Solvent Extraction of Tc(VII) by Calixarenes Bearing Pyridino Groups

Rainer Ludwig\textsuperscript{b,a} and Nguyen Thi Kim Dzung\textsuperscript{a,b}

\textsuperscript{a}Freie University Berlin, Inst. of Chem./IAC, Fabeckstr. 34-36, 14195 Berlin, Germany
\textsuperscript{b}ITRRE, Vietnam Atomic Energy Commission, 48 Lang Ha Street, Hanoi, Vietnam

Received: July 31, 2005; In Final Form: September 30, 2005

The distribution of Tc(VII) between aqueous acidic solutions and organic solutions of nitrogen-containing extractants is investigated in order to establish the relationships between extraction power towards the Tc(VII)-anion, ligand molecular structure, influence of the media and the speciation in this ion-pair extraction. Ligands belonging to the class of calixarenes, derivatized with pyridino groups were newly developed for these purposes. The number of substituents per calixarene, the position of the nitrogen and the size of calixarene are varying structural parameters. The nitrogen atoms are either protonated or quaternized to allow ion-pair extraction. The aqueous phase contains nitrate or chloride at varying or constant ionic strength. Experiments using Aliquat-336 were conducted under similar conditions.

1. Introduction

In the course of studies on coupled transport of Tc(VII) and U(VI) by conventional extractants such as TBP\textsuperscript{8}, we realized the impact of anion competition and acid extraction on the separation of Tc(VII). Furthermore, we were interested whether macrocyclic ligands are able to extract Tc(VII) more selectively compared with non-cyclic or monodentate ligands. Among the latter, qua- termary amines such as Aliquat-336 extract Tc(VII) well.\textsuperscript{2,3} An example for the macrocyclic effect with cooperative binding to the anion was recently observed with respect to HPO\textsubscript{4}\textsuperscript{2−}.\textsuperscript{4}

Among macrocyclic ligands, calixarenes\textsuperscript{9} gained significant interest in the field of ion recognition. So far, few publications on the extraction of Tc(VII) by nitrogen-containing crowned calixarenes were published, in which the relationship between ligand structure and binding was thoroughly investigated.\textsuperscript{6,7} Metalated cyclotriveratrylenes extract anionic Tc(VII) and Re(VII) as ion pairs, the selectivity over competing anions improves when the ligand has a deep cavity.\textsuperscript{9} Calix[4]arenes with carbonyl groups\textsuperscript{9} were reported to complex alkali ions and extract Tc(VII) as ion-pair at high pH. Crowned calixarenes turned out to be effective in co-extracting Tc(VII) with Cs\textsuperscript{+} as counterion from both acidic and alkaline media.\textsuperscript{10}

In terms of this work we synthesized a series of calix[n]arenes (n = 4, 6, 8) bearing methylypyridyl (picolyl) groups and investigated their ligating behaviour by means of solvent extraction from acidic media. The calixarene acts as molecular backbone to which the pyridino groups are anchored via methoxy groups in vicinity to each other. The purpose of the pyridino groups in protonated state is to provide an anion exchange functionality, and the possibility of backextraction by deprotonating them. The number of pyridino groups per molecule, the position of the nitrogen atom as well as the size and flexibility of the calixarene vary among the ligands in order to systematically study the influence of the molecular structure. In addition, ligands bearing quaternized pyridinium groups were synthesized. All these substituents were attached to the so-called ‘lower rim’ of the calixarene. In the distribution experiments, the concentration and kind of acid as well as the ionic strength were varied. Extraction by Aliquat-336\textsuperscript{a} under the same conditions was done for reasons of comparison.

