Detection and Quantification of Vitamins in Microliter Volumes of Biological Samples by LC-MS for Clinical Screening

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A method for simultaneous determination of water-soluble vitamins B1, B2, B3 (nicotinamide), B5, B6 (pyridoxamine), B9 and fat-soluble vitamins E (α-tocopherol) and K1 in tears, and B1, B2, B3, B5, B6, B9, A (retinol), and E in blood serum is described. Liquid chromatography-mass spectrometry (LC-MS) was used with a ternary mobile phase of water and acetonitrile containing 0.1% formic acid and methanol containing 5 mM ammonium formate. Vitamins were quantified using an internal standard method. Using 25 μL injection volumes, the limits of detection were in the range of 0.066-5.3 ng in tear, and 0.087-1.1 ng in serum with linear responses for all vitamins. Intra- and inter-day precision and recoveries were satisfactory. This is the first study to demonstrate simultaneous vitamin detections in microliters of biological samples which has distinct advantages in many diagnostic applications with limited available fluids (e.g., tears; elderly anemic blood) or sampling small subjects (e.g., rodents).  © 2018 American Institute of Chemical Engineers AIChE J, 64: 3709–3718, 2018

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Introduction

Essential biochemical functions of vitamins in the human body include important roles in protein metabolism, maintenance of blood glucose levels, regulation of cell growth, and cell differentiation.1 Thirteen different vitamins are currently recognized by the USDA2 and are classified into two groups according to their solubility; water-soluble vitamins include all B vitamins and vitamin C whereas the fat-soluble vitamins include vitamins A, D, E, and K. Water-soluble vitamin deficiencies can cause permanent tissue damage and debilitating effects in humans while fat-soluble vitamins assist in anabolic and catabolic pathways in the body and are a current source of interest to nutritionists and clinicians.

All living tissues require vitamins and nutrients. The cornea is the outermost, transparent layer of living cells in the eye that helps focus light and protect the complex network of nerves and tissues in the eye.3 The metabolism of the cornea requires a constant supply of amino acids, vitamins, and other nutrients; no blood vessels extend to the cornea,3,4 so tears likely supply these nutrients. Previously we demonstrated the determination of water-soluble vitamins B1, B2, B3, B5, and B9 and fat-soluble vitamin E in tears and blood serum via two separate LC-MS methods.5,6 Water-soluble and fat-soluble vitamins have diverse chemical structures and properties, making their determination from a single chromatography assay challenging.7 Previously reported vitamin detection methods predominantly focused on determining individual vitamins or a subset of vitamins with similar polarities. For example, HPLC methods exist for a subset of water-soluble vitamins in blood serum, multivitamin tablets, and food.8-11 HPLC assays for several fat-soluble vitamins in blood serum, tablets, and daily products have been reported as well.12-17 There are also methods for simultaneous extraction of water-soluble and fat-soluble vitamins except they eventually used separate analytical methods for analysis.16,19 Although simultaneous detection of water-soluble and fat-soluble vitamins in a single chromatography run from a
single aliquot of sample has been reported, these methods are only reported to determine vitamin contents in uncomplicated, non-biological matrices, such as pharmaceutical preparations,20-23 foodstuff,20,22,24,25 and beverages.26 The method by Ferreiro-Vera et al.27 was able to determine only two vitamins with different polarities in the blood serum while unified supercritical fluid and liquid chromatography method by Taguchi et al.28 was only validated with standard solutions. Determination of tear vitamins A,29,30 D,31 and C32,33 are also separately reported in the literature. To date, no simultaneous vitamin detection method is available for extraction and detection of multiple vitamins with different polarities from biological samples.

In this article, we report a new, ternary solvent LC method using electrospray ionization (ESI) mass spectrometry for the identification and quantification of water-soluble and fatsoluble vitamins. The present LC-MS method is able to capture all seven B vitamins and five fat-soluble vitamins (including two forms of vitamins D); while B1, B2, B3 (nicotinamide), B6, B9, B12 (pyridoxamine), B10 (retinol), and E (α-tocopherol) were successfully detected in blood serum and B1, B2, B3 (nicotinamide), B5, B6, E (α-tocopherol), and K1 were successfully detected in human tear samples.

