

Article

Hydrometallurgical Recovery of Cobalt(II) from Spent Industrial Catalysts

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Abstract: The work presents studies on the application of hydrometallurgical recovery of cobalt(II) from solutions after leaching spent industrial catalysts used in process of hydrodesulfurization. A four-stage process was proposed, which consists of: leaching, precipitation of metal hydroxides accompanying Co(II), extraction of Co(II) with *bis*(2,4,4-trimethylpentyl)phosphinic acid and Co(II) stripping from the organic phase. The results indicate that by using the proposed method it is possible to leach Co(II) and Mo(VI) from spent catalyst, and remove main impurities such as Al(III), Fe(III) in hydroxide precipitation step and separate Co(II) from Mo(VI) by extraction and stripping.

Keywords: spent catalyst; cobalt recovery; hydrodesulfurization catalyst; leaching; hydrometallurgy; liquid-liquid extraction; Cyanex 272

1. Introduction

The search for new, alternative metal sources is not only driven by scientific curiosity but nowadays also an industrial necessity. The obligation is conditioned by economic and financial need, as well as, a conversion from the, so-called, linear economy to the world circular economy. Such metal sources include, among others electronic wastes (known as waste electrical and electronic equipment WEEE or e-waste, including waste printed circuit boards (WPCBs)), spent automotive catalysts, electroplating wastes, or a wide range of catalysts, especially from the chemical and petrochemical industry [1–4].

It should be noted that over 60% of all chemicals are produced using catalysts in 90% of all industrial chemical processes [5]. Wherein 80% of catalysts are in the solid form, less common are homogeneous and biocatalysts, about 17% and 3%, respectively [6]. Due to the widespread use of catalysts, as well as their limited lifetime and the possibility of regeneration [7], the problem of their utilization is widely discussed in the literature [8–10]. Especially, that according to the United State Environmental Protection Agency (EPA) catalysts are classified as “hazardous waste as one posing a substantial or potential hazard to human health and the environment” [11]. Besides, many studies are carried out on the recovery of valuable elements from spent catalyst, especially the platinum-group metals (PGMs) [12], the rare-earth elements (REEs) [13], and the variety of transition metals [14]. Several methods are proposed and successfully used in academic and industrial practice. The conventional approaches used in metal reclamation as pyrometallurgy [15] and hydrometallurgy (leaching, ion exchange, extraction) [16–18], as well as new methods, include bioleaching [19]. As was mentioned above such methods are applied also in commercial solutions, i.e., the Dutch company

Moxba-Metrex [20], the Germany company Nickelhütte Aue GmbH [21], or the Belgian company Sadaci N.V. [22], etc., described in detail in the work [23]. The selection of the right method depends on many factors [24]. Generally, the pyrometallurgical processes are appropriate for the treatment of high-grade ores, while hydrometallurgical processes for the treatment of low-grade or complex ores. It should be emphasized that in the case of high content of metals both methods could be used, although pyrometallurgy is more economic. At the same time, this method is unsuitable either for low content of metals in the material because a large amount of energy is required to melt associated gangues, or for complex ores due to difficulties in separation of the components. Moreover, there is no possibility to separate chemically similar metals in contrast to hydrometallurgy. Since catalysts are usually secondary ores containing small amounts of metals, most of the techniques proposed are based on hydrometallurgical processes [25,26]. Additionally, hydrometallurgy has some advantages such as low cost requirements, possible recovery of leachants and no air pollution, in contrast to emission of hazardous gases (SO_2 , dioxins, furans) during pyrometallurgical processes. For sustainable development, bioleaching of metals is also recommended [27]. It seems to be promising due to its' environmental friendliness, and low cost and low energy requirements. However, the use of microorganisms, bacteria or fungi means that this process requires a longer time in comparison to the conventional chemical leaching, and strongly depends on atmospheric conditions.

Although the recovery of metals from spent catalysts has been known for many years, new recovery methods are still being sought, considering various industrial catalysts, as well as operating conditions. Therefore, taking into account the advantages and disadvantages of the available recovery methods, the work aims in the investigation of the possibility of the hydrometallurgical recovery of cobalt(II) from spent industrial hydrodesulfurization catalyst, with the optimal selection of leaching and extraction processes applied.

