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1. Introduction

Room temperature ionic liquids (ILs) have been paid attention as environmentally benign media, because they have attractive properties such as thermal stability, nonflammability, high ionic conductivity, and wide electrochemical potential windows (Earle & Seddon, 2000; Rogers et al., 2000; Wasserscheid & Welton, 2003). In the nuclear industry field, ILs are expected to be applied as media for reprocessing of spent nuclear fuels and treatment of radioactive wastes contaminated with radioactive nuclides (Bladley et al., 2002; Rogers et al., 2002; Giridhar et al., 2006; Cocalia et al., 2006; Giridhar et al., 2007; Binnemans, 2007).

In this chapter, our feasibility studies on applications of ILs as the media of pyro-reprocessing processes and the treatment method of radioactive wastes contaminated with uranium will be introduced.

2. Investigation on application of ILs as media of the pyro-reprocessing processes

We studied electrochemical properties of uranyl species in 1-butyl-3-methylimidazolium (BMI) based ILs (BMICl, BMIBF₄, and BMINfO (NfO = nonafluorobutanesulfonate)) to examine their feasibility as alternatives to conventional molten salts as media for pyro-reprocessing processes for spent nuclear fuels. BMICl (Kanto Chemical Co., Inc.) was used without further purification and BMIBF₄ (Kanto) was purified by using activated carbon. BMINfO was synthesized as follows: 1-Butyl-3-methylimidazole was dissolved into tetrahydrofuran (THF) and stirred vigorously. After that, 1-bromo butane was dropped slowly with a dropping funnel, and the resulting solution was refluxed. After refluxing, THF phase was separated and stirred with ethylacetate (EA). Crude 1-butyl-3-methylimidazolium bromide (BMIBr) was obtained by removing THF and EA in vacuo. The crude BMIBr was dissolved into distilled water and
stirred vigorously. To this solution, KNfO was added. The resulting solution was refluxed with stirring at 70 °C. The BMINfO phase was separated from aqueous one and mixed with activated carbon for removing organic impurities. After filtration, the filtrate was mixed with the distilled water for striping inorganic impurities. In order to remove water and volatile impurities, all ILs used were kept for more than 3 h under reduced pressure at 120 °C.

Sample solutions for electrochemical experiments were prepared by dissolving Cs₂UO₂Cl₄ or UO₂Cl₂·nH₂O (n = 1~3) into ILs. Cs₂UO₂Cl₄ and UO₂Cl₂·nH₂O (n = 1 ~ 3) were synthesized according to the reported procedures (Cordfunke, 1969; Denning et al., 1979). Cyclic voltammograms (CV) of sample solutions controlled at appropriate temperatures were measured by using an electrochemical analyzer (BAS, ALS model 660B) in glove box under an Ar atmosphere. A glassy carbon and a Pt wire were used as a working electrode and a counter electrode, respectively. As a reference electrode, an Ag/AgCl electrode was used and connected with a cyclic voltammetry cell by a liquid junction filled with BMIBF₄ or BMINfO. All potentials reported here are vs. Ag/AgCl. In the CV measurements, potential was swept to cathodic direction initially.

