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Manganese Removal from Sulfuric Acid Leach Solutions of Nickel Laterite

Ores

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DECLARATION

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ABSTRACT

The growing demand for nickel since the 1950s and the dwindling nickel sulphide resources have resulted in an increasing need to process nickel laterite ores as an alternative source. Currently, pressure acid leaching (PAL) is the most widely used leaching technique for these ores. It involves leaching the ore with sulfuric acid at high temperature (250 °C) yielding high nickel and cobalt dissolution (>95%) but also substantial co-dissolution of other components of the ore, and thus generating a highly contaminated pregnant leach solution (PLS). Among the contaminants, manganese constitutes the bulk of the impurities in the PLS after iron, together with aluminium and chromium, has been removed and therefore affects the efficiency of the downstream processing, whether intermediate precipitation, mixed hydroxide precipitation (MHP) or mixed sulphide precipitation (MSP), or direct solvent extraction (DSX) is chosen as the purifying technique. These purification techniques have proven to be unsuccessful in separating manganese from nickel and cobalt, owing to its co-precipitation in intermediate precipitation or co-extraction in DSX. Moreover, intermediate nickel and cobalt products, which can either be as mixed sulfides or mixed hydroxides coming from intermediate precipitation, require further processing through consecutive re-leaching and solvent extraction steps before nickel and cobalt can be recovered as final saleable products and therefore increasing the operational costs associated to processing nickel laterite ores. Clearly, removing manganese prior to the purifying step would enhance the efficiency of the step regardless of the route taken and hence, this proposed study.

The purpose of the present work was to explore the removal of manganese(II) from the PLS generated from PAL of nickel laterite ores by oxidative precipitation with potassium permanganate. In pursuing this aim, screening experiments using unreplicated full factorial design were performed to determine the variables that significantly affect the oxidative precipitation of manganese(II) using synthetic partially neutralised and post-partially neutralised PLS. Afterwards, the variables determined to have a significant effect on the oxidative precipitation of manganese(II) by potassium permanganate were optimised before further applying the technique to actual partially neutralised and post-partially neutralised PLS.

The results of the screening experiments revealed that pH and molar ratio between moles of permanganate ion and total moles of iron(II) and manganese(II) (denoted as mol MnO4⁻: total mol Fe(II) and Mn(II)) for partially neutralised PLS and between moles of permanganate ion and moles of manganese(II) (denoted as mol MnO4⁻:mol Mn(II)) for post-partially neutralised PLS significantly affect the oxidative precipitation of manganese(II). Optimisation experiments using synthetic PLS revealed that the oxidative precipitation of manganese(II) can be carried out in acidic conditions, specifically at a pH less than or equal to pH 4. The results also showed that the amount of precipitated manganese(II) increased with an increase in molar ratio. For a partially neutralised PLS, pH 3 and a stoichiometric molar ratio of 0.50 were found to be optimum where the amount of precipitated manganese(II) was 85% with minimal losses for nickel(II) and cobalt(II) at 2% and 7%, respectively. As for the optimisation experiments using post-partially neutralised PLS, a pH maintained between pH 1.66 and pH 1.71 and molar ratio of less than or equal to 0.70 were optimum. The amount of precipitated manganese(II) was nearly 100% with losses of nickel(II) between 3% and 10% while for cobalt(II), it was between 50% and 60%. Application of the optimised pH and molar ratio for the oxidative precipitation of manganese(II) from actual partially neutralised and post-partially neutralised PLS resulted in nearly 100% of precipitated manganese(II). Nickel(II) losses were approximately 27% while for cobalt(II), it was approximately 50% for the actual partially neutralised PLS. As for the actual post-partially neutralised, nickel(II) losses were approximately between 3% and 10% while for cobalt(II), it was approximately between 43% and 54%. Based on these results, it can be concluded that the right combination of pH and molar ratio is important in order to achieve maximum removal of manganese(II) with minimum nickel(II) and cobalt(II) losses.

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Chapter 1

INTRODUCTION

1.1 Background of the Study

The global production of nickel has been increasing by approximately four percent per annum since the 1950s (Dalvi et al. 2004, Goonan et al. 2017). This is owing to nickel's industrially desirable properties, mainly corrosion resistance, heat resistance, elasticity, and hardness, and hence its increasing industrial use in building and construction, transportation, and electronics, to name a few. According to Mudd (2010), approximately 58% of nickel's end use application is in stainless steel production while the rest is used for nickel-based alloys, casting and alloy steels, electroplating, and rechargeable batteries. Except for stainless steel production, all these applications require high purity nickel. The growing demand for nickel drives the need to improve existing processes as well as develop new processes that can maximise the nickel that can be obtained from its ores.

Nickel ores can either be sulfidic or lateritic. Approximately 70% of the nickel ore deposits are lateritic while the rest are sulfidic. Traditionally, nickel sulfide ores have been the source of nickel since they are amenable to beneficiation by concentrating the valuable metals and rejecting the gangue in the ore, and this accounts for approximately 60% of the total global nickel production (Dalvi et al. 2004, Kyle 2010, Mudd 2010). As a result, nickel sulfide ore resources are being consumed and thus the need to process nickel laterite ores has been increasing. By increasing the global production coming from nickel laterite ores, it can help sustain the increasing demand for nickel. In 2009, the shift to processing nickel laterite ores resulted in an increase in its global proportion by 54% and the growth is projected to increase close to 72% by the year 2022 (Oxley et al. 2016).

Unlike nickel sulfide ores that can directly be beneficiated by conventional methods, such as flotation, before further extracting the valuable metals, nickel laterite ores are of lower grade and not amenable to beneficiation. The concentrations of the valuable metals in nickel laterite ores—nickel and cobalt—relative to the impurities are much lower and their minerals have similar properties to those of the gangue, making their separation difficult (Dalvi et al. 2004, Kyle 2010). Moreover, in these ores, the nickel is primarily associated in the goethite (80%) phase and no free nickel mineral is present in the ore (Zhu et al. 2012). Hence, upgrading of the ore is difficult (Quast et al. 2015). It follows that the bulk of the ore, including the impurities, needs to be processed and consequently, can be more expensive and complex to do than those that can be upgraded, such as the nickel sulfide ores.

1.2 Overview of Occurrence and Characteristics of Nickel Laterite Ores

Nickel laterite ores are mostly distributed in tropical and subtropical regions around the world, such as Australia, New Caledonia, Philippines, and Indonesia, as shown in Figure 1 (Brand et al. 1998).



Figure 1. World distribution of nickel laterite ore resources by country (Brand et al. 1998). *Content is removed due to copyright restrictions.*

The formation of nickel laterite ores is a result of chemical weathering through the natural leaching of peridotite, which consists mainly of the mineral olivine or its hydrated derivative, serpentine (Golightly 1979, Alcock 1988). As peridotite undergoes weathering, the presence of water in the ground, which is usually affected by the frequency and amount of rainfall, allows the movement of soluble elements

from peridotite that results in the development of different nickel laterite ore profiles. As such, the profile developed in dryer climate will be different from those in humid climate due to the difference in the frequency and amount of rainfall. Figure 2 shows three types of nickel laterite ore profiles based on dryer and humid climatic conditions. Each profile is further classified into three distinct zones from bottom to top. The bottom zone is classified as the saprolite zone while the middle and top zones are nontronite zone and limonite zone, respectively.



Figure 2. Typical nickel laterite ore profiles in dry and humid climates (Alcock 1988). *Content is removed due to copyright restrictions.*

The saprolite and nontronite zones develop in dryer climatic conditions as opposed to the limonite zone, which develops in humid climatic conditions. The saprolite zone is enriched in soluble elements, such as nickel and magnesium, that decrease as it moves up the profile towards the limonite zone, which contains mostly insoluble elements, primarily iron together with aluminium and chromium. The silica-enriched nontronite zone only develops when the water circulation is sporadic during weathering in areas of dryer climatic conditions and it is said to be non-economic when compared with the composition of the saprolite zone (Alcock 1988). Elias (2002) also highlighted in his study that the composition and development of a nickel laterite profile into the different zones during weathering can vary from one deposit to another or even within the same deposit due to varying climatic and geological factors. Clearly, having the right understanding of both the occurrence and characteristics of nickel laterite ores, including the different minerals associated with the valuable metals, has a significant implication for the selection of the most suitable processing route for nickel laterite ores. Table 1 shows the typical chemical composition ranges of nickel laterite ores while Table 2 shows the nickel content in significant minerals that can be found in the different zones of a nickel laterite ore profile.

| | Limonitic | Saprolitic |
|--------------------------------|-----------|------------|
| Ni | 0.12-3.0 | 1.0-4.0 |
| Со | 0.05-0.28 | 0.05-0.08 |
| MgO | 0.2–5.0 | 25–38 |
| CaO | 0.6-1.0 | 1.0-2.0 |
| Al ₂ O ₃ | 4.0–18 | 1.0-3.9 |
| Cr ₂ O ₃ | 1.5–4.5 | 1.0-3.0 |
| Fe ₂ O ₃ | 50–85 | 10–25 |
| MnO | 0.3–2.5 | 0.5–1.0 |
| SiO ₂ | 1.3–6.0 | 40–55 |

Table 1. Chemical Composition Ranges (%) of Nickel Laterite Ores (Alcock 1988).

Table 2. Nickel Content in Significant Minerals of Nickel Laterite Ores (Alcock1988).

| Zone | | Formula | % Ni |
|--------------|--|---|---------|
| Saprolite | 'olite Nickeliferous (Mg,Fe,Ni) ₃ Si ₂ O ₅ (OH) ₄ Serpentine | | 1–10 |
| | Garnierite | (Ni,Mg) ₃ Si ₄ O ₁₀ (OH) ₂ | 10–24 |
| Intermediate | Nontronite | (Ca,Na,K) _{0.5} (Fe ^{3+,} Ni,Mg,Al) ₄ (Si,Al) ₈ O ₂₀ (OH) ₄ | 0–5 |
| | Quartz | SiO ₂ | 0 |
| Limonite | Goethite | (Fe,Al,Ni)OOH | 0.5–1.5 |
| | Asbolite | Mn,Fe,Co,Ni Oxide | 1–10 |

1.3 Processing Routes for Nickel Laterite Ores

The processing of nickel laterite ores is via one of two major routes: pyrometallurgical or hydrometallurgical. The pyrometallurgical route, as shown in a generalised

flowsheet in Figure 3, which is suitable for saprolitic ores, is carried out in a series of steps by drying, heating, reducing, and smelting of the ore before the final recovery of the valuable metals (Alcock 1988, Davis 2000, Dalvi et al. 2004, Kyle 2010). However, the pyrometallurgical route produces only either ferronickel or nickel matte. Ferronickel or nickel matte are only suitable if the end use application does not require high purity nickel, such as in stainless steel production. Moreover, cobalt as a by-product cannot be separated if this route is chosen.





Alternatively, if the aim is to produce high purity nickel from nickel laterite ores that is suitable for end use application, such as in battery manufacturing and metal plating, the only option available is the hydrometallurgical route. The hydrometallurgical route, as shown in a generalised flowsheet in Figure 4, which is suitable for limonitic ores, is carried out by leaching the ore, partially neutralising the pregnant leach solution (PLS), and purifying the PLS by removing any remaining undesirable major impurities before further recovering the valuable metals. One advantage of this route over the pyrometallurgical route is the separation of cobalt as a valuable by-product (Alcock 1988, Davis 2000, Dalvi et al. 2004, Kyle 2010). This advantage will enable the maximisation of returns through the hydrometallurgical route. As a result of the accompanying advantage in choosing the hydrometallurgical route, it has been apparent that there is an upward trend in the development of nickel laterite ore processing projects since the 1990s focusing towards increasing the global hydrometallurgical capacity (Dalvi et al. 2004). Currently, there are three available leaching processes under the hydrometallurgical route, and these are the Caron process, pressure acid leaching (PAL), and atmospheric leaching (AL) (Dalvi et al. 2004, Kyle 2010).



Figure 4. Generalised flowsheet for the hydrometallurgical route.

The Caron process, which was developed during the 1920s and commercialised in the early 1940s (Caron 1950), involves consecutive energy-intensive drying and reductive

roasting of limonite ore (Fe content >40%) followed by atmospheric ammoniacal leaching of the nickel and cobalt. Although this process is reasonably selective in leaching nickel and cobalt from the ore and it allows recycling of excess ammonia, lowering its consumption, the dissolutions of nickel and cobalt are less than 90% and 60%, respectively (Crundwell et al. 2011). The overall low recovery of the Caron process is due to the low dissolution of nickel and cobalt in the reduced ore. During the reductive roasting step, the ore is only partially reduced to convert portions of nickel and cobalt into free metals while the rest remains bound in the iron matrix (Chander and Sharma 1981, Valix and Cheung 2002, Kyle 2010, Senanayake et al. 2010). Because of the issues underlying the dissolution of nickel and cobalt from the reduced ore, the commercial application of the Caron process is waning.

The PAL process, which was first commercially used in the late 1950s (Carlson and Simons 1960), involves leaching the ore with sulfuric acid (H_2SO_4) in a titanium-lined autoclave at 250 °C. Unlike the Caron process, this process does not require the drying and reductive roasting of the ore, thereby eliminating the cost of two energy-intensive processes. Additionally, the dissolution of both nickel and cobalt is greater than 95% while the precipitation reactions involved in the process allow the removal of significant amounts of co-dissolved iron and the regeneration of the initially consumed sulfuric acid, as shown in Equations 1-1 to 1-3.

