracted from white blood cells. Genotyping of the TET1-rs3998860 (p.Ile1123Met) and TET2-rs2454206 (p.Ile1762Val) was performed using a TaqMan genotyping assay (dbSNP rs3998860 assay C_2719530_10 and dbSNP rs2454206 assay C_11566753_20; Applied Biosystems, Carlsbad, CA) according to the manufacturer’s instructions.

Using the CaTS power calculator for genetic association studies and assuming a prevalence of NAFLD of 0.30, an minor allele frequency of ~0.3, and a relative risk of ~2, our sample had 100% power for the additive genetic model and 98% for the dominant 1. Complete details are provided in the Supplemental Content, http://links.lww.com/MD/A400.

Quantification of Liver mtDNA Copy Number
An assay based on real-time quantitative PCR was used for both nuclear DNA (nDNA) and mtDNA quantification using SYBR Green as a fluorescent dye (Invitrogen, Carlsbad, CA 92008, USA) as previously described. Primer sequences are shown in Supplemental Content, http://links.lww.com/MD/A400, along with further details on quantification of liver mtDNA copy number.

Statistical Analysis
Quantitative data were expressed as mean ± standard deviation (SD) unless otherwise indicated. Because significant differences in variance were observed between the groups in most of the variables and the distribution was significantly skewed in most cases, we chose to be conservative and assessed the differences between the groups by using nonparametric Mann–Whitney U or Kruskal–Wallis tests. As indicated, in some comparisons, log-transformed variables were compared by analysis of variance and Neuman–Keuls test for mean
differences, as in the case of liver 5-hmC levels, liver methylation and mRNA expression of PPARGC1A, and serum CK-18 levels. Univariate correlations were obtained via the Spearman’s rank correlation test. For association tests between genotypes or allele frequencies and phenotypes, such as liver disease and Type 2 diabetes, Cochran–Armitage test for trend, or logistic regression were employed.

The CSS/Statistica program package version 6.0 (StatSoft, Tulsa, OK) was employed in the aforementioned analyses.

RESULTS

Clinical and biochemical features of the entire population in each of the study phases are shown in Tables 1–3. Overall, patients with NAFLD had all the features of the MetS, including abdominal obesity, insulin resistance, hypertriglyceridemia, and arterial hypertension.

Global Liver DNA Hydroxymethylation Levels Are Significantly Associated With Liver mtDNA Content and the Abundance of PPARGC1A mRNA

First, we explored the association between liver 5-hmC levels and NAFLD as a disease trait. The findings revealed no differences in the overall mean levels of global DNA hydroxymethylation in subjects with NNLH (0.100 ± 0.016) and patients with NAFLD (0.095 ± 0.034), P = NS. The overall median (interquartile range, IQR) levels of global DNA hydroxymethylation were 0.101% (IQR: 0.035) and 0.094% (IQR: 0.052) for subjects with NNLH and NAFLD, respectively.

The analysis of global liver 5-hmC levels and NAFLD disease severity did not show significant differences either (NAFL: 0.090 ± 0.035% vs NASH: 0.100 ± 0.032%, mean, SD, P = NS), and the same conclusion was reached for liver fibrosis (F0–F1: 0.10 ± 0.032% vs F2–F4: 0.094 ± 0.034%, P = NS). To explore the localization pattern of 5-hmC, we further evaluated the liver specimens (stained by immunohistochemistry [IHQ], by a specific antibody) of NAFLD patients and subjects with NNHL. While the immunoreactivity product of 5-hmC was preferentially localized in the nucleus of hepatocytes in both controls and NAFLD patients, with no significant differences in staining intensity, there were remarkable differences in both the pattern of nonnuclear localization and the immunostaining counts. The immunoreactivity product of 5-hmC in subjects with NNHL was randomly located in nonnuclear (cytoplasmic) areas of hepatocytes, showing a granular pattern. Conversely, in patients with NAFLD, 5-hmC was preferentially localized at the edge of lipid-laden hepatocytes (Figure 1). Notably, liver specimens of patients with NAFLD were examined by EM and showed mitochondria in close physical association with large lipid droplets (Figure 1)—a finding that supports a putative localization of the nonnuclear immunoreactivity product of 5-hmC in mtDNA. Further

TABLE 1. Clinical and Biochemical Characteristics of Subjects With Near Normal Liver Histology (NNLH) and Patients With NAFLD Included in the Exploration of Liver Global Levels of DNA 5-Hydroxymethylation

| Variables (Mean ± SD) | NNLH | NAFL | NASH | P Value* (P<) | P Value** (P<) | P Value*** (P<) |
|----------------------|------|------|------|--------------|--------------|--------------|
| Number of subjects   | 23   | 31   | 36   |              |              |              |
| Age, y               | 47 ± 9 | 51 ± 8 | 48.0 ± 8.7 | NS           | NS           | NS           |
| BMI, kg/m²            | 25.5 ± 3.9 | 31.5 ± 5.0 | 32.3 ± 6.0 | 0.00008      | 0.000032     | NS           |
| Waist circumference, cm | 91 ± 16 | 104.0 ± 8.4 | 107 ± 13 | 0.027 | 0.004 | NS           |
| Fasting plasma glucose, mg/dL | 88.4 ± 11.7 | 98 ± 13 | 104 ± 12 | 0.014 | 0.000036 | NS           |
| Fasting plasma insulin, μU/mL | 5.9 ± 1.4 | 45.10 ± 10.0 | 15.0 ± 9.5 | 0.00001 | 0.000036 | NS           |
| HOMA-IR index         | 1.2 ± 0.3 | 3.4 ± 2.0 | 3.9 ± 2.6 | 0.000005 | 0.000014 | NS           |
| SABP, mm Hg           | 116 ± 19 | 124 ± 13 | 126.0 ± 13.5 | 0.06 | 0.04 | NS           |
| DABP, mm Hg           | 72 ± 11 | 77 ± 13 | 78.4 ± 8.5 | NS           | NS           | NS           |
| Total cholesterol, mg/dL | 202 ± 41 | 204 ± 36 | 218 ± 49 | NS           | NS           | NS           |
| Triglycerides, mg/dL  | 145 ± 97 | 153 ± 68 | 209 ± 136 | NS           | 0.03 | NS           |
| AST, U/L              | 48.5 ± 37.0 | 36 ± 16 | 39 ± 22 | NS           | NS           | NS           |
| ALT, U/L              | 74 ± 70 | 59 ± 39 | 59 ± 36 | NS           | NS           | NS           |
| GGT, U/L              | 110 ± 106 | 77 ± 83 | 61 ± 39 | 0.02 | 0.01 | NS           |
| AP, U/L               | 259 ± 169 | 232 ± 123 | 171 ± 68 | NS           | 0.06 | 0.04 |
| Histological features |      |      |      |              |              |              |
| Degree of steatosis, % | —   | 45 ± 27 | 60.6 ± 21.0 | —   |         | 0.01 |
| Lobular inflammation (0–3) | —   | 0.5 ± 0.6 | 0.9 ± 0.6 | —   |         | 0.008 |
| Portal inflammation (0–2) | —   | 0    | 1.5 ± 0.6 | —   |         | 0.000001 |
| Hepatocellular ballooning (0–2) | —   | 0    | 1.00 ± 0.46 | —   |         | 0.000001 |
| Fibrosis stage         | —   | 0    | 1.5 ± 0.6 | —   |         | 0.000001 |
| NAS                   | —   | 2.4 ± 1.3 | 5.8 ± 1.8 | —   |         | 0.000001 |

