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Identification of unusual oxysterols and bile acids with 7-oxo or 3β,5α,6β-trihydroxy functions in human plasma by charge-tagging mass spectrometry with multistage fragmentation

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Running title: Identification of unusual oxysterols and bile acids

Abbreviations: ChOx, cholesterol oxidase; CTX, cerebrotendinous xanthomatosis; CYP, cytochrome P450; GP, Girard P; LALD, lysosomal acid lipase deficiency; LC-MS, liquid chromatography – mass spectrometry; MRM, multiple reaction monitoring; MS3, mass spectrometry with multistage fragmentation; NP, Niemann-Pick; RIC, reconstructed ion-chromatogram; SLOS, Smith-Lemli-Opitz syndrome; SPE, solid phase extraction; 3βH,7o-CA, 3β-hydroxy-7-oxocholest-5-en-(25R)26-oic acid; 3βH,7o-Δ5-BA, 3β-hydroxy-7-oxochol-5-enoic acid; 3βH,7,24-diO-CA, 3β-hydroxy-7,24-bisoxocholest-5-en-26-oic acid; 3β,5α,6β-triHBA, 3β,5α,6β-trihydroxycholanoic acid; 3β,5α,6β-triHCA, 3β,5α,6β-trihydroxycholestan-(25R)26-oic acid; 3β,5α,6β-triH,H; 3β,5α,6β-trihydroxycholestan-26-oic acid; 3β,5α,6β-triH-triol; cholestane-3β,5α,6β-triol; 3β,5α,6β,24-tetraHCA, 3β,5α,6β,24-tetrahydroxycholestan-26-oic acid; 3β,5α,6β-triH,24O-CA, 3β,5α,6β-trihydroxy-24-oxocholestan-26-oic acid; 3β,5α,6β,26-tetrol, cholestane-3β,5α,6β,(25R)26-tetrol; 3β,7α-diHCA, 3β,7α-dihydroxycholesterol-5-en-(25R)26-oic acid; 3β,7α-diHCA(25S), 3β,7α-dihydroxycholesterol-5-en-(25S)26-oic acid; 3β,7α,12α-triHCA, 3β,7α,12α-trihydroxycholesterol-5-en-
(25R)26-oic acid; 3β,7α,24S-triHCA, 3β,7α,24S-trihydroxycholest-5-en-(25R)26-oic acid; 3β,7α,25-triHCA, 3β,7α,25-trihydroxycholest-5-en-26-oic acid; 3β,7β-diHCA(25S), 3β,7β-dihydroxycholest-5-en-(25S)26-oic acid; 3β,22,25-triH,24O-C, 3β,22,25-trihydroxycholest-5-en-24-one; 3β,24-diH,7O-CA, 3β,24-dihydroxy-7-oxocholest-5-en-26-oic acid; 5,6-EC, 3β-hydroxycholestan-5,6-epoxide; 6β-HC, cholest-4-ene-3β,6β-diol; 7-DHC, cholesta-5,7-dien-3β-ol; 7-OC, 3β-hydroxycholest-5-en-7-one; 7α-HC, cholest-5-en-3β,7α-diol; 7α-HCO, 7α-hydroxycholesta-4-ene-3-one; 7αH,3O-CA, 7α-hydroxy-3-oxocholest-4-en-(25R)26-oic acid; 7αH,3O-Δ4-BA, 7α-hydroxy-3-oxocholesta-4-ene-3-one; 7α,12α-diHCO, 7α,12α-dihydroxycholesta-4-ene-3-one; 7α,24S-diHCO, 7α,24S-dihydroxycholesta-4-ene-3-one; 7α,24S-diH,3O-CA, 7α,24S-dihydroxy-3-oxocholest-4-en-(25R)26-oic acid; 7α,25-diH,3O-CA, 7α,25-dihydroxy-3-oxocholest-4-en-26-oic acid; 7α,25-diHC, cholest-5-ene-3β,7α,25-triol; 7α,25-diHCO, 7α,25-dihydroxycholesta-4-ene-3-one; 7α,26-diHCO, 7α,(25R)26-dihydroxycholest-4-en-3-one; 22R-HCO, 22R-hydroxycholest-4-en-3-one; 24R/S-HC, cholest-5-en-3β,24R/S-diol; 25H-VitD3, 9,10-secocholesta-5Z,7E,10(19)-triene-3S,25-diol; 25H,7O-C, 3β,25-dihydroxycholest-5-en-7-one; 26H,7O-C, 3β,(25R)26-dihydroxycholesta-5-en-7-one.
Abstract

7-Oxocholesterol (7-OC), 5,6-epoxycholesterol (5,6-EC) and its hydrolysis product cholestane-3β,5α,6β-triol (3β,5α,6β-triol) are normally minor oxysterols in human samples, however, in disease their levels may be greatly elevated. This is the case in plasma from patients suffering from some lysosomal storage disorders e.g. Niemann Pick disease type C, or the inborn errors of sterol metabolism e.g. Smith-Lemli-Opitz syndrome and cerebrotendinous xanthomatosis. A complication in the analysis of 7-OC and 5,6-EC is that they can also be formed ex vivo from cholesterol during sample handling in air causing confusion with molecules formed in vivo. When formed endogenously 7-OC, 5,6-EC and 3β,5α,6β-triol can be converted to bile acids. Here, we describe methodology based on chemical derivatisation and liquid chromatography – mass spectrometry with multistage fragmentation (MS^n) to identify the necessary intermediates in the conversion of 7-OC to 3β-hydroxy-7-oxochol-5-enoic acid and 5,6-EC and 3β,5α,6β-triol to 3β,5α,6β-trihydroxycholanoic acid. Identification of intermediate metabolites is facilitated by their unusual MS^n fragmentation patterns. Semi-quantitative measurements are possible, but absolute values await the synthesis of isotope-labelled standards.

