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Tantalum Recycling by Solvent Extraction: Chloride Is Better than Fluoride

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Abstract: The recycling of tantalum (Ta) is becoming increasingly important due to the criticality of its supply from a conflict mineral. It is used extensively in modern electronics, such as in capacitors, and so electronic waste is a potentially valuable secondary source of this metal. However, the recycling of Ta is difficult, not least because of the challenges of its leaching and subsequent separation from other metals. In this work, we show that Ta(V) halides, such as TaCl$_5$ and TaF$_5$, which can potentially be accessed from Ta metal upon acid halide leaching, can be recovered by solvent extraction using a simple primary amide reagent. The need for high halide concentrations in the aqueous phase implies the formation of the hexahalide salts [TaX$_6$]$^-$ (X = F, Cl) and that an anion-swing mechanism operates. While extraction of the fluorides is poor (up to 45%), excellent extraction under chloride conditions is found (>99%) and presents an alternative route to Ta recycling.

Keywords: tantalum; solvent extraction; WEEE; recycling; chemical separation

1. Introduction

The third-row transition metal element tantalum is critical to modern consumer technologies, finding extensive use as a component in capacitors for the microelectronics industry. Approximately 40% of the world’s tantalum production is used in this application, but its recycling is limited, at only 1% [1]. This is surprising, as electronic waste (e-waste) is becoming one of the fastest growing waste categories worldwide [2,3]. Furthermore, it was found that metal recovery from secondary resources is gradually becoming more cost-effective than virgin mining, especially as the volume of e-waste is projected to increase substantially; over the next few decades, new technologies will become cheaper and more accessible, driving up the rate of device obsolescence [4,5]. With the best estimates suggesting that only 20% of e-waste is properly recycled [5], the development of sustainable, efficient hydrometallurgical techniques for the recovery of these critical resources offers huge potential to reduce the environmental impact of e-waste, lower energy costs, and move towards a more circular economy [6]. However, this depends on the availability of inexpensive reagents for both leaching and separation processes.

The status of tantalum as a metal derived from a conflict mineral [7], combined with its low abundance in the earth’s upper crust means there is a need to design a sustainable closed-loop process to recycle tantalum from waste capacitors. Due to its very low solubility in typical mineral acids such as HCl, H$_2$SO$_4$, and HNO$_3$ compared with other d-block elements, current hydrometallurgical processes for tantalum involve the use of highly corrosive and toxic solutions of concentrated HF, combined with H$_2$SO$_4$ or HCl in order to generate soluble anionic metalates, such as [TaF$_6$]$^-$ and [TaF$_7$]$^{2-}$ [8–11].
As such, there is a need to develop more sustainable processes which can not only efficiently recycle tantalum from secondary sources but do so using milder reagents [12–15].

Recent efforts towards fluoride-free tantalum recovery processes have focused on alkaline matrices to generate polyoxotantalates that are recovered at elevated temperatures by solvent extraction [16–18]. Ionic liquids (ILs) have also been investigated as potential extractants for tantalum recovery. Methylisobutylketone (MIBK)-derived ILs were shown to efficiently extract [TaF7]5− from concentrated sulfuric acid solutions with excellent selectivity for tantalum over other, potentially interfering metals [19]. In this case, it was thought that this enhanced extraction compared with MIBK alone could be due to greater solubility of water in the IL phase, overcoming the prerequisite to displace the hydration sphere prior to transport of the metalate into nonpolar solvents.

To our knowledge, there are very few reports that describe the extraction of tantalum from chloride-based media; acids such as HCl were tested in the extraction of TaF5, but in these cases, significant solubility issues were seen [20]. Despite the limited speciation data available for tantalum in halide acids, previous theoretical studies have postulated that TaCl5 could behave in a similar manner to TaF5 in aqueous solution, so forming halidometalates such as TaCl5− in the presence of excess chloride ion [21,22].

We have shown previously that the primary amide, L (Figure 1), forms hydrophobic supramolecular assemblies with metalates, including [AuCl4]−, which allows for e-waste separations using anion-exchange solvent extraction processes [23]. In this work, we describe the transport of tantalum as its halometalate TaCl5− into a toluene organic phase using the simple primary amide reagent L from acidic solutions in the presence of chloride. Subsequent back extraction (stripping) of the metalate into a fresh aqueous phase is possible with water or dilute HCl.

![Figure 1. Structure of 3,5,5-trimethylhexanamide L.](image)

2. Experimental

2.1. Materials and Instruments

Unless otherwise stated, all solvents and reagents were purchased from Sigma-Aldrich, Fisher scientific UK, Alfa Aesar (Heysham, UK), Acros Organics (Geel, Belgium), or VWR international (Lutterworth, UK) and used without further purification. Deionised water was produced using a Milli-Q purification system. The syntheses of 3,5,5-trimethylhexanamide, L and [H(C11H23NO)2]2[SnCl6] were carried out according to the literature [23,24].