ALT and AST = serum alanine and aspartate aminotransferase, respectively; AP = alkaline phosphatase; BMI = body mass index; GGT = gamma-glutamyltransferase; HOMA-IR = homeostatic model assessment-insulin resistance; NAFL = nonalcoholic fatty liver or simple steatosis; NAFLD = nonalcoholic fatty liver disease; NNLH = near-normal liver histology and abnormal liver enzymes; NS = nonsignificant; SABP and DABP = systolic and diastolic arterial blood pressure, respectively; SD = standard deviation.

*NNLH vs NAFL comparisons.
**Comparisons between NNLH vs NASH.
***Comparisons between NAFL and NASH subjects. The P value pertains to the statistical significance calculated using Mann–Whitney U test.
TABLE 2. Clinical and Biochemical Characteristics of Control Subjects and Patients With NAFLD Included in the Discovery Group of the Exploration of TET 1–3 Genetic Variation by Next Generation Sequencing

| Variables (Mean ± SD) | Control | NAFL | NASH | P Value* (P<) | P Value** (P<) | P Value*** (P<) |
|-----------------------|---------|------|------|---------------|---------------|---------------|
| Number of subjects    | 32      | 32   | 32   | —             | —             | —             |
| Age, yr               | 48.0 ± 7.4 | 51.9 ± 9.8 | 51.2 ± 11.0 | NS            | NS            | NS            |
| BMI, kg/m²             | 23.0 ± 2.4 | 31.6 ± 4.7 | 36.0 ± 5.6 | 1.0 × 10⁻⁸     | 1.0 × 10⁻⁸     | 0.003         |
| Waist circumference, cm| 80.0 ± 9.1 | 101.0 ± 8.6 | 112 ± 14 | 1.0 × 10⁻⁸     | 1.0 × 10⁻⁸     | 0.001         |
| Fasting plasma glucose, mg/dL | 81.3 ± 7.2 | 96.5 ± 19.0 | 127 ± 49 | 0.0002         | 1.0 × 10⁻⁸     | 0.002         |
| Fasting plasma insulin, μU/mL | 5.3 ± 2.4 | 12.4 ± 6.5 | 20.9 ± 13.6 | 1.0 × 10⁻⁸     | 1.0 × 10⁻⁸     | 0.0007        |
| HOMA-IR index         | 1.0 ± 0.5 | 3 ± 2 | 6.2 ± 4.6 | 1.0 × 10⁻⁸     | 1.0 × 10⁻⁸     | 0.0001        |
| SABP, mm Hg           | 116 ± 9.2 | 125.0 ± 11.5 | 133.0 ± 16.4 | 0.021         | 0.0008         | NS            |
| DABP, mm Hg           | 72.5 ± 8.6 | 79 ± 8 | 79.0 ± 14.6 | 0.025         | 0.01           | NS            |
| Total cholesterol, mg/dL| 219 ± 51 | 208 ± 59 | 207 ± 46 | NS            | NS            | NS            |
| Triglycerides, mg/dL  | 87 ± 30 | 149 ± 77 | 221 ± 142 | 0.01          | 0.008         | NS            |
| AST, U/L              | 21.5 ± 4.4 | 38 ± 19 | 49.4 ± 33.0 | 0.001         | 0.0007        | NS            |
| ALT, U/L              | 21.3 ± 5.0 | 69 ± 101 | 60 ± 35 | 0.001         | 0.0007        | NS            |
| GGT, U/L              | 36.2 ± 26 | 48.0 ± 29.5 | 86 ± 82 | NS      | 0.039         | 0.03          |
| AP, U/L               | 160 ± 67 | 220 ± 89 | 269 ± 92 | 0.04          | 0.01          | 0.03          |
| CK18 levels           | 129 ± 90 | 243 ± 202 | 355 ± 284 | NS            | 0.03          | NS            |

**Historical features**

| Degree of steatosis, % | — | 49 ± 23 | 60 ± 20 | — | — | — | 0.05 |
| Lobular inflammation (0–3) | — | 0.6 ± 0.5 | 1.4 ± 0.5 | — | — | — | 0.001 |
| Portal inflammation (0–2) | — | 0 | 1.7 ± 0.7 | — | — | — | 1.0 × 10⁻⁸ |
| Hepatocellular ballooning (0–2) | — | 0.7 ± 0.6 | — | — | — | — | 0.002 |
| Fibrosis stage | — | 0.1 ± 0.5 | 1.9 ± 1.2 | — | — | — | 1.0 × 10⁻⁸ |
| NAS | — | 2.4 ± 1.0 | 6.4 ± 1.6 | — | — | — | 1.0 × 10⁻⁸ |

ALT and AST = serum alanine and aspartate aminotransferase, respectively; BMI = body mass index; GGT = gamma-glutamyltransferase; AP = alkaline phosphatase; HOMA-IR = homeostatic model assessment-insulin resistance; NAFL = nonalcoholic fatty liver or simple steatosis; NAFLD = nonalcoholic fatty liver disease; NASH = nonalcoholic steatohepatitis; NS = nonsignificant; SABP and DABP = systolic and diastolic arterial blood pressure, respectively; SD = standard deviation. 

