Chemotyping the distribution of vitamin D metabolites in human serum

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Most studies examining the relationships between vitamin D and disease or health focus on the main 25-hydroxyvitamin D3 (25(OH)D3) metabolite, thus potentially overlooking contributions and dynamic effects of other vitamin D metabolites, the crucial roles of several of which have been previously demonstrated. The ideal assay would determine all relevant high and low-abundant vitamin D species simultaneously. We describe a sensitive quantitative assay for determining the chemotypes of vitamin D metabolites from serum after derivatisation and ultra-high performance liquid chromatography-electrospray ionisation-tandem mass spectrometry (UHPLC-ESI-MS/MS). We performed a validation according to the ‘FDA Guidance for Industry Bioanalytical Method Validation’. The proof-of-concept of the method was then demonstrated by following the metabolite concentrations in patients with chronic liver diseases (CLD) during the course of a vitamin D supplementation study. The new quantitative profiling assay provided highly sensitive, precise and accurate chemotypes of the vitamin D metabolic process rather than the usually determined 25(OH)D3 concentrations.

A debate over healthy levels of vitamin D has been waging in the scientific and popular press in recent years. While the vital role of vitamin D in bone health is clearly established1, its functional involvement in other diseases such diabetes, cancer, multiple sclerosis, depression, hepatic, renal and cardiovascular diseases is currently the subject of intense research2,3,4,5, particularly because vitamin D receptor is present in virtually all organs. Typically, the 25-hydroxyvitamin D (25(OH)D) metabolite is used for assessing vitamin D status, because it can be readily measured due to its long half-life and high concentration in blood, as well as the fact that the 25(OH)D level represents the optimum marker for substrate availability rather than the tightly regulated levels of other downstream metabolites6. In the photosynthesis and metabolism of vitamin D, the substrate vitamin D is transported to the liver, where it undergoes the first hydroxylation step at C-25 (catalysed by CYP2R1 and CYP27A1) to give 25(OH)D. This is followed by CYP27B1 conversion to the active form 1,25(OH)2D in the kidneys1. 1,25(OH)2D suppresses gene transcription by binding to vitamin D receptor (VDR), with CYP3A4 being the most important cytochrome P450 species supressed by 1,25(OH)2D7. It has been shown that patients, who were supplemented with vitamin D, exhibited increased clearance of atorvastatin (substrate of CYP3A4)8. While CYP24A1 activity dominates the 1,25(OH)D/25(OH)D catabolism in the healthy kidney, CYP3A4 dominates in the liver and small intestine, because of the higher level of basal and induced enzyme expression9. A correlation of plasma levels of vitamin D metabolites and intestinal CYP3A4 activity showed only low levels of vitamin D metabolites in patients with cirrhosis as a result of the reduction of CYP3A protein expression9. Intra and inter-individual differences of circulating vitamin D levels and associated intestinal CYP3A4 activities may also contribute to variability of oral bioavailability of vitamin D; utilizing these effects may allow customised supplementation of vitamin D in the future, if the CYP genotype status of an individual is known. Recently, Binkley et al.10 nicely demonstrated a mathematical model using baseline and 6 months metabolite levels of vitamin D3, 25(OH)D3 and 24,25(OH)2D3 to allow efficient “treat-to-target” prediction of 25(OH)D3 levels10.

In this simplified assessment, however, dynamic effects of downstream metabolites are overlooked. For example, after Holick et al.11 discovered 24,25(OH)2D3 as clearance product of vitamin D, it was initially assumed that 24,25(OH)2D3 is merely a catabolute without biological activity. Today, we know that 24,25(OH)2D3 plays crucial roles in intra-membrane/endochondral bone formation and bone fracture repair12. Increased activity of 24,25(OH)2D3 has also been associated with increased risk of mortality in renal patients13; it has also been

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suggested that 24,25(OH)D$_3$ is biologically active because it is recognised by the kidneys and converted to 1,24,25(OH)$_3$D$_3$ and 24-oxo-25(OH)D$_3$ in rats\textsuperscript{14}.

