

Published in final edited form as:

Clin Biochem. 2014 June ; 47(9): 860–863. doi:10.1016/j.clinbiochem.2014.04.017.

A useful multi-analyte blood test for cerebrotendinous xanthomatosis

Andrea E DeBarber^{a,*}, Jenny Luo^a, Roberto Giugliani^{b,c}, Carolina FM Souza^c, John (Pei-Wen) Chiang^d, Louise S Merkens^e, Anuradha S Pappu^e, and Robert D Steiner^{e,f,g}

^a Department of Physiology & Pharmacology, Oregon Health & Science University (OHSU), Portland, Oregon 97239, United States

^b Medical Genetics Service, Hospital de Clinicas de Porto Alegre, Brazil

^c Department of Genetics, Federal University of Rio Grande do Sul, Porto Alegre, Brazil

^d Casey Molecular Diagnostic Laboratory, OHSU

^e Department of Pediatrics, OHSU

^f Department of Molecular & Medical Genetics, Child Development & Rehabilitation Center and Doernbecher Children's Hospital, OHSU

^g Marshfield Research Clinic Foundation, Marshfield, Wisconsin 54449, United States

Abstract

Objectives—Cerebrotendinous xanthomatosis (CTX) is a rare genetic disorder of bile acid (BA) synthesis that can cause progressive neurological damage and premature death. Blood (normally serum or plasma) testing for CTX is performed by a small number of specialized laboratories, routinely by gas chromatography-mass spectrometry (GC-MS) measurement of elevated 5 α -cholestanol. We report here on a more sensitive biochemical approach to test for CTX particularly useful for confirmation of CTX in the case of a challenging diagnostic sample with 5 α -cholestanol that, although elevated, was below the cut-off used for diagnosis of CTX (10 μ g/ml or 1.0 mg/dL).

Design and Methods—We have previously described liquid chromatography-electrospray ionization-tandem mass spectrometry (LC-ESI-MS/MS) methodology utilizing keto derivatization to enable the sensitive quantification of plasma ketosterol BA precursors that accumulate in CTX. We have expanded this methodology to perform isotope dilution LC-ESI-MS/MS quantification of a *panel* of plasma ketosterol BA precursors, with internal standards readily generated using isotopically-enriched derivatization reagent.

Results—Quantification of plasma ketosterol BA precursors (7 α -hydroxy-4-cholesten-3-one, 7 α , 12 α -dihydroxy-4-cholesten-3-one and 7 α , 12 α -dihydroxy-5 β -cholestan-3-one) in a single LC-

© 2014 The Canadian Society of Clinical Chemists. Published by Elsevier Inc. All rights reserved.

* Corresponding author (debarber@ohsu.edu; phone: 503-494-3154; fax: 503-494-4352).

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

ESI/MS/MS test provided better discrimination between a CTX-positive and negative samples analyzed (n=20) than measurement of 5 α -cholestanol alone.

Conclusions—Quantification of plasma ketosterol BA precursors provides a more sensitive biochemical approach to discriminate between CTX negative and positive samples. A multiplexed LC-ESI-MS/MS test quantifying a panel of plasma ketosterols, with simple sample preparation, rapid analysis time and readily available internal standards, can be performed by most clinical laboratories. Wider availability of testing will benefit those affected with CTX.

Keywords

Cerebrotendinous xanthomatosis; CYP27A1; bile acids; ketosterols; cholestanol

1. Introduction

Cerebrotendinous xanthomatosis (CTX; OMIM#213700) is an autosomal recessive neurodegenerative disorder associated with deficient sterol 27-hydroxylase (CYP27A1), a mitochondrial enzyme important in conversion of cholesterol to bile acids (BA). Childhood-onset symptoms can include diarrhea and juvenile cataracts. Adolescent to adult-onset symptoms can include tendon and cerebral xanthomas associated with neurological symptoms. CTX is difficult to diagnose, often there are many years between the age of first symptom onset and the age at diagnosis, which often occurs only after significant neurological involvement. As the disorder progresses, affected individuals can become incapacitated with motor dysfunction with premature death occurring due to advancing neurological deterioration. Although only around three hundred cases of CTX have been described worldwide [1], relatively large series of patients has been described by physicians with experience in recognizing the disorder.

