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Extraction of 25-hydroxy Vitamin D from Serum Using ISOLUTE[®] PLD+ Prior to LC-MS/MS Analysis

This application note describes the extraction of 25-hydroxy vitamin D from serum, prior to LC-MS/MS analysis.

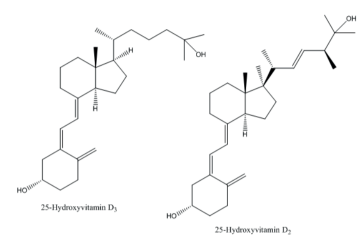


Figure 1. Structures of 25-hydroxy Vitamin D.

Introduction

ISOLUTE® PLD+ Protein and Phospholipid Removal plates offer a substantial improvement in extract cleanliness compared to traditional protein precipitation techniques for bioanalytical sample preparation.

This application note describes a simple, effective ISOLUTE[®] PLD+ protocol for the extraction of 25-hydroxy vitamin D from serum, demonstrating high, reproducible analyte recoveries with low protein and phospholipid content in the extracts.

Analytes

25-hydroxy vitamin D2, 25-hydroxy vitamin D3 and d6-25hydroxy vitamin D3 as the internal standard.

Sample Preparation Procedure

Format: ISOLUTE[®] PLD+ Protein and Phospholipid Removal plate, part number 918-0050-P01

Sample Pre-treatment

Add 10 μL of ISTD (equivalent to 30 ng/mL) to the serum sample. Mix. Allow to stand for ~1 hour for binding to occur.

Solvent Application

Apply 400 μL of Acetonitrile (MeCN) to each well of the ISOLUTE° PLD+ plate.

Sample Application

Add 100 μL of serum with ISTD and mix thoroughly via repeat aspirate/dispense steps.

Analyte Elution

Apply vacuum -0.2 bar or 3 psi positive pressure for approximately 5 minutes. For highly particulate laden samples increased pressure or vacuum conditions may be required.

Post Extraction

Dry the extract in a stream of air or nitrogen using a Biotage[®] SPE Dry (40 °C at 40 L/min) or TurboVap[®] (40 °C at 1.0 bar).

Reconstitution

Apply 100 μL of 30/70 2 mM Ammonium Formate, 0.1% formic acid aq/MeOH.

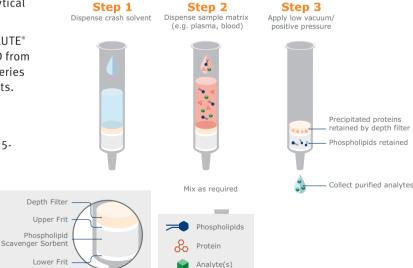


Figure 2. Typical ISOLUTE® PLD+ procedure



UPLC Conditions

Instrument

Waters Acquity UPLC (Waters Assoc., Milford, MA, USA)

Column

ACE EXCEL 2 C18-PFP, 100 mm x 2.1 mm id 2 µm, (ACT, UK)

Mobile Phase

A: 2 mM ammonium formate/0.1% formic acid (aq) B: 2 mM ammonium formate/0.1% formic acid/MeOH

Flow Rate

o.4 mL/min

Table 1. UPLC Gradient Conditions.

Time	%A	%B	Curve	
0	25	75	1	
3	0	100	6	
4	25	75	11	

Curve 11: Conditions in line initiated immediately once time passed. i.e. 25:75 resumed at 4 minutes.

Curve 6: Linear Gradient

Injection Volume

15 µL (partial loop with overfill)

Sample Temperature

20 °C

Column Temperature

40 °C

Note: alternative UPLC conditions may be suitable. Check for good chromatographic separation of isobaric interferences to ensure accurate analyte quantitation

Mass Spectrometry Conditions

Instrument

Quattro Premier XE triple quadrupole mass spectrometer (Waters Assoc., Manchester, UK) equipped with an electrospray interface for mass analysis

Desolvation Temperature

450 °C

Ion Source Temperature

150 °C

Collision Cell Pressure

3.76 e⁻³ mbar

Positive ions acquired in the multiple reaction monitoring (MRM) mode:

 $\ensuremath{\textbf{Table 2.}}\xspace$ Qualifier on details shown in parenthesis).