* Comparison between NNLH vs NAFL. 
** Comparison between NNLH vs NASH. 
*** Comparisons between NAFL and NASH subjects. The P value pertains to the statistical significance calculated using Mann–Whitney U test.

Analysis on nonnuclear immunostaining scores showed significant differences between patients with NAFLD (0.4 ± 0.5) and controls (1.8 ± 0.8), P = 0.028.

We next examined crude linear trends in the association of global liver DNA hydroxymethylation with continuous phenotypic variables, including age, body mass index (BMI), percent body fat, waist circumference, homeostatic model assessment-insulin resistance (HOMA-IR), and ALT and AST (serum alanine and aspartate aminotransferase, respectively) liver enzyme levels, as well as continuous molecular traits of interest associated with NAFLD, including expression levels of PPARC1A-mRNA and liver mtDNA copy number. Of note, we found a positive and significant correlation between liver 5-hmC levels and liver mtDNA content (Spearman R = 0.50, P = 0.000382) (Figure 1), and an inverse and significant association with the abundance of liver PPARC1A-mRNA (Spearman R = −0.57, P = 0.04). As previously reported, patients with NAFLD have a significantly lower liver mtDNA copy number, expressed as an mtDNA/nDNA ratio (62.5 ± 41.0), in comparison with subjects with NNLH (103.5 ± 88.0).

In addition, a significant and inverse correlation of liver 5-hmC levels was found with ALT (Spearman R = −0.24, P = 0.02) and AST (Spearman R = −0.28, P = 0.01) serum levels, and BMI (Spearman R = −0.27, P = 0.01).

Analysis of Missense Variation in TET1 and TET2 Loci Shows Association With NAFLD-Disease Biology and Type 2 Diabetes

TET proteins play a significant role in the generation of 5-hmC; in fact, these enzymes mediate the conversion of 5-mC to 5-hmC utilizing oxygen and 2-ketoglutarate as substrates.

To understand the potential contribution of genetic variation in TET1–3 loci in NAFLD-associated molecular traits, including changes in promoter methylation levels of PPARC1A, we followed an NGS approach, which was applied to all the exons, exon–intron unions and 5’ and 3’ untranslated regions of these genes. Furthermore, based on the previous knowledge of TET-mutations are able to mediate cell death, we explored the role of variants on surrogate markers of hepatocellular apoptosis (release of caspase-cleaved cytokeratin-18, CK-18) and the disease severity.

In the exploratory study, we generated more than 2 million reads with an average length of 145 bases of quality equal to or exceeding Q20, and a coverage on average higher than 100×. Overall, analysis of sequence data yielded 28 SNPs in TET1 gene, 37 in TET2, and 24 in TET3; according to the information pertaining to the dbSNP or the 1000 genomes, among these, 23 were novel variants.
TABLE 3. Clinical and Biochemical Characteristics of Control Subjects and Patients With NAFLD in the Independent Replication Study of the Exploration of TET 1–3 Genetic Variation

| Variables (Mean ± SD) | Control Subjects | NAFL | NASH | P Valuec (P<) | P Value** (P<) | P Valuet (P<) |
|-----------------------|------------------|------|------|--------------|--------------|--------------|
| Number of subjects    | 135              | 103  | 142  | —            | —            | —            |
| Age, yr               | 48 ± 12          | 53.6 ± 10.0 | 51.8 ± 11.0 | NS          | NS          | NS          |
| BMI, kg/m²            | 25 ± 4           | 31.6 ± 5.5  | 33.0 ± 5.7   | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | 0.009        |
| Waist circumference, cm | 84 ± 16          | 103.0 ± 15.4 | 108 ± 13     | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | 0.001        |
| Fasting plasma glucose, mg/dL | 81.0 ± 13.6    | 98 ± 20      | 130 ± 122    | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  |
| Fasting plasma insulin, μU/mL | 7.0 ± 4.7     | 12.9 ± 8.0   | 16.4 ± 11.0  | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | 0.003        |
| HOMA-IR index         | 1.4 ± 1.0        | 3.1 ± 2.0    | 5.3 ± 6.8    | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | 0.00008      |
| SABB, mm Hg           | 115 ± 14         | 125.4 ± 14.0 | 128 ± 16     | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | NS           |
| DABP, mm Hg           | 71.5 ± 9.4       | 77 ± 11      | 79 ± 12      | 1.0 × 10⁻⁸  | 1.0 × 10⁻⁸  | NS           |
| Total cholesterol, mg/dL | 209 ± 42        | 205 ± 48     | 210 ± 43     | NS          | NS          | NS           |
| Triglycerides, mg/dL  | 114 ± 79         | 152 ± 74     | 192 ± 121    | 0.0005      | 1.0 × 10⁻⁸  | 0.04         |
| AST, U/L              | 18.7 ± 24.0      | 35 ± 17      | 51 ± 33      | 0.00001     | 1.0 × 10⁻⁸  | 0.00001      |
| ALT, U/L              | 17.45            | 57 ± 63      | 72 ± 55      | 0.000001    | 1.0 × 10⁻⁸  | 0.0003       |
| GGT, U/L              | 24 ± 69          | 66 ± 58      | 86 ± 86      | 0.0002      | 1.0 × 10⁻⁸  | 0.04         |
| AP, U/L               | 140 ± 118        | 233 ± 101    | 227 ± 113    | 0.000006    | 0.0001      | NS           |
| CK18 levels           | 136 ± 93         | 230 ± 231    | 312 ± 259    | NS          | 0.03        | 0.04         |

Histological features

| Degree of steatosis, % | — | 48 ± 26 | 60 ± 21 | — | — | 0.0003 |
| Lobular inflammation (0–3) | — | 0.6 ± 0.6 | 1.2 ± 0.6 | — | — | 1.0 × 10⁻⁸ |
| Portal inflammation (0–2) | — | 0 | 1.50 ± 0.66 | — | — | 1.0 × 10⁻⁸ |
| Hepatocellular ballooning (0–2) | — | 0 | 0.8 ± 0.6 | — | — | 1.0 × 10⁻⁸ |
| Fibrosis stage | — | 0 | 1.5 ± 1.3 | — | — | 1.0 × 10⁻⁸ |
| NAS | — | 2.7 ± 1.1 | 5.9 ± 1.5 | — | — | 1.0 × 10⁻⁸ |

ALT and AST = serum alanine and aspartate aminotransferase, respectively; AP = alkaline phosphatase; BMI = body mass index; GGT = gamma-glutamyl-transferase; HOMA-IR = homeostatic model assessment-insulin resistance; NAFL = nonalcoholic fatty liver or simple steatosis; NAFLD = nonalcoholic fatty liver disease; NASH = nonalcoholic steatohepatitis; NS = nonsignificant; SABB and DABP = systolic and diastolic arterial blood pressure, respectively; SD = standard deviation.