Key Words: Sterols; cholesterol/metabolism; Niemann-Pick type C; oxidised lipids; tandem mass spectrometry
Introduction

Until recently 7-oxocholesterol (7-OC), 5,6-epoxycholesterol (5,6-EC) and its hydrolysis product cholestane-3β,5α,6β-triol (3β,5α,6β-triol) were regarded by many as artefacts generated by sample handling of cholesterol rich material in air (1-4). For a list of sterol abbreviations see Supplemental Table S1. That view has changed with the discovery that 7-OC can be generated enzymatically from the cholesterol precursor 7-dehydrocholesterol (7-DHC) by cytochrome P450 (CYP) 7A1 (5) and is abundant in plasma of patients with Smith-Lemli-Opitz Syndrome (SLOS), where levels of 7-DHC are high, and cerebrotendinous xanthomatosis (CTX) where CYP7A1 is highly expressed (6, 7). Furthermore, in patients with lysosomal storage disorders Niemann-Pick (NP) disease types C and B and lysosomal acid lipase deficiency (LALD), 3β,5α,6β-triol is elevated in plasma as is 7-OC, despite apparently normal levels of 7-DHC (8-13). For an up-to-date review see reference (14). Importantly, recent reports by Clayton and colleagues in London (15) and Ory and colleagues in St Louis (16) have documented the presence of the unusual bile acid 3β,5α,6β-trihydroxycholanoic acid (3β,5α,6β-triHBA) in plasma of NPC patients, while Alvelius et al and Maekawa et al have reported the presence of the sulphuric acid and glycine conjugates of 3β-hydroxy-7-oxochol-5-enoic acid (3βH,7O-Δ5-BA) in urine and plasma of NPC patients (17, 18). The observation of these unusual bile acids associated with NPC and of 3βH,7O-Δ5-BA with other disorders e.g. SLOS (19-21) strongly suggests that their precursors 7-OC and 5,6-EC are formed in vivo and are not (only) ex vivo artefacts generated through sample handling in air. Both 7-OC and 5,6-EC are dietary oxysterols (22, 23), while 5,6-EC may also be formed by environmental ozone in lung (24), representing alternative sources of these molecules in healthy individuals. In fact Lyons et al showed that 7-OC was rapidly metabolised by the liver in rats and excreted into the intestine mainly as aqueous soluble metabolites, presumably bile acids (22). Pulfer and Murphy showed that 5,6-EC was the major cholesterol-derived product formed in the reaction of ozone with lung surfactant and that 3β,5α,6β-triol, and more abundant levels of an unexpected metabolite, 3β,5α-dihydroxycholestan-6-one were formed from 5,6-EC by lung epithelial cells (24).

To investigate how 7-OC is metabolised in vivo into 3βH,7O-Δ5-BA and 5,6-EC and 3β,5α,6β-triol into 3β,5α,6β-triHBA, as reported in the accompanying manuscript (Griffiths W.J. et al), we have optimised a charge-tagging methodology to specifically identify 7-oxo containing sterols and sterols with a 3β,5α,6β-
triol function using chemical derivatisation and liquid chromatography - mass spectrometry (LC-MS) with multistage fragmentation (MS^n). The resultant method is described below.
Materials and Methods

Human Samples

Plasma was from patients diagnosed with lysosomal storage disorders, their siblings or parents. All participants or their parents provided written informed consent in accordance with the Declaration of Helsinki and the study was conducted with institutional review board approval (REC08/H1010/63). NIST standard reference material (SRM1950, Gaithersburg, MD), a pooled plasma sample representative of the US population (25), was used as a reference.

Materials

Oxysterols and C27 bile acids were from Avanti Polar Lipids Inc. (Alabaster, AL, USA); C24 bile acids were a kind gift from Professor Jan Sjövall, Karolinska Institutet, Stockholm, Sweden; 3β,5α,6β-triHBA was a kind gift from Professor Douglas F. Covey, Washington University School of Medicine. See Supplemental Table S1 for a list of oxysterols and bile acid with their common and systematic names, abbreviations, LipidMaps ID and suppliers. Cholesterol oxidase (ChOx) enzyme from Streptomyces sp was from Sigma Aldrich Ltd (Dorset, UK), [2H7]Girard P ([2H7]GP) reagent was from TCI Europe (Zwijndrecht, Belgium), [2H5]GP was synthesised as described in (26). Reversed-phase Certified Sep-Pak C18 (200 mg) and Oasis HLB (60 mg) solid phase extraction (SPE) columns were from Waters Ltd (Elstree, Herts, UK).

Sample Preparation

ethanol with 0.35 mL of water, sonicated and centrifuged to remove precipitated matter. To separate bile acids and oxysterols from cholesterol and similarly hydrophobic sterols, the sample solution was applied to a 200 mg Sep-Pack C18 column, cholesterol was absorbed while oxysterols and bile acids eluted in the flow-through and column wash (SPE1-Fr1, 7 mL 70% ethanol). After a further column wash (SPE1-Fr2, 4 mL 70% ethanol), cholesterol, and sterols of similar hydrophobicity, were then eluted in a separate fraction with absolute ethanol (SPE1-Fr3, 2 mL). The oxysterol/bile acid (SPE1-Fr1) and cholesterol-rich (SPE1-Fr3) fractions were then each divided into two equal aliquots (A) and (B) and lyophilised. After reconstitution in propan-2-ol (100 µL), cholesterol oxidase (ChOx, 0.26 units) in 50 mM phosphate buffer (1 mL), pH 7, was added to sub-fractions (A). After 1 hr at 37 °C the reaction was quenched with methanol (2 mL). Sub-fractions (B) were treated in an identical manner but in the absence of cholesterol oxidase. Glacial acetic acid (150 µL) was added to each sub-fraction followed by [3H5]GP as the bromide salt (190 mg) to sub-fractions (A) and [3H5]GP as the chloride salt (150 mg) to sub-fractions (B). The derivatisation reactions were left to proceed overnight in the dark at room temperature. Excess derivatisation reagent was removed by SPE using a re-cycling method. Each sub-fraction (now 3.25 mL, 69% organic) was applied to a 60 mg Oasis HLB column and was washed with 70% methanol (1 mL) and 35% methanol (1 mL). The combined effluent was diluted to 35% methanol and re-cycled through the column. This was repeated with dilution to 17.5% methanol, a further re-cycling and the column was finally washed with 10% methanol (6 mL). At this point all oxysterols/bile acids, or more hydrophobic sterols, were absorbed on the column while unreacted GP reagent elute to waste. Oxysterols/bile acids were then eluted with 100% methanol (2 mL, SPE2-Fr1+2) while more hydrophobic sterols with 3 mL of 100% methanol (SPE2-Fr1+2+3). Just prior to LC-MS(MSn) analysis equal volumes of sub-fractions (A) and (B) were combined and diluted to 60% methanol ready for injection. This allowed the simultaneous analysis of sub-fractions A and B.