An effective oral therapy for CTX is available in the form of chenodeoxycholic acid (CDCA), the main BA deficient in CTX. Treatment with CDCA has been shown to normalize the biochemical phenotype and halt progression of disease [2,3]. In many cases treatment of patients with advanced neurological disease does not reverse the impairment [3], therefore it is essential to diagnose and treat CTX as early as possible.

Biochemical tests for CTX include screening blood samples using gas chromatography-mass spectrometry (GC-MS) measurement of elevated 5 α -cholestanol [4,5]. The upper normal range for plasma 5 α -cholestanol has been reported to range from 7-11 μ g/ml [4,5]. Diagnostic confirmation is routinely performed using fast atom bombardment (FAB)-MS measurement of more specific markers for CTX, bile alcohol glucuronides in urine [6]. We have utilized keto- moiety derivatization to enable sensitive liquid chromatography-electrospray ionization-tandem MS (LC-ESI-MS/MS) measurement of ketosterol BA precursors that accumulate in CTX [7-9]. Free (non-hydrolyzed) plasma 7 α -hydroxy-4-cholesten-3-one (7 α C4) possessed improved utility over 5 α -cholestanol as a marker for CTX [7]. We describe here quantification of a *panel* of ketosterols in a plasma sample from a CTX affected individual with a plasma 5 α -cholestanol concentration above the concentration range we determined for normal samples (n=20) but below our cut-off for diagnosis of CTX (10 μ g/ml).

2. Materials and methods

2.1 Human subject research considerations

Blood was obtained from participants enrolled in studies at OHSU under Institutional Review Board (IRB) approved protocols. Informed consent was obtained from all study participants. De-identified plasma samples submitted to the OHSU diagnostic laboratory for CTX testing using GC-MS measurement of elevated plasma 5 α -cholestanol were used with IRB approval. For CTX positive samples diagnostic confirmation was by molecular genetic testing.

2.2 Chemicals and reagents

7 α C₄, 7 α , 12 α -dihydroxy-4-cholesten-3-one (7 α ,12 α C₄) and 7 α ,12 α -dihydroxy-5 β -cholestan-3-one (7 α ,12 α C₅ β) were from Toronto Research Chemicals (Toronto, Ontario). 5 α -Cholestanol was from Steraloids (Newport, RI) and epicoprostanol from Sigma-Aldrich (St. Louis, MO). BSTFA reagent was from Thermo Scientific (Bellefonte, PA). Human plasma and double charcoal stripped (DCS) plasma were from Golden West Bio (Temecula, CA). Methanol and water (GC-MS grade) were from Burdick and Jackson (Muskegon, MI). Formic acid (90%) was J.T.Baker brand, and glacial acetic acid (99.99%) was from Aldrich. Quaternary ammonium (QAO) reagent (O-(3-trimethylammoniumpropyl) hydroxylamine) bromide is commercially available as Amplifex™ Keto reagent from <http://www.sciex.com>. The QAO-d₃ reagent was provided by AB SCIEX.

2.3 Preparation of calibrators and samples for GC-MS measurement of 5 α -cholestanol

GC-MS measurement of elevated plasma 5 α -cholestanol has been described [7,9]. In brief, internal standard (epicoprostanol) was added to plasma samples or calibrants generated using 5 α -cholestanol. Sterols were saponified by the addition of ethanol/KOH and the aqueous phase was extracted with hexane. Dried sterols were derivatized with BSTFA and the trimethylsilyl ether derivative of 5 α -cholestanol was measured using GC (splitless injection) performed with a ZB1701 column (30m, 0.25mmID, 0.25 μ m film thickness, Phenomenex, Torrance, CA) coupled to a mass spectrometer (Agilent GC 6890N and MS 5975; Santa Clara, CA). Mass spectra were collected in selected ion mode (with m/z = 355 and 370 ions monitored for epicoprostanol and m/z = 306 and 305 ions for 5 α -cholestanol).

