RSC Advances

PAPER

Cite this: RSC Adv., 2017, 7, 2002

Received 18th October 2016 Accepted 1st December 2016

DOI: 10.1039/c6ra25412h

www.rsc.org/advances

1. Introduction

The separation of metal ions from industrial and waste solutions is extremely important in industrial applications.^{1,2} Efficient and economical methods for the separation and recovery of metal ions have drawn great attention due to the increasing demand on metal production.³⁻⁵ Several technologies are used for the separation and recovery of metal ions, including chemical precipitation, solvent extraction, ion exchange, liquid membranes, electrodialysis etc.⁵⁻¹¹ Cadmium and its compounds are important resources and used in different industries such as electroplating, pigments, synthetic chemicals, ceramics, metallurgical and photographic products, electronics and other industries.¹² Cadmium is a relatively rare metal, and primarily produced as a by-product from mining, smelting and refining of ore concentrates of zinc sulfide. It is also present in the metallurgical processing of other metals such as copper and lead.¹³ On the other hand, cadmium is one of the most toxic heavy metals, which can cause various health problems, such as choking, anemia, abdominal pain, dysfunction $etc.$ ^{14,15} Therefore, cadmium is both a resource and a very common heavy metal toxicant. As a resource, it is necessary to purify and remove copper impurities therein; as a heavy metal toxicant, it is necessary to remove cadmium from other metals in smelting, such as Cu, Zn and Ni. It is significant to develop effective methods for the separation of cadmium from other commonly associated metals.

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Cadmium and its compounds are important resources in different industries; on the other hand, cadmium is one of the most toxic heavy metals which can cause various health problems. Therefore it is important to develop effective methods for the separation of cadmium from other commonly associated metals from the stance of both resource recycling and environmental purification. Lime neutralization (Ca(OH)₂) and ferrite are widely used to precipitate heavy metals. Limestone (calcium carbonate: $CaCO₃$) is too stable to be used directly for this purpose. Mechanochemical activation was introduced to increase the activity of CaCO₃. Fe(II)sulfate heptahydrate (FeSO₄·7H₂O) was used as a selective precipitation agent. As a result, Cu(II) was preferentially precipitated as (Fe_x,Cu_y)O while the Cd(II) remained in the solution. The residual of Cu(II) ions in solution could be controlled at less than 0.1%, meanwhile more than 90% of $Cd(II)$ ions remained in aqueous solution. Then, $Cu(II)$ – $Cd(II)$ separation was achieved by a simple solid– liquid separation. PAPER

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The separations of $Zn(\text{II})-Cd(\text{II})$ and $Ni(\text{II})-Cd(\text{II})$ have been studied extensively.^{13,16-20} However, few documents have discussed the separation of Cu(II)–Cd(II). In this regard, the Cu(II)– $Cd(n)$ system was chosen for this investigation. The conventional chemical precipitation was effective and the formed precipitates can be separated from the water by sedimentation or filtration. Hydroxide precipitation is the most widely used chemical precipitation technique. For example, the lime milk $(Ca(OH₂)$ neutralization is a widely practiced technique in the hydroxide precipitation.^{21,22} However, the process of Ca(OH)₂ neutralization cannot achieve the selective precipitation of metal ions. Limestone $(CaCO₃)$ may be used as an adsorbent for the removing of heavy metals at very low concentration, due to its easily available, low-cost and insoluble in water. However, $CaCO₃$ is too chemically stable to neutralize the salts to precipitate heavy metals at an amount of stoichiometric ratio for a complete reaction. Thus, an activation process should be introduced for the application of $CaCO₃$ on the precipitation of metal ions. Mechanochemical activation (MA) is well-known in the expanding field of mechanochemistry and applied in many fields, such as materials syntheses, surface modification, hydrometallurgy, resource recycling and so on.^{23,24} In our previous studies we have investigated the separation of copper from nickel and cobalt in the sulfate solutions using $CaCO₃$ based on MA.^{25,26} In that system, the activated CaCO₃ reacted with copper sulfate to form precipitate of copper compounds, but not with nickel sulfate or cobalt sulfate, achieving the separation of them. However, in the effort to separate Cu from Cd, we noted that it was easy to form the co-precipitation of

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 $\text{Cu}(n)$ –Cd(n) compound by milling the mixture of Cu(n)sulfate,
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 $Cd(n)$ sulfate and $CaCO₃$ in aqueous solution. So the selective precipitation agent should be used in Cu–Cd–Ca system for $Cu(n)-Cd(n)$ separation.