**LC-MS(MSn)**

Analysis was performed on an Orbitrap Elite (Thermo Fisher Scientific, Hemel Hempstead, UK) exploiting electrospray ionisation. Chromatographic separation was achieved with a reversed-phase Hypersil Gold column (1.9 µm particle size, 50 x 2.1 mm, Thermo Fisher) using an Ultimate 3000 LC system (now Thermo Fisher Scientific) with the mobile phase and gradient described in (26, 27). To separate some closely eluting
oxysterols/bile acids the usual 17 min gradient was extended to 37 min. The mobile phase composition was initially at 80% A (33.3% methanol, 16.7% acetonitrile, 0.1% formic acid), 20% B (63.3% methanol, 31.7% acetonitrile, 0.1% formic acid) for 10 min, changed to 50% A, 50% B over the next 10 min, maintained at this composition for 6 min, then changed to 20%A, 80% B over the next 3 min. The mobile phase composition was held at 20% A, 80% B for a further 3 min before returning to 80% A, 20% B in 0.1 min and reconditioning the column for a further 4.9 min.

For each injection five scan events were performed: one high resolution scan (120,000 full-width at half maximum height definition at m/z 400) in the Orbitrap analyser in parallel to four MS3 scan events in the linear ion-trap. Quantification was by isotope dilution or by using isotope-labelled structural analogues.
Results

7-Oxo Containing Sterols

7-OC is a $\alpha,\beta$-unsaturated ketone (5-en-7-one) and unlike oxysterols/bile acids without an oxo group will react with GP reagent in the absence of cholesterol oxidase, hence this compound and its metabolites possessing a 7-oxo group will be found in the (B) sub-fraction derivatised with the $[^{2}H_{6}]GP$ reagent (Figure 1B). Like most GP derivatives, 7-oxo compounds give an intense $[M-Py]^+$ ion upon MS$^{2}$ (MS/MS, Figure 1B), however, MS$^{3}$ ([M]$^{+}$→[M-Py]$^{+}$→) fragmentation patterns for the 7-oxo derivatives are unlike those from compounds with GP derivatisation at position C-3 (cf. Figure 2B & C). In contrast to 3-oxo compounds, 7-oxo compounds show a prominent pattern of fragment-ions corresponding to $[M-Py-43]^+$, $[M-Py-59]^+$, $[M-Py-90]^+$ and $[M-Py-98]^+$ (m/z 412.4, 396.3, 365.3 and 357.3). The suggested structures of these ions are shown in Figure 3 and Supplemental Figure S1A. As can be seen in Figure 3 the neutral-losses are associated with the unsaturated diazacyclohexanone ring and water, this is evident by the invariant nature of the neutral-loss upon variation of the sterol side-chain or by the incorporation of deuterium atoms in the side-chain e.g. as in $[25,26,26,26,27,27,27-^{2}H_{7}]7$-OC (cf. Figure 2B & Supplemental Figure S2A). Definitive identification of the neutral-losses was achieved by utilising $[^{13}C^{15}N]$ and $[^{13}C_{2}]$ isotope-labelled GP reagents (Figure 3). Neutral-losses, $[M-Py-18]^+$, $[M-Py-28]^+$ and $[M-Py-61]^+$ common to 7-oxo-5-ene and 3-oxo-4-ene derivatives are illustrated in Figure 3 and Supplemental Figure S3A.

7-OC is well resolved from its isomer 7α-hydroxycholesterol-4-en-3-one (7α-HCO) in the 17 min chromatographic gradient and is readily quantified by isotope dilution against $[^{3}H_{2}]7$-OC (Figure 2A), but 26-hydroxy-7-oxocholesterol (26H,7O-C), its CYP27A1 metabolite (28, 29), is only partially resolved from its isomer 7α,26-dihydroxycholesterol-4-en-3-one (7α,26-diHCO) by the 17 min chromatographic gradient, although it is well resolved from 25-hydroxy-7-oxocholesterol (25H,7O-C), 7α,12α-dihydroxycholesterol-4-en-3-one (7α,12α-diHCO), 7α,24S-dihydroxycholesterol-4-en-3-one (7α,24S-diHCO) and 7α,25-dihydroxycholesterol-4-en-3-one (7α,25-diHCO), four further metabolites identified in human plasma. Extending the chromatographic gradient to 37 min provides almost base line separation of 26H,7O-C from 7α,26-diHCO while maintaining resolution from the other isomers (Figure 2D). In the absence of an ideal isotope-labelled
standard, approximate quantification of 26H,7O-C is performed against the internal standard \( ^2H_7 \)22R-HCO, taking into account relative response factors.

The down-stream CYP27A1 metabolite of 26H,7-OC, 3\beta\)-hydroxy-7-cholesterol-5-en-26-oic acid (3\betaH,7O-CA) (29) is not resolved from its isomer 7\alpha\)-hydroxy-3-cholesterol-4-en-26-oic acid (7\alphaH,3O-CA) in either the 17 min or 37 min chromatographic gradients (Figure 4A). However, 3\betaH,7O-CA gives a MS\(^3\) fragment-ion at m/z 426.3 ([M-59]\(^+\)) that is not present in the MS\(^3\) spectrum of co-eluting 7\alphaH,3O-CA (Figure 4C - 4E). Thus, by generating a reconstructed-ion chromatogram (RIC) for m/z 426.3±0.3 from the MS\(^3\) spectra, 3\betaH,7O-CA is revealed (Figure 4B). Therefore, multiple reaction monitoring (MRM), [M]\(^+\)→[M-Py]\(^+\)→[M-Py-59]\(^+\), can be used to resolve 3\betaH,7O-CA from its co-eluting isomer 7\alphaH,3O-CA. The fragment-ion at m/z 421.3 ([M-Py-64]\(^+\)) is present in the MS\(^3\) spectrum of 7\alphaH,3O-CA (Figure 4E) but not 3\betaH,7O-CA (Figure 4C), hence the MRM [M]\(^+\)→[M-Py]\(^+\)→[M-Py-64]\(^+\) can identify 7\alphaH,3O-CA. By necessity, semi-quantification of 3\betaH,7O-CA is performed utilising the MRM transition 564.4→485.3→426.3 and reference to an external standard.

