EXTRACTION OF Ni(II) IONS INTO CHCl₃ SOLUTION OF N,N'-ETHYLENEBIS(4-BUTANOYL-2,4-DIHYDRO-5-METHYL-2-PHENYL-3H-PYRAZOL-3-ONE IMINE) SCHIFF BASE

J. Godwin[a], F. C. Nwadire[b] and B. A. Uzoukwu[c]

Keywords: Extraction, Nickel(II), N,N’-Ethylenebis(4-butanol-2,4-dihydro-5-methyl-2-phenyl-3H-pyrazol-3-one imine), Effect of pH and anions

The extraction of Ni(II) ions in aqueous media was studied using chloroform solution of N,N’-ethylenebis(4-butanol-2,4-dihydro-5-methyl-2-phenyl-3H-pyrazol-3-oneimine) (H₂BuEtP) Schiff base as the organic extractant. The synergistic effect of 4-butanol-2,4-dihydro-5-methyl-2-phenyl-3H-pyrazol-3-one (HBuP) in these extractions were also investigated. The pH₃ of extraction of Ni(II) was significantly lowered from a near neutral pH 7.14 ± 0.10 to an acidic region of 5.51 ± 0.10 when a mixture of the ligands was used. The partition coefficient (log Kₐ₉) 1.89 ± 0.05 (H₂BuEtP) was same as (log Kₐ₉) 1.89 ± 0.02 (H₃BuEtP·HBuP) while the extraction constant (log Kₐ₉) -12.39 ± 0.64 (H₂BuEtP) was slightly less than (log Kₐ₉) -10.57 ± 0.10 (H₃BuEtP·HBuP). Data analysis indicated that Ni(II) distributed slightly better into chloroform solution of H₂BuEtP/HBuP as Ni(HBuEtP)(BuP) than into a solution of H₃BuEtP as Ni(HBuEtP)₂BuP. Mineral acids show a masking effect in the extractions in both ligand alone and mixed ligands systems while mixed ligands system was shown to be a better extractant system for Ni(II) ions in the presence of anions and auxiliary complexing agents studied. Acetate, fluoride, and phosphate ions gave the best percentage extractions and the extraction of Ni(II) ions with the ligand is more efficient at near neutral to basic pH.

Introduction

Schiff bases are compounds that contain the carbon-nitrogen double bond traditionally connected to an aryl or alkyl group. Earlier studies on isolation and characterization of metal complexes of Schiff bases have shown that they are capable of forming stable metal complexes with Cu(II) and Ni(II) and unstable metal complexes with Mo(VI), Co(II) and Cd(II).¹² Derivatives of 1-Phenyl-3-methyl-4-acylpyrazolone-5 are known for their complexation reactions with transition metals forming metal complexes with interesting coordination chemistry.³⁴⁵ Most of the current systems used for extracting metals work best mostly at acidic pH. However, natural waters are at near neutral to basic pH, and much of the currently stored wastes are at very caustic conditions. Thus, there is high demand for ligands that can extract over a wide range of pH.

In continuation of our earlier work on the synthesis, characterization⁷ of 1-phenyl-3-methyl-4-acylpyrazolone-5 derivatives and extraction of metal ions with N,N’-ethylenebis(4-butanol-2,4-dihydro-5-methyl-2-phenyl-3H-pyrazol-3-oneimine) Schiff base, which has been successfully applied in the extraction of Lead(II)⁸ and U(VI) ions², we report the application of the present Schiff base derivative of 4-butanolpyrazolone as a potential extractant for Nickel(II) ions. The synthesis provided an opportunity for a N=C-C-OH bonding moiety and extended the scope of coordination to involve tetradeptate ligands from the initial bidentate 4-acylpyrazolone.

Materials and Methods

Reagents and Apparatus

N,N’ - Ethylenebis(4 - butanolyl - 2, 4 - dihydro - 5 - methyl-2-phenyl-3H-pyrazol-3-oneimine) (H₂BuEtP) was synthesized by methods reported earlier.⁷ The ligand’s purity after recrystallization from aqueous ethanol was established by elemental analysis for C, H and N; analysis of IR and NMR spectral data at the Institute for Inorganic Chemistry, University of Dresden, Germany.