2.4 Preparation of calibrators and samples for LC-MS/MS measurement of ketosterols

We have previously described the approach used for LC-ESI-MS/MS measurement of elevated plasma ketosterols [9]. In brief, method calibrators were generated using dilutions of authentic standard in methanol spiked into DCS plasma. QAO reagent solution (210 μ g in methanol plus 5% acetic acid, v/v) was added to calibrators or plasma samples (4 μ l). After 2 hours at RT QAO derivatization was complete. Previously prepared QAO-d₃ tagged ketosterol internal standards (300 pg) in 10 μ l methanol were added [9].

2.5 LC-ESI-MS/MS method

LC-ESI-MS/MS analyses were performed using a QTRAP® 5500 triple-quadrupole hybrid mass spectrometer with linear ion trap functionality (AB SCIEX, Framingham, MA),

equipped with a TurboIonSpray® ESI source. The ionization interface was operated in the positive mode and multiple reaction monitoring (MRM) transitions monitored for quantification of ketosterols were as follows: QAO 7 α C4 m/z 515.7→152.2, for QAO 7 α 12 α C4 m/z 531.7→152.2, for QAO 7 α 12 α C5 β m/z 533.7→145.0, for QAO-d₃ 7 α C4 m/z 518.7→152.3, for QAO-d₃ 7 α 12 α C4 m/z 534.7→152.1 and for QAO-d₃ 7 α 12 α C5 β m/z 536.7→145.0. The QTRAP® 5500 was coupled to a Shimadzu UPLC system (Columbia, MD) composed of a SIL- 20ACXR auto-sampler and two LC-20ADXR LC pumps. QAO derivatives were resolved using a 50×2.1(i.d.) mm, 5.0 μ m Luna C₈-HPLC column with guard (Phenomenex; Torrance, CA). The gradient mobile phase was delivered at a flow rate of 0.8 ml/min and the water:acetonitrile:0.1% formic acid mobile phase [9]. The column temperature was kept at 35°C using a Shimadzu CTO-20AC column oven. The sample injection volume was 10 μ l.

2.6 Data analysis and method performance

Calibration curves were generated by performing a least-squares linear regression for peak area ratios (QAO-d₀ ketosterol analyte/QAO-d₃ ketosterol internal standard) plotted against specified calibrant concentration in plasma (ng/ml) [9]. The lower limit of quantification (LLOQ) was determined as the lowest spiked concentration in matrix for which the signal-to-noise (S/N) ratio was 5 and the within- and between-day reproducibility of the peak area was 20% relative standard deviation (RSD). Potential method interference was evaluated by examination of the peak shape, peak shoulder, and peak area ratio of two MRM transitions (quantifier and qualifier) acquired for each analyte and by analysis of possible method interferents. The matrix effect was determined by comparing the QAO ketosterol peak area for matrix-based against non-matrix calibrators.

3. Results

As we have previously described, DCS plasma calibration curves for QAO tagged 7 α C4 and 7 α 12 α C4, with QAO-d₃ tagged 7 α C4 and 7 α 12 α C4 internal standards respectively, demonstrated acceptable linearity (correlation coefficients across the range 20-250 ng/ml possessed r^2 values >0.990) [9]. This was also the case for a novel ketosterol marker for CTX we quantified with isotope dilution LC-ESI-MS/MS, 7 α 12 α C5 β . DCS plasma calibration curves for QAO tagged 7 α 12 α C5 β with QAO-d₃ tagged 7 α 12 α C5 β internal standard demonstrated acceptable linearity (correlation coefficients across the range 20-250 ng/ml possessed r^2 values >0.990). Satisfactory between and within-run accuracy and precision data was also obtained for calculated concentrations of 7 α 12 α C5 β (<20% RSD at the LLOQ and <15% RSD between 50 and 250 ng/ml). The LLOQ for all ketosterols in plasma (including 7 α 12 α C5 β) was 20 ng/ml.

The sensitivity of ketosterols as plasma markers for CTX compared to 5 α -cholestanol was highlighted by analysis of a plasma sample from a possible CTX-affected individual with a clinical history consistent with the disorder (see *Figure 1*). GC-MS analysis revealed a plasma 5 α -cholestanol concentration of 8.4 g/ml, close to reported upper normal concentrations of 7- 11 g/ml [4,5]. The plasma ketosterol concentrations determined with LC-ESI-MS/MS were 795, 1548 and 1004 ng/ml for 7 α C4, 7 α 12 α C4 and 7 α 12 α C5 β

respectively (with upper normal concentrations of 22 and 0.8, and 29 ng/ml determined; for reference ranges see *Table 1*). Molecular genetic testing confirmed the individual possessed a homozygous R474Q *CYP27A1* gene mutation, previously reported to be a causative mutation for CTX [10].

