Characterization of Vitamin A Metabolome in Human Livers With and Without Nonalcoholic Fatty Liver Disease

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ABSTRACT

Retinoids are essential endogenous compounds involved in regulation of critical biologic processes, including maintenance of metabolic homeostasis in the liver. Much of the knowledge of altered retinoid homeostasis in human disease states is derived from changes in indirect markers such as mRNA expression of retinoid-related genes and circulating concentrations of retinol or its binding protein RBP4. We hypothesized that in the human liver, concentrations of the active retinoid all-trans-retinoic acid (atRA) correlate with the concentrations of retinyl palmitate (RP), the storage form of atRA, retinol, the inactive vitamin A, and the mRNA expression of retinoid-related genes. On the basis of existing knowledge of altered vitamin A homeostasis in metabolic syndrome, we also predicted that in human livers with nonalcoholic fatty liver disease (NAFLD) retinoid concentrations would be decreased. Using novel liquid chromatography–tandem mass spectrometry methods, the hepatic vitamin A metabolome was quantified in normal human livers (n = 50) and 22 livers from donors with NAFLD. The hepatic concentrations of RP, atRA, 13-cisRA, and 4-oxo-atRA were significantly decreased in NAFLD samples in comparison with normal liver samples, whereas retinol levels remained unchanged. The concentrations of atRA were positively correlated with RP and 13-cisRA but not with retinol or the relative mRNA expression of LRAT, ALDH1A1, CYP26A1, RARα, and RARβ. An active metabolite of atRA, 4-oxo-atRA was, for the first time, detected in human tissues at comparable concentration with RA isomers, suggesting this retinoid may contribute to retinoid signaling in humans.

SIGNIFICANCE STATEMENT

This study shows that in NAFLD liver vitamin A homeostasis is disrupted potentially contributing to disease progression. The results show that interpretation of retinoid homeostasis on the basis of indirect markers such as retinol concentrations or mRNA data is probably misleading when evaluating human disease processes, and analysis of the broader retinoid metabolome is needed to characterize disease effects on retinoid signaling.

Introduction

Retinoids, which are structural analogs of the inactive vitamin A (retinol, Fig. 1), are essential endogenous compounds involved in regulation of various biologic processes, including adipogenesis and lipid metabolism (Shirakami et al., 2012). Mouse studies have suggested that all-trans-retinoic acid (atRA) plays a role in enhanced lipid oxidation and reduced fat accumulation in the liver (Kang et al., 2007; Amengual et al., 2010; Kim et al., 2014). On the other hand, it is generally believed that obesity and development of nonalcoholic fatty liver disease (NAFLD) are associated with disturbed vitamin A homeostasis (Shiota, 2005; Graham et al., 2006; Ziouzenkova et al., 2007; Liu et al., 2015; Trasino et al., 2015). In humans, serum retinol concentrations have been shown to inversely correlate with the progression of NAFLD in a number of studies (Suano de Souza et al., 2008; Villaca Chaves et al., 2008; Botella-Carretero et al., 2010; Chaves et al., 2014; Liu et al., 2015). As such, understanding liver vitamin A homeostasis, its correlation with retinol concentrations, and relationship with NAFLD is important.

The liver plays a key role in regulating vitamin A homeostasis, as retinol is stored via esterification by lecithin retinol acyltransferase (LRAT) as retinyl esters in hepatic stellate cells (O’Byrne and Blaner, 2013). The liver concentrations of the active retinoid all-trans-retinoic acid (atRA) and the main circulating retinoid, retinol, are believed to be regulated via a complex network of metabolic steps and enzymes (Fig. 1) (Napoli, 2012; Kedishvili, 2016). However, how this network is altered in disease states is not well understood. There is a lack of qualitative and quantitative information about retinoid concentrations and the expression of vitamin A-metabolizing enzymes in human healthy and diseased tissues. In fact, most clinical data related to altered vitamin A homeostasis are derived from indirect measurements of circulating or tissue...
retinol concentrations and evaluation of changes in mRNA expression of retinoid-related genes. For example, several studies have shown that circulating retinol and retinol binding protein RBP4 plasma concentrations are significantly altered in diabetes and in NAFLD (Graham et al., 2006; Suano de Souza et al., 2008; Botella-Carretero et al., 2010), but retinoid concentrations, especially at RA concentrations, in the liver in these disease states have not been directly measured. In addition, at present no comprehensive characterization of the vitamin A metabolome in human livers has been conducted. As such, the interindividual variability and biologically relevant concentrations in human livers are unknown.

The lack of information on tissue retinoid concentrations and homeostasis is partially a result of the bioanalytical challenges in analyzing retinoids. Retinoids are light-sensitive and have poor mass spectrometric fragmentation. Retinol and retinyl esters have mainly been measured using liquid-liquid extraction and liquid chromatography-UV absorption (LC-UV) (Gundersen and Blomhoff, 2001; Schmidt et al., 2003; Kane and Napoli, 2010). However, retinoids are endogenous compounds and it is impossible to obtain “blank” biologic tissues devoid of retinoids. Therefore, analytical recovery, selectivity, and lack of endogenous interference cannot be easily confirmed. Of special concern is that nearly always the methods have used structurally related but chemically different internal standards, which may not have been adequate to correct for variability in extraction efficiency from different matrices (Schmidt et al., 2003; Obrochta et al., 2014). This is of particular concern when the role of vitamin A homeostasis is assessed in disease states such as NAFLD that involve considerable changes in tissue composition. Fibrosis and fat infiltration for example may alter extraction efficiency and matrix effects and confound the quantification of retinoids. To address these issues, we developed and validated a set of