2. Results and Discussion

CoMo catalysts are prepared by successive impregnation steps of the alumina support with metal solutions. As a result of the impregnation, Co exists as a thin layer between the Mo layer and the Al support. Usually cobalt is deposited on alumina support first, and then molybdenum (in the oxidized form of Mo(VI)) is impregnated on the Al-Co material [28]. Spent hydrodesulfurization (HDS) catalysts can be treated as a valuable source of cobalt. Thus, hydrometallurgical recovery of Co(II) from CoMo catalyst was proposed in this work in a four-step process shown in Figure 1.



Figure 1. Four-step process for hydrometallurgical recovery of Co(II) from CoMo catalyst of hydrodesulfurization.

2.1. Leaching of Co(II)

The first step of spent catalyst treatment was leaching of metals from a ground CoMo catalyst with sulfuric acid solutions or mixtures of sulfuric acid and hydrogen peroxide. It should be emphasized that, although some authors recommend leaching from HDS catalysts with mixtures of mineral acids (e.g., $\text{HNO}_3/\text{H}_2\text{SO}_4/\text{HCl}$ [29] or $\text{HNO}_3/\text{H}_2\text{SO}_4$ [30]), in this case sulfuric acid was chosen as a leaching agent to provide appropriate solution for the third step of the planned process, i.e., extraction with acidic extractant Cyanex 272. It should be noted that most data in the literature refer to sulfate solutions, while the application of Cyanex 272 in chloride and nitrate solutions is possible but less effective [31]. The results of leaching efficiency (LE), for different sulfuric acid concentrations, are shown in Figure 2.

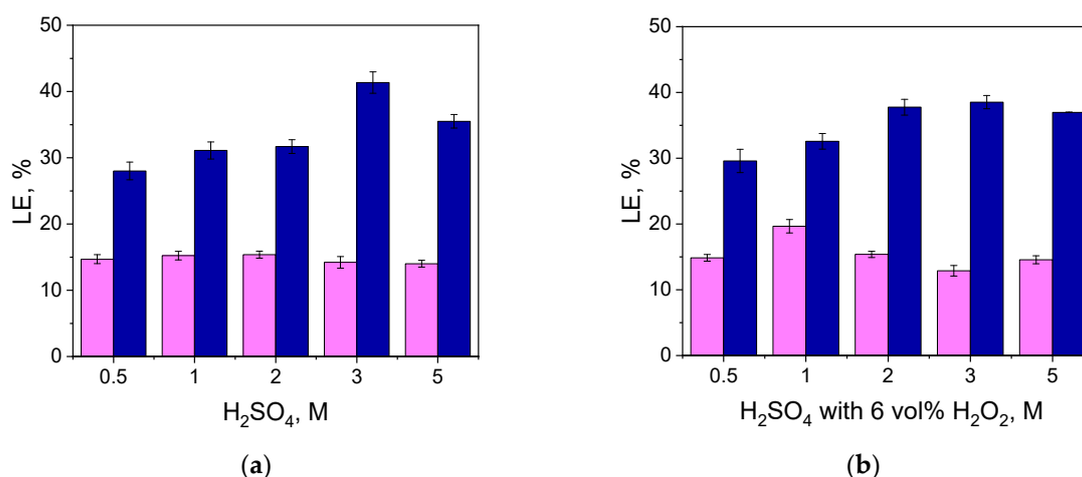


Figure 2. Efficiency of Co(II) (■) and Mo(VI) (■) leaching from spent industrial catalyst with (a) sulfuric acid solutions, (b) sulfuric acid solutions with addition of 6 vol% of 30% H₂O₂ (50 °C, S/L = 1/20 g/cm³, particle size of catalyst <45 μm, stirring 300 rpm).

The leaching efficiency (LE) of cobalt(II) and molybdenum(VI) was calculated according to the following equation:

$$LE = \frac{m_{leach}}{m_0} \cdot 100\% \quad (1)$$

where m_0 , m_{leach} stand for mass of metal ions in the spent catalyst before leaching and in the leach solution after leaching, respectively. Initial mass of metal in the catalyst was estimated on the base of metal content (u_M) according to XRF analysis ($u_{Co} = 0.124$, $u_{Mo} = 0.359$) and mass of the sample taken to leaching (m_{cat}) as follows:

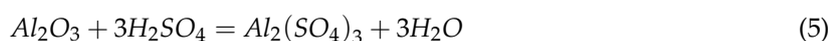
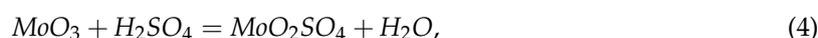
$$m_0 = m_{cat} \cdot u_M \quad (2)$$

The efficiency of Co(II) leaching with the solutions of sulfuric acid (in the studied range of concentrations) without and with the addition of H₂O₂ reached as maximum 20% (Figure 2). The leaching agents were more effective for Mo(VI) leading to the dissolution of 30–40% of this metal (Figure 2). An increase in acid concentration (between 0.5 and 5 M) did not affect significantly the amount of the leached Co(II), while it increased the amount of Mo(VI) leached.