The importance of dynamic changes of vitamin D concentrations and intra/inter-individual variability after administration of fixed supplementation doses has also been demonstrated in several recent studies. For example, Bosworth \textit{et al.} observed decreased vitamin D catabolism and 1,25(OH)$_2$D$_3$ production in chronic kidney disease (CKD)\textsuperscript{13}. Stubbs \textit{et al.} showed CKD patients’ altered ability to increase serum 24,25(OH)$_2$D$_3$ after vitamin D$_3$ therapy, suggesting decreased 24-hydroxylase activity\textsuperscript{15}. de Boer \textit{et al.} investigated associations of estimated glomerular filtration rate (GFR) with circulating 24,25(OH)$_2$D$_3$ levels across populations with wide range of GFRs and found that lower estimated GFR was associated strongly with reduced vitamin D catabolism, as measured by 24,25(OH)$_2$D$_3$\textsuperscript{16}. Berg \textit{et al.}\textsuperscript{17} suggested the ratio of serum 24,25(OH)$_2$D$_3$ to 25(OH)D$_3$ (vitamin D metabolite ratio, VMR) as a new candidate biomarker for vitamin D status, while Wagner \textit{et al.}\textsuperscript{18} demonstrated that serum 24,25(OH)$_2$D$_3$/25(OH)D$_3$ ratios were predictive of 25(OH)D$_3$ response to supplementation\textsuperscript{18}. Finally, no significant differences for sera concentrations of 25(OH)D$_3$ was observed by Duan \textit{et al.}\textsuperscript{19} between a cohort of healthy subjects and multiple sclerosis patients, but levels of 1,25(OH)$_2$D$_3$ and 24,25(OH)$_2$D$_3$ were significantly lower in the patient group as compared to the healthy group.

Correlations of vitamin D metabolite distributions (chemotypes) and disease/health phenotypes therefore are an interesting topic to study, requiring metabolomics techniques for vitamin D metabolites for measurement. Of course, these assays would need to determine all relevant high and low-abundant vitamin D species simultaneously with equal analytical figures of merit. Commonly applied techniques for vitamin D are immunoassays and liquid chromatography-tandem mass spectrometry (LC-MS/MS)\textsuperscript{20}. These methods are often restricted to 25(OH)D$_3$ because of limited sensitivity and specificity\textsuperscript{20,21}. Fortunately, recent technical advances have permitted analysis of metabolites in the metabolic cascade beyond 25(OH)D$_3$, including lower abundant species\textsuperscript{21,22}.

In the present study, we developed a new assay for quantitative measurement of vitamin D chemotypes based on ultra-high performance liquid chromatography (UHPLC)-MS/MS. We performed a validation according to the U.S. Food and Drug Administration (FDA)’s ‘Guidance for Industry Bioanalytical Method Validation’ and demonstrated the assay’s performance by following patients with chronic liver diseases (CLD) during a vitamin D supplementation study.

**Results**

Important features of the new vitamin D chemotyping assay were: (1) it required only 50μL of human serum and was conducted in parallel 96-well plate format; (2) metabolites were derivatised for enhanced response. The derivatisation protocol was developed for considerably improving the reagent’s performance by eliminating the formation of multiple diastereomers; (3) stable isotope standards were used for all investigated metabolites, to correct for systematic errors from matrix interferences and differential ionisation properties of metabolites.