Fig. 1. Metabolic network of retinoid homeostasis shown with chemical structures of retinoids and known metabolic enzymes involved in each step. ADH, alcohol dehydrogenase; BCMO1, β-carotene monoxygenase 1; RDH, retinol dehydrogenase; REH, retinyl ester hydrolase.
new liquid chromatography–mass spectrometry (LC-MS/MS) methods to quantify the vitamin A metabolome in human liver using isotope-labeled internal standards and LC-MS/MS-based quantification. Using these quantification methods, our goal in this study was to test the hypothesis that the commonly used indirect markers of atRA concentrations, retinol and retinyl ester concentrations and retinoid-related mRNA expression, correlate with atRA concentrations in human livers from normal and NAFLD donors. We aimed to determine whether measurements of indirect markers or altered atRA concentrations in disease states could be used to define how retinoid signaling networks and metabolism are altered in disease states. Our results show that although NAFLD results in a significantly decreased concentrations of atRA and retinyl esters in human livers, measurement of retinol or retinoid-related mRNA expression fails to accurately capture these changes in retinoid homeostasis. This study shows that to identify the processes that lead to altered vitamin A homeostasis in disease states, the accessible vitamin A metabolome should be characterized.

Materials and Methods

Chemicals and Reagents. atRA and 13-cisRA were purchased from MilliporeSigma (St. Louis, MO). atRA-d5, 13-cisRA-d5, 4-oxo-atRA, 4-oxo-13-cisRA, 4-oxo-atRA-d5, all-trans-retinyl palmitate, and retinol were purchased from Toronto Research Chemicals (North York, ON). All-trans-retinyl palmitate-d4 (RP-d4) and all-trans-retinol-d4 (ROL-d4) were purchased from Cambridge Isotopes (Andover, MA). Blank human serum (DC MASS SPECT GOLD MSG4000) was purchased from Golden West Diagnostics (Temecula, CA). All solvents used in LC-MS/MS analysis were Optima LC-MS grade from Fisher Scientific (Pittsburg, PA).

Liver Samples. To characterize the hepatic vitamin A metabolome in human subjects, 50 de-identified human liver samples were obtained from Corning (Corning, NY). The livers were confirmed by Corning as healthy livers with no major pathology. Relevant donor histories are listed in Supplemental Table S1. This set of samples is referred to as normal samples in this study. The livers were from 39 males and 11 females with donor ages ranging from 21 to 80 years old. Donors included 41 Caucasians, five Asians, three African Americans, and one Hispanic. To investigate if the hepatic vitamin A metabolome is changed in NAFLD, 22 human liver samples were obtained from the Liver Tissue Cell Distribution System (Supplemental Table S2), including nine steatosis (ethnicity unknown; four males and five females) and 13 nonalcoholic steatohepatitis (NASH) samples (10 Caucasians, one Hispanic, two ethnicity unknown; eight males and five females). The diagnosis of steatosis and NASH in these livers was confirmed by histopathology (Supplemental Table S2). The donor ages ranged from 32 to 70 years old. The samples with either steatosis or NASH are together referred to as NAFLD samples in the study. Mouse livers were obtained from C57BL/6×129 mice used in a previous study (Stevison et al., 2017) and all liver samples were from the vehicle control groups in the previous study. All samples were stored in –80°C until analysis.

Sample Preparation. All sample preparation steps for retinoid measurements were performed under yellow light to prevent retinoid isomerization and degradation. As retinyl palmitate is the primary retinyl ester stored in the liver, the hepatic concentration of RP was measured to represent retinyl esters in human liver. To measure RP and retinol, human liver tissue (40–60 mg) was homogenized in a 7:1 ratio of 0.9% saline to tissue weight using Omni Bead Ruptor 24 Bead Mill homogenizer with ceramic beads in methanol-dry ice. The liver homogenate (240 µl) was transferred to a 1.7-ml Eppendorf tube and spiked with 15 µl of internal standard mixture containing 400 µM RP-d4 and 20 µM ROL-d4. After vortexing, the sample was left on ice for 10 minutes, followed by addition of 1 ml ice-cold acetonitrile (ACN) to precipitate protein. The sample was vortexed and placed on ice for 20 minutes before centrifugation. After the sample was centrifuged at 18,000g at 4°C for 30 minutes to pellet liver protein, the supernatant was collected. A 50-µl aliquot of the supernatant was diluted with 950 µl of 80% ACN for the LC-MS/MS analysis of RP. The rest of the supernatant was transferred to amber LC-MS vials for the LC-MS/MS analysis of retinol.