4. Discussion

5 α -Cholestanol can be elevated in a number of liver diseases and concerns have been raised regarding the specificity of this disease marker for CTX [11]. Although not routinely used for diagnosis, other markers elevated in CTX include cholesterol precursors such as 7-dehydrocholesterol and 8-dehydrocholesterol [12]. Ketosterol BA precursors and bile alcohol glucuronides also accumulate in CTX [7-9,13-15]. We have previously described an approach using keto derivatization to incorporate a permanent charge and improve sensitivity for LC-ESI- MS/MS detection of ketosterol BA precursors [7-9]. We report here on expansion of this methodology to allow isotope dilution quantification of a *panel* of plasma ketosterol BA precursor markers to test for CTX. The sensitivity of the panel to detect disease was highlighted by analysis of a CTX-positive sample that possessed a 5 α -cholestanol concentration below the cut-off used for diagnosis of CTX (10 μ g/ml). Although GC-MS analysis of the sample indicated elevated 7- and 8-dehydrocholesterol suggestive of CTX [12], these sterols are not ideal disease markers as their quantification can be problematic. They are relatively unstable, and stable-isotope labeled internal standard analogues are not readily available. Isotope dilution LC- ESI-MS/MS quantification of plasma ketosterols allowed for ready discrimination between the sample we describe here and CTX negative samples (n=20), such that it serves as an improved blood test for CTX. The availability of LC-ESI-MS/MS methodology utilizing QAO derivatization with simple and rapid sample preparation amenable to automation, analysis times of 4-6 min using conventional LC instrumentation and buffers, and readily available stable-isotope labeled internal standard analogues, allows performance by most clinical laboratories. Wider availability of testing will benefit those affected with this disorder.

Acknowledgments

The authors would like to thank the Bioanalytical Shared Resource at OHSU for providing technical assistance and access to analytical instrumentation. AED has been supported as a KL2 awardee by the Oregon Clinical and Translational Research Institute (OCTRI), grant number (KL2TR000152) from the National Center for Advancing Translational Sciences (NCATS) at the National Institutes of Health (NIH) and also as a training grant awardee by the Sterol and Isoprenoid Diseases (STAIR) consortium. STAIR is part of the NIH Rare Diseases Clinical Research Network (RDCRN). Funding and/or programmatic support for this project has been provided by a grant (1U54HD061939) from the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) and the NIH Office of Rare Diseases Research (ORDR).

Abbreviations

CTX	cerebrotendinous xanthomatosis
BA	bile acid
GC-MS	gas chromatography- mass spectrometry
ESI- MS/MS	liquid chromatography-electrospray ionization-tandem MS

CDCA	chenodeoxycholic acid
FAB	fast atom bombardment
7αC4	7 α -hydroxy-4-cholesten-3-one
IRB	Institutional Review Board
7α12αC5β	α ,12 α -dihydroxy-4-cholesten-3-one (7 α 12 α C4) and 7 α ,12 α -dihydroxy-5 α -cholestan-3-one
DCS	double charcoal stripped
QAO	quaternary amonoxy
MRM	multiple reaction monitoring
LLOQ	limit of quantification
S/N	signal-to-noise
RSD	relative standard deviation

