Time-of-day-dependent variation of the human liver transcriptome and metabolome is disrupted in MASLD



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Background & Aims: Liver homeostasis is ensured in part by time-of-day-dependent processes, many of them being paced by the molecular circadian clock. Liver functions are compromised in metabolic dysfunction-associated steatotic liver disease (MASLD) and metabolic dysfunction-associated steatohepatitis (MASH), and clock disruption increases susceptibility to MASLD progression in rodent models. We therefore investigated whether the time-of-day-dependent transcriptome and metabolome are significantly altered in human steatotic and MASH livers.

Methods: Liver biopsies, collected within an 8 h-window from a carefully phenotyped cohort of 290 patients and histologically diagnosed to be either normal, steatotic or MASH hepatic tissues, were analyzed by RNA sequencing and unbiased metabolomic approaches. Time-of-day-dependent gene expression patterns and metabolomes were identified and compared between histologically normal, steatotic and MASH livers.

Results: Herein, we provide a first-of-its-kind report of a daytime-resolved human liver transcriptome-metabolome and associated alterations in MASLD. Transcriptomic analysis showed a robustness of core molecular clock components in steatotic and MASH livers. It also revealed stage-specific, time-of-day-dependent alterations of hundreds of transcripts involved in cell-to-cell communication, intracellular signaling and metabolism. Similarly, rhythmic amino acid and lipid metabolomes were affected in pathological livers. Both TNF α and PPAR γ signaling were predicted as important contributors to altered rhythmicity.

Conclusion: MASLD progression to MASH perturbs time-of-day-dependent processes in human livers, while the differential expression of core molecular clock components is maintained.

Impact and implications: This work characterizes the rhythmic patterns of the transcriptome and metabolome in the human liver. Using a cohort of well-phenotyped patients (n = 290) for whom the time-of-day at biopsy collection was known, we show that time-of-day variations observed in histologically normal livers are gradually perturbed in liver steatosis and metabolic dysfunction-associated steatohepatitis. Importantly, these observations, albeit obtained across a restricted time window, provide further support for preclinical studies demonstrating alterations of rhythmic patterns in diseased livers. On a practical note, this study indicates the importance of considering time-of-day as a critical biological variable which may significantly affect data interpretation in animal and human studies of liver diseases.

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Introduction

Normal tissue homeostasis requires a precisely timed expression of genes and proteins around the clock and its alignment with cycles of light/dark exposure, feeding periods and physical activity.

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The central clock, located in the suprachiasmatic nucleus, is lightentrained and connects with peripheral tissues to synchronize clock oscillators in these tissues. However, peripheral tissue clocks can operate autonomously, *i.e.* independently of the central hypothalamic clock. For example, the major *Zeitgeber* ("time giver") setting the liver clock is food intake/nutrient availability rather than daylight.^{1,2} Studies of molecular mechanisms controlling these circadian regulations in nocturnal rodents generated a global picture defining a universal molecular clock machinery.³ This cell-autonomous circadian core clock is made of two autoregulatory loops comprising 14 transcription factors, encoded by so-called core clock genes (CCGs). Heterodimeric BMAL1 (*ARNTL*)



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and CLOCK (or NPAS2) transcriptional activators and PER and CRY transcriptional repressors, along with the nuclear receptors RORs, REV-ERB α and β constitute interlocked transcriptional-transla tional feedback loops. These interlocked transcriptional-transl ational feedback loops define a cell-autonomous clock machinery which controls clock output genes,⁴ in turn regulating multiple cellular functions.³

Because most primates (including humans) are diurnal, there are likely important differences from rodents in circadian regulation that have yet to be explored. A limited number of human time-of-day-resolved transcriptomes is available, especially for internal organs. Transcriptomes from whole blood,^{5,6} peripheral blood mononuclear cells,⁷ skin,⁸ subcutaneous white adipose tissue,^{9,10} heart,¹¹ or skeletal muscle^{12–14} were analyzed for a relatively low number of individuals (n <30). A gene expression study in subcutaneous white adipose tissue and skin from 625 healthy volunteers enabled the identification of time-of-day-regulated genes strongly enriched in CCGs.¹⁵

Circadian rhythm dyssynchrony is observed in, and likely causative of, various diseases such as obesity and its complications like metabolic dysfunction-associated steatotic liver disease (MASLD), formerly known as non-alcoholic fatty liver disease (NAFLD). MASLD is a spectrum of liver conditions characterized by hepatic steatosis combined with varying degrees of necroinflammation and excluding excessive alcohol consumption.¹⁶ Its more severe, yet generally asymptomatic form, termed metabolic dysfunction-associated steatohepatitis (MASH; formerly known as NASH, non-alcoholic steatohepatitis), may evolve towards liver fibrosis, cirrhosis and hepatocellular carcinoma. Large-scale patient cohort studies have reported differences in gene expression between disease stages without timeof-day information.¹⁷⁻²⁰ While hepatic metabolomes and transcriptomes are deregulated in rodent models of MASH and fibrosis,^{21,22} whether this occurs similarly in human MASLD remains unknown. A 24 h-circadian transcriptome atlas of 64 tissues from healthy baboons identified only a small set of robustly cycling genes in the liver, surprisingly not including CCGs.²³ Rhythmic gene expression patterns were inferred from the analysis of tissues collected post mortem from 600 human donors. Relatively few genes (n = 648), including only a few CCGs, exhibited predicted time-of-day-dependent expression.²⁴ Thus, both ethical and technical hurdles hinder the thorough investigation of time-of-day-dependent processes in healthy human liver and the deregulation thereof in MASLD. Importantly, this conclusion extends to the hepatic metabolome, which is clockcontrolled and disturbed in various liver dysfunction models.^{22,25–28} Disturbances in the hepatic chronometabolome observed in rodent MASLD models have not yet been reported in humans.²⁹