To measure atRA, 13-cisRA, and 4-oxo-atRA in human liver, a liquid-liquid extraction was adapted and modified from a published method (Obropta et al., 2014). Liver tissues (80–100 mg) were homogenized in a 2-ml Potter-Elvehjem glass homogenizer (Kimbel Glass, Vineland, NJ) on ice with a ratio of 5:1 of 0.9% saline to tissue weight. The Bead Mill homogenizer was not used for liquid-liquid extraction because the ceramic beads used for homogenization caused technical difficulties for the performance of liquid-liquid extraction. The homogenate was transferred with a Pasteur pipette to a 15-ml glass culture-tube on ice and was spiked with 10 µl of internal standard (IS) mixture containing 500 nM atRA-d5, 13-cisRA-d5, and 4-oxo-atRA-d5. To this mixture, 2 ml of 25 mM KOH in ethanol was added, and the sample was vortexed. Ten milliliters of hexanes were then added, and the sample was vortexed and centrifuged at 300g for 2 minutes to separate organic phase from aqueous phase. The top hexane layer was removed using a Pasteur pipette, and 120 µl of 4 M HCl was added to the aqueous layer of the sample. The sample was quickly vortexed, followed by the addition of 10 ml of hexanes to extract atRA and 13-cisRA. After centrifugation at 300g for 2 minutes, the top hexane layer containing RA isomers was removed using a Pasteur pipette to a new glass culture tube on ice. Ethyl acetate (10 ml) was then added to the aqueous sample phase followed by 2 ml of water to enhance phase separation to extract 4-oxo-atRA. After centrifugation at 300g for 15 minutes, the top ethyl acetate layer containing 4-oxo-atRA was transferred to a new glass culture tube on ice using a Pasteur pipette. The organic layers were dried down at 35°C under a gentle N2 flow and then placed on ice. Residues containing RA isomers (from hexane extraction) or 4-oxo-atRA (ethyl acetate extraction) were resuspended into 60 µl of ACN and transferred to amber mass spectrometry vials with glass inserts using Pasteur pipettes. Right before the LC-MS/MS analysis, water (40 µl) was added to the ACN resuspensions to improve chromatographic resolution.

LC-MS/MS Analysis. Three LC-MS/MS methods for the analysis of RP, retinol, RA isomers, and 4-oxo-RA isomers were developed. All LC-MS/MS analyses were performed under positive ion mode using atmospheric pressure chemical ionization and nitrogen as the collision and curtain gases. The analytical column (2.1 × 150 mm, 2.7 µm) and guard column (2.1 × 5 mm, 2.7 µm) used for analyte separation were both Ascentis Express RP-amide columns from MilliporeSigma. The autosampler was set at 6°C. To measure RA isomers and 4-oxo-RA, the LC-MS/MS analysis was performed on an AB Sciex 5500 QTRAP Mass Spectrometer (AB Sciex LLC, Framingham, MA) coupled with an Agilent 1290 UHPLC (Agilent Technologies, Santa Clara, CA). RP and retinol were measured using AB Sciex 6500 QTRAP Mass Spectrometer coupled with Shimadzu UPLC XR DGU-20A5 (Shimadzu Corporation, Kyoto, Japan). For all the LC-MS/MS analyses, mobile phases A and B were water with 0.1% formic acid (FA) and ACN with 0.1% FA, respectively. To measure retinol and RP, the injection volume, LC flow rate, and column temperature were set at 4 µl, 0.4 ml/min, and 40°C. The LC condition started at 60% B for 2 minutes, followed by a gradient to 66% B by 9.2 minutes, and then to 100% B by 13 minutes, and then kept at 100% B for 10 minutes before the return to starting conditions. To measure RA isomers, the injection volume, LC flow rate, and column temperature were set at 20 µl, 0.4 ml/min, and 25°C. The LC condition started at 70% B for 3 minutes, then increased linearly to 78% B by 12 minutes, then changed to 100% B, and was kept at that for 3 minutes before return to starting condition. For 4-oxo-RA...
In MS3 mode, which was set up to monitor RA isomers, product ions exit potential 10 V. All other MS parameters are listed in Table 1. LC-MS/MS methods were: temperature 350°C, nebulizer current in parallel for measurement of RA isomers and for confirmation multiple reaction monitoring (MRM), whereas MS/MS/MS (MS3, were analyzed and the concentrations quantified on the basis of measurements, the injection volume, LC flow rate, and column temperature were set at 20 μl, 0.5 ml/min, and 40°C. The LC conditions started at 40% B for 2 minutes, followed by a gradient to 67% B by 9 minutes, then changed to 100% B, and maintained for 2 minutes before the return to the starting conditions. All retinoids were analyzed and the concentrations quantified on the basis of multiple reaction monitoring (MRM), whereas MS/MS/MS (MS3, second generation production ion monitoring) data were collected in parallel for measurement of RA isomers and for confirmation of MS parameters used in all three MS-MS/MS methods: temperature 350°C, nebulizer current 5 μA, curtain gas 35 psi, collision-associated dissociation –2 psi and cell exit potential 10 V. All other MS parameters are listed in Table 1. In MS3 mode, which was set up to monitor RA isomers, product ions [M+H]⁺ derived from the precursor ion of the MRM transition were collected, ranging from m/z 109 to 220 Da. The Q1 resolution was set to low and the Q0 trapping was turned on. The Q3 entry barrier was 8 V and the scan rate 10,000 Da/s. The linear ion trap (LIT) fill time at 100 milliseconds, Q1 at low resolution, and Q3 entry barrier at 8 V. The Q0 trap was turned on. The LC conditions and other MS parameters were the same as those described above for the measurement of 4-oxo-RA isomers.

**Method Validation.** The methods used for retinoid concentration measurements were validated following the FDA guidance on bioanalytical method validation (FDA, 2018), adapted for endogenous compounds. Owing to the lack of blank retinoid-free liver tissue, charcoal-treated blank human serum free of retinoids was used as a blank matrix. For method development, the blank human serum was diluted with 5× volume of 0.9% saline, and used to prepare standard-curve samples and three quality control samples (QC1–3) spiked with the retinoids in low, low-medium, or medium-high concentration. The range of standard curves and QC concentrations for all the analyzed retinoids is listed in Table 2. As a further validation to assess extraction variability in the liver tissue matrix, human liver samples from six donors were homogenized with 5:1 0.9% saline/tissue weight, the homogenates pooled, and stored as aliquots at −80°C for analysis as liver QC samples (LQCs). During validation, diluted serum or 540 μl of pooled liver homogenate per analysis were used to measure RA isomers and 4-oxo-αRA, and 240 μl per analysis were used for retinol and RP measurements. All samples were processed and analyzed with corresponding LC-MS/MS methods as described above. The peak areas of analyte and IS were integrated using Analyst software version 1.6.3 (AB Sciex LLC). To build standard curves, the peak area ratio of analyte to IS was plotted as a function of serum retinoid concentration. All QCs and LQCs were prepared in six replicates per day and validation assays were repeated on three separate days to determine intraday and interday coefficient of variation (%CV). Retinoid standards, including RA and 4-oxo-RA dissolved in 60% acetonitrile, and retinol and RP dissolved in 80% acetonitrile, were measured.