Also, other researchers have indicated that direct leaching of the spent catalyst with sulfuric acid was not effective, whereas the combination of sulfuric and nitric acid significantly enhanced the recovery of Mo(VI) and Co(II) [30]. However, they reported that at S/L = 1/10, 50 °C and 400 rpm cobalt(II) was better extracted than Mo(VI), and the leaching efficiency with 1.5 M H₂SO₄ reached 55.3 and 19.3% after 4 h of reaction time. The mixture of HNO₃/H₂SO₄ caused an increase in metal leaching up to almost 80% of Mo(VI) and more than 90% of Co(II) (after 5 h). Furthermore, it was reported by other researchers [29] that the leaching efficiency of Mo(VI) and Co(II) with mixtures of HNO₃/H₂SO₄/HCl and HNO₃/H₂SO₄ acids are 86 and 87% for Mo(VI) and 91 and 93% for Co(II), respectively.

Although higher efficiency of the leaching could be obtained with the addition of HNO₃, in this work only sulfuric acid was chosen as the leaching agent because of further steps of Co(II) recovery, i.e., extraction and stripping that provide purified Co(II) electrolyte which can be used for electrowinning of metallic cobalt.

The reactions considered during leaching of these metals are as follows [28,32]:



The addition of an oxidant, i.e., H_2O_2 , did not cause a significant increase in the leaching efficiency. Either in the presence of hydrogen peroxide or without it, Co(II) concentration in the leach solutions and leaching efficiency of Co(II) were comparable. In the case of Mo(VI) leaching, the addition of H_2O_2 increased efficiency by 1.5 to 6 percentage points. As no significant effect of the H_2O_2 addition was observed, investigation of the temperature and time influence on metal leaching was carried out with sulfuric acid alone. Moreover, the use of H_2O_2 is disadvantageous due to the high price of the oxidant.

To achieve efficient Co(II) leaching, the operation was carried out in various temperatures (Figure 3). A slight increase in ion content of both metals (Co(II) and Mo(VI)) in the solution was observed with increasing temperature of leaching within the range from 23 to 95 ± 2 °C.

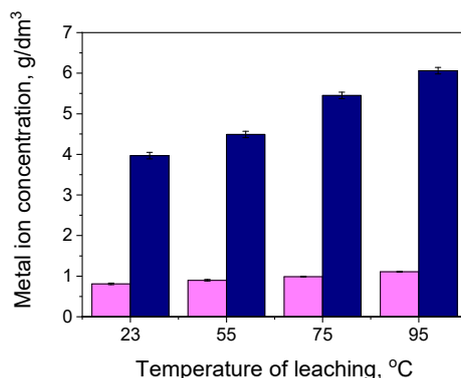


Figure 3. Dependence of Co(II) (■) and Mo(VI) (■) concentration in the leach solutions on temperature of leaching for 3 h with 0.5 M H_2SO_4 at $\text{S/L} = 1/20$ g/cm³.

The change in concentration of various metals in the leaching solution during 3 h of leaching with 0.5 M H_2SO_4 at 95 °C and $\text{S/L} = 1/20$ g/cm³ is shown in Figure 4. As Al_2O_3 is a support for the active metals, and the content of aluminium in the leached material is the highest of all the metals, the concentration of the dissolved Al(III) is also the highest (almost 10 g/dm³). However, concerning the initial content (Table 7), the loss of aluminium from the solid material of the catalyst is not as large as of cobalt, iron or copper (Table 1). Also, low efficiency of Al(III) leaching (about 10%) in contrast to 70–86% for Ni from spent catalysts ($\text{NiO}/\text{Al}_2\text{O}_3$) was obtained by Miazga and Mulak [33]. Moreover, Hamza et al. [28] reported poor leaching of Al(III) (about 5%) and they attributed it to worse accessibility of the catalyst support to the leaching agent. However, they mainly leached raw, not ground, material, while in our case aluminium is well-accessed because the catalyst is in the form of fine powder (particle size below 45 μm). Thus, not only a thin layer of cobalt and molybdenum but also aluminium are dissolved by the leaching agent.