Considering these knowledge gaps, we asked whether the hepatic time-of-day-dependent transcriptome and metabolome are affected during MASLD progression. We leveraged a large cohort of morbidly obese patients undergoing bariatric surgery from whom liver biopsies were taken peri-operatively (Hôpital Universitaire de Lille [HUL] cohort¹⁸). Hepatic transcriptomes and metabolomes were obtained from a sub-cohort of 290 patients whose biopsies were histologically identified as either normal, steatotic, or MASH livers and for whom the exact time-of-day at biopsy was known (Fig. 1A). In an original approach integrating multiple statistical tests, we provide the first-ever robust analysis of time-of-day-dependent gene expression and

tissue metabolite abundance, as well as changes associated with the different stages of MASLD, in human livers.

Materials and methods

Liver biopsies from the HUL cohort

The HUL cohort, also known as the Biological Atlas of Severe Obesity (ABOS) cohort, was established in 2006 by the University Hospital of Lille, France (ClinicialTrials.gov: NCT01129297) and comprised severely and morbidly obese patients visiting the Obesity Surgery Department. The study protocol conforms to the ethical guidelines of the 1975 Declaration of Helsinki, All patients of the cohort fulfilled criteria for, and were willing to undergo, bariatric weight-loss surgery (for details, see¹⁸). Written informed consent was obtained from each patient included in the study. The protocol required that patients fast from midnight until the time of surgery. During the surgical procedure, wedge biopsies were taken from the liver to be immediately snapfrozen and the exact time of the biopsy was noted. A total of >1,500 patients are currently included in the HUL cohort, amongst whom 319 were selected to build a sub-cohort with complete clinical, biometric parameters and a robust histological MASLD classification of quality-controlled biopsies eliminating all intermediary MASLD stages (see¹⁸ and Fig. 1B for more details). Both transcriptomes and metabolomes were obtained for these 319 patients with biopsy mass >100 mg. Out of these, the time-of-day at biopsy (ranging from 8am to 4pm) was known for 290 patients, who were included in this study. The main clinical and histological characteristics are provided in Table 1. Patient clinical data shown in Table 1 were analyzed using the package "gtsummary" (v1.5.0).

Total RNA sequencing and data processing

A detailed procedure can be found in the supplementary materials and methods.

Liver metabolomics by liquid chromatography-mass spectrometry

All tissue samples were flash frozen and maintained at -80 °C until processing. Sample preparation was carried out as described previously³⁰ at Metabolon, Inc. (Morrisville, NC, USA). A detailed procedure can be found in the supplementary materials and methods.

Bioinformatic analysis

Bulk RNA sequencing

All analyses were carried out using RStudio (v1.4.1106) with R (v4.1.0). Data processing for differential expression as a function of time-of-day can be found in the supplementary materials and methods.

Single-cell RNA sequencing

All analysis has been made under R (v 4.2.0). A detailed procedure for data extraction and processing can be found in the supplementary materials and methods.

Enrichment analysis

Time-dependent gene lists were analyzed for enrichment of Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and gene ontology biological process terms using Metascape 3.5³¹ with default settings (https://metascape.org/). A detailed

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Fig. 1. Timed liver biopsies from a large cohort of humans with obesity. (A) Overall experimental strategy. (B) Decision tree to stratify the HUL sub-cohort. "HN" (histologically normal), "steatosis" (benign steatosis only) or "MASH" (steatosis+inflammation). (C) Inter-sample variation in gene expression. ANOVA was used to reveal the main sources of overall inter-sample variation. (D) Biopsy daytime distribution. (E) Gene expression analysis. A volcano plot was generated by comparing gene expression using DEseq2 from samples collected in the morning (AM) or in the afternoon (PM) regardless of the pathological state. X axis: log₂ (fold change), Y axis: -log₁₀(*p* values). (F) KEGG pathway enrichment analysis. Biological term enrichment was carried out using the 1,660 genes whose expression was significantly different as determined in (E) (FC >1.2, FDR <0.05). (E, F) Font size was adjusted for clarity purpose. FC, fold change; FDR, false discovery rate; HN, histologically normal; KEGG, Kyoto Encyclopedia of Genes and Genomes; MASH, metabolic dysfunction-associated steatohepatitis.

procedure for data processing can be found in the supplementary materials and methods.