To improve upon assays that only measure a subset of vitamins, this combined method was developed using commonly available and robust analytical tools to provide a more complete nutritional status with reduced material and chemical demands, reduced instrument preparation and run times, smaller sample volumes, and shortened active technician time. Furthermore, small sample requirements improve the ability to detect vitamins in infants, sample limited situations (e.g., tears), elderly patients, or those with anemia. This article describes the methodology used in a larger clinical study. Results from an extension of this method applied to tears and blood serum of 45 infant/parent pairs will be published in a subsequent article. Here, we describe the technical aspects of the combined analytical strategy, which enables simultaneous determination of most water-soluble and fat-soluble vitamins without the need for separate assays.

Materials and Methods

Materials and chemicals

The purchased standard water-soluble and fat-soluble vitamins were: thiamine hydrochloride (reagent grade, ≥99%, HPLC), (−)-riboflavin (≥98%), nicotinamide (≥98%, TLC), d-pantothenic acid hemicalcium salt (≥98%, TLC), pyridoxamine dihydrochloride (≥98%), biotin (≥99%, TLC), folic acid (≥97%), retinol (synthetic, ≥95% [HPLC], crystalline), cholecalciferol (pharmaceutical secondary standard), 25-hydroxycholecalciferol (≥98%, HPLC), (±α)-tocopherol (pharmaceutical secondary standard), and phylloquinone (analytical standard) from Sigma-Aldrich (St. Louis, MO). Vitamin metabolites were selected based on the clinical assays34,35 and availability in biological samples36,37 (details in Supporting Information Section 2).

Stable isotope internal standards (IS) of vitamins B1 (thiamine-[4-methyl-13C-thiazol-5-yl-13C3] hydrochloride), E (α-tocopherol-[ring-5,7-dimethyl-d10]), and K1 (5,6,7,8-d4, 2-methyl-d3) were purchased from Sigma-Aldrich (St. Louis, MO). Vitamin metabolites were selected based on the clinical assays34,35 and availability in biological samples36,37 (details in Supporting Information Section 2).

For extraction of water-soluble and fat-soluble vitamins simultaneously from blood, 800 μL of MeOH/ACN/Acetone, 1:1:1 (v/v/v), containing 1 μM of each water-soluble and fat-soluble vitamin IS and 200 μL of 1.5 g/L BHT solution were added to 200 μL plasma and vortexed. The mixture was incubated at 4°C for 10 min to precipitate proteins, then vortexed, and centrifuged for 10 min at 896 relative centrifugal force (rcf) to separate plasma. The plasma was removed and stored at −20°C in a glass container.

For extraction of water-soluble and fat-soluble vitamins simultaneously from tears, 700 μL of MeOH, 400 μL of MeOH/ACN/Acetone, 1:1:1 (v/v/v), containing 1 μM of each water-soluble and fat-soluble vitamin IS and 200 μL of 1.5 g/L BHT solution were added to 200 μL plasma and vortexed. The mixture was incubated at 4°C for 10 min to precipitate proteins, then vortexed, and centrifuged for 10 min at 896 rcf. The supernatant (serum) was dried under nitrogen and analytes were reconstituted in 100 μL of 0.1% FA in water/MMeOH, 9:1 (v/v). Simultaneous extraction of water-soluble and fat-soluble vitamins from tears was accomplished from the two tear strips (70 μL tears) via the addition of 400 μL MeOH, ACN, acetone (1:1:1 by volume). A 2 μM water-soluble and fat-soluble IS was added along with 70 μL BHT solution for vitamin stabilization. The vial containing sample and solvents was incubated at 4°C for 10 min, then centrifuged at 896 rcf for 10 min. Supernatant was dried under a gentle stream of nitrogen and reconstituted in 100 μL of 0.1% FA in water/MMeOH, 9:1 (v/v).
LC-MS/MS analysis

LC-MS/MS was performed using an Accela LC quaternary pump coupled with an autosampler and an LQI Fleet MS with an electrospray (ESI) probe (Thermo Scientific, Waltham, MA). Separation was performed using a Waters (Milford, MA) Atlantis T3 column, 2.1 mm × 150 mm, packed with 3 μm C18 silica and 100 Å pore size coupled with a guard column (Atlantis T3 Sentry, 2.1 mm × 10 mm). A 1.5 ratio post-column binary fixed flow splitter (20% to MS, 80% to waste) was used to increase the analyte ionization efficiency (this is unnecessary with a heated ESI probe.).