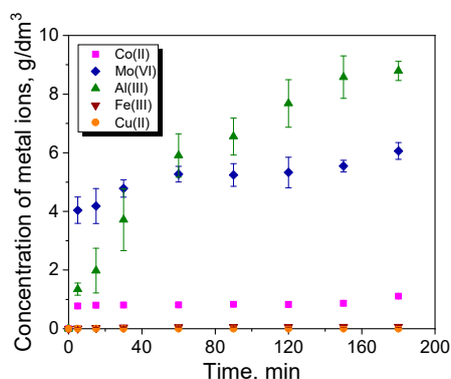


Figure 4. Change in concentration of metal ions in the leach solution during leaching for 3 h with 0.5 M H_2SO_4 at 95 °C, $\text{S/L} = 1/20$ g/cm³.

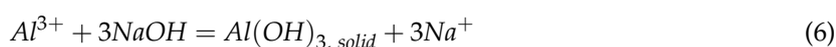
Table 1. An exemplary composition of the spent catalyst after leaching with 1 M H₂SO₄ at 55 °C, S/L = 1/20 g/cm³.

Chemical Element	Percentage Composition after Leaching, %	Loss in Element Content after Leaching, %
Al	43.6	19.0
Mo	35.3	14.9
Co	4.54	68.3
Fe	0.20	76.2
Cu	0.02	79.4

The results of XRF analysis of the samples after leaching with sulfuric acid confirmed depletion of more than 60% of cobalt, almost 60% of copper and 80% of iron. However, the concentration of Cu(II) or Fe ions in the leaching solution is low (Figure 4) because the initial content of these metals in the catalyst is small (Table 7).

2.2. Precipitation of Impurities

After leaching the solid residue was separated from the leach solution. The aqueous phase containing ions of various metals (i.e., Al(III), Fe(III), Cu(II), Ni(II), Co(II), Mo(VI)) was treated with NaOH solution to precipitate hydroxides of the metals accompanying Co(II), according to the reaction:



Metal content in the deposits precipitated from various leach solutions is presented in Table 2.

Table 2. Metal content in the selected deposits after precipitation with 30% NaOH solution.

Leaching Solutions	Contents in the Sediment after Precipitation, mg				
	Co(II)	Mo(VI)	Ni(II)	Cu(II)	Fe(III)
1 M H ₂ SO ₄	0.768	0.160	0.004	0.019	0.223
1 M H ₂ SO ₄ + H ₂ O ₂	0.857	0.180	0.004	0.016	0.106
5 M H ₂ SO ₄	0.182	0.620	0.000	0.002	0.013
5 M H ₂ SO ₄ + H ₂ O ₂	0.172	1.015	0.000	0.004	0.028

Along with Mo(VI) and Fe(III), also Co(II) is precipitated which is disadvantageous for Co(II) recovery due to loss of this metal. The main aim of the precipitation step is the removal of Al(III) before the extraction of Co(II). The smallest loss of Co(II) in the deposit (0.100 mg) is noted from the leachate of 0.5 M H₂SO₄, probably because the final pH of precipitation was equal to 3.5, while Co(II) precipitation increases with increasing pH. As Huang et al. [34] indicated, major cobalt precipitation occurred within a pH range of 5–7. However, the pH increase from 4 to 5 enhances substantially also molybdenum(VI) precipitation from the solutions [34]. Consequently, various ratios of Co(II) and Mo(VI) in the precipitate can result from changes in composition of the deposit, for example, from cobalt hydroxide (Co(OH)₂), through cobalt molybdenum oxide hydrate (CoMoO₆·0.9H₂O) to sodium cobalt molybdenum oxide (NaCo₂.31(MoO₄)₃) [34]. Thus, as Co(II) loss was the lowest, 0.5 M H₂SO₄ was selected as the most appropriate for the four-step process. Additionally, the use of low concentrated acid is advantageous from the environmental protection and economic point of view. An important aspect of precipitation is Al(III) removal from the leachate because at pH 5 (extraction pH) Al(III) would start to precipitate and disturb Co(II) extraction. Content of Al(III) in the solutions before and after precipitation with 30% NaOH is shown in Table 3.