In bile acid biosynthesis side-chain shortening occurs in the peroxisome and proceeds through 24-hydroxylation of the C\(_{27}\) acid, dehydrogenation to a 24-carbonyl group then beta-oxidation to a C\(_{26}\) acid (30). The appropriate metabolites from 7-OC are 3\beta,24-dihydroxy-7-cholesterol-5-en-26-oic acid (3\beta,24-dih,7O-CA), 3\beta-hydroxy-7,24-bisoxocholesterol-5-en-26-oic acid (3\betaH,7,24-dio-CA) and 3\betaH,7O-\( \Delta^5 \)-BA. Although authentic standards are not available for these metabolites, by analogy to the MS\(^3\) spectra of 7-OC, 26H,7-OC and 3\betaH,7O-CA where standards are available, prominent patterns of distinguishing fragment-ions corresponding to the neutral-losses [M-Py-43]\(^+\), [M-Py-59]\(^+\), [M-Py-90]\(^+\) and [M-Py-98]\(^+\) are predicted to be present in the MS\(^3\) spectra of 3\beta,24-dih,7O-CA, 3\betaH,7,24-dio-CA and 3\betaH,7O-\( \Delta^5 \)-BA. Similar to the analysis of 3\betaH,7O-CA, a MS\(^3\) RIC for [M-Py-59]\(^+\) i.e. m/z 442.3±0.3, should reveal 3\beta,24-dih,7O-CA (Figure 4F). A fragment-ion at m/z 442.3 is absent from the MS\(^3\) spectra of 7\alpha,24S-dihydroxy-3-cholesterol-4-en-(25R)26-oic (7\alpha,24S-dih,3O-CA) and 7\alpha,25-dihydroxy-3-cholesterol-4-en-26-oic (7\alpha,25-dih,3O-CA) acids, two commercially available isomers of 3\beta,24-dih,7O-CA (Supplemental Figure S2I & S2J). While for most plasma samples we have analysed in recent times the MS\(^3\) RIC channel for m/z 442.3±0.3 is empty (27), when patient samples containing high levels of 7-OC from diseases such as NPC are analysed a peak is
evident in the MRM chromatogram m/z 580.4→501.3→442.3 i.e. [M]+→[M-Py]+→[M-Py-59]+ (Figure 4F) and the underlying MS³ spectrum is compatible with that predicted for 3β,24-diH,7O-CA (Figure 4G). In the extended chromatographic gradient of 37 min, presumptively identified 3β,24-diH,7O-CA is resolved from isomeric dihydroxy-3-oxocholest-4-en-26-oic acids (Supplemental Figure S2K). Semi-quantitative measurement of 3β,24-diH,7O-CA are made using the extended chromatographic gradient assuming the same response factor as for 7α,24-diH,3O-CA and using the internal standard [²H₇]22R-HCO.

To-date in none of the plasma samples we have analysed have we observed any chromatographic peaks compatible with 3βH,7,24-diO-CA. In contrast, as with 3β,24-diH,7O-CA, an MS³ RIC for [M-Py-59]+ (m/z 384.3) reveals 3βH,7O-Δ⁵-BA in plasma samples from patients with elevated 7-OC (Figure 5B). 3βH,7O-Δ⁵-BA is clearly resolved from its isomer 7α-hydroxy-3-oxochol-4-enoic acid (7αH,3O-Δ⁴-BA) in the 17 min chromatographic gradient (Figure 5A). Semi-quantitative measurement are made for 3βH,7O-Δ⁵-BA using the extended chromatographic gradient assuming the same response factor as for 7αH,3O-Δ⁴-BA and using the internal standard [²H₇]22R-HCO.

While GP-derivatised sterols with a 7-hydroxy-3-oxo-4-ene structure give a characteristic pattern of ring fragment ions at m/z 151.1 (*b₁-12), 177.1 (*b₂), 179.1 (*b₃-28) and 231.1 (*c₁+2-18) (Figures 2C, 2E, 4E, 5C) (31), sterols with a 3β-hydroxy-7-oxo-5-ene structure give a minor fragment-ion at m/z 157.1 which probably consists of the unsaturated diazacyclohexanone ring and remnants of the B-ring (Figures 2B, 4C, 4D, 5D). This ion is only minor and of limited diagnostic value.

Semi-Quantitative Measurements

Using the methodology described, other than for 7-OC where an isotope labelled standard is available i.e. [²H₇]7-OC, we can only make approximate or semi-quantitative measurements. However, as all the 7-oxo compounds, except 3βH,7O-CA, are resolved from their 3-oxo isomers in either the 17 min or 37 min chromatographic gradients, quantification is possible using the isotope-labelled internal standard [²H₇]22R-HCO. In the absence of authentic standards, 3β,24-diH,7O-CA and 3βH,7O-Δ⁵-BA were quantified assuming the same response factors as for their structural analogues 7α,24-diH,3O-CA and 7αH,3O-Δ⁴-BA. As 3βH,7O-
CA was not chromatographically resolved from 7αH,3O-CA, the MRM [M+]→[M-Py]+→[M-Py-59]+ was used for quantification of the former isomer.

