Incorporation Solvation With Cloud Point Extraction Method for Separation and Spectrophotometric Determination of Cerium (III)

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ABSTRACT

Through joined two sensitive methods solvation and cloud point extraction separated and determined micro amount of Ce(III) from aqueous solutions by using organic reagent 2,4-Dimethyl pentan-3-one (2,4-DMP) as extractant and non-ionic surfactant 1% TritonX-100, the experimental study shows the solvation species of Ce$^{3+}$ extracted into cloud point layer (CPL) have a wavelength for maximum absorbance was $\lambda_{max}=295$ nm and the laboratory experiments appear needed as optimum concentration of salting out KNO$_3$ was 0.6 M in the presence 100µg Ce$^{3+}$/10 mL and $1x10^{-4}$ M (2,4-DMP) so that 0.5 mL at % tritonX-100 as well as after heating the solution at 90°C for 15 minutes the study involved effect of different kind and concentration of salting out, and interferences so that effect of different organic reagent as extractants, also stoichiometry study demonstrated the more probable structure of solvation extracted species was 1:1 2,4-DMP: Ce(NO$_3$)$_3$ with spectrophotometric determination.

Keywords: Cerium(III) , Solvation method, Cloud point extraction method, Preconcentration, Salting out agent.

INTRODUCTION

Extraction experiments for Ce$^{3+}$ ion from aqueous phase by new laboratory prepared Azo derivation as complexation agent 4- [N-(5-methyl isoxazol-3-yl)benzene sulfonamide azo]-1- Naphthol (AMBN) shows the optimum conditions for this extraction method was (pH= 9) (10 minutes) shaking time and 100µg (1.5x10-4M) concentration of Ce$^{3+}$ ion in aqueous phase. Organic solvents effect study$^1$. Extraction ability of 8-HQ in the countenance of DB18C6, DCH18C6, and DB18C6 crown ethers for chosen Ln(III) ions from acidic media to DMSO/ chloroform solvent system. The study effect of HNO$_3$ concentrations on the selectivity of extraction, the effects of solvent, pH, Crown ether organic reagent and concentrations of crown ethers$^2$. Separation and extraction of Zn(II) and Cd(II) as a neutral compound by used methyl stearate as extractant dissolved in chloroform. Applications of separation
and extraction according to solvation method from nitrate aqueous solution by used methyl stearate which is binding coordinately with nitrate salt of metal under study Zn(II) and Cd(II). The extracted species have \( \lambda_{\text{max}} = 263 \) nm for Zn\(^{2+}\) and \( \lambda_{\text{max}} = 346 \) nm for Cd\(^{2+}\). The studies about limitation optimum condition for extraction show 0.3 M salting out effect KNO\(_3\) for Zn\(^{2+}\) and 0.7 M KNO\(_3\) for Cd\(^{2+}\) in present 100 µg Zn\(^{2+}\)/5 mL and 50 µg Cd\(^{2+}\)/5 mL with optimum shaking time 15 min. for both ions\(^{[3]}\). N-octylaniline in dissolved xylene was used as complexing agent for the extraction separation of Cerium (IV) from salicylate medium. Ce(IV) was stripped from the organic phase with [2.0M] HCl as well as determined spectrophotometrically by Arsenazo III. We studied some effect of parameters such as of pH, effect of metal ion, equilibration time, stripping agents and diverse ions on the extraction of Ce(IV). This study stoichiometric of the extracted species was obtained from slope analysis. The method applicable for the determination of Ce(IV) in semiconductor films, synthetic mixture and monazite sand\(^{[6]}\).

**EXPERIMENT**

Biochrom model (80-7000-11) Libra S60 Cambridge CB40FJ spectrophotometer with 1.00 cm quartz cell was used for recording the absorbance spectra and absorbance, controlled the temperature of heating so that the necessary time of heating needed to used electrostatic water bath (WNB7-45) (England) was used to maintain the temperature and time.

