Microfluidic Analytical System with On-Line Luminol Chemiluminescence Detection Based on Annular Flow of Phase Separation Multiphase Flow

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Abstract

Microfluidic analytical system was developed based on annular flow of phase separation multiphase flow with a ternary water-hydrophilic/hydrophobic organic solvent solution. The analytical system was combined with on-line luminol chemiluminescence detection for catechin analysis. The water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate mixed solution (3:8:4, volume ratio) containing 60 μM luminol and 2 mM hydrogen peroxide as a carrier was fed into the capillary tube (open-tubular fused-silica, 75 μm inner diameter, 110 cm effective length) at a flow rate of 1.0 μL·min⁻¹. The carrier solution showed stable chemiluminescence as a baseline on the flow chart. Eight catechins were detected as negative peaks for their antioxidant potential with different detection times. The system was applied to analyze the amounts of catechin in commercially available green tea beverages.

Keywords

Microfluidic Analytical System, Phase Separation Multiphase Flow, Catechin, On-Line Luminol Chemiluminescence Detection

1. Introduction

Fluidic flow in a tube can be typically classified into one of two groups: homogeneous single-phase flows, which do not have a liquid-liquid interface, and immiscible multiphase flows, which have a liquid-liquid interface [1]. Electroosmotic and laminar flows generated in a microspace are basically classified...
as homogeneous single-phase flows. Over the last century, capillary electrophoresis [2], micellar electrokinetic chromatography [3], and hydrodynamic chromatography [4], were developed by manipulating electroosmotic and laminar flows. Various types of immiscible multiphase flows that generate liquid-liquid interfaces using water and hydrophobic organic solvents have been reported [5] [6]. These reports show droplet, slug, parallel, or annular flows depending on the conditions employed. Investigation concerning a liquid-liquid interface in microfluidic flow has been interesting and attractive in Lab-on-a-Chip and μ-TAS research area [7] [8] [9]. However, to our knowledge, there have been no reports regarding capillary chromatography taking advantage of liquid-liquid interfaces based on such multiphase flows.

We have recently developed a method of generating a multiphase flow using two-phase separation mixed solvent solutions [10] [11] [12] [13]. When mixed solvent solutions, such as a water-hydrophilic/hydrophobic organic solvent solution, are delivered into a microspace, such as a capillary tube and a microchannel on a microchip, phase transformation of the mixed solvent solution occurs in the microspace with changes in temperature and/or pressure, leading to a kinetic liquid-liquid interface. These new types of multiphase flows are called “phase separation multiphase flows”. We can also observe droplet, slug, parallel, and annular flows under different conditions in phase separation multiphase flow. We are particularly interested in annular flows, including inner and outer phases, generated in a phase separation multiphase flow. The specific microfluidic phenomenon and flow are termed the “tube radial distribution phenomenon” (TRDP) and “tube radial distribution flow” (TRDF), respectively [14] [15].

We have experimentally and theoretically investigated phase separation multiphase flows from the viewpoints of the mechanism of generation and functional appearance [14] [15] [16] [17]. Open-tubular capillary chromatography, designated as tube radial distribution chromatography (TRDC), has been developed based on the annular flow or TRDF from a phase separation multiphase flow. Various types of analyte were chromatographically separated through the open-tubular capillary tube on the TRDC system, where the inner and outer phases in the TRDF act as the mobile and pseudostationary phases, respectively [14] [15].

In this study, we first examined luminol chemiluminescence (CL) behavior in a ternary mixed solvent solution of water-acetonitrile-ethyl acetate, and the mixed solvent solution was found to enhance luminol CL in time and intensity under certain conditions. Based on these new experimental data, we combined the TRDC system with on-line luminol CL detection for catechin analysis. The present fluidic analytical system was used to detect eight catechins with each detection times on the CL profile and further analyze the amounts of catechin in commercially available green tea beverages.

2. Experimental

2.1. Reagents and Materials

Water was purified with an Elix UV 3 system (Millipore Co., Billerica, MA). All
reagents used were commercially available and of analytical grade. Luminol, hydrogen peroxide, acetonitrile, ethyl acetate, antioxidant reagents, and eight catechins as well as catechin mixture from green tea were purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). Fused-silica capillary tubes with an inner diameter of 75 μm were purchased from GL Science Co. (Tokyo, Japan).