Hematite:
$$Fe_2(SO_4)_3 + 3 H_2O \rightarrow Fe_2O_{3(s)} + 3 H_2SO_4$$
 (1-1)

Alunite: 1.5
$$Al_2(SO_4)_3 + 7 H_2O \rightarrow (H_3O)Al_3(SO_4)_2(OH)_{6(s)} + 2.5 H_2SO_4$$
 (1-3)

The higher dissolution of the valuable metals that is achievable using PAL is a distinct advantage over the Caron process. The PAL-generated PLS, however, has much more impurities than that of Caron's owing to the higher solubilities of magnesium, calcium, manganese, aluminium, and chromium in acidic media, which can affect the downstream purification process, resulting in additional cost (Georgiou and Papangelakis 1998, Whittington and Muir 2000, Kyle 2010, Ucyildiz and Girgin 2017).

Atmospheric leaching (AL), which was adopted by BHP Billiton in the early 2000s as part of its Ravensthorpe nickel project, involves leaching the ore with sulfuric acid at temperatures between 90 °C to 100 °C in an open vessel. Operating at lower temperatures does not necessarily reduce the energy cost of processing the ore because it also reduces the leaching kinetics and thereby requiring longer residence time. Consequently, the resulting PLS is contaminated with highly soluble iron that further affects the downstream processing (McDonald and Whittington 2008, Kyle 2010). High concentration of soluble iron is present in the AL-generated PLS due to the absence of iron hydrolysis that occurs in PAL at high operating temperature (250 °C). The occurrence of iron hydrolysis in PAL allows the bulk of the dissolved iron to precipitate right after dissolution as opposed to AL. The differences between the concentration of soluble iron in AL and PAL are reported by the studies of Liu et al. (2004) and White et al. (2006). In the study completed by Liu et al. (2004), the resulting AL-generated PLS from a limonitic nickel laterite ore contained as high as 80 g/L of iron. In contrast, White et al. (2006) observed that using PAL on a limonitic nickel laterite ore only resulted in an iron concentration of 2.2 g/L. Therefore, the presence of considerable amounts of soluble iron in AL-generated PLS compared to PAL-generated PLS will make the subsequent downstream purification process challenging.

1.4 Purification of the Pregnant Leach Solution from Pressure Acid Leaching

After leaching the ore, the PAL-generated PLS is partially neutralised in order to remove largely the remaining iron, aluminium, and chromium that are present in the solution. The resulting PLS will further undergo a purification step before recovering nickel and cobalt as separate products. The purification of the PLS is necessary in order to separate nickel and cobalt from each other and any remaining impurities, particularly manganese, which can range between 1 g/L and 3 g/L (Cheng and Urbani 2005). Purification of the PAL-generated PLS is carried out through either intermediate precipitation or direct solvent extraction (DSX). Mixed sulfide precipitation (MSP) and mixed hydroxide precipitation (MHP) are two processes available under intermediate precipitate. The intermediate product needs to undergo

further processing through consecutive re-leaching and solvent extraction steps to recover nickel and cobalt as high purity products. Conversely, DSX does not involve producing an intermediate product of mixed nickel and cobalt, but instead, it separates nickel and cobalt directly from the partially neutralised PLS through consecutive solvent extraction steps.

1.4.1 Intermediate Precipitation

Intermediate precipitation has been widely used in various nickel laterite operations, such as Moa Bay (Carlson and Simons 1960), Murrin Murrin (Motteram et al. 1996), Cawse (Kyle and Furfaro 1997, Mason et al. 1997, Grassi et al. 2000, White 2009), Ravensthorpe (White et al. 2006, White 2009), Coral Bay (Llerin et al. 2011), and Taganito (Shibayama et al. 2016), to further purify the PAL-generated PLS after the partial neutralisation step. Precipitation allows the separation of nickel and cobalt as intermediate products from any remaining major impurity, which in this case is manganese. Precipitation in the said operations is carried out through the addition of a suitable reagent to form an intermediate product of mixed nickel and cobalt precipitate.

Mixed sulfide precipitation (MSP) was first applied in the nickel laterite operation in Moa Bay (Carlson and Simons 1960) and later on in Murrin Murrin (Motteram et al. 1996), Coral Bay (Llerin et al. 2011), and Taganito (Shibayama et al. 2016) using hydrogen sulfide. The PAL-generated PLS undergoes partial neutralisation to precipitate iron, chromium, and aluminium using coral mud (Moa Bay), calcrete (Murrin-Murrin) or limestone (Coral Bay and Taganito) by gradually increasing the pH. In-between washing in a counter current decantation (CCD) is carried out before hydrogen sulfide (H₂S) gas is introduced to the solution to induce sulfide precipitation of nickel and cobalt (Carlson and Simons 1960, Motteram et al. 1996, Llerin et al. 2011, Shibayama et al. 2016). The application of MSP has proven to be selective in separating nickel and cobalt over the impurities associated with them in the PALgenerated feed, which includes manganese. Fundamentally, the solubility product constant of manganese sulfide is greater than nickel sulfide and cobalt sulfide. This suggests that it is unlikely for manganese to co-precipitate with the intermediate products of nickel and cobalt and therefore avoiding any contamination. In the mentioned operations (Carlson and Simons 1960, Motteram et al. 1996, Llerin et al. 2011, Shibayama et al. 2016), sulfide precipitation precipitates up to 99% of nickel and cobalt from the PLS as metal sulfides while impurities such as manganese that are more soluble than nickel and cobalt in the presence of sulfide ions remain dissolved in the solution. However, the application of MSP can only be carried out at elevated pressure and temperature, which would require the use of high-specification equipment, such as brick-lined pressure vessels (White 2009). Additionally, there are risks in using hydrogen sulfide due to its toxicity and therefore requires proper use, handling, and control of the reagent in a pressurised vessel, which would demand highly trained personnel (Dickson 2000, Willis 2007, Kyle 2010), making sulfide precipitation less attractive. The intermediate product coming from MSP would still require re-leaching and solvent extraction to ultimately recover nickel and cobalt. Despite the advantages of using MSP, it has not been adopted as the main purification process in nickel laterite operations due to the drawbacks mentioned.

Mixed hydroxide precipitation (MHP) was first applied in Cawse (Kyle and Furfaro 1997, Mason et al. 1997, Grassi et al. 2000, White 2009) and later in Ravensthorpe (White et al. 2006, White 2009) nickel laterite operations for the purification of the PAL-generated PLS. After the ore undergoes PAL, the PLS is partially neutralised using limestone to raise the pH before proceeding to the CCD. At this stage, most of the ferric ions would precipitate, leaving manganese as the main impurity in the solution and some ferrous ion, which are subsequently removed after the CCD. After the removal of the remaining ferrous ion, the solution is subjected to hydroxide precipitation using magnesia (MgO) to precipitate nickel and cobalt (Kyle and Furfaro 1997, Mason et al. 1997, Grassi et al. 2000, White 2009). The application of MHP can be carried out in atmospheric pressure using standard equipment, such as open top tanks and agitators (White 2009), which can be advantageous when compared with MSP. However, the use of MHP in Cawse and Ravensthorpe operation was found to be challenging due to its poor selectivity in separating nickel and cobalt from manganese. The poor selectivity of the process was attributed to the co-precipitation of manganese together with nickel and cobalt, which affects the purity and grade of the intermediate product (Kyle and Furfaro 1997, Grassi et al. 2000, White et al. 2006). Analysis of the intermediate product of mixed nickel-cobalt hydroxide precipitate contained as high as 12% manganese (Kyle and Furfaro 1997); therefore, the precipitate needs to undergo consecutive re-leaching and solvent extraction steps similar to MSP in order to separate nickel and cobalt as pure saleable products. Fundamentally, since the solubility product constants of nickel hydroxide and cobalt hydroxide are close to that of manganese hydroxide, the probability of manganese co-precipitating is high. Therefore, removing manganese prior to the precipitation of nickel and cobalt as MHP would substantially enhance the efficiency of the process and reduce the potential contamination of the intermediate products.

1.4.2 Direct Solvent Extraction

DSX involves the use of organic extractants to separate nickel and cobalt directly from the PAL-generated PLS. The commercial application of the process was pioneered in the Bulong nickel laterite operation. In this process, the neutralised PAL-generated PLS is first fed into the cobalt extraction circuit (CoSX) with a dialkyl phosphinic acid, Cyanex 272, as the extractant. Cobalt, together with the impurities such as zinc, iron, copper, and manganese, is extracted leaving the nickel in the raffinate, which is then fed to the nickel extraction circuit (NiSX) where nickel is extracted with a neodecanoic acid, Versatic 10. The extracted nickel is then stripped with barren electrolyte from the tank house, which is then fed back for the electrowinning of nickel (Griffin 2000, O'Callaghan 2003, Donegan 2006). Meanwhile, the cobalt together with the coextracted impurities is stripped and put through a separate purifying stream, which involves precipitation, re-leaching, and solvent extraction to separate the cobalt from all the co-extracted impurity metals (Griffin 2000, Donegan 2006). Manganese is the major impurity and constitutes the bulk of the PLS. Moreover, when manganese is entrained in the organic solution used in solvent extraction, there is a possibility that it will be transferred to the loaded nickel electrolyte and oxidise to a higher oxidation state during the electrowinning step (Cheng et al. 2000). Clearly, removing manganese prior to CoSX would considerably enhance the efficiency of the whole DSX process.

1.5 Statement of the Problem

Although the Caron process is a well proven technology and AL provides lower operating costs and uses less expensive equipment compared with PAL, the latter has a distinct advantage as it allows higher nickel and cobalt dissolution and with much lower impurities in the PLS than AL. PAL therefore provides opportunity to maximise the recoveries of these metals from nickel laterite ores, which are abundant. One of its major drawbacks, however, is the large amount of manganese in the PLS, which remains even after the partial neutralisation step. An effective technique of removing this metal from the post-partially neutralised PLS or, better, if in conjunction with the partial neutralisation step, is therefore highly desirable as it would make the purifying step much more efficient, whether it is carried out by intermediate precipitation (MHP or MSP) or DSX. In order to find a solution in solving the problem, it has been postulated by D.C. Ibana (2017, May) in a personal communication that the removal of manganese(II) from PAL-generated PLS may be achieved through oxidative precipitation using potassium permanganate, which will not introduce any new ions to the highly contaminated PLS, and if successful, it may allow the recovery of manganese as a valuable by product¹.

1.6 Aim and Scope of the Study

This project was aimed to explore the removal of manganese from partially neutralised and post-partially neutralised PLS of nickel laterites ores as manganese dioxide by oxidative precipitation with potassium permanganate as a method of purifying the PLS with regard to this impurity. Specifically, it was aimed to:

- review the aqueous chemistry of the manganous and permanganate ions, including their redox reaction;
- determine the variables that significantly affect the oxidative precipitation of manganese;

¹ D. C. Ibana, Associate Professor, Curtin University – Western Australian School of Mines, Personal Communications, May 2017

- optimise the variables with significant effects determined in the previous step by applying the technique using synthetic PLS, both partially neutralised and post-partially neutralised; and
- 4. apply the optimised variables using actual PLS, both partially neutralised and post-partially neutralised.

1.7 Significance of the Study

This project will contribute to the much-needed development of knowledge in the field of process metallurgy of nickel laterite ores, which is quite new, and will benefit the fledgling nickel laterite industry including researches in the field. If successful, a commercial application is within easy reach as Australia has the world's largest economic nickel resource (25%) where 69% occur as nickel laterite ores and thus, a major beneficiary of the work (Geoscience Australia 2012, McRae 2018). In addition, several countries in the region, including Papua New Guinea, New Caledonia, Indonesia and the Philippines, also host large deposits of nickel laterite ores and should therefore benefit from this work. This resource is also plentiful in South America, Europe, and Africa. Advances in its processing therefore is truly of global interest.

Chapter 2

REVIEW OF STUDIES ON THE REMOVAL OF MANGANESE BY

OXIDATIVE PRECIPITATION

2.1 Manganese in Nickel Laterite Ores and their PAL Leach Solutions

Manganese is contained in many minerals that are often found in nickel laterite ores. Among these are asbolane, chalcophanite, cryptomelane, ernienickelite, lithiophorite, and todorokite (Brand et al. 1998), where the general formulae representing these minerals are shown in Table 3. Depending on the ore, nickel and cobalt are often associated with some of the manganese-bearing minerals in Table 3.

| Mineral | General Formulae | |
|----------------|--|--|
| Asbolane | Mn ⁴⁺ (O,OH) ₂ ·(Co,Ni,Mg,Ca) _x (OH) _{2x} ·nH ₂ O | |
| Chalcophanite | ZnMn ⁴⁺ ₃ O ₇ ·3H ₂ O | |
| Cryptomelane | K(Mn ⁴⁺ 7Mn ³⁺)O ₁₆ | |
| Ernienickelite | NiMn ⁴⁺ ₃ O ₇ ·3H ₂ O | |
| Lithiophorite | (Al,Li)(Mn ⁴⁺ ,Mn ³⁺)O ₂ (OH) ₂ | |
| Todorokite | (Na,Ca,K,Ba,Sr) _{1-x} (Mn,Mg,Al) ₆ O ₁₂ ·3-4H ₂ O | |

 Table 3. General formulae of different manganese-bearing minerals in nickel

 laterite ores.

Note. General formulae obtained from the approved list of International Minerological Association (2018). The general formulae are only used to represent the major elements found in the mineral; therefore, the overall net charge is not zero. Actual formula may vary due to the occurrence of elemental substitution within the mineral structure.