**TABLE 1**

MS parameters used in LC-MS/MS and LC-MS/MS/MS analysis

<table>
<thead>
<tr>
<th>Analyte</th>
<th>MRM (MS2)</th>
<th>DP (V)</th>
<th>EP (V)</th>
<th>CE (V)</th>
<th>GS1</th>
<th>GS2</th>
</tr>
</thead>
<tbody>
<tr>
<td>ROL and RP</td>
<td>m/z 269 &gt; 93, 95</td>
<td>35</td>
<td>4</td>
<td>30</td>
<td>70</td>
<td>70</td>
</tr>
<tr>
<td>ROL-d8</td>
<td>m/z 277 &gt; 98, 102</td>
<td>35</td>
<td>4</td>
<td>30</td>
<td>70</td>
<td>70</td>
</tr>
<tr>
<td>RP-d4</td>
<td>m/z 273 &gt; 94, 98</td>
<td>35</td>
<td>4</td>
<td>30</td>
<td>70</td>
<td>70</td>
</tr>
<tr>
<td>αRA and 13-cisRA</td>
<td>m/z 301 &gt; 205</td>
<td>60</td>
<td>10</td>
<td>17</td>
<td>80</td>
<td>0</td>
</tr>
<tr>
<td>αRA-d3 and 13-cisRA-d3</td>
<td>(m/z 301 &gt; 205 &gt; 159)</td>
<td>80</td>
<td>10</td>
<td>17</td>
<td>80</td>
<td>0</td>
</tr>
<tr>
<td>4-oxo-αRA and 4-oxo-13cisRA</td>
<td>m/z 315 &gt; 137, 147, 241</td>
<td>50</td>
<td>10</td>
<td>29, 26, 20</td>
<td>60</td>
<td>70</td>
</tr>
<tr>
<td>4-oxo-αRA-d3</td>
<td>m/z 318 &gt; 137</td>
<td>50</td>
<td>10</td>
<td>29</td>
<td>60</td>
<td>70</td>
</tr>
</tbody>
</table>

CE, collision energy; DP, declustering potential; EP, entrance potential; GS, gas.

**TABLE 2**

Method validation data for the analyzed retinoids

The measured concentrations are the mean concentrations of interday measurements.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>αRA</th>
<th>13-cisRA</th>
<th>4-oxo-αRA</th>
<th>Retinol</th>
<th>RP</th>
</tr>
</thead>
<tbody>
<tr>
<td>QC1</td>
<td>1-1000 nM</td>
<td>1-1000 nM</td>
<td>2.7-444 nM</td>
<td>0.15-37.5 μM</td>
<td>3-75 μM</td>
</tr>
<tr>
<td>QC2</td>
<td>25.9 nM</td>
<td>25.9 nM</td>
<td>47.4 nM</td>
<td>1.6 μM</td>
<td>31.3 μM</td>
</tr>
<tr>
<td>QC3</td>
<td>222 nM</td>
<td>222 nM</td>
<td>296 nM</td>
<td>6.3 μM</td>
<td>125 μM</td>
</tr>
<tr>
<td>Liver QC</td>
<td>139.5 pmol/g</td>
<td>31.3 pmol/g</td>
<td>188.5 pmol/g</td>
<td>2.2 nmol/g</td>
<td>66.2 nmol/g</td>
</tr>
</tbody>
</table>

CV, coefficient of variation.
used to determine on-column limit of detection (LOD; signal-to-noise >3 to 1) and lower limit of quantitation (LLOQ; signal-to-noise >5 to 1). Owing to the lack of “blank” liver tissue, LOD and LLOQ in matrix were not measured. To confirm the validity of the approach of using blank serum as a reference matrix instead of liver to construct standard curves, mouse liver homogenates (540 µl per analysis for RA and 4-oxo-RA isomers and 240 µl per analysis for retinol and RP) were also spiked with retinoids to build standard curves and samples were processed and analyzed as described above in parallel with serum standard curves. The slopes of standard curves for each retinoid were compared between the two matrices to test for a systematic matrix effect and to verify similar concentration-response in the two matrices.

mRNA Expression in Human Liver Samples. Total RNA was extracted from human liver tissues using TRIzol reagent (Invitrogen, Carlsbad, CA), and was reverse-transcribed into cDNA using High-Capacity cDNA Reverse Transcription Kit (Life Technologies, CA). Real-time PCR was performed using StepOnePlus Real-Time PCR System with PrimeTime probes (Integrated DNA Technologies, Coralville, IA). The mRNA expression of aldehyde dehydrogenase 1A1 (ALDH1A1), LRAT, CYP26A1, retinoic acid receptor (RARα, and RARβ was measured in 48 out of the 50 normal human livers. GAPDH was used as the housekeeping gene. The mRNA expression of ALDH1A1, LRAT, CYP26A1, RARβ, and RBP4 was also measured in 22 NAFLD samples and in parallel in nine randomly selected normal liver samples with GAPDH as the housekeeping gene. The following primers purchased from Integrated DNA Technologies were used: ALDH1A1 (Hs.PT.56a.38450309), CYP26A1 (Hs.PT.58.2905296), LRAT (Hs.PT.58.39384986), RARα (Hs.PT.58.2437218), RARβ (Hs.PT.58.364456), RBP4 (Hs.PT.58.38653618), and GAPDH (Hs.39a.22214836). The relative mRNA expression level normalized to the housekeeping gene was calculated using the $2^{-ΔΔCt}$ method and comparing the expression in individual donors to the mean of all livers (Livak and Schmittgen, 2001).