2.2. CL Measurement in a Batch System

The CL profile was examined with a batch-type detection system (CLA-FS4; Tohoku Electronic Industrial Co. Ltd., Sendai, Japan). To the water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate mixed solution (3:8:4, volume ratio; 2.5 mL) containing 0.1 μM luminol in a petri dish was added 15 mM hydrogen peroxide solution (50 μL) with constant stirring to obtain the CL profile. To examine the suppressive effects of various antioxidant reagents on CL, antioxidant reagents dissolved in ternary mixed solvent solution (250 μM, 0.1 mL) were added to the solutions in petri dishes that emitted around the maximum CL intensity, and the decreases in CL intensities due to their antioxidant potential were measured.

2.3. Fluidic Analytical System with On-Line CL Detection

Figure 1 shows a schematic diagram of the present analytical system with on-line CL detection, comprised of an open-tubular fused-silica capillary tube (75 μm inner diameter and 140 cm length (effective length, 110 cm)), a micro-syringe pump (MF-9090; Bioanalytical Systems, Inc., West Lafayette, IN), and a CL detector (Kimoto Electronic Co. Osaka, Japan). The tube temperature was controlled by dipping the capillary tube in water maintained at a definite temperature in a beaker with stirring. Water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate mixture (3:8:4 volume ratio) solution containing 60 μM luminol and 2 mM hydrogen peroxide was used as a carrier solution. Analyte solutions were prepared with the carrier solution.

The analyte solution was introduced directly into the capillary inlet side by the gravity method (20 cm height for 15 s). After analyte injection, the capillary inlet was connected through a joint to a micro-syringe. The syringe was placed on the micro-syringe pump. The carrier solution was fed into the capillary tube at a

Figure 1. Schematic diagram of the present microfluidic analytical system with on-line CL detection.
flow rate of 1.0 μL-min⁻¹ under laminar flow conditions. Negative peaks observed from the CL baseline were detected by the CL detector.

3. Results and Discussion
3.1. CL Profile with the Ternary Mixed Solvent Solution

The phase diagram of water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate is shown in Figure 2; the dotted line indicates the boundary or solubility curve. The CL profiles were examined for ternary mixed solvent solutions with (a) organic solvent-rich and (b) water-rich component ratios, with a batch-type CL measurement system. Mixed solutions of (a) and (b) are most commonly used as carrier solutions in the TRDC system [14] [15]. In the TRDC system the specific homogeneous mixed solutions are required for generating TRDF; the component ratios of them are positioned near the boundary curve on the phase diagram, such as (a) and (b), and when they become away from the curve, TRDF cannot be observed because phase transformation does not occurs. The obtained CL profiles are shown in Figure 3, taken together with the CL profile obtained with 10 mM phosphate buffer (pH 7.3) solution. From the obtained CL profiles, the component ratio (a) showed the greater and longer lasting CL, and was selected as the carrier solution for use in the system. It was completely confirmed that the CL lasted more than 8 hour with good reproducibility. There was little information of CL in the ternary water-hydrophilic/hydrophobic organic solvent mixed solution, and the reason why the ternary mixed solution enhanced CL is not yet clear. However, the ternary homogeneous solution near the boundary

![Figure 2. Phase diagram for the ternary water-acetonitrile-ethyl acetate mixed solvent solution. The dotted curve in the diagram indicates the homogeneous-heterogeneous solution boundary curve. The component ratios of the solvents, (a) and (b), are water (10 mM phosphate buffer pH 7.3)-acetonitrile-ethyl acetate of 3:8:4 and 15:3:2 volume ratio, respectively.](image-url)
3.2. Effects of Antioxidants on CL Profiles

The effects of antioxidants on the CL profiles were examined using a batch-type CL measurement system. The analytical conditions were described in the Experimental section. 2-Propanol and nordihydroguaiaretic acid that acted as an antioxidant for hydroxyl radicals (·OH) decreased the CL intensity by 80% and 100%, respectively, compared to the maximum CL intensity in the CL profiles. Other antioxidants, ascorbic acid for superoxide radical anion (·O₂⁻), singlet oxygen (¹O₂), and hydrogen peroxide (H₂O₂), sodium azide for ¹O₂, and nitroblue tetrazolium for ·O₂⁻, did not decrease CL intensity. These observations indicated that ·OH mainly acted as an oxidant for the luminol CL reaction in the present ternary mixed solvent solution. The researchers have been interested in ·OH action in CL reaction and CL analysis [18] [19]. Generally, ·OH has a very short life time and is one of the strongest oxidants. It is interesting that ·OH was generated gradually from the present mixed solvent solution.