Data visualization and illustrations

Graphs were generated as *.svg files using R packages mentioned above. Data were imported in CorelDraw2020 to assemble figures. Drawings in Fig. 1A are from Renée Gordon, Victovoi, and Mikael Häggström, M.D. and were made available to the public domain via Wikimedia Commons with no restriction of use. Bubbleplots were generated in R studio using the ggplot2, plotly, reshape2, rcpp, and tidyverse packages as described in.³²

Results

Time-of-day is a major factor affecting gene expression in human liver

Human liver biopsies were collected from patients with obesity and undergoing bariatric surgery, for which the exact daytime of

Table 1. Biometric and biochemica	parameters of the HUL sub-cohort.
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		MASLD group			p values		
Variable	Ν	HN, n = 89	St., n = 122	MASH, n = 79	HN vs. St.	HN vs. MASH	St. vs. MASH
Sex Female Male	290	75/89 (84%) 14/89 (16%)	80/122 (66%) 42/122 (34%)	49/79 (62%) 30/79 (38%)	0.003	0.008	0.7
Age, yr	290	34.6 ± 11.4	41.8 ± 10.9	46.4 ± 10.4	<0.001	< 0.001	0.050
BMI	290	45.4 ± 6.8	47.3 ± 7.9	46.2 ± 7.9	0.2	0.5	0.2
Steatosis score 0 1 2 3	290	89/89 (100%) 0/89 (0%) 0/89 (0%) 0/89 (0%)	0/122 (0%) 0/122 (0%) 71/122 (58%) 51/122 (42%)	0/79 (0%) 18/79 (23%) 29/79 (37%) 32/79 (41%)	<0.001	<0.001	<0.001
Inflammation score 0 1 2	290	89/89 (100%) 0/89 (0%) 0/89 (0%) 0/89 (0%)	70/122 (57%) 44/122 (36%) 8/122 (6.6%) 0/122 (0%)	0/79 (0%) 55/79 (70%) 23/79 (29%) 1/79 (1.3%)	<0.001	<0.001	<0.001
Ballooning score 0 1 2	290	89/89 (100%) 0/89 (0%) 0/89 (0%)	115/122 (94%) 7/122 (5.7%) 0/122 (0%)	0/79 (0%) 57/79 (72%) 22/79 (28%)	0.54	<0.001	<0.001
Fibrosis score (Kleiner) 0 1 2 3 4	282	79/89 (89%) 9/89 (10%) 0/89 (0%) 1/89 (1.1%) 0/89 (0%)	78/118 (66%) 28/118 (24%) 7/118 (5.9%) 5/118 (4.2%) 0/118 (0%)	12/75 (16%) 22/75 (29%) 13/75 (17%) 25/75 (33%) 3/75 (4.0%)	0.009	<0.001	<0.001
NAS score 0 2 3 4 5 6 7 8	290	89/89 (100%) 0/89 (0%) 0/89 (0%) 0/89 (0%) 0/89 (0%) 0/89 (0%) 0/89 (0%)	0/122 (0%) 40/122 (33%) 51/122 (42%) 26/122 (21%) 5/122 (4.1%) 0/122 (0%) 0/122 (0%) 0/122 (0%)	0/79 (0%) 0/79 (0%) 11/79 (14%) 19/79 (24%) 29/79 (37%) 17/79 (22%) 2/79 (2.5%) 1/79 (1.3%)	<0.001	<0.001	<0.001
HOMA-IR	279	3.8 ± 4.8	13.5 ± 58.8	24.2 ± 55.2	< 0.001	< 0.001	< 0.001
Biopsy time	290				0.13	0.3	0.4
AM		62/89 (70%)	70/122 (57%)	46/79 (58%)			
PM		27/89 (30%)	52/122 (43%)	33/79 (42%)			

The main biometric, biochemical and liver histological features of selected patients are indicated. Continuous values are expressed as mean ± SD. Inter-group comparisons were performed using the unpaired Wilcoxon test for continuous variables (age, BMI, HOMA-IR) and Fisher's exact test for the remaining categorical variables. HN, histologically normal; HOMA-IR, homeostatic model assessment for insulin resistance; MASH, metabolic dysfunction-associated steatohepatitis; NAS, NAFLD/MASLD activity score; St., steatosis.

the liver biopsy was recorded (Fig. 1A). Key clinical parameters of the 290 patients are summarized in Table 1. Based on histological features of liver biopsies (steatosis, hepatocyte ballooning, lobular inflammation), patients were grouped according to MASLD stages and labelled as histologically normal (HN), steatotic or MASH liver following the decision tree shown in Fig. 1B. The proportion of men in this cohort increased with MASLD severity, rising from 16% (HN) to 38% (MASH) with an average Clinical Research Network NAFLD activity score rising from 0 to 5, respectively. Since sex is an important biological variable in this context,¹⁸ this was considered during further analysis (see below). Patients in the steatosis and MASH groups were slightly older than those in the HN group, and expectedly also had higher insulin resistance on average.

The source of variation in gene expression levels was estimated by a multivariate analysis of variance (Fig. 1C). The F-ratio (ratio of the between-group variance to the within-group variance) not only confirmed sex and group (*i.e.* MASLD stage) as the main sources of variation as previously reported,¹⁸ but very interestingly identified biopsy time (AM vs. PM) as the third most significant source of variation (Fig. 1C). Age made only a minor contribution to signal variation. The distribution of biopsy times (Fig. 1D) revealed a daytime window of about 8 h. Biopsies were predominantly (>60%) taken in the morning with a first peak around 9:30 AM and a second, lower peak around 2:30 PM, due to the logistical schedule of surgical interventions. There was, however, no significant difference in biopsy time distribution between histological groups (Table 1). Since exclusion of biopsies collected between 11am and 1pm did not modify the outcomes of preliminary statistical analysis, the "AM" subgroup was defined as biopsies taken after noon.