**MATERIALS AND SOLUTIONS**

All chemical received from sedate commercial company and used as received without any furthermore purification stock solution of Ce\(^{3+}\) ion (1000 µg/mL) prepared by dissolved 0.309 g of Ce(NO\(_3\))\(_3\) \(\cdot\)6H\(_2\)O in 100mL distilled water contain 1mL of nitric acid HNO\(_3\) and other working solutions prepared by dilution with distilled water and in all preparation of solutions used volumetric flask, so that 1% 8-Hydroxy quinolone dissolved in ethanol, as well \(1\times10^{-2}\) M 2,4-DMP prepared by dissolved 0.1142 g in 100 mL distilled water and other working solution prepared by dilution with distilled water in suitable volumetric flask.

For determination quantity of Ce\(^{3+}\) in the aqueous phase, we have followed 8-HQ Spectrophotometric method\(^{[7]}\) and with the following equation:

\[
y = 0.0117x + 0.0005 \\
R^2 = 0.9998
\]

![Fig. 1: Calibration curve for Ce\(^{3+}\) by 8-HQ method](image-url)
performance of this method preparing calibration curve by making an experiment on many solutions contain rising quantity of Ce$^{3+}$ the results were as in Figure (1):

**Comprehensive Method**

Preparing 10 mL aqueous solutions to contain 100 µg of Ce$^{3+}$ ion with an optimum concentration of salting out KNO$_3$ and 1×10$^{-4}$ M from 2,4-DMP and 0.5 mL % Triton X-100, heating this solution at 90 °C for 15 min. until complete formation cloud point layer (CPL) with smaller volume and higher density, than separated the cloud point layer (CPL) from aqueous solution, afterward dissolved CPL in 5 mL ethanol and measure the absorbance of ethanolic solution at $\lambda_{max}$ of solution species extracted against blank prepared in the same manner without metal ion, but the aqueous solution treated according to 8-HQ spectrophotometric method and after return to the calibration curve determined the reminder quantity of Ce$^{3+}$ in aqueous phase after extraction, then subtraction the reminder quantity from the original quantity to determine the transferred quantity of Ce$^{3+}$ to the CPL as solvation species afterward calculate the distribution ratio D, according to relative below:

$$D = \frac{[\text{Ce}^{3+}]_{\text{CPL}}}{[\text{Ce}^{3+}]_{\text{aq}}}$$

We are could be determine the quantity of Ce$^{3+}$ transferred to the CPL by stripping method involved shaking ethanolic solution of CPL with two 5ml portion of 1 M HNO$_3$ aqueous solution to destroy the solvation species and transferred Ce$^{3+}$ ion into aqueous acid solution and the by application 8-HQ spectrophotometric method to determination Ce$^{3+}$ ion in acidic aqueous solution but by performed some experimental stripping method fined the transferred quantity of Ce$^{3+}$ ion. Determined by stripping method was equal to some quantity determined subtraction then we are followed the subtraction method to determine the transferred quantity in all experiments because easier and faster.

**RESULTS AND DISCUSSION**

The spectrophotometric study involved preparing 10 mL aqueous solutions contain 100µg Ce$^{3+}$ and 0.5 M KNO$_3$, 0.5 mL of % Triton X-100 and 1×10$^{-4}$ M (2,4-DMP) heated the solution at 90 °C for 15 min. until complete appearance cloud point layer (CPL) separated this layer and dissolved in 5 mL ethanol and taking the UV-Visible spectrum for it against blank prepared in the same manner in absence Ce$^{3+}$, the result as in Figure (2):

The spectrum appears wavelength for maximum absorbance was $\lambda_{max}$=295 nm.