A study by Watling et al. (2011), which involved analysing the minerals found in more than 50 Western Australian nickel laterite ores, revealed that the manganese-bearing minerals can range up to 7% by mass and manganese from 0.05 to 1.9% by mass. Carlson and Simons (1960), Mayze (1999), and Önal and Topkaya (2014) provided elemental composition analysis (Table 4) for nickel laterite ores used in different nickel laterite operations indicating the presence of manganese in each of these ores. The findings by Carlson and Simons (1960), Mayze (1999), Watling et al. (2011), and Önal and Topkaya (2014) demonstrate the presence of manganese in nickel laterite ores.

| | Operations | | | | |
|---------|------------|-------|---------------|--------|---------|
| Element | | | | | |
| | Bulong | Cawse | Murrin Murrin | Caldag | Moa Bay |
| Ni | 1.11 | 1 | 1.24 | 1.215 | 1.35 |
| Со | 0.08 | 0.07 | 0.09 | 0.078 | 0.146 |
| Fe | 20.80 | 18 | 22 | 32.7 | 47.5 |
| Mg | 4.62 | 1.58 | 4 | 1.62 | 1 |
| Al | 2.75 | 1.71 | 2.5 | 1.66 | 4.5 |
| Mn | 0.36 | 0.17 | 0.40 | 0.349 | 0.76 |
| Cr | 0.60 | 0.92 | 0.88 | 1.01 | 2.01 |
| Са | 0.03 | 0.03 | 0.53 | 0.60 | na |

 Table 4. Elemental compositions (%) of nickel laterite ores used in different nickel laterite operations.

na, not available. Note. Data for Bulong, Cawse, and Murrin Murrin from Mayze (1999), for Caldag from Önal and Topkaya (2014), and for Moa Bay from Carlson and Simons (1960).

No studies on the solubility of specific manganese-bearing minerals in nickel laterite ores during the PAL process were found in literature. However, there have been studies by several authors on the leaching behaviour of metals in nickel laterite ores during PAL. It has been reported (Georgiou and Papangelakis 2009, Guo et al. 2011, Önal and Topkaya 2014, Ucyildiz and Girgin 2017) that the leaching of nickel and cobalt from the ore can be greater than 90% and the co-leaching of manganese is up to 90%. The leaching behaviour of metals is attributed to the operating parameters used in PAL, such as acid to ore ratio, temperature, and leaching time, which not only favour the dissolution of nickel and cobalt from nickel laterite ores, but also other metals including manganese (Georgiou and Papangelakis 2009, Guo et al. 2011, Önal and Topkaya 2014, Ucyildiz and Girgin 2017). The dissolution in sulfuric acid of the minerals found in nickel laterite ores can be represented by the simplified equation shown in Equation 2-1, where M_mO_n represents the metal oxide of minerals in nickel laterite ores that reacts with the proton (H^+) released from the acid, including those bearing nickel, cobalt, and manganese, m and n are stoichiometric coefficient, and z is the valency of the metal M.

$$M_m O_n + 2n H^+ \rightarrow m M^{z+} + n H_2 O$$
(2-1)

The elemental composition of PLS from PAL (Table 5) is indicative of the solubility of the manganese-bearing minerals due to the presence of manganese in the PLS.

Clearly, processing nickel laterite ores using sulfuric acid in PAL will generate a PLS containing large amounts of impurities including manganese, which needs to be removed prior to the downstream processing.

| Flomont | | Operations | |
|---------|---------|---------------|--------------|
| Element | Moa Bay | Murrin Murrin | Ravensthorpe |
| Ni | 5.95 | 5.02 | 11.80 |
| Со | 0.64 | 0.37 | 0.41 |
| Mn | 1.98 | 1.62 | 2.20 |
| Fe | 0.80 | 1.53 | 2.20 |
| Mg | 2.76 | 17.2 | 21.0 |
| Al | 2.30 | 1.63 | 0.055 |
| Cr | 0.30 | 0.10 | 0.020 |

Table 5. Elemental compositions (g/L) of PAL-generated PLS from Moa Bay, Murrin Murrin, and Ravensthorpe nickel laterite operations.

Note. Data for Moa Bay from Carlson and Simons (1960), for Murrin Murrin from Motteram et al. (1996), and for Ravensthorpe from White et al. (2006).

In summary, the presence of manganese in PAL-generated PLS from nickel laterite ores is due to the dissolution of manganese-bearing minerals, which is attributed to the operating parameters used in PAL. The removal of manganese as an impurity is therefore particularly important in order to avoid affecting the efficiency of the downstream processing.

2.2 Manganese as an Impurity and its Oxidation States and Stability in Acidic Leach Solutions

After leaching the ore, the major impurities present in the PAL-generated PLS that can form insoluble hydroxides are removed by different nickel laterite operations through partial neutralisation by using alkaline reagents, such as limestone (Kyle and Furfaro 1997, Mason et al. 1997, Grassi et al. 2000, O'Callaghan 2003, White et al. 2006, Llerin et al. 2011, Shibayama et al. 2016), coral mud (Carlson and Simons 1960), and calcrete (Motteram et al. 1996), in order to adjust the pH to between 2.4 and 5.0. Partial neutralisation of the PLS causes iron, aluminium, and chromium to precipitate out of the solution as hydroxides; therefore, leaving manganese as the main impurity in the PLS, which can affect the downstream processing. A few studies report that manganese can either co-precipitate with nickel and cobalt as a hydroxide

during the intermediate precipitation step (Grassi et al. 2000, Oustadakis et al. 2006, Oustadakis et al. 2007) or oxidise to a higher oxidation state during the electrowinning step (Cheng et al. 2000, Donegan 2006) when present in sulfate leach solutions during the downstream processing. The co-precipitation of manganese with nickel and cobalt as a mixed hydroxide will require additional re-leaching and solvent extraction steps in order to separate nickel and cobalt as final products (Grassi et al. 2000, Oustadakis et al. 2006, Oustadakis et al. 2007) while the oxidation of manganese to a higher oxidation state in the electrowinning step will lead to the potential degradation of the organic when the spent electrolyte is recycled back to the solvent extraction circuit (Cheng et al. 2000, Donegan 2006). The likely reason for the co-precipitation or oxidation of manganese to occur during the downstream processing is due to the existence of manganese in different oxidation states and its stability in acidic leach solutions.

Manganese, which can exist as either dissolved or solid species, exhibits different oxidation states depending on the redox potential and pH of the solution. Figure 5 shows a diagram of the electrochemical stability for the different redox species of manganese. The diagram (Figure 5) identifies specific conditions where different redox species of manganese are stable at molar concentrations of 10^{-6} , 10^{-4} , 10^{-2} , and 10^{0} . Each molar concentration is represented by its respective base-10 power in the diagram, -6, -4, -2, and 0. The lines with encircled numbers in the diagram represent the equilibria between two manganese species. Horizontal lines represent pure redox reactions or reactions that are independent of pH, while vertical lines represent a combination of both redox and acid–base reactions, which depends on both the redox potential and pH of the solution.

Figure 5. Potential–pH equilibrium (Pourbaix) diagram for the system of manganese–water, at 25 °C, considering β -MnO₂ (Pourbaix 1966). Encircled numbers in the diagram represent the equilibria between two manganese species while molar concentration is represented by its respective base-10 power in the diagram, -6, -4, -2, and 0. *Content is removed due to copyright restrictions*.

The oxidation states of known dissolved species of manganese include +2, +3, +6, and +7. In highly acidic conditions (~ pH 2) within the stability region of water, which match PAL-generated PLS, only manganous ions (Mn^{2+}) exist as stable dissolved species. In contrast, +3, +6, and +7 oxidation states are unstable in highly acidic conditions. Manganic ion (Mn^{3+}) disproportionates into Mn^{2+} and MnO_2 , as shown in Equation 2-2, while for the +6 oxidation state, manganate ion (MnO_4^{2-}), it reduces to

 Mn^{2+} , as shown in Equation 2-3. In the case of the +7 oxidation state, permanganate ion (MnO_4^{-}) , which is the highest oxidation state of manganese and a strong oxidising agent, is unstable and reacts with water to form manganese dioxide (MnO_2) , as shown in Equation 2-4. Thus, Mn^{3+} , MnO_4^{2-} , and MnO_4^{-} are unlikely to exist as stable dissolved species in PAL-generated PLS.

$$2 \operatorname{Mn}_{(aq)}^{3+} + 2 \operatorname{H}_2 O \leftrightarrow \operatorname{Mn}_{(aq)}^{2+} + \operatorname{MnO}_{2(s)} + 4 \operatorname{H}_{(aq)}^{+}$$
(2-2)

$$MnO_{4(aq)}^{2-} + 8 H_{(aq)}^{+} + 4 e^{-} \leftrightarrow Mn_{(aq)}^{2+} + 4 H_2O$$
(2-3)

$$4 \text{ MnO}_{4(aq)}^{-} + 2 \text{ H}_2\text{O} \leftrightarrow 4 \text{ MnO}_{2(s)} + 4 \text{ OH}_{(aq)}^{-} + 3 \text{ O}_{2(g)}$$
(2-4)

The only solid species of manganese that can exist in highly acidic condition is manganese dioxide (MnO_2), which is the +4 oxidation state of manganese after undergoing hydrolysis. Manganese dioxide (MnO_2) is a dark brown to black solid substance, which can be either anhydrous or hydrous (Pourbaix 1966). In highly acidic conditions, manganese dioxide (MnO_2) can either form from the oxidation of Mn^{2+} (Equation 2-5) or reduction of MnO_4^- (Equation 2-6).

$$Mn_{(aq)}^{2+} + 2 H_2O \leftrightarrow MnO_{2(s)} + 4 H^+ + 2 e^-$$
 (2-5)

$$MnO_{4(aq)}^{-} + 4 H^{+} + 3 e^{-} \leftrightarrow MnO_{2(s)} + 2 H_{2}O$$
(2-6)

Other solid species of manganese, such as $Mn(OH)_2$, Mn_2O_3 , and Mn_3O_4 , are not stable under highly acidic conditions. Each will dissociate to form manganous ion (Mn^{2+}) , as shown in Equations 2-7 to 2-9.

$$\mathsf{Mn}(\mathsf{OH})_{2(s)} \leftrightarrow \mathsf{Mn}^{2+}_{(aq)} + 2 \mathsf{OH}^{-}$$
(2-7)

$$Mn_2O_{3(s)} + 6 H^+ + 2 e^- \leftrightarrow 2 Mn_{(aq)}^{2+} + 3 H_2O$$
 (2-8)

$$Mn_3O_{4(s)} + 8 H^+ + 2 e^- \leftrightarrow 3 Mn_{(aq)}^{2+} + 4 H_2O$$
 (2-9)

Manganese remains as the main impurity in the PLS even after the partial neutralisation step. The presence of manganese in the PLS can affect the efficiency of the downstream processing due to its stability in highly acidic conditions and at the same time its existence in different oxidation states; therefore, the removal of manganese will be beneficial to the process.

2.3 Removal of Manganese from Acidic Leach Solutions by Oxidative Precipitation

Oxidative precipitation of Mn²⁺ in an aqueous solution at highly acidic conditions (~ pH 2) involves the use of a strong oxidising reagent, which must have a standard reduction potential higher than 1.23 V (redox couple Mn^{4+}/Mn^{2+}) that will result in the in-situ formation of a higher oxidation state manganese, Mn⁴⁺, which undergoes hydrolysis to form manganese dioxide (MnO₂) (Equation 2-5). Manganese has different oxidation states (Section 2.2) where the stability is dependent on the pH and potential of the solution. In multi-component solutions such as PAL-generated PLS, one important consideration in the application of oxidative precipitation is to ensure that Ni²⁺ and Co²⁺ will not oxidise together with Mn²⁺. Careful examination of the potential-pH equilibrium (Pourbaix) diagram of manganese (Figure 5) in comparison with those of nickel (Figure 6) and cobalt (Figure 7) shows that at highly acidic conditions, there is a firm thermodynamic basis for the selective precipitation of manganese compared with nickel and cobalt at a high solution potential (e.g. pH 2 and $E^{\Theta} = 1.23 \text{ V}$) where Mn²⁺ oxidises to form MnO₂ while nickel and cobalt remain in the +2 oxidation state and soluble in solution. This is a clear indication that it can be possible to oxidise manganese by oxidative precipitation, even in the presence of nickel and cobalt at highly acidic conditions.