The chemotyping method for each serum sample comprised extraction of 50μL of serum by supported liquid extraction (SLE), ultra-high-performance liquid chromatography separation using a fluorinated HPLC column and detection and quantification by tandem mass spectrometry using specific multiple reaction monitoring (MRM) transitions for each investigated metabolite (see Table S1, Supplementary Information, for optimised conditions). A representative LC-MS/MS chromatogram is shown in Fig. 1. Importantly, baseline chromatographic resolution was achieved for all species, including the 25(OH)D$_3$ epimers; previously, baseline resolution was not achieved for the PTAD-derivatised 25(OH)D$_3$ epimers\textsuperscript{23}. The assay was then applied to a cohort of CLD patients. The measured chemotypes for all investigated patients are summarised in Figure S1 (Supplementary Information).
Information). For the FDA-compliant validation of the chemotyping assay24, vitamin D-free blank serum samples were spiked with varying amounts of the investigated vitamin D compounds. Tables S1-S4 (Supplementary Information) summarise the full validation results.

Analyte recovery from the SLE material was determined by spiking vitamin D-free human serum with 25(OH)D3, 25(OH)D2 and 3-epi-25(OH)D3 at 50 ng/mL; and 1,25(OH)2D3, 1,25(OH)2D2 and 24,25(OH)2D3 at 50 pg/mL. After performing the extraction, the corresponding isotope standards were added to the extracts (n = 3). The individual recoveries were: 25(OH)D3, 74%; 25(OH)D2, 70%; 3-epi-25(OH)D3, 72%; 1,25(OH)2D3, 64%; 1,25(OH)2D2, 62%; 24,25(OH)2D3, 68%.

Precision of the assay was excellent, between 1.9–3.9% RSD (intra-day) and 4.1–6.0% (inter-day). The upper and lower limits of the calibration range for all metabolites were chosen, so that all patient samples fell into the target range. Linear regression analysis demonstrated excellent linearity over the calibration range (10 pg/mL–100 ng/mL, R² > 0.998). Limits of quantification (LOQ) for all compounds were between 10 pg/mL and 1 ng/mL, as determined by the implemented lowest calibrators. Limits of detection (LOD) were several orders of magnitude lower (Table 1) and demonstrated that quantitative measurements are readily possible at much lower levels, after extending the calibration range to lower concentrations. Six in-house calibration standards in vitamin D-free serum were measured using five determinations per concentration; none of these measurements deviated by more than 11% (20% is acceptable24). Quality control samples at five concentrations were measured in duplicate and showed only small deviations between 5–8% (15% is acceptable24).

Ion suppression effects were negligible (between 0.01–1.7% at 10 ng/mL, depending on the individual metabolites).

Stability experiments under different storage conditions (room temperature, 4 °C, −80 °C) demonstrated only limited sample degradation (< 6.8%) under all studied conditions (Table S4, Supplementary Information; for more information).

Unequivocal assignment of analyte peaks in the MRM chromatograms of samples to the investigated metabolites, in particular for the closely-eluting dihydroxylated isomers 1,25(OH)2D3 and 24,25(OH)2D3, was achieved by matching the retention times of the extracted ion chromatograms (using the quantifier ions) with those of the isotope standards from a separate calibrator sample. These numbers were always within ± 0.15 min or smaller.

In addition, the quantification results for 25(OH)D3 were compared to results from a routine clinical vitamin D assay for accuracy assessment. Bland-Altman analysis for 25(OH)D3 exhibited minor concentration-dependent bias at low concentrations (< 10 ng/mL) in comparison to the Diasorin chemiluminescence assay, showing that the difference between assays was higher at very low concentrations of 25(OH)D3.

### Table 1. Limits of quantification (LOQ), limits of detection (LOD), linear dynamic range as well as intra- and inter-day precision. S/N = 3.

| Metabolite       | LOQ (serum) (ng/mL) | LOD (serum) (pg/mL) | Linearity (ng/mL) | Precision intra-day (CV, %) | Precision inter-day (CV, %) |
|------------------|---------------------|---------------------|-------------------|-----------------------------|----------------------------|
| 25(OH)D3         | 1                   | 0.05                | 1–100             | 2.1                         | 5.3                        |
| 25(OH)D2         | 1                   | 0.05                | 1–100             | 2.4                         | 5.1                        |
| 3-epi-25(OH)D3   | 0.25                | 0.04                | 0.25–65           | 1.9                         | 4.1                        |
| 1,25(OH)2D1      | 0.01                | 0.04                | 0.01–0.5          | 3.9                         | 5.4                        |
| 1,25(OH)2D2      | 0.01                | 0.04                | 0.01–0.5          | 3.6                         | 6.0                        |
| 24,25(OH)2D3     | 0.1                 | 0.05                | 0.1–25            | 3.2                         | 4.3                        |