Fe(II)sulfate heptahydrate (FeSO₄ $·$ 7H₂O) is often used as a precipitation agent to remove heavy metals from waste solution, the so-called ferrite process (FP). In the ferrite process, Fe(II) can be oxidized to Fe³⁺ and transformed into iron (Fe(II)/ Fe(III)) compounds, such as Fe₃O₄ (FeO \cdot Fe₂O₃), FeOOH and Fe(OH)₃.²⁷ For yielding high quality ferrite and suppressing the formation of the other crystal phases, the solution pH needs to be controlled at 9–11 and reaction temperature over 70 $^{\circ}$ C. A dosage of $Fe²⁺ 5-10$ times higher than the gross metal content in solution is required to ensure the removal efficiency.²⁸ The heavy metal ions can be removed by the incorporation into the structure through co-precipitation when they coexist with $Fe(\pi)$.

In this work, we present a detailed study on the $Cu(II)-Cd(II)$ separation using CaCO₃ assisted by adding Fe (n) sulfate heptahydrate. MA is used to increase the reactivity of $CaCO₃$. As the reactivity of $CaCO₃$ increased at a level, Fe(π) is transformed into iron (Fe²⁺/Fe³⁺) compounds, at the same time, Cu(II) is preferentially precipitated by the FP while $Cd(n)$ still exists in aqueous solution. The residual of $Cu(II)$ ions in solution could be controlled at less than 0.1%, meanwhile more than 90% of the $Cd(n)$ ions remained in aqueous solution, though a simple solid–liquid separation. This result is important for the development of new environment-friendly processes of various metals separation.

2. Materials and methods

2.1 Materials

Cu(II)sulfate pentahydrate CuSO₄.5H₂O, Cd(II)sulfate 8/ 3hydrate $CdSO_4 \cdot 8/3H_2O$, Fe(II)sulfate heptahydrate (FeSO₄- $-7H₂O$) and CaCO₃ from Sinopharm Chemical Reagent Co., Ltd in analytical grade (purity > 99%) were used as received without further purification.

2.2 Methods

With a separate case, 0.01 mol $CuSO_4·5H_2O$ or 0.01 mol $C dSO₄·8/3H₂O$ and equimolar $C a CO₃$ with 2 mL distilled water (designated as Cu–Ca and Cd–Ca system), were put into a zirconia pot of 45 mL inner volume with 7 zirconia balls of 15 mm in diameter. A laboratory scale planetary mill (Pulverisette-7, Fritsch, Germany) was used to conduct the milling operation with a milling speed of 400 rpm and milling time of 60 min (10 min grinding and 5 min pause in turn). Then the mixture solutions containing 0.0075 mol CuSO₄- \cdot 5H₂O, 0.0025 mol CdSO₄ \cdot 8/3H₂O and 0.01 mol CaCO₃ (designated as Cu–Cd–Ca system) or 0.005 mol FeSO₄.7H₂O, 0.005 mol $CuSO_4 \cdot 5H_2O$ and 0.01 mol $CaCO_3$ (designated as Cu–Fe–Ca system) were conducted at same grinding operation by fixing 2 mL distilled water. And then the $FeSO_4 \cdot 7H_2O$ was gradually added in the above-mentioned Cu–Cd–Ca system, setting the ratio of $Fe(n)/Cu(n)$ at 0.2, 0.3, 0.5 and 1. All the products were dried at 80 \degree C for 24 h for phase identification. X-ray diffraction (XRD: RU-200B/D/MAX-RB RU-200B, Japan)