2. Experimental

The structures of the extractants developed for this purpose are shown in Scheme 1. In addition to those, di- and tri-substituted 2PyOC4 were investigated in terms of work. The abbreviation

\begin{scheme}
Scheme 1. Extractants developed for this work.
\end{scheme}

© 2005 The Japan Society of Nuclear and Radiochemical Sciences
Published on Web 12/28/2005
2PyOC4 for example means octyl calix[4]arene functionalized with four 2-pyridyl methoxy groups. The compounds 2PyOC4, 4PyOC4$^*$ and 2PyBuC6$^*$ were reported previously. The synthesis of the other ligands involves treatment of the corresponding parent calix[n]arene (n = 4, 6, 8) with a base, followed by reaction with picolylchlorides and work-up by crystallization or column chromatography, similar to References 13 and 14. Quaternization was done by reaction with methyliodide. The compounds were identified by $^1$H/$^13$C-NMR, elemental analysis, MS and thin-layer chromatography. Their analytical data and the synthesis details will be reported elsewhere.

Solvent extraction was carried out batchwise in sealed PE-vials at 295 ± 2 K and equal phase volumes of 3 mL. Magnetic stirring was carried out overnight. It has not been tested which time period is sufficient to reach equilibrium. In previous anion extraction$^{15}$ it was two minutes, while in Ag(I) extraction a slow kinetics was observed.$^{11}$ The organic phase contained 0.002 M calixarene or 0.008 M Aliquat-336 in chloroform. The aqueous phase contained 0.00005 M $^{99}$Tc(TcO$_7^{2-}$) (3 kBq) and either nitrate or chloride: HNO$_3$ (0.003 to 4 M) or a mixture of HNO$_3$ and Na$_2$NO$_3$ (3 M NO$_3$); or HCl (0 to 5 M). For weakly acidic solutions, the pH at equilibrium was measured with a glass electrode. One mL of the aqueous phase was subjected to liquid scintillation $\beta$-counting. The activity in the organic phase was calculated from the difference to the initial one.

3. Results and Discussion

**Effect of substituent number.** Figure 1a shows the distribution coefficients ($D = c_{aq}/c_{org}$) for Tc(VII) in the extraction by calix[4]arenes bearing 2, 3, or 4 pyridino (py) substituents at the lower rim. In this series, the nitrogen atom is located ortho to the anchor group (2PyOC4), thus being located ‘inside’ the cavity. The extractability decreases when the nitric acid concentration increases, an effect which is discussed later.

There is a significant increase in log $D$ when the number of py-groups changes from two to three. Introducing a fourth py-group contributes little to the extractability. It is concluded that three nitrogen atoms are mainly involved in binding to TcO$_7^{2-}$ by means of ion-ion, cation-dipole and hydrogen bond interactions. The calix[4]arene backbone may be too rigid to allow the fourth nitrogen atom to rearrange into a position suitable for interaction.

Calix[6]arenes are characterized by a higher molecular flexibility and the possibility to introduce up to 6 functional groups. In Figure 1b, data for the 2PyOC6 are shown. In both Figures 1b and 1a, the nitrogen is located at ortho position. Results for nitrogen in 3-position (3PyBuC6 and 3PyOC6) are given in Figures 1c and 1d, respectively. However, the anchoring of 3, 4, or 6 py-groups does not significantly improve the extraction of Tc(VII) as compared with calix[4]arenes as seen in Figures 1b to 1d. The extraction improves only slightly when more py-groups are contained within the molecule. The reasons may be (i) that the short methoxy spacers do not allow re-orientation of the py-groups according to an ‘induced fit’ mechanism, and (ii) entropy loss of the ligand during complexation. Data for the hexa-substituted butylcalix[6]arene are not included, because the ligand precipitates upon complete protonation. Octyl groups render the ligands sufficiently hydrophobic even in the protonated state and this was the reason why for this study the majority are octylcalixarenes.

**Position of the nitrogen atom in the substituents.** Figure 2 shows results for calix[4]arenes where the nitrogen atom in the py-groups is located at the 4-position (4PyOC4), ‘outside’ the cavity. As in Figure 1a, the values for log $D$ are higher for the tetra-substituted ligand. Changing the acid concentration in the aqueous phase over a wider range shows two regions for the distribution coefficient: an increase at the weakly acidic
region results from the protonation of the nitrogen atoms. Although the protonation constants were not yet measured, we assume that protonation occurs stepwise due to cooperativity, as does deprotonation in carboxylic acid derivatives. The distribution coefficient decreases at higher acidity nearly linearly on a half-logarithmic scale.