3β,5α,6β-Trihydroysterols

Sterols with a 3β-hydroxy group and a planar A/B ring system are substrates for cholesterol oxidase, these include cholest-5-en-3β-ols and 5α-cholestan-3β-ols (32). 3β,5α,6β-triol is planar and becomes oxidised at C-3 and also dehydrated through elimination of the 5α-hydroxy group. The dehydration reaction does not go to completion under our experimental conditions, so after derivatisation the GP derivatised triol is observed as both [M]+ and [M-H2O]+ ions (Figure 1A) (33), the [M-H2O]+ ion giving the stronger signal and more informative MS3 spectrum (Figure 6B). In fact, the MS3 ([M-H2O]+→[M-H2O-Py]+→) spectrum of the 3β,5α,6β-triol is identical to the MS3 ([M]+→[M-Py]+→) spectrum of cholest-4-ene-3β,6β-diol (6β-HC) confirming dehydration through loss of the 5α-hydroxy group. An unusually prominent fragment-ion observed in the MS3 ([M-H2O]+→[M-H2O-Py]+→) spectrum of 3β,5α,6β-triol is at m/z 383.3, corresponding to [M-H2O-Py-72]+ (Figure 6B, see also Figure 7). A second unusual neutral-loss [M-H2O-Py-100]+ gives a fragment-ion at m/z 355.3. Both fragment-ions are elevated by 7 Da in the spectrum of the [25,26,26,27,27,27-2H7] analogue as is the [M-H2O-Py-90]+ fragment-ion (see Supplemental Figures S4A, S5C & S5D). The availability of [3H2]3β,5α,6β-triol allows quantification by isotope dilution utilising RICs for [M-H2O]+ ions. Similar to 3β,5α,6β-triol, 3β,5α,6β-triHBA, the end product of 3β,5α,6β-triol metabolism (15, 16), gives [M]+ and [M-H2O]+ ions after GP derivatisation, the latter of which is dominant. The MS3 ([M-H2O]+→[M-H2O-Py]+→) spectrum of 3β,5α,6β-triHBA shows a prominent [M-H2O-Py-72]+ fragment-ion at m/z 371.3, [M-H2O-Py-90]+ at m/z 353.2 and the unusual neutral-loss [M-H2O-Py-100]+ at m/z 343.3 (Figure 6D, see also Supplemental Figures S4B & S5I). The particularly prominent neutral loss fragment-ion [M-H2O-Py-72]+ is common to MS3 spectra of both 3β,5α,6β-triol and 3β,5α,6β-triHBA and can potentially be used to identify further metabolites with a 3β,5α,6β-trihydroxy structure via [M-H2O]+→[M-H2O-Py]+→[M-H2O-Py-72]+ MRM chromatograms (see below). In the absence of an isotope-labelled standard, approximate quantification of 3β,5α,6β-triHBA is made against the internal standard [3H2]24R/S-HC, taking into account relative response factors.
In plasma samples high in 3β,5α,6β-triol (e.g. NPC), a new peak is evident in the RIC for the [M-H2O]+ ion of cholestanetetrols (m/z 555.4317 ± 5 ppm, 5.19 min in Figure 6E) which is not seen in control plasma. With both the 17 min and 37 min chromatographic gradients this peak is only partially resolved from the [M]+ ion of 7α,25-dihydroxycholesterol (7α,25-diHC) which has an identical mass. However, chromatographic resolution is sufficiently good to generate an MS3 (m/z 555.4→471.4→) spectrum from the apex of the new peak which is entirely compatible with that expected for the [M-H2O]+ ion of cholestan-3β,5α,6β,26-tetrol (3β,5α,6β,26-tetrol), showing a prominent fragment-ion at m/z 399.3 i.e. [M-H2O-Py-72]+, a distinct ion at m/z 381.3 i.e. [M-H2O-Py-90]+ and a minor fragment at m/z 371.3 i.e. [M-H2O-Py-100]+ (Figure 6G, see also Figure 7). The MS1 (m/z 555.4→471.4→) spectrum of closely eluting 7α,25-diHC does not give a fragment-ion at m/z 399.3 or 371.3 (Supplemental Figure S5L), so by generating a RIC for the fragment-ion m/z 399.3 from the MS3 (m/z 555.4→471.4→) chromatogram, 3β,5α,6β,26-tetrol ([M-H2O]+) is resolved from 7α,25-diHC ([M]+, Figure 6F). Other isomers of 7α,25-diHC ([M]+) i.e. the dihydroxycholesterols (diHC) 7α,12α-diHC, 7α,24S-diHC, 7α,26-diHC, 7β,25-diHC and 7β,26-diHC are all chromatographically resolved from 3β,5α,6β,26-tetrol ([M-H2O]+). It is only possible to make semi-quantitative measurement of 3β,5α,6β,26-tetrol in the absence of an authentic standard and its incomplete chromatographic resolution from 7α,25-diHC. Semi-quantification is made based on the [M-H2O]+ of 3β,5α,6β,26-tetrol against the internal standard [2H6]7α,25-diHC.

CYP27A1 is the enzyme likely to introduce the (25R)26-hydroxy group to the sterol side-chain. This enzyme could then oxidise the primary alcohol to a carboxylic acid to give 3β,5α,6β-trihydroxycholestan-(25R)26-oic acid (3β,5α,6β-triHCa). As discussed above, peroxisomal side-chain shortening of C27 acids proceeds through C-24 hydroxylation, C-24 dehydrogenation and β-oxidation, to generate 3β,5α,6β-triHBA as the ultimate product. The relevant pathway intermediates would be 3β,5α,6β,24-tetrahydroxycholestan-26-oic acid (3β,5α,6β,24-tetraHCa) and 3β,5α,6β-trihydroxy-24-oxocholestan-26-oic acid (3β,5α,6β-triH,24O-Ca). By analogy to 3β,5α,6β-triol, 3β,5α,6β,26-tetrol and 3β,5α,6β-triHBA; 3β,5α,6β-triHCa, 3β,5α,6β,24-tetraHCa and 3β,5α,6β-triH,24O-Ca should give [M]+ and [M-H2O]+ products upon cholesterol oxidase treatment and GP derivatisation and the MS3 spectra of the [M-H2O]+ ions ([M-H2O]+→[M-H2O-Py]+→)) are predicted to show neutral-loss fragment-ions [M-H2O-Py-72]+, [M-H2O-Py-90]+ and [M-H2O-Py-100]+.
For 3β,5α,6β-triHCA the [M-H₂O-Py-72]⁺ ion has an m/z of 413.3 (see Supplemental Figure S4B). A RIC for the MRM transition [M-H₂O]⁺→[M-Py-H₂O]⁺→[M-H₂O-Py-72]⁺ (m/z 569.4→485.3→413.3) reveals a new chromatographic peak in samples where the concentration of the 3β,5α,6β-triol is high (e.g. NPC) which is at, or below, the detection limit in normal plasma samples (Figure 8B). The MS³ ([M-H₂O]⁺→[M-H₂O-Py]⁺→) spectrum underlying the new chromatographic peak in plasma samples rich in 3β,5α,6β-triol is compatible with that predicted for 3β,5α,6β-triHCA, showing the predicted neutral-loss fragment-ions [M-H₂O-Py-72]⁺, [M-H₂O-Py-90]⁺ and [M-H₂O-Py-100]⁺ and is thus assigned to this acid (Figure 8C, see also supplemental Figure S4B and S5N). This new chromatographic peak does, however, co-elute with that of the [M]⁺ ion of 3β,7β-dihydroxycholest-5-en-(25S)26-oic acid (3β,7β-diHCA(25S)) and a second oxysterol with a probable 3β,22,25-trihydroxycholest-5-en-24-one structure, but neither compound gives a fragment ion at m/z 413.3 in their MS³ ([M]⁺→[M-Py]⁺→) spectra (Supplemental Figure 5O), unlike well resolved 3β,7α-dihydroxycholest-5-en-(25S)26-oic (3β,7α-diHCA(25S)) and 3β,7α-dihydroxycholest-5-en-(25R)26-oic (3β,7α-diHCA(25R)) acids where this ion is more abundant (Supplemental Figure S5P). In fact, the 25R and 25S epimers give identical MS³ spectra, but are chromatographically resolved. In the absence of an authentic standard of 3β,5α,6β-triHCA only semi-quantification is possible. This can be made using the RIC for the [M-H₂O]⁺ ion in samples where co-eluting compounds 3β,7β-diHCA(25S) and 3β,22,25-triH,24O-C are minor (and assuming a similar response factor to 3β,5α,6β-triHBA), then determining an appropriate response factor for the MRM chromatogram 569.4→485.3→413.3 and using this for quantification of other samples.

For 3β,5α,6β,24-tetraHCA the [M-H₂O-Py-72]⁺ fragment ion has an m/z of 429.3 (Figure S4B). In the RIC for the MRM transition [M-H₂O]⁺→[M-Py-H₂O]⁺→[M-H₂O-Py-72]⁺ (m/z 585.4→501.3→429.3) from samples where 3β,5α,6β-triol is abundant new peaks appears in both the 17 min (Figure 8E) and 37 min chromatograms. The underlying MS³ spectrum of the new peak at 3.5 min in an NPC plasma sample (Figure 8G, see also Supplemental Figure S5Q), shows a minor fragment-ion at m/z 429.3 and a more prominent ion at m/z 383.3, the former corresponding to [M-H₂O-Py-72]⁺ and the latter [M-H₂O-Py-118], a dehydration product of [M-H₂O-Py-100]⁺ (see Supplemental Figure S4B). Both fragment ions are essentially absent from MS³ spectra of the closely eluting peak (Figure 8D) annotated as 3β,7α,12α-trihydroxycholest-5-en-(25R)26-
oic acid (3β,7α,12α-triHCA, Supplemental Figure S5R) and its isomers 3β,7α,24-trihydroxycholest-5-en- (25R)26-oic (3β,7α,24S-triHCA) and 3β,7α,25-trihydroxycholest-5-en-26-oic acid (3β,7α,25-triHCA). The MRM chromatogram [M-H₂O]^+→[M-Py-H₂O]^+→[M-H₂O-Py-118]^+ provides even clearer definition of 3β,5α,6β,24-tetraHCa (Figure 8F). The chromatographic peak for the [M-H₂O]^+ ion of 3β,5α,6β,24-tetraHCa is sufficiently resolved in the 37 min gradient to allow semi-quantification against the internal standard [²H７]24R/S-HC and by assuming a response factor similar to 3β,5α,6β-triHBA.

Semi-Quantitative Measurements

Other than for 3β,5α,6β-triol itself, for which the [²H7] analogue is available, we can only make approximate or semi-quantitative measurements of 3β,5α,6β-triol containing sterols. An authentic standard of 3β,5α,6β-triHBA is available, thus can be used to give approximate quantification. Other than 3β,5α,6β-triHCA, all of the 3β,5α,6β-triol containing sterols are sufficiently chromatographically resolved from similarly derivatised sterols to allow semi-quantification against added internal standards.
Discussion

When 7-OC or 3β,5α,6β-triol is abundant in a sample, whether formed enzymatically or through radical reactions, the analyst should consider the possibility of the presence of down-stream metabolites. An absence may indicate that the primary metabolites are formed ex vivo, while a presence will indicate formation in vivo or perhaps from the diet or environment. By taking plasma samples from patients with the lysosomal storage disorder NPC as an example we illustrate how metabolites of 7-OC and 3β,5α,6β-triol leading to bile acids 3βH,7O-Δ5-BA and 3β,5α,6β-triHBA, respectively, can be identified.

Considering metabolites with a 7-oxo-5-ene structure, the fragment-ion resulting from an [M-Py-59]+ neutral-loss (Figure 3) and appearing in MS3 ([M]*→[M-Py]*→) spectra is characteristic and valuable for metabolite identification via appropriate MRM ([M]*→[M-Py]*→[M-Py-59]*) RICs (Figure 4B & 4F & 5B). The [M-Py-59]* fragment-ion is likely a radical cation stabilised by delocalisation across a conjugated system from C3 – C7 and two nitrogen atoms (Figure 3). With respect to sterols containing a 3β,5α,6β-triol structure, treatment with cholesterol oxidase and GP derivatisation leads to dehydration through loss of the 5α-hydroxy group and formation of an [M-H2O]+ ion. MS3 ([M-H2O]*→[M-H2O-Py]*→) leads to a characteristic neutral-loss fragment-ion [M-H2O-Py-72]* (Figure 7). Again, the appropriate MRM, [M-H2O]*→[M-H2O-Py]*→[M-H2O-Py-72]* can lead to the identification of 3β,5α,6β-triol-containing metabolites (Figure 6F & 8B & 8E). The [M-H2O-Py-72]* fragment-ion is likely stabilised through delocalisation of positive charge across the conjugated double bonds in the A-ring (Figure 7). Besides giving the [M-H2O]+ ion, both 3β,5α,6β-triol and 3β,5α,6β-triHBA give an [M]+ ion. However, the absence of A/B-ring unsaturation leads to MS3 ([M]*→[M-Py]*→) spectra which are less structurally characteristic making identification of intermediate metabolites difficult (Supplemental Figures S5A & S5H).

Although the primary aim of this work was identification of 7-oxo-5-ene- and 3β,5α,6β-triol containing metabolites, approximate or semi-quantitative measurements can also be made. Accurate quantification, however, awaits further synthesis of authentic standards and their isotope-labelled analogues.
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Footnote

Swansea Innovations Ltd have licensed derivatisation technology described in this paper to Avanti Polar Lipids Inc and Cayman Chemical.
Figure 1. (A) Oxidation of 3β-hydroxysterols with cholesterol oxidase (ChOx) then derivatisation with [3H]GP and MS² fragmentation. (B) Derivatisation of oxosterols with [3H]GP and MS² fragmentation.
Figure 2. Chromatographic separation and MS$^3$ ([M]$^+$→[M-Py]$^+$→) spectra of 7-OC, 7α-HCO, 26H,7O-C and 7α,26-diHCO from an NPC plasma sample derivatised with [H$_2$]GP reagent. (A) Reconstructed ion chromatograms (RICs, m/z 534.4054 ± 5 ppm) demonstrating the separation of 7-OC from 7α-HCO in a plasma sample from an NPC (upper panel) patient and in the NIST reference material (lower panel). MS$^3$ spectra of (B) 7-OC and (C) 7α-HCO from the NPC plasma sample. (D) RIC (550.4003 ± 5 ppm) demonstrating chromatographic separation of 26H,7O-C from 7α,26-diHCO and other isomers in the NPC (upper panel) and NIST (lower panel) plasma samples using the 37 min chromatographic gradient. The chromatograms in (D) were recorded on different days resulting in an offset in retention time of 0.2 – 0.3 min in the earlier peaks. MS$^3$ spectra of (E) 7α,26-diHCO and (F) 26H,7O-C from the NPC sample. Measured concentrations of 7-OC and 26H,7-OC are given in the right-hand corners of the chromatograms (A) and (D), respectively. Note, 7α-HCO, 7α,25-diHCO and 7α,26-diHCO give twin peaks corresponding to syn and anti conformers of the derivative. MS$^3$ spectra of authentic standards of [H$_2$]7-OC, 7-OC, 7α-HCO, 26H,7O-C, 7α,26-diHCO, 7α,25-diHCO and 7α,24S-diHCO can be found in Supplemental Figure S2A-G.
Figure 3. Patterns of MS² ([M]+→[M-Py]+→) neutral-losses which distinguish between, or are common to, [²H₃]GP derivatised 7-oxo-5-ene and 3-oxo-4-ene sterols. Structures of R groups are shown within brackets, in the lower left-hand box. Isotope-labelled [¹³C₂]GP and [¹³C¹⁵N]GP reagents used to determine the composition of the fragment-ions are shown in the lower right-hand box. An asterisk indicates a heavy isotope label.
Figure 4. MRM chromatograms $[M]^+ → [M-Py]^+ → [M-Py-59]^+$ reveal 3βH,7O-CA and 3βH,24-diH,7O-CA in NPC plasma. (A) The authentic standard 3βH,7O-CA (lower panel) co-elutes with 7αH,3O-CA (NPC plasma, upper panel; NIST plasma, central panel) in the 17 min chromatographic gradient. (B) MRM chromatogram ($m/z$ 564.4→485.3→426.3) from an NPC plasma sample (upper chromatogram), the NIST control sample
(central chromatogram) and an authentic standard of 3βH,7O-CA (lower chromatogram). Measured concentrations of 7αH,3O-CA + 3βH,7O-CA and of 3βH,7O-CA alone are given on the right-hand side of the chromatograms (A) and (B), respectively. 25S and 25R epimers of 7αH,3O-CA each gives twin peaks corresponding to syn and anti conformers of the derivative as seen in (A). The twin peaks observed in (B) from NPC and NIST samples probably correspond to 25S and 25R epimers of 3βH,7O-CA. MS³ spectra of (C) 3βH,7O-CA authentic standard, (D) 3βH,7O-CA from a NPC plasma sample, and (E) 7αH,3O-CA(25R) authentic standard. (F) MRM chromatogram (m/z 580.4→501.3→442.3) from an NPC plasma sample (upper panel) and the NIST plasma sample (lower panel) generated with the 17 min gradient. Measured concentrations of 3β,24-diH,7O-CA are given in the right-hand corners of the chromatograms. ND, not detected. (G) MS³ ([M]+→[M-Py]+→) spectrum underlying the major peak at 2.35 min in (F) from NPC plasma, identified as of 3β,24-diH,7O-CA. Structures of fragment ions are shown in Supplemental Figure S1B. MS³ spectra of 7αH,3O-CA(25R) from NIST plasma, 7α,24S-diH,3O-CA and 7α,25-diH,3O-CA authentic standards are shown in Supplemental Figure S2H-J.
Figure 5. The MRM chromatogram [M]+→[M-Py]+→[M-Py-59] + reveals 3βH,7O-Δ5-BA in samples rich in 7-OC. (A) RICs for m/z 522.3326 ± 5 ppm corresponding to 3βH,7O-Δ5-BA and its isomer 7αH,3O-Δ4-BA in NPC (upper panel) and NIST (lower panel) plasma samples. (B) MRM m/z 522.3→443.3→384.3 chromatograms from an NPC (upper panel) and NIST (lower panel) plasma sample. Measured concentrations of 3βH,7O-Δ5-BA are given in the right-hand corners of the chromatograms. The chromatograms were recorded on different days resulting in an offset in retention time of 0.2 min in the early eluting peaks. MS3 ([M]+→[M-Py]+→) spectra of the compounds underlying the chromatographic peaks eluting at (C) 1.96 min (7αH,3O-Δ4-BA) in chromatogram (A) and (D) 2.16 min (3βH,7O-Δ5-BA) in chromatogram (B).
Figure 6. 3β,5α,6β-triHBA and 3β,5α,6β,26-tetrol give [M-H$_2$O-Py-72]$^+$ and [M-H$_2$O-Py-100]$^+$ neutral-loss fragment-ions in their MS$^2$ ([M-H$_2$O]$^+$$\rightarrow$[M-H$_2$O-Py]$^+$) spectra. (A) RIC, m/z 539.4368 ± 5 ppm, demonstrating chromatographic separation of 3β,5α,6β-triHBA ([M$^+$] ions) and 5,6-EC ([M$^+$] ions) in NPC (upper panel) and NIST (lower panel) plasma samples. Measured
concentrations of 3β,5α,6β-triol are given in the right-hand corners of the chromatograms. Monohydroxycholesterols give syn and anti conformers of the GP derivative, resulting in twin peaks. (B) MS³ ([M-H₂O]⁺→[M-H₂O-Py]⁺→) spectrum of 3β,5α,6β-triol from an NPC plasma sample. (C) RIC of m/z 527.3640 ± 5 ppm demonstrating chromatographic separation of 3β,5α,6β-triHBA [M-H₂O]⁺ from 3β,7β-diH-Δ⁵-BA ([M]⁺ ions) and 3β,7α-diH-Δ⁵-BA ([M]⁺ ions) in NPC (upper panel) and NIST (lower panel) plasma samples. Measured concentrations of 3β,5α,6β-triHBA are given in the right-hand corners of the chromatograms. Both diH-Δ⁵-BA isomers give twin chromatographic peaks. The chromatograms were recorded on different days resulting in a retention time shift of ~0.2 min. (D) MS³ ([M-H₂O]⁺→[M-H₂O-Py]⁺→) spectrum of 3β,5α,6β-triHBA in an NPC plasma sample. (E) RIC for m/z 555.4317 corresponding to the [M-H₂O]⁺ ion of cholestanetetrols and the [M]⁺ ion of dihydroxycholesterols from NPC (upper panel) and NIST (lower panel) plasma samples. Measured concentrations of 3β,5α,6β,26-tetrol are given in the right-hand corners of the chromatograms. (F) RIC for the MRM transition m/z 555.4→471.4→399.3 corresponding to [M-H₂O]⁺→[M-H₂O-Py]⁺→[M-H₂O-Py-72]⁺ for cholestanetetrols from the NPC (upper panel) and NIST (lower panel) plasma samples. (G) MS³ spectrum of the peak eluting at 5.2 min in the NPC plasma sample. See Supplemental Figure S4A & S4B for assignment of fragment-ions. MS³ spectra of authentic standards of [²H₇]3β,5α,6β-triol, 3β,5α,6β-triol and 5α,6-EC are shown in Supplemental Figure S5C – S5E.
Figure 7. Patterns of MS³ neutral-losses which distinguish between, or are common to, 3β,5α,6β-triol-containing and 3β,7-dihydroxy-5-ene sterols. The pattern of neutral-losses shown in the red box distinguish between [M-H₂O]⁺ ions of 3β,5α,6β-triols from [M]⁺ ions of 3β,7-dihydroxy-5-ene sterols of identical mass. Neutral-losses/fragment-ion shown in the blue box are common to both structures. Structures of R groups are shown within brackets, in the lower left-hand box.
Figure 8. Identification of 3β,5α,6β-triHCA and 3β,5α,6β,24-tetraHCA in plasma samples rich in 3β,5α,6β-triol. (A) RIC of m/z 569.4110 ± 5 ppm corresponding to [M-H2O]+ and [M]+ ions of 3β,5α,6β-triHCA and dihydroxycholestenolic acids, respectively, from plasma from a patient with NPC (upper panel) and the NIST
control plasma (lower panel). 3β,7α-diHCA and 3β,7β-diHCA appear as 25S and 25R epimers and both give twin peaks due to syn and anti conformers of the GP derivative. The NPC plasma was analysed on a different day to the NIST plasma samples resulting in a 0.1 - 0.2 min offset in the earlier eluting chromatographic peaks. (B) RIC for the MRM transitions 569.4→485.3→413.3 in plasma from a patient with NPC (upper panel) and the NIST control plasma (lower panel). Measured concentrations of 3β,5α,6β-triHCA are given on the right-hand side of the chromatograms. (C) MS^3 ([M-H_2O]^+→[M-Py-H_2O]^+) spectrum from the compound underlying the chromatographic peak at 4.69 min in the NPC chromatogram in (B). See Supplemental Figure S4B for a description of fragment ions. (D) RIC of m/z 585.4059 ± 5 ppm corresponding to [M-H_2O]^+ of 3β,5α,6β,24-tetraHCA and [M]^+ of trihydroxycholestenoic acids in NPC (upper panel) and NIST control plasma (lower panel). MRM chromatograms (E) m/z 585.4→501.3→429.3 and (F) m/z 585.4→501.3→383.3 revealing 3β,5α,6β,24-tetraHCA in NPC (upper panels), but not the NIST control plasma (lower panels). Measured concentrations of 3β,5α,6β,24-tetraHCA are given on the right-hand side of the chromatograms. The NPC plasma was analysed on a different day to the NIST plasma samples resulting in a 0.1 - 0.2 min offset in the earlier eluting chromatographic peaks. (G) MS^3 ([M-H_2O]^+→[M-H_2O-Py]^+) spectrum of 3β,5α,6β,24-tetraHCA in NPC plasma.