2.1 Electrochemical study on uranyl chloride in BMICl

Figure 1(a) shows the UV-visible absorption spectrum of the solution prepared by dissolving Cs₂UO₂Cl₄ into BMICl at 80°C. This absorption spectrum is found to exhibit remarkable vibrational fine structure, which is similar to that of [UO₂Cl₄]²⁻ in AlCl₃/EMIC (EMIC = 1-ethyl-3-methylimidazolium chloride) (Dai et al., 1997), BMITf₂N, MeBuNTf₂N, and C₅minTf₂N (Tf₂N = bis(trifluoromethanesulfonyl)imide, MeBuN = tri-n-butylmethylammonium, C₅min = 1-hexyl-3-methylimidazolium) (Sornein et al., 2006; Nockemann et al., 2007). Nockemann et al. have reported that the fine structure is typical for the [UO₂Cl₄]²⁻ with D₅₀ coordination symmetry (Nockemann et al., 2007). The molar absorption coefficient (ε) at 429.8 nm (maximum peak) is 16.3 M⁻¹ cm⁻¹ (M = mol dm⁻³), and is almost same as those reported previously (Sornein et al., 2006; Nockemann et al., 2007). We also measured the UV-visible absorption spectrum of the solution obtained by dissolving UO₂Cl₂·nH₂O in BMICl at 80 °C (see Fig. 1(b)), where the concentration of uranium was determined by ICP-AES. A similar absorption spectrum to that in Fig. 1(a) was observed. The ε value of maximum peak at 428.6 nm was 16.3 M⁻¹ cm⁻¹. These results indicate that the uranyl species in solutions prepared by dissolving Cs₂UO₂Cl₄ or UO₂Cl₂·nH₂O into BMICl is [UO₂Cl₄]²⁻.

Slight differences in the ε values and the wavelength of peak maxima in Fig. 1(a) and (b) might be due to the effects of Cs⁺ ions in BMICl system obtained by dissolving Cs₂UO₂Cl₄ and residual water in the sample solutions.

Based on spectrophotometric data, to examine the electrochemical behavior of [UO₂Cl₄]²⁻ in BMICl, CVs of the sample solutions prepared by dissolving Cs₂UO₂Cl₄ or UO₂Cl₂·nH₂O into BMICl (abbreviated as Cs₂UO₂Cl₄/BMICl system and UO₂Cl₂·nH₂O/BMICl system) were measured at 80 °C in the potential range of -1.0 ~ -0.4 V at various scan rates (ν = 10 ~ 50 mV s⁻¹). A typical result for the Cs₂UO₂Cl₄/BMICl system is shown in Fig. 2 (a). As seen from this figure, two peaks corresponding to one redox couple were observed around -0.72 (Epc) and -0.65 V (Epa). The potential differences between two peaks (ΔEpa) are 75 and 81 mV at 10 and 50 mV s⁻¹, respectively, and close to the theoretical value (67 mV) for the reversible one electron transfer reaction at 80 °C. Furthermore, the values of (Epc + Epa)/2 is constant, -0.687 ± 0.005 V, regardless of ν (see Table 1). Similar results were also obtained from the CVs

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for the UO₂Cl₂·nH₂O/BMICl system at 80 °C as shown in Fig. 2 (b), that is, one redox couple was observed around -0.73 (Epc) and -0.66 V (Epa), the ΔEp values are 69 mV at 10 mV/s and 77 mV at 50 mV/s, the values of (Epc + Epa)/2 is constant, -0.693 ± 0.001 V (see Table 1). From these results, it is suggested that [UO₂Cl₄]²⁻ in BMICl is reduced to [UO₂Cl₂]³⁻ quasi-reversibly and that the formal redox potential (E⁰) is -0.690 V in the present system. Hence, it should be concluded that BMICl is not applicable as the medium of the pyro-reprocessing process, because the uranyl species in BMICl are not reduced to UO₂.

Fig. 1. UV-visible absorption spectra of the solutions prepared by dissolving uranyl chloride complexes into BMICl at 80 °C. (a): Complex = Cs₂UO₂Cl₄ [UO₂²⁺] = 1.47 x 10⁻² M. (b): Complex = UO₂Cl₂·nH₂O [UO₂²⁺] = 1.47 x 10⁻² M.
Fig. 2. Cyclic voltammograms of the solutions prepared by dissolving uranyl chloride complexes into BMICl measured in the potential range from -0.1 to -0.4 V at different scan rates ($v = 10 \text{ to } 50 \text{ mV s}^{-1}$). (a): Complex = Cs$_2$UO$_2$Cl$_4$; [UO$_2^{2+}$] = 1.47 x 10$^{-2}$ M. (b): Complex = UO$_2$Cl$_2$·nH$_2$O; [UO$_2^{2+}$] = 1.47 x 10$^{-2}$ M. Temp. = 80 °C. Initial scan direction: cathodic.
2.2 Electrochemical study on uranyl chloride in BMIBF₄