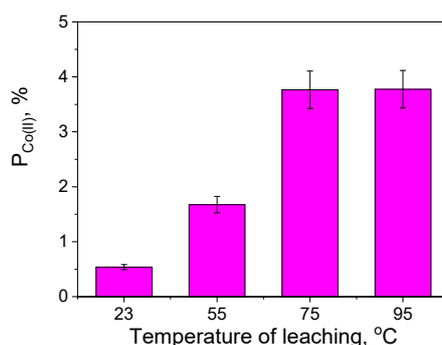
Table 3. Al(III) content in solution after leaching with 0.5 M H₂SO₄ and after precipitation with 30% NaOH.

Temperature of Leaching, °C	Al(III) Content in Solution, mg	
	After Leaching	After Precipitation
23	34.4	0.8
55	77.3	0.6
75	212.0	1.2
95	296.3	2.1

As shown in Table 3, precipitation with 30% NaOH leads to effective removal of Al(III) ions from the solution. However, it is not a selective operation because hydroxides of other elements are also precipitated. To estimate the loss of Co(II) during the precipitation, the efficiency of precipitation (P) was calculated as follows:

$$P = \frac{m_1}{m_2} \cdot 100\% \quad (7)$$

where: m_1 —mass of the precipitated Co(II), m_2 —mass of Co(II) before the precipitation. The efficiency of Co(II) precipitation depending on the temperature in which the previous step, i.e., leaching, was carried out, is shown in Figure 5.

**Figure 5.** The efficiency of Co(II) precipitation from solutions after leaching with 0.5 M H₂SO₄ at various temperatures.

The efficiency of Co(II) precipitation was calculated for the mass of the whole precipitate obtained during the precipitation. The highest Co(II) losses (about 4%) were found in the samples after leaching at 75 and 95 °C (Figure 5). In the samples after leaching at low temperatures (23 and 55 °C), the amount of precipitated Co(II) is low and does not exceed 2%. Additional XRF analysis (the results not presented in this work) showed that almost 60% of the mass of the precipitate is sulfur, which probably derives from sulfates from the leaching solution, and confirms that Co(II), Mo(VI), and Al(III) are deposited not only as hydroxides (Equation (6)) but also as mixed compounds [34,35], e.g., Co(SO₄)_i(OH)_j·kH₂O, MoO₂SO₄, 2Al₂O₃·SO₃·11H₂O, respectively. The XRF analysis confirmed that some amount of Co(II) is co-precipitated which causes loss of Co(II) from the solution forwarded to the next step—extraction.

2.3. Extraction of Co(II)

Various organic compounds were reported as extractants for Co(II) or Mo(VI) separation from leachates. For example, basic tertiary amine (Alamine 304) was proposed to extract Mo(VI) from H₂SO₄ at pH 1.8 [32]. Also, two-step extraction with neutral tributylphosphate (TBP) or basic *tris*-2-ethylhexylamine (TEHA) was shown to be effective for removal of Mo(VI) or Co(II), respectively, from HCl [36]. Another combination of extractants was proposed, i.e., trioctylphosphine oxide (TOPO) and tertiary amine Alamine 308, to extract Mo(VI) and Co(II), respectively, from 3 M HCl [37]. However, in this work, *bis*(2,4,4-trimethylpentyl)phosphinic acid (Cyanex 272) was chosen to separate Co(II)

from the leachate and next to transfer it to sulfate stripping solution. On the one hand, extraction with Cyanex 272 makes possible generate Co(II) electrolyte for further electrowinning, on the other hand, only one-step extraction reduces the amount of the organics used in the process. Our previous works have shown that Cyanex 272 or its sodium salt can selectively extract Co(II) in the presence of Ni(II) [38,39]. An effective Co(II) extraction from solution containing a hundredfold excess of Ni(II) has been confirmed also by other authors [40].

Thus, after removal of Al(III) and Mo(VI) from the aqueous phase, liquid-liquid extraction with 0.4 M Cyanex 272 to recover Co(II) and separate it from ions of other metals. Percentage extraction of Co(II) was calculated as follows:

$$E = \frac{m_{org}}{m_{aq,0}} \cdot 100\% \quad (8)$$

where $m_{aq,0}$, m_{org} stand for metal mass in the aqueous phase before extraction and in the organic phase after extraction, respectively. Results of Co(II) extraction are compared with the conditions (acid concentration) of the leaching stage, and shown in Figure 6.