Figure 6. Potential–pH equilibrium (Pourbaix) diagram for the system of nickel– water, at 25 °C (Pourbaix 1966). Encircled numbers in the diagram represent the equilibria between two nickel species while molar concentration is represented by its respective base-10 power in the diagram, –6, –4, –2, and 0. *Content is removed due to copyright restrictions.*

Figure 7. Potential-pH equilibrium (Pourbaix) diagram for the system of cobaltwater, at 25 °C (Pourbaix 1966). Encircled numbers in the diagram represent the equilibria between two cobalt species while molar concentration is represented by its respective base-10 power in the diagram, -6, -4, -2, and 0. *Content is removed due to copyright restrictions.*

2.4 Oxidants for the Oxidative Precipitation of Manganese

Oxidative precipitation has proven to be effective for the removal of manganese using various oxidants (Zhang and Cheng 2007). Since MnO_2 is a strong oxidant with a standard reduction potential (E^{Θ}) of 1.23 V, it would need a stronger oxidant in order to oxidise Mn^{2+} to MnO_2 . Various oxidants that have been applied in the

oxidation of Mn^{2+} are ozone (O₃), Caro's acid (H₂SO₅), and peroxydisulfuric acid $(H_2S_2O_8)$, hypochlorite (ClO⁻), and chlorate (ClO₃⁻) (Zhang and Cheng 2007). These oxidants have either high cost, are highly corrosive resulting in difficulty with control and handling (Zhang and Cheng 2007), or introduce foreign ions that are undesirable and may form toxic by-products; therefore, their use in the oxidative precipitation of Mn²⁺ from PAL-generated PLS can be challenging and unattractive. A few low-cost oxidants are available for the oxidative precipitation of manganese and these include sulfur dioxide and oxygen (SO_2/O_2) gas mixture and potassium permanganate (KMnO₄). However, even if using SO_2/O_2 mixture is low cost, the downside of using the gas mixture is its dependence on the mass transfer and diffusion of the gases to the solution during oxidation, which is affected by the ratio between the dissolved S^{4+} species coming from SO₂ to the dissolved O₂ (Zhang et al. 2010). Gas dispersion and flow rate would also be critical as O_2 is less soluble than SO_2 , making it difficult to maintain oxidising conditions in the solution. In order to effectively oxidise manganese using an SO_2/O_2 mixture, an optimum ratio between SO_2 and O_2 must be maintained with proper dispersion so as not to limit the rate of manganese oxidation (Zhang et al. 2010). The difficulty in using and handling SO_2/O_2 gas mixture will only add to the already challenging purification step of existing nickel laterite operations. This would leave potassium permanganate as a possible option for the oxidative precipitation of manganese from acidic leach solutions. Other than being low cost, another advantage of potassium permanganate is that it has been widely used in water treatment (Environmental Protection Agency 1999) and at the same time it will not introduce any new ions to the highly contaminated PLS.

2.4.1 Potassium Permanganate as Oxidant

Most studies conducted on the oxidative precipitation of manganese using potassium permanganate focused on applications used for water treatment and industrial waste solutions, such as those from mineral processing. There is limited information available on applications in solutions similar to PAL-generated PLS.

The literature review revealed no studies on the application of potassium permanganate (KMnO₄) to PAL-generated PLS. Potassium permanganate is widely

used in water treatment for the oxidation of iron and manganese as well as other compounds that affect the taste and odour of water (Environmental Protection Agency 1999). The net ionic reaction for the oxidative precipitation of Mn^{2+} with permanganate (MnO_4^{-}) is shown in Equation 2-26, which shows that Mn^{2+} will precipitate as MnO_2 in the presence of MnO_4^{-} .

$$3 \operatorname{Mn}_{(aq)}^{2+} + 2 \operatorname{MnO}_{4(aq)}^{-} + 2 \operatorname{H}_{2}O \leftrightarrow 5 \operatorname{MnO}_{2(s)} + 4 \operatorname{H}^{+}$$
(2-26)

2.5 Variables Influencing the Oxidative Precipitation of Manganese(II) by Potassium Permanganate

The oxidative precipitation of Mn²⁺ using potassium permanganate is influenced by pH (Adams 1960, Phatai et al. 2010, Freitas et al. 2013, Elsheikh et al. 2017) and oxidant amount (Adams 1960, Heviankova and Bestova 2007, Phatai et al. 2010, Freitas et al. 2013, Phatai et al. 2014, Macingova et al. 2016, Elsheikh et al. 2017). Much of the current literature using potassium permanganate for the oxidative precipitation of manganese concerns water treatment, but it has been applied as well to acid mine water, which is sometimes known as acid mine drainage (AMD). The aim in the oxidative precipitation of manganese from either raw water or acid mine water is to ensure that the concentration of manganese will be within the acceptable limit set by various international legislation. For instance, raw water should have less than 0.10 mg/L of manganese based on global standards (Phatai et al. 2014), while for acid mine water, it should be less than 1.0 mg/L, as in the case of Brazilian legislation (Freitas et al. 2013). If the present study could achieve a comparable concentration of manganese close to the acceptable limit in either raw water (<0.10 mg/L) or acid mine water (<1.0 mg/L), it will be low enough so as to not to affect the efficiency of downstream processing of PAL-generated PLS.

2.5.1 Influence of pH

It is known that the oxidative precipitation of manganese is influenced by changes in pH of the solution (Figure 5). In addition, changing the pH of the solution can cause a change in the standard reduction potential (E^{Θ}) of permanganate when potassium permanganate is used as an oxidant. At highly acidic conditions, permanganate has a

standard reduction potential (E^{Θ}) of 1.51 V and as the pH changes to basic conditions, E^{Θ} decreases to 0.59 V. At basic conditions, manganese does not only form MnO₂ but can also form other stable solid species, such as Mn₂O₃, Mn₃O₄, and Mn(OH)₂, even at a low solution potential.

Studies investigating the influence of pH on the oxidative precipitation of manganese using potassium permanganate are mostly found for water treatment (Adams 1960, Phatai et al. 2010, Elsheikh et al. 2017), but it has also been applied for the treatment of AMD (Freitas et al. 2013). A study on the oxidative precipitation of manganese from raw water by Adams (1960) showed that soluble manganese in high concentrations can rapidly and completely oxidise at a high pH range. The pH range required to completely oxidise less than 0.5 mg/L of manganese is between pH 7.2 and pH 7.4, but as the concentration of manganese increases to between 2 and 4 mg/L, the pH range for effective manganese oxidation was between pH 8.0 and pH 8.3.

A similar study investigating the influence of pH on the oxidative precipitation of manganese from MnSO₄ solutions (90 – 150 mg/L Mn²⁺) and actual AMD solutions at pH 3, 5, and 7 was carried out by Freitas et al. (2013) as well as from synthetic ground water at pH 8 to 9 by Phatai et al. (2010) and at pH 7 to 9 by Elsheikh et al. (2017). Freitas et al. (2013) reported nearly complete removal of manganese to less than 0.1 mg/L of the acceptable limit from MnSO₄ solutions at pH 5 and 7 while at pH 3, the concentration remained at 0.3 mg/L. The increase in manganese removal was attributed to the increase in the Mn^{2+} oxidation rate as the pH increases (Van Benschoten et al. 1992, Freitas et al. 2013). In addition, the same trend was observed by Freitas et al. (2013) using actual AMD solutions. In one of the solutions (102.6 mg/L Mn, 2.8 mg/L Fe, 3.2 mg/L Zn, 9.5 mg/L U, 251.9 mg/L Ca, 194.8 mg/L Al, and 65.2 $mg/L F^{-}$), almost complete removal of manganese (99.7%) was obtained at pH 7 while only 85.4% at pH 3 (Freitas et al. 2013). The results in the investigation of Elsheikh et al. (2017) and Phatai et al. (2010) using synthetic ground water were also consistent with the trend, despite using different test solutions and pH ranges. Manganese removal increased from 75% to 85% (Elsheikh et al. 2017) and from 80% to almost complete removal (Phatai et al. 2010) when the pH was increased from pH 8 to 9.

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Clearly, operating at basic conditions will be beneficial in enhancing the oxidative precipitation of manganese. However, what is not yet clear is the influence of pH on the oxidative precipitation of manganese when a solution is maintained at highly acidic conditions, as in the case of PAL-generated PLS. The reason for this is that operating at basic conditions will not be beneficial when applied to PAL-generated PLS due to the possibility for nickel and cobalt to co-precipitate with manganese by forming metal hydroxides. Raising the pH to basic conditions will promote the precipitation of nickel and cobalt as metal hydroxides due to the decrease in the solubility of nickel hydroxide and cobalt hydroxide. It is for this reason that, in the present study, it is of interest to determine whether the oxidative precipitation of manganese from PAL-generated PLS using potassium permanganate will still be an effective technique to selectively separate manganese from nickel and cobalt by taking into consideration the firm thermodynamic basis for the oxidative precipitation of the manganese based on potential–pH equilibrium (Pourbaix) diagram of the manganese–water system (Figure 5) at highly acidic pH conditions.

2.5.2 Influence of Oxidant Amount

Another variable that is known to have an influence on the oxidative precipitation of manganese is the amount of potassium permanganate used since it will affect the availability of permanganate ion (MnO_4^-) needed for the oxidative precipitation of Mn^{2+} (Equation 2-26). The stoichiometric amount of MnO_4^- required to completely oxidise manganese is 2:3 (Equation 2-26). Therefore, for 1 mg (1.82 x 10⁻⁵ mol) of manganese to completely oxidise, it will require 1.92 mg (1.21 x 10⁻⁵ mol) of potassium permanganate. Several studies have investigated the influence of potassium permanganate amount on the oxidative precipitation of manganese from water treatment (Phatai et al. 2014, Elsheikh et al. 2017) and acid mine water treatment (Heviankova and Bestova 2007, Macingova et al. 2016).

A study by Phatai et al. (2014) on the oxidative precipitation of manganese and iron from synthetic groundwater using potassium permanganate investigated the influence of the amount of potassium permanganate in single (Mn = 0.314 mg/L) and dual (Mn = 0.314 mg/L and Fe = 0.048 mg/L) metal systems at pH 8. The stoichiometric amount of potassium permanganate required on the oxidative precipitation of the single component system was 0.603 mg/L while for dual component system, it was 0.648 mg/L (Phatai et al. 2014). Phatai et al. (2014) reported that for the single component system, the removal of manganese increased from 84.6% to 92.2% when the amount of potassium permanganate was increased from 0.603 mg/L to 0.648 mg/L. The result was attributed by Phatai et al. (2014), to the increase in conversion of Mn²⁺ to MnO₂ when higher amount (0.648 mg/L) of potassium permanganate was used instead of 0.603 mg/L to oxidise 0.314 mg/L of manganese. The ratio of reacting MnO₄⁻ to Mn²⁺ will increase as more potassium permanganate is added to the solution. A comparable trend was also reported by Phatai et al. (2014) in the dual component system. Phatai et al. (2014) noted that the result was higher in the dual component system compared with the single component system due to the precipitation of iron as Fe(OH)₃, which provided surface for the autocatalytic oxidation of Mn²⁺, and therefore enhancing the oxidative precipitation of Mn²⁺.

In the same vein, Elsheikh et al. (2017), in their study on the oxidative precipitation of iron (1.50 mg/L) and manganese (1.0 mg/L) from a simulated groundwater at pH 7, yielded the same increasing trend. As the potassium permanganate amount increased from 1 mg/L to 3 mg/L, which was near the theoretical stoichiometric amount, the oxidative precipitation of manganese increased from 50% to 80%. The removal, however, started to decrease to approximately 70% when the amount was increased to 4 mg/L, which is roughly 1.2 times the theoretical stoichiometric amount required. The drop in the amount of manganese precipitated was consistent with Equation 2-27, where reduction of permanganate resulted in the generation of Mn²⁺, which was also the reason attributed by Elsheikh et al. (2017) for the decrease in oxidative precipitation of manganese; therefore, this suggests that the amount of potassium permanganate must carefully be considered in order to ensure the complete oxidative precipitation of Mn²⁺ while avoiding the generation of additional Mn^{2+} due to the reduction of permanganate. In the potassium permanganate amounts investigated, up to 97% iron removal was consistently attained by Elsheikh et al. (2017).

$$MnO_{4(aq)}^{-} + 8 H_{(aq)}^{+} + 5 e^{-} \leftrightarrow Mn_{(aq)}^{2+} + 4 H_2O$$
(2-27)

Studies by Heviankova and Bestova (2007) and Macingova et al. (2016) also investigated the influence of the amount of potassium permanganate, but for the oxidative precipitation of manganese from acid mine water. In the studies by Heviankova and Bestova (2007) and Macingova et al. (2016) on the oxidative precipitation of manganese from acid mine water containing 5.33 mg/L of Mn^{2+} and 16.50 mg/L of Mn^{2+} , respectively, at pH 7, they observed that there was a limit to the amount of potassium permanganate that could effectively oxidise Mn²⁺. Beyond the limit, the complete oxidative precipitation of Mn²⁺ was affected due to the generation of Mn²⁺, which is supported by the observation of Elsheikh et al. (2017) using simulated groundwater. Although the observations by Heviankova and Bestova (2007) and Macingova et al. (2016) using acid mine water were consistent with those observed by Elsheikh et al. (2017) for groundwater, the values reported for the limiting amount of potassium permanganate varied. The amounts of potassium permanganate reported by Macingova et al. (2016) and Elsheikh et al. (2017) were only stoichiometric while in the case of Heviankova and Bestova (2007), the limiting value was 1.2 times the stoichiometric amount. The differences in the reported values of the various authors may be attributed to the presence of different metal components that were present together with manganese in the feed solutions used. It is for this reason that the present study will determine the amount of potassium permanganate that will be needed to ensure the complete oxidative precipitation of manganese from PAL-generated PLS.

2.7 Summary of the Review

The major findings of this review may be summarised as follows:

- The dissolution of manganese-bearing minerals in nickel laterite ores is attributed to the operating parameters used in PAL, which results in a large amount of manganese in PAL-generated PLS.
- At highly acidic conditions (~pH 2), manganese can exist either as manganese(II) or manganese dioxide (MnO₂).