Adding UO₂Cl₂-nH₂O to BMIBF₄ precipitates were formed. Hence, supernatant solutions containing UO₂⁺ were used for electrochemical experiments. The UV-visible absorption spectrum of supernatant is shown in Fig. 3, and is found to be similar to those of [UO₂Cl₂]⁻ in Fig. 1. This result suggests that the uranyl species in the supernatant exists as [UO₂Cl₂]⁻.

The CV measurements were continuously repeated five times in the range of –1.0 ~ 1.0 V. The results are shown in Fig. 4. An irreversible reduction peak was observed around –0.7 V and gradually decreased with the repetition of the potential sweep. By wiping off the surface of working electrode, the reduction peak was appeared again. This suggests that the surface of the electrode is covered by insoluble film produced by redox reaction. Similar phenomenon was reported by Chagnes et al., that is, they observed the formation of a blocking film on the graphite electrode in CV measurements in BMIBF₄ (Chagnes et al., 2005).

Table 1. Cyclic voltammetric data for solutions prepared by dissolving Cs₂UO₂Cl₄ (a) and UO₂Cl₂-nH₂O (b) into BMICl

<table>
<thead>
<tr>
<th>System</th>
<th>ν / (mV·s⁻¹)</th>
<th>Epc / V</th>
<th>Ep a / V</th>
<th>ΔE / V</th>
<th>ipc / A</th>
<th>ipa / A</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a)</td>
<td>10</td>
<td>–0.729</td>
<td>–0.654</td>
<td>0.075</td>
<td>–7.21x10⁻²</td>
<td>6.59x10⁻²</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>–0.727</td>
<td>–0.653</td>
<td>0.074</td>
<td>–1.08x10⁻⁶</td>
<td>9.38x10⁻⁷</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>–0.725</td>
<td>–0.646</td>
<td>0.079</td>
<td>–1.32x10⁻⁶</td>
<td>1.13x10⁻⁶</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>–0.726</td>
<td>–0.647</td>
<td>0.079</td>
<td>–1.52x10⁻⁶</td>
<td>1.29x10⁻⁶</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>–0.723</td>
<td>–0.642</td>
<td>0.081</td>
<td>–1.68x10⁻⁶</td>
<td>1.42x10⁻⁶</td>
</tr>
<tr>
<td>(b)</td>
<td>10</td>
<td>–0.752</td>
<td>–0.682</td>
<td>0.070</td>
<td>–7.36x10⁻²</td>
<td>4.37x10⁻²</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>–0.752</td>
<td>–0.678</td>
<td>0.074</td>
<td>–1.06x10⁻⁶</td>
<td>5.95x10⁻⁷</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>–0.754</td>
<td>–0.679</td>
<td>0.075</td>
<td>–1.22x10⁻⁶</td>
<td>7.09x10⁻⁷</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>–0.759</td>
<td>–0.679</td>
<td>0.080</td>
<td>–1.43x10⁻⁶</td>
<td>8.15x10⁻⁷</td>
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<tr>
<td></td>
<td>50</td>
<td>–0.762</td>
<td>–0.676</td>
<td>0.086</td>
<td>–1.72x10⁻⁶</td>
<td>9.18x10⁻⁷</td>
</tr>
</tbody>
</table>

(a): [UO₂²⁺] = 1.47 x 10⁻² M. (b): [UO₂⁴⁺] = 1.47 x 10⁻² M

Fig. 3. UV-visible absorption spectrum of the supernatant solution prepared by adding UO₂Cl₂-nH₂O into BMIBF₄ at 80 °C. [UO₂²⁺] = 5.1x10⁻³ mol/kg.
Judging from the above results, BMIBF$_4$ is concluded to be not applicable as the medium of the pyro-reprocessing process.

2.3 Electrochemical study on uranyl chloride in BMINfO

Figure 5 shows the UV-visible absorption spectrum of the solution prepared by dissolving UO$_2$Cl$_2$·nH$_2$O into BMINfO at 80°C. As seen from this figure, the absorption spectrum is different from those of [UO$_2$Cl$_2$]$^{2-}$ shown in Fig. 1. Although we do no have exact data on the structure yet, it seems likely that the uranyl species in BMINfO is present as [UO$_2$Cl$_2$(NfO)$_n$]$^{n-}$.

Fig. 5. UV-visible absorption spectrum of the solution prepared by dissolving UO$_2$Cl$_2$·nH$_2$O into BMINfO at 80°C. [UO$_2^{2+}$] = 8.7 x 10$^{-3}$ mol/kg.
Figure 6 shows CVs of neat BMINfO and uranyl species in BMINfO. As seen from this figure, three irreversible reduction peaks (i, ii, iii) and a sharp oxidation one (iv) appear in the range of $-0.6 \sim -0.2$ V and around 0.85 V, respectively. It is known that a sharp oxidation peak (iv) is due to the oxidative dissolution of reduction products deposited on the electrode (Shirai et al., 1998). Therefore, the sharp oxidation peak at 0.85 V is considered to corresponds to oxidative dissolution of U(IV) compounds deposited on the electrode. Thus, the reduction peaks should be assigned to multistep reduction of U(VI) to U(IV) as follows.

i. $\text{U(VI)} + e \rightarrow \text{U(V)}$

ii. $\text{U(V)} + e \rightarrow \text{U(IV)}$

iii. $\text{U(VI)} + 2e \rightarrow \text{U(IV)}$

Fig. 6. Cyclic voltammograms of neat BMINfO and uranyl species in BMINfO ($80^{\circ}$C, $[\text{UO}_2^{2+}] = 8.7 \times 10^{-3}$ mol/kg, sweep rate = 50 mV/s).

2.4 Bulk electrolysis of uranyl species in BMINfO

Judging from the results of the CV measurements described above, it might be possible to recover UO$_2$ by electrochemical reduction of UO$_2^{2+}$ in BMINfO. Thus, bulk electrolysis of UO$_2^{2+}$ (0.3 mol·kg$^{-1}$) in BMINfO was carried out at $-1.0$ V by using cell for bulk electrolysis shown in Fig. 7. As a result, the deposits were produced on a carbon electrode as cathode and a part of such deposits was fallen to the bottom of the electrolysis cell. The photograph of recovered deposits is shown in Fig. 8. After the electrolysis, the IL on the carbon electrode was washed away with acetone and dichloromethane, and then the surface of the carbon electrode was analyzed by the scanning electron microscope (SEM) and the energy dispersive X-ray spectrometer (EDX). The micrograph and distribution of elements on the electrode surface are shown in Fig. 9. The distribution of carbon is attributable to the carbon electrode. Consequently, it was found that the deposits are uranium compounds including chlorine components such as uranium oxides and uranium oxychlorides.
2.5 Summary of electrochemical properties of uranyl species in BMI based ionic liquids

Electrochemical properties of uranyl species in BMICl, BMIBF₄, and BMINfO were examined by using cyclic voltammetry. And based on such investigations, the applicability of ILs as the media of pyro-reprocessing processes was also examined. The results are summarized as follows.

- In BMICl, the reversible redox couple was observed in CV. This suggests that the redox couple corresponds to the redox couple of UO₂²⁺/UO₂⁺ with one electron transfer.
- In BMIBF₄, an irreversible reduction peak was observed around −0.7 V and gradually decreased by the repetition of the potential sweep. This phenomenon is caused by the formation of insoluble reduction products on the surface of the electrode.

