The Utilization of Triacontyl-Bonded Silica Coated with Imidazolium Ions for Capillary Ion Chromatographic Determination of Inorganic Anions

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Abstract
This paper describes the separation and detection of five inorganic anions (I-, BrO3-, NO2-, Br-, and NO3-) by capillary ion chromatography using 1-alkyl-3-methylimidazolium-coated columns and UV detection at 210 nm. The salts of imidazolium (IM) ions (C2-, C4-, C6-, C8-, C10-, and C18-IMs) are a family of ionic liquids and were newly examined as permanent coating reagents on a triacontyl (C30)-bonded silica reversed phase column. When 40 mM NaCl aqueous solution was used as an eluent, the elution time of these anions was increased with the length of alkyl chain between C2- and C10-IMs, but turned to decrease with C18-IM. Good and faster separations were attained using a C18-IM-coated column with 150 mM NaCl in 25%(v/v) acetonitrile-75%(v/v) water. The method was successfully applied to the determination of nitrite, bromide, and nitrate in seawater and river water.

Keywords: Ion chromatography; Quaternary ammonium; Water analysis; Ionic liquid; Inorganic anions

1. Introduction
Since its advent in 1975, ion chromatography using a separation column of low ion exchange capacity in conjunction with a suppressor column to remove the background conductance of the eluent has been the dominant method for the determination of inorganic anions. To eliminate the suppressor column, anion exchange resins with a sufficiently lower exchange capacity (ca. 0.005-0.10 meq/g) have been developed and used with a very dilute eluent [1].

Apart from such fixed-site ion-exchange materials, anion exchange stationary phases can be prepared by permanently coating a variety of stationary phases with quaternary ammonium salts. The majority of permanent coating was performed on octadecyl silica-based reversed-phase materials (particles [2-23] and monoliths [24-28]). Jiang et al. utilized conventional silica columns to modify the particle surface with cetyltrimethylammonium ions [29]. In addition, polystyrene-divinylbenzene polymers [2-6,30,31] and graphitized carbon [23,32-35] were utilized. The quaternary ammonium salts included the salts of different tetraalkylammonium [9-18,21,24-27,29,30,32-34], n-cetylpyridinium [2-5,8,18,19,27,30,34], cationic dyes [6,7], and polyethyleimine [36]. A nonionic polymer surfactant was infrequently used for permanent coating of reversed-phase materials [37].

The advantages of using such permanently coated columns are the ability to vary the ion exchange capacity and the attainment of high column efficiency [29]. In addition, ion chromatography of anions can be practiced in laboratories where only conventional HPLC systems are available.

In this study, triacontyl (C30) silica reversed-phases were permanently coated with 1-alkyl-3-methylimidazolium ion and examined as a stationary phase for ion chromatography of inorganic anions. The salts of imidazolium (IM) ion are a member of the family of materials known as ionic liquids, which have recently gained wide popularity in almost all fields of chemistry, including synthesis, catalysis and electrochemistry. In the field of separation, ionic liquids have been used as stationary phases for GC and HPLC, eluent additives for HPLC and CE, and capillary-wall coating
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materials for CE (see a review [38] regarding this issue). An IM salt was covalently immobilized on silica particles and used for separation of inorganic and organic anions [39-42]. IM ionic liquids were also used as the mobile phase additives in reversed phase liquid chromatography for analysis of anions [43-45]. In the present study, the preparation of IM-based stationary phases was accomplished by a permanent coating procedure; its simplicity is a distinct advantage over the synthetic methods.

2. Experimental

2.1. Reagents

Acetonitrile was of HPLC grade from Merck (Darmstadt, Germany). Sodium chloride, used as an eluent additive, and chemicals used in the preparation of standard sample solutions were of analytical reagent grade and were purchased from FUJIFILM Wako Pure Chemical Corporation (Osaka, Japan) and Kanto Chemical (Tokyo, Japan). All eluents and standard mixture solutions were prepared using deionized water which was produced by a Model WG-23 water purification system (Yamato Scientific, Tokyo, Japan), and prior to use, filtrated with a 0.45-μm membrane filter. Stock solutions of anions were prepared at a concentration of 100 mM and diluted or mixed for calibration purposes. All standard solutions were stored in polyethylene containers to maintain the concentration.