Data Analysis. For both normal and NAFLD liver samples, retinoid concentrations in liver and the mRNA expression of retinoid homeostasis-related genes were first log-transformed and Grubb’s test was used to test for the presence of outliers in each retinoid group and each gene group. Spearman’s rank-order correlation was done to analyze correlations of the hepatic atRA concentration with other retinoid concentrations and the mRNA expression of retinoid homeostasis-related genes in normal livers. Because multiple comparisons were done in the correlation analyses, a Bonferroni
adjustment was used to adjust P value (0.05/9) and a P value less than 0.005 was considered significant. To compare NAFLD livers with normal liver samples, significant differences in retinoid concentrations and the relative mRNA expression of retinoid-related genes among the normal, steatosis, and NASH group were analyzed by Kruskal-Wallis test followed by Dunn’s multiple comparison test. The Spearman’s rank-order correlation was used to analyze correlations of atRA concentrations with other retinoid concentrations in all measured human liver samples. A P value less than 0.01 (Bonferroni adjusted P value 0.05/4) was considered significant. Spearman’s rank-order correlation was also used in an exploratory analysis to test for correlation between donor age and retinoid concentrations. Differences in retinoid concentrations between male and female donors were tested using two-tailed students t test.

Results

Detection of Vitamin A Metabolome in Human Livers. All measured retinoids in human liver were separated and detected using the corresponding LC-MS/MS methods. The developed methods allowed specific and sensitive LC-MS/MS-based detection and quantification of retinol, RP, atRA, and 13-cisRA in human livers (Figs. 2 and 3). In addition, for the first time, 4-oxo-atRA and 4-oxo-13-cisRA were detected in the human liver (Fig. 4). The retention times of all analytes detected in human liver matched with those of synthetic standards and isotope-labeled internal standards. The retention times and relative signal intensities of atRA and 13-cisRA peaks identified on the basis of the MRM scans corresponded to those observed in MS3 scans, indicating correct identification and lack of interference in the peaks and mass transitions selected to quantify atRA and 13-cisRA (Fig. 3). In addition, peaks with retention times around 7.9 and 9.1 minutes were detected in the human liver MRM chromatograms in the same transition as RA (Fig. 3), which appear to be nonretinoid-related interfering peaks on the basis of the considerably lower signal intensity in the RA-specific MS3 transition compared with atRA and 13-cisRA (Fig. 3). This finding is similar to what has been reported previously for mouse tissues and human testis (Kane et al., 2008; Arnold et al., 2015).

To confirm the identity of 4-oxo-atRA and 4-oxo-13-cisRA in human liver, MS/MS spectra were collected for the tentative peaks and compared with the spectrum collected for the reference standard (Fig. 4). The collected spectra showed a fragmentation pattern of the tentative 4-oxo-atRA in human liver that matched with the spectrum of 4-oxo-atRA synthetic standard, confirming that the identified peak was indeed 4-oxo-atRA. 4-oxo-13-cisRA was also detected in human liver and an MS/MS spectrum was collected for this analyte as well. Owing to the low signal intensity of 4-oxo-13-cisRA in human liver (Fig. 4), high background noise was detected in the spectrum of the peak eluting at the same retention time as 4-oxo-13-cisRA standard, making the identification of this peak equivocal. However, product ions of m/z 137, 147, 159, and 241 were detectable, suggesting that the peak is 4-oxo-13-cisRA (Fig. 4). In mouse liver, 4-oxo-atRA was detected but its concentration was below LLOQ, and 4-oxo-13-cisRA was not detected (Fig. 4).

Method Validation. On the basis of preliminary experiments, the concentrations of RP and retinol were >1000-fold higher than those of RA isomers and 4-oxo-atRA within the human liver. RP was present in pooled liver homogenates (n = 6) at a concentration of 66 nmol/g of liver and retinol was about 2 nmol/g of liver, and the concentrations of atRA and 13-cisRA were about 140 and 39 pmol/g of liver, respectively. Hence, it is impossible to quantify the vitamin A metabolome in a single analytical LC-MS/MS run owing to the dynamic

![Fig. 3. Detection of atRA and 13-cisRA in human liver.](http://example.com/image3.png)
range limitations of modern LC-MS/MS instruments. In this study, the high-concentration retinoids (retinol and RP) were analyzed separately after protein precipitation with acetonitrile, and the acidic metabolites (RA isomers and 4-oxo-RA) were analyzed in two separate methods owing to the requirement of different organic extraction limits.
The methods for quantification of vitamin A and its metabolites were validated using isotope-labeled retinoids as internal standards. On-column LOD and LLOQ of measured retinoids (in the absence of matrix) were 13 and 27 fmol for arRA, 19 and 38 fmol for 13-cisRA, 26 and 49 fmol for 4-oxo-arRA, 15 and 30 fmol for retinol, and 30 and 60 fmol for RP, respectively. The slopes of the standard curves were different between the two matrices, demonstrating the confounding effect of using nonidentical chemical internal standards (4-oxo-atRA-d₃) for normalization. It is probable that the chromatographic separation of 13-cis- and all-trans-isomers results in different matrix effects for the two isomers, confounding the quantification of 4-oxo-13-cisRA when 4-oxo-atRA-d₃ is used as the internal standard. Thus, in the current study, 4-oxo-13-cisRA was not quantified even though the analyte was detectable in all liver samples.