Gene expression profiles were thus compared between morning (AM) vs. afternoon (PM) samples. Differentially expressed genes were identified using DEseq2 independent of the MASLD status but correcting for sex. Transcript counts for 1,660 genes were significantly different (Benjamini-Hochberg adjusted *p* value <0.05) between AM or PM biopsies (Fig. 1E). Among the 100 top hits were most of the CCGs (*PER3, ARNTL/ BMAL1, NPAS2, NR1D1, NR1D2, PER2, CRY1, PER1*), clock-related genes (*CIART, DBP, NFIL3*) (Fig. 1E) as well as circadian-regulated genes involved in lipid metabolism (*PPARD, LIPG, LPIN2 ...*). Because patients were fasting from midnight irrespective of surgery time, genes implicated in hepatic gluconeogenesis (*G6PC, PCK1, SGK2 ...*) were, as expected, expressed at a higher level in PM samples (Fig. 1E and Table S1). Contrary to nocturnal rodents, genes from the negative limb of the clock displayed lower expression in the afternoon (*PER3, NR1D1, NR1D2, CIART ...*), whereas genes from the positive clock limb displayed higher expression in the afternoon (*ARNTL/BMAL1, NPAS2 ...*) (Fig. 1E). Globally, genes displaying AM vs. PM differential expression were significantly enriched for the KEGG pathways "circadian rhythm", "PPAR signaling pathway", carbohydrate and lipid metabolic pathways, as well as cellular architecture and communication, among others (Fig. 1F).

Thus, an 8-hour time frame allowed for the detection of significant changes in time-of-day-dependent liver gene expression, with a large proportion of transcripts functionally related to circadian rhythmicity.

Time-dependent genes vary between MASLD stages

We next examined whether time-of-day-dependent distributions of gene expression would differ between the histological states "HN", "steatosis" and "MASH". In order to achieve statistical power and obtain robust and exhaustive lists of timedependent gene expression over the available daytime window, we used three complementary statistical methods analyzing different aspects of gene expression distribution (differential expression, partial Spearman correlation, Kolmogorov-Smirnov test), the results of which were agglomerated by a Fisher test to yield a combined *p* value for each gene (Fig. 2A-C and Figs S1 and S2). The three types of analyses are graphically exemplified for the CCG ARNTL/BMAL1 (Fig. 2A-C), which served as a positive control to validate our approach, as it is among the most highly time-dependent genes regardless of the histological group. First DEseq2, which relies on a negative binomial distribution of gene expression, was used to identify differential gene expression between two conditions (AM vs. PM as in Fig. 1) (Fig. 2A). Second, partial Spearman correlation was computed between ARNTL gene expression and time-of-day (Fig. 2B). Both approaches integrated sex as a confounding factor. Third, the Kolmogorov-Smirnov test was employed to determine whether AM and PM ARNTL expression distribution followed a similar law and thus were similar in shape (Fig. 2C). Finally, the Fisher combined probability test or "Fisher's method" was used as a meta-analysis method for *p* value combination: individual raw *p* values resulting from each statistical test were agglomerated into a single *p* value per group (Fig. 2D). The detected expression profile of ARNTL, of other CCGs (Fig. S1) and of all other transcripts (Fig. S2), clearly confirmed that the available time window was sufficient for robust time-of-day analysis of gene expression. A total of 1,427 genes with an absolute fold change greater than 1.2 (AM vs. PM) (false discovery rate [FDR] <0.01) was identified (Fig. 2E). The vast majority of these timedependent genes (TDGs) were strikingly distinct when comparing the three patient groups. Less than 10% (132 genes) were indeed common to all three groups ("common TDGs") (Fig. 2E) and notably included most CCGs (ARNTL, NR1D1/2, NPAS2, CRY1, PER1/2/3, DBP, CIART) (Table S1). TDG repartition outside of this core set was strongly unequal between groups, with $\approx 50\%$ (558) of non-shared TDGs found in HN, $\approx 35\%$ (392) in steatotic and less than 15% (177) in MASH livers. Along the

same line, we found that AM to PM fold changes of common TDGs were, on average, decreased in steatotic and even more in MASH livers when compared to HN livers (Fig. 2F). These differences are illustrated for a selection of TDGs with AM-PM differences either decreasing (*PCAT18, ARMC4, SIK1B*) or increasing in MASH (*CYP4Z1, RHOBTB1, PPIAP71*) (Fig. 3A,B).

Common TDGs were collectively enriched for KEGG terms like "circadian rhythms" as expected from the content in transcripts coding for CCGs, and for metabolic regulatory pathways like the PPAR and FoxO pathways (Fig. 4, Fig. S3), illustrated by genes such as S1PR1, G6PC1, PCK1, SGK2, FASN, AQP7 and PPARD. TDGs unique to the HN group were also enriched, albeit to a lesser extent, for pathways linked to circadian rhythm (notably including CLOCK) as well as to fatty acid and amino acid metabolism. The most highly represented pathway was "gap junctions" (Figs 4A and S3), characterized by genes such as PDGFB. MAP2K1 and transcripts encoding for tubulins TUBA1C/8, TUBB, TUBB1/2B (Table S1), suggesting that homeostasis of epithelial barrier integrity/permeability, which is known to be disturbed in MASLD,³³ requires an oscillating expression of these genes in healthy conditions. Genes unique to the steatosis group (Table S1) were mostly linked to metabolism of lipids and fatty acids (DGKG, PLA2G4B/5, LPIN2/3, ETNK2, ETNPPL, PLPP4, FADS1/2, CYP2C8. GDPD1. SCD) and also to metabolism of peptides and amino acids (DNMT3B, GCLM, SDS, PSAT1, GNMT, ALDOC, GPT2, CSAD, UPB1) (Figs 4A and S3). Lastly, TDGs specific to the MASH group (Table S1) were highly enriched for signaling by calcium, cAMP or neurotransmitters (ADRB2, DRD1, GRM1, NTRK1, NTSR1, P2RX7, RYR2, CACNA1H, SSTR5, TBXA2R, FFAR2, SUCNR1), as well as for lipolysis (ADRB2, IRS1, PNPLA2) (Figs 4A and S3). The temporal pattern of gene expression in homeostatic conditions is thus strongly affected by the disease state and indicative of compromised cellular communication and metabolic pathways.