Tetrafluoroborate salts of 1-ethyl-3-methylimidazolium (C2-IM), 1-butyl-3-methylimidazolium (C4-IM), and 1-methyl-3-octylimidazolium (C8-IM) were obtained from FUJIFILM Wako Pure Chemical Corporation. 1-Decyl-3-methyl imidazolium tetrafluoroborate (C10-IM) was obtained from Tokyo Chemical Industry (Tokyo, Japan). 1-Hexyl-3-methyl imidazolium tetrafluoroborate (C6-IM) was purchased from Sigma-Aldrich Japan (Tokyo, Japan). 1-Methyl-3-Octyl decylimidazolium (C18-IM) hexafluorophosphate was obtained from IoLiTec (Denzlingen, Germany). These imidazolium salts were used for preparation of the coating solutions as received.

2.2. Apparatus

Eluents were delivered at 4.0 μL/min by a Model 8301 micro-feeder (L. TEX Co., Tokyo, Japan) equipped with a 0.5-μL gas-tight syringe. Sample injections were made by a Model M435 micro-injection valve (internal dead volume, 50 nL; Upchurch Scientific, Oak Harbor, WA, USA) equipped with a fused silica capillary loop of 50-μm i.d. and 76 mm length, providing the sample loading of 0.20 μL. Detection was performed at 210 nm with a Model 870 UV detector (Jasco, Tokyo, Japan) equipped with a flow cell made of a 100-μm i.d. fused silica capillary. The detection window was located at 50 mm from the inlet end of the flow cell capillary which was directly connected to a separation column. The data were acquired by using a Chromato Monitor (Nippon Filcon, Tokyo, Japan). A Model VSR-50R ultrasonic bath (As One, Osaka, Japan) and a Model FB-4000 centrifuge (Kurabo, Osaka, Japan) were used for the preparation of real samples.

2.3. Column preparation and coating procedure

Columns were made of a piece of fused silica tubing of 0.32 mm i.d., 0.45 mm o.d., and 10 cm length, obtained from GL Sciences (Tokyo, Japan). The columns were slurry-packed with Develosil C-30-UG-5 (5-μm particle diameter, monomeric C30-silica) obtained from Nōmura Chemical (Seto, Japan). The stationary phase was coated by passing a 10 mM solution of an IM salt at a flow rate of 4.0 μL/min for 50 min; the coating solution was prepared in pure water except for the C18-IM solution which was prepared from 50% (v/v) acetonitrile in water. The columns were then conditioned by passing an appropriate solution.

2.4. Estimation of the amount of the immobilized C18-IM on the column

The amount of the C18-IM immobilized on a triacetylated silica reversed phase column was roughly estimated in the following manner. The coated column was flushed with air and then pure acetonitrile was pumped through the coated column at 2.0 μL/min until acetonitrile reached the outlet end of the column. After the column was equilibrated as such for 30 min, the coated column was flushed with acetonitrile at 16 μL/min for 33 min: the complete release of the C18-IM immobilized on the triacetylated silica from the column was confirmed by the descending absorption followed by the level-off of absorption at 210 nm. The column effluent (530 μL volume in total) was collected in a 0.6 mL microcentrifuge tube. Then the column was demounted from the chromatographic system and the micro-injection valve was connected directly to the flow cell of the UV detector, followed by directing a 0.20 μL portion of the collected solution with a flow of acetonitrile at 4.0 μL/min to the detector cell where absorbance was monitored at 210 nm. In order to determine the concentration of the C18-IM in the collected solution, a calibration curve (plots of peak area versus C18-IM concentration) was established using a set of C18-IM solutions of known concentration with the same experimental set-up. As a result, the concentration of C18-IM was estimated to be 1.30 mg/L. Subsequently, the packing material was pushed out of the column and dried well before it was weighed; the weight of the packing was 5.82 mg. Finally, it was calculated that the capacity of the coated column would be 0.35 meq/g.

2.5. Preparation of real samples

Surface seawater was collected on the coast of
Hamamatsu facing the Pacific Ocean and subjected to the ion chromatography analysis after filtration through a 0.45-μm membrane filter. River water was collected from the Magome River that flows through the city of Hamamatsu. The river water was centrifuged at 12000-rpm for 15 min and then filtrated with a 0.45-μm membrane filter.