The ternary mobile phases were (a) 0.1% FA in water, (b) 0.1% FA in ACN, and (c) 5 mM ammonium formate in MeOH. The gradient was 0 min, 100% A; 7 min, 100% A; 12 min, 50% A/50% B; 16 min, 20% A/80% B; 16.01 min, 100% C; 34 min, 100% C. ESI and MS parameters were optimized over six segments with 4 kV spray voltage and 275°C capillary temperature for all vitamins. The capillary and tube lens voltages were optimized over time and after instrument maintenance and were in the range of 1-46 and 60-115 V, respectively. The electrospray probe was operated in the positive ion mode in segments 1-6 and the sheath gas flow rate was set to 10 and 20 (arbitrary units), respectively, for water-soluble and fat-soluble vitamins. A 25 μL sample with full loop injection was introduced at a flow rate of 0.2 mL/min for the first 16 min and 0.4 mL/min for the last 18 min. The column was re-equilibrated between runs with 20 column volumes of mobile phase A for 20 min. Autosampler and column temperatures were fixed at 5 and 25°C, respectively. Nitrogen was used as a nebulizing gas. All data acquisition was done with Xcalibur 2.3 (version C; Thermo Fisher Scientific, Waltham, MA).

Linearity and limit of detection

For vitamin quantifications, signals from triplicate analysis of the six calibration solutions were measured and calibration curves were built by plotting the ratio of analyte peak area to the area of IS vs. concentration using the least-squares regression method.

Limits of detection (LOD) and limits of quantification (LOQ) were determined based on the standard deviation method. The LOD and LOQ were respectively determined based on the standard deviation of the replicate injections (n = 7) of a low-level sample (standard solution, tear, or blood serum) and m is the slope of the linear calibration curve. The standard deviations were calculated for concentrations lower than the LOQ and the RSD were <20%.

Results and Discussion

In our related publication, we developed separate HPLC methods for the determination of water-soluble and fat-soluble vitamins in tears and blood serum. These methods required 18 min for the water-soluble vitamins separation and 25 min for the fat-soluble vitamins separation and each method needed 28 min for column re-equilibrium. Furthermore, separate extractions were required and the total blood serum volume was 500 μL. We used these methods as a starting point for the development of a combined method, which is described here. Original implementations of the combined method took >60 min, but flow rate and gradient optimization enabled run times to be cut in half. The combined method described herein can detect 12 water-soluble and fat-soluble vitamins in <34 min using three mobile phases which reduced the total sample preparation and analysis time by 42% compared to our separate methods. Eight of these vitamins were successfully extracted and detected from tears and blood serum using a single extraction step in <30 min with a total sample volume of 200 μL blood serum and 70 μL tears. This method therefore enables detection of a majority of water-soluble and fat-soluble vitamins simultaneously from limited sample volumes.

Optimization of LC-MS/MS conditions

Using the ternary gradient elution and ESI-MS/MS conditions described in “LC-MS/MS Analysis” section, water-soluble and fat-soluble vitamins eluted from the LC column in <34 min as shown in Figure 1. The chromatography run was divided into six segments with respect to the analyte retention times to allow the ion trap mass analyzer to scan the precursor ions (listed in Table 1) using selected ion scanning mode. For quantifications, MS/MS specifications were used with selected reaction monitoring. Thus, specific fragment ions of each precursor ion were captured in order to increase the resolution and selectivity.

Figure 1 shows the chromatograms achieved by a standard mixture solution containing the 12 water-soluble and fat-soluble vitamins under described LC-ESI-MS/MS conditions. Chromatograms were generated from the signals for the MS/MS fragment ions listed in Table 1. Peak areas were used for quantification.