- Potassium permanganate, SO₂/O₂ gas mixture, ozone, Caro's acid, and peroxydisulfuric acid are strong oxidants available for the oxidative precipitation of manganese(II). Among the available oxidants, however, SO₂/O₂ gas mixture and potassium permanganate are the options available as inexpensive oxidants compared with ozone, Caro's acid, and peroxydisulfuric acid.
- Between SO₂/O₂ gas mixture and potassium permanganate, the latter provides more interest to test the suitability of the technique using PALgenerated PLS compared with the former, which is sensitive to mass transfer, diffusion, gas dispersion, and flow rate that would require higher operational costs due to the need to constantly control different variables.
- Two of the critical variables that have an influence on the oxidative precipitation of manganese(II) by potassium permanganate are pH and amount of potassium permanganate. Operating at pH greater than seven (pH >7) will enhance the oxidative precipitation of manganese(II) by increasing the oxidation rate and promote, not only the formation of MnO₂ but also those of Mn₂O₃, Mn₃O₄, and Mn(OH)₂. The amount of potassium permanganate influences the oxidative precipitation of manganese(II) by increasing the conversion of manganese(II) to MnO_2 due to an increase in the ratio of available MnO_4^- that reacts with manganese(II). It was concluded, therefore, that a need exists to determine whether the oxidative precipitation of manganese(II) from PAL-generated PLS using potassium permanganate will still be an effective technique to selectively separate manganese(II) from nickel(II) and cobalt(II) while the pH is at highly acidic conditions and at the same time determine the amount of potassium permanganate needed to ensure the complete oxidative precipitation of manganese(II) from PALgenerated PLS.

Chapter 3

MATERIALS AND METHODS

3.1 Reagents

The reagents used in the experimental work are listed in Table 6. The reagents used were of various chemical grades: ACS Reagent grade, Laboratory Reagent (LR) grade, Analytical Reagent (AR) grade, and Extra Pure (EP).

| Reagents | Formula | Grade | Purity (%) | Supplier |
|-----------------------------------|--------------------------------------|----------------------------|--------------------|-------------------------------------|
| Nickel sulfate hexahydrate | NiSO₄·6H₂O | ACS | 98.00 | Acros Organics |
| Cobalt sulfate heptahydrate | CoSO ₄ ·7H ₂ O | LR | 96.00 | Chem-Supply |
| Manganese sulfate monohydrate | MnSO₄·H₂O | AR | 99.80 | Chem-Supply |
| Magnesium sulfate heptahydrate | MgSO₄·7H₂O | EP | 99.00 | Acros Organics |
| Calcium sulfate dihydrate | $CaSO_4 \cdot 2H_2O$ | AR | 98.00 | Chem-Supply |
| Ferrous sulfate heptahydrate | FeSO ₄ ·7H ₂ O | AR | 99.00 | Chem-Supply |
| Potassium permanganate | KMnO4 | LR | 99.00 | Chem-Supply |
| Sodium hydroxide | NaOH | AR | 98.00 | Rowe Scientific |
| Sulfuric acid | H_2SO_4 | AR | 98.0 | Ajax Finechem |
| Nitric acid | HNO ₃ | (Univar) AR (Univar) | m/m 70.0 m/m | Pty Ltd Ajax Finechem Pty Ltd |

Table 6. Reagents used in the experimental work.

3.2 Test Solutions

The following test solutions were used in the current study: sulfuric acid solution (1 mol/L H_2SO_4), nitric acid solution (4% m/m HNO₃), sodium hydroxide (1 mol/L and 5 mol/L NaOH), and potassium permanganate (0.2 mol/L and 0.4 mol/L KMnO₄). Both

the sulfuric acid and nitric acid solutions were prepared by dilution from concentrated reagents using deionised (DI) water following standard procedures. Sodium hydroxide and potassium permanganate were prepared by dissolving the pellets and crystals, respectively, in a sufficient amount of deionised water to ensure complete dissolution before the volumetric flasks were filled to the mark.

3.3 Preparation of Synthetic and Actual Pregnant Leach Solutions

Synthetic pregnant leach solutions (PLS) that simulated those produced from the pressure acid leaching (PAL) process in terms of the concentration of the major elements both at the partial neutralisation and post-partial neutralisation steps were prepared by dissolving the respective sulfate salts of the metals in a minimum amount of deionised water until the salts were completely dissolved. The volume was then adjusted to nearly that of the volumetric flask but still allowing the adjustment of the pH of the solution to approximately pH 2 by adding sulfuric acid (1 mol/L) and then made to the volume using deionised water. Typical elemental compositions of the synthetic solutions prepared to simulate the PLS solutions, both partially neutralised and post-partially neutralised, are shown in Table 7. The former refers to the solution that still contains residual iron, mainly iron(II), after the removal of iron(III) during the first iron removal step. The latter refers to the solution after all iron has been removed by oxidising the iron(II) to iron(III) before precipitating by adjusting the pH to approximately pH 3.5. The major impurities in the synthetic partially neutralised PLS were manganese(II) and iron(II), which simulated the composition of the PLS after the removal of iron(III), aluminium(III), and chromium(III). In the case of the synthetic post-partially neutralised PLS, the solution simulated an iron-free solution where only manganese(II) remained as the major impurity.

| _ | Concentration (g/L) | | | | |
|---------|---------------------|----------------|--|--|--|
| Element | Partially | Post-Partially | | | |
| | Neutralised | Neutralised | | | |
| Ni | 3.40 | 3.50 | | | |
| Со | 0.30 | 0.30 | | | |
| Mn | 2.00 | 2.50 | | | |
| Mg | 17.10 | 10.00 | | | |
| Са | 0.70 | 0.50 | | | |
| Fe | 2.60 | - | | | |

Table 7. Typical compositions of synthetic feed solution.

The actual PLS was supplied by a nickel laterite processing plant located in Western Australia. The PLS available was only partially neutralised and additional pretreatment of the PLS was carried out prior to the experimental work in order to prepare a post-partially neutralised PLS. Typical elemental compositions of the actual partially neutralised and post-partially neutralised PLS are shown in Table 8.

| | Concentration (g/L) | | | | |
|---------|---------------------|----------------|--|--|--|
| Element | Partially | Post-Partially | | | |
| | Neutralised | Neutralised | | | |
| Al | 0.73 | 0.42 | | | |
| Ca | 0.44 | 0.54 | | | |
| Со | 0.33 | 0.31 | | | |
| Cr | 0.06 | 0.004 | | | |
| Cu | 0.01 | 0.01 | | | |
| Fe | 2.46 | 0.28 | | | |
| Mg | 19.49 | 16.62 | | | |
| Mn | 1.90 | 1.66 | | | |
| Ni | 3.25 | 2.93 | | | |
| Zn | 0.04 | 0.08 | | | |

Table 8. Typical compositions of actual PLS

The concentrations of the metals in both synthetic and actual PLS were determined using inductively coupled plasma optical emission spectrophotometer (ICP-OES) (Agilent 5100).

3.4 Experimental Design

The main experimental response investigated in this work was the amount of precipitated manganese(II). In addition, since nickel and cobalt are the valuable

products, their responses were also included. In case of the partially neutralised PLS, the amount of precipitated iron(II) was also investigated since the residual iron(II) that is present together with manganese(II) in the PLS will readily oxidise and be removed along with manganese(II). The reason for the oxidation is that, besides MnO_2 (Equation 3-1), the standard reduction potential (E^{Θ}) for the redox couple Fe(III)/Fe(II) (Equation 3-2) is also lower than that of MnO_4^- ; therefore, in the presence of MnO_4^- , which has an E^{Θ} of 1.69 V (Equation 2-6), Equation 3-1 and Equation 3-2 will proceed in the reverse direction (i.e. Fe^{2+} to Fe^{3+} and Mn^{2+} to MnO_2). However, due to the much lower E^{Θ} of Fe³⁺ compared with MnO_2 , the reverse of Equation 3-2 will proceed first before the reverse of Equation 3-1.

$$MnO_{2(s)} + 4 H^{+} + 2 e^{-} \leftrightarrow Mn_{(aq)}^{2+} + 2 H_2O$$
 $E^{\theta} = 1.23 V (3-1)$

$$Fe_{(aq)}^{3+} + e^- \leftrightarrow Fe_{(aq)}^{2+} + 2 H_2O$$
 $E^{\theta} = 0.77 V (3-2)$

Screening experiments were performed using an unreplicated full factorial design to evaluate the variables that may have significant effects on the responses (amount of precipitated metal ions) investigated in the current study. The advantage of using unreplicated full factorial design over one-variable-at-a-time (OVAT) for screening experiments is that it allows the determination of the main effects of each variable to the response as well as any interaction effects that may exist among the various variables using a fixed number of experimental runs even with limited resources and time. The variables, which often affect chemical reactions, chosen for investigation that may influence the oxidative precipitation of manganese(II) were temperature, molar ratio, pH, and agitation speed. The variables investigated for the oxidative precipitation experiments were evaluated at two levels: low (-1) and high (+1). The levels used for each variable are detailed in Table 9. These levels were then systematically changed for each experimental run of the unreplicated full factorial design by following the experimental design matrix shown in Table 10.

Table 9. Variables and selected levels in unreplicated 2⁴ full factorial design.

| Variables | Name | Low (–1) | High (+1) | Unit |
|-----------|-----------------|-------------------------|-----------------------------|------|
| А | Temperature | Ambient ^a | 35 | °C |
| В | Molar ratio | 85% ^b | Stoichiometric ^c | - |
| С | рН | Unadjusted ^d | 3.00 | - |
| D | Agitation speed | 50 | 200 | rpm |

^aActual ambient temperature was noted down based on the temperature probe reading of the PLS during the experimental work.

^bLow level of 85% refers to 85% of the stoichiometric molar ratio required for complete oxidation.

^cHigh level refers to the stoichiometric molar ratio required for complete oxidation.

^dUnadjusted refers to the natural pH inherent to the reaction between the synthetic PLS and the oxidant.

| Temperature (A) | Molar ratio (B) | рН (С) | Agitation speed (D) |
|-----------------|-----------------|--------|---------------------|
| -1 | -1 | -1 | -1 |
| +1 | -1 | -1 | -1 |
| -1 | +1 | -1 | -1 |
| +1 | +1 | -1 | -1 |
| -1 | -1 | +1 | -1 |
| +1 | -1 | +1 | -1 |
| -1 | +1 | +1 | -1 |
| +1 | +1 | +1 | -1 |
| -1 | -1 | -1 | +1 |
| +1 | -1 | -1 | +1 |
| -1 | +1 | -1 | +1 |
| +1 | +1 | -1 | +1 |
| -1 | -1 | +1 | +1 |
| +1 | -1 | +1 | +1 |
| -1 | +1 | +1 | +1 |
| +1 | +1 | +1 | +1 |

Table 10. Experimental matrix of unreplicated 2⁴ full factorial design.

Temperature was considered as one of the variables that may affect the oxidative precipitation of manganese(II) since most chemical reactions are influenced by temperature. The low and high levels of temperature (A) were set to ambient and 35°C, respectively. The difference between these two temperatures should be sufficient to produce observable difference in the experimental response if temperature is a significant variable. Since the oxidative precipitation of manganese(II) is dependent on the amount of permanganate ion (MnO_4^-) available for complete oxidation, their molar ratio (B) was also considered. In the experiments wherein the test sample was the post-partially neutralised PLS, the molar ratio was the ratio between the total amount in moles of MnO_4^- and manganese(II) (denoted

as mol MnO₄⁻: mol Mn(II)). The low and high levels were set to 0.57 and 0.67, respectively. The high level was maintained at the stoichiometric molar ratio between MnO₄⁻ (2 mol) and manganese(II) (3 mol) based on Equation 2-26, while the low level was 85% of the high level molar ratio to produce an observable difference in the experimental response. In the case of partially neutralised PLS, this ratio, however, incorporated the amount in moles of iron(II) together with the amount in moles of manganese(II) (denoted as mol MnO₄⁻: total mol Fe(II) and Mn(II)) to take into consideration the oxidation of iron(II) (Equation 3-3) before manganese(II) due to a much lower standard reduction potential (E^{Θ}). The low and high levels were set to 0.43 and 0.50, respectively. The high level was the stoichiometric molar ratio between the total moles of MnO₄⁻ (3 mol) required to completely oxidise the total moles of iron(II) and manganese(II) (6 mol) based on Equation 2-26 and Equation 3-3 with the overall equation shown in Equation 3-4, while the low level was 85% of the high level molar ratio.

$$3 \operatorname{Fe}_{(\operatorname{aq})}^{2+} + \operatorname{MnO}_{4(\operatorname{aq})}^{-} + 4 \operatorname{H}^{+} \leftrightarrow 3 \operatorname{Fe}_{(\operatorname{aq})}^{3+} + \operatorname{MnO}_{2(\operatorname{s})} + 2 \operatorname{H}_{2} \operatorname{O}$$
(3-3)

$$3 \operatorname{Fe}_{(\operatorname{aq})}^{2+} + 3 \operatorname{Mn}_{(\operatorname{aq})}^{2+} + 3 \operatorname{MnO}_{4(\operatorname{aq})}^{-} \leftrightarrow 3 \operatorname{Fe}_{(\operatorname{aq})}^{3+} + 6 \operatorname{MnO}_{2(\operatorname{s})}$$
(3-4)

Another variable considered was pH as it has an influence in most redox reactions and this can be seen in the potential–pH equilibrium (Pourbaix) diagram for the manganese–water system (Figure 5). The low level of pH (C) was the natural pH of the solution, meaning no pH adjustment was carried out during the experiment while the high level was set at pH 3, which was a convenient pH to operate at for the oxidative precipitation of manganese(II) as it avoided the precipitation of nickel(II) and cobalt(II). Lastly, agitation speed (D) was considered to ensure homogeneous mixing of the solution during the experimental work. The low and high levels used for agitation speed were set to 50 and 200 rpm, respectively, and these values ensured that there was still noticeable mixing of the solution while the experiment was performed. The effect of agitation in nucleation and particle size, however, was not investigated, but may be relevant during the characterisation of the precipitate in future studies. Prior to the implementation of the unreplicated full factorial design for the screening experiments, an exploratory experiment was conducted using a constantly agitated (50 rpm) synthetic post-partially neutralised PLS at ambient temperature (19.5 °C). The pH of the solution was not adjusted throughout the experiment and the amount of potassium permanganate used was at stoichiometric molar ratio (0.67). Samples for assay were taken at 1 and 2 hours. The results indicated that only 1.51% (from 96.88% to 98.39%) additional manganese(II) was precipitated in the second hour but at the expense of losing 28.57% (from 4.56% to 33.13%) additional nickel(II) and 17.82% (from 48.79% to 66.61%) additional cobalt(II). Because of the increased losses of nickel(II) and cobalt(II) over time, subsequent experiments were limited to 1 hour.

