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A Simplified and Automated Extraction Method for the Determination of 25-OH Vitamin D₂/D₃ in Human Serum Using a Strata[®] RP On-Line SPE Column

Xianrong (Jenny) Wei, Matthew Brusius, and Sean Orłowicz
Phenomenex, Inc., 411 Madrid Ave., Torrance, CA 90501 USA



Xianrong (Jenny) Wei
Senior Scientist
Jenny is a Senior Scientist in the Phenomenex PhenoLogix applications laboratory.

Overview

In this technical note, we explore the effectiveness of an automated on-line Solid Phase Extraction (SPE) method with a Thermo Cohesive system ran in TX mode coupled with MS/MS analysis for the characterization of 25-OH Vitamin D₂/D₃ from human serum. The relationship between the on-line extraction and LC column was investigated in the context of Aria[®] based software. The assay includes a shorter 30 x 3.0mm Kinetex[®] 2.6µm C18 LC analytical column and a 50 x 0.5mm Strata RP on-line SPE column, which reduces the total run time to under 6 minutes including the system equilibration. The assay shows the accuracy and precision, including LLOQ (n=6), and the resulting method is assessed for a linear dynamic range from 2-100ng/mL. **Figure 1** displays the Aria based software overview, while **Table 1** provides the on-line SPE conditions and **Table 2** shows the LC methodology.

Materials

25-OH Vitamin D₂ and D₃ standards were purchased from Ceriliant[®] (Round Rock, TX). Double charcoal stripped human serum was purchased from BioreclamationIVT[®] (Westbury, NY) All other reagents and chemicals were obtained from Sigma - Aldrich[®].

Experimental Conditions

Sample Pre-treatment

1. Dilute 150µL of human serum* with 200µL of Precipitating Reagent**
2. Add 10µL of 25-OH Vitamin-D₃-₂H₆ (1 µg/mL), mix for one minute
3. Centrifuge at 14,000 RPM for 10 minutes
4. Transfer 200µL supernatant to autosampler injection vial

*Double Charcoal-stripped human serum was used to prepare all Standards and QCs

**Precipitating Reagent prepared as (5:2:1) Methanol/Acetonitrile/Zinc Sulfate

On-Line Solid Phases Extraction

On-line SPE Column: Strata RP
Dimensions: 50 x 0.5 mm
Part No.: 00B-S326-AF

LC Conditions

Column: Kinetex[®] 2.6µm C18
Dimensions: 30 x 3.0 mm
Part No.: 00A-4462-YO
Guard Column: SecurityGuard[™] ULTRA Cartridges
Guard Part No.: AJ0-8775
Mobile Phase: A: 0.1 % Formic acid in Water
B: 0.1 % Formic acid in Methanol
Gradient: See Table 2
Flow Rate: See Table 2
Needle Wash 1: Methanol/Water (50:50)
Needle Wash 2: 0.1 % Formic acid in Water
Instrument: Cohesive System run in TX Mode: Agilent[®] 1260 with Leap Technologies PAL autosampler LX-2
Instrument: MS/MS (SCIEX 4000 QTRAP[®]) APCI+

Figure 1.

Screenshot of Method from Aria Software
Extraction column conditions (blue)
Analytical column conditions (pink)

| Step | Start | Sec | Flow | Grad | %A | %B | %C | %D | Val | Loop | Flow | Grad | %A | %B | Comments |
|------|-------|-----|------|------|-------|------|----|----|------|------|------|------|------|------|---|
| 1 | 0.00 | 30 | 0.75 | Step | 30.0 | 70.0 | | | **** | out | 0.75 | Step | 30.0 | 70.0 | Extract sample |
| 2 | 0.50 | 5 | 0.35 | Step | 30.0 | 70.0 | | | **** | out | 0.35 | Step | 30.0 | 70.0 | Slow down pumps |
| 3 | 0.58 | 60 | 0.35 | Step | 2.0 | 98.0 | | | T | in | 0.35 | Step | 30.0 | 70.0 | Transfer analytes |
| 4 | 1.58 | 30 | 1.50 | Step | 100.0 | | | | **** | out | 0.70 | Step | 10.0 | 90.0 | Separate analytes, wash extraction column |
| 5 | 2.08 | 90 | 1.50 | Step | 100.0 | | | | **** | out | 0.70 | Step | 10.0 | 90.0 | Elute analytes, wash extraction column |
| 6 | 3.58 | 30 | 0.75 | Step | 30.0 | 70.0 | | | **** | in | 0.70 | Step | 10.0 | 90.0 | Wash columns & valves |
| 7 | 4.08 | 30 | 0.75 | Step | 30.0 | 70.0 | | | **** | in | 0.75 | Step | 30.0 | 70.0 | Fill transfer loop, equilibrate HPLC column |
| 8 | 4.58 | 60 | 0.75 | Step | 30.0 | 70.0 | | | **** | in | 0.75 | Step | 30.0 | 70.0 | Equilibrate columns |

Table 1.

On-line SPE Conditions

| Step | Time (min) | Flow Rate (mL/min) | 0.1 % Formic acid in Water (A) | 0.1 % Formic acid in Methanol (B) | Comments |
|------|------------|--------------------|--------------------------------|-----------------------------------|---|
| 1 | 0 | 0.75 | 30 | 70 | Extract sample |
| 2 | 0.5 | 0.35 | 30 | 70 | Slow down pumps |
| 3 | 0.58 | 0.35 | 2 | 98 | Transfer analytes |
| 4 | 1.58 | 1.50 | 0 | 100 | Separate analytes, wash extraction column |
| 5 | 2.08 | 1.50 | 0 | 100 | Elutes analytes, wash extraction column |
| 6 | 3.58 | 0.75 | 30 | 70 | Wash columns and valves |
| 7 | 4.08 | 0.75 | 30 | 70 | Fill transfer loop, equilibrate HPLC column |
| 8 | 4.58 | 0.75 | 30 | 70 | Equilibrate columns |

Table 2.

LC Conditions

| Step | Time (min) | Flow Rate (mL/min) | 0.1 % Formic acid in Water (A) | 0.1 % Formic acid in Methanol (B) |
|------|------------|--------------------|--------------------------------|-----------------------------------|
| 1 | 0 | 0.75 | 30 | 70 |
| 2 | 0.5 | 0.35 | 30 | 70 |
| 3 | 0.58 | 0.35 | 30 | 70 |
| 4 | 1.58 | 0.70 | 10 | 90 |
| 5 | 2.08 | 0.70 | 10 | 90 |
| 6 | 3.58 | 0.70 | 10 | 90 |
| 7 | 4.08 | 0.75 | 30 | 70 |
| 8 | 4.58 | 0.75 | 30 | 70 |

Table 3.

MRM Transitions

| ID | Q1 Mass (DA) | Q3 Mass (DA) | Dwell (msec) | CE |
|--|--------------|--------------|--------------|----|
| 25-OH D ₂ 1 | 395.4 | 209 | 100 | 36 |
| 25-OH D ₂ 2 | 395.4 | 269.1 | 100 | 28 |
| 25-OH D ₃ 1 | 383.6 | 257.2 | 100 | 23 |
| 25-OH D ₃ 2 | 383.6 | 229.4 | 100 | 28 |
| D ₆ -25-OH D ₃ 1 | 389.5 | 263.3 | 100 | 23 |
| D ₆ -25-OH D ₃ 2 | 389.5 | 229.4 | 100 | 28 |

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Table 4.
Assay Recovery

| Serum Spiked at 100 ng | 25-OH Vitamin D ₂ | 25-OH Vitamin D ₃ |
|--------------------------|------------------------------|------------------------------|
| Average Area Ratio (n=4) | 4.48E-01 | 3.01E-01 |
| % CV (n=4) | 0.74 | 1.28 |

| Neat Solution at 100 ng | 25-OH Vitamin D ₂ | 25-OH Vitamin D ₃ |
|--------------------------|------------------------------|------------------------------|
| Average Area Ratio (n=4) | 4.69E-01 | 3.23E-01 |
| % CV (n=4) | 3.09 | 2.10 |

| Assay Recovery | 25-OH Vitamin D ₂ | 25-OH Vitamin D ₃ |
|----------------------------|------------------------------|------------------------------|
| Spiked Serum/Neat Solution | 95.5 | 93.2 |

Figure 2.
Representative of chromatograms of LLOQ at 2 ng/mL in human serum

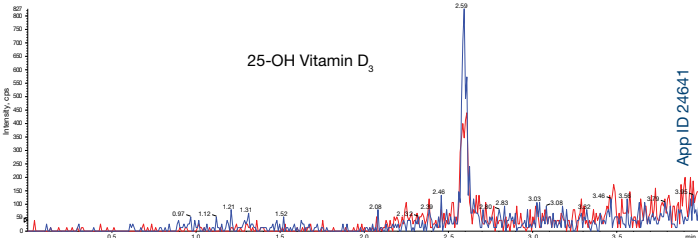
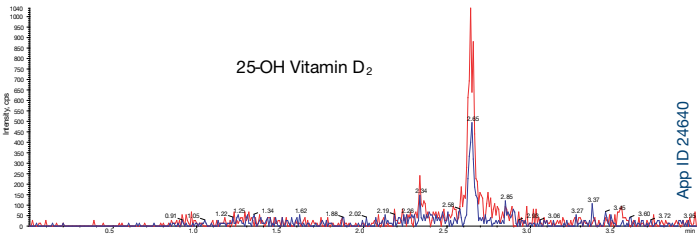


Figure 3.
Representative of chromatogram of ULOQ at 100 ng/mL in human serum

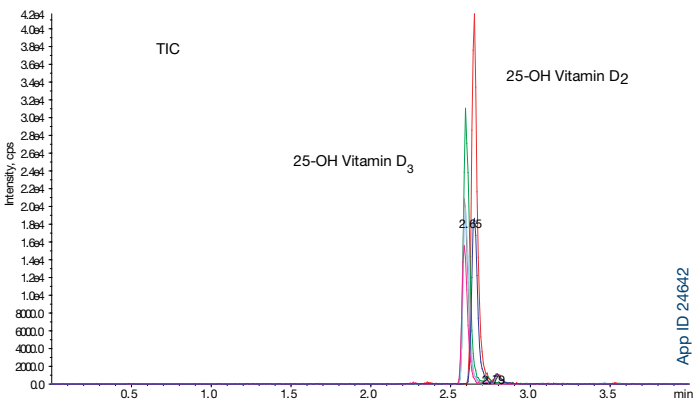


Figure 4.
Representative chromatograms of blank human serum matrix

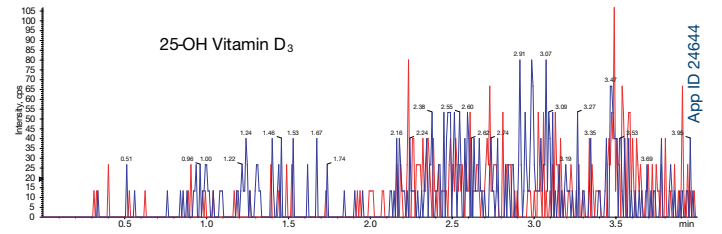
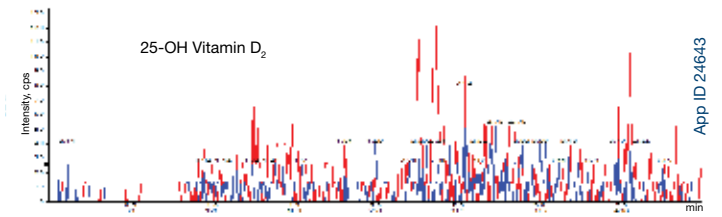
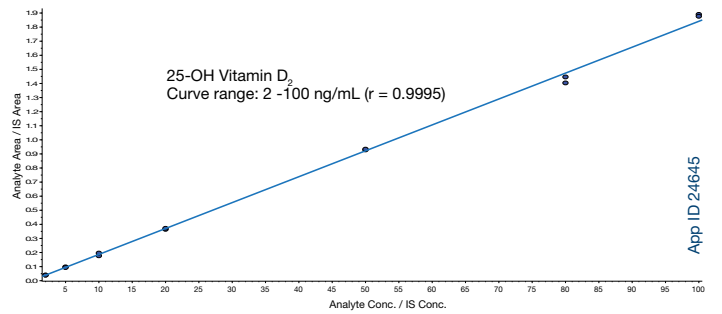


Figure 5.
Representative of the linearity of the curve (n=2)



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Table 5.
Accuracy and Precision

| | LLOQ | QCL | QCM | QCH |
|------------------------------------|------|------|------|------|
| Target Concentration (ng/mL) | 2 | 6 | 50 | 80 |
| 25-OH Vitamin D₂ | | | | |
| Mean Concentration Found (ng/mL) | 2.05 | 6.26 | 50.1 | 82.5 |
| % CV (n=6) | 8.8 | 3.42 | 3.63 | 3.5 |
| % Accuracy (n=6) | 103 | 104 | 100 | 103 |
| 25-OH Vitamin D₃ | | | | |
| Mean Concentration Found (ng/mL) | 1.83 | 6.05 | 50 | 84.4 |
| % CV (n=6) | 9.78 | 3.56 | 3.11 | 3.84 |
| % Accuracy (n=6) | 92 | 101 | 100 | 106 |

Results and Discussion

Sample Preparation and Cohesive System Set-Up

The on-line extraction is supplemented with an off-line protein precipitation. The precipitating reagent, Methanol/Acetonitrile/Zinc sulfate (5:2:1), has been optimized for both efficient protein removal and acceptable analyte recovery. Protein precipitation is required to prevent 25-OH Vitamin D₂/D₃ from binding to proteins in solution which would otherwise significantly reduce overall method sensitivity.

Table 1 displays the experimental details for the on-line SPE column. Contrary to off-line solid phase extraction, Step 1 (Blue) serves as both the loading and washing steps in this method and as such, the sample is loaded onto the extraction column under 70 % organic, (i.e. 70 % Mobile Phase B). A flow rate of 0.75 mL/min provides a good solvent volume that thoroughly washes the extraction column while the bed length is suitable for analyte retention.

Step 3 (Blue) is the elution of analytes from the extraction column and subsequent transfer to the analytical column. The flow rate for the elution is reduced to 0.35 mL/min and the organic content is increased to 98 % Mobile Phase B to maximize recovery off

the extraction column. For **Table 2**, in the same step (3 in Pink), the elution solvent mixes with a “dilution solvent” of 30 % mobile phase A flowing at 0.35 mL/min prior to reaching the analytical column. This mixing effectively serves as a dilution to mitigate against strong solvent effect to analytical column, since the analytical method (Step 4 in Pink) starts at 90 % Mobile Phase B.

Because the Cohesive system is engineering with focus mode, the dilution solvent and elution solvent's flow rates are additive, so it is important to reduce their combined flow rates equal to or below the starting 0.75 mL/min flow rate for the analytical method in Step 4 (Pink).

Steps 4-8 (Blue) serve to simultaneously wash and re-equilibrate the Strata RP On-line SPE column, while the isocratic LC method runs to completion of the separation of the analytes (Pink).

Assay Performance

To assess the performance of the method, an assay recovery was evaluated in **Table 4**, which is calculated by the peak area ratio of analytes spiked into serum divided by the response for analytes in a neat solution that is passed through both the extraction and analytical columns (TX Mode). This data shows that for both 25-OH Vitamin D₂ and D₃ recovery is greater than or equal to 93.3 %, while the RSD is less than or equal to 1.28 %.

The linear dynamic range of this method was tested with seven calibrators (n=2) from 2-100 ng/mL and the linearity of curve is shown in **Figure 5**, which shows an r=0.9995. Chromatograms of LLOQ and ULOQ are shown in **Figure 2** and **Figure 3**, respectively. The chromatogram for the blank matrix, double charcoal-stripped serum, is shown **Figure 4**, indicating that there is no detectable levels of 25-OH Vitamin D₂/D₃, so it was selected for standards and QCs preparation during the assay evaluation.

This assay was subsequently evaluated using four different level QC's, including LLOQ at n=6 for each sample set. The accuracy and precision is shown in **Table 5**, and it meets GLP environment regulations, respectively.

Conclusion

The combined Strata[®] RP On-Line SPE and LC analytical method result in a method with a total runtime of less than six minutes. The speed of this method associated with its accuracy and precision make it ideal for the high-throughput research environment.



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Ordering Information Kinetex[®] Core Shell LC Columns

| 2.6 µm MidBore™ Columns (mm) | | | | | | SecurityGuard™ ULTRA Cartridges [†] |
|------------------------------|-------------|-------------|-------------|-------------|-------------|---|
| Phases | 30 x 3.0 | 50 x 3.0 | 75 x 3.0 | 100 x 3.0 | 150 x 3.0 | 3/pk |
| C18 | 00A-4462-YO | 00B-4462-YO | 00C-4462-YO | 00D-4462-YO | 00F-4462-YO | AJ0-8775 for 3.0 mm ID |

[†] SecurityGuard ULTRA Cartridges require holder, Part No.: AJ0-9000.

Ordering Information Strata[®] On-Line SPE

| On-Line Extraction Columns (mm) | | | |
|---------------------------------|-------------|-------------|-------------|
| Phase | 50 x 0.5 | 50 x 1.0 | 20 x 2.1 |
| Strata RP | 00B-S326-AF | 00B-S326-AO | 00M-S326-AN |

Australia

t: +61 (0)2-9428-6444
f: +61 (0)2-9428-6445
auinfo@phenomenex.com

Austria

t: +43 (0)1-319-1301
f: +43 (0)1-319-1300
anfrage@phenomenex.com

Belgium

t: +32 (0)2 503 4015 (French)
t: +32 (0)2 511 8666 (Dutch)
f: +31 (0)30-2383749
beinfo@phenomenex.com

Canada

t: +1 (800) 543-3681
f: +1 (310) 328-7768
info@phenomenex.com

China

t: +86 400-606-8099
f: +86 (0)22 2532-1033
phen@agela.com

Denmark

t: +45 4824 8048
f: +45 4810 6265
nordicinfo@phenomenex.com

Finland

t: +358 (0)9 4789 0063
f: +45 4810 6265
nordicinfo@phenomenex.com

France

t: +33 (0)1 30 09 21 10
f: +33 (0)1 30 09 21 11
franceinfo@phenomenex.com

Germany

t: +49 (0)6021-58830-0
f: +49 (0)6021-58830-11
anfrage@phenomenex.com

India

t: +91 (0)40-3012 2400
f: +91 (0)40-3012 2411
indiainfo@phenomenex.com

Ireland

t: +353 (0)1 247 5405
f: +44 1625-501796
eireinfo@phenomenex.com

Italy

t: +39 051 6327511
f: +39 051 6327555
italiainfo@phenomenex.com

Luxembourg

t: +31 (0)30-2418700
f: +31 (0)30-2383749
nlinfo@phenomenex.com

Mexico

t: 01-800-844-5226
f: 001-310-328-7768
tecnicomx@phenomenex.com

The Netherlands

t: +31 (0)30-2418700
f: +31 (0)30-2383749
nlinfo@phenomenex.com

New Zealand

t: +64 (0)9-4780951
f: +64 (0)9-4780952
nzinfo@phenomenex.com

Norway

t: +47 810 02 005
f: +45 4810 6265
nordicinfo@phenomenex.com

Portugal

t: +351 221 450 488
f: +34 91-413-2290
ptinfo@phenomenex.com

Spain

t: +34 91-413-8613
f: +34 91-413-2290
espinfo@phenomenex.com

Sweden

t: +46 (0)8 611 6950
f: +45 4810 6265
nordicinfo@phenomenex.com

Switzerland

t: +41 61 692 20 20
f: +41 61 692 20 22
swissinfo@phenomenex.com

United Kingdom

t: +44 (0)1625-501367
f: +44 (0)1625-501796
ukinfo@phenomenex.com

USA

t: +1 (310) 212-0555
f: +1 (310) 328-7768
info@phenomenex.com

All other countries Corporate Office USA

t: +1 (310) 212-0555
f: +1 (310) 328-7768
info@phenomenex.com



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