ICP-OES analysis was carried out on a Perkin Elmer Optima 8300DC Inductively Coupled Plasma Optical Emission Spectrometer. Samples in 1-methoxy-2-propanol were taken up by peristaltic pump at a rate of 2.0 mL min−1 into a Gem Tip cross flow nebuliser and a glass cyclonic spray chamber. Argon plasma conditions were: 1500 W RF forward power; argon gas flows of 17, 1.0, and 0.50 L min−1 for plasma, auxiliary, and nebuliser flow, respectively. ICP-OES calibration standards were obtained from SCP science.

Karl-Fischer water content determinations were carried out on a Metrohm 831 KF Coloumeter with Hydranal Coulomat AG as the reagent mixture.

2.2. Preparation of TaCl5 and TaF5 Solutions

A 0.12 M stock solution of TaX5 (X = F, Cl) was prepared in 12 M HCl at 25 °C. This stock solution was then diluted to 0.01 M TaX5 in 1 M HCl with varying solutions of lithium chloride in deionised water. A colourless precipitate formed over several weeks in dilute HCl solutions of TaCl5, and so solutions were used immediately after diluting from the 12 M stock solution. A colourless precipitate
formed overnight from stock solutions of TaF_5 in 12 M HCl, and so solutions were therefore diluted and used immediately.

2.3. General Solvent Extraction Procedure

An aqueous solution of TaX_5 (0.01M, X = F, Cl) in 1 M HCl, 0-11 M LiCl (2 mL) was contacted with a toluene organic phase (2 mL) containing 3,5,5 trimethylhexanamide L (0.1 M) and stirred (1 h, 1000 rpm, 25 °C). The phases were separated physically, and samples from the organic phase taken and diluted with 1-methoxy-2-propanol for ICP-OES analysis. Samples of the stock solution were analysed by ICP-OES to confirm mass balance and percentage extraction. Samples of the post extraction aqueous phase could not be analysed by ICP-OES due to the high salt content suppressing the tantalum wavelengths. Samples from relevant phases were also taken for water concentration determination and NMR analysis as required.

2.4. NMR Data for [H(C_{11}H_{23}NO)]_2[SnCl_6].

^1H NMR (CDCl_3, 500 MHz, ppm): δ_H 10.17 (s, broad, 1H), 3.27 (s, 6H, NCH_3), 2.76 (dd, 2H, CH_2CO, J = 14.2, 6.7 Hz), 2.65 (dd, 2H, CH_2CO, J = 14.2, 8.7 Hz), 2.18-2.08 (m, 2H, CH(CH_3)), 1.32 (dd, 2H, CH(CH_3), J = 14.1, 3.3 Hz), 1.21 (dd, 2H, CH(CH_3), J = 14.1, 7.1 Hz), 1.05 (d, 6H, CH(CH_3), J = 6.6 Hz), 0.90 (s, 18H, C(CH_3)_3).

^13C{^1H} NMR (C_6D_6, 126 MHz, ppm): δ_C 176.34, 50.65, 41.33, 38.90, 31.09, 29.99, 28.12, 22.39.

3. Results and Discussion

3.1. Solvent Extraction of Tantalum Pentachloride

The dissolution of TaCl_5 in HCl only occurs at high concentrations of HCl (12 M). Initial solvent extraction procedures were therefore developed by contacting a 0.1 M solution of L in toluene with a 0.01 M solution of TaCl_5 in 12 M HCl. However, at this high HCl concentration, no appreciable tantalum extraction is seen, and it is likely that the amide L is instead protonated and transported into the aqueous phase (Figure 2). Upon addition of LiCl to the TaCl_5 solution at 12 M HCl, the concentration of Cl^- is increased further, although this results in the out-gassing of HCl from the solution. Under these conditions, low to moderate quantities (30%-50%) of Ta are transferred to the organic phase (Figure 3). As tantalum begins to be extracted at 3–6 M LiCl, small quantities of a precipitate forms which dissolve at higher LiCl concentrations. Quantitative ^1H NMR analysis of this third phase dissolved in deuterated dimethyl sulfoxide shows that no L from the organic phase is present, and ICP-OES analysis shows it comprises tantalum and lithium only, so is potentially LiTaCl_6 or a similar complex. Furthermore, L does not transfer significant quantities (<10 ppm) of Li into the organic phase, as the concentrations of L or LiCl are varied.
Figure 2. Extraction of tantalum from aqueous solutions of TaCl₅ in 12 M HCl with varying concentrations of LiCl into a toluene solution of L. Conditions: TaCl₅ (0.01 M) in 12 M HCl with 0–3 M LiCl (2 mL) stirred with L (0.1 M) in toluene (2 mL); phases contacted for 1 h at RT with magnetic stirring. Experiments performed in duplicate and reported as an average.