An unreplicated full factorial design is a saturated experimental design. This means that only the main and interaction effects as well as the general mean can be estimated. There are no degrees of freedom left in a saturated experimental design to estimate the experimental error. A workaround in this limitation is to analyse the unreplicated full factorial design on the assumption of the sparsity-of-effects principle and then plot the estimated effects in a normal probability plot (Daniel 1959, Montgomery 2013). The sparsity-of-effects principle states that main effects (single variable) and low-order interaction effects (two-variable) are dominant in a factorial design and any higher order interactions (three-variable and up) are negligible (Montgomery 2013). The normal probability plot will assist in screening the variables that may have significant effect on the responses being investigated. Negligible or not significant effects are normally distributed and will fall along the normal distribution line in the normal probability plot. In contrast, significant effects are not normally distributed and will fall outside the normal distribution line. In order to generate normal probability plots, the data gathered were encoded in Minitab® 18 and the significance of the variables was tested against a significance level (α) of 0.05. The significance of the variables was examined from the normal probability plots before further experimental work was carried out to optimise the variables using the OVAT approach. Afterwards, the optimum values determined from the OVAT approach were applied to an actual PLS as test solution.

3.5 Experimental Set-up for the Oxidative Precipitation Test Procedure

All oxidative precipitation tests were performed inside a fume hood to control any exposure to hazardous substances produced during the experiment. The tests were carried out in a 150 mL five-neck glass flask (Pine Research RRPG022) immersed in a water bath. Heating and agitating for the experiment were facilitated using a hotplate with magnetic stirrer (VELP Scientifica AREX heating and magnetic stirrer F20500163). The temperature of the PLS was monitored using a temperature probe connected to a pH meter to ensure that the temperature throughout the experiment was maintained within the required operating temperature of the experimental run. Additionally, the pH of the PLS was monitored using a pH meter equipped with pH probe (TPS WP-80 pH meter, lonode IJ44 pH probe). A 50 mL burette was used for the addition of the potassium permanganate (KMnO4). Figure 8 shows the experimental set-up used in the experimental work.



Figure 8. Experimental set-up for the oxidative precipitation tests.

3.6 Oxidative Precipitation Test Procedure

Oxidative precipitation tests were conducted using the experimental set-up described in Section 3.5. The tests were carried out by transferring 50 mL of the PLS into a constantly agitated 150 mL five-neck glass flask. The temperature of the PLS inside the five-neck glass flask was adjusted using a water bath until thermal equilibrium was reached. The temperature of the PLS was constantly monitored using the temperature probe and to ensure that the required temperature was maintained, the temperature controller of the hot plate was manually adjusted. Potassium permanganate (KMnO₄) was then added in a drop-wise manner using the 50 mL burette to minimise any local concentration effect in the feed solution. Afterwards, the pH of the solution was adjusted by adding sodium hydroxide until the desired equilibrium pH was reached and monitored using the pH meter with the addition of sodium hydroxide as required to maintain a constant pH value. The same test procedure was performed for the experimental work using the OVAT approach for the optimisation of the variables that were identified to be significantly affecting the oxidative precipitation of manganese(II).

3.7 Assay of Metals

The concentrations of the metal ions in the solution were analysed using ICP-OES. The samples (10 mL) for assay were collected during the oxidative precipitation tests, vacuum filtered (5.5 cm no. 1 Whatman qualitative filter paper) and then diluted with nitric acid (4% m/m) to quench the reaction and prevent further precipitation. The addition of nitric acid also ensured that the samples had the same acid matrix as the ICP-OES calibration standards during analysis. No washing of the solids produced from the vacuum filtered 10 mL samples were carried out as it was observed that the amounts of solids were very small.

Chapter 4

RESULTS AND DISCUSSION

4.1 Screening of Effects of Experimental Variables on the Oxidative Precipitation of Manganese(II)

An unreplicated 2⁴ full factorial design (Section 3.4) was implemented in order to screen the main and interaction effects of the four selected variables in the oxidative precipitation of manganese(II) from partially neutralised and post-partially neutralised PLS. The four variables selected were temperature (A), molar ratio (B), pH (C), and agitation speed (D). A set of oxidative precipitation experiments was carried out following the experimental procedure outlined in Section 3.6. Synthetic solutions simulating partially neutralised and post-partially neutralised PLS with typical concentrations enumerated in Section 3.3 were used for the oxidative precipitation experiments instead of an actual PLS since the elemental compositions of synthetic solutions were better defined. Moreover, using a synthetic PLS enabled a straightforward identification of the significant variables that may affect the oxidative precipitation of manganese in the absence of any unassayed components that might be present in an actual PLS.

The operating conditions for each experimental run were defined using the selected levels and experimental matrix described in Section 3.4. The experimental plan as well as the percent precipitated metal ions for the partially neutralised and post-partially neutralised PLS are shown in Table 11 and Table 12, respectively. The percent of precipitated metal ions were calculated from the initial minus the final metal ion concentration after the dilution effect had been factored in due to the addition of the reagents.

| Tomporatura | Malar | | Agitation | | | | |
|----------------------|--------------------|-------------------------|-----------|--------|----------|----------|--------|
| (°C) | ratio ^a | рН | speed | Precip | itated N | letal Io | ns (%) |
| | | | (rpm) | | | | |
| Α | В | С | D | Mn | Fe | Ni | Со |
| Ambient ^b | 0.43 | Unadjusted ^c | 50 | 75.64 | 61.21 | 1.03 | 5.00 |
| 35 | 0.43 | Unadjusted | 50 | 74.80 | 63.29 | 0.27 | 4.29 |
| Ambient | 0.50 | Unadjusted | 50 | 86.44 | 63.88 | 6.93 | 11.01 |
| 35 | 0.50 | Unadjusted | 50 | 83.20 | 64.55 | 0.00 | 4.42 |
| Ambient | 0.43 | 3 | 50 | 77.38 | 98.98 | 3.10 | 7.72 |
| 35 | 0.43 | 3 | 50 | 76.07 | 99.51 | 0.00 | 3.82 |
| Ambient | 0.50 | 3 | 50 | 84.96 | 99.33 | 1.18 | 6.62 |
| 35 | 0.50 | 3 | 50 | 85.54 | 99.73 | 1.76 | 9.28 |
| Ambient | 0.43 | Unadjusted | 200 | 76.44 | 60.41 | 0.83 | 4.59 |
| 35 | 0.43 | Unadjusted | 200 | 74.86 | 62.65 | 0.01 | 4.15 |
| Ambient | 0.50 | Unadjusted | 200 | 83.57 | 62.97 | 0.00 | 3.15 |
| 35 | 0.50 | Unadjusted | 200 | 84.42 | 66.72 | 6.90 | 11.95 |
| Ambient | 0.43 | 3 | 200 | 75.13 | 99.01 | 0.00 | 2.50 |
| 35 | 0.43 | 3 | 200 | 75.67 | 99.66 | 0.00 | 2.58 |
| Ambient | 0.50 | 3 | 200 | 84.78 | 99.66 | 1.04 | 6.74 |
| 35 | 0.50 | 3 | 200 | 85.82 | 99.66 | 2.20 | 9.29 |

Table 11. Experimental plan and responses of unreplicated 2⁴ full factorialdesign using a partially neutralised PLS as feed.

^aMolar ratio for a partially neutralised PLS is denoted as mol MnO₄⁻: total mol Fe(II) and Mn(II); ^bAmbient temperature range =

23.9–26.7 °C ; Unadjusted pH range = 1.71–1.85

| Temperature (°C) | Molar ratio ^a | рН | Agitation speed (rpm) | Precipitated Metal Ions (%) | | |
|----------------------|-----------------------------|-------------------------|-----------------------------|--------------------------------|-------|-------|
| Α | В | С | D | Mn | Ni | Со |
| Ambient ^b | 0.57 | Unadjusted ^c | 50 | 96.60 | 5.02 | 41.05 |
| 35 | 0.57 | Unadjusted | 50 | 94.86 | 9.18 | 44.53 |
| Ambient | 0.67 | Unadjusted | 50 | 99.32 | 9.71 | 60.14 |
| 35 | 0.67 | Unadjusted | 50 | 99.45 | 16.42 | 67.13 |
| Ambient | 0.57 | 3 | 50 | 97.36 | 8.01 | 63.98 |
| 35 | 0.57 | 3 | 50 | 96.07 | 12.91 | 62.31 |
| Ambient | 0.67 | 3 | 50 | 99.76 | 18.39 | 84.20 |
| 35 | 0.67 | 3 | 50 | 99.83 | 14.79 | 87.20 |
| Ambient | 0.57 | Unadjusted | 200 | 96.01 | 9.66 | 43.59 |
| 35 | 0.57 | Unadjusted | 200 | 95.53 | 16.01 | 48.22 |
| Ambient | 0.67 | Unadjusted | 200 | 99.80 | 16.52 | 61.72 |
| 35 | 0.67 | Unadjusted | 200 | 99.30 | 13.80 | 67.30 |
| Ambient | 0.57 | 3 | 200 | 97.51 | 13.88 | 65.14 |
| 35 | 0.57 | 3 | 200 | 96.08 | 13.68 | 62.74 |
| Ambient | 0.67 | 3 | 200 | 99.73 | 19.31 | 81.34 |
| 35 | 0.67 | 3 | 200 | 99.75 | 19.54 | 85.77 |

Table 12. Experimental plan and responses of unreplicated 2⁴ full factorialdesign using a post-partially neutralised PLS as feed.

^aMolar ratio for a post-partially neutralised PLS is denoted as mol MnO₄⁻:mol Mn(II); ^bAmbient temperature range = 19.0– 24.5 °C ;^cUnadjusted pH range = 1.65–1.71

4.1.1 Synthetic Partially Neutralised PLS as Feed Solution in Determining Effects of Experimental Variables on Oxidative Precipitation of Manganese(II)

The effects of the variables were estimated by constructing normal probability plots for the precipitated metals (%) of manganese(II) and iron(II) versus standardized effects, as shown in Figures 9 and 10, respectively. On these plots, a significant effect is indicated by a square while a nonsignificant effect is indicated by a circle.





Figure 9 shows that the effect of molar ratio (B) on the precipitation of manganese(II) was significant. The significance of the effect of molar ratio (B) with a standardized effect (SE) of 17.664 was probably owing to the change in the number of moles of manganese(II) that underwent oxidative precipitation produced by the change in molar ratio, which manifested in the amount of precipitated manganese(II). Increasing the molar ratio will cause an increase in available MnO_4^- that is being consumed for the oxidative precipitation of manganese(II) and therefore resulting in an increase in amount of precipitated manganese(II).



Figure 10. Normal probability plot of the main and interaction effects for the precipitation of iron(II) (%) from a synthetic partially neutralised PLS ($\alpha = 0.05$).

In the case of the precipitation of iron(II), Figure 10 shows that the effects of temperature (A), molar ratio (B), and pH (C) as well as the interaction effects between temperature and pH (AC) and molar ratio and pH (BC) were significant. The significance of the effect of temperature (A) with SE of 3.757 was probably owing to the considerable difference in the change in solubility of the iron compound produced by the change in temperature, which manifested in the amount of precipitated iron(II). The solubility of most insoluble compounds such as ferric hydroxide tends to change as temperature changes, which can be predicted by the Second Law of Thermodynamics. The Second Law of Thermodynamics predicts that changing the temperature allows insoluble substances to shift towards a more disorientated state by increasing the average velocity of the particles in order to achieve thermal equilibrium of the system. The significance of the effect of molar ratio (B) (SE = 4.291) was probably owing to the change in the number of moles of iron(II) that underwent oxidative precipitation produced by the change in molar ratio, which manifested in the amount of precipitated iron(II). Increasing the molar ratio caused an increase in the available MnO₄-consumed for the oxidation of iron(II) to iron(III), which then precipitated to form ferric hydroxide. The significance of the

effect of pH (C) was also evident in Figure 10. Among the effects that were significant, it had the largest standardized effect (SE) of 105.60. A possible explanation for the significance in the effect of pH (C) is the dependence of iron precipitation on pH as in the case of ferric hydroxide. It can be seen in Equation 4-1 that an increase in concentration of the hydroxide ion during pH adjustment will result in the shift of the reaction forward to form ferric hydroxide following Le Chatelier's principle.

$$Fe_{(aq)}^{3+} + 3 OH_{(aq)}^{-} \leftrightarrow Fe(OH)_{3(s)}$$

$$(4-1)$$

The effects of the interaction between temperature and pH (AC) and molar ratio and pH (BC) were also identified as significant based on Figure 10. However, the significance of the interaction effects AC and BC had the smallest SE among the effects that were significant since the points are relatively close to the normal probability line. The SE of AC was -2.607 while the SE of BC was -3.400. This suggested that the interaction effects had a smaller influence in the amount of precipitated iron(II) than the individual effects.

The effect of interactions AC and BC was further evident when the interaction plots were constructed, as shown in Figures 11 and 12, respectively. An interaction plot shows the impact of the interacting variables to the response being investigated. The more parallel the lines in an interaction plot, the less the interaction between the two variables.



Figure 11. Interaction plot between temperature (A) and pH (C) with respect to the mean of precipitated iron(II) (%) where -1.0 and 1.0 represent the low level and high levels of the variables used in the experimental plan (Table 11).





In the case of either Figure 11 or Figure 12, the lines were relatively parallel to each other with only a slight increase in the solid line, which means that the interaction between temperature and pH (AC) and molar ratio and pH (BC) was weak and

therefore, their significance in the precipitation of iron(II) can be considered to be negligible. A possible reason for the interaction between temperature and pH (AC) is due to the influence of the temperature on the pH. A change in temperature alters the activity of all the ions in the solution. An increase in temperature will increase the activity of hydrogen ion and cause the pH to decrease. The decrease in pH will result in an increase in pOH and decrease the activity of hydroxide ion; therefore, it hinders iron precipitation (Equation 4-1). The interaction between molar ratio and pH (BC), indicated that the effect of pH (C) on the amount of iron(II) precipitated was influenced by the molar ratio (B). For iron(II) to precipitate as ferric hydroxide upon pH adjustment, all iron(II) in the partially neutralised PLS must first be oxidised to iron(III) with enough MnO_4^- .

No significant effects were shown in the normal probability plot for precipitated (%) nickel(II) (Figure 13) and cobalt(II) (Figure 14). Any losses in nickel(II) and cobalt(II) were probably due to the occurrence of co-precipitation, which usually occurs when the solution is multi-component. This can either be due to inclusion, occlusion, or surface adsorption in the precipitate formed (Harvey 2000). However, another mechanism may have occurred, which requires further studies to confirm the reason behind the losses in nickel(II) and cobalt(II).

Taking into consideration the magnitude of the SE among the significant main and interaction effects, it was determined that for a synthetic partially neutralised PLS, molar ratio (B) had the largest SE for the precipitation of manganese(II) while pH (C) had the largest SE for the precipitation of iron(II); therefore, OVAT tests concentrated on the optimisation of molar ratio (B) and pH (C). The results are discussed in Section 4.2.



Figure 13. Normal probability plot of the main and interaction effects for the precipitation of nickel(II) (%) from a synthetic partially neutralised PLS ($\alpha = 0.05$).



Figure 14. Normal probability plot of the main and interaction effects for the precipitation of cobalt(II) (%) from a synthetic partially neutralised PLS ($\alpha = 0.05$).

4.1.2 Synthetic Post-Partially Neutralised PLS as Feed Solution in Determining the Effects of Experimental Variables on the Oxidative Precipitation of Manganese(II)

The effects of the variables were estimated by constructing a normal probability plot for precipitated metal (%) of manganese(II) versus the standardized effect, as shown in Figure 15. In this plot (Figure 15), a significant effect is indicated by a square while an insignificant effect is indicated by a circle.



Figure 15. Normal probability plot of the main and interaction effects for the precipitation of manganese(II) (%) from a synthetic post-partially neutralised PLS ($\alpha = 0.05$).

Figure 15 shows that temperature (A), molar ratio (B) and pH (C) as well as interaction between temperature and molar ratio (AB) were significant. The significance of the effect of temperature (A) with SE of -3.898 is owing to the considerable difference in the change in solubility of the manganese compound (MnO₂) produced by the change in temperature, which manifested in the amount of precipitated manganese(II). As described in Section 4.1.1, the solubility of most insoluble compounds tends to change as temperature changes following the Second Law of Thermodynamics, which can result in a change in the amount of precipitated metal ions. A possible explanation for the significance of the effect of molar ratio (B) (SE = 20.132) is similar to that given in Section 4.1.1, where the change in molar ratio produced a change in the number of moles of manganese(II) that underwent oxidative precipitation, which manifested in the amount of precipitated manganese(II). Increasing the molar ratio resulted in an increase in available MnO_4^- that was consumed for the oxidative precipitation of manganese(II) (Equation 2-26) from the synthetic post-partially neutralised PLS and therefore resulting in an increase in amount of precipitated manganese(II). The effect of pH (C) (SE = 3.901) was also significant, as can be seen in Figure 15. This result may be explained by the influence of pH (C) on the oxidative precipitation of manganese(II), which can be seen in the potential–pH equilibrium (Pourbaix) diagram (Figure 5) for the manganese–water system. Although MnO_2 can readily form at highly acidic conditions, increasing the pH results in an increase in the oxidation rate of manganese(II) (Van Benschoten et al. 1992, Freitas et al. 2013) and may result in the formation of other insoluble manganese compounds, such as Mn_2O_3 , Mn_3O_4 , and $Mn(OH)_2$.

The effect of the interaction between temperature and molar ratio (AB) (SE = 3.478) was also identified to be significant, based on the data of Figure 15. This may indicate that the effect of molar ratio (B) on the precipitation of manganese(II) is influenced by temperature (A). It is possible that the interaction between temperature and molar ratio (AB) is due to the change in the number of moles of manganese(II) that underwent oxidative precipitation produced by the change in molar ratio and since temperature (A) affects the solubility of the insoluble manganese compound (MnO₂), as in the case of the individual effect, its interaction with molar ratio (B) manifested through considerable changes in the amount of precipitated manganese(II).

The effect of interaction AB was further evident when the interaction plot was constructed, as shown in Figure 16. It can be seen in Figure 16 that changes in temperature only occurred in instances when the molar ratio was at the low level. At a high level of molar ratio, the mean of precipitated manganese(II) was relatively constant as the temperature changed, which may be beneficial since the amount of precipitated manganese(II) was at a maximum. This suggested that the individual effect of molar ratio is stronger compared with the interaction effect between temperature and molar ratio at the high-level molar ratio. By operating at high-level

molar ratio, the interaction effect will be minimised while the amount of precipitated manganese(II) is maximised.



Figure 16. Interaction plot between temperature (A) and molar ratio (B) with respect to the mean of precipitated manganese(II) (%) where –1.0 and 1.0represent the low level and high levels of the variables used in the experimental plan (Table 12).

In the case of the precipitation of nickel(II), Figure 17 shows that the effects of molar ratio (B) and agitation speed (D) were significant. The significance of the effect of molar ratio (B) (SE = 3.982) suggests that nickel(II) may have also undergone precipitation as the molar ratio (B) changed, which manifested in the amount of precipitated nickel(II). However, this finding was unexpected since, based on the potential–pH equilibrium (Pourbaix) diagram (Figure 6) for the nickel–water system, nickel should remain soluble under the conditions used for the present study. A possible reason for this is that the solution behaviour being multi-component contains different interacting ions that may have caused nickel(II) to co-precipitate due to the changes in molar ratio and therefore causing a considerable difference in the amount of precipitated nickel(II). As for the significance of the effect of agitation speed (D) (SE = 2.776), it is possible that the change in agitation speed produced a change in the collision frequency between the different ions in the solution, which affected the precipitation mechanism (Misra 2016), resulting in a change in the

amount of precipitated nickel(II). Misra (2016) reported that agitation affected the nucleation, growth, and agglomeration of solids formed during precipitation and may also lead to attrition and breakage of the solids when the agitation speed is high.



Figure 17. Normal probability plot of the main and interaction effects for the precipitation of nickel(II) (%) from a synthetic post-partially neutralised PLS ($\alpha = 0.05$).

Figure 18 shows that the effects of temperature (A), molar ratio (B), and pH (C) as well as the interactions effects between temperature and molar ratio (AB), temperature and pH (AC), molar ratio and agitation speed (BD), and pH and agitation (CD) were significant for the precipitation of cobalt(II). The significance of the effect of temperature (A) with SE of 6.215 was possibly due to the considerable difference in the change in solubility of the cobalt compound produced by the change in temperature, which manifested in the amount of precipitated cobalt(II). However, it is not clear what cobalt compound formed and may require further investigation to be undertaken to examine the precipitate since this is beyond the scope of the present work. Similarly, the reason behind the significance of the effect of molar ratio (B) with SE of 42.193 is also not clear at this point. Based on the potential–pH equilibrium (Pourbaix) diagram (Figure 7) for the cobalt–water system, cobalt should remain soluble under the conditions used for the present study. However, the

significance of molar ratio (B) suggests that cobalt(II) may have also undergone precipitation as the molar ratio (B) changed. A possible explanation for this may be similar to that for nickel(II), where the multi-component behaviour of the solution and the interaction between the different ions may have driven the co-precipitation of cobalt(II) to occur during the oxidative precipitation of manganese(II) due to the changes in molar ratio. Further studies may need to be undertaken to understand the mechanism by which molar ratio affects the precipitation of cobalt(II) during the oxidative precipitation of cobalt(II) during the oxidative precipitation of cobalt(II) during the oxidative precipitation of manganese(II) by potassium permanganate. In the case of the effect of pH (C) (SE = 41.098), its significance was probably owing to the dependence of different cobalt compounds on pH, as what can be seen from the potential–pH equilibrium (Pourbaix) diagram (Figure 7) for the cobalt–water system. Increases in pH can enhance the precipitation of cobalt(II), which may result in the formation of insoluble cobalt compounds such as Co(OH)₃, Co₃O₄, and Co(OH)₂.



Figure 18. Normal probability plot of the main and interaction effects for the precipitation of cobalt(II) (%) from a synthetic post-partially neutralised PLS ($\alpha = 0.05$).

The effects of the interaction between temperature and molar ratio (AB) (SE = 4.127), temperature and pH (AC) (SE = -4.477), molar ratio and agitation speed (BD) (SE =

-2.675), and pH and agitation (CD) (SE = -2.761) were also identified as significant, based on Figure 18. However, interaction effects AB, AC, BD, and CD had the smallest SE among the effects that were significant since the points are relatively close to the normal probability line. This suggested that the interaction effects would have a smaller influence on the precipitation of cobalt(II) than the individual effects.

The effects of interactions AB, AC, BD, and CD were further evident when the interactions plots were constructed as shown in Figures 19, 20, 21, and 22, respectively.





The significance of the effect of interaction AB may indicate that the effect of molar ratio (B) to the precipitation of cobalt(II) is influenced by temperature (A). The reason for this is similar to that of manganese(II). The interaction between temperature and molar ratio (AB) is owing to the change in the number of moles of cobalt(II) that underwent precipitation produced by the change in molar ratio. Since temperature (A) affects solubility, its interaction with molar ratio (B) manifested through considerable changes in the amount of precipitated cobalt(II). Figure 19 shows that

at the high level of molar ratio, there was an increase in the amount of mean precipitated cobalt(II) as temperature changed from low to high level. It can also be seen that at the low level of molar ratio, the mean of precipitated cobalt(II) was relatively constant. Since the interaction was only evident at a high-level molar ratio, this is probably the reason why the SE for the effect of interaction AB was small and may indicate that there is a weak interaction between temperature and molar ratio and therefore, its significance in the precipitation of cobalt(II) can be considered to be negligible.

Turning now to the significance of the effect of AC, Figure 20 indicates that the effect of pH (C) on the precipitation of cobalt(II) is influenced by temperature (A). The significance of effect AC can be explained by the fact that temperature has an influence on pH. A change in temperature will alter the activity of all the ions in the solution. An increase in temperature will increase the activity of hydrogen ions and cause the pH to decrease; therefore, it may cause some changes in the precipitation of cobalt(II).



Figure 20. Interaction plot between temperature (A) and pH (C) with respect to the mean of precipitated cobalt(II) (%) where -1.0 and 1.0 represents the low level and high levels of the variables used in the experimental plan (Table 12).

Figure 20 shows that at the low level of pH, there was an increase in the amount of mean precipitated cobalt(II) as temperature changed from low to high level. The

amount of mean precipitated cobalt(II), however, remained relatively constant when the pH was at the high level as temperature changed from low to high level, which is probably the reason why the SE for the effect of interaction AC on the precipitation of cobalt(II) was small and may indicate a weak interaction between temperature and pH and therefore, its significance in the precipitation of cobalt(II) can also be considered to be negligible.

The lines in either Figure 21 or Figure 22, respectively, were relatively parallel to each other with only a slight decrease in the solid line at the low level of either molar ratio for interaction effect BD or pH for interaction effect CD. This indicated that the interaction between molar ratio and agitation speed (BD) and pH and agitation speed (CD) was weak and therefore, their significance in the precipitation of cobalt(II) can be considered to be negligible. It was also evident from Figures 21 and 22 that the agitation speed did not cause much change in the mean of precipitated cobalt(II). The change in the mean of precipitated cobalt(II) was primarily influenced by either molar ratio or pH, which suggested that the individual effects were strong compared with the interaction effects. The interaction of agitation speed to either molar ratio or pH may possibly due to a similar reason as discussed in the earlier portion of Section 4.1.2 concerning nickel(II).



Figure 21. Interaction plot between molar ratio (B) and agitation speed (D) with respect to the mean of precipitated cobalt(II) (%) where -1.0 and 1.0 represents the low level and high levels of the variables used in the experimental plan (Table 12).



Figure 22. Interaction plot between pH (C) and agitation speed (D) with respect to the mean of precipitated cobalt(II) (%) where –1.0 and 1.0 represents the low level and high levels of the variables used in the experimental plan (Table 12).

Taking into consideration the magnitude of the SE among the significant main and interaction effects on the oxidative precipitation of manganese(II), it was determined

that for a synthetic post-partially neutralised PLS, molar ratio (B) had the largest SE for the precipitation of manganese(II); therefore, OVAT tests concentrated on the optimisation of molar ratio (B). The results are discussed in Section 4.2. Significant effects for the precipitation of either nickel(II) and cobalt(II) were not considered for the OVAT tests since the aim of the current study was focused on optimising the variables that had significant effect on the oxidative precipitation of manganese(II).

4.2 Optimisation of Variables with Significant Effect on Oxidative Precipitation of Manganese(II)

Optimisation experiments using synthetic partially neutralised and post-partially neutralised PLS were carried out using the same experimental procedure outlined in Section 3.6. As determined from the screening experiments (Section 4.1.1) using partially neutralised PLS as test solution, molar ratio (B) and pH (C) were considered for optimisation using the OVAT approach since both variables had the largest standardized effect in comparison with all the other variables with significant effect. In the case of the post-partially neutralised PLS, it was determined from the screening experiments (Section 4.1.2) that molar ratio (B) had the largest standardized effect in comparison with significant effect on the precipitation of manganese(II) and it was considered for optimisation using the OVAT approach. The variations of molar ratio (B) and pH (C) were investigated in separate sets of experiments to explore whether higher amounts of manganese(II) could be precipitated. Optimisation of pH (C) was only applicable to synthetic partially neutralised PLS due to the presence of iron(II), which required pH adjustment for its oxidative precipitation.

4.2.1 Effect of pH on Oxidative Precipitation of Manganese(II)

The effect of pH on the oxidative precipitation of manganese(II) was examined by varying the pH in the range of 1.77 to 4.0. Afterwards, the amount of precipitated metal ions (%) was plotted against pH to observe any trend. Similar to Section 4.1, the percent of precipitated metal ions were calculated from the initial minus the final metal ion concentration after the dilution effect had been factored in due to the

addition of the reagents. Figure 23 shows the effect of pH on the amount of precipitated metals ions (%).



Figure 23. The effect of pH on the amount of precipitated metals ions (%) from synthetic partially neutralised PLS. Initial pH = 2.12–2.15; ambient T = 23.9–26.0 °C; agitation speed = 50 rpm; molar ratio = 0.5 (stoichiometric); t = 1 h.

From the data in Figure 23, there was an increasing trend in the amount of precipitated manganese(II), iron(II), nickel(II), and cobalt(II) (%) as the pH increased. The increase in the amount of precipitated manganese(II) was observed when the pH was greater than or equal to 3 ($pH \ge 3$). This suggested that operating at a higher pH will enhance the oxidative precipitation of manganese(II) that will result in an increase in the amount of precipitated manganese(II). A possible explanation for this increase is the tendency for manganese(II) to form other stable solid species such as Mn_2O_3 , Mn_3O_4 , and $Mn(OH)_2$ when operating at higher pH conditions, as can be seen in the potential-pH equilibrium (Pourbaix) diagram for the manganese-water system (Figure 5). Additionally, the result is consistent with the trend reported in earlier studies on the oxidative precipitation of manganese(II) for water treatment (Adams 1960, Phatai et al. 2010, Elsheikh et al. 2017) and acid mine water treatment (Freitas et al. 2013) despite using different manganese-containing solutions and operating at different pH conditions (neutral to basic conditions). A similar trend was also observed with the amount of precipitated iron(II) (%). The trend can be attributed to the fact that the precipitation of ferric hydroxide would require the adjustment of pH to greater than 2 (pH > 2). Operating at a higher pH (pH > 2), however, was unfavourable to nickel(II) and cobalt(II) due to the increase in their precipitation (Figure 23). It is possible that the increase was due to the co-precipitation (Harvey 2000) of nickel(II) and cobalt(II) either as nickel hydroxide or cobalt hydroxide as the pH increases or a different mechanism that requires further investigation in order to confirm this observation.

An issue that may arise from the increase in the amount of precipitated nickel(II) and cobalt(II) was that there will be losses of the valuable metals that goes together with the manganese precipitate (MnO₂). Given these considerations, the most appropriate approach is to minimise the losses of nickel(II) and cobalt(II) by operating at the minimum pH needed to maximise both iron(II) and manganese(II) precipitation, which was at pH 3. At a pH of 3, iron(II) precipitation was complete while manganese(II) precipitation was close to 85%. Moreover, at this pH, the amounts of nickel(II) and cobalt(II) precipitated were close to 2% and 7%, respectively. However, if the desire is to push the amount of precipitated manganese(II) beyond 85% by operating at pH greater than 3 (pH > 3), it would be at the expense of losing nickel(II) and cobalt(II), which reached as high as 28% and 43%, respectively.

Taken together, these results suggest that the oxidative precipitation of manganese(II) can be carried out in acidic conditions, but can be enhanced at a pH > 3, which is beneficial for the oxidative precipitation of iron(II) when present together with manganese(II) in the partially neutralised PLS. However, inevitable losses in nickel(II) and cobalt(II) was observed at a pH >3. Hence, the pH value applied on the oxidative precipitation of manganese(II) from an actual partially neutralised PLS was 3 to ensure the precipitation of the oxidized iron(II) before manganese(II). The results of the experiments are discussed in Section 4.3. As for nickel(II) and cobalt(II), further study needs to be conducted in order to understand the mechanism by which the ions are affected by pH control during the oxidative precipitation of manganese(II) in a partially neutralised PLS.

4.2.2 Effect of Molar Ratio on Oxidative Precipitation of Manganese(II)

The molar ratios used in the experimental work were at 85% (0.43), 100% (0.50), 105% (0.53), and 115% (0.58) of the stoichiometric ratio (0.50) in order to observe

the effect of molar ratio on the oxidative precipitation of manganese(II). Afterwards, the amount of precipitated metal ions (%) was plotted against molar ratio to observe any trend. The percent of precipitated metal ions were calculated from the initial minus the final metal ion concentration after the dilution effect had been factored in due to the addition of the reagents. Figure 24 shows the effect of molar ratio on the amount of precipitated metal ions (%).





From the data in Figure 24, the amount of precipitated iron(II) (%) was relatively complete and constant at the molar ratios (0.43, 0.50, 0.53, and 0.58) investigated, indicating that there was sufficient available permanganate ion (MnO_4^-) to ensure the complete oxidative precipitation of the iron(II) in the solution. This result may be attributed to the fact that any iron(II) present in the partially neutralised PLS will preferentially be oxidised by the available MnO_4^- before manganese(II) due to a much lower standard reduction potential (E^{Θ}) (Section 3.4). In contrast, the result was not the same for the amount of precipitated manganese(II). Figure 24 shows that the amount of precipitated manganese(II) increased with increasing molar ratio. A possible explanation for this result is that by increasing the molar ratio, there was an increase in the available MnO_4^- that reacted not only with iron(II), but also with manganese(II); therefore, resulting in more manganese(II) is consistent with

previous studies on the oxidative precipitation of manganese(II) reported by Phatai et al. (2014) and Elsheikh et al. (2017) using synthetic groundwater and Heviankova and Bestova (2007) and Macingova et al. (2016) using acid mine water despite using different manganese-containing solutions and operating at different pH conditions (neutral to basic conditions) in comparison with the present study. However, it should be noted down that although the trend of the present study is consistent with those reported from previous studies, the amount of potassium permanganate consumed for the oxidative precipitation of manganese(II) differed from that of the various authors (Heviankova and Bestova 2007, Phatai et al. 2014, Macingova et al. 2016, Elsheikh et al. 2017) mentioned. The difference in the amount of potassium permanganate used by various authors is probably owing to the dissimilar composition of the test solution used, which may have resulted in different solution behaviour and interaction among the metal ions in the solution that caused some interference in the oxidative precipitation by KMnO₄.

It is also apparent from Figure 24 that there was an increase in the amount of precipitated nickel(II) and cobalt(II) as the molar ratio increased to greater than 0.50. Operating at molar ratios greater than 0.50 showed complete oxidative precipitation of manganese(II), but the downside of this is that nickel(II) and cobalt(II) were lost in the manganese precipitate due to the observed increase in amount of precipitated nickel(II) and cobalt(II). Although it was observed from the potential-pH equilibrium (Pourbaix) diagram (Figure 6 and Figure 7) that it is unlikely for nickel(II) and cobalt(II) to undergo oxidative precipitation at the conditions used for the present study, it is possible that the increase is due to the co-precipitation (Harvey 2000) of nickel(II) and cobalt(II). The co-precipitation mechanism whether by inclusion, occlusion, or surface adsorption (Harvey 2000), however, is not clear at this point and may require further investigation to understand the mechanism in which nickel(II) and cobalt(II) coprecipitates during the oxidative precipitation of manganese(II). Additionally, a different mechanism may have also occurred other than co-precipitation that requires further investigation in order to confirm this observation. Given these considerations, the most probable approach to minimise the losses in nickel(II) and cobalt(II) is to operate at stoichiometric molar ratio (0.50) while maximising the
precipitation of iron(II) and manganese(II). At stoichiometric molar ratio, iron(II) precipitation was complete while manganese(II) was close to 85%. Moreover, the amounts of precipitated nickel(II) and cobalt(II) were close to 2% and 7%, respectively. However, if the desire is to obtain higher amounts of precipitated iron(II) and manganese(II), using 5% more of stoichiometric molar ratio (0.53) resulted in the complete precipitation of both iron(II) and manganese(II) at the expense of higher nickel(II) and cobalt(II) losses. At a molar ratio of 0.53, the amounts of precipitated nickel(II) and cobalt(II) were close to 20% and 30%, respectively.

Turning now to the effect of molar ratio on the oxidative precipitation of manganese(II) from synthetic post-partially neutralised, the molar ratios used in the experimental work were at 85% (0.57), 100% (0.67), 105% (0.70), and 115% (0.77) of the stoichiometric ratio (0.67). The amount of precipitated metal ions (%) was plotted against molar ratio to observe any trend. Figure 25 shows that there was an increase in the amount of precipitated manganese(II) (%) from a molar ratio of 0.57 to 0.70, but a slight decrease was observed at 0.77.



Figure 25. The effect of molar ratio on the amount of precipitated metal ions (%) from synthetic post-partially neutralised PLS. Unadjusted pH = 1.66–1.71; ambient T = 18.5–23.0 °C; agitation speed = 50 rpm; t = 1 h.

A possible explanation for the increase in the amount of precipitated manganese(II) from 0.57 to 0.70 is due to the increase in the available permanganate ion (MnO_4^-) and therefore, resulting in more manganese(II) being oxidised and then precipitated

from the solution. This finding was in agreement with the results obtained using partially neutralised PLS, where an increase in molar ratio also resulted in an increase in amount of precipitated manganese(II). However, it should be noted down that the presence of iron(II) in the partially neutralised PLS as opposed to that of a postpartially neutralised PLS would require more permanganate ion to oxidise iron(II) before manganese(II). The increase in the amount of precipitated manganese(II) as the molar ratio increased is also consistent with previous studies on the oxidative precipitation of manganese(II) reported by Phatai et al. (2014) and Elsheikh et al. (2017) using synthetic groundwater and Heviankova and Bestova (2007) and Macingova et al. (2016) using acid mine water despite using different manganesecontaining solutions and operating at different pH conditions (neutral to basic conditions) in comparison with the present study. Turning now to the slight decrease in the amount of precipitated manganese(II) observed at a molar ratio of 0.77, there was probably an overdose of permanganate ion in the solution, which may have generated additional manganese(II) according to Equation 4-1. This was not observed from the results obtained from the experimental work using partially neutralised PLS, which indicated that the presence of iron(II) possibly caused some interference during the oxidative precipitation of manganese(II) that prevents the generation of additional manganese(II) when there is an overdose of permanganate ion in the solution.

$$MnO_{4(aq)}^{-} + 8 H_{(aq)}^{+} + 5 e^{-} \leftrightarrow Mn_{(aq)}^{2+} + 4 H_2O$$
(4-1)

In the case of either the amount of precipitated nickel(II) or cobalt(II), Figure 25 shows a relatively similar trend except that the percent of precipitated cobalt(II) was at a higher range in comparison with nickel(II). This result was contrary to the results obtained from nickel(II) and cobalt(II) in Figure 24 especially with the amount of precipitated cobalt(II), which was already high even at low molar ratio. The reason for this is not clear at this point but it may have something to do with the multicomponent behaviour of the solution, which have caused the variability in the amount of precipitated nickel(II) and cobalt(II) due to a similar reason as discussed in the earlier portion of Section 4.2.2 for Figure 24. Given these considerations, the most probable approach is to operate at a molar ratio where the generation of additional manganese(II) from permanganate will be avoided. In the case of the present study, this would be less than or equal to a molar ratio of 0.70.

Overall, the results for both partially neutralised and post-partially neutralised PLS indicated that an increase in molar ratio resulted in an increase in the amount of precipitated manganese(II). However, special care should be taken into consideration for choosing the amount of potassium permanganate to ensure that the ratio of available MnO₄⁻ will be sufficient for the oxidative precipitation of manganese(II) from a multi-component solution and to avoid the overdose of permanganate ion, which can lead to the generation of additional manganