Plasma triacylglycerols are biomarkers of β-cell function in mice and humans

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ABSTRACT

Objectives: To find plasma biomarkers prognostic of type 2 diabetes, which could also inform on pancreatic β-cell deregulations or defects in the function of insulin target tissues.

Methods: We conducted a systems biology approach to characterize the plasma lipidomes of C57Bl/6J, DBA/2J, and BALB/cJ mice under different nutritional conditions, as well as their pancreatic islet and liver transcriptomes. We searched for correlations between plasma lipids and tissue gene expression modules.

Results: We identified strong correlation between plasma triacylglycerols (TAGs) and islet gene modules that comprise key regulators of glucose- and lipid-regulated insulin secretion and of the insulin signaling pathway, the two top hits were Gck and Abhd6 for negative and positive correlations, respectively. Correlations were also found between sphingomyelins and islet gene modules that overlapped in part with the gene modules correlated with TAGs. In the liver, the gene module most strongly correlated with plasma TAGs was enriched in mRNAs encoding fatty acid and carnitine transporters as well as multiple enzymes of the β-oxidation pathway. In humans, plasma TAGs also correlated with the expression of several of the same key regulators of insulin secretion and the insulin signaling pathway identified in mice. This cross-species comparative analysis further led to the identification of PITPNC1 as a candidate regulator of glucose-stimulated insulin secretion.

Conclusion: TAGs emerge as biomarkers of a liver-to-β-cell axis that links hepatic β-oxidation to β-cell functional mass and insulin secretion.

Keywords Triacylglycerols; β-cell function; Systems biology; Type 2 diabetes; PITPNC1; Biomarkers

1. INTRODUCTION

Type 2 diabetes (T2D) is a chronic hyperglycemic condition characterized by reduced glucose-stimulated insulin secretion (GSIS) and increased insulin resistance of insulin target tissues [1,2]. Multiple initiating pathogenic mechanisms can affect any tissue primarily involved in glucose homeostasis, including the pancreatic islet β-cells, liver, muscle, adipose tissue, or the autonomic nervous system [3]. The deregulations in insulin secretion or in insulin action then lead to defects in inter-organ communications causing imbalanced glucose homeostasis, thus T2D. For developing better prevention strategies or treatment options, identification of the primary tissue defects that underlie the appearance of T2D in individual patients will be very useful. The presence of myriads of metabolites in the plasma represents a potential source of information about the function and deregulation of individual tissues [3,4]. Thus, their quantitative measurements could, in principle, inform the deregulations of specific tissue metabolic pathways. In addition, as circulating metabolites can modulate the function of various cell types, their identification could help define modes of interorgan communication that regulate glucose homeostasis.
In our previous study, we initiated a search for circulating biomarkers candidates for the susceptibility to T2D using preclinical models followed by replication of the mouse data in two pre-diabetes cohorts [5]. We first analyzed the correlation between plasma lipids and various glucose homeostasis phenotypes in mice from different genetic backgrounds and fed regular chow or a high-fat diet (HFD) for different periods. This analysis led to the identification of dihydroceramides as candidate biomarkers for T2D development. In human cohorts, we found that the same plasma dihydroceramides were elevated in the plasma of individuals who would develop T2D up to nine years later. Thus, preclinical studies in mice can provide valuable information to identify novel human T2D prognostic biomarkers.

In the present study, we attempted to further exploit this experimental mouse paradigm to investigate whether plasma lipid biomarkers could be identified that predict the function of pancreatic β-cells and whether we could identify tissue metabolic pathways involved in the regulation of the plasma concentrations of these lipids. Thus, we performed a new set of experiments using mice from three different genetic backgrounds fed for different periods with regular chow (RC) or HFD and searched for a correlation between plasma lipids and islets and liver gene co-expression modules. This analysis revealed strong positive and negative correlations between plasma TAGs and islets gene modules that comprise major regulators of insulin secretion such as glucokinase and the K<sub>ATP</sub> channel or mRNAs encoding lipid metabolic enzymes that modulate GSIS. In the liver, we found that plasma TAGs showed a strong correlation with a gene module enriched in fatty acid β-oxidation genes. Together our data indicate that plasma TAGs reflect the activity of a liver-to-β-cell axis that may regulate β-cell function. Similar observations were made in humans linking plasma TAGs and islet genes controlling β-cell mass and function.

2. METHODS

2.1. Mouse phenotyping

Eight-week-old male C57Bl/6J, DBA/2J, and BALB/cJ mice were fed ad libitum with a high fat, high sucrose diet (SAFE 235F, with 46% fat expressed in Kcal/kg) or a regular diet (SAFE A04). Oral glucose tolerance tests (OGTT, 0.5 UI/kg) were performed in five-hour fasted mice on days 2, 10, and 30, as described by Cruciani-Guglielmacci et al. [6]. Analysis of the ITTs revealed basal glycaemia, using a glucometer (A. Menarini Diagnostics, France), and insulin resistance calculated as the area under the curve of glycaemia (AUC; mg/dL* t) measured at 0, 15, 30, 45, 60, 90, and 120 min after insulin administration. From the OGTTs we also obtained basal insulinemia, using an Ultra-Sensitive Mouse Insulin ELISA Kit (Crystal Chem Inc., #90080); stimulated insulinemia calculated as the AUC of insulinemia (mg/mL* t) measured at 0, 15, and 90 min after insulin administration, and glucose intolerance calculated as the AUC of the glycaemia (mg/ dL* t) measured at 0, 15, 30, 45, 60, 90, and 120 min after glucose administration. The number of mice used in these phenotyping experiments ranged between 185 and 195.