3.3. TRDF Observation

We confirmed the TRDF, where the organic solvent-rich major inner phase (pe-rylene, blue) and the water-rich minor outer phase (Eosin Y, green) were generated, in the capillary tube of the system using a fluorescence microscope-CCD camera. The obtained fluorescence photograph is shown in Figure 4; analytical conditions are also described in the figure legend. Phase separation multiphase

**Figure 3.** CL profiles obtained with the batch-type CL detector. To 0.1 µM luminol solution (2.5 mL) prepared with (a) water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate (3:8:4 volume ratio), (b) water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate (15:3:2 volume ratio), or (c) 10 mM phosphate buffer (pH 7.3) was added 15 mM hydrogen peroxide solution (50 µL) prepared with 10 mM phosphate buffer (pH 7.3).
Figure 4. Fluorescence photograph of TRDF observed with a fluorescence microscope-CCD camera system. (a) Non-TRDF at 20˚C and (b) TRDF at 5˚C. Conditions: Capillary tube, 140 cm (observation point: 110 cm from the inlet) of 75 μm i.d. fused silica; carrier, water (10 mM phosphate buffer, pH 7.2)-acetonitrile-ethyl acetate (3:8:4 volume ratio) containing 0.1 mM perylene and 1 mM Eosin Y, and flow rate, 1.0 μL·min⁻¹.

flow occurred with the ternary mixed solvent solution in the capillary tube at lower temperature through phase transformation; the annular flow or TRDF as shown in the figure could be observed under certain conditions.

3.4. Fluidic Analytical System with On-Line CL Detection

We developed various types CL detection system for flow-injection analysis, high-performance liquid chromatography, and capillary electrophoresis [20] [21]. However, all were off-line CL detection systems, where it was necessary to mix the eluate with reagent at the end of the tube or capillary. To our knowledge, there have been no reports regarding an on-capillary CL detection system. In this study, luminol CL reaction was shown to continue for several hours, maintaining sufficient CL with the ternary mixed solvent solution. The CL phenomenon could lead to the development of on-capillary CL detection systems without a mixing process at the end of the capillary, where analytes as inhibitors for CL reaction are detected as negative peaks from the baseline CL intensity.

The antioxidant reagents were subjected to the system with on-line CL detection. 2-Propanol and nordihydroguaiaretic acid were detected with negative peaks due to their antioxidant potential for luminol CL, but other antioxidants (superoxide radical anion, sodium azide, and nitroblue tetrazolium) were not different from baseline. The CL quenching tendency in the system was consistent with that observed with the batch-type CL detector mentioned above.

3.5. Chromatographic Information of Catenchin on Fluidic Analytical System

Catechin is a well-known antioxidant, and is one of the polyphenol derivatives,
where the hydroxyl groups work as antioxidant-sites, that react with radical species, such as hydroxyl radicals. Eight catechins were analyzed with the system, and all showed the negative peaks. The catechin structures were classified according to the number of hydroxyl groups containing diol and triol structures. The relative CL peak areas as negative peaks are summarized with the numbers of hydroxyl groups in Table 1. As shown here, the negative CL peak areas increased with increasing number of hydroxyl groups.

The elution times of catechin became later with increasing hydrophilicity expressed as log P (octanol/water partition coefficient) [22] and log K (distribution coefficient) (Table 2). The value of log P acts as an index of hydrophobicity. The value of K was examined as a distribution coefficient of upper phase (organic solvent-rich) and lower phase (water-rich) for each catechin that was obtained through phase transformation of the ternary mixed solvent solution in a batch vessel. The values of both increased with increasing hydrophobicity of the compounds.

In the present system, the organic solvent-rich major inner phase acts as a mobile phase, while the water-rich minor outer phase acts as a pseudostationary phase. That is, as the outer phase moves very slowly according to parabolic flow

Table 1. Relative CL peak areas as negative peaks for catechins with diol and/or triol hydroxyl groups estimated based on 100 for epigallocatechingallate.

<table>
<thead>
<tr>
<th>Antioxidants</th>
<th>Hydroxyl group</th>
<th>Relative CL peak area</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Diol</td>
<td>Triol</td>
</tr>
<tr>
<td>Catechin</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Epicatechin</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Gallocatechin</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Epigallocatechin</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Catechin gallate</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Epicatechin gallate</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Gallocatechin gallate</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Epigallocatechin gallate</td>
<td>0</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 2. Detection times of CL peaks for catechins with values of logP and logK.