Stock solutions of 0.05 M H₂BuEtP were prepared by dissolving appropriate amount of the ligand in CHCl₃. Stock solutions of 0.05 M 4-butanolyl-2,4-dihydro-5-methyl-2-phenyl-3H-pyrazol-3-one (HBuP) were also prepared by

---

Corresponding author
Email: godwino2012@gmail.com
Tel.: +2348033401456
[b] Department of Pure and Industrial Chemistry, University of Port Harcourt, Choba, Port Harcourt, Rivers State. Nigeria, Email: nwadirefc@yahoo.com
[c] Department of Pure and Industrial Chemistry, University of Port Harcourt, Choba, Port Harcourt, Rivers State, Email: uzoukwupob331@yahoo.co.uk

dissolving appropriate amount of the ligand in CHCl₃. Stock solutions of 1.704 x 10⁻² M (1,000 mg/L) of Ni(II) were prepared by dissolving appropriate amount of ammonium nickel(II) tetraoxosulphate(VI) hydrate (2:1:2:6), (NiSO₄·(NH₄)₂SO₄·6H₂O) in 0.1 mL of 10 M HNO₃ and making up to mark in a 50 ml volumetric flask with deionized water. Buffer solutions were prepared with 0.1M HCl/0.1 M NaCl (pH 1.0-2.9), 0.1 M acetic acid/0.1 M NaCl (pH 3.0-3.5), 0.1 M acetic acid/0.1 M Na-acetate (pH 3.6-5.6), and 0.1 M KH₂PO₄/0.1 M NaOH (pH 5.7-10.0). pH of the buffered solutions were determined with a Labtech Digital pH meter. Solutions of 0.001-3.0 M mineral acid, 0.001 - 1.0 M anions or 0.0005 M - 0.5 M auxiliary complexing agent concentrations were prepared by diluting appropriate volumes of stock solutions of mineral acids or sodium salts of anions or auxiliary complexing agent. All experiments were performed at ionic strength of 0.1 M (NaClO₄).

### Extraction Procedure

2 mL aliquot of a buffer solution containing 8.52 x 10⁻⁴ M (50 mg/L) of Ni(II) ions and the desired pH of solution was prepared in a 10 ml extraction container. For extraction studies involving mineral acid or complexing agents, 2 ml aliquot of solution containing 8.52 x 10⁻⁴ M of Ni(II) ions and the desired mineral acid or complexing agent concentration was prepared in a 10 ml extraction container. An equal volume (2 mL) of chloroform solution of 0.05 M concentration of H₂BuEtP or 0.05 M H₂BuEtP : 0.05 M HBuP (9:1 ratio by volume) was added and the mixture shaken mechanically for 30 minutes at room temperature of about 30 °C. A shaking time of 30 minutes was found suitable enough for attaining the equilibration. The phases were allowed to settle and separated. Concentration of Ni(II) ion in aqueous phase was determined with a Buck Scientific Atomic Absorption Spectrophotometer (AAS) 205 at wavelength of 232.0 nm. Ni(II) ion concentration extracted into the organic phase was determined by the difference between the concentration of Ni(II) ion in aqueous phase before and after the extraction. Distribution ratio \( D \) was calculated as the ratio of metal ion concentration in the organic phase \( (C_o) \) to that in the aqueous phase \( (C) \). Thus \( D = C_o/C \).

### Results and Discussion

#### Effect of buffer solutions

Extraction of Ni(II) from aqueous phase into an organic phase \((o)\) containing the ligand H₂BuEtP can be represented by equation

\[
Ni^{2+} + H_2BuEtP_{(o)} \rightleftharpoons Ni(BuEtP)_{(o)} + 2H^+ \]  

(1)

where H₂BuEtP is a tetradeinate ligand. It shows that the reaction should take place in the metal:ligand mole ratio of 1:1. Thus the extraction constant \( K_{ex} \) can be represented by,

\[
K_{ex} = \frac{[Ni(BuEtP)_{(o)}][H^+]^2}{[Ni^{2+}][H_2BuEtP_{(o)}]} \]  

(2)

The distribution ratio \( D_o \) can be calculated as the ratio of metal ion concentration in the organic phase \( (C_o) \) to that in the aqueous phase \( (C) \). Thus \( D_o = C_o/C \).

\[
Ni^{2+} + 2H_2BuEtP_{(o)} \rightleftharpoons Ni(HBuEtP)_{2(o)} + 2H^+ \]  

(4)

The distribution ratio \( D \) is calculated as the ratio of metal ion concentration in the organic phase \( (C_o) \) to that in the aqueous phase \( (C) \). Thus \( D = C_o/C \).

\[
K_{next} = \frac{[Ni(HBuEtP)_{2(o)}][H^+]^2}{[Ni^{2+}][H_2BuEtP_{(o)}]^2} \]  

(5)

The distribution ratio \( D_o \) can be calculated as the ratio of metal ion concentration in the organic phase \( (C_o) \) to that in the aqueous phase \( (C) \). Thus \( D_o = C_o/C \).

\[
lgD_{Ni} = lgK_{next1} + 2log[H_2BuEtP] + 2pH \]  

(6)
more acidic region indicating that extraction of Ni$^{2+}$ ions at lower pH values was more effective in chloroform solutions containing HBuP as a synergist. Uzoukwu et al$^{10}$ obtained a similar result from their studies on the effect of chloride ion (Cl$^{-}$) concentration on Cu(II) and Ni(II) ions using 1-phenyl-3-methyl-4-trichloroacetylpyrazol-5-one. The partition coefficient $K_{NiD2}$ is given by the equation of $K_{NiD2} = [Ni(HBuEtP)(BuP)_{o}]_{o}/[Ni(HBuEtP)(BuP)]$, for which a value of 1.89 ± 0.02 that was statistically same with $\log K_{NiD1}$ was determined from the graph in Fig. 1b.