*NNLH vs NAFL comparisons.

**Stands for comparisons between NNLH vs NASH.

***Comparisons between NAFL and NASH subjects. The P value pertains to the statistical significance calculated using Mann–Whitney U test.

Information of genetic variation took precedence over nonsynonymous SNPs (those that change the amino acid sequence). Distribution of the variants’ function is depicted in the Supplemental Content, http://links.lww.com/MD/A400. As can be seen, a high percentage of genetic variation of TET1 (35.5%) and TET2 (27%) were missense variants. The main features of variants selected for the replication study, including chromosome location and predicted functional consequences, are shown in the Supplemental Content, http://links.lww.com/MD/A400.

The association analysis in the exploratory study showed that TET1-rs3998860 (p.Ile1123Met) was significantly associated with serum levels of CK-18 (P = 0.003; Figure 2). Replication analysis performed on the larger validation set confirmed the initial findings, indicating that hepatocytes of NAFLD patients who are homozygous for the G allele are more susceptible to apoptosis compared with carriers of the ancestral A allele (P = 0.001). Accordingly, an allelic test showed that the variant was significantly associated with the disease severity (Cochran–Armitage test for trend P = 0.005); generalized odds ratio (OR): 1.47, 95% confidence interval (CI): 1.10 to 1.97 for G vs A allele as a reference. Genotype distribution was in Hardy–Weinberg equilibrium (P = 0.49).

To gain insights into the putative mechanisms by which this variant might influence the disease phenotype, we next used IHQ to explore the level of liver TET1 expression according to rs3998860 genotypes. We observed that TET1 immunoreactivity product significantly (P = 0.036) increased in subjects carrying at least 1 copy of the A allele (AA and AG), in comparison with carriers of the rs3998860-GG genotype (Figure 2).

On the other hand, we aimed to analyze whether genetic variation in TET1–3 loci plays any role in the modulation of liver PPARGC1a transcriptional activity. The association analysis of sequence variation in the exploratory study showed that p.Ile1762Val substitution (TET2-rs2454206), located in exon 11, was significantly associated with the liver promoter methylation status of PPARGC1a (methylated DNA/unmethylated DNA ratio; AA: 0.61 ± 0.1 vs AG + GG: 1.6 ± 0.4, mean ± SD, P = 0.03 on log transformed values), indicating that G allele carriers have higher levels of promoter methylation compared with homozygous AA.

To further assess whether this variant has any effect on liver gene expression of PPARGC1a, we explored the abundance of the transcript according to the variant genotypes in a sub-sample of the replication set that had available isolated liver mRNA (n = 66). In line with the findings observed in the discovery stage, we found that rs2454206 was significantly associated with the relative abundance of liver PPARGC1a-mRNA in both the dominant (P = 0.005) and additive (P = 0.018) model of inheritance (Figure 3), suggesting that
carriers of the G allele have decreased liver expression of PPARGC1α. Collectively, our findings suggest that the p.Ile1762Val substitution would be involved in the modulation of liver PPARGC1α methylation profile, thereby regulating gene transcription.

To gain insights into possible clinical implications of this finding, and based on previous knowledge showing that liver expression of PPARGC1α modulates insulin resistance,6 we explored the association between the variant and the presence of Type 2 diabetes in the entire replication set. Notably, we observed a significant association between the rs2454206 and Type 2 diabetes, chi-square 10.37, $P = 0.00128$ (OR 1.60 per G allele, 95% CI: 1.01–2.51; Figure 3). The genotype distribution was in Hardy–Weinberg equilibrium ($P = 0.91$).

**DISCUSSION**

Epigenetic modifications, including changes in the DNA methylation/demethylation rate, are involved in the pathogenesis of complex diseases, including NAFLD. It is known that tissue levels of 5-hmC are variable, and are higher in embryonic pluripotent stem cells and cells of neuronal lineages, compared to adult tissue.33–35 In addition, high 5-hmC levels are a feature of rapidly proliferating cell populations. Nevertheless, 5-hmC level in the liver of patients with NAFLD presently remains unknown.

In this study, we aimed to explore global levels of 5-hmC in the liver of patients with NAFLD at different stages of disease severity, aiming to better understand its role in the disease biology. In agreement with a recent study, whereby the authors measured global 5-hmC levels in nonneoplastic liver tissue of patients with hepatocellular carcinoma,36 the amount of 5-hmC in the liver of patients with NAFLD, as well as in the liver of subjects with NNLH, was relatively low (up to 0.1%). These findings were confirmed by immunohistostaining of liver sections with specific anti-5hmC antibody.

Furthermore, while no differences in the global level of liver 5-hmC between patients and controls were observed, a significant association was found with molecular traits of interest, suggesting the possibility of an involvement of hydroxylation of mC in the development of NAFLD. For instance, we could demonstrate a significant association between liver 5-hmC and liver mtDNA copy number. Previous work in which the authors used mtDNA immunoprecipitation by an antibody directed against 5-hmC supports our findings, showing substantial 5-hmC levels in the mtDNA.25,26 It is noteworthy to highlight that a major limitation of all current methodologies that measure global DNA hydroxymethylation stems from their
inability to provide information on whether 5-hmC occurs in the nuclear or mt genome. Moreover, none provides information about locus localization in the genome. Nevertheless, we were able to show that oxidation of mC to hmC is a process that occurs not only in the nucleus of liver cells but outside them. In fact, the cytosolic localization of liver 5-hmC by immunostaining strongly suggests mitochondrial localization as extrachromosomal DNA, which is any DNA that is found outside of the nucleus of a cell, corresponds exclusively to mtDNA. While our immunohistochemical analysis revealed no differences in the nuclear 5-hmC staining scores between patients with NAFLD and controls, we observed that NAFLD was associated with a significant decrease of nonnuclear 5-hmC suggesting that mtDNA has different hydroxymethylation levels according to the disease status.