3. Results and discussion
3.1. Coating of C30-silica with IMs

It is not the first time to use C30-silica for permanent coating of quaternary ammonium ion: Rong et al. used hexadimethrine bromide (cationic polymer) coated on C30 silica particles for determination of iodine in seawater [46]. In this study, ten mM solutions of 1-alkyl-3-methyl imidazoliums possessing various alkyl chain lengths (C2-, C4-, C6-, C8-, and C10-IMs) were prepared in pure water, while that of C18-IM was prepared in a 50:50(v/v) water-acetonitrile mixture. The coating solutions were pumped through a C30-silica packed column at 4.0 μL/min for a period of 50 min, with the column effluent directed through the UV detector set at 210 nm. The column was then flushed with 40 mM NaCl in water until the baseline was stabilized. Sample injection was made at this point. The columns coated with C2-, C4-, and C6-IMs provided stable baseline within 20 min, while the C8- and C10-IM-coated columns took approximately 1.5 and 2.0 h, respectively, before the baseline was stabilized. The C18-IM-coated column gave a stable baseline within 10 min. Such differences in time required for equilibration may be due to the differences in the amounts of individual IMs accumulated on the bonded-phase and due to the difference of solubilities of individual IMs in the NaCl aqueous solution.

The first chromatographic evaluation was performed using a test mixture of inorganic anions composed of iodate, bromate, nitrite, bromide and nitrate. As can be seen in Fig.1, these anions are more strongly retained on the column as the alkyl chain becomes longer from ethyl to decyl in the IM, but the elution times of the anions decreased on the octadecyl-IM column as compared with that on the decyl-IM column. The complete separation of individual anions was accomplished on the C10-IM-coated column, whereas bromate and nitrite coeluted on the C18-IM-coated column. What was worse, we found that retention of each anion changed with the running time. On the C10-IM-coated column the elution time of every anion initially increased and after a certain point of time, turned to decline (Fig. 2(A)). The decrease in the retention of anions on the C10-IM coated column may signify that the IM was released from the reversed-phase material in the column. On the other hand, the retention of anions continued to increase on the C18-IM coated column (Fig. 2(B)).

![Fig. 1. Separations of inorganic anions on the C30-silica columns coated with imidazoliums (IMs) possessing different alkyl chain lengths. (A) C2-IM, (B) C4-IM, (C) C6-IM, (D) C8-IM, (E) C10-IM, (F) C18-IM. Peaks: 1, iodate; 2, bromate; 3, nitrite; 4, bromide; 5, nitrate. Eluent: 40 mM NaCl aqueous solution at 4.0 μL min⁻¹. Sample injection volume: 0.20 μL. Detection: UV at 210 nm. Analyte concentration: 0.5 mM.](image1)

![Fig. 2. Variation of the elution times of anions on (A) C10-IM and (B) C18-IM coated reversed-phase columns with running time. Analytes: (1) iodate; (2) bromate; (3) nitrite; (4) bromide; (5) nitrate. Chromatographic conditions as in Fig.1.](image2)
According to the paper by Li et al. [47], it follows that water promotes the ionic dissociation of IMs significantly compared with acetonitrile because of the high dielectric constant and proton-donating ability of water. It seems therefore that as acetonitrile surrounding the IM sorbed on the reversed-phase material was replaced with water, the effective charge on the IM available to interact with the analytes increased, thereby increasing the retention.

3.2. Effect of addition of acetonitrile into the eluent

Acetonitrile was added into the eluent containing 40 mM NaCl in an attempt to keep the analyte retention under control. Anion separations were performed using an eluent containing 40 mM NaCl in 25% (v/v) acetonitrile-75% (v/v) water on the C18-IM-coated column (Fig. 3(B)), where individual anions eluted earlier compared to when 40 mM NaCl in water (Fig. 3(A)) was used as the eluent. Interestingly, bromate and nitrite ions were successfully separated with this particular eluent. Although at the present stage it is difficult to clarify the effect of added acetonitrile on the their elution behavior, it is known that various organic solvents can exert a variety of effects on the analyte ions [48].