Figure 2. Bland-Altman analysis for the newly developed UHPLC-MS/MS method versus the Diasorin chemiluminescence assay for the 25(OH)D3 levels using all investigated patient samples. The analysis exhibited minor concentration-dependent bias at low concentrations (<10 ng/mL) in comparison to the Diasorin chemiluminescence assay, showing that the difference between assays was higher at very low concentrations of 25(OH)D3.
For the CLD patient samples, circulating concentrations of 25(OH)D$_3$ were between 7–60 ng/mL and values for 1,25(OH)$_2$D$_3$ were in the range of 10–100 pg/mL; levels of 24,25(OH)$_2$D$_3$ were approximately an order of magnitude lower than 25(OH)D$_3$ for most patients. The vitamin D$_2$ metabolites 25(OH)D$_2$ and 1,25(OH)$_2$D$_2$ were not found in the investigated CLD patient samples. The 3-epi-25(OH)D$_3$ species was present in all samples between 0.1–4.5 ng/mL, demonstrating the importance of separating 3-epi-25(OH)D$_3$ from 25(OH)D$_3$ to avoid overestimation of vitamin D status (Fig. 1).

**Discussion**

In the present study, we developed profiling techniques for capturing the dynamic chemotypes of vitamin D metabolites in human biofluids. Serum 25(OH)D$_3$ levels are routinely used as a measure for vitamin D status$^{27}$, due to 25(OH)D$_3$’s long half-life, relatively high concentration and clear relation to serum vitamin D$_3$$^{28}$. Here, the goal was to measure multiple high and low abundant vitamin D metabolites with the same performance as that obtained for 25(OH)D$_3$. The new developed method was specific, sensitive and precise for simultaneous quantification of six vitamin D metabolites using a serum volume of only 50 µL in parallel 96-well plate format. The newly developed cold-temperature derivatisation combined with high resolution chromatographic separation was very simple to perform and avoided previous problems with diastereomer formation and epimer resolution$^{25,29,30}$. Stable isotope standards were used for all investigated metabolites, to correct for systematic errors from matrix interferences and differential ionisation properties of metabolites.

One important further aspect of metabolite derivatisation is that the converted metabolites showed similar electrospray ionisation (ESI) response factors (0.93–0.99). This was not surprising considering the permanently-charged quaternary ammonium function, which was expected to dominate the remainder of the hydrophobic structure. It provides a response-leveling effect. Because of this response equalisation, other vitamin D metabolites, conjugates or yet undiscovered vitamin D compounds can be readily quantified with analytical figures of merit expected to be similar to those described above even without using dedicated stable isotope standards.

We applied the assay to a subset of patients for the validation of the assay and preliminary characterisation. Importantly, these patients served as their own controls as we compared all vitamin D values before, during and after a six-month supplementation. This procedure provided patient specific chemotypes of vitamin D metabolites at each of the investigated time points and allowed both intra-individual as well as inter-individual comparisons. For example, the separate determination of 3-epi-25(OH)D$_3$ from 25(OH)D$_3$, provided accurate vitamin D status, but, more importantly, it reveals considerable differences in the 3-epi-25(OH)D$_3$ to 25(OH)D$_3$ ratios. The absolute concentration levels for 3-epi-25(OH)D$_3$ mentioned in the previous section hide the fact that these levels corresponded to strongly varying amounts of the 3-epimer relative to 25(OH)D$_3$. In the investigated patients, these values corresponded to between 1.6 and 12% of the individuals’ 25(OH)D$_3$. Similar subject-specific ratios have also been observed in previous studies$^{31}$.