Inferring upstream regulatory cues or altered biological processes may be achieved by comparing differentially expressed gene lists to consensus gene expression patterns induced by a given perturbagen (Fig. 4B-D). Speed2 (Signaling Pathway Enrichment using Experimental Datasets³⁴) analysis enables gene lists to be probed against ranked gene signatures for 16 signaling pathways, with the aim of identifying upstream signaling mediators. Ranked signatures suggested that cues in homeostatic (HN) conditions could be TGFB, TNFa, oxidative stress, TLR and estrogen (Fig. 4B). In steatosis and MASH conditions, this pattern shifted towards a more limited signaling pathway panel with similar statistical significance, which included either TLR and VEGF (steatosis) or TNF α and TLR (MASH) (Fig. 4C,D, respectively). Although causative links cannot be proven, this data could reflect a loss of physiological rhythmic function(s) in steatotic and MASH livers, which in turn gain rhythmic functions associated to pathogenic immune and proliferative stimuli and responses.

MASLD stages correlates with time-of-day changes in liver metabolites

Our results suggested that metabolic pathways are altered in a time-of-day-dependent manner as MASLD progresses, with an enrichment in amino acid- and lipid metabolism-regulating genes (Fig. S3). Therefore, an unbiased tissue metabolomic study by liquid chromatography-mass spectrometry was performed on the same 290 liver samples. Similar to the gene expression analysis, a global approach was initially employed to evaluate overall time-of-day dependence of tissue metabolite

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Fig. 2. A multi-test method identifies time-of-day-dependent genes. *BMAL1/ARNTL* expression as a function of time and of the liver histological grade (A-C). (A) A violin plot is shown to illustrate DEseq2 results to compare AM vs. PM expression and correcting for sex as a confounding factor. (B) A dot plot with linear tendency lines is shown to illustrate partial Spearman correlation analysis between gene expression and biopsy daytime. (C) A density plot is shown to illustrate the output of the Kolmogorov-Smirnov test for comparison of AM or PM gene expression distributions. (D) Fisher's agglomeration method. Frequency histograms of uncorrected Fisher *p* values are shown for each group and the first (colored) bar of each group histogram indicates genes with *p* <0.05. (E) TDG distribution within histological groups. A Venn diagram shows the unequal distribution of 1,427 TDGs (FC >1.2, FDR <0.01) amongst groups. Numbers indicate the numbers of identified transcripts. (F) Gene expression variation of 132 "common TDGs". CCGs and other transcripts are indicated. *Indicates *p* <0.05 using ANOVA and Fisher's LSD *post hoc* test. CCGs, core clock genes; FC, fold change; FDR, false discovery rate; TDGs, time-dependent genes.

levels regardless of the MASLD status. This global analysis identified \approx 220 metabolites whose amounts were significantly different in AM and PM biopsies (DEseq2 corrected for sex, FDR <0.1) (Fig. 5A, Table S2). Visual inspection of the volcano plot highlighted intermediates of lipid β -oxidation (carnitine derivatives), amino acids (kynurenate, oxo-arginine ...) as differentially detected in AM *vs.* PM livers (Fig. 5A). It also confirmed the more marked fasting status of "PM" patients exhibiting an increased hepatic content in 3-hydroxybutyrate. A biological term enrichment analysis confirmed that the majority of the

identified metabolites belonged to amino acid, lipid and fatty acid metabolic pathways (Fig. 5B).

To highlight a possible time-of-day differential representation of metabolites between MASLD groups, we again combined the three statistical approaches as described for gene expression analysis (DEseq2, Spearman correlation, Kolmogorov-Smirnov test) followed by Fisher's agglomeration for a robust identification of time-dependent metabolites (TDMs) (Fig. 5C). A total of 251 TDMs were identified using this method (combined FDR <0.1), out of which only 14 (6%) were common to all three MASLD

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Fig. 3. Example distributions of TDGs altered in MASH. (A) Violin plots showing AM vs PM gene expression variations for representative TDGs losing their timedependency in MASH. (B) Violin plots showing AM vs. PM gene expression variations for representative TDGs gaining time-dependency in MASH. Statistical tests were Kruskal-Wallis tests followed by unpaired Wilcoxon *post hoc* test for AM-PM comparisons in each group (**p* <0.05, ***p* <0.01, ****p* <0.005, *****p* <0.001). MASH, metabolic dysfunction-associated steatohepatitis; TDGs, time-dependent genes; VST, variance-stabilizing transformation.

groups (Fig. 6A,B). These common TDMs included amino acids such as proline and threonine, several fatty acids and the ketone body component β -hydroxybutyrate (Figs 6B and S4A-F). KEGG metabolic pathway enrichment analysis revealed that these common TDMs were most significantly associated with the metabolism of amino acids (arginine, proline, threonine) (Figs 6B and S4A-C). In agreement with the identification of the PPAR pathway based on gene expression patterns (Fig. 4) and the detection of 3-hydroxybutyrate (Fig. S4D), synthesis of ketone bodies was also identified as a relevant term (Fig. 6B).