3.3. Speed up of conditioning

When the C18-IM-coated column was conditioned with an eluent containing 40 mM NaCl in 25% (v/v) acetonitrile-75% (v/v) water, it took more than 20 h until each anion gave a constant elution time. In order to speed up the conditioning, 1M NaCl aqueous solution was passed through each column for a short period of time. Fig. 4 shows the change in the elution time of the last peak (nitrate) on the columns which were conditioned over the different periods of time. It is obvious that almost constant elution times of anions are achieved by passing 1M NaCl aqueous solution through the column for about one hour.

![Fig. 3. Effect of acetonitrile in eluent on the separation of anions.](image)

![Fig. 4. Effect of time periods for conditioning on the elution time of nitrate.](image)

![Fig. 5. Separation of inorganic anions on a C18-IM-coated column with 40 mM (A) and 150 mM (B) NaCl in 25% (v/v) acetonitrile-75% (v/v) water.](image)
an eluent of 40 mM NaCl in 25%(v/v) acetonitrile-75% (v/v) water is also shown. It is seen that by use of the higher-salinity eluent, the separation can be completed within 7 min. The elution order of five anions is the same as that obtained with silica columns modified with cetyltrimethylammonium ion [29]. The reproducibility of analysis for the inorganic anions is demonstrated in Fig. 6, where ion chromatograms obtained every hour are presented. It appears that the C18-IM is strongly retained on the reversed phase material through hydrophobic interaction.

3.5. Effect of salinity in the eluent on the retention factor

The influence of salinity on the retention of the analytes was investigated using eluents containing NaCl within the range of 40 mM to 300 mM in 25%(v/v) acetonitrile-75%(v/v) water. Increasing the salinity tended to decrease the retention factor ($k$). As can be seen in Fig. 7, there is a linear relationship between the logarithm of the retention factor ($\log k$) and the logarithm of the NaCl concentration in the eluent. The slopes for the analytes were $-0.92$, $-1.08$, $-1.05$, $-1.04$ and $-1.06$ for iodate, bromate, nitrite, bromide, and nitrate, respectively. Theoretically, the value of slope is determined by the ratio of the valence of analyte ion to the valence of the chloride ion in the eluent. Therefore, the slopes should be equal to $-1$: the separation of the analytes is based on ion-exchange mechanism.

![Fig. 6. Long-term stability of a C18-IM-coated column after conditioned with 1M NaCl aqueous solution for 1h. Eluent: 150 mM NaCl in 25%(v/v) acetonitrile-75%(v/v) water at 4.0 $\mu$L min$^{-1}$. Sample injection volume: 0.20 $\mu$L. Detection: UV at 210 nm. Peak identifications as in Fig. 5.](image)

![Fig. 7. The logarithm of the retention factor versus the logarithm of NaCl concentration (mM). Eluent: 40 mM to 300 mM NaCl in 25%(v/v) acetonitrile-75%(v/v) water. Separation and detection conditions as in Fig. 6. Peak identification as in Fig. 5.](image)

3.6. Analytical performance

Reproducibility with respect to retention time, peak area and peak height for different anions present in a single mixture was studied under the established operating conditions, which are summarized in Table 1. The results obtained (Table 2) show a maximum RSD of 1.10% for the retention time, 4.21% for the peak area and 3.38% for the peak height. As compared with conventional silica column modified with cetyltrimethylammonium bromide (CTAB) [29], the RSDs in retention times obtained in the present method are smaller except for iodate ion, while the RSD values in peak area and peak height are almost comparable. Furthermore the analysis time achieved in our method is shorter than that obtained with the CTAB-modified silica column.

The limits of detection (LOD) for the anions detected at 210 nm ranged from 3 to 5 mM (from 0.14 to 0.82 ppm), with the exception of bromate that more weakly absorbs at this particular wavelength compared with other anions. The LOD values achieved with the present method are worse than those obtained with the CTAB-modified silica column coupled to a conventional UV detector set at 210 nm [29]. A good linear relationship between analyte concentration and peak area was obtained over the concentration range studied (from 62.5 $\mu$M to 1.00 mM) as revealed by the values of the square of the correlation coefficient ($R^2$) ranging from 0.997 to 0.999.

The stability of the IM-coated stationary phase was examined. The retention time ($t_R$) remained insignificantly changed over a period three months: the $t_R$ of iodate ranged from 1.88 to 1.98 min.
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3.7. Analysis of real samples

Seawater and river water samples were collected, filtered through a 0.45-μm membrane filter, and without dilution or concentration, injected into the C18-IM-coated column. The resultant chromatograms are shown in Fig. 8. Nitrite and nitrate are detected in both samples without any interference by matrix ions. A large peak of bromide is observed in the seawater sample. In general, column capacity must be considered when real samples are analyzed using such a small column. It may be noticed that the retention times of these anions for the real samples are the same as those for the standard sample (Fig. 8(A)): this illustrates that the column capacity was sufficiently high to cope with the different sample matrix. The estimated capacity of the C18-IM coated column (0.35 meq/g) is smaller than that of classical ion-exchange resins (3-5 meq/g), but higher than that of low-capacity ion-exchangers (0.005-0.10 meq/g) which have been popular in recent years. The analytical results are summarized in Table 3. Bromide in seawater was quantified at a level of 43.8 mg/L which falls within the range of the reported values [49]. It is well known that the concentration levels of nitrite and nitrate in environmental water can vary considerably depending on the extent of human-caused or natural eutrophication, and in fact, a wide range of nitrite and nitrate concentration values are indicated in the literatures [11,13,16,26]. Therefore, the concentration level of nitrite and nitrate obtained for our water samples appears to be quite probable.

![Fig. 8.](image)

**Table 1.** Operating chromatographic conditions.

| Column | Devosil C30-UG-5 column (0.32 mm i.d., 10 cm length) coated with 200 μL of 10 mM C18-IM in 50:50, v/v water/acetonitrile |
|---|---|
| Conditioning | 1M NaCl at 4.0 μL/min for 1h |
| Detection | Direct absorption at 210 nm |
| Column temperature | 25°C |
| Eluent | 150 mM NaCl in 25%(v/v) acetonitrile-75%(v/v) water |
| Flow rate | 4.0 μL/min |
| Sample injection volume | 0.20 μL |

**Table 2.** The RSDs of the retention time (t_r), peak area and peak height, LOD at S/N=3 of anions, and the coefficient of determination (R^2) of the calibration curves.

| Analyte | RSD, % (n=30) | LOD μM (ppm) | R^2 | t_r min |
|---|---|---|---|---|
| Iodide | 1.10 | 3.35 | 1.20 | 5 (0.82) | 0.9972 | 1.96 |
| Bromate | 1.05 | 4.06 | 3.30 | 12(1.54) | 0.9990 | 3.24 |
| Nitrite | 1.00 | 2.76 | 2.59 | 3 (0.14) | 0.9992 | 3.59 |
| Bromide | 0.90 | 4.21 | 3.38 | 4 (0.33) | 0.9991 | 5.05 |
| Nitrate | 0.87 | 2.59 | 2.54 | 3 (0.16) | 0.9987 | 6.56 |

**Table 3.** Analytical results of nitrite, bromide and nitrate in river water and seawater.

| Analyte | Seawater | River water |
|---|---|---|
| | Concentration, RSD, n=5 | Concentration, RSD, n=5 |
| Nitrite | 5.5 (0.25) | 3.69 | 5.8 (0.27) | 2.47 |
| Bromide | 548 (43.8) | 0.81 | ND^a | - |
| Nitrate | 7.8 (0.48) | 2.73 | 82.4 (5.1) | 1.26 |

^a Not detected. Concentrations were determined based on the peak area.
4. Conclusion
Moderately high-capacity anion-exchange columns were developed by sorption of IM on reversed-phase stationary phases. A C30-silica packed capillary column which was permanently coated with 10 mM C18-IM in a (50:50, v/v) water-acetonitrile mixture allowed separation and detection of five inorganic anions in < 7 min, using an eluent of 150 mM NaCl in 25%(v/v) acetonitrile-75%(v/v) water and UV detection at 210 nm. The proposed method seems to be more convenient and effective than other methods [39-42] for preparing columns with moderately high ion exchange capacity.

To illustrate how the present method has real application potential, the method was applied to the determination of nitrite and nitrate in both a local river water and seawater. Bromide was determined in seawater. Due to its moderately high exchange capacity, the IM-coated column can be applied to samples of considerable ionic strength such as seawater.

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