In Figures 3a and 3b the position of the nitrogen atom, either ortho or para with respect to the anchor group (2PyOC4 and 4PyOC4, respectively) is compared for partially (3a) and fully substituted (3b) calix[4]arenes. We observe significant differences: In the di-substituted ligands, the protonation of the nitrogen dominates up to 0.3 M HNO₃ and causes an increase in log D. In tetra-substituted calixarene, the 2PyOC4 starts with high log D-values, while 4PyOC4 first passes the protonation stage before it can extract Tc(VII). We interpret this difference in terms of a cooperative effect in 2PyOC4, which promotes protonation even at low acidity. On the other hand, 4PyOC4 offers the advantageous possibility of Tc-backextraction with diluted acid.

Figure 4 compares ortho- and meta-position of the nitrogen (2PyOC6 and 3PyOC6, respectively). As for the calix[4]arenes on the previous picture, in these calix[6]arenes the nitrogen in 2-position (ortho) appears to be protonated all over the investigated range. The 3Py-derivative (meta position) lies in-between the ortho- and para ones, showing a region where it is not yet fully protonated (< 0.1 M HNO₃). We therefore conclude that in these fully-substituted macrocycles, the protonation constants follow the order 2Py > 3Py > 4Py, with possible stabilization due to intramolecular hydrogen bonding.

**Calixarene size.** Next, we compared the fully-substituted calix[n]arenes with n = 4, 6, or 8 in order to find an optimum extraction performance. In this series, 4, 6, or 8 pyridino groups respectively are attached to one molecular backbone with their nitrogen atoms in ortho position. The results are shown in Figure 5. Surprisingly, the ligands show a very similar behaviour, with the exception of the calix[8]arene at lower acidity. The latter is ascribed to the high molecular flexibility of this large macrocycle, which may hinder the intramolecular stabilization of the protonated form. However, above 0.1 M HNO₃ the differences between the three ligands level off and the values of log D decrease with the acid concentration linearly on a half-logarithmic scale. The question arises why does a compound with 4 ligating groups performs similar to its homologues containing 6 or 8 of these groups? We conclude that not all of the available nitrogen atoms in the larger calixarenes contribute to the anion binding. In other words, these ligand appear to represent not yet the optimum molecular structure to match the anion coordination sphere.

**Acid competition.** In the previous Figures the distribution coefficient decreased at higher acidity. In order to answer the question if this caused by the competitive extraction of nitrate or by the protonation of pertechnetate, we varied the composition of the aqueous phase. Figure 6 depicts the results. At constant
Anion competition in the extraction of Tc(VII). In the di-substituted compound 4PyOC4Di-Q, the hydrophobic character dominates and it is not soluble in water.

**4. Conclusions**

It turned out that (i) the extraction improves with more pyridino groups per calixarene molecule; (ii) the number of substituents and the position of the nitrogen atom in the pyridine ring influence the ligand protonation and thus the extraction behaviour towards Tc(VII) at lower acid concentration which follows the order 2Py > 3Py > 4Py; and (iii) molecular flexibility, cavity size and the number of pyridino groups in fully substituted calix[n]arenes (n = 4, 6, 8) compensate each other which causes a similar extractability towards Tc(VII) among them.

By varying the composition of the aqueous phase, it was shown that nitrate competes with pertechnetate. The competition by chloride is weaker. Protonation of TcO$_4^-$ contributed negligibly to the decrease of extraction up to 4 M acid in the aqueous phase. Quaternized nitrogen is a better ligand than protonated one in weakly acidic media. A macrocyclic effect was observed. The binding is explained by electrostatic and hydrogen bonding interactions.

**Acknowledgement.** The authors thank Cognis, Mining- & Ion-Transfer Chemicals Technology for a free sample of Aliquat-336.

Financial support by the National Science Council/Fundamental Research Projects of Vietnam is appreciated.
Reference