2.2. Quantitative lipidomics analysis

Plasma lipid concentrations were measured by mass spectrometry at the Lipotype shotgun lipidomics platform. Samples processing, lipid extraction, spectra acquisition, data processing, and normalization were as described in Sumra et al., 2015 [7]. A principal component analysis (PCA) was performed using the ‘prcomp’ function in R package.

2.3. RNA-Seq and bioinformatics analysis

Complementary DNA (cDNA) libraries were prepared from RNA isolated from mice tissues using the Illumina TruSeq protocol. RNA-Seq was performed on the Illumina HiSeq platform to generate ~40Mio 125 nt single-end reads per sample. Reads were mapped and quantified with STAR-2.5.3a software [6] using M.musculus-mm10 as reference genome and GRCh38.83 from ENSEMBL as the reference annotation index. For each sample, quality control included verification of the total number of reads, percent of uniquely mapped reads, number of detected expressed genes, gene body coverage, and cumulative gene diversity. The resulting counts per gene from different samples were integrated to construct a single count matrix for each tissue that was filtered, excluding those genes with <1 count per million with ‘edgeR’ [9]. We excluded three clear outliers identified by PCA and hierarchical clustering in the islets data set. The count matrix was normalized using the trimmed mean (TMM) normalization method. Differentially expressed genes (DEGs) comparing HF and RC, and the different strains were detected using the Limma package in R [10]. P-values were adjusted for multiple comparisons with the Benjamini-Hochberg procedure, and those genes with adjusted p-value of <0.05 were considered as differentially expressed.

Weighted gene correlation network analysis (WGCNA) was performed on the RNA-Seq dataset from all time points, mouse strains, and diets to generate modules of co-expressed genes. Co-expression networks for each tissue were constructed by calculating signed adjacency matrices using a soft-thresholding power of six and a pair-wise Pearson correlation among all genes. A signed topological overlap matrix (TOM) was then calculated from each adjacency matrix, converted to distances, and clustered by hierarchical clustering using average linkage clustering. Modules were identified in the resulting dendrogram by the Dynamic Hybrid tree cut with a cut height of 0.995 and a minimum module size of 20 genes. The correlation between the resulting modules and plasma lipids was calculated in two steps for the liver data set, and the three steps for the islets data set are as follows: first, a PCA was calculated for each module in each data set using only module constituent genes. Second, the Spearman correlation coefficient was calculated between the summarized values of expression of each module (the first principal component or eigenvalue) and plasma lipids. In the islets data set, an intermediate step was added because plasma lipidomics and islet transcriptomics data were derived from different mouse individuals and direct correlation was not possible. Thus, 18 mouse groups were defined by the three strains, two diets, and three-time points of harvesting. Module eigenvales and plasma lipid concentrations were summarized per mouse group using the mean, which yielded 18 pairs of values to be considered per lipid and module. Student asymptotic p-values were calculated for the given correlations and were adjusted for multiple comparisons using the Benjamini-Hochberg procedure [11]. The functional enrichment analysis was performed using the R library “clusterProfiler” [12]. Pathways from KEGG and GO databases were searched using a hypergeometric test to examine the over-representation of the terms within the functional annotation and to determine the p-values using the hypergeometric distribution. P-values were adjusted for multiple comparisons by the Benjamini-Hochberg procedure. Terms with an adjusted p-value ≤0.05 were considered as overrepresented.

2.4. Correlation analysis in partially pancreatectomised patients

The correlation analysis was performed with 60 samples that included T2D, impaired glucose tolerance (IGT), pancreaticogenic diabetes
(T3cD), and control patients (18, 21, 16, and 4 cases respectively). Pairwise correlations between the pancreatic islets transcriptome and TAGs plasmatic concentrations were performed with Spearman correlation coefficients. For the correlations, Student asymptotic p-values were calculated.

2.5. Cell culture
EndoC-βH1 cells [13] were cultured in 5.6 mM glucose Dulbecco’s Modified Eagle’s Medium (DMEM, Thermo Fisher Scientific), supplemented with 2% BSA fraction V (Roche-Diagnostics), 50 μM β-mercaptoethanol (Sigma–Aldrich), 5.5 μg/mL transferrin (Sigma–Aldrich), 6.7 ng/mL sodium selenite (Sigma–Aldrich), 10 mM nicotinamide (Calbiochem), 100 μg/mL streptomycin and 100 U/mL penicillin (Thermo Fisher Scientific). Cells were seeded on plates coated with 1.2% Matrigel/3 μg/mL fibronectin (Sigma–Aldrich), and cultured at 37 °C and in 5% CO₂.

MIN6B1 cells [14] were cultured in 25 mM glucose DMEM (Thermo Fisher Scientific), supplemented with 15% heat-inactivated fetal bovine serum (Sigma–Aldrich), 71 mM β-mercaptoethanol, and maintained at 37 °C and 5% CO₂ (24).

2.6. Small interfering RNA transfection of EndoC-βH1 cells
EndoC-βH1 cells were seeded on 12-well plates and transfected with siRNA using Lipofectamine RNAiMAX (Thermo Fisher Scientific) 24 h later. The PITPNC1 specific siRNA (siPITPNC1 #1 and #2) sequences were CCACAGACGCACCCGAAUU and CGAUGAAAUUCCAGAGCGC (Microsynth), respectively. Briefly, siPITPNC1 or siCTRL (negative control siRNA, Microsynth) were diluted in OptiMEM (Thermo Fisher Scientific); then Lipofectamine RNAiMAX was added. After 10 min of incubation, the lipid-siRNA complex was added to cells to obtain a control siRNA, Microysnth) were diluted in OptiMEM (Thermo Fisher Scientific) and RNeasy Plus Micro kit (Qiagen) and RNeasy Mini kit (Qiagen), respectively. cDNAs were synthesized using SuperScript II Reverse Transcriptase (Invitrogen), according to the manufacturer’s instructions. Expression of target genes was measured by real-time quantitative PCR (qRT-PCR) using the 7500 Fast Real-Time PCR System (Applied Biosystems). qRT-PCR was run in a final volume of 10 μL containing 2 μL of cDNA and 8 μL of Power SYBR Green PCR mix (Applied Biosystems), in the presence of forward and reverse primers (sequences in Key resources table), GsuB transcript levels were used to normalize each sample. All the primers were obtained from Microsynth.

2.8. Glucose-stimulated insulin secretion
Forty eight hours after siRNA transfection, EndoC-βH1 cells were starved in 0.5 mM glucose DMEM, supplemented with 2% BSA fraction V, 50 μM β-mercaptoethanol, 5.5 μg/mL transferrin, 6.7 ng/mL sodium selenite, 10 mM nicotinamide, 100 μg/mL streptomycin, and 100 U/mL penicillin. After 24-h of starvation, cells were washed and pre-incubated in Krebs–Ringer bicarbonate HEPES buffer (KRHB) containing 0.2% BSA fraction V and 0 mM glucose for 1 h. Insulin secretion was measured following a 4-h incubation with KRHB containing 0.2% BSA fraction V and 0 or 20 mM glucose, in the presence or absence of 45 μM 3-isobutyl-1-methylxanthine (IBMX, Sigma–Aldrich). At the end of the incubation, supernatants were collected; cells lysed for 1 h on ice in TETG buffer: 20 mM Tris HCl pH 8, 1% Triton X-100, 10% glycerol, 137 mM NaCl, 2 mM EGTA; supplemented with protease inhibitor cocktail (complete Tablets Mini EDTA-free, Roche). Insulin secretion and cellular insulin content were measured by ELISA according to the manufacturer’s protocols using the Human Insulin kit (Mercodia).

2.9. Glucose-stimulated insulin secretion statistics
Data were analyzed using R software (v. 3.6.1) and are presented as the means ± S.E.M. from three independent experiments. Comparisons were performed using two-sample unpaired t-tests, and the p-values adjusted for multiple comparisons. Statistically significant differences are indicated with asterisks (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001).

3. RESULTS

3.1. Metabolic, lipidomic, and transcriptomic phenotyping

3.1.1. Metabolic phenotyping
To search for potential correlations between plasma lipids and islet genes controlling β-cell function, we explored the natural diversity of mice with different genetic backgrounds. In a previous study of the differential metabolic adaptation of six different strains of mice [6], we found that different genetic backgrounds directed various insulin secretion capacities and sensitivities to insulin of target tissues, and to different metabolic adaptations to HFD. Here, we decided to work with C57Bl/6J, BALB/cJ, and DBA/2J mice because of their distinct metabolic characteristics [6]. Groups of 8-week-old mice were fed with an RC or an HFD and phenotyped at 2, 10, and 30 days. They were then sacrificed at respectively, 5, 13, and 33 days, for transcriptomic and plasma lipidomic analysis (Figure 1A). These short periods of feeding were selected because we previously showed that important changes in gene expression that predict the long-term (>3 months) establishment of different obesity/diabetes phenotypes occur early after the initiation of the HFD [6,15,16]. In addition, time-series analysis provides additional power to establish meaningful correlations between physiological and omics data. Mouse phenotyping included the assessment at days 2, 10, and 30 of body weight, five hours fasted glycemia and insulinemia, stimulated insulinemia (AUC of insulinemia measured during an OGTT), glucose intolerance, and insulin resistance (n = 185—195 mice, Figure S1). PCA of the physiological data revealed clear separation of mice by strain (Figure 1B) and a weak effect of diet, as indicated by the shift in the average position of each strain of mice, represented by large triangles and circles in the biplot of Figure 1B. This biplot also indicates that “glucose intolerance” was the main driver separating BALB/cJ mice and that “basal insulinemia” and “basal glycemia” drove the separation of DBA/2J mice.

3.1.2. Lipidomic analysis
Four mice from each group were sacrificed in the random fed state at days 5, 13, and 33 of RC or HFD feeding. Their quantitative lipidomic measurements were obtained for 71 out of 72 plasma samples. The total amount of lipids extracted from each plasma sample was in the optimal range for lipidomic measurements (5200–12900 pmol; Figure S2A and Suppl. Table 1) [7]. We identified ~130 lipid molecules from 11 classes of lipids (Table 1). The percentage of total lipids accounted for by each class of lipids (Figure S2B) showed that the major classes were cholesterol esters (CEs), phosphatidylcholines (PCs), cholesterol (Chol), triacylglycerols (TAGs), lysophosphatidylcholines (LPCs), and phosphatidylinositol(s) (Pis). The other lipid
classes formed only small proportions of the total plasma lipids. The percentage of plasma lipids contributed by TAGs showed significant variation across different mouse strains and feeding conditions whereas the contribution of the other lipid class was relatively constant.