**Effect of salting out concentration**

Preparing 10 mL aqueous solutions each solution contain 100 µg Ce$^{3+}$ with rising concentration of salting out KNO$_3$ and 0.5 mL of % Triton X-100 in the presence, 1×10$^{-4}$ M (2,4-DMP) heating this solution at 90°C for 15 min. then complete the waking

![Fig. 2: UV-Vis spectrum for solvation species of Ce$^{3+}$](image-url)
as in comprehensive method the results were as in
Figure (3 and 4):

The results shows there was increasing
the extraction efficiency of Ce$^{3+}$ by increasing

![Graph](image1)

**Fig. 3: Effect of salting out concentration on
formation and stability of solvation species**

 Concentration of salting out KNO$_3$ until reach to
the maximum at optimum concentration value 0.6
M KNO$_3$ then decrease extraction efficiency by
increasing concentration of KNO$_3$ more than optimum
value the results appear the increasing of salting out
KNO$_3$ effected to increase the forward direction of
thermodynamic equilibrium for formation solvation
species and the maximum rate of formation with
higher concentration of solvation species obtained
at 0.6 M KNO$_3$ of concentration more than optimum
value effect to decrease the rate of formation
direction with decrease the concentration of solvation
species extracted according to mass action law.

**Effect of Ce$^{3+}$ concentration**

Extracted Ce$^{3+}$ ion according to the
comprehensive method from 10mL aqueous
solutions contain a different quantity of metal ion the
results were as in Figures (5 and 6):

The results there is a linear relative for effect
of metal ion concentration whereas the concentration
of metal ion considers as a thermodynamic parameter

![Graph](image2)

**Fig. 5: Effect of metal ion concentration on
solvation species formation and stability**

![Graph](image3)

**Fig. 6: Effect of metal ion concentration on
extraction efficiency**

and the increasing concentration of metal ion to give
rise to increasing the rate of forward direction for
thermodynamic equilibrium that is a mean increase
in concentration and stability of solvation species extracted.

**Variation 2,4-DMP concentration**

By application the comprehensive method
for extraction Ce$^{3+}$ ion in the presence rising
concentration of organic reagent 2,4-DMP the results
were as in Figures (7 and 8).

The results appear there is a linear relation between extraction efficiency and molar
concentration of organic reagent 2,4-DMP, the
increasing of extraction efficiency indicate to behave
2,4-DMP as thermodynamic parameter effect with
increasing its concentration to increase the rate
of forward thermodynamic direction and increase
solvation species concentration and stability
Effect of variation surfactant volume

According to comprehensive method extracted Ce\(^{3+}\) ion from 10mL aqueous solution at optimum conditions by using the different volume of surfactant 1% Triton X-100, the results were as in Figures (9 and 10):

The results appear 0.5mL 1% Triton X-100 was optimum volume of surfactant giving higher extraction efficiency because at this volume reached to the critical micelles concentration CMC of the surfactant to obtain best cloud point layer with suitable volume and density to extracted higher concentration of solvation species according to thermodynamic equilibrium any volume less than optimum not allow to reach favorite equilibrium so that volume more than optimum giving decrease in extraction efficiency because effect of increasing diffusion micelles and produce cloud point layer with larger volume and less density.

Thermodynamic study

By application the procedure detailed in the comprehensive method at optimum conditions extracted Ce\(^{3+}\) at different temperature of heating in an electrostatic water bath for 15minutes the results were as in Figures (11 and 12):

After calculated extraction constant K\(_{ex}\) by application the relation below and the diagram between logarithm values of K\(_{ex}\) against 1/T K the results demonstrate in Figure (13).

Then from the slope straight line relation in Figure (13) and application relation below
Fig. 12: Effect of temperature on extraction efficiency and D values

![Graph showing the effect of temperature on extraction efficiency and D values.]

Table 1: Thermodynamic data for extraction Cerium

<table>
<thead>
<tr>
<th>ΔSex kJ/mol</th>
<th>ΔGex kJ/mol</th>
<th>ΔHex kJ/mol</th>
</tr>
</thead>
<tbody>
<tr>
<td>184.520</td>
<td>-67.100</td>
<td>0.1198</td>
</tr>
</tbody>
</table>

The results appear 90°C was the optimum temperature for extraction solvation species of Ce³⁺, this temperature was the favorite temperature to formation cloud point layer and giving best extraction according to solvation method any temperature more than optimum effect to decrease density and increase volume of cloud point layer by increasing diffusion for micelles, the minimization value of enthalpy ΔHex interpret to approach one another in solvation species extracted so that the large positive value of entropy ΔS_ex reflect the importance of entropy in extraction method and the method of extraction was entropic in region.