Analyte	MRM Transition	Cone V	Collision Energy eV
25-OH D ₂	395.5 > 269.5 (395.5 > 119.2)	30	18 26
25-OH D ₃	383.5 > 257.5 (383.5 > 107.2)	30	17 25
d6-25-OH D ₃	389.6 > 263.5	30	16

Results

Chromatography

Good chromatographic separation of 25-hydroxy vitamin D2 and D3 was achieved in less than 3 minutes, as shown in **Figure 3**.

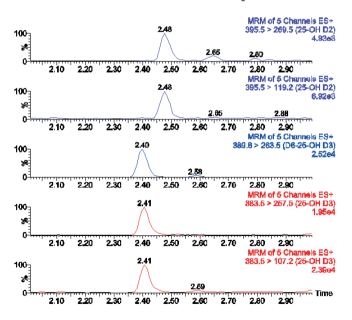


Figure 3. Chromatographic separation of 25-hydroxy vitamin D2 and D3 from Chromsystems calibrated serum at 14.8 and 19.6 ng/mL respectively. ISTD at 30 ng/mL.



Recovery

Serum and stripped serum was spiked at various concentrations from 2–100 ng/mL. High reproducible recoveries > 70% with corresponding RSDs < 10% were demonstrated. Typical recovery data is shown in **Figure 4**.

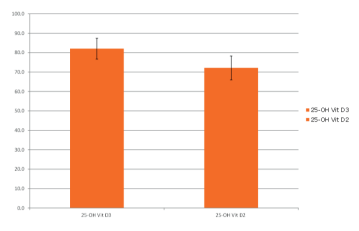


Figure 4. Recovery profile for 25-hydroxy vitamin D extracted at 50 ng/mL.

PBS/BSA Calibration Curves

Calibration curves were generated using PBS/BSA spiked at concentrations from 1–100 ng/mL. Good coefficients of determination were obtained for 25-hydroxy vitamin D2 and D3 ($r^2 > 0.99$).

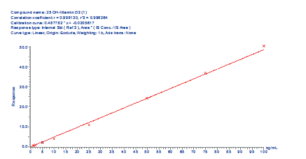
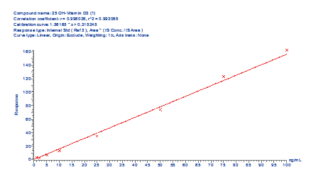
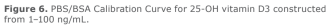


Figure 5. PBS/BSA Calibration Curve for 25-OH vitamin D2 constructed from 1–100 $\rm ng/mL.$





Chromsystems Calibration Curves

Curves were also generated using calibrated serum standards (obtained from Chromsystems) spiked at concentrations from o-69 ng/mL. Good coefficients of determination were obtained for 25-hydroxy vitamin D2 and D3 ($r^2 > 0.99$).

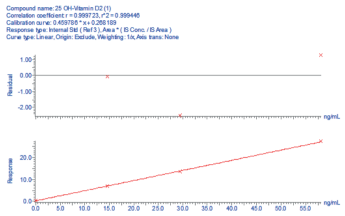


Figure 7. Chromsystems Calibrated Serum Curve for 25-OH vitamin D2 constructed from 4-69 ng/mL.

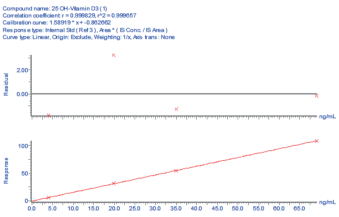


Figure 8. Chromsystems Calibrated Serum Curve for 25-OH vitamin D3 constructed from 0-58 ng/mL.



DEQAS External Quality Assessment Scheme

Final method testing was performed for 5 DEQAS serum samples extracted alongside the Chromsystems calibrators using the optimized method. The DEQAS criteria for acceptable performance is that at least 80% of results should fall within + or - 25% of the All Laboratory Trimmed Mean. Method performance is shown in **Table 3**. Units are quoted as ng/mL. All values fall within the accepted criteria.

 Table 3. DEQAS 25-OH vitamin D results obtained using optimum method.

DEQAS Sample I.D.	DEQAS LC/MS Mean	ISOLUTE° PLD+
451	12.9	14.5
452	46.7	49.1
453	26.6	28.9
454	21.4	25.3
455	22.2	23.7

Additional Notes

Buffer Preparation

- 1. 2 mM ammonium formate/0.1% formic acid (aq): Weigh 0.12612 g and dissolve in H_2O . Add 1 mL of formic acid and make up to 1 L in H_2O .
- 2 mM ammonium formate/0.1% formic acid/MeOH: Weigh 0.12612 g and dissolve in MeOH. Add 1 mL of formic acid and make up to 1 L in MeOH.

Processing Conditions

Positive Pressure: Process at approximately 3 psi.

Vacuum Processing: Process at approximately -0.2 bar.

Ordering Information

Part Number	Description	Quantity
918-0050-P01	ISOLUTE [®] PLD+ Fixed Well Plate	1
121-9600	Biotage® VacMater™-96 Sample Processing Manifold	1
PPM-96	Biotage [®] PRESSURE+ 96 Positive Pressure Manifold	1
SD-9600-DHS-EU	Biotage [®] SPE Dry Sample Concentrator System 220/240 V	1
SD-9600-DHS-NA	Biotage® SPE Dry Sample Concentrator System 100/120 V	1
C103263	TurboVap®96, Evaporator 100/120V	1
C103264	TurboVap® 96, Evaporator 220/240V	1



Appendix Biotage® Extrahera™ Settings

The method described in this application note was automated on the Biotage[®] Extrahera, using ISOLUTE PLD+ Protein and Phospholipid Removal plates. Method performance was comparable.

This appendix contains the software settings required to configure Extrahera to run this method.

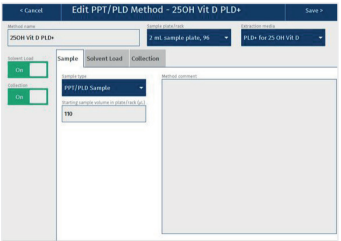
An importable electronic copy of this method for Extrahera can be downloaded from www.biotage.com

Biotage[®] Extrahera[™] Data

Analyte	25-OH Vitamin D2	25-OH Vitamin D3
Recovery (n=8) at 50 ng/mL	92.0	81.1
%RSD	4.0	7.4
Linearity (r ²)	0.999	0.999
LLOQ	<4 ng/mL	<4 ng/mL

Method name:	AN842 Biotage® Extrahera™ 25OH Vit D PLD+
Sample plate/rack:	2 mL sample plate, 96
Extraction Media:	ISOLUTE [®] PLD+ plate

Screenshot





Settings

"Sample" Tab Sample Type: Starting Sample Volume (µL): Method Comment:

PPT/PLD Sample 110

Transfer 100 μL of plasma to a PLD plate containing 400 μL acetonitrile. A 100 μL volume is mixed 5 times before a 0.4 bar pressure is applied for 5 minutes

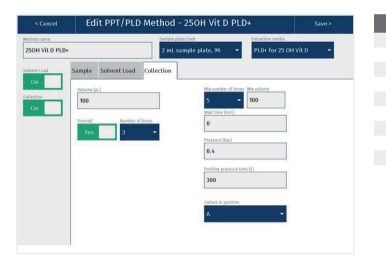


Screenshot

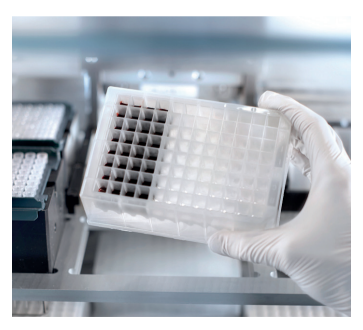
< Cancel Edit	PPT/PLD Method - 250H	Vit D PLD+	Save >
thod name SOH Vit D PLD+	Sample plate/rack 2 mL sample plat	Extraction media	vit D 👻
Ivent Load On Sample So Solvent Acetonitii Volume (L) 400	e		Dispose tips?

Settings

	Solvent	Load	Activated	
	Dispose	tips	No	
	Solvent	:		
1	Acetonit	rile		
		1		
Volu	me (µL)	400		



Collection	Activated
Volume (µL)	100
Premix	Yes
Number of times	3
Mix number of times	5
Mix volume (µL)	100
Wait time (min)	0
Pressure (bar)	0.4
Positive pressure time (s)	300
Collect in position	A





Solvent Properties

	Solvent Description
1	Acetonitrile
2	
3	
4	
5	
6	
7	
8	
9	
10	



Solvent	1	2	3	4	5	6	7	8	9	10
Reservoir type		Refi	llable				N	on Refillab	le	
Capacity	N/A									
Aspiration flow rate (mL/min)	10									
Dispense flow rate (mL/min)	20									
Lower air gap flow rate (mL/min)	20									
Lower air gap volume (µL)	5									
Upper air gap flow rate (mL/min)	120									
Upper air gap volume (µL)	100									
Upper air gap dispense pause	300									
Conditioning?	Yes									
Conditioning number of times	2									
Conditioning flow rate (mL/min)	20									
Chlorinated	No									
Serial dispense	No									



Screenshot

Sample	Air Gap	
Sample name	Lower air gap flow rate (mL/min)	
PPT/PLD Sample	20	
Sample description	Lower air gap volume (µL)	
Sample for PPT/PLD method:	5	
Aspiration flow rate (mL/min)	Upper air gap flow rate (mL/min)	
50	120	
Dispense flow rate (mL/min)	Upper air gap volume (µL)	
220	700	
	Upper air gap dispense pause (ms)	
	1000	

"Sample" Screen	
Sample name	PLD/PPT Sample
Sample description	PLD/PPT Sample
Aspiration flow rate (mL/min)	50
Dispense flow rate (mL/min)	220
Lower air gap flow rate (mL/min)	20
Lower air gap volume (µL)	5
Upper air gap flow rate (mL/min)	120
Upper air gap volume (µL)	700
Upper air gap dispense pause	1000

Extraction Media	Pipetting Height	
Name	Solvent dispensation height (mm)	
PLD+ for 25 OH Vit D	-135.0	
Manufacturer	Sample dispensation height (mm)	
Biotage	-135.0	
Part number	Aspiration height (mm)	
918-0050-P01	-151.0	
Sorbent load (mg)		
0	Tune Pipetting Heights	
Capacity volume (µL)		
0		
Format		
96 -		
Comment		

Sample Plate/Rack Pipeting Height Name 2 mL sample plate, 96 Creative volume (st) -%2.0 1800 -%2.0 Format -96 Tune Pipetting Heights...

"Extraction Media" Screen

Name	ISOLUTE® PLD+
Manufacturer	Biotage
Part number	918-0050-P01
Sorbent load (mg)	0
Capacity volume (µL)	0
Format	96
Comment	N/A
Solvent dispensation height (mm)	-135.0
Sample dispensation height (mm)	-135.0
Aspiration height (mm)	-151.0

"Sample Plate/Rack" Screen

Name	2 mL Sample Plate, 96
Capacity volume (µL)	1800
Format	96
Aspiration height (mm)	-162.0
Pre-treatment dispensation height (mm)	-128.0



< cancet Edit Pipette Tip - 1000 μL Biotage tip	Save >
Pipette Tip Nime 1000 µL Biotage tip Manufacturer Biotage Part number 414141 Capacity (jsl.) 1000 Length (mm) 95	

"Pipette tip" Screen

Name	1000 µL Biotage Tip
Manufacturer	Biotage
Part number	414141
Capacity (µL)	1000
Length (mm)	95

Additional Comments

In this application the ISOLUTE® PLD+ extraction media default setting are edited such that the tip mixes the very top 100 μ L of the 500 μ L mixture. If the default ISOLUTE[®] PLD+ settings are used there is a greater risk of tip blockage during mixing.

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