The separation mechanism selected for detection of water-soluble and fat-soluble vitamins was based on the structure of the compounds. An extensive literature review was conducted to minimize the time and cost for the method development. Water-soluble vitamins are polar compounds while fat-soluble vitamins are relatively less polar than water-soluble vitamins. Thus, a reversed phase C18 column was selected because it is known to be an appropriate method for separation of compounds that differ by polarity. Reverse phase separations start with a high aqueous mobile phase then increase the solvent composition throughout the gradient. Acetonitrile and methanol are common solvents used in reversed phase separations. The LC separation was optimized by changing compositions of the mobile phases in order to achieve the best and fastest separation (Supporting Information Figure S1). For the mass spectrometry detection, the polar functional groups in water-soluble vitamin structures enable easy ionization by protonation. Ion formation was enhanced by 0.1% FA added to the mobile phases A and B (mass spectra are shown in Figure 2). Fat-soluble vitamins are considerably less polar than water-soluble vitamins and lack functional groups that readily accept or donate electrons, so mobile phase additives were necessary to facilitate their ionization with adduct formation using the hydroxyl (vitamin A, D3, 25[OH]D3, and E) or oxygen (vitamin K1) groups in their structures. Mobile phase additives that have been used to promote ion formation for fat-soluble vitamins include formic acid, silver perchlorate, ammonium acetate, ammonium formate, or cesium acetate. We systematically tested these additives at different concentrations. Cationic adducts from ammonium acetate, silver perchlorate, and cesium acetate did not yield sufficient spectral intensity (Figure 3). However, the hydrogen adduct peak height generated by 5 mM ammonium formate demonstrated sufficiently enhanced ionization concurrent with increasing mobile phase C pH to ~6 (below pKa of 9.25 for ammonium ion) and was thus chosen as the third mobile phase modifier that elutes fat-soluble vitamins. The mass spectra for
all 12 vitamins and IS are shown in Figure 2 with the base peaks representing the protonated vitamins (Table 1).

**Analysis of standard vitamins with optimized LC-MS**

Using the identified LC-MS parameters, the water-soluble and fat-soluble vitamin retention times were determined from three replicate injections of the 0.3-10 μM concentrations as documented in Table 1. A summary of the chromatography and mass spectrometry parameters for detection of water-soluble and fat-soluble vitamins simultaneously is provided in Table 1. All 12 vitamins were detected in positive ESI mode. Vitamin B_{12} and its corresponding IS (thiamine-[13C4]) precursor ions were observed as molecular ions, [M]^{+}. The precursor ions of vitamin A and its IS (retinol-[d5]) resulted from the dehydration of the protonated molecule, [M + H-H_{2}O]^{+}. Peaks at m/z 429 and 435 were observed for vitamin E and its IS (tocopherol-[d6]), which are produced from dehydrogenation of the protonated molecule to yield [M + H-H_{2}]^{+}.
other vitamins and their IS, precursor ions were generated from the protonated molecule, [M + H]\(^+\). The time periods of the six segments are also shown in Table 1. For quality assurance, Table 1 also lists some fragment ions produced with the optimized collision energies along with ones used for quantification.

**Linearity, LOD, LOQ, and precision**

The LOD and LOQ were calculated by the method described in the “Linearity and limit of detection” section and were determined in standard solutions for all 12 vitamins, and in tears and serum samples for detectable vitamins. Table 2 reports the standard curves and \( R^2 \) values for each of the 12 vitamins. The instrument response was linear for all vitamins with correlation coefficients >0.99. The ranges of linearity for vitamins B\(_5\) and B\(_6\) were up to 200 \( \mu \)M, B\(_2\), B\(_3\), B\(_9\), 25(OH)-D\(_3\), D\(_3\), E, and K\(_1\) were up to 100 \( \mu \)M, B\(_1\), and A were up to 50 \( \mu \)M and B\(_7\) up to 10 \( \mu \)M. These values indicate sufficiently high reliability that is consistent with other published techniques.\(^{13}\)