Among the 197 stage-specific TDMs, 31% were specific to HN, 47% to steatosis and 20% to MASH (Fig. 6A and Table S2). TDMs specific to HN livers, and thus lost at the steatosis and MASH stages, were mainly associated with the metabolism of sphin-golipids (Fig. 6C and Table S2) such as CDP-choline and sphin-ganine (Fig. S4G, H). Glycerophosphoethanolamines, glycerophosphocholines as well as derivatives of cholesterol, amino acids and pyrimidine were also identified as time-dependent in normal livers (Table S2 and Fig. S4I-L).

A similar enrichment analysis identified amino acid metabolic pathways (branched-chain, sulfur-containing, arginine, taurine) (Figs 6D and S5A,B and Table S2) as time-dependent in steatotic livers. Visual inspection of steatosis TDMs also identified a carnitine precursor (N6,N6,N6 trimethyl-lysine, Fig. S5C) and derivatives (Table S2 and Fig. S5D,E) which could reflect altered fatty acid oxidation activity. Finally, MASH-specific TDMs were enriched mainly for vitamin, glycan and glycosylphosphatidyl inositol metabolic intermediates (Figs 6E and S5, and Table S2).

Taken together, these analyses highlight the disruption of time-of-day-dependent bioactive phospholipid metabolism and of amino acid biotransformation pathways during MASLD progression. Intriguingly, PPAR γ ligands of the linoleic acid class (9,10-dihydroxy-9-octadecenoic acid [DiHOME]³⁵ and 9- and 13-hydroxyoctadecadienoic acid [HODE]³⁶) displayed a differential abundance in AM *vs.* PM steatotic and MASH livers, with estimated concentrations in the 10-100 μ M range which are sufficient to activate PPAR γ (Fig. S5F,J).

Integrative analysis of time-dependent genes and metabolites We next performed an integrative analysis of the transcriptomic and metabolomic data at the pathway level using the KEGG database. This analysis combined TDGs and TDMs specific to either HN or MASH stages and common TDGs and TDMs, irrespective of their relative time-of-day direction of change, to identify associated transcriptomic and metabolomic conditions operating in normal and MASH livers (Fig. 7A). TDGs and TDMs characterizing the HN stage were enriched for metabolic pathways related to lipid and amino acid metabolism, while most of them were not detected at the MASH stage, or with a decreased

significance (arginine [Arg] and proline [Pro] metabolism,

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Fig. 4. Term enrichment analysis of TDGs and upstream regulatory pathway prediction. (A) Biological term enrichment analysis. TDGs identified in Fig. 2 were enriched for GO terms related to BP using Metascape. Significantly enriched GO-BP clusters were manually collapsed for visualization purposes and top hits are indicated. (B-D) Pathway activity ranking. Speed2 TDG enrichment for pathway signature genes. Each pathway is represented as a bar showing the mean rank of the query list. The "bar code" plot shows the distribution of genes from the query list in the ranked reference signatures. BP, biological process; GO, gene ontology; TDGs, time-dependent genes.

glycerophospholipid metabolism). Linoleic metabolism was associated with the MASH stage (Fig. 7B,C). HN- or MASHenriched pathways (glycerophospholipid and linoleic pathways, respectively) were further probed for daytime variation of associated TDGs and TDMs. The glycerophospholipid pathway was characterized by an increased abundance in AM livers of three out of four detected diacylglycerol (DAG) species specifically at the HN stage. Glycerophosphocholine (GPC) intermediates (CDP-choline, GPC) displayed stage-specific time-ofday variations that were not correlated to the occurrence of glycerophospholipid species (X-GPC) (Fig. 7D). Higher abundance of DAG species in the morning did not correlate with HN TDG expression changes in transcripts encoding enzymes involved in this metabolic pathway, with the exception of the PNPLA3 gene. PNPLA3/adiponutrin has acyltransferase activity, increasing the formation of phosphatidic acid from lysophosphatidic acid, which may lead to more DAG synthesis. It might

also reflect the hydrolytic activity of PNPLA3 on triacylglycerol molecules, favoring DAG accumulation. Along the same line, we compiled TDG and TDM data related to linoleic acid metabolism (Fig. 7E). The increased abundance of the PPAR γ ligands 9- and 13-HODE in the afternoon at the MASH stage was mirrored by the gene expression of *ALOX15*, a dioxygenase catalyzing the synthesis of these two hydroxyoctadecadienoic acids which was higher in the morning. A similar lack of correlation was observed between 9,10-DiHOME hepatic content and the expression of the linoleic acid-converting *CYP2C8* at the steatosis stage. These results suggest that time-dependent metabolite variation in these pathways is delayed with respect to gene regulation, and/or controlled by post-transcriptional processes.