PCA showed that plasma lipidomics was influenced by both strains and feeding conditions (Figure 1C). A detailed description of the relative concentration of each lipid species in all the plasma samples analyzed is presented in the heat map of Figure 1D. This illustrates that the concentrations of individual lipids were strain- and diet-dependent. In addition, feeding the HFD, which comprises mostly C16:0, C 18:0, C18:1 and C18:2 fatty acids [17] led to a general tendency to increase the concentrations of lipids containing these fatty acyl side chains (see examples: open arrows in Figure 1D). Nevertheless, several lipid species were also increased that contained one of these C16—C18 fatty acyl side chains and one longer, and more unsaturated fatty acyl side chain (C20:2—C20:4; double arrow in Figure 1D), suggesting that increased activity of various elongases and desaturases also contribute to the changes in lipid species induced by HFD feeding [18]. Figure 1D shows that the plasma concentrations of TAGs were reduced by HFD feeding in the three strains of mice. This observation can be explained by the coordinated induction by HFD feeding of several hepatic β-oxidation genes (See Suppl. Table 2 and further results below).

### 3.1.3. Islet transcriptomic analysis

Islet transcriptomic data were obtained from islets isolated from 3 to 6 randomly fed mice for each of the 18 experimental groups. A total of 69 islet transcriptomic data sets were used for the subsequent analysis as three samples were rejected after quality control.

PCA showed that the islets transcriptomic was influenced mostly by the mouse genetic backgrounds and to a lower extent by the feeding conditions (Figure 1E). The Venn diagrams of Figure S3A show that differential gene expression between islets from RC- and HFD-fed mice was dependent on strains and period of HFD feeding (Suppl. Table 3). The highest number of differentially regulated genes was found at day 5 of HFD feeding. Islets from C57Bl/6J mice displayed the highest number of differentially expressed genes (354 vs. 113 for DBA/2J mice and 43 for BALB/cJ mice). Analysis of islet genes differentially expressed (adj. p < 0.05) between strains, and considering all feeding conditions, revealed that of the 13,780 genes considered, 9,104 genes were differentially expressed between DBA/2J and C57Bl/6J islets,
8,941 when comparing BALB/cJ with C57Bl/6J islets, and 6,268 when comparing BALB/cJ with DBA/2J islets (Figure S2B and Suppl. Table 4).

Together the above multiomics data sets form the basis for the subsequent, unbiased search for salient correlations between plasma lipids and islet gene expression.

3.2. Correlation between islet gene expression modules and plasma lipids

Islet transcriptomic data were subjected to weighted gene co-expression network analysis (WGCNA) [19] to identify groups of genes that were co-regulated across all mouse strains and feeding conditions (gene modules). The Spearman correlation between the summarized value of the expression (eigenvalue) for each of these modules and the concentration of individual lipids was then calculated as described in the Materials and Methods section. These results are presented as a heat map in Figure 2A and the Suppl. Table 5. This shows that some lipid classes correlated homogeneously, positively or negatively, with gene modules identified by a color code. For instance, triacylglycerols (TAGs) were negatively correlated (|ρ| > 0.4), with the light green, turquoise, dark turquoise, and green modules and positively with the blue, cyan, and black modules (Figure 2A). Sphingomyelins, as a class, also correlated with unique modules (magenta, salmon, red, and brown) and the modules also correlated with TAGs (turquoise, blue, and cyan). In other classes of lipids, only a small number of individual species correlated with gene modules. This was the case for PCs, where only the indicated molecules (dashed lines in Figure 2A) showed correlations (|ρ| > 0.2) with some gene modules. Several of these modules were shared with the TAGs (light green, turquoise, dark turquoise, and cyan modules) or SMs (red module). Because TAGs and SMs correlated as classes with islet gene modules, we focused our further analysis on these lipids.

3.3. Correlation of TAGs and SMs with islet gene modules and functional pathways

We performed KEGG and Gene Ontology (GO) analysis of the genes included in the islet modules, thus showing the strongest correlations with plasma TAGs and SMs. The results of this analysis are presented in Figure S4 (for TAGs) and S5 (for SMs). The modules that negatively correlated with TAGs were enriched in “insulin secretion” genes (green module), “insulin signaling pathway” genes (turquoise module), and “histone modification” genes (dark turquoise module). The positively correlated modules were enriched for “phospholipid metabolic process” genes (black module) and “oxidative phosphorylation” genes (blue and cyan modules). No overrepresented terms were found among the light green module genes.

The modules that negatively correlated with SMs included not only the same “insulin signaling pathway” genes (turquoise module) but also “mRNA processing/histone modification” (magenta module). The positively correlated modules were enriched in “vesicular ER/Golgi
transport” genes (salmon module), “histone modification/autophagy” genes (brown module), and “response to wounding” genes (red module).