It is currently unknown whether members of the TET family are able to target liver mtDNA, as in vitro explorations in HeLa and 3T3-L1 cells found TET1 and TET2, or only TET1, respectively, in the mitochondrial fraction, suggesting that the TET-mtDNA interaction is very much cell-type dependent.26 Thus, we cannot rule out the possibility of an interaction between 5-hmC and liver mtDNA involving a large regulatory network, including genes that regulate mitochondrial biogenesis, replication and transcription. The significant association between liver levels of 5-hmC and liver mtDNA content allows us to hypothesize that epigenetic editing by hydroxymethylation is involved in the regulation of liver mtDNA copy number. Hydroxymethylated cytosines within the human mtDNA control region (D-loop), as demonstrated in previous work,26 strongly support the biological plausibility of our findings.

In addition, we observed that global 5-hmC levels in the liver were negatively associated with the abundance of liver PPARγC1A-mRNA. This finding suggests that the balance between hypomethylation and hypermethylation in this locus modulates its transcriptional activity, as we previously reported by exploring targeted regions of the gene promoter.6 In fact, the capacity to convert 5-mC into 5-hmC is very much influenced by the initial amount of 5-mC, which was shown to be higher in the liver of NAFLD patients.6

On the other hand, we observed that a missense p.Ile1123-Met variant located in TET methylcytosine dioxygenase 1 (rs3998860) was significantly associated with serum levels of the CK-18 cell death biomarker. In addition, empirical evidence indicated that caspase-mediated cleavage of the keratin 18 reflects degradation of the intracellular cytoskeleton when cells undergo apoptosis.37 This finding was replicated in the validation sample, in which the rs3998860-G allele was significantly associated with the disease severity, suggesting an involvement of TET1 locus in the modulation of apoptosis and liver injury in NAFLD. Exploration of a putative functional role of rs3998860 revealed that immunostaining scores of liver TET1 were significantly reduced in homozygous carriers of the G allele. While previous evidence indicated that the effects of TET1 overexpression on gene transcription are unrelated to its 5-mC dioxygenase function33 TET1-mediated DNA demethylation provides protection against oxidative stress in neuronal cells.28 In effect, knockdown of tet1 in cerebellar granule cells

![FIGURE 2. Association analysis of TET1-rs3998860 (p.Ile1123Met) and serum CK-18 levels. Upper panels: Results of the association analysis of rs3998860 and serum CK-18 cell death biomarker levels in the discovery and replication stage, left and right, respectively; caspase-mediated cleavage of the CK-18 reflects degradation of the intracellular cytoskeleton when cells undergo apoptosis. Each bar represents the mean ± standard error value. P values pertain to log-transformed serum CK-18 levels, analyzed by analysis of variance. Lower panels: Representative specimens of immunostaining for liver TET1 protein expression evaluated by immunohistochemistry, according to TET1-rs3998860 genotypes. Counterstaining was performed using hematoxylin. Original magnification: ×400.](image-url)
in mice significantly increases apoptosis by promoting neuronal cell death induced by oxidative stress.28

Finally, we observed that the p.Ile1762Val substitution (TET2-rs2454206) was significantly associated with liver PPARGC1A methylation and transcriptional levels. This finding suggests that the TET2 locus might be involved in the modulation of the liver PPARGC1A methylation/demethylation balance, putatively in response to the metabolic cellular environment. More importantly, this molecular trait would influence the development of Type 2 diabetes, suggesting new avenues in the understanding of the genetic susceptibility to diseases that cluster in the metabolic syndrome.

Of note, we were not able to find any significant association between global liver 5-hmC levels and genetic variation in TET-locus; neither we were able to find an association between TET-variants and mtDNA copy number. Nevertheless, previous evidence suggest that the dynamics of DNA methylation/demethylation is governed by a complex interaction of proteins, including isocitrate dehydrogenases (IDHs) that catalyze the oxidative decarboxylation of isocitrate to 2-oxoglutarate, which is a substrate of TET proteins.38 For instance, it was recently reported that the expression levels of both IDH1 and IDH2 regulate the total 5-hmC content in the adult liver.38 Furthermore, as explained before, it cannot be ascertained whether mitochondrial 5-hmC is a product of the same enzymatic reaction that takes place in the nucleus, as there is not conclusive evidence of the expression of TET proteins in the mitochondria. Hence, further research is required to guarantee a deep understanding of the regulatory mechanisms by which 5-hmC levels are modulated in NAFLD, and so, what are the mechanisms by which this epigenetic modification regulates gene expression. Remarkably, IDH-proteins localize to the mitochondrial matrix and have activity as NADP-dependent IDH either cytosolic or peroxisomal; the latter of particular significance as NAFLD severity is associated with peroxisome proliferation.15

In conclusion, our data provide evidence of the involvement of 5-hmC in the modulation of NAFLD phenotype by regulating liver mitochondrial biogenesis and the transcriptional activity of the co-activator PPARGC1A. In addition, our findings revealed a putative role of TET1-p.Ile1123Met-mediated regulation of hepatocyte apoptosis, suggesting the possibility of an “epigenetic” regulation of programmed liver-cell death. Using sequencing and replicated data, we also identified signatures of TET2-p.Ile1762Val-mediated regulation of liver PPARGC1A expression.