Of note, mapping of human transcript expression to liver cell types using a reference single-cell RNA sequencing dataset³⁷ suggested that the identified enzymatic pathways may follow a cell-specific expression pattern. They appear mainly restricted,

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Fig. 5. Identification of TDMs measured by liquid chromatography-mass spectrometry. (A) Volcano plot of metabolite time-of-day-dependent differential abundance in liver. Fold change values and corresponding *p* values obtained using DEseq2 considering sex as a confounding factor. (B) Enrichment of KEGG metabolic pathways. A biological term enrichment against the KEGG database was run using the differentially detected metabolites (TDMs) identified in (A) (DEseq2 adjusted *p* value <0.1). (C) *p* value agglomeration by Fisher's method. As for gene expression analysis (see Fig. 2), *p* value agglomeration from three separate statistical tests (DEseq2, partial Spearman correlation, Kolmogorov-Smirnov) was carried out using Fisher's method and identified TDMs within each group. The resulting uncorrected Fisher *p* values are shown as frequency histograms and the first (colored) bar of each group histogram indicates metabolites with *p* <0.1. TDMs, time-dependent metabolites.

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Fig. 6. Distribution and characterization of TDMs. After correction for multiple testing, 238 metabolites with a Fisher FDR<0.1 were considered as robustly time-dependent. (A) Distribution of TDMs among liver histological groups. (B-E) KEGG metabolic pathway enrichment of common (central intersection) or group-specific TDMs using the online metabolomics analysis platform MetaboAnalyst. FDR, false discovery rate; TDMs, time-dependent metabolites.

but not limited to, hepatocytes. As an example, *ALOX15* is detected in dendritic cells, whereas *ALOX5* is also detected in monocytes, neutrophils and basophils (Fig. S6). Therefore time-dependent metabolite variation may occur in either identical or distinct cell types, reflecting a functional compartmentalization. Yet these observations confirmed the presence of time-of day variation in hepatic gene expression and the hepatic

metabolome, and identified several new oscillating metabolites at the MASH stage (Fig. 7 and Table S2).

Discussion

Chronobiological studies require multiple replicates at 2-hour intervals over a total period of 24 or even 48 h, within a contr

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olled environment including timed exposure to light and food. These conditions are not achievable in human studies, precluding the analysis of cyclic processes and particularly in internal organs.³⁸ While human circadian rhythms are appreciated by genome-wide association studies and the phenotypic manifestation of disturbed cyclic processes such as sleep, light exposure and eating patterns,³⁹ their study in healthy or pathological conditions is indeed hindered by ethical and technical constraints. Despite controlled experimental setup with regard to sleep behavior and food intake in previous studies, ^{5–14} a limited number of samples of mostly healthy individuals were collected, thereby limiting data interpretation due to high inter-sample variability.

Herein, we report the first-ever robust analysis of the time-ofday-resolved human liver transcriptome and associated liver tissue metabolites, using a 290-patient cohort. Although the available time window of liver biopsies was only about 8 h. this temporal window was sufficient to robustly identify TDGs and TDMs. Time-dependency of genes and metabolites was distinct between histologically defined MASLD groups. However, a small proportion of genes was identified as time-dependent in all three patient groups and included CCGs, indicating that the molecular clock is rather robust in pathological conditions. In contrast, the alignment of rhythmic biological processes such as intercellular communication (gap junctions) and metabolic regulations is disrupted upon MASLD progression. Interestingly, none of the detected TDGs in patients with MASH belonged to the human and mouse core set of MASH/fibrosis-associated genes,²⁰ underlining the need to consider time as an important biological variable. Interestingly, the number of stage-specific TDGs (Fig. 2E) decreases from the HN to the MASH stage, and concomitantly enriched pathways lessen (Fig. S3), hinting at a loss of functional adaptability/(metabolic) flexibility. In high-fat diet-fed mice, transcripts gaining rhythmicity when compared to chow diet-fed mice are strongly enriched for glycerophospholid metabolism,²² as in steatotic patients (Fig. S3), indicating convergent mechanisms for liver adaptation to dietary imbalance as often occurring in MASLD. At the cellular level, Ca²⁺ fluxes are submitted to ultradian variations and coupled to metabolic regulations,⁴⁰ and mishandled intracellular Ca²⁺ stores in MASH can significantly impact parenchymal and nonparenchymal liver cellular functions.⁴¹ A number of genes encoding for Ca²⁺ channel components or involved in intracellular Ca²⁺ signaling (PKD1L1, TRPC1 P2RX7, FFAR2, ADRB2, CAC-NA1H, GRM1, NTRK1) exhibited differential AM vs. PM expression specifically in MASH livers. Thus, in addition to the lipid-induced dysfunction of endoplasmic reticulum Ca²⁺ transport,⁴² de novo oscillation of the calcium handling process accompanies progression to MASH. CLOCK gene deletion affected metabolite oscillations in mouse livers²⁵ and we detected time-of-daydependent differential expression of CLOCK exclusively in HN livers, which display a more diverse metabolic activity than MASH livers (Fig. 7B). While hinting at a possible role of (the loss of) CLOCK, the examination of individual metabolites nevertheless showed little overlap between mouse CLOCK-dependent metabolites and HN-specific metabolites. This lack of clear concordance can be explained by species-specific mechanisms, distinct effects of gene deletion vs. loss of time-dependent expression and/or technical bias.

While examining the coherence between MASLD state-specific liver transcriptomes and metabolomes, we observed little correlation between enzyme-encoding genes and metabolite abundance. This disconnection is not unprecedented and was also observed in a highly standardized mouse study which minimizes the variability typically observed in human samples.²² The narrow time window of our study may explain in part this lack of correlation as transcripts are likely to precede metabolite production, hence affecting our statistical approach for the PM sub-cohort. It may also indicate significant time-of-day-dependent translational control⁴³ as well as post-translational modifications regulating enzyme activity which are not captured by our analysis.