The relationship between lipidomic data, islet gene modules, and functional pathways is represented in the hive plot of Figure 2B. This shows that the lipid classes with the highest correlations to gene expression modules were TAGs and SMs and that these correlated with essential β-cell functional pathways, such as insulin secretion, oxidative phosphorylation, phospholipid metabolic processes, and insulin signaling. The heat maps of Figure 2C,D shows the correlation of individual TAGs and SMs with the most important β-cell functional pathways identified. Most individual species showed identical correlations with the islet gene modules.

Collectively, the above data showed that the plasma concentrations of TAGs and SMs, showed a significant correlation with islet gene pathways controlling key β-cell functions. Because of the variability of plasma TAGs concentrations across mouse strains and feeding conditions we next focused on the genes characterizing the functional terms correlated with TAGs.

### 3.3.1. Islet modules negatively correlates with TAGs

Analysis of the “insulin secretion” term showed that it included a large number of mRNAs encoding key regulators of GSIS (Figure 3A). The highest correlation with plasma TAGs was found for the Gck mRNA ($r = -0.70; p = 6.52E-09$), which encodes the rate-controlling enzyme in GSIS [20] (Figure 3B). Other highly correlated mRNAs encoded the subunits of the KATP channel (Abcc8 and KCNJ11), the alpha subunit of the Na$^+$/K$^+$ ATPase (Atp1a1), the voltage-gated Ca$^{2+}$ channel subunit alpha-1 (CaCna1d). They also included regulators of the gluco-incretin signaling pathway, which depends on cyclic adenosine monophosphate (cAMP) production (Adcy6, Adcy8, Pik3ca, Creb3l2, Pcdb) and other regulators of insulin granule exocytosis (Camk2b, Snap25).

The top correlated “signaling” terms (turquoise module, Figure S4B) included “MAPK signaling”, “mTOR signaling”, and “Foxo signaling”. The complete list of genes forming these terms and their correlation with TAGs are presented in Suppl. Table 6. The mRNAs most highly correlated to plasma TAGs encoded most of the components of the insulin receptor signaling pathway, Insr, Irs1, Ptk3ca, Ptk3r1, Pten, Pdpk1, Akt3, Foxo1, Foxo3, Mapk1, Mapk9, Sos2, Braf, Mras, and Kras (Figure 3C). The correlation of each of these genes with TAGs is presented in Figure 3D.

The dark turquoise module contained only 29 genes; a significant association of these genes with the “histone modification” term was identified and eight mRNAs supported this association (Suppl. Table 7). These were the acetylated epigenetic mark readers (Brd1, Brd3); a
component of a histone acetyltransferase complex (Ep400); a histone methyltransferase (Setd1b) and demethylase (Pth1r), and Hcfc1 and Arid1a, which are part of chromatin remodeling complexes.

3.3.2. Islet modules positively correlated with TAGs

The GO analysis of the positively correlated black module showed enrichment in “phospholipid metabolic process” and other lipid metabolism-related terms (Figure S4D). These included genes controlling the metabolism of phospholipids (Pipp4r, Pala21z, Bmoa1t, Apgat5, Apgat4), of phosphatidyl-inositols (Pitpn1, Pip4p2, Pip4kc2, Tlc7b, Gpd1f), and of sphingolipids (Cni3, Galc, Cerk, Degs2, St3gala1, Neu3, HexA), with each lipid class playing a modulatory role in insulin secretion [21–24] (Figure 4A). The highest positive correlation was with Abhd6 (r = 0.72, and p = 8.49E-10), encoding monoacylglycerol lipase alpha/beta hydrolase Domain 6, an endocannabinoid degrading enzyme that negatively controls insulin secretion [25].

The blue and cyan modules were enriched in terms (“Thermogenesis”, “Huntington”, “Alzheimer”, “Parkinson”, and “Oxidative phosphorylation”), which primarily comprised oxidative phosphorylation mRNAs (Figures S4E,F). The scheme of Figure 4B shows that most of these genes belong to the OXPHOS chain up to the ATP synthase step and that most of these genes control the metabolism of phospholipids (blue), phosphatidylinositol (green), and sphingolipids (orange) metabolic pathways.

Among the negatively correlated modules, only the black module showed enrichment in “sterol biosynthetic process” (red), “fatty acid degradation” (cyan), “related to endoplasmic stress” (light yellow), and “response to endoplasmic stress” (light yellow); no pathway enrichments were found in the saddle brown module.

Four modules, enriched in specific GO terms showed a positive correlation with TAGs: “sterol biosynthesis process” (dark orange), “cell cycle phase transition” (red), “lipid transport” (brown 4), and “regulation of sterol biosynthetic process” (black). The correlation between individual TAGs and these GO terms is presented in Figure S6A. The scatter plot of Figure S6A shows the separation of the transcriptomic profiles by strain and, in part, by diet, particularly in BALB/cJ mice. These data were then used for WGCN analysis and for identification of gene co-expression modules.

The heat map of Figure S6B shows a correlation between the plasma lipids and liver gene modules. Four modules showed strong negative correlations with plasma TAGs and were enriched in specific GO terms: “fatty acid degradation” (cyan), “vagotomy vesicle transport” (bisque4), “response to endoplasmic stress” (light yellow); no pathway enrichments were found in the saddle brown module.