Taken together, our findings might contribute as a proof-of-concept study to the understanding of the role of DNA-hydroxymethylation and TET proteins in the molecular tissue-specific quantitative phenotypes associated with NAFLD pathogenesis and disease severity. To our knowledge, this is the first report of the influence of liver 5-hmC and TET-mediated

FIGURE 3. Association analysis of TET2-rs2454206 (p.Ile1762Val) and liver methylation and the transcriptional status of the coactivator PPARGC1A (peroxisome proliferator-activated receptor gamma coactivator 1α). Upper panel: Methylated DNA/unmethylated DNA ratio for the liver PPARGC1A promoter according to rs2454206 and liver abundance of PPARGC1A mRNA evaluated by quantitative real-time PCR genotypes in the dominant model of inheritance. Bottom panel: Liver abundance of PPARGC1A mRNA evaluated by quantitative real-time PCR according to rs2454206 genotypes in the additive model of inheritance, and the distribution of the variant genotypes according to the presence or absence of Type 2 diabetes in the entire replication set. Each bar represents the mean ± standard error value. In each sample, the abundance of target genes was normalized to the amount of cyclophilin to carry out comparisons between the groups. P values apply to log-transformed values of liver PPARGC1A promoter methylation and mRNA levels. PCR = polymerase chain reaction.
genetic susceptibility of nonmalignant complex diseases. Our results suggest the importance of early intervention once NAFLD is suspected/detected, as epigenetic modifications are potentially reversible and the altered “liver epigenome” can be reversed.\textsuperscript{13,15} Emerging hypotheses remain to be explained by the use of novel technologies, including high-resolution mapping of 5-hmC regions in the mitochondrial genome and sequencing of additional candidate genes that regulate the liver methylome.

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Supplemental Figure F1

*Figure illustrates genetic variation of TET1 in the whole population explored by next-generation sequencing: distribution of the predicted function of the variants. Information of variants annotation was extracted from dbSNP (http://www.ncbi.nlm.nih.gov/SNP/); information on predicted function was based on the bioinformatic tool variant Effect Predictor (VEP) (http://useast.ensembl.org/Homo_sapiens/Tools/VEP) and SnpEff platform (http://snpeff.sourceforge.net/) using University of California, Santa Cruz (UCSC) transcripts*
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Supplemental Figure F2

Figure illustrates genetic variation of \textit{TET2} in the whole population explored by next-generation sequencing: distribution of the predicted function of the variants. Information of variants annotation was extracted from dbSNP (http://www.ncbi.nlm.nih.gov/SNP/); information on predicted function was based on the bioinformatic tool variant Effect Predictor (VEP) (http://useast.ensembl.org/Homo_sapiens/Tools/VEP) and SnpEff platform (http://snpeff.sourceforge.net/) using University of California, Santa Cruz (UCSC) transcripts
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Supplemental Figure F3

Figure illustrates genetic variation of TET3 in the whole population explored by next-generation sequencing: distribution of the predicted function of the variants. Information of variants annotation was extracted from dbSNP (http://www.ncbi.nlm.nih.gov/SNP/); information on predicted function was based on the bioinformatic tool variant Effect Predictor (VEP) (http://useast.ensembl.org/Homo_sapiens/Tools/VEP) and SnpEff platform (http://snpeff.sourceforge.net/) using University of California, Santa Cruz (UCSC) transcripts.
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Supplemental Table S1

| Gene            | Forward primer 5’→3’ | Reverse primer 5’→3’ | Size, bp |
|-----------------|----------------------|----------------------|----------|
| **The primer sequences for mRNA gene expression** |
| PPARGC1A        | CCTGCATGAGTCTGTGCTCT | GCAAGAGGCTGGTCTTCAC  | 164      |
| PPIA            | TCCAGGGTTTATGTGTCAG  | CTCATCTTCAATTTCTCC   | 89       |
| **The primer sequences for mtDNA and nDNA amplification** |
| Mt F3212        | CACCCCAAGAAGAGGTTTG | TGGCCATGGGTATGTTGTTA | 108      |
| 18S rRNA        | TAGAGGGACAAGTGCGTTC | CGCTGAGCCAGTCA-GTGT  | 105      |
| **The primers sequences for methylation-specific polymerase chain reaction** |
| PPARGC1A-M      | ATTTTTTATTGTATGGGGTAGTC | AAAATATTTAAAAACGCAAACGAA | 143 |
| PPARGC1A-U      | TTTTATTGTATGGGGGTAGTTGA  | AAAAATATTTAAAAACAAACAACAAA | 141 |

Table that illustrates PCR primers used for exploration of liver PPARGC1A mRNA abundance and promoter methylation status, and liver mtDNA copy number

PPARGC1A=peroxisome proliferator-activated receptor gamma, coactivator 1 alpha; PPIA=cyclophilin. M: methylated, U: unmethylated.
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Supplemental Table S2

| Gene symbol | Chromosome Location | Variant | db SNP | Alleles | MAF* | Predicted functional consequences | Ancestral allele |
|-------------|---------------------|---------|--------|---------|------|----------------------------------|-----------------|
| **TET1**    | 10:68646098 (forward strand) | p.Ile1123Met Isoleucine (I) to Methionine (M) at position 1123 | rs3998860 | A/G | 0.27 (A) | Missense variant | A |
| **TET2**    | 4:105275794 (forward strand) | p.Ile1762Val Isoleucine (I) to Valine (V) at position 1762 | rs2454206 | A/G | 0.24 (G) | Missense variant | A |

Table that illustrates the main features of variants identified in TET locus by next generation sequencing and selected for following up in the replication study

* MAF indicates minor allele frequency in the 1000 Genome Project.
TET1: Ten-Eleven Translocation-1 or tet methylcytosine dioxygenase 1, TET2: Ten-Eleven Translocation-2 or tet methylcytosine dioxygenase 2.
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SUPPLEMENTAL METHODS

1a-Physical, anthropometric, and biochemical evaluation

1b. Liver biopsy and histopathological evaluation

1c-Liver Immunohistochemistry (IHQ)

1d-RNA preparation and real-time RT-PCR for quantitative assessment of mRNA expression

1e-Bisulfite treatment of DNA and methylation-specific polymerase chain reaction

1f-Quantification of mtDNA

1g-Next generation sequencing (NGS)

1h-Variant calling, estimation of quality control, data analysis and prediction of variant/mutation effect

1i-Replication study: Genotype and association analysis, power and sample size calculation, and population stratification

1a. Physical, anthropometric, and biochemical evaluation

The body mass index (BMI) was calculated as weight/squared height (kg/m²) and was used as an index for relative weight. The waist and hip circumferences were also measured. Elevated blood pressure was defined as systolic arterial blood pressure (SABP) ≥ 130 mm Hg and/or diastolic (DABP) ≥ 85 mm Hg, or evidence of antihypertensive treatment.