Inter-day (\( n = 7 \)) and intra-day (\( n = 6 \)) precision were evaluated with replicate injections of samples. RSD values were in the range of 1.6-12\% for tear vitamin B\(_9\) (Table 3). Tear B\(_9\) recovery was low (29\% and 57\% RSD) and as such, calculated amounts in serum samples to be less reliable than other vitamins. Precisions, except for vitamin B\(_9\) were sufficient for use in subsequent assays and consistent with other vitamin techniques.\(^{13,22}\) B\(_9\) recovery was also tested using a stable B\(_9\) isotope, B\(_9\)-[\(^{13}\)C\(_5\)], as an internal standard. Low recoveries of ~10\% were still obtained from the serum. This result is likely due to the combination of instability and low concentrations for this tested form of vitamin B\(_9\) in serum samples. According to the Mayo Clinic,\(^{45}\) more sensitive methods such as competitive binding assays are required for reliable detection of vitamin B\(_9\). In these assays, folate is measured as an indicator of all folic acid derivatives, which in serum is almost entirely present as N-(5)-methyl tetrahydrofolate.\(^{46}\)

**Tears and serum analysis**

To test the combined method performance on complex biological samples, vitamins were extracted from tears and blood serum of five human subjects under procedures described in the “Blood and tear preparation” section, and analyzed with the LC-MS/MS combined detection method for water-soluble and fat-soluble vitamins. Vitamin concentrations resulting from the five individuals are summarized in Table 4. The combined method detected vitamins B\(_1\), B\(_2\), B\(_3\), B\(_5\), B\(_6\), B\(_8\), A, and E in blood serum and vitamins B\(_1\), B\(_2\), B\(_3\), B\(_5\), B\(_6\), B\(_8\), and E in tears. Serum B\(_9\) recovery and tear B\(_9\) precision were low as described in the “Linearity, LOD, LOQ, and precision” section, thus the detected amounts are not reported in Table 4. This method was also tested on newborn tears resulting in detection of vitamin K\(_1\). The detectable vitamin K\(_1\) concentrations in newborns can be explained by the vitamin K shot that they receive after birth. This data will be reported in a future article. Figure 4 shows the vitamin chromatograms achieved from analysis of a tear sample (Figure 4a) and a serum sample (Figure 4b) under the described combined method. Our combined method, compared to our previously published method, is capable of detecting most water-soluble and fat-soluble vitamins simultaneously in human blood and it is also the first demonstration of simultaneous detection of these two groups of vitamins in human tears. A longitudinal study will be published with the application of this combined method for vitamin detections in infants and parents.

Our combined method was also tested using a triple quadrupole MS with higher sensitivity at our co-author’s laboratory in the Kellogg Eye Institute and the same vitamins remained undetectable. Undetected K\(_1\) in serum was attributed to the very low sample concentrations (0.0004-0.002 \( \mu \)M\(^2\)) which were lower than our method LOD. K\(_1\) was also undetectable in serum of newborns which lead us to hypothesize that vitamin K is probably higher in tears than serum of newborns. The undetected B\(_7\) and vitamin D metabolites were tested for potential matrix effects. Standard solutions of these vitamins were spiked into pooled serum and tear samples such that precision and
Figure 2. ESI-mass spectra for water-soluble vitamins (with 0.1% formic acid in mobile A and B) and fat-soluble vitamins (with 5 mM ammonium formate in mobile phase C) and their corresponding internal standards (stable isotope substituted analytes) generated from a vitamin standard solution.

Precursor ions of the analytes are labeled on the spectra. The fragment ions for each of these precursor ions are reported in Table 1 and were used for quantification. The mass spectra of vitamin D₃ and E are shown in Figure 3.
recoveries were calculated. B7 and D vitamins recovery was 91.6–103% with inter- and intra-day precision of 3.4–9.0%.

Thus, the undetected vitamins were not due to matrix effects. This is not surprising since the presence of IS would elucidate matrix effects. However, B7 and vitamin D are both protein-bound vitamins,48,49 such that a proteolysis step is required to break the protein bond and release the vitamins prior to protein precipitation. The body requires extremely small concentrations of B7 which can efficiently be recycled, and food sources with B7 are abundant,50 thus the inability to detect B7 is not a major concern. Deficiency of vitamin K is also quite rare because intestinal bacteria produce this vitamin, which is also abundant in many foods.51 The biologically active form of vitamin D is the 25(OH)D considered for nutritional health diagnosis.52 Thus, only one of the critical vitamins for nutritional health, 25(OH)D, was undetectable in the time and resource-efficient combined method presented herein.

Comparisons of results with the literature revealed additional efficiencies with our combined method. Liquid-liquid extractions (LLE) with hazardous organic solvents (e.g., hexane) or solid-
phase extractions were primarily used to pre-concentrate vitamin analytes prior to HPLC.\textsuperscript{10,27,53} Our previously published separate water-soluble and fat-soluble vitamin methods\textsuperscript{5} included three LLE steps with hexane for fat-soluble vitamins. In the combined method herein, both groups of vitamins were extracted under a single extraction step using MeOH/ACN/Acetone. Chatzimichalakis et al.\textsuperscript{10} published a simultaneous method for determination of only B-complexes (thiamine, riboflavin, nicotinic acid and nicotinamide, pyridoxine, folic acid, and cyanocobalamin) in blood serum and pharmaceuticals with solid-phase extraction and total analysis time of 27 min. While in our combined method, six different B vitamins were extracted and eluted in <15 min with a flow rate of four times less and lower detection limits along with two fat-soluble vitamins.

In separate LC-MS methods, which are perceived to be the rapid standard, analysis takes 5 min with at least 15 min of active technician time for sample preparation for each vitamin. In the method published by Papadoyannis et al.\textsuperscript{53}, solid-phase extraction cartridges were necessary to separate water-soluble and fat-soluble vitamins from 500 μL of blood plasma. Further, vitamin detection required two different HPLC columns.

Our combined method analyzed eight vitamins in <30 min with 3 min or less active technician time. Thus, our total assay time was 33 min for eight vitamins compared to 20 min per sample for a total time of 160 min for eight vitamins. Further, since our combined method requires only one column cleaning and stabilization cycle, while the separate assays require one each, solvent utilization decreases and instrument utilization time can increase by 42%. Other sensitive quantification methods, such as enzyme linked immunosorbent assay (ELISA), require 2-24 h assay time and ~30 min active technician time per vitamin assay. When compared to the combined method presented herein, active technician time is reduced by a factor of ~5. Thus, the method presented herein demonstrates advantages beyond prior protocols for determination of water-soluble and fat-soluble vitamins in serum samples. To the best of our knowledge, simultaneous detection of water-soluble and fat-soluble vitamins in complex biological samples has not been previously reported in the literature. This combined method provides a resource-lean and efficient means to simultaneously detect most water-soluble and fat-soluble vitamins in complex biological samples.

### Table 4. Vitamin Concentrations (μM) in Tears and Blood Serum of Five Tested Individuals

<table>
<thead>
<tr>
<th>Vitamins</th>
<th>Sample 1 Tear</th>
<th>Sample 2 Tear</th>
<th>Sample 3 Tear</th>
<th>Sample 4 Tear</th>
<th>Sample 5 Tear</th>
<th>Sample 1 Serum</th>
<th>Sample 2 Serum</th>
<th>Sample 3 Serum</th>
<th>Sample 4 Serum</th>
<th>Sample 5 Serum</th>
</tr>
</thead>
<tbody>
<tr>
<td>B\textsubscript{1}, thiamine</td>
<td>0.022</td>
<td>0.054</td>
<td>0.038</td>
<td>0.14</td>
<td>N.D.</td>
<td>0.13</td>
<td>0.027</td>
<td>0.013</td>
<td>N.D.</td>
<td>0.053</td>
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<tr>
<td>B\textsubscript{2}, riboflavin</td>
<td>0.018</td>
<td>0.034</td>
<td>N.D.</td>
<td>0.036</td>
<td>0.060</td>
<td>0.034</td>
<td>0.032</td>
<td>N.D.</td>
<td>0.015</td>
<td>0.018</td>
</tr>
<tr>
<td>B\textsubscript{3}, niacinamide</td>
<td>4.0</td>
<td>1.9</td>
<td>1.3</td>
<td>0.92</td>
<td>5.0</td>
<td>2.4</td>
<td>1.3</td>
<td>1.1</td>
<td>6.9</td>
<td>0.35</td>
</tr>
<tr>
<td>B\textsubscript{5}, pantothenic acid</td>
<td>0.49</td>
<td>0.44</td>
<td>0.12</td>
<td>0.12</td>
<td>0.78</td>
<td>0.46</td>
<td>0.23</td>
<td>0.13</td>
<td>0.21</td>
<td>0.16</td>
</tr>
<tr>
<td>B\textsubscript{6}, pyridoxamine</td>
<td>N.D.</td>
<td>0.62</td>
<td>0.083</td>
<td>0.71</td>
<td>0.12</td>
<td>1.3</td>
<td>N.D.</td>
<td>0.51</td>
<td>0.22</td>
<td>0.42</td>
</tr>
<tr>
<td>A, retinol</td>
<td>–</td>
<td>1.1</td>
<td>–</td>
<td>2.9</td>
<td>–</td>
<td>2.7</td>
<td>–</td>
<td>1.4</td>
<td>–</td>
<td>1.0</td>
</tr>
<tr>
<td>E, α-tocopherol</td>
<td>0.13</td>
<td>14</td>
<td>0.090</td>
<td>16</td>
<td>0.42</td>
<td>20</td>
<td>0.055</td>
<td>9.2</td>
<td>0.12</td>
<td>7.1</td>
</tr>
</tbody>
</table>

N.D., not detected.

Figure 4. LC-MS/MS chromatograms of water-soluble and fat-soluble vitamins detected in (a) a tear sample and (b) a blood serum sample using the <30 min, reduced materials/chemicals combined vitamin method presented herein.
Conclusions

This article describes the first demonstration of simultaneous determination and quantification of eight water-soluble and fat-soluble vitamins from clinically obtained human tears and blood serum samples. Our simultaneous protocol was able to capture 12 water-soluble and fat-soluble vitamins in <34 min from standard solutions, while among these vitamins, B1, B2, B3 (nicotinamide), B5, B6 (pyridoxamine), B9, E (α-tocopherol), and K1 were simultaneously extracted and detected in tears and B1, B2, B3 (nicotinamide), B5, B6 (pyridoxamine), B9, A (retinol), and E (α-tocopherol) were simultaneously extracted and detected in blood serum in <30 min. Previously published methods have not demonstrated simultaneous extraction and detection of these vitamins from biological samples. The combined method presented herein optimized extraction solvents combined with tuned LC properties such as column, ternary mobile phase, and eluent modifiers to produce sufficient sensitivity and peak resolutions that are consistent or better than separate methods. Also, isotopically labeled versions of the target analytes utilized as IS reduced sample preparation errors and compensated for matrix effects and recoveries.

Compared to separate methods for water-soluble and fat-soluble vitamins, our combined method decreased instrument preparation and run time, reduced active technician time by a factor of 5, reduced material and chemical demands, and reduced sample demands. Sample preparation time was also shortened as a single extraction step efficiently extracted both water-soluble and fat-soluble vitamins. This combined method is highly beneficial for applications with limited availability of samples (e.g., infant tears; elderly anemic blood) or sampling small subjects (e.g., rodents). Furthermore, this combined method detects all but one of the critical water-soluble and fat-soluble vitamins for clinical screening and provides substantial time and resource savings for nutritional assessments from biofluids.

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Notation

ACN acetoniitrile
BHT butylated hydroxytoluene
D2O deuterium oxide
DMSO dimethyl sulfoxide
ESI electrospray ionization
FA formic acid
IS internal standard
LC liquid chromatography
LOD limits of detection
LOQ limits of quantification
MeOH methanol
MS mass spectrometry
rcf relative centrifugal force
RSD relative standard deviation
RT retention time

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multivitamin preparations by qNMR. J Agric Food Chem. 2015; 63(12): 3135-3143.


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