The assay was able to capture serum concentrations of patients with severe vitamin D deficiency (i.e., serum 25(OH)D$_3$ ≤ 10 ng/mL), which was present in nine (30%) of patients included in the analysis. A clear inverse linear correlation was observed between baseline 25(OH)D$_3$ and response to supplementation: patients with the lowest 25(OH)D$_3$ at baseline exhibited the largest increase in 25(OH)D$_3$ ($P < 0.001$; Fig. 3a). Additionally, lower baseline 25,25(OH)$_2$D$_3$, correlated with larger changes to 25(OH)D$_3$ concentrations ($P = 0.005$; Fig. 3b). This finding however, cannot be generalised to the entire sample. Three women with severe vitamin D deficiency had much higher baseline 25,25(OH)$_2$D$_3$, though these concentrations were not detected in men (Fig. 3c).

Interestingly, these three supplemented women attained serum 25(OH)D$_3$ levels within the normal range, which coincided with decreased 24,25(OH)D$_3$ concentrations. In contrast, patients with lower baseline 24,25(OH)D$_3$ demonstrated increases in response to increasing 25(OH)D$_3$ levels. Additionally, patients with a greater response to vitamin D supplementation tended to have lower concentrations of 3-epi-25(OH)D$_3$, as shown in Fig. 3d ($P < 0.001$). Importantly, the measured chemotypes include accurate C-3 epimer contributions for 3-epi-25(OH)D$_3$ and are not skewed by inter-epimer response variations as reported before$^{22}$.

In conclusion, we developed a highly sensitive, precise and accurate assay for simultaneous determination of six relevant vitamin D metabolites in human serum over a wide dynamic range. The assay provided chemotypes of the vitamin D metabolic process rather than the usually determined 25(OH)D$_3$ concentration; the latter gives an incomplete picture that may or may not correlate with function or disease. The chemotypes can be extended to include other vitamin D compounds without compromise as the implemented procedure provides similar performance for all metabolites. Furthermore, the assay can be readily extended to quantitative measurements at concentration levels several orders of magnitude lower than described here after appropriate re-validation; the limits of quantification reported here were determined by calibration solutions chosen for the specific application. In summary, the new metabolomics technique is believed to be a useful tool with the potential to aid the understanding of the pathobiological function of vitamin D in health and disease, and to predict individual response to vitamin D supplementation.

**Methods**

**Materials.** Standards of 25(OH)D$_2$, 3-epi-25(OH)D$_3$, 1,25(OH)$_2$D$_3$, and 24,25(OH)$_2$D$_3$ and methanol (HPLC grade) were purchased from Sigma-Aldrich (Steinheim, Germany). Stable isotope-labelled d$_6$-25(OH)D$_3$ and d$_6$-1,25(OH)$_2$D$_3$ were from Chemaphor (Ottawa, ON, Canada); d$_6$-3-epi-25(OH)D$_3$, d$_6$-25(OH)D$_2$, d$_6$-1,25(OH)$_2$D$_3$, and d$_6$-24,25(OH)$_2$D$_3$ from Endotherm (Saarbrücken, Germany). Stock solutions were prepared in methanol (0.1 mg/mL); working solutions were obtained prior to use by dilution. Organic-free water was generated by a Millipore (Bedford, MA, USA) Direct-Q8 purification system. Lyophilised ClinCal and ClinCheck 25-OH-Vitamin D$_3$/D$_2$ serum calibrators (level 0–III) and quality control sera (level I, II) were from Recipe (Munich, Germany) and reconstituted in water prior to use. Vitamin D-free human serum as calibration and
quality control matrix was purchased from Golden West Biologicals (Temecula, CA, USA). Supported liquid extraction 96-well AC micro-extraction plates were from Tecan (Männedorf, Switzerland). Amplifex reagent was obtained from Sciex (Concord, ON, Canada).