analysis was performed to identify the phases in the ground samples. For the separation test, a Cu–Cd–Ca–Fe system was conducted at same grinding operation with five parameters changed of Fe-Cu ratio, dosage of CaCO₃, milling time, milling speed and the ratio of $Cd/(Fe + Cu)$. Detailed information was specified in the following section of Experimental results. Then, the prepared samples were agitated in 200 mL distilled water at room temperature for 60 min by a magnetic stirrer, respectively. And 10 mL solution was centrifuged at the 8933 \times g for 5 min. The concentrations of Cu, Cd and Fe in the supernatant were determined by inductively coupled plasma (ICP) spectrophotometry (ICP-AES: Optima-3300SYS, Perkin Elmer, USA). In addition, the pH of the sample from different Fe–Cu ratio was detected at different conditions by a pH meter (METTLER TOLEDO FE20-FiveEasy™, Switzerland). Paper

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Mild milling experiment was conducted for pH measurement for simulating the high-energy ball milling operation. The test was carried out in a vertical type stirred media mill. The net volume of the milling chamber is 200 mL. The mixture of 0.0075 mol CuSO₄ $5H_2O$ and 0.0025 mol CdSO₄ $8/3H_2O$ was put into the chamber with 100 mL of distilled water. The $FeSO₄·7H₂O$ and $CaCO₃$ were added in the Cu–Cd solution. The molar ratios of $Fe(\text{II})/Cu(\text{II})$ were set at 0, 0.2, 0.3, 0.5, 1 and 2. The molar ratio of $Ca/(Fe,Cu)$ was fixed at 1. After adding $CaCO₃$, the mixture was milled with 100 g zirconia balls of 2 mm in diameter at rotational speed of 250 rpm by a stirred mill for 30 min.

3. Results and discussion

3.1 Differences between Cu–Ca, Cd–Ca, Cu–Cd–Ca and Cu– Fe–Ca system

The metal sulfate was separately milled with $CaCO₃$ in aqueous solution. Fig. 1(a) displays the XRD pattern of milling the mixture containing copper sulfate and calcium carbonate. CaSO₄ 2H₂O (dihydrate gypsum) (JCPDS-ICDD card no. 33-0311) and $Cu₂CO₃(OH)₂$ (malachite) (JCPDS-ICDD card no. 41-1390) were determined in the sample, indicating that the copper sulfate and calcium carbonate were transformed into basic copper carbonate and gypsum by the mechanochemical processing. Fig. 1(b) shows the case of milling the mixture of $CdSO₄$ and $CaCO₃$. $CaCO₃$ (JCPDS-ICDD card no. 05-0586) and $CdSO₄·H₂O$ (JCPDS-ICDD card no. 53-0841) were determined in the XRD pattern, indicating that the raw materials still existed in the sample. Although the existence of $CaSO₄·2H₂O$ as a reaction product suggested a partial reaction between CaCO₃ and CdSO4, the raw materials of calcium carbonate and cadmium sulfate remained as the main phases. The reaction might be as eqn (1). Since the $\overline{{\rm CO_3}^{2-}}$ easily hydrolyzed in water as shown in eqn (2), it was probable that the generation of $CdCO₃$ might be hindered.

$$
CaCO3 + CdSO4 + 2H2O \rightarrow CaSO4 \cdot 2H2O \downarrow + CdCO3\downarrow (1)
$$

$$
CO_3^{2-} + H_2O \rightarrow HCO_3^- + OH^-
$$
 (2)

Fig. 1 XRD patterns of the samples from milling the metal sulfates with CaCO₃: (a) CuSO₄ + CaCO₃; (b) CdSO₄ + CaCO₃; (c) CuSO₄ + FeSO₄ + CaCO₃; (d) CdSO₄ + CuSO₄ + CaCO₃