<table>
<thead>
<tr>
<th>Antioxidants</th>
<th>LogP</th>
<th>LogK</th>
<th>Detection Time/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catechin</td>
<td>0.31</td>
<td>−0.11</td>
<td>6.2</td>
</tr>
<tr>
<td>Epicatechin</td>
<td>0.11</td>
<td>−0.17</td>
<td>6.2</td>
</tr>
<tr>
<td>Gallocatechin</td>
<td>−0.32</td>
<td>−0.44</td>
<td>6.3</td>
</tr>
<tr>
<td>Epigallocatechin</td>
<td>−0.50</td>
<td>−0.55</td>
<td>6.5</td>
</tr>
<tr>
<td>Catechin gallate</td>
<td>1.55</td>
<td>1.03</td>
<td>5.5</td>
</tr>
<tr>
<td>Epicatechin gallate</td>
<td>1.06</td>
<td>0.88</td>
<td>5.7</td>
</tr>
<tr>
<td>Gallocatechin gallate</td>
<td>0.92</td>
<td>0.79</td>
<td>5.8</td>
</tr>
<tr>
<td>Epigallocatechin gallate</td>
<td>0.39</td>
<td>0.47</td>
<td>6</td>
</tr>
</tbody>
</table>
under laminar flow conditions, the outer phase acts as a pseudohydrophilic-stationary phase on chromatographic separation mode. The hydrophilic compounds were distributed into the outer phase or pseudohydrophilic-stationary phase, and then eluted slowly, as shown in Table 2. Figure 5 shows the separation of catechingallate and epigallocatechin with the system. Catechingallate had the largest values of logP and logK, and epigallocatechin had the smallest values. Naturally, catechingallate eluted earlier than epigallocatechin, as shown in Figure 5. They were separated and detected with a base-line separation. The TRDF has been reported in a microspace, such as a capillary tube and a microchannel on a microchip, with various types of two-phase separation mixed solutions [14] [15]. The present system has a potential as chromatographic instrument, which was described as TRDC in the introduction section. The system will improved to separate more several compounds chromatographically in the future.

3.6. Catechin Analysis

We examined catechin (catechin mixture from green tea) using the present system. The most common species of catechin included in green tea is epigallocatechingallate, which usually accounts for more than 40% of the total amount of catechin. Although the catechin mixture from green tea was not separated by the present system under the conditions used, a negative CL peak was observed on the flow profile at around 7 minutes.

We attempted to estimate the total amounts of catechin in commercial green tea beverages using the present system. First, the relationship between the catechin concentration of the above catechin mixture from green tea and the percentage

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**Figure 5.** Flow profile of a mixture of catechingallate and epigallocatechin obtained by the microfluidic analytical system with on-line CL detection. Conditions: Capillary tube, 140 cm (effective length: 110 cm) of 75 μm i.d. fused silica; carrier, water (10 mM phosphate buffer, pH 7.3)-acetonitrile-ethyl acetate (3:8:4 volume ratio) containing 60 μM luminol and 2 mM hydrogen peroxide; sample injection, 20 cm height (gravity) × 15 s; flow rate, 1.0 μL·min⁻¹; tube temperature, 15°C; and catechin concentration, 250 μM each.
inhibition due to the negative CL peak was examined. The molecular weight of epigallocatechingallate (458.37) was used as that of the mixture. The negative CL peak responded over the range of 0.05 - 1 mM catechin with a detection limit of 0.05 mM (S/N = 3). A linear relationship was observed for the concentration range of 0.05 - 1 mM (correlation coefficient, 0.985). Tenfold-diluted commercial green tea beverage samples were analyzed using this system. Negative CL peaks were also observed on the CL profile at a detection time of ca. 7 minutes. Using the above-mentioned calibration curve, the commercial green teas, A, B, and C, were found to include catechin at concentrations of 363, 186, and 116 μM, (average values from 4 - 7 measurements), respectively. The data obtained with the present method were consistent with the manufacturers’ reported catechin concentrations for A, B, and C (337, 175, and 109 μM, respectively).

4. Conclusions

Microfluidic analytical system was developed with CL detection based on TRDF from phase separation multiphase flow. The simple microfluidic system consisted of single delivery line with on-capillary CL detection. The ternary mixed solvent solution of water-acetonitrile-ethyl acetate (3:8:4 volume ratio) was used as the carrier solution, because it was shown to act as an enhancer of the luminol-hydrogen peroxide CL reaction. The antioxidants for hydroxyl radicals, 1,2-propanol and nordihydroguaiaretic acid, quenched the CL intensity. Eight catechins that acted as antioxidants for hydroxyl radicals were detected as negative CL peaks due to their antioxidant potential with different detection times on the flow profile. The system could estimate the amounts of catechins in commercially available green tea.

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