Fig. 7. Cell for bulk electrolysis.

Fig. 8. A photograph of the deposits recovered by electrochemical reduction of uranyl species in BMINfO.
In BMINfO, three irreversible reduction peaks and a sharp oxidation one were observed in the range of $-0.6 \sim -0.2$ V and around 0.85 V, respectively. This suggests that the redox reactions consist of the multi step reduction of U(VI) to U(IV) and the oxidative dissolution of U(IV) as reduction products.

Electrochemical reduction of uranyl species in BMINfO was performed by bulk electrolysis. As a result, deposits were observed on the cathodic electrode. From the SEM-EDX analyses, it was confirmed that the deposits are uranium compounds including chlorine components such as uranium oxides and uranium oxychlorides.

These results indicate that the uranyl species in IL can be recovered electrolytically as uranium compounds. Hence, from the electrochemical viewpoint it is expected that ILs can be used as media of pyro-reprocessing processes.
A new pyro-reprocessing method shown in Fig. 10 should be proposed. This method consists of three processes, i.e., first one is the dissolution of spent nuclear fuels using oxidant such as Cl₂, second one is the recovery of UO₂ by electrochemical reduction, and third one is the electrochemical deposition of UO₂/PuO₂ mixed oxide.

Fig. 10. Schematic diagram of proposed processes of the pyro-reprocessing by using IL as media

3. Investigation on application of ILs as electrolytic media for treating wastes contaminated with uranium

Most of metal and bed materials generated from uranium enrichment facilities or uranium refining and conversion plants are contaminated by uranium fluorides such as UF₄. These wastes are mainly classified as the medium-level wastes. Hence, it is desired to recover uranium as much as possible from such wastes. Moreover, if these wastes are decontaminated up to the clearance level, the resulting decontaminated materials should be reused. As one of effective decontamination methods of metal wastes, wet chemical decontamination processes using inorganic or organic acids have been developed (Ikeda et al., 2002; Enda et al., 2006). However, from such wet processes, a relatively large amount of secondary wastes should be generated with treating spent acid solutions, because base metal part of wastes is dissolved by acid with the dissolution of contaminated part. And also it is reported that uranium of spent adsorbents can be recovered by electrolysis in sodium chloride molten salt (Amamoto et al., 2005). However, this method must be performed under high temperature (672°C). Decontamination methods carried out under milder conditions must be preferable. Ionic liquids are expected to meet such demands. Hence, we investigated the solubility of UF₄ in ILs and the electrochemical properties of uranium species dissolved into ILs.

Uranium tetrafluoride (UF₄) was synthesized from yellow cake according to the reported method (Higgins et al., 1958). Synthesis of UF₄ was confirmed using a X-Ray diffractometer (Rigaku, RAD-rPC). Impurities in UF₄ were detected using an ICP-MS (Thermo electron Co., ELEMENT). Purity of UF₄ was 97%. BMI-Cl was used as an IL. Water containing BMI-Cl was removed by heating in vacuo. Water content in BMI-Cl after drying was determined to be 0.1 wt % using a Karl Fischer moisture content meter (Metrohm, 737 KF Coulometer).
3.1 Dissolution behavior of UF₄ powders

Dissolution experiments were carried out at 100°C under the atmosphere with stirring at 100 rpm. The UF₄ powders (0.1 g, 3.2×10⁻⁴ mol) were weighed accurately and dissolved in BMICl solution (1.0 ml) in a beaker. After dissolution, the residual solid phases were filtered off and the uranium concentrations in the filtrates were measured ICP-MS. Figure 11 shows a plot of dissolution ratios vs. time. As seen from this figure, the UF₄ powders do not dissolve in BMICl easily.