The method validation results are shown in Table 2. Considering the biologic variability of retinoid concentrations in human tissues, the ranges of standard curves were designed to be wide enough to cover retinoid concentrations in all measured liver samples. All standard curves were linear within the ranges analyzed, with $R^2$ values higher than 0.99. The percentage of error of all QCs was less than 15% compared with true values. Interday and intraday CV% of all QCs and LQCs including the pooled liver sample showed that variability within day and between days was less than 15%, confirming the reproducibility of the method.

**The Vitamin A Metabolome in Normal Human Livers.** Hepatic concentrations of each measured retinoid in 50 normal liver samples showed log-normal distribution, with high variability but with no identified outliers in the dataset (Fig. 5). Retinol $(n = 50)$, RP $(n = 50)$, arRA $(n = 50)$, 13-cisRA $(n = 50)$, and 4-oxo-arRA $(n = 48)$ were all detected in the liver samples analyzed. Hepatic concentrations of retinol and RP were in the range of nmol/g of liver, much higher than other measured retinoids, which were in the range of picomoles per gram of liver. On the basis of the median concentrations (Fig. 5; Table 3), hepatic RP concentration was about 19-fold higher than retinol. Compared with arRA concentrations in the liver, 13-cisRA had lower median concentration with less interindividual variability. The hepatic 4-oxo-arRA concentration was consistently lower than arRA (45 of 48 livers) with only two liver samples having higher 4-oxo-arRA concentrations than arRA and one sample having similar arRA and 4-oxo-arRA concentrations. The arRA concentration in normal human livers was positively correlated with the concentration of RP, retinol, 13-cisRA,

<table>
<thead>
<tr>
<th></th>
<th>Normal $(n = 50)$</th>
<th>Steatosis $(n = 9)$</th>
<th>NASH $(n = 13)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>RP (nmol/g)</td>
<td>Median</td>
<td>Range</td>
<td>Median</td>
</tr>
<tr>
<td></td>
<td>276</td>
<td>12.6–4282</td>
<td>59.7</td>
</tr>
<tr>
<td>Retinol (nmol/g)</td>
<td>14.1</td>
<td>1.2–161</td>
<td>8.1</td>
</tr>
<tr>
<td>arRA (pmol/g)</td>
<td>42.6</td>
<td>14.0–579.5</td>
<td>29.8</td>
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<tr>
<td>13-cisRA (pmol/g)</td>
<td>8.3</td>
<td>1.4–43.5</td>
<td>5.1</td>
</tr>
<tr>
<td>4-oxo-arRA (pmol/g)</td>
<td>18.3*</td>
<td>1.8–378.3*</td>
<td>2.5</td>
</tr>
</tbody>
</table>

*The RP concentration in one NASH sample was below LOD. Hence, the RP concentration in that sample was excluded.

*4-Oxo-arRA concentrations were measured in 48 samples in the normal group.
The group of normal livers includes \( n = 50 \) livers free of known NAFLD pathologies. The normal and NAFLD group includes all the livers in the normal group and the additional livers with pathologically characterized fatty liver and cirrhosis (Supplemental Tables S1 and S2).

### TABLE 4

Spearman correlation of \( a \)-RA concentration with other retinoid concentrations and the mRNA expression of retinoid-related genes in human livers

<table>
<thead>
<tr>
<th>Retinoid</th>
<th>Normal only (( n = 50 ))</th>
<th>Normal and NAFLD (( n = 72 ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>RP</td>
<td>0.48</td>
<td>0.48</td>
</tr>
<tr>
<td>Retinol</td>
<td>0.43</td>
<td>0.48</td>
</tr>
<tr>
<td>13-cis-RA</td>
<td>0.83</td>
<td>0.82</td>
</tr>
<tr>
<td>4-oxo-RA</td>
<td>0.82</td>
<td>0.71</td>
</tr>
<tr>
<td>LRAT</td>
<td>0.26</td>
<td>0.067</td>
</tr>
<tr>
<td>ALDH1A1</td>
<td>−0.29</td>
<td>0.046</td>
</tr>
<tr>
<td>CYP26A1</td>
<td>0.14</td>
<td>0.25</td>
</tr>
<tr>
<td>RAR( \alpha )</td>
<td>−0.14</td>
<td>0.34</td>
</tr>
<tr>
<td>RAR( \beta )</td>
<td>0.02</td>
<td>0.92</td>
</tr>
</tbody>
</table>

\( ^a \)P < 0.005; \( ^b \) Bonferroni adjusted \( P \) value of 0.005 (0.05/9) was used to indicate significance for normal liver samples.

\( ^c \)P < 0.01; \( ^d \) Bonferroni adjusted \( P \) value of 0.01 (0.05/4) was used to test for significance of correlation for normal and NAFLD samples.

and 4-oxo-\( a \)-RA, with \( P \) values < 0.005 (Table 4). The correlation between hepatic concentration of retinol and RP was also analyzed using Spearman rank correlation, and showed an \( r \) of 0.36 (\( P = 0.011 \)). In an exploratory analysis of sex differences in retinoid metabolome no differences were observed between male and female donors for any of the retinoids (data not shown). Likewise, no correlation was observed between donor age and retinoid concentrations in the liver. This lack of correlation and sex differences may be owing to the retrospective nature of the analysis and lack of ability to adjust for covariates in the analysis.