Biochemical determinations: Blood was drawn from 12-hour fasting subjects that had been in a supine resting position for at least 30 min. Laboratory evaluation included serum ALT and AST, gamma glutamyl transferase (γGT), alkaline phosphatase (AP), glucose and insulin, total cholesterol and plasma triglycerides (TG). All biochemical determinations were measured using a Hitachi-912 Autoanalyzer (Roche, Diagnostic, Buenos Aires, Argentina) or Immulite 1000 (DPC, Buenos Aires, Argentina). Homeostasis Model Assessment (HOMA-IR) was used to evaluate an insulin resistance index and was calculated as follows: Fasting serum insulin (µU/ml) × Fasting plasma glucose (mmol/l) / 22.5.

1b. Liver biopsy and histopathological evaluation

The degree of steatosis was assessed according to the system developed by Kleiner et al.¹ based on the percentage of hepatocytes containing macrovesicular fat droplets: grade 0 = <5%; grade 1 = 5-33%; grade 2 = 34% to 66% and grade 3 = >66%. NASH was defined as steatosis, accompanied by mixed inflammatory-cell infiltration, hepatocyte ballooning and necrosis,
glycogen nuclei, Mallory’s hyaline, and any stage of fibrosis, including absent fibrosis \(^2\). Intra-acinar (lobular) inflammation was defined according to Brunt \(^2\) as presence of cellular components of inflammation (polymorphonuclear leukocytes, lymphocytes and other mononuclear cells, eosinophils and microgranulomas) located in sinusoidal spaces, surrounding Mallory’s hyaline or in hepatocellular necrosis. It was graded 0–3 and was defined as 0 (absent) = no foci; 1 = < 2 foci per 200 × field; 2 = 2 to 4 foci per 200 × field; and 3 = >4 foci per 200 × field. Ballooning was scored as: 0 = none; 1 = rare or few; and 2 = many. The severity of fibrosis was expressed on a 4-point scale, as follows: 0 = none; 1 = perivenular and/or perisinusoidal fibrosis in zone 3; 2 = combined pericellular portal fibrosis; 3 = septal/bridging fibrosis; and 4 = cirrhosis.

\[1c\]-Liver Immunohistochemistry

Four-micrometer sections were mounted onto silane-coated glass slides to ensure section adhesion through subsequent staining procedures. Briefly, sections were deparaffinized, rehydrated, and washed in phosphate buffer solution (PBS), before being treated with 3% \(\text{H}_2\text{O}_2\) in PBS for 20 min at room temperature to block endogenous peroxidase. Following microwave heat-induced epitope retrieval in 0.1 M citrate buffer at pH 6.0 for 20 min, the slides were incubated with a dilution of 1:100 of TET1 (PAB21054, Abnova Taipei, Taiwan) and 1:500 of 5-hmc (Active Motif 39770, CA, USA). Immunostaining was performed using the VECTASTAIN Elite ABC Kit (Vector Lab. CA, USA) detection system. Subsequently, slides were immersed in a 0.05% 3,3’-diaminobenzidine solution in 0.1 M Tris buffer, pH 7.2, containing 0.01% \(\text{H}_2\text{O}_2\). After brown color developed, slides were removed and the reaction was arrested by immersion in PBS. Hematoxylin was used as a counterstain for TET1 and eosin for 5-hmc. Immunostaining was semi-quantitatively evaluated in a blinded fashion regarding any of the histological and clinical characteristics of the patients by two independent observers. The extent of staining was scored according to its amount and intensity, using a 4-point scoring system, as follows: 0 = no staining; 1 = positive staining in less than 20% of cells; 2 = 21-50% of positive cells, and 3 = positive staining in more than 50% of cells. Scoring included nuclear and cytoplasm staining and the sections were observed by bright field microscopy, using an Axiostar plus (Carl Zeiss, Germany) microscope at a 400× magnification.

\[1d\]-RNA preparation and real-time RT-PCR for quantitative assessment of mRNA expression

For RT-PCR, 1 to 3 \(\mu\)g of total RNA was reverse-transcribed using random hexamers and Moloney Murine Leukemia Virus (MMLV) reverse transcriptase (Promega, Wisconsin, USA). Real-time PCR was performed for quantitative assessment of mRNA expression in a StepOne Plus Real-Time PCR Systems (Applied Biosystems, CA 94404, USA). All the real-time PCR reactions were run in triplicate. The mRNA abundance of target genes was normalized to the amount of a housekeeping gene (cyclophilin A) to carry out comparisons between the groups. The selection of the housekeeping gene was based on the exploration of the most stable reference gene for testing liver mRNA expression among other housekeeping genes (\(\beta\)-actin, TATA box binding protein, cyclophilin A and glyceraldehyde-3-phosphate dehydrogenase) tested before starting this experiment. The geNorm program \(^3\) was used to identify the appropriate reference control in our samples. The mRNA levels were expressed as the ratio of the estimated amount of the target gene relative to the cyclophilin A mRNA levels using fluorescence threshold cycle values (\(\text{Ct}\)) calculated for each sample, and the estimated efficiency of the PCR for each product was expressed as the average of all sample efficiency values obtained \(^4\). The specificity of
amplification and absence of primer dimers were confirmed using the melting curve analysis at the end of each run.

**1e-Bisulfite treatment of DNA and methylation-specific polymerase chain reaction**

DNA bisulfite modification was based on bisulfite treatment of genomic DNA, thereby converting all the unmethylated cytosines to uracils, while conserving the methylated cytosines, by the EZ DNA Methylation Kit, according to the manufacturer’s protocol (Zymo Research Corporation, Orange, CA, USA). The chemically modified DNA was subsequently used as a template for a methylation-specific polymerase chain reaction (MS-PCR) to determine the promoter methylation status of the selected CpG dinucleotides in the *PPARGC1A* promoter. The assay was based on a real-time quantitative PCR in an iCycler thermocycler (BioRad Hercules, CA, USA) using SYBR Green (Invitrogen, Buenos Aires, Argentina) as a fluorescent dye. For the MS-PCR experiment, two pairs of primers were used: one pair was specific for bisulfite-modified methylated DNA (M primers) and the other pair was specific for bisulfite-modified unmethylated DNA (U primers). Thus, for each sample studied, two PCRs were performed simultaneously using the M primer and U primer pairs. Successful amplification from the M primers and U primers indicated methylation and unmethylation, respectively. For primer design, a sequence starting 2000 bp upstream from the transcriptional start site (TSS) of *PPARGC1A* was used in the MethPrimer program (http://www.urogene.org//methprimer/index1.html) to search for regions with potentially methylated CpG sites. For maximal discrimination between methylated and unmethylated alleles, M and U primers were designed to contain at least one CpG site at the 3'.