![Graph showing dissolution ratio vs. time](image)

**Fig. 11. A plot of dissolution ratios vs. time for the dissolution of UF₄ powders (0.1 g) in BMICl (1.0 ml) at 100 ºC.**

The color of the dissolution solution was green after about 1 h and the powders were completely dissolved after around 6 h. The color of BMICl solution after complete dissolution of UF₄ was yellowish green. However, its absorption spectrum did not show characteristic bands assigned to the U⁴⁺ species (Rodden, 1964). The yellowish green solution was further heated for 10 h under the atmosphere. As a result, the color of the solution changed from yellowish green to yellow. Figure 12 shows the absorption spectrum of the resulting solution and is similar to that of UO₂Cl₂⁺ shown in Fig. 1. The ε value at maximum peak of 422 nm is 13.1 M⁻¹ cm⁻¹ and almost the same as that (about 14 M⁻¹ cm⁻¹) at maximum peak of 429 nm reported (Sornein et al., 2006). This result indicates that the species generated with the dissolution of UF₄ powders are oxidized to uranyl(VI) by O₂ under the present conditions. The relative slow dissolution of UF₄ in BMICl should be due to that the oxidation process of U(IV) with O₂ is slow.

Wipff et al. have reported that in BMICl dissolving uranyl triflate (TfO⁻) or uranyl perchlorate, Cl⁻ ions interact with uranyl(VI) more strongly than ClO₄⁻ and TfO⁻, and that the uranyl(VI) species mainly exist as UO₂Cl₂⁻ (Gaillard et al., 2007). And also they have proposed from the results of molecular dynamics (MD) and quantum mechanical (QM) calculations that F⁻ ions coordinate to uranyl(VI) more strongly than Cl⁻(Gaillard et al., 2007; Chaumont & Wipff, 2005). Based on these reports, it is suggested that the uranyl(VI) species with the mixed ligands of F⁻ and Cl⁻ should be formed in the BMICl solution dissolving UF₄.
Fig. 12. UV-visible absorption spectrum of the solution prepared by dissolving UF₄ into BMICl ([UO₂²⁺] = 8.0 x 10⁻² M)

3.2 Electrochemistry of sample solutions prepared by dissolving UF₄ into BMICl

The cyclic voltammograms of the sample solutions prepared by dissolving UF₄ (0.52 g) into BMICl (30 ml) were measured at 80 °C in the potential range -2.0 – 0.95 V at 50 mV/s. In these experiments, a glassy carbon wire, a Pt wire, a Ag/AgCl electrode (BAS, RE-1B) with a liquid junction filled with BMICl were used as working, counter, and reference electrode, respectively. The result is shown in Fig. 13. As seen from this figure, one uncoupled reduction peak and one uncoupled oxidation peak are observed around -0.93 and 0.18 V, respectively, and the current value of the oxidation peak is smaller than that of the reduction peak. This result is different from that of Cs₂UO₂Cl₄ system mentioned in 2.1. In the BMICl system dissolving Cs₂UO₂Cl₄, the uranyl(VI) species were confirmed to be present as UO₂Cl₂⁺, and one quasi-reversible redox couple assigned as UO₂Cl₂⁺ + e⁻ = UO₂Cl₃⁻ was observed around -0.72 and -0.65 V. These support the above suggestion that the uranyl(VI) species with the mixed ligands of F⁻ and Cl⁻ are formed in the BMICl solution dissolving UF₄, and suggest that the reduction product of the uranyl(VI) complexes with the mixed ligands of F⁻ and Cl⁻ are less stable than that of UO₂Cl₂⁻, i.e., UO₂Cl₄²⁻. Sornein et al. have reported that the uncoupled reduction peak should correspond to the following reduction processes (Sornein et al., 2006).

\[
\text{Uranyl(VI)} + e^- \rightarrow \text{Uranyl(V)}
\]

\[
\text{Uranyl(V)} + e^- \rightarrow \text{UO}_2
\]

From these results, it is expected that the uranium component can be recovered electrolytically from the solutions generated in the decontamination treatments of the wastes contaminated with UF₄ in BMICl.
Fig. 13. Cyclic voltammograms of the solution prepared by dissolving UF₄ in BMICl ([UO₂²⁺] = 8.0 x 10⁻² M) and neat BMICl at 80 °C. Initial scan direction: cathodic.