The mixture powders of copper sulfate, ferrous sulfate and calcium carbonate were milled in aqueous solution, as shown in Fig. 1(c). The CaSO₄ \cdot 2H₂O was clearly identified in the sample. In addition to the peaks of $CaSO₄·2H₂O$, the rest of the peaks were consistent with FeO (JCPDS-ICDD card no. 46-1312) and Cu₂O (JCPDS-ICDD card no. 05-0667). In the FP, the Fe(π) can be occupied by Cu (n) , leading to the incorporation of Cu (n) in FeO. So the rest of peaks might be attributed to the compound (Fe_x, Cu_v) O. In the case of milling the mixture of copper sulphate, cadmium sulphate and calcium carbonate, the phase of hydrated basic sulfate of copper and cadmium with the molecular formula of $CdCu_4(SO_4)_2(OH)_6.4H_2O$ (niedermayrite) (JCPDS-ICDD card no. 50-1550) was identified in the XRD pattern, as shown in Fig. 1(d). It demonstrated that the coprecipitation of $Cu(\pi)$ and $Cd(\pi)$ occurred in the milling process. CaCO₃ reacted with CuSO₄ by liquid assisted milling operation, and the reactions were represented as eqn (3) and (4) as follows:

$$
2CuSO4 + 5H2O + 2CaCO3 \rightarrow Cu2CO3(OH)2\n+ 2CaSO4·2H2O\downarrow + CO2\uparrow
$$
\n(3)

$$
4CuSO_4 + CdSO_4 + 3CaCO_3 + 13H_2O \rightarrow
$$

CdCu₄(SO₄)₂(OH)₆·4H₂O₊ + 3CaSO₄·2H₂O₊ + 3CO₂† (4)

Although there was a significant difference between $Cu(II)$ and $Cd(n)$ sulfates when they were individually milled with CaCO3, the separation of them could not be achieved by only using the CaCO₃ due to the co-precipitation of Cu(π) and Cd(π) in the forms of $CdCu₄(SO₄)₂(OH)₆·4H₂O$.

3.2 Effect of $Fe(n)$ on phase transition and Cu–Cd separation

The existence of $Fe(n)$ effect on the Cu–Cd–Ca system was investigated by changing the molar ratio of $Fe(\pi)/Cu(\pi)$ from 0, 0.2, 0.3, 0.5 to 1, with other parameters fixed: $Cu(n)-Cd(n)$ ratio at 3 : 1; the dosage of CaCO₃ equal to the mole of $Fe(n) + Cu(n)$; the milling time at 60 min; rotating speed at 400 rpm; solid-liquid ratio at $1:1$. As shown in Fig. 2,

Fig. 2 XRD patterns of the samples from Cu–Cd–Ca–Fe system with the increasing of Fe(II)–Cu(II) ratio: (a) 0; (b) 1/5; (c) 1/3 (d) 1/2; (e) 1.

 $CdCu₄(SO₄)₂(OH)₆·4H₂O$, $CaSO₄·2H₂O$, $(Fe_x,Cu_y)O$ and $CaCO₃$ were determined in the samples. From the pattern a to e, the intensity of the peaks (020) and (040) belonging to $CdCu₄($ $SO_4)_2(OH)_6 \cdot 4H_2O$ was decreased with the increase of Fe(II) ions. In contrast, the intensity of the peaks from (Fe_x, Cu_y) O became stronger correspondingly. As a result, the existence of $Fe(\pi)$ ions had effectively prevented the generation of $CdCu₄(SO₄)₂(OH)₆·4H₂O.$

In addition, $CaCO₃$ was activated through the milling operation. Many ${CO_3}^{2-}$ were released into water and hydrolysis occurred as eqn (2). Meanwhile, the Fe^{2+} combined with hydroxyl groups as eqn (5) . After milling, the samples were stirred in beakers for duration of time. The stirring operation provided oxygen from air for chemical oxidation as eqn (6). Thus, the ferrite formation was assisted by the operation of milling and stirring, without high pH (9–11) and high temperature (over 70 \degree C).

$$
\text{Fe}^{2+} + 2\text{OH}^- \rightarrow \text{Fe}(\text{OH})_2 \tag{5}
$$

$$
4Fe(OH)_2 + O_2 + 2H_2O \rightarrow 4Fe(OH)_3 \downarrow
$$
 (6)

The $Cu(II)$ was preferentially precipitated by the FP in the Cu– Cd–Ca–Fe system and the Cd (n) remained in aqueous solution. In this regards, $Cu(n)$ had been separated from the $Cd(n)$ solution by solid–liquid separation.

Fig. 3 shows the effect of Fe(π) on the residual ratio of Cd(π) in aqueous solution. The residual ratio of $Cd(n)$ increased from 28.2% to 74.7% with the ratio of $Fe(n)/Cu(n)$ increased from 0 to 1, clearly indicating that the existence of $Fe(II)$ was beneficial to maintain the solubility of $Cd(n)$. The result was consistent with the results of XRD analysis in Fig. 2, which demonstrated that the $Fe(II)$ ions prevented the formation of $CdCu₄(SO₄)₂(OH)₆·4H₂O$ while promoting the generation of $(Fe_x, Cu_y)O.$

Therefore, the residual ratio of $Cd(n)$ in aqueous solution increased with the increase of $Fe(\pi)$ addition. Considering this phenomenon, the Cu(π)–Cd(π) separation might be achieved by regulating parameters.

Fig. 3 Effect of Fe(II) ions on the residual ratio of $Cd(II)$ in aqueous solution.

Fig. 4 Effect of speed on the residual ratio of Cd, Cu and Fe in aqueous solution.

3.3 Separation of Cu(II) and Cd(II)

3.3.1 Effect of rotational speed. The effect of the speed of milling operation was investigated and the results are shown in Fig. 4, with the solid-liquid ratio at $1:1$, Fe-Cu ratio at $1:1$, Cu–Cd ratio at $3:2$ and Ca–(Fe + Cu) ratio at 0.9. The rotational speed of the planetary mill was set at 200, 300, 400, 500 and 600 rpm. As shown in Fig. 4, the residual ratio of Fe and Cu decreased from 22.1% and 17.3% to 0 with the speed increased from 200 to 600 rpm, while the value of remaining Cd in the solution decreased from 96.4% to 86.3%. Considering the separation efficiency, 400 rpm was an optimal rotational speed.

3.3.2 Effect of CaCO₃. The effect of the molar ratio of Ca/(Fe + Cu) on the separation efficiency is shown in Fig. 5, with the solid–liquid ratio at 1 : 1, Fe–Cu ratio at 1 : 1, Cu–Cd ratio at 3 : 2, milling time at 60 min and grinding rotational speed at 400 rpm. The molar ratio of Ca/(Fe + Cu), as a variable factor, was set at 0.5, 0.9, 1.3, 1.7, 2.1 and 2.5. The residual ratio of three metal ions was decreased with the increase of $CaCO₃$. The declining tendency was relatively slow for Cd, compared with

Fig. 5 Effect of $CaCO₃$ on the residual ratio of Cd, Cu and Fe in aqueous solution.

Fig. 6 Effect of milling time on the residual ratio of Cd, Cu and Fe in aqueous solution.

that of Fe and Cu. As the molar ratio of $Ca/(Fe + Cu)$ increased from 0 to 2.5, the residual ratio of Cd decreased from 98% to 54% with 47.66% to 0 for Cu and 17.3% to 0 for Fe. It was noteworthy that the residual ratio of Cd was 90.7% when the molar ratio of Ca/(Fe + Cu) at 0.9, accompanying with only 5.8% Fe and 3.5% Cu remain in aqueous solution. The results revealed that the effective separation of $Cu(II)$ and $Cd(II)$ could be achieved using $CaCO₃$ with Fe(π)sulfate heptahydrate.

3.3.3 Effect of milling time. The separation efficiency affected by the milling time was investigated with the solid– liquid ratio at 1 : 1, Fe–Cu ratio at 1 : 1, Cu–Cd ratio at 3 : 2, Ca– (Fe + Cu) ratio at 0.9, and grinding rotational speed at 400 rpm. The milling time was set at 15, 30, 60, 90 and 120 min. The results are shown in Fig. 6. It was seen that the residual ratio of Cd, Cu and Fe in aqueous solution decreased with the increasing of milling time. The residual ratio of Cd decreased from 92.4% to 76.6% as the milling time increased from 15 to 120 min, with 12% to 0 and 7.7% to 0 for Fe and Cu respectively. The results indicated that there existed an optimal grinding time for high separation efficiency.

3.3.4 Effect of Cd. The effect of Cd on the residual ratio of Cd, Cu and Fe in aqueous solution was shown in Fig. 7. The ratio of Fe–Cu and Ca– $(Fe + Cu)$, the solid–liquid ratio, the grinding rotational speed and milling time were set at 1 and 1, 1, 400 rpm and 60 min, respectively. The molar ratio of Cd/(Fe + Cu), as a variable factor, was adjusted at 1/3, 1/2, 1, 3 and 6. As shown in Fig. 7, the residual ratio of Cd had a significant rise with the increasing of Cd –(Fe + Cu) ratio. The ratio increased from 69.1% to 99.9% as the Cd–(Fe + Cu) ratio was raised from 1/3 to 6. If the Cd–(Fe + Cu) ratio was more than 1, the residual ratio of Cd would be higher than 99% in the solution. Meanwhile, the residual ratio was not more than 3.3% and 1.5% for Cu and Fe respectively. It can be concluded that the separation of Cu–Cd is effective by regulating the operating conditions.

3.4 Mechanism discussion

The Fe(II) addition prevents the generation of CdCu₄(SO₄)₂(- $OH)_{6}$ · 4H₂O precipitation and promotes the formation of (Fe_x,-Cu_y)O phase. The Cu(II) is precipitated in the form of (Fe_x,Cu_y)O

Fig. 7 Effect of Cd on the residual ratio of Cd, Cu and Fe in aqueous solution.

while the $Cd(n)$ remains in the solution. Then the separation of $Cu(II)$ from Cd(II)sulfate solution is achieved by a simple solid– liquid separation. The mechanism of the process can be interpreted in two aspects as the following: (a) the acidic reaction environment is conducive to prevent the precipitate of $Cd(n)$; (b) Fe^{2+} is oxidized to Fe^{3+} and transformed into iron (Fe(II)/Fe(III)) compounds. The ferrite has an essential role in the precipitation of $Cu(_{II})$.

Mild milling experiment was conducted to simulate the process of high-energy ball milling. The pH was monitored as shown in Fig. 8(A). It can be seen that the pH value of Cu–Cd–Ca system is higher than that of Cu–Cd–Ca–Fe system. Acidic pH in the Cu–Cd–Ca system is derived from the hydrolysis of Cu (n) . Considering the relatively weaker hydrolysis of Fe (n) than Cu (n) , the pH value of Cu–Cd–Ca system should be theoretically lower than that of Cu–Cd–Ca–Fe system at the same molar metal ions. However, the hydrolysis of $Fe(m)$ is stronger than Cu(π). The lower value from the Cu–Cd–Ca–Fe system might be caused by the superimposed effect of the hydrolysis of Cu(π) and Fe(π). The theoretical solubility of cadmium hydroxide vs. pH is shown in Fig. $8(B)$ ²⁹ It demonstrated that the range of pH value leading to cadmium precipitate is 8–11. So from the aspect of pH, the additional $Fe(II)$ is not conducive to the precipitation of cadmium.

On the other hand, the ferrite has an essential role in the precipitation of $Cu(II)$. The XRD pattern of the sample formed by milling $FeSO₄$ and $CaCO₃$ in solution is shown in Fig. 9. The $FeSO₄$ and $CaCO₃$ were transformed into $FeO(OH)$ (goethite) (JCPDS-ICDD card no. 29-0713) and $CaSO₄·2H₂O$. It indicated that the Fe(π) was oxidized to Fe(π) by milling with CaCO₃ and transformed into $Fe(m)$ compounds, such as $FeO(OH)$. So in the Cu–Cd–Ca–Fe system, a part of $Fe(\text{II})$ also is oxidized to $Fe(\text{III})$ and transformed into iron compounds. The ferrite preferentially precipitate Cu(π) rather than Cd(π) may be ascribed to the difference in their ionic radii. The ionic radius of $Fe(n)$, Cu (n) and Cd(π) are 0.77, 0.71 and 0.97 Å, respectively.^{16,30} The ionic radii of $Cu(\pi)$ is close to that of Fe(π) but with larger difference from that of $Cd(n)$. The smaller the ionic radius, the greater its

Fig. 8 pH values effected by the additional Fe(II) (A) and theoretical solubility of cadmium hydroxide vs. pH (B)

Fig. 9 XRD pattern of the sample formed by milling $FeSO₄$ and $CaCO₃$ in solution.

affinity to occupy the sites of the Fe(π). Thus the Cu(π) preferentially occupies the position of $Fe(II)$ in the process of FP. This is consistent with the description in Fig. 2. The $Fe(II)$ ions prevented the formation of $CdCu₄(SO₄)₂(OH)₆·4H₂O$. And the compound (Fe_x,Cu_v)O was facilitated by the increasing of Fe(π). As a result, a part of $Cu(n)$ is co-precipitated with iron, leaving almost all $Cd(n)$ in solution.

The selective precipitation of $Cu(II)$ can be achieved by adding Fe(π) in the Cu–Cd–Ca system. FeSO₄ $7H_2O$ is the most common flocculant used in the removal of heavy metals. The sources of $CaCO₃$, such as calcite, limestone and shells, are around the world and very low-cost. In this process, the $CaCO₃$ and FeSO₄ \cdot 7H₂O are used to separate Cu(II) and Cd(II) and the result is feasible. The dosage of $FeSO₄·7H₂O$ in this process is less than 20% that of traditional FP processing under the same processing capacity. In addition, the neutral sediments are environmentally friendly.

4. Conclusion

Mechanochemical activation technique is efficiently used to the activation of CaCO₃. The FeSO₄ \cdot 7H₂O is used as a selective precipitation agent. The process allows the reaction of calcium

carbonate and ferrous sulfate with copper sulfate to form precipitate of copper compounds, but without similar reaction with cadmium sulfate. The separation of copper from cadmium is feasible by a simple solid–liquid separation as filtering, with 99% copper in solid and 90% of cadmium remaining in aqueous solution at optimized conditions. The experimental results may serve as the basis to treat various types of solutions containing Cu–Cd ions to develop an easy and environment-friendly separation process. The concept developed in this report may be also applied to other metals separation when some large differences in specific property exist as the case here.

References

- 1 S. E. Kentish and G. W. Stevens, Chem. Eng. J., 2001, 84, 149– 159.
- 2 R. S. Juang and S. W. Wang, Water Res., 2000, 34, 3795–3803.
- 3 A. R. Mahdavian and M. A.-S. Mirrahimi, Chem. Eng. J., 2010, 159, 264–271.
- 4 F. J. Alguacil, M. Alonso and A. M. Sastre, Chem. Eng. J., 2002, 85, 265–272.
- 5 F. J. Alguacil, A. Cobo and M. Alonso, Chem. Eng. J., 2002, 85, 259–263.
- 6 M. Sadrzadeh, T. Mohammadi, J. Ivakpour and N. Kasiri, Chem. Eng. J., 2008, 144, 431–441.
- 7 P. Cañizares, Á. Pérez and R. Camarillo, *Desalination*, 2002, 144, 279–285.
- 8 X. J. Yang, A. G. Fane and C. Pin, Chem. Eng. J., 2002, 88, 37– 44.
- 9 D. Pathania, G. Sharma and R. Thakur, Chem. Eng. J., 2015, 267, 235–244.
- 10 R. Kumar, M. Kumar, R. Ahmad and M. A. Barakat, Chem. Eng. J., 2013, 218, 32–38.
- 11 M. Laatikainen and K. Laatikainen, Chem. Eng. J., 2016, 287, 74–82.
- 12 M. S. Safarzadeh, M. S. Bafghi, D. Moradkhani and M. O. Ilkhchi, Miner. Eng., 2007, 20, 211–220.
- 13 B. Ramachandra Reddy, D. Neela Priya and J. Rajesh Kumar, Hydrometallurgy, 2004, 74, 277–283.
- 14 W. Wei, J. K. Bediako, S. Kim and Y.-S. Yun, J. Taiwan Inst. Chem. Eng., 2016, 61, 188–195.
- 15 J. Li, C. Chen, K. Zhu and X. Wang, J. Taiwan Inst. Chem. Eng., 2016, 59, 389–394.
- 16 M. Horsfall Jr and A. A. Abia, Water Res., 2003, 37, 4913–4923.
- 17 K. Takeshita, K. Watanabe, Y. Nakano and M. Watanabe, Hydrometallurgy, 2003, 70, 63–71.
- 18 P. R. Danesi, R. Chiarizia and A. Castagnola, J. Membr. Sci., 1983, 14, 161–174.
- 19 L. B. Chaudhari and Z. V. P. Murthy, J. Hazard. Mater., 2010, 180, 309–315.
- 20 M. S. Safarzadeh and D. Moradkhani, Sep. Purif. Technol., 2010, 73, 339–341.
- 21 D. R. Jenke, G. K. Pagenkopf and F. E. Diebold, Environ. Sci. Technol., 1983, 17, 217–223.
- 22 J. G. Dean, F. L. Bosqui and K. H. Lanouette, Environ. Sci. Technol., 1972, 6, 518–522.
- 23 S. L. James, C. J. Adams, C. Bolm, D. Braga, P. Collier, T. Friscic, F. Grepioni, K. D. Harris, G. Hyett, W. Jones,
	- A. Krebs, J. Mack, L. Maini, A. G. Orpen, I. P. Parkin,

W. C. Shearouse, J. W. Steed and D. C. Waddell, Chem. Soc. Rev., 2012, 41, 413–447.

- 24 P. Balaz, M. Achimovicova, M. Balaz, P. Billik, Z. Cherkezova-Zheleva, J. M. Criado, F. Delogu, E. Dutkova, E. Gaffet, F. J. Gotor, R. Kumar, I. Mitov, T. Rojac, M. Senna, A. Streletskii and K. Wieczorek-Ciurowa, Chem. Soc. Rev., 2013, 42, 7571–7637. **SC Advances**

14 W. Wei, J. K. Beilaho, S. Kim, A. This article. Published on 12 January 2017. Downloaded on 12 January 2017. Downloaded on 5/22/2025 8:43:45 AM. This article is licensed under a Creative Commons Attributi
	- 25 X. W. Li, Z. W. Lei, J. Qu, H. M. Hu and Q. W. Zhang, Sep. Purif. Technol., 2017, 172, 107–112.
	- 26 X. W. Li, Z. W. Lei, J. Qu, Z. Li and Q. W. Zhang, Sep. Sci. Technol., 2016, 51, 2772–2779.
	- 27 J. C. Lou and C. K. Chang, Sep. Purif. Technol., 2007, 57, 513– 518.
	- 28 Y. J. Tu, C. K. Chang, C. F. You and S. L. Wang, J. Hazard. Mater., 2012, 209, 379–384.
	- 29 D. M. Ayres, A. P. Davis and P. M. Gietka, Engineering Research Centre Report, 1972, pp. 518–522.
	- 30 S. T. Breviglieri, E. T. G. Cavalheiro and G. O. Chierice, ´ Thermochim. Acta, 2000, 356, 79–84.