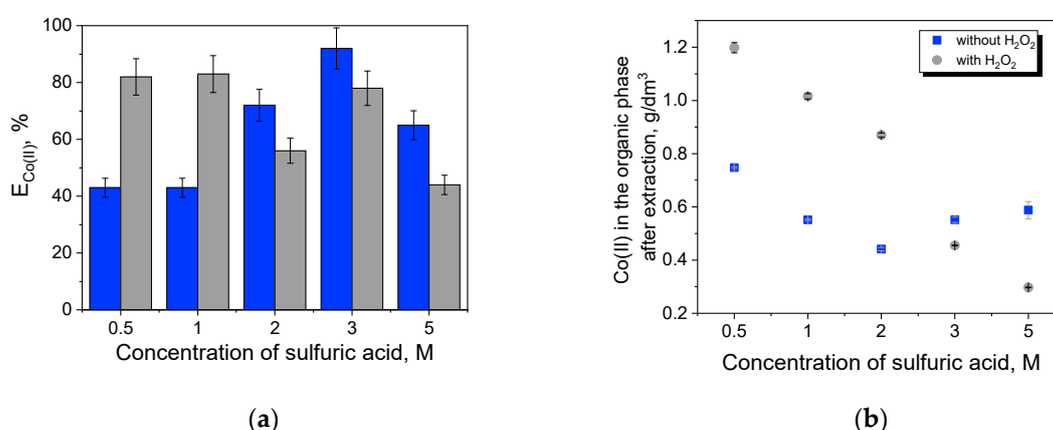


Figure 6. Dependence of (a) percentage extraction of Co(II), (b) concentration of Co(II) in the organic phase after extraction from solutions after leaching with (■) sulfuric acid or (●) sulfuric acid with H₂O₂ addition.

It was shown in our previous work that the best conditions for Co(II) extraction are at pH 5.2–5.5 [38]. Also, manufacturer of Cyanex 272 reports that at pH 5.5 maximum extraction of cobalt(II) from sulfate solution can be obtained, while ions of other metals present in our system extract efficiently at different pH values (Al at 2.8, Fe pH 2.1, Cu pH 4, Ni pH 7.5) [41]. Moreover, it was important to remove Mo(VI) and Al(III) before Co(II) extraction because they start to precipitate at pH 3 and the formation of the deposit becomes substantial at pH 5 [28,34]. The highest Co(II) extraction efficiency is noted when leaching was carried out with 3 M H₂SO₄, while for leach solutions with the addition of 6 vol% of 30% H₂O₂ percentage extraction of Co(II) exceeded 80% when 0.5, 1 and 3 M H₂SO₄ solutions were used. However, it must be emphasized that Co(II) concentrations in the organic phase after extraction are the highest when 0.5 M H₂SO₄ solution, either with H₂O₂ or without it, was applied (Figure 6b).

2.4. Stripping of Co(II)

After selective extraction of Co(II) to the organic phase, an effective stripping of Co(II) from the loaded Cyanex 272 to 1 M sulfuric acid solution was accomplished to form sulfate electrolyte solution. The final concentrations of Co(II) and Mo(VI) in the stripping aqueous phases in the selected experiments are shown in Table 4.

Table 4. Co(II) and Mo(VI) concentrations in the stripping phase (after the fourth step of leaching-precipitation-extraction-stripping process) for various conditions of the first step—leaching.

Leaching Conditions	Temperature of Leaching, °C	Final pH during Precipitation	Concentration of Mo(VI), g/dm ³	Concentration of Co(II), g/dm ³
2 M H ₂ SO ₄ + H ₂ O ₂	55	3.5	0.00	0.77
1 M H ₂ SO ₄	55	4	0.00	1.56
		5	0.00	2.23
1 M H ₂ SO ₄ + H ₂ O ₂	55	5	0.00	1.44
0.5 M H ₂ SO ₄	55	3.5	0.00	2.69
		5	0.07	1.93
0.5 M H ₂ SO ₄	55	5	0.05	2.22
		75	0.07	2.53
		95	0.34	3.63
0.5 M H ₂ SO ₄ + H ₂ O ₂	55	3.5	0.00	1.25

In almost all solutions after stripping no Mo(VI) was detected. Very low concentration of Mo(VI) was noted after the four-stage process in which the temperature was changed during the first step, i.e., leaching. Based on the results presented in Table 4 it can be concluded that in most cases Co(II) was efficiently separated from Mo(VI) by extraction and stripping.