3.3 Application to decontamination of the steel wastes contaminated with uranium

Samples of steel waste were prepared from the dismantled carbon steel cylinders which had been used for storing UF₆ (see Fig. 14). The chemical forms of uranium species adhered on the steel wastes were confirmed to be UF₄ by measuring XRD. Chemical forms of iron species on surfaces of contaminated steel wastes were confirmed to be FeF₃ and Fe₂O₃ by X-ray Photoelectron Spectroscopy (JEOL, JPS-9000MC) using Mg Kα radiation of 1253.6 eV.

Fig. 14. A photograph of sample of steel waste
The contaminated steel wastes were cut into the quarter sector (28mm Φ × 6 mm thick, central angle 90°), and soaked in BMICl (2.0 ml) at 100 ºC under the atmosphere. After decontamination, the ILs remained on surfaces of steel wastes were washed off with ethanol. Uranium concentrations (Bq/g) of decontaminated steels were evaluated as the ratio of radioactivity due to U of samples to total weight of samples. Figure 15 shows a plot of U concentrations (Bq/g) against time in the dissolution of adhered uranium by soaking the contaminated steel waste into BMICl. The U concentrations are found to drop below the temporary proposed clearance level (1.0 Bq/g) within 3 h under the present conditions (IAEA, 2004).

![Figure 15](image)

**Fig. 15.** A plot of uranium concentrations of steel waste vs. soaking time for the dissolution in BMICl at 100 ºC under the atmosphere

Furthermore, the XPS spectra for the top surface of the steel waste were measured after decontamination treatment. As a result, the peaks due to UF₄ and FeF₃ were found to disappear. This indicates that the FeF₃ component is also dissolved with the dissolution of UF₄ in BMICl.

As mentioned in 3.2, it is suggested that the uranium component can be recovered electrolytically from the BMICl solution dissolving UF₄. Hence, it should be possible to recover only uranium component from the solutions after decontamination of the steel wastes in BMCl by controlling electrolytic potential.

### 3.4 Summary for application of ILs to the treatment of wastes contaminated with uranium

Dissolution behaviour of UF₄ in BMICl and the electrochemical properties of dissolved uranium species were investigated. Based on such basic studies, the feasibility of decontamination of steel wastes contaminated uranium using BMICl as medium was also examined. The results are summarized as follows.

- UF₄ can be dissolved completely in BMICl by heating under the atmosphere.
From the UV-visible absorption spectra of dissolution solutions, it was found that the dissolved uranium species are oxidized to uranyl(VI) by O$_2$ and that the resulting uranyl(VI) species are the complex with the mixed ligands of F and Cl.

The CV measurements suggest that the resulting uranyl(VI) species with the mixed ligands of F and Cl are reduced to UO$_2$ electrochemically.

The steel wastes contaminated with UF$_6$ can be decontaminated below the temporarily proposed clearance level (1.0 Bq/g) within 3 h by soaking in BMICl at 100 ºC.

It should be possible to recover only uranium component from the solutions after decontamination of the steel wastes in BMICl by controlling electrolytic potential.

4. References


This book is the second in the series of publications in this field by this publisher, and contains a number of latest research developments on ionic liquids (ILs). This promising new area has received a lot of attention during the last 20 years. Readers will find 30 chapters collected in 6 sections on recent applications of ILs in polymer sciences, material chemistry, catalysis, nanotechnology, biotechnology and electrochemical applications. The authors of each chapter are scientists and technologists from different countries with strong expertise in their respective fields. You will be able to perceive a trend analysis and examine recent developments in different areas of ILs chemistry and technologies. The book should help in systematization of knowledges in ILs science, creation of new approaches in this field and further promotion of ILs technologies for the future.

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