Four modules, enriched in specific GO terms showed a positive correlation with TAGs: “sterol biosynthesis process” (dark orange), “cell cycle phase transition” (red), “lipid transport” (brown 4), and “regulation of sterol biosynthetic process” (black). The correlation between individual TAGs and these GO terms is presented in Figure S6C. Among the negatively correlated modules, only the “fatty acid degradation” cyan module contained metabolism-related genes. These included a large number of β-oxidation mRNAs (Acad1, Acadvl, Eci1, Eci2, Hadha, Hadhb and Acad11) and two mRNAs encoding a plasma membrane (Slc22a5) and a mitochondrial (Slc25a20) carnitine transporter. The position of the encoded proteins in the overall fatty acid uptake and catabolism pathway is described in the scheme of

Figure 4: Plasma TAGs correlate with islet lipid signaling and oxidative phosphorylation pathways. A. Genes of the “lipid signaling” term of the black module and their Spearman correlation coefficients with plasma TAGs (shown only for correlations with adjusted p-values < 0.05 and |rho| > 0.4). These genes encoded enzymes contributing to the phospholipids (blue), phosphatidylinositol (green), and sphingolipids (orange) metabolic pathways. B. Scheme of the oxidative phosphorylation pathway with, in blue boxes, the genes of the “oxidative phosphorylation” term of the black and cyan modules. C. Spearman correlation coefficients between expression of the “oxidative phosphorylation” genes and plasma TAGs.
Figure 5A. The individual correlations between these mRNAs with plasma TAGs are listed in Figure 5B. The expression of these mRNAs was increased by HFD in the livers of the three mouse strains, as mentioned above (Suppl. Table 2).

Among the positively correlated modules, the dark orange module was enriched with mRNAs encoding enzymes involved in cholesterol biosynthesis (Figure S7). The brown module included 4 mRNAs in the GO term "lipid transport" (Aqp8, Lipc, Cyp2j5, Atp8b4). The red module

Figure 5: Plasma TAGs anti-correlate with the liver β-oxidation pathway. A. Scheme of the liver fatty acid and carnitine uptake and β-oxidation pathways with, in blue, the genes present in the liver "fatty acid degradation term" of the cyan module. B. Spearman correlations between expression of the fatty acid metabolism-related genes of (A) and plasma TAGs. C. Plasma TAGs link liver β-oxidation to β-cell pathways controlling insulin secretion and functional mass. Pathways in blue are anti-correlated with plasma TAGs, those in red are positively correlated with TAGs.

Figure 6: Plasma TAGs also correlate with key β-cell transcripts in humans. Heatmap of the Spearman correlation coefficients between individual TAGs and human islet mRNAs. The human islet mRNAs analyzed were homologous of the mRNAs showing the highest correlation with plasma TAGs in mice. The human data were from paired islet transcriptomic and plasma lipidomic analysis generated from biosamples obtained from partially pancreatectomized patients.
was enriched in biological processes related to cell cycle regulation (Suppl. Table 8). The black module also contained five mRNAs involved in cholesterol biosynthesis (Srebp1f, Abcg1, Eelin1, Eelin2, Scp2). Together, the above data showed a striking inverse correlation between liver ω-oxidation and plasma TAGs concentrations, which conforms with the established relationship between these two parameters [30]. It further suggests a link between hepatic ω-oxidation, plasma TAGs, and β-cell function (Figure 5C).

3.5. Correlation between plasma TAGs and islet transcripts in humans

We next assessed whether similar correlations between plasma TAGs and key functional β-cell genes could be observed in humans. To this end, we analyzed RNA-Seq data generated from laser capture microdissected islets and quantitative plasma lipidomic data obtained from a cohort of 60 partially pancreatectomized patients [31]. For correlation analysis, we selected the aforementioned human orthologues of the mouse islet mRNAs that encode components of the GSIS pathway, lipid metabolic enzymes, and proteins of the insulin signaling pathway. Out of the 86 considered mouse genes, 85 had human orthologs that were expressed in the human islets data set. Figure 6 shows the heatmap of the correlation between plasma TAGs and 44 of the most highly correlated human islet mRNAs. Strikingly, as in mice, the top negative correlation was found for GCK. The correlation was also found with CREBP and KCNB1, important regulators of the insulin secretory function (Figure 5C).

We found that plasma TAGs, as a class, correlate strongly with islet mRNAs encoding regulators of GSIS, enzymes producing AβH06, and for HSD17B11. Key elements of the insulin signaling pathway showed significant correlation with TAGs: IRS2, PI3KR1, AKT3, FOXO1, FOXO3 (see Figure 6B).

3.6. PITPNC1 as a novel regulator of insulin secretion

Because islet PITPNC1, which codes for a phosphatidyl-inositol transfer protein [32], showed a strong correlation with TAGs both in mice and humans, we performed preliminary experiments to assess whether this gene could be a so-far uncharacterized regulator of insulin secretion. We first measured its expression in the mouse MIN6B1 and the human EndoC-bH1 insulin cell lines. MIN6B1 cells have an almost undetectable level of the Pitpnc1 mRNA (Ct ~ 30). In contrast, the two splice variants of PITPNC1 [32] were robustly expressed in EndoC-bH1 cells (Ct ~ 25; Figure 7A,B). We, therefore, investigated whether silencing PITPNC1 in EndoC-bH1 would impact GSIS. Figure 7C shows that transfecting the cells with two distinct siRNAs, which target both splice variants, reduced PITPNC1 expression by ~60–70%. This led to a significant reduction in insulin secretion in basal conditions as well as in the presence of 20 mM glucose or of 20 mM glucose and IBMX, a phosphodiesterase inhibitor (Figure 7D).