Figure 3. Transport of tantalum from aqueous solutions of TaCl₅ in 1 M HCl with varying concentrations of LiCl into a toluene solution of L. Conditions: 0.01 M TaCl₅ diluted in 1 M HCl, 1–11 M [LiCl] (2 mL), stirred with L (0.1 M) in toluene (2 mL); phases contacted for 1 h at RT with magnetic stirring. Interpolation used to aid the eye only. Experiments performed in duplicate and reported as averages.

The need for high levels of LiCl to promote Ta extraction suggests that either the formation of the anion TaCl₆⁻ is a prerequisite for transport into the organic phase, or a salting-out effect is operating. To understand this, experiments were undertaken which vary the concentration of BF₄⁻ (using NaBF₄) as a non-coordinating anion instead of Cl⁻ and show that negligible Ta transfer to the organic phase occurs. This confirms that TaCl₅ is not simply “salting-out” to the organic phase and that a source of chloride is necessary to drive [TaCl₆]⁻ formation; even so, the high salt content may aid extraction by reducing the concentration of free water in the aqueous phase [25]. Importantly, Ta was readily back-extracted (~80%) from the organic phase to a fresh aqueous phase with either water or 1 M HCl; a white precipitate forms with water, while no precipitate is immediately observed in the strip solution with 1 M HCl.
3.2. Structure Elucidation

From the data above, it is apparent that Ta is transported into the organic phase as its ion pair [HL][L]n[TaCl6]. To further interrogate the organic-phase speciation, slope analysis of Log D against Log [L] was carried out (Figure 4), which results in a L:Ta ratio of approximately 2 (slope of 1.71 ± 0.13), suggesting the formation of an ion pair such as [HL2][TaCl6] in the organic phase. This type of behaviour is similar to that seen previously for Au extraction using L in which experimental and computational analysis showed that two amides chelate a proton through the oxygen atoms to form a charge-diffuse cation that can interact with diffusely charged metalate anions through classical (N-H···X) and non-classical (C-H···X) hydrogen bonds [23,24].

![Figure 4. Slope analysis for the transport of Ta with L. Conditions: 2 mL 0.01 M TaCl5 in 1 M HCl, 11 M LiCl, contacted with 2 mL of 0.01 to 0.1 M L in toluene for 1 h at RT with magnetic stirring.](image)

Direct structural characterisation of the extracted species has so far proven inconclusive, although 1H NMR studies provide an indication as to how the receptor [HL2]+ is interacting with the metalate (Figure 5). As the concentration of tantalum in the organic phase increases, the two signals at 4.61 and 6.46 ppm associated with the NH2 protons coalesce and shift downfield to 9.63 ppm. Variable temperature NMR spectra (Figure S1) reveal an additional signal that is attributable to associated water or hydronium, which coalesces with the amide protons upon Ta loading. There is also a large downfield shift of the signals at 1.59 and 1.87 ppm, associated with the diastereotopic CH2 protons adjacent to the amide group, to 2.49 and 2.56 ppm respectively. These changes in the 1H NMR spectra for Ta-loaded L are similar to those seen for the solid third phase formed between [SnCl6]2− and a tertiary amide version of L (Figures S2 and S3). In this latter case, the solid-state structure showed that the two amide ligands chelate the single proton through the amido oxygen atoms, with the cation subsequently interacting with the outer-sphere of the [SnCl6]2− octahedron through non-classical C-H hydrogen bonds [24,26,27]. The similarity in the 1H NMR spectra for Ta/L, and also a relatively large downfield shift of the carbonyl carbon from 174 ppm to 180 ppm in the 13C NMR spectrum (Figure S4) upon Ta loading strongly suggest that a similar structure is formed in the organic phase, with no direct interaction of the amido-oxygen donor with the Ta centre; i.e., an amide complex such as TaCl5(L) is not formed.
Karl-Fischer titrations were undertaken to determine the role of water in the extracted species. It is seen that as more tantalum transfers to the organic phase, the concentration of water does not significantly increase, ruling out a micellar extraction mechanism (Figure 6). However, at even higher concentrations of L where the concentration of tantalum in the organic phase does not change, more water is transported, suggesting that water transport into the organic phase is associated with the concentration of L.

Figure 5. $^1$H NMR spectra of (a) 0.1 M L in C$_6$D$_6$; then 0.1 M L in C$_6$D$_6$ after contact with 0.01 M TaCl$_5$ in 1 M HCl and: (b) 5 M LiCl; (c) 6 M LiCl; (d) 7 M LiCl; (e) 8 M LiCl; (f) 9 M LiCl.