**Vitamin A Homeostasis Is Disrupted in Liver Samples with NAFLD Compared with Normal Liver Samples.**

Intended retinoids were detected in all NAFLD samples (nine steatosis and 13 NASH) except one NASH liver sample in which RP concentration was below LOD (in three repeated, independent analyses). The RP concentration in this liver was excluded from the data shown in Fig. 6A. Hepatic \( a \)-RA, 13-cis-RA, and RP concentrations were significantly lower in NASH liver samples than in normal liver samples and there was a trend toward lower concentrations in steatosis samples, but this difference did not reach statistical significance (Fig. 6; Table 3). In addition, compared with liver samples in the normal group, both steatosis and NASH samples had significantly lower concentrations of 4-oxo-\( a \)-RA (Fig. 6). Together these data demonstrate significant retinoid depletion in NASH and indicate a progressive depletion of retinoids with NAFLD severity. No significant differences in hepatic ROL concentrations were observed between the normal and steatosis livers or between the normal and NASH group, indicating that ROL concentrations are maintained in the liver despite depletion of other retinoids (Fig. 6B; Table 3). The positive correlation between hepatic \( a \)-RA and other retinoid concentrations (RP, 13-cis-RA, and 4-oxo-\( a \)-RA) observed in normal livers remained unchanged when NAFLD samples were included. However, the correlation between \( a \)-RA and retinol was no longer significant when NAFLD samples were included (Table 4), suggesting that independent measurements of the different retinoids are needed to assess retinoid status in disease states.

The Relative mRNA Expression of Retinoid-Related Genes in Normal and NAFLD Human Livers. The relative mRNA expression of several retinoid-related genes was measured in 48 normal liver samples (Fig. 5). This included analysis of genes responsible for hepatic retinol esterification (LRAT), for RA biosynthesis (ALDH1A1) and elimination (CYP26A1), and genes of nuclear receptors that RA and 4-oxo-RA isomers bind to and induce (RAR\( \alpha \) and RAR\( \beta \)). The relative mRNA expression of measured genes was variable in livers except for LRAT, for which the expression was fairly consistent. Surprisingly, none of the target gene mRNA expressions showed a significant correlation with the hepatic \( a \)-RA concentration (Table 4), demonstrating that
mRNA expression should not be analyzed as a surrogate for retinoid concentrations. In the comparisons among the normal, steatosis, and NASH groups, the relative mRNA expression of ALDH1A1 was significantly higher in the NASH livers than normal livers, whereas RBP4 mRNA was significantly lower in the NAFLD livers compared with control (Supplemental Fig. S2). The relative mRNA expression of LRAT, CYP26A1, and RARβ was similar in the NAFLD and NASH groups compared with the normal (Supplemental Fig. S2).

Discussion

The goal of this study was to determine whether atRA concentrations in normal human livers and livers from donors with NAFLD correlate with commonly used markers of altered vitamin A homeostasis such as liver retinol and RP concentrations and retinoid-related mRNAs. Novel LC-MS/MS methods were developed for quantification of vitamin A metabolome in human liver. It is well known that different biologic matrices result in distinct interferences affecting analyte identification, recovery, matrix effects, and quantification. In addition, interferences from same tissue can vary with disease stages and between individuals. For example, NAFLD liver samples contain higher lipid content than normal liver samples, and liver tissue becomes fibrotic as the disease progresses. This may affect analyte recovery, ionization, and detection. Hence, accurate characterization of effects of NAFLD on hepatic vitamin A metabolome relies on good LC-MS/MS methods and appropriate internal standards to normalize extraction recovery and matrix effects. Validating bioanalytical methods is, however, particularly challenging for vitamin A metabolome as retinoids are endogenous molecules, and blank tissues do not exist for method development. It has been suggested that exposing tissues to light can be used to generate retinoid-depleted matrix such as serum (Arnold et al., 2012), but we did not observe a significant change in RA isomer concentrations in human liver samples after they were exposed to visible light for 1 hour (data not shown). This was probably the result of the low matrix transparency of the liver and high protein binding of retinoids. Therefore, to allow normalization for extraction recovery of the retinoids using isotope-labeled internal standards, LC-MS/MS methods for vitamin A metabolome were developed using retinoid-free charcoal-treated serum as a reference matrix and deuterated internal standards. The validity of this approach was confirmed by the determination that the slopes of the standard curves from liver and charcoal-treated serum were equal. Recovery of hepatic retinol and retinyl esters using the acetonitrile protein precipitation method developed here was confirmed by also measuring the concentrations of retinol and RP in 49 of 50 normal livers by the traditional hexane-extraction method (Obrochta et al., 2014; Grizotte-Lake et al., 2018) combined with isotope-labeled internal standards (Supplemental Fig. S3). No significant differences were observed between these methods.