4. DISCUSSION

In the present study, we aimed to determine whether plasma lipids could be the indicators of β-cell function and whether the tissues and pathways controlling the production of these lipids could be identified. We found that plasma TAGs, as a class, correlate strongly with islet gene co-expression modules. The most correlated modules are enriched in mRNAs encoding regulators of GSIS, enzymes producing

![Figure 7: PITPNC1 as a novel regulator of insulin secretion. Analysis of a regulatory role of PITPNC1 in glucose-stimulated insulin secretion.](image-url)
various lipids that modulate insulin secretion, and transducers of the insulin signaling pathway. In the liver, the highest correlation was with the \( \beta \)-oxidation pathway. In humans, analysis of paired plasma lipid and islet transcriptomic data revealed a similar pattern of correlation between plasma TAGs and key regulators of insulin secretion as found in mice. This cross-species correlation analysis led to the identification of PITPN1C1 as a new candidate regulator of insulin secretion. Thus, plasma TAGs emerge as biomarkers of \( \beta \)-cell function both in mice and humans and as potential effectors of a liver-\( \beta \)-cell axis.

It is usually considered that T2D appears when pancreatic \( \beta \)-cells can no longer compensate for the insulin resistance of the liver, adipose tissue, and muscle [1,33]. However, whether the primary defects lie in insulin secretion or in insulin action may vary among patients. Assessing insulin sensitivity and appearance of insulin resistance or determining the insulin secretion capacity of an individual during the progression of pre-diabetes or T2D is a complex process that relies on glucose or insulin tolerance tests or on insulinemic/glycemic clamp techniques. Therefore, finding prognostic plasma biomarkers for T2D, which could also give information about primary defects in pancreatic \( \beta \)-cells and/or in insulin target tissues would pave the way for improved preventive or therapeutic options adapted to individual patients.

Several previous studies led to the identification of prognostic biomarkers for T2D or which can help T2D patients’ stratification [34] and to the identification of islet gene modules characteristic of diabetic \( \beta \)-cell deregulations [31,35–37]. Plasma metabolomic biomarkers include various classes of hydrophilic and lipophilic molecules [34,38–45]. In a previous study, we identified dihydroceramidase as prognostic biomarker candidates for T2D in mice and humans [5]. Plasma TAGs showed a correlation in mice in our previous study with fasting insulinemia and glucose intolerance. Here, we exploited the vast diversity of metabolic traits of mice with different genetic backgrounds to search, in an unbiased manner, for salient correlations between plasma lipids and tissue gene expression modules.

We found that plasma TAGs and SMs displayed high correlations with islet gene modules that regulate critical \( \beta \)-cell physiological functions. The highest correlations with plasma TAGs were found for mRNAs that encoded key regulators of insulin secretion. These included the negative correlations with Gck and the subunits of the K\( _{ATP} \) channel, Abcc8, and Kcnj11, and the positive correlation with Abhd6, a negative regulator of insulin secretion [25]. Thus, during the initial period of HFD feeding, when plasma TAGs are reduced, increased expression of the Gck gene and the K\( _{ATP} \) channel and lower levels of Abhd6 represent a mechanism for \( \beta \)-cell compensation and insulin hypersecretion.

A negative correlation between plasma TAGs and a large number of genes involved in the insulin signaling pathway was also documented. The insulin/IGF1R signaling pathway in \( \beta \)-cells participates in the control of \( \beta \)-cell proliferation, differentiation, and protection against apoptosis [26,46–50]. Thus, we suggest a strong association between the activity of this signaling pathway and GSIS in our studied mouse models. Contrastively, no correlation between plasma TAGs and transducers or modulators of the insulin/IGF1R signaling pathway was established in the liver. This suggests that the insulin signaling pathway may be differentially regulated in \( \beta \)-cells and the liver. Transcriptional control of the expression of the various signaling components may predominate in the pancreas whereas phosphorylation events play a major role in modulating the insulin signaling cascade in the liver [51].

Correlation between SMs and islet gene modules overlapped, in part, with that between gene modules and plasma TAGs but also included other modules, characterized by “vesicular ER/Golgi transport”, “histone modification/autophagy” or “response to wounding” genes. These were less directly related to the processes of acute GSIS and control of \( \beta \)-cell mass and function; therefore, we focused on the modules correlated with plasma TAGs.

High levels of plasma TAGs can result from increased liver de novo lipogenesis, increased hepatic uptake of fatty acids originating from the adipose tissue or the ingested food, or reduced hepatic \( \beta \)-oxidation [30]. Additionally, we found that plasma TAGs displayed a strong inverse correlation only with a liver gene module enriched in \( \beta \)-oxidation genes (Acadl, Acadvl, Eci1, Eci2, Hadha, Hadhb, and Acad11). This suggests that in our experimental model increased hepatic fatty acid catabolism is the main regulator of plasma TAG concentrations. This is consistent, for instance, with the observation that fenofibrate-induced hepatic \( \beta \)-oxidation reduces hypertriglyceridemia [30,52]. The same gene co-expression module also included genes for the plasma membrane (Slc22a5) and mitochondrial membrane (Slc25a20) carnitine transporters. Carnitine is required for transporting fatty acids into mitochondria by carnitine palmitoyl-transferase for their subsequent \( \beta \)-oxidation. The appearance of two carnitine transporters in our correlation analysis indicates that they regulate the availability of carnitine for free fatty acid degradation. Carnitine nutritional supplementation can reduce plasma TAGs levels in rodents and humans [53–55].