Figure 6. Karl-Fischer water measurements (1.0 mL samples from organic phase analysed for H$_2$O; concentrations of Ta and H$_2$O are on the same scale). Interpolation used to aid the eye only. Experiments performed in duplicate and reported as an average.
During the slope analysis and Karl-Fischer titrations, a solid precipitate is seen at concentrations of L below 0.02 M, with more precipitate forming at lower concentrations of L. As with above, quantitative analysis of the organic phases by $^1H$ NMR spectroscopy shows that the ligand concentration remains constant (i.e., no loss of L to solid 3rd phase or the aqueous phase); furthermore, this phenomenon was irreproducible in the absence of either TaCl$_5$ or L.

3.3. Solvent Extraction of Tantalum Fluoride

As current commercial processes for tantalum recovery involve the use of hydrogen fluoride to leach the metal as its fluoride salts, it is important to compare how the chloride process described above would compare with its fluoride equivalent. However, all attempts to extract TaF$_5$ instead of TaCl$_5$ were unsuccessful under the conditions used, with negligible extraction seen between 0–12 M HCl and 0–18 M H$_2$SO$_4$ with L; at high acid concentrations L is lost to the aqueous phase. When the HCl concentration is 1 M and the concentration of LiCl is varied, optimal extraction of the tantalum is 45% at 5 M LiCl before trending downwards upon further increase of LiCl (Figure 7).

Figure 7. Extraction of tantalum from aqueous solutions of TaF$_5$ in 1 M HCl with varying concentrations of LiCl into a toluene solution of L. Conditions: 0.01 M TaF$_5$ diluted in 1 M HCl, 1–11 M [LiCl] (2 mL), stirred with L (0.1 M) in toluene (2 mL); phases contacted for 1 h at RT with magnetic stirring. Interpolation used to aid the eye only. Experiments performed in duplicate and reported as averages.

In these experiments, no solid third phase forms at any stage, and quantitative $^1H$ NMR experiments confirm that no amide is lost to the aqueous phase. These data contrast with the experiments using TaCl$_5$, perhaps because potential fluorometalates such as [TaF$_6$]$^-+$ or [TaF$_5$Cl]$^-+$ that would be formed in the aqueous phase, are smaller and have more charge dense anions with respect to [TaCl$_6$]$^-+$, and therefore will have higher hydration energies. Poor Ta extraction at high LiCl concentrations is likely to result from the competitive transport of chloride by L into the organic phase.

4. Conclusions

The simple primary amide, L, shows excellent performance for tantalum recovery by solvent extraction under high chloride conditions. The extraction mechanism is identified as similar to that seen for the recovery of gold by L, in which charge-diffuse protonated receptors are formed that preferentially interact with the charge diffuse monoanionic metalate [AuCl$_4$]$^-+$ [23,24]. $^1H$ and $^{13}$C NMR data are consistent with a strong outer-sphere interaction between [HL$_2$]$^+$ and [TaCl$_6$]$^-+$. In contrast, amide L is a poor reagent of choice when attempting to transport TaF$_5$ under high chloride conditions, likely due to a combination of competitive chloride extraction and significantly weaker interactions.
between [HL$_2$]$^+$ and more charge-dense species in the aqueous phase, such as [TaF$_5$Cl]$^-$. This latter aspect is evident from a comparison of the X-ray crystal data for [TaCl$_6$]$^-$ and [TaF$_6$]$^-$, which shows a 1 A decrease in the diameter of the Ta complex moving from chloride to fluoride [28,29]. It is therefore anticipated that L would be a poor extractant for Ta when employed under solely fluoride conditions, in which species such as [TaF$_5$]$^-$, and even the dianion [TaF$_7$]$^{2-}$ may be present. Even so, the process described here represents an alternative, fluoride-free route to recycling Ta from waste electronics, using milder reagents than the current commercial methods.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2075-4701/10/3/346/s1. Figure S1: Variable temperature $^1$H NMR spectra of 0.1 M L in d$_6$-toluene after contact with 0.01 M TaCl$_5$ in 1 M HCl and 9 M LiCl from ~40 °C to 60 °C. Presence of additional water or hydronium signal is revealed amongst the signals assigned to the amide protons (between 8.5 ppm and 10 ppm) when the sample is cooled. Y-axes are offset for clarity. Figure S2: $^1$H NMR spectra of a C$_{11}$H$_{23}$NO (top) and [H(C$_{11}$H$_{23}$NO)$_2$][SnCl$_4$] (bottom) in CDCl$_3$. Figure S3: $^1$C($^1$H) NMR spectra of a C$_{11}$H$_{23}$NO (top) and [H(C$_{11}$H$_{23}$NO)$_2$][SnCl$_4$] (bottom) in CDCl$_3$. Figure S4: $^{13}$C($^1$H) NMR spectra of 0.1 M L in C$_6$D$_6$ (top) and 0.1 M L in C$_6$D$_6$ after contact with 0.01 M TaCl$_5$ in 1 M HCl and 9 M LiCl (bottom).

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