The level of methylated DNA is expressed as the ratio of the estimated amount for methylated DNA to the unmethylated DNA levels, calculated for each sample using the fluorescence threshold cycle (Ct) values for a previously estimated efficiency, according to a previous report ⁵. We estimated the efficiency for each single sample tube using the slope of the exponential phase, as described by Ramakers et al. ⁶. Furthermore, as expected, controls for unmethylated (a purified native amplicon) and fully methylated DNA (the same amplicon treated with the DNA methylase M.SssI (New England Biolabs, Ipswich, MA, USA) gave 0 and 100% DNA methylation patterns.

All the experiments were carried out in triplicate. The CV% was observed to be less than 5%. The specificity of amplification and the absence of primer dimers were confirmed by melting curve analysis at the end of each run and agarose electrophoresis. To ensure the specificity of the method and to avoid variability in the results because of the presence of two CpG dinucleotides in the reverse primer, we designed a degenerated reverse primer that introduced a mismatch in the second CG site, and observed the absence of amplification, regardless of the target DNA (data not shown), indicating that primers recognize the status of both the CpG dinucleotides simultaneously.

**If-Quantification of mtDNA**

An assay based on real-time quantitative PCR was used for both nuclear DNA (nDNA) and mitochondrial DNA (mtDNA) quantification using SYBR Green as a fluorescent dye (Invitrogen) as previously described ⁷. Results were presented as the mtDNA/nDNA ratio. Real-time quantitative PCR was carried out in a BioRad iCycler (Bio-Rad). The calculation of DNA copy number involved extrapolation from the fluorescence readings in the mode of background subtracted from the BioRad iCycler according to Rutledge et al. ⁸. Specificity of
amplification and the absence of primer dimers were confirmed by the melting curve analysis at the end of each run. The two target amplicon sequences (mtDNA and nDNA) were visualized in agarose 2% and purified by Qiagen Qiaex II, Gel extraction Kit (Tecnolab, Buenos Aires, Argentina), and dilutions of purified amplicons were used as the standard curve. The inter-assay variation coefficient was less than 20%. Specificity of the method was evaluated as was previously described. 8,9

Ig-Next generation sequencing (NGS)

DNA was isolated from whole blood, as previously described 10 and was quantified by a Qubit DNA high-sensitivity assay kit. Library preparation for each sample was performed using the IT AmpliSeq 2.0 Beta kit following the manufacturer's instructions. Briefly, 10 ng of DNA was used as a template to generate the amplicon library for sequencing variation in the three genes. Genomic regions of interest were PCR amplified prior to sequencing, and the sequencing adaptors with short stretches of index sequences (96 barcodes) that enabled sample identification were ligated to the amplicons using the IT Xpress barcode adaptor kit. The prepared library was quantified using the Ion library TaqMan Quantitation Kit. Sequencing template preparation (emulsion PCR and beads-enrichment) from sequencing libraries was carried out using an Ion OneTouch Template Kit and Ion OneTouch system (Ion OneTouch Instrument and Ion OneTouch ES, Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s protocol. Prepared templates were sequenced using Ion Sequencing Kit v2. Ion Torrent Suite software v4.2.1 (Life Technologies, Carlsbad, CA, USA) was used for converting raw signals into base calls and extracting FASTQ files of sequencing reads. Number of nucleotide flows during sequencing was set to 500 cycles.

1h-Variant calling, estimation of quality control, data analysis and prediction of variant/mutation effect

The data obtained from the Ion Torrent PGM were processed using the Ion Torrent Suite Software v 4.2.1 (Life Technologies, Carlsbad, CA, USA). Variants were annotated with dbSNP (http://www.ncbi.nlm.nih.gov/SNP/) IDs using SnpSift. In silico analysis, aimed at predicting gene and transcript functional consequences was performed by the bioinformatic tool variant Effect Predictor (VEP) (http://useast.ensembl.org/Homo_sapiens/Tools/VEP), employing ENSEMBL transcripts and Snpeff platform (http://snpeff.sourceforge.net/) using University of California, Santa Cruz (UCSC) transcripts. CellBase available at http://docs.bioinfo.cipf.es/projects/cellbase was used to annotate variants with the phenotype information from HGMD, ClinVar, UNIPROT and COSMIC. Annotation in http://www.ncbi.nlm.nih.gov/SNP was also used to determine whether variants were novel or already associated with a phenotype.

Read alignment to the hg19 Human genome reference sequence was performed using torrent mapping alignment program (TMAP) included in PGM. The Ion Torrent variant caller plug-in was used to perform the variant calling with the germline low stringency settings. In order to filter erroneous base callings, control quality filtering steps were performed using proprietary Perl scripts. At the variant coverage depth > 20, we inspected the presence of strand bias by checking the number of bases covered in both strands for the variant, the reference =/ > 4 reads in both strands, and the quality for variant calling > 20. The retained variants were visually
examined using Integrative Genomics Viewer (IGV) software (http://www.broadinstitute.org/igv/) to check for any inconsistency in the base calls.

**1i-Replication study: Genotype and association analysis, power and sample size calculation, and population stratification**

To ensure genotyping quality, we included DNA samples as internal controls, hidden samples of known genotype, and negative controls (water). The overall genotype completion rate was 100%. To account for possible population stratification, we used a collection of 13 SNPs (rs6830727, rs12639788, rs1282807, rs1947745, rs7162312, rs12951674, rs7212346, rs1934869, rs9542666, rs11843545, rs9725124, rs2798659, and rs2199940) at different loci (located in chromosomes 4, 15, 17, 13, 1, and 3) and then analyzed the data with the Structure program Version 2. We found no evidence of stratification in our sample because Q values were similar for the cases and the controls. Moreover, the Structure program assigned a similar distance to clusters and no further improvement in the fitting model was achieved by adding up to four clusters (the ln of likelihood was maximum for K = 1).

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