Together, our lipidomic—transcriptomic correlation analysis suggests the existence of a liver-to–\( \beta \)-cell axis whereby hepatic \( \beta \)-oxidation, by regulating plasma TAGs levels, may influence the expression of key regulators of GSIS. Although the mechanistic link between TAGs and \( \beta \)-cell insulin secretion capacity remains poorly understood, it is well known that exposure of islets to high glucose, high free fatty acids, or a combination of both lead to increased basal insulin secretion [56–58] and defects in \( \beta \)-cell function. \( \beta \)-cells express the enzyme lipoprotein lipase, which releases free fatty acids from circulating TAGs for cellular uptake [59,60]. Free fatty acids are required to maintain insulin secretion capacity in the fasted state [61]. However, in the long term, elevated concentrations of free fatty acids increase basal- and reduce glucose-stimulated insulin secretion [62–64]. We found that the lipoprotein lipase (Lppl) mRNA, which is expressed in \( \beta \)-cells and is required for normal GSIS [59], is expressed at a similar level in all islet preparations (not shown). Thus, differential free fatty acid uptake may primarily be determined by the circulating levels of TAGs. The mentioned detrimental effects of free fatty acids on \( \beta \)-cell function may explain why elevated plasma TAGs can be causal biomarkers of \( \beta \)-cell dysfunction and T2D susceptibility [65,66]. An alternative or complementary explanation is that plasma TAGs are secreted by the liver as part of lipoprotein particles; studies have shown that low-density lipoproteins (LDL) have a detrimental effect on GSIS, whereas high-density lipoproteins (HDL) protect the \( \beta \)-cell functional mass [67]. The fact that higher circulating LDL/HDL ratios may increase diabetes susceptibility is in agreement with the observation that single nucleotide polymorphisms leading to reduced expression of LPL in humans are associated with an increased risk of T2D development [68].

Our study further unveiled similar correlations between plasma TAGs and islet genes involved in glucose and lipid regulation of insulin secretion and insulin signaling in mice and humans. This is the case for ABHD6, a negative regulator of insulin secretion [25], or the vitamin D receptor (VDR), which has also been suggested to control the glucose competence of \( \beta \)-cells [69,70]. But this cross-species analysis also identified several highly correlated genes whose function in regulating insulin secretion remains undefined. Examples include PITPN1C1, the mRNA that showed the second highest correlation with TAGs in mice and the highest correlation in humans. PITPN1C1 encodes a
phosphatidylinositol transfer protein, which associates with the Golgi complex through its binding to resident PI4P and which recruits Rab1b to enhance protein secretion [71,72]. Single nucleotide polymorphism in PITPNC1 has also been linked to an increased risk of T2D [73]. Other potentially interesting mRNAs are HSD17B11 and HSD17B12 genes, which encode, respectively, a short-chain fatty acid dehydrogenase/reductase and an enzyme that converts estrone into estradiol and which also has a fatty acid elongase activity. Further work is, however, needed to determine the exact mechanism underlying the regulation of insulin secretion by PITPNC1.

Collectively, our data show that plasma TAGs display high correlations with islet gene co-expression modules encoding key regulators of β-cell function and strong correlation, in the liver, with a module that contains mRNAs encoding fatty acid transporters, carnitine transporters, and β-oxidation genes. These correlations indicate that lower hepatic β-oxidation activity leads to increased plasma TAGs, which then may negatively impact the expression of genes controlling GSIS. Therefore, TAGs can be considered not only as biomarkers of type 2 diabetes [74] but also as indicators of the correlation that exists between liver fatty acid degradation and β-cell pathways that control these cells’ integrity and secretion capacity. Finally, this research provides a rationale for reducing plasma TAGs, for instance through carnitine supplementation or by fenofibrate-like molecule treatments, to prevent type 2 diabetes development.

**AUTHOR CONTRIBUTIONS**

B.T., C.M., M.I., and C.C.G. conceived the experiments. C.C.-G., J.L., J.D. and N.K. performed the mouse phenotyping and collected and processed the samples. C.R. and K.S. performed the lipidomic analysis. A.R.S.-A. performed the analysis and multi-omics integration in the pre-clinical model and the human cohort of PPP. L.W. and F.M. provided lipidomics quality controls and phenotypic measures quality controls and statistical tests. C.R. performed the experimental validation in the mouse cell lines. M.S., M.B., J.W., M.D., and M.I. contributed human transcriptomic data. B.T. and A.R.S.-A. wrote the manuscript with contributions from M.I. who also supervised the bioinformatics analysis. B.T. coordinated and managed the research project with support from C.M. and M.I. All authors have critically read the manuscript and approved it for publication.

**DATA AVAILABILITY**

The data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus (GEO) database with the accession number GSE140369 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE140369) and liver expression with accession number GSE164672 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE164672).

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