To summarize the process, mass of Co(II) and Mo(VI) in the solutions after leaching and in the solutions after stripping is shown in Table 5. The yield (Y) of the four-stage Co(II) separation process was calculated as follows:

$$Y = \frac{m_{strip}}{m_{leach}} \cdot 100\% \quad (9)$$

where: m_{leach} —mass of Co(II) or Mo(VI) after leaching, m_{strip} —mass of Co(II) or Mo(VI) after stripping.

Table 5. Recovery yield of Co(II) and Mo(VI) in the four-step process (leaching-precipitation-extraction-stripping).

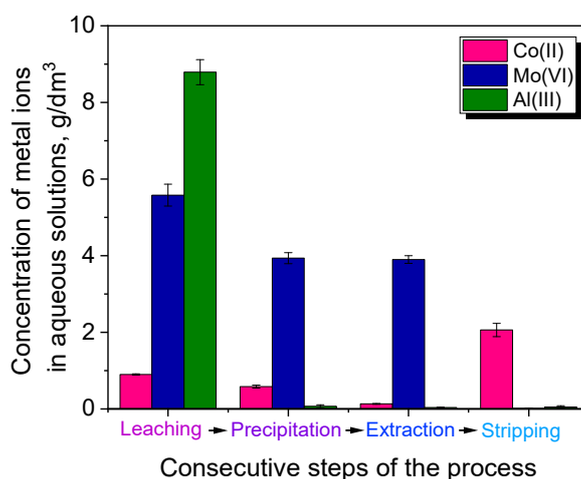
Temperature of Leaching with 0.5 M H ₂ SO ₄ , °C	Concentration in the Leach Solution, g/dm ³		Concentration in the Stripping Solution, g/dm ³		Y, %	
	Co(II)	Mo(VI)	Co(II)	Mo(VI)	Co(II)	Mo(VI)
23	0.81	3.97	1.93	0.07	16.5	0.08
55	0.90	4.49	2.22	0.05	18.6	0.07
75	0.99	5.45	2.53	0.07	19.8	0.12
95	1.11	6.06	3.63	0.34	18.8	0.33

Generally, the efficiency of Co(II) separation from solution after leaching did not exceed 20%. The efficiency of Mo(VI) stripping did not exceed 1%, thus, it was proven that Co(II) can be effectively separated from Mo(VI). However, further research should be carried out to increase the total yield of Co(II) recovery. As Al(III) is leached in a great amount from the catalyst, also the content of Al(III) in various solutions of the four-step process is shown in Table 6.

Table 6. Al(III) content in four-step process: leaching-precipitation-extraction-stripping (leaching solution: 0.5 M H₂SO₄ at 95 °C).

Al(III) Content in Solution, mg					
Leaching, min			After Precipitation	Raffinate after Extraction	Stripping Solution
5	90	180			
40.1	234.8	296.3	2.1	0.6	0.5
Al(III) Concentration in Solution, g/dm ³					
1.33	7.82	9.87	0.12	0.05	0.28

The amount of Al(III) in the stripping solution is comparable with Mo(VI) concentration and tenfold lower than Co(II) concentration. It means that cobalt(II) sulfate solution obtained after four steps of the process contains significantly less accompanying metal ions compared to the leaching solution. Finally, Co(II) recovery from spent industrial CoMo catalyst was carried out in six parallel leaching-precipitation-extraction-stripping processes. The results are shown in Figure 7 as concentrations of Co(II), Mo(VI) and Al(III) (in the aqueous solutions) after the consecutive steps of the process.

**Figure 7.** Flow of Co(II) (■), Mo(VI) (■) and Al(III) (■) in consecutive steps of the proposed process.

Compared to other processes developed to recover metals from spent catalysts [28,36,42], the advantage of our process lies in the fact that (1) the final solution contains cobalt(II) sulfate and is enriched in Co(II) related to the leaching solution, (2) separation of Mo(VI) is carried out in one extraction step, not two-step as proposed by others [36,37].

3. Materials and Methods

Sample of spent industrial CoMo catalyst was provided by a Polish waste treating company. The percentage content of elements determined by XRF analysis is presented in Table 7. Acidic extractant, i.e., *bis*(2,4,4-trimethylpentyl)phosphinic acid (Cyanex 272) supplied by Cytec Industries Inc. (Woodland Park, New Jersey, USA) Solvay Group was used as an extractant (0.4 M) and was dissolved in Exxsol D80 (ExxonMobil Chemical, Irving, Texas, USA) with the addition of modifier (i.e., 5 vol% decanol). The organic phase was regenerated after the previous extraction with sulfuric acid and scrubbed with water, and recycled to extraction.