Effect of heating time

According to comprehensive method extracted Ce³⁺ from 10mL aqueous solution at optimum conditions with different times of heating the results were as in Figures (14 and 15).

The results demonstrate 15minutes was the optimum heating time in extraction method whereas the time of heating represent the kinetic side of extraction method by cloud point method, at this time reached to the maximum kinetic energy help to aggregation the micelles at best state with
higher dehydration to form CPL with best properties at less time the kinetic energy was not enough, so that heating time more than optimum giving increase in kinetic energy which effect of increasing diffusion of micelles.

Effect of kinds salting out

Extracted Ce$^{3+}$ ion according to the procedure detailed in the comprehensive method by using different kinds of salting out with different concentration for each one the results were as in Figures (16 and 17):

The results appear there is different extraction efficiency and solvation species formation and stability with different kinds and concentration of salting out because the different salting out have different behavior in aqueous solutions that is cloud point layer by effect on the dehydration of micelles.

Interferences

From 10mL aqueous solutions extracted Ce$^{3+}$ at optimum conditions in the presence of foreign ions the results were as at Table (2).

The results show all the foreign ions presence in the aqueous solution with Ce$^{3+}$ ion giving interference effect to decline the extraction efficiency of Ce$^{3+}$ with different degrees, according to the behavior and activity of the foreign ion in aqueous phase, the interfere included participation of each foreign ion to formation solvation species with different concentration this behavior effect to decrease salting out concentration KNO$_3$ and organic reagent 2,4-MP from the optimum concentration necessary to formation solvation species for Ce$^{3+}$ which mean decreasing in extraction efficiency.

Effect of organic reagent structure

Extracted Ce$^{3+}$ ion as in the procedure detailed in the comprehensive method at optimum conditions by using different organic reagent the results were as in Table 3.

The results demonstrate there is different extraction efficiency for Ce$^{3+}$ with each organic reagent is one of the thermodynamic parameters according to its structure which is the effect on the thermodynamic equilibrium as well as the rate of formation solvation species extracted.

Stoichiometry

By using two spectrophotometric method slope analysis and slope ratio methods according to the procedure detailed in the comprehensive
Table 3: Effect of organic reagent structure no extraction efficiency of Cerium ion.

<table>
<thead>
<tr>
<th>Organic reagent</th>
<th>$\lambda_{\text{max}}$</th>
<th>Absorbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isobutyl methyl ketone</td>
<td>294</td>
<td>0.5</td>
</tr>
<tr>
<td>Butanol</td>
<td>282</td>
<td>0.88</td>
</tr>
<tr>
<td>Tributyl phosphato acetone</td>
<td>305</td>
<td>0.11</td>
</tr>
<tr>
<td>Acetophene</td>
<td>287</td>
<td>0.7</td>
</tr>
</tbody>
</table>

method the results were demonstrated in Figures (18→20).

This two method demonstrated the more probable structure of solvation species structure extracted to CPL was 1:1 organic reagent: metal ion (NO$_3$)$_3$, 2,4-DMP: Ce(NO$_3$)$_3$.

Spectrophotometric determination of Ce$^{3+}$

For determination Ce$^{3+}$ ion in different aqueous samples by application comprehensive method at optimum conditions performed the calibration curve for that goal as in Figure (20):

![Figure 21: Calibration curve for spectrophotometric determination of Ce$^{3+}$ in different samples](image)

**Fig. 21: Calibration curve for spectrophotometric determination of Ce$^{3+}$ in different samples**

$$y = 0.1142x + 0.0024$$

$R^2 = 0.9997$

**Fig. 20: Slope ratio effect of Cerium concentration on extraction efficiency**

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