Plots of log D against pH in the mixed ligands system in figure 1b gave a slope of 2 indicating that 2 moles of hydrogen were displaced during the extraction process. Interaction between Ni(II) and H$_2$BuEtP in the presence of HBuP at constant pH 6.0, 6.5 and 7.0 in fig 3 and fig 4 all gave a slope of 1 confirming 1 mole of both ligands were involved in the interactions. Combining these results with metal variation plots in the presence of synergist in figure 5b, the probable reaction during the extraction process can therefore be represented as:

$$Ni^{2+} + H_2BuEtP(o) + HBuP(o) \leftrightarrow Ni(HBuEtP)(BuP)(o) + 2H^+ \quad (7)$$

where [HBuP] is constant and incorporated in $K_{Niex2}$. The distribution ratio $D_{Ni^{2+}} = [(Ni(HBuEtP).BuP(o))/[Ni^{2+}]]$, on substitution into Eqn. (8) gives,

$$\log D_{Ni^{2+}} = \log K_{Niex2} + \log[H_2BuEtP] + 2pH \quad (9)$$

In this study, it was observed that the extraction of Ni(II) ion increased with an increase in pH and reached a maximum at pH of 8.78 after that further increase in pH resulted in a decrease in the percentage extraction of Ni$^{2+}$. Quantitative percentage extraction of 98.87% was obtained at pH of 8.78. The partition coefficient $K_{NiD1}$ is given by $K_{NiD1} = [Ni(HBuEtP)]_{o}/[Ni(HBuEtP)]$ for which a value of 1.89 ± 0.05 was determined from the graph. The pH$_{50}$ was also found to be 7.14 ± 0.01. Data of all the extraction processes are recorded in Table 1.

**Effect of addition of HBuP on the distribution of Ni(II)**

Quantitative percentage extraction of 99.11% at pH of 7.85 was obtained and there was a shift of pH$_{50}$ to 5.51 ± 0.10, a

Figure 2. kgD-lg [H$_2$BuEtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions at constant pH of 6.7, 7.3, and 7.8 in the absence of synergist

Figure 3. kgD-lg[H$_2$BU EtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions into chloroform solutions of ligands with [HBuP] kept constant (5 x 10$^{-3}$ M)

Figure 4. lgD-lg[H$_2$BuEtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions into chloroform solution of ligands with [H$_2$BuEtP] was kept constant (2.5 x 10$^{-2}$ M)

Figure 5. lg D-log [Ni(II)] plot of extraction of Ni(II) from buffer solutions into chloroform solutions of ligand 0.05 M H$_2$BuEtP (b) mixture of 0.05 M H$_2$BuEtP and 0.05 M HBuP in a 9:1 ratio

---

**Figure 2.** kgD-lg [H$_2$BuEtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions at constant pH of 6.7, 7.3, and 7.8 in the absence of synergist

**Figure 3.** kgD-lg[H$_2$BU EtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions into chloroform solutions of ligands with [HBuP] kept constant (5 x 10$^{-3}$ M)

**Figure 4.** lgD-lg[H$_2$BuEtP] plot of extraction of 8.52 x 10$^{-4}$ M of Ni(II) from buffer solutions into chloroform solution of ligands with [H$_2$BuEtP] was kept constant (2.5 x 10$^{-2}$ M)

**Figure 5.** lg D-log [Ni(II)] plot of extraction of Ni(II) from buffer solutions into chloroform solutions of ligand 0.05 M H$_2$BuEtP (b) mixture of 0.05 M H$_2$BuEtP and 0.05 M HBuP in a 9:1 ratio
Table 1. Extraction data of the influence of pH of aqueous phase on the extraction of \(8.52 \times 10^{-4}\) M of Ni(II) into organic solutions of 0.05 M H\(_2\)BuEtP and mixture of 0.05 M H\(_2\)BuEtP-0.05 M HBuP (9:1) ratio at room temperature of 30 °C (ionic strength of 0.1 M NaClO\(_4\))

<table>
<thead>
<tr>
<th>Organic Phase</th>
<th>(\lg K_{ex})</th>
<th>pH(%)</th>
<th>(\lg K_D)</th>
<th>Species extracted</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05 M H(_2)BuEtP</td>
<td>-12.39 ± 0.64</td>
<td>7.14 ± 0.10</td>
<td>1.89 ± 0.05</td>
<td>Ni(HBuEtP)(_{2(o)})</td>
</tr>
<tr>
<td>0.05 M H(_2)BuEtP-0.05 M HBuP (9:1) mixture</td>
<td>-10.57 ± 0.52</td>
<td>5.51 ± 0.10</td>
<td>1.89 ± 0.02</td>
<td>Ni(HBuEtP)(BuP)(_{9(o)})</td>
</tr>
</tbody>
</table>

Effect of mineral acids, anions and auxiliary complexing agents on the distribution of Ni(II)

Results presented in fig 6 show that extraction of nickel is masked in the presence of mineral acids as percentage extraction was less than 10% even in the presence of the HBuP at all concentrations of the various mineral acids used in this study as was observed in the extraction of Pb(II) with most mineral acids with same ligand system.\(^8\) This can be attributed to the formation of unextractable hydrophilic ion pair nickel anionic species in acidic media even in the presence of H\(_2\)SO\(_4\) which could be responsible for poor percentage extractions in the acidic regions in plots of extraction of Ni(II) ions in buffered media shown in Fig 1.

**Figure 6.** Plot of % extraction of \(8.52 \times 10^{-4}\) M Ni(II) from mineral acid solutions into (a) 0.05 M H\(_2\)BuEtP solution (b) 0.05 M H\(_2\)BuEtP-0.05 M HBuP (9:1) solution

The effect of anions on the distribution of Ni(II) ions in Fig. 7 indicate that chloride ion show the least enhancing effect (with an highest % extraction of 30) in the extraction of Nickel with the ligand H\(_2\)BuEtP while phosphate ion gave the highest percentage extraction (99.43%) of Ni(II). In most cases, at anion concentrations above 0.01M the percentage extraction starts to decrease. This trend was also observed in the mixed ligands organic phase. The results in fig 7b clearly show that a mixture of H\(_2\)BuEtP- HBuP/CHCl\(_3\) was a better extractant than H\(_2\)BuEtP/CHCl\(_3\) for Ni(II) ions, as the following increases in percentage extraction was observed; Cl\(^-\) (29.18% to 58.71%), I\(^-\) (56.8% to 97.5%), PO\(_4^2-\) (95% to 99.9%), SO\(_4^{2-}\) (41.83% to 67.05%), NO\(_3^-\) (47.68% to 95.2%) and CH\(_3\)COO\(^-\) (85% to 90.59%) at 0.01M concentration of ions. Most of the results on the effect of these anions follow trends observed in the distribution of Pb(II) using the same ligand system.\(^8\)

From the results presented in Fig 7, the influence of auxiliary complexing agents on the extraction of Ni(II) ions was not appreciable as all percentage of extraction was less than 50% at all concentrations of complexing agents used for the study except for F\(^-\) ion at 0.5M when H\(_2\)BuEtP only was used. There was an increase in % extraction when a mixed ligand H\(_2\)BuEtP/HBuP organic phase was used as > 50% extraction of Ni(II) was obtained in all complexing agents except EDTA. The % extraction decreases at concentrations above 0.005M of complexing agents in both type of ligand system, except for fluoride and tartrate ions. For fluoride ion, the % extraction increases steadily from 44.1% at 0.0005M to 99.6% at 0.5M whilst, for tartrate, from 52.5% at 0.0005M to 70.3% at 0.1M.
we conclude that extraction of Ni(II) ions with the ligand alone or its mixture with HBuP is better in alkaline medium.

From results of the effect of anions and auxiliary complexing agents on the extraction of Ni(II) ions, we may confirm that the mixture of ligands H₂BuEtP/HBuP was a better extractant for Ni(II) ion than H₂BuEtP alone collaborating the calculated physicochemical parameter values from distribution of Ni(II) ions in buffered media.

Though extraction of Ni(II) ions can be obtained in acidic pH between 4 – 6 with a mixture of the ligands, the optimal conditions for the extraction of Ni(II) ions was in the mixture of ligands H₂BuEtP/HBuP organic phase in the presence of phosphate ion at ≤ 0.01M, acetate ion at ≤ 0.1M and fluoride ion at ≤ 0.5M in near neutral to alkaline condition were ≥ 99% extraction of Ni(II) ions was obtained.

Acknowledgement
The authors wish to thank the University of Port Harcourt, Alexander von Humboldt Stiftung, Germany and Prof. Karsten Gloe for research assistance to BAU.

References

Received: 01.10.2012.
Accepted: 15.10.2012.