Table 7. Percentage content of elements in spent industrial CoMo catalyst.

Element	Al	Si	P	S	Cl	K	Ca	Mn
Content, %	46.6	0.58	1.24	0.83	0.94	0.05	0.32	0.01
Element	Fe	Co	Cu	As	Se	Mo	Pb	-
Content, %	0.84	12.4	0.06	0.12	0.05	35.9	0.01	-

The solution of 30% sodium hydroxide (Avantor, Gliwice, Poland) was used to precipitate metal hydroxides from the solution after leaching, and then to adjust pH during extraction with Cyanex 272. Sulfuric acid (Avantor, Gliwice, Poland) in the range from 0.5 to 5 M was used as a leaching agent, in some cases, the addition of 30% H₂O₂ (Avantor, Gliwice, Poland) was applied. Sulfuric acid also was employed as a stripping phase (1 M) in all the cases, as it is the most convenient electrolyte for further electrowinning of metallic cobalt.

3.1. Leaching Procedure

The spent catalyst was ground, then sieved and a fraction of particle size below 45 µm was separated. 1.5 g of the ground catalyst was placed in a reactor, and 30 cm³ of H₂SO₄ or 25 cm³ of H₂SO₄ with the addition of 5 cm³ of H₂O₂ was added. The solid to liquid (S/L) ratio was kept equal to 1/20 g/cm³. The mixture of catalyst and the leaching agent was heated and mixed at 300 rpm, and the leaching was carried out in the range of temperature between 23 and 95 ± 2 °C. Leaching was investigated for 3 h and 1 cm³ sample was taken after 5, 15, 30, 60, 90, 120, 150, 180 min from the beginning of the leaching.

3.2. Precipitation Procedure

Precipitation of hydroxide deposit was carried out by slow addition of 30% NaOH to the leach solution up to pH 5. The precipitation was separated by centrifugation, and the liquid phase was taken to the next step, i.e., liquid-liquid extraction.

3.3. Extraction-Stripping Procedure

Extraction from model solutions was carried out in a typical batch way: aqueous feeds containing metal ions were mechanically shaken with the organic phase 0.4 M Cyanex 272 (volume ratio w/o = 2) for 15 min at 23 ± 2 °C in glass separation funnels, and then allowed to stand for phase separation. The pH of the aqueous phase was controlled by the addition of drops of 30% NaOH solution and kept equal to 5–5.2. Stripping of metal ions from the organic phases loaded was carried out for 5 min with 1 M H₂SO₄ at o/w = 3.

Atomic absorption spectroscopy (Contraa300, Analytik Jena, Jena, Germany) was applied to determine Co(II), Cu(II), Fe ions, Ni(II) concentrations in the aqueous solutions, microwave plasma-atomic emission spectroscopy (4210 MP AES, Agilent, Santa Clara, California, USA) was applied for Al(III), and Mo(VI). The content of elements in the spent catalyst was determined using XRF analysis (Malvern PANalytical Epsilon 1, Malvern, United Kingdom).

Standard deviations calculated for four repetitions of each step of the process did not exceed 5% for leaching, 8% for precipitation and extraction, and 8.5% for stripping.

4. Conclusions

Studies on the recovery of Co(II) from the leaching solutions prove that the proposed four-step hydrometallurgical process involving leaching of metals, precipitation of metal hydroxides, Co(II) extraction with *bis*(2,4,4-trimethylpentyl)phosphinic acid, and Co(II) stripping from the organic phase can be used to separate cobalt ions from molybdenum from sulfate solutions. During purification of the leaching solution by precipitation of Al(III) and Mo(VI) hydroxides, cobalt(II) losses are noted, as Co(II)

co-precipitates also as hydroxide or mixed compounds, e.g., $\text{Co}(\text{SO}_4)_i(\text{OH})_j \cdot k\text{H}_2\text{O}$ or $\text{CoMoO}_6 \cdot 0.9\text{H}_2\text{O}$, and passes into the sludge. This phenomenon is unfavorable for further stages of cobalt(II) recovery. Generally, the yield of Co(II) recovery from solution after leaching did not exceed 20%. The yield of Mo(VI) recovery did not exceed 1%, thus, it is proven that Co(II) can be effectively separated from Mo(VI). However, further research should be carried out to increase the total yield of Co(II) recovery.

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