Using the novel LC-MS/MS methods, human hepatic vitamin A metabolome was characterized for the first time in a relatively large sample of normal livers. The median concentrations of human hepatic retinol and RP reported in the current study are comparable with those measured previously by LC-UV methods in smaller samples (Schmitz et al., 1991; Schmidt et al., 2003). However, the analysis of human liver vitamin A metabolome discovered several species differences between human and mouse liver. The concentrations of atRA (14.0–579.5 pmol/g) and 13-cis-RA (1.4–43.5 pmol/g) in human livers were 3- to 100-fold higher than those reported in mouse livers, in which atRA concentrations are about 5–20 pmol/g and 13-cis-RA is below the limit of detection (Obrochta et al., 2014; Grizotte-Lake et al., 2018). This difference may have biologic significance, as laboratory mice may be more prone to RA deficiency than humans owing to the lower baseline RA concentrations. In addition, 4-oxo-atRA, an active retinoid that binds to retinoic acid receptors, was for the first time detected in the liver. This is a significant finding as 4-oxo-atRA has been shown to be more potent retinoid than atRA in anteroposterior axis formation and bind to RARs with affinity similar to atRA (Pijnappel et al., 1993; Idres et al., 2002; Topletz et al., 2015). In the human liver the concentrations of 4-oxo-atRA were comparable to that of atRA and 13-cis-RA. However, the concentrations of 4-oxo-atRA were below the limit of quantitation in mouse livers and 4-oxo-13-cis-RA was undetectable (Fig. 4). The lack of detection of 4-oxo-RA isomers in the liver previously is probably a result of this lack of 4-oxo-RA in mouse liver and the poor extraction recovery of 4-oxo-RA isomers by the common hexane extraction used for RA isomers in various tissues. The differences in RA and 4-oxo-atRA concentrations between human and mouse livers are surprising, as the retinyl esters and retinol concentrations in human livers (retinol 14.1 nmol/g, RP 276 nmol/g) were similar to what have been reported in mouse livers (9.6 nmol/g retinol and 560 nmol/g retinyl esters) (Kane et al., 2008). However, the variability in retinyl ester concentrations in human livers was much greater than that in mouse livers. Further studies are needed to address the apparent species differences, the biochemical processes that drive these differences, and the consequences in retinoid signaling in various disease models.

Serum retinol and RBP4 concentrations, and liver retinol and retinyl ester concentrations, together with mRNA expression of retinoid-related genes are often used as indirect measures of altered atRA concentrations and signaling, as dietary vitamin A deficiency has been shown to lead to decreased concentrations of these retinoids and decreased retinoid signaling (Tanumihardjo, 2012). We aimed to test whether, in a sufficiently large set of human livers, these findings of correlations between retinoid concentrations and signal-
expression is insufficient to reveal potential alterations in vitamin A metabolism. The data also suggest that atRA-induced changes in mRNA expression of these genes may only be captured within a specific time-window after an intervention. Of note, in the NAFLD samples, ALDH1A1 mRNA was significantly increased, whereas atRA concentrations were decreased, potentially owing to feedback regulation of ALDH1A1 mRNA in the liver. This finding is also in agreement with the weak negative correlation between atRA concentrations and ALDH1A1 mRNA in normal livers although causality of these observations cannot be defined.

Circulating concentrations of retinoids have also been used as markers of altered retinoid signaling in disease states. However, the data collected here from human liver suggest that serum retinoids do not reflect tissue retinoid profiles. The median liver concentrations of atRA and 13-cisRA were 2- to 10-fold higher than what has been reported in human systemic circulation (Arnold et al., 2012). In addition, although the median concentration of 13-cisRA in human serum is higher than atRA, the liver atRA concentrations were about 5-fold higher than those of 13-cisRA. While these findings are not from matched donors and hence may be confounded, they strongly suggest that if circulating retinoid profiles are used in clinical studies as biomarkers of retinoid homeostasis, more research is needed to establish the biochemical processes that regulate the distribution of retinoids between tissues and systemic circulation. It is possible that interconversion kinetics between isomers in various tissues affect RA isomer distribution. Likewise, differences in the binding affinity to cellular (and plasma)-binding proteins may drive differences in tissue-to-serum concentration ratios. Previous studies have shown that atRA exhibits much higher binding affinity to cellular retinoid acid binding proteins (K<sub>D</sub> < 20 nM) than does 13-cisRA (K<sub>D</sub> > 150 nM) (Fiorella and Napoli, 1991; Fiorella et al., 1993). These differences may drive higher tissue partitioning of atRA compared with 13-cisRA and cause higher cellular unbound fractions of 13-cisRA than do the all-trans isomers.

The findings shown here of depleted vitamin A stores and atRA concentrations in NAFLD are in agreement with previous findings in mouse models. Progression to vitamin A deficiency in NAFLD is believed to be associated with hepatic stellate cell differentiation and reduced retinyl ester storage (Shirakami et al., 2012; Saeed et al., 2017). The current study strongly supports the model that during disease progression in NAFLD, hepatic retinyl ester, atRA, 13-cisRA, and 4-oxo-atRA concentrations are progressively diminished. However, unexpectedly, despite the significant depletion of retinyl esters and atRA, hepatic retinol concentrations were unchanged regardless of the disease progression. It is important to note that these observations were not collected as a prospective study, and prospective studies designed and powered on the basis of these observations are needed to further establish the progression and the extent of depletion of vitamin A storage in NAFLD.

In conclusion, the human hepatic vitamin A metabolome was quantified using novel LC-MS/MS methods and deteriorated internal standards, providing a foundational understanding of the retinoid homeostasis in normal human liver. Additionally, the alterations of hepatic vitamin A metabolome were also defined in NAFLD liver samples to reveal the disease effects on vitamin A deficiency. These data show that vitamin A metabolome is profoundly altered in NAFLD, and that the complete vitamin A metabolome should be analyzed to characterize disease effects on retinoid signaling and concentrations.

**References**


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