Reference List

- Gallus GN, Dotti MT, Mignarri A, Rufa A, Da PP, Cardaioli E, Federico A. Four novel CYP27A1 mutations in seven Italian patients with CTX. *Eur J Neurol*. 2010; 17:1259–1262. [PubMed: 20402754]
- Berginer VM, Salen G, Shefer S. Long-term treatment of cerebrotendinous xanthomatosis with chenodeoxycholic acid. *N Engl J Med*. 1984; 311:1649–1652. [PubMed: 6504105]
- Mondelli M, Sicurelli F, Scarpini C, Dotti MT, Federico A. Cerebrotendinous xanthomatosis: 11-year treatment with chenodeoxycholic acid in five patients. An electrophysiological study. *J Neurol Sci*. 2001; 190:29–33. [PubMed: 11574103]
- Leitersdorf E, Safadi R, Meiner V, Reshef A, Bjorkhem I, Friedlander Y, Morkos S, Berginer VM. Cerebrotendinous xanthomatosis in the Israeli Druze: molecular genetics and phenotypic characteristics. *Am J Hum Genet*. 1994; 55:907–915. [PubMed: 7977352]
- Salen G. Cholesterol deposition in cerebrotendinous xanthomatosis. A possible mechanism. *Ann Intern Med*. 1971; 75:843–851. [PubMed: 5134895]
- Egestad B, Pettersson P, Skrede S, Sjøvall J. Fast atom bombardment mass spectrometry in the diagnosis of cerebrotendinous xanthomatosis. *Scand J Clin Lab Invest*. 1985; 45:443–446. [PubMed: 4035280]
- DeBarber AE, Connor WE, Pappu AS, Merckens LS, Steiner RD. ESI-MS/MS quantification of 7 α -hydroxy-4-cholesten-3-one facilitates rapid, convenient diagnostic testing for cerebrotendinous xanthomatosis. *Clin Chim Acta*. 2010; 411:43–48. [PubMed: 19808031]
- DeBarber AE, Sandlers Y, Pappu AS, Merckens LS, Duell PB, Lear SR, Erickson SK, Steiner RD. Profiling sterols in cerebrotendinous xanthomatosis: Utility of Girard derivatization and high resolution exact mass LC-ESI-MS(n) analysis. *J Chromatogr B Analyt Technol Biomed Life Sci*. 2011; 879:1384–1392.
- DeBarber AE, Luo J, Star-Weinstock M, Purkayastha S, Geraghty MT, Chiang J, Merckens LS, Pappu AS, Steiner RD. *J Lipid Res*. 2013 in press.
- Kuriyama M, Fujiyama J, Yoshidome H, Takenaga S, Matsumuro K, Kasama T, Fukuda K, Kuramoto T, Hoshita T, Seyama Y. Cerebrotendinous xanthomatosis: clinical and biochemical evaluation of eight patients and review of the literature. *J Neurol Sci*. 1991; 102:225–232. [PubMed: 2072121]
- Koopman BJ, van der Molen JC, Wolthers BG, de Jager AE, Waterreus RJ, Gips CH. Capillary gas chromatographic determination of cholesterol/cholesterol ratio in biological fluids. Its potential

- usefulness for the follow-up of some liver diseases and its lack of specificity in diagnosing CTX (cerebrotendinous xanthomatosis). *Clin Chim Acta*. 1984; 137:305–315. [PubMed: 6421514]
12. de Sain-van der Velden MG, Verrips A, Prinsen BH, de BM, Berger R, Visser G. Elevated cholesterol precursors other than cholestanol can also be a hallmark for CTX. *J Inherit Metab Dis*. 2008; 2:S387–393. [PubMed: 18949577]
 13. Bjorkhem I, Oftebro H, Skrede S, Pedersen JI. Assay of intermediates in bile acid biosynthesis using isotope dilution--mass spectrometry: hepatic levels in the normal state and in cerebrotendinous xanthomatosis. *J Lipid Res*. 1981; 22:191–200. [PubMed: 7017048]
 14. Clayton PT, Verrips A, Sistermans E, Mann A, Mieli-Vergani G, Wevers R. Mutations in the sterol 27-hydroxylase gene (CYP27A) cause hepatitis of infancy as well as cerebrotendinous xanthomatosis. *J Inherit Metab Dis*. 2002; 25:501–513. [PubMed: 12555943]
 15. Honda A, Salen G, Matsuzaki Y, Batta AK, Xu G, Leitersdorf E, Tint GS, Erickson SK, Tanaka N, Shefer S. Differences in hepatic levels of intermediates in bile acid biosynthesis between Cyp27(-/-) mice and CTX. *J Lipid Res*. 2001; 42:291–300. [PubMed: 11181760]

Highlights

1. Elevated cholestanol is used to test for cerebrotendinous xanthomatosis (CTX)
2. We describe a useful test for a case with cholestanol below the diagnostic cut-off
3. Plasma ketosterols allowed for discrimination between the case sample and controls
4. Rapid isotope-dilution LC-ESI-MS/MS testing can be performed by most laboratories
5. Wider availability of plasma testing will benefit those affected with CTX

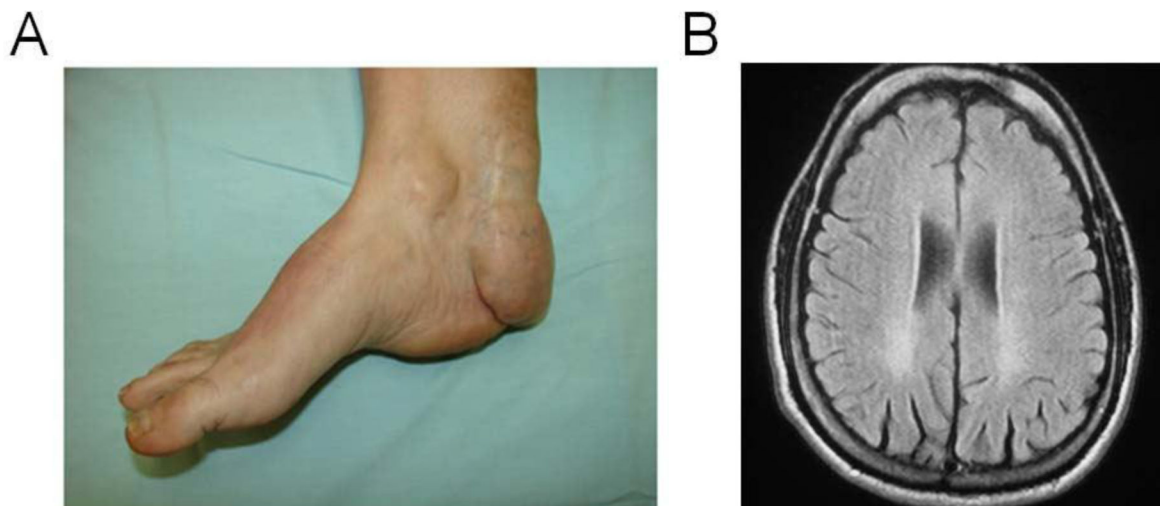


Figure 1. the plasma sample analyzed was from the daughter of a non-consanguineous couple, with two possibly affected siblings that died at 24 and 55 years old with neurological symptoms suggestive of CTX. The daughter had developed cataracts at age 22 years old that were surgically corrected and Achilles tendon xanthomas from age 13 (*left panel*) that could not be alleviated with surgery, resulting in difficulty walking and intense pain in the feet and knees. There was spastic gait present (pyramidal tract syndrome) with no ataxic manifestations, extrapyramidal signs or dementia, although an MRI of the brain showed bilateral occipital periventricular white matter signal hyperintensity on T2 and Flair (*right panel*).

Table 1

Plasma ketosterol concentrations

	7 α -hydroxy-4-cholesten-3-one ^{a,b} (ng/ml)	7 α 12 α -dihydroxy-4-cholesten-3-one ^{a,b} (ng/ml)	7 α 12 α -dihydroxy-5 β -cholestan-3-one ^a (ng/ml)	5 α -cholestano1 ^{b,c} (μ g/ml)
CTX-case described plasma	795	1548	1004	8.4
<i>Untreated CTX-affected adult plasma (n=10)</i>	1174 \pm 711 ^d [204-1828]	1545 \pm 1144 ^d [57-2420]	498 \pm 310 ^d [100-1004]	31 \pm 19 [8.4-66]
<i>Unaffected adult plasma (n=20)</i>	6.4 \pm 5.3 ^e [1.6-22]	0.4 \pm 0.3 ^e [0.1-0.8]	10 \pm 5.5 ^e [0-29]	1.3 [0.8-1.8] ^f

The mean concentration \pm S.D. and [range of results] are given

^a non-hydrolyzed free sterol.

^b this data has been reported previously [9].

^c hydrolyzed total sterol; quantification by GC-MS.

^d CTX plasma samples with concentrations $>$ 250 ng/ml were diluted and re-analyzed.

^e Calculated outside the quantifiable range.

^f Upper 5 α -cholestano1 normal cut-off reported as between 7-11 μ g/ml [4,5].