Finally, another point of convergence between our study and mouse studies is the differential abundance of linoleic acid derivatives and PPAR_{γ} ligands 9,10-DiHOME, 9-HODE and 13-HODE in steatotic and MASH livers, respectively. The PPAR_{γ}-encoding gene *NR1C3/PPARG* itself did not display significant oscillatory expression in the human liver, contrasting with high-fat diet-fed mouse livers.²² Whether the estimated concentrations of these compounds are indeed sufficient to differentially activate human liver PPAR_{γ}, which is mostly expressed in endothelial cells, hepatocytes and macrophages (Fig. S6), and play a causative role in hepatic transcriptional reprograming requires further in-depth investigation.

On the one hand, it is remarkable that many of the biological processes and pathways shown to be affected by MASLD in other studies, identified on the basis of changes in gene expression or metabolite abundance levels regardless of time, also display altered time-dependent expression profiles in our study. On the other hand, we identified many novel potential links between genes with deregulated timed expression and MASLD pathogenesis, which were not previously considered by standard analyses. It is probably the combination of both types of deregulations that underlies the deeply disturbed liver functions once MASH is declared. Conversely, some of the changes detected when time-of-day information is absent or ignored may turn out to be artefacts, as time-of-day as a biological variable might not be equilibrated between groups.

Considering that the time-of-day dependence of transcript/ protein/metabolite measurements was previously neglected or ignored in nearly all human MASLD studies, our findings reveal a significant impact of time-of-day on many relevant pathogenic processes. As such, differences found between sample groups in cohort studies might reflect a previously under-appreciated bias in sampling time between groups. Further investigation is needed in this regard, particularly when studying human material. In any case, a new generality should be that sampling daytimes (or *Zeitgeber* times) must be carefully recorded and included in *post hoc* analyses whenever possible.

This study has both strengths and limitations. This first-of-itskind study revealed time-of-day transcriptomic and metabolomic alterations in human livers as a function of the histologically proven MASLD stage. It used a large cohort allowing both the selection of biopsies to adequately encompass healthy control, steatosis and MASH cases and robust statistical analysis. There are however a number of limitations inherent to the observational nature of the study. The narrow time window for biopsy collection (8 h) precludes the assessment of a 24 h diurnal rhythmicity and hence of the integrity of the molecular clock. The unavoidable difference in fasting duration could also be a confounding factor.

Our bulk RNA sequencing approach provided full coverage of the transcriptome, but precluded the identification of liver cell types expressing TDGs. We thus calculated a cell specificity index of TDGs in human resident parenchymal (PC, hepatocytes) and non-parenchymal (NPC) CD45⁻ cells. The tau (τ) index, calculated by a robust and simple method to assess cell type specificity, was used as a metric.⁴⁴ Using a reference single-cell RNA sequencing dataset for human livers collected at unknown times,³⁷ τ could be calculated for 584 transcripts out of the 1,481 identified TDGs (Table S1). Using this metric, only 51 genes displayed a cell typerestricted expression pattern in CD45⁻ cells (τ >0.85), which was not limited to hepatocytes (Fig. S7). The remaining 533

Abbreviations

CCGs, Core clock genes; DAG, Diacylglycerol; DiHOME, Dihydroxy-9octadecenoic acid; FDR, False discovery rate; GPC, Glycerophosphocholine; HODE, Hydroxyoctadecadienoic acid; HUL, Hôpital universitaire de lille; KEGG, Kyoto encyclopedia of genes and genomes; MASLD, Metabolic dysfunction-associated steatotic liver disease; MASH, Metabolic dysfunction-associated steatohepatitis; TDGs, Time-of-day-dependent genes; TDMs, Time-of-day-dependent metabolites.

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Conflicts of interest

The authors of this study declare that they do not have any conflict of interest.

Please refer to the accompanying ICMJE disclosure forms for further details.

Authors' contributions

Conceptualization: MJ, JTH, JV, FP, BS, PL; Software: MJ, JTH, JV, MD; Validation: MJ, JTH, JV, BD, VR, JE, BS, PL; Formal analysis: MJ, JTH, JV, JE, PL; Investigation: MJ, JTH, JV, BD, MD, AB; Resources: AB, PF, FP, BS, PL; Data curation: MJ, JTH, JV, BD, MD, VR, PL; Writing: MJ, JTH, JV, BS, PL; Visualization: MJ, JTH, JV, PL; Supervision: BS, PL; Project administration: BS, PL; Funding acquisition: FP, BS, PL, BD, VR.

Data availability statement

Further information and requests for resources and data sets should be directed to and will be fulfilled by the corresponding author. Data exploration of the HUL cohort is still ongoing and restrictions apply to clinical data availability.

Disclaimer

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Supplementary data

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transcripts including CCGs could be mapped to two or more cell types. Of note, 15 out of the 51 cell-restricted transcripts were mostly expressed in dendritic or Kupffer cells (Fig. S6).

Taken together, our bulk RNA sequencing approach detected a higher number of TDGs than previous efforts, including many pathways never previously reported as being time-dependent. This resource will serve as an important basis for further investigations that also consider the cell type specificity in time-ofday gene expression variation in human MASLD.

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