CLD patient samples. Serum samples of patients with chronic liver diseases (CLD) of various etiologies were used from a study at the Department of Medicine II, Saarland University Medical Center (Homburg, Germany). Blood samples were collected, the serum was extracted, aliquoted and frozen at −80 °C. Haemolysis of blood samples was minimised as outlined in the sample collection, handling and transport SOP of the department’s laboratory. The samples were neither subjected to prolonged light exposure, nor to freeze/thaw cycles. One aliquot per patient was thawed out shortly before conducting the LC-MS/MS analyses. Furthermore, studies have shown that the vast majority of haemolysed specimens (~95%) only slightly haemolyse under typical collection, transport, handling and storage conditions. All patients had their serum 25(OH)D3 levels assessed at baseline (t0). Those with vitamin D deficiency (25(OH)D3 < 30 ng/mL) were supplemented with vitamin D3 at 20,000 IU/week (Dekristol, Jenapharm, Jena, Germany) for six months. All other patients were monitored as controls. Serum 25(OH)D3 and metabolite levels were measured at different time points during the study (t1, t2 and t3, at 3, 6 and 12 months, respectively). All patients provided written informed consent and the study was approved by the local research ethics committee (Ärztekammer des Saarlandes, ref. 57/11). The study was conducted in accordance with the good clinical practice guidelines as defined in the Declaration of Helsinki. A subset of 30 patients (120 samples) was used for the validation of the assay and preliminary characterisation.

Sample preparation. Serum calibrators, quality control and patient samples were extracted in parallel by means of supported liquid extraction (SLE) using 96-well micro-extraction plates (AC extraction plates). The extraction method for 25(OH)D3, 3-epi-25(OH)D3, 1,25(OH)2D3 and 24,25(OH)2D3 was adapted from our previously reported technique for 25(OH)D3. The following internal standards and buffer solutions were used: (A) internal standard: d6-25(OH)D3 (50 ng/mL), d6-3-epi-25(OH)D3 (25 ng/mL), d6-1,25(OH)2D3 (0.20 ng/mL) and d6-24,25(OH)2D3 (15 ng/mL) in acetonitrile; (B) extraction buffer (0.2 M sodium carbonate/sodium hydroxide 1:1 (v/v) in water/acetonitrile 95:5 v/v); (C) washing buffer (water/methanol 90:10 v/v); and (D) elution buffer (water/methanol 10:90 v/v). A mixture of internal standard and extraction buffer (A/B 1:2 v/v) was used for the validation of the assay and preliminary characterisation.
(150 μL) was transferred to each extraction well and 50 μL of serum was added. To release vitamin D metabolites from the binding proteins, horizontal shaking (Eppendorf Thermomixer, Hamburg, Germany) was performed (10 min); the supernatant was discarded, and the extraction phase washed with the washing buffer (200 μL) by horizontal shaking (2 min). Finally, all vitamin D metabolites were eluted with elution buffer (200 μL) by horizontal shaking (5 min). Drying of serum sample extracts was performed using an Eppendorf concentrator (model 5301) under vacuum conditions at 30 °C; the drying time depended on the number of samples (5 min-2h).

**Derivatisation reaction.** The reagent used for derivatisation of the vitamin D compounds was Amplifex Diene Reagent Kit, consisting of two components, Diene Reagent and Diene Dilution. For derivatisation, 900 μL of Diene Diluent solution was added to the Diene reagent vial and the mixture was vortexed. 50 μL of the derivatisation mixture was added to the dried serum sample extracts, followed by vortexing for 20 s and incubation for 30 min in a fridge at 4 °C. A difficulty arose during the derivatisation from the two possible sides of attack of the cis-diene moiety of vitamin D compounds (Fig. 4)[25,34. The reagent can link to the molecule from both α and β sides of vitamin D compounds, giving two epimeric products for each compound. As a result, two peaks may be expected for each compound in the MRM ion chromatograms. We observed a kinetic effect and utilised it to improve the reaction: derivatisation under cold conditions at 4 °C directed the reaction favourably into a single epimeric species. To our knowledge, derivatisation at low temperatures and forcing the reaction into predominiantly a single epimeric species has not yet been described. Because dedicated stable isotope standards were utilised for all metabolites, this differential formation had no impact on the accuracy of the determination. Hedman et al. developed a highly sensitive assay for 1,25(OH)2D3 using Amplifex labeling without specifically mentioning the epimeric products[25. Other derivatisation procedures for vitamin D also showed epimeric products, e.g. after PTAD derivatisation[25,34 and usually either the major product was used for quantification or the flow rate was reduced to try eluting both species in a single peak. Importantly, derivatisation shifted the molecular weights to higher m/z regions with less isomeric and isobaric interference as compared to the native molecules[36,37. The final step comprised termination of the reaction by adding 50 μL of methanol/ deionised water (50:50 v/v) and vortexing for 20 s, followed by transfer to the HPLC autosampler vials.

**Liquid chromatography-mass spectrometry.** Five microliter of each sample was injected in triplicate into a Dionex Ultimate 3000 UHPLC system (Thermo-Fisher, Bremen, Germany). Separations of the six vitamin D metabolites were performed on a Phenomenex (Torrance, CA, USA) Kinexet PFP 100 A column (100 × 2.1 mm, d = 2.6 μm) at 25 °C and flow rate of 0.4 mL/min using gradient elution. The mobile phases were: (A) water + 0.1% formic acid and (B) methanol + 0.1% formic acid. The gradient started at 50% B and was held for 0.3 min, then increased to 70% and held there for 7 min, and further increased to 85% B and held for 1.2 min. Finally, B was increased to 90% and held for 4 min before returning to the initial conditions and re-equilibration for 5 min. The UHPLC system was coupled to a Scienx QTRAP 5500 quadrupole-linear ion trap (QqLIT) mass spectrometer via electrospray ionisation (ESI) operated in positive ion mode. Ion source and MS parameters were as follows: source temperature: 550 °C, curtain gas: 50 psi, gas 1: 45 psi, gas 2: 45 psi, collision exit potential: 20 V and collision energy: 40–43 V declustering potential: 0–130 V, optimised for each of the individual vitamin D metabolites (Table S1, Supplementary Information). Quantification of 25(OH)D3, 1,25(OH)2D3 and 24,25(OH)2D3 was performed by multiple reaction monitoring (MRM) using the [M + H]+ → [(M + H)–(Ni(CH3)3)]+ transitions for all measured vitamin D metabolites with 200 ms dwell time per transition (Table S1, Supplementary Information). The overall chromatographic run time including column wash/equilibration was 15 min per sample. Data acquisition was performed using Analyst software 1.6 (Sciex) and MultiQuant 2.1.1 (Sciex) for quantification of all vitamin D metabolites. Linear regression analysis was performed by the least-squares method to evaluate the calibration curve of each analyte as a function of its serum concentration.

**Chemiluminescence immunoassay.** Serum levels of 25(OH)D3 of the CLD patients samples were determined in Saarland University Medical Center’s central laboratory (Department of Clinical Chemistry and Laboratory Medicine) using the Liaison 25-OH Vitamin D chemiluminescence immunoassay (DiaSorin, Dietzenbach, Germany).
FDA Guidance for Industry-Bioanalytical Method Validation. We performed the method validation in accordance with guidelines of the United States Food and Drug Administration (FDA)\textsuperscript{24};

Cross-validation. assay was compared to validated reference method (chemiluminescence assay) using incurred samples.

Selectivity. blank samples of appropriate biological matrix (serum) were obtained from different sources (human serum, vitamin D-free human serum, horse serum).

Reproducibility. reproducibility was assessed by replicate measurements using the new assay, including quality controls and incurred samples (Table S1, Supplementary Information).

Calibration curve. calibration curves were generated for each analyte in the sample in the same biological matrix as the samples in the intended study by spiking the matrix with known concentrations of the analyte. Six concentrations of standards were chosen on the basis of the concentration range expected in the conducted study; analysed in triplicate in each run. (Tables S2 and S3, Supplementary Information).

Accuracy, precision and recovery. a minimum of three concentrations in the range of the expected study sample concentrations was performed and measured using a minimum of five determinations per concentration (results should not deviate by more than 20%). Recovery experiments were performed by comparing analytical results for extracted samples at three concentrations (low, medium, high) with non-extracted standards that represent 100% recovery (Table S3, Supplementary Information).

Quality control samples. three concentrations (low level, medium level, high level) in triplicate were incorporated into each run. At least 67% of them (in total) should be within 15% of their respective theoretical values and 50% (at each level) should be within 15% of their theoretical concentrations (Table S3, Supplementary Information).

Matrix effects (ion suppression). Matrix effects were evaluated for five different serum samples at three different concentration levels: high (50 ng/mL), medium (30 ng/mL), low (10 ng/mL), as described by Matuszewski et al.\textsuperscript{38} The experimental strategy comprised a first set of serum samples, which were fortified after SLE with analytes and internal standards at three different concentration levels. The second set of samples consisted of pure analytes and internal standards at three different levels in mobile phase without performing SLE. The absolute matrix effect was calculated by dividing the mean peak areas of each metabolite from the first set (with SLE) by the mean of each metabolite peak area of the second set (without SLE). The obtained values (expressed in percent) were subtracted from 100% to give the amount of ion suppression in the presence of matrix.

Stability study. Stability was tested using vitamin D-free blank serum (n = 5) spiked with the metabolites of interest at three concentration levels: high (50 ng/mL), medium (30 ng/mL), low (10 ng/mL). Short-term temperature stability was evaluated for 24 h at room temperature, for 24 h at 4 °C within the autosampler, and for 24 h at −80 °C in the freezer. Additionally, three freeze/thaw cycles at −20 °C and a long-term stability experiment (20 d) were performed (Table S4, Supplementary Information). On the day of the LC-MS/MS analysis, the internal standard was added and analysis performed in an identical manner to the patient samples as described above; LC-MS/MS experiments were performed in duplicate for each sample. The calculated concentrations for each stability experiment were compared to the measured sample concentration of the day of preparation.

Ruggedness. The robustness of the method was examined by systematically modifying several experimental parameters. The HPLC flow rate was varied between 0.3–0.5 mL/min (regular 0.4 mL/min) using two different mixtures of vitamin D metabolites at different levels: 10 and 40 ng/mL for 25(OH)D\textsubscript{3}, and 3-epi-25(OH)D\textsubscript{3}, and 10 and 40 pg/mL for the dihydroxylated species 1,25(OH)\textsubscript{2}D\textsubscript{3} and 24,25(OH)\textsubscript{2}D\textsubscript{3}. Additionally, the column temperature was investigated between 20–30 °C (regular 25 °C). In these experiments, one parameter was held constant, while the other was varied: first the HPLC flow rate was set to 0.3 mL, while the column temperature was set to 20, 25 and 30 °C. This procedure was repeated for the flow rates of 0.4 and 0.5 mL/min. In all ruggedness-testing experiments, the differences of peak areas in comparison to the standard settings were always less than 4%.

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Author Contributions
M.J.J. performed all analytical experiments, analysed and interpreted the data, and drafted the manuscript. C.S.S. designed the supplementation study, collected patient samples and contributed to interpretation of the data. F.L. contributed to the design of the supplementation study and interpretation of the data. D.A.V. conceived, designed and coordinated the study, analysed and interpreted the data, and drafted the manuscript. All authors revised the manuscript for important intellectual content and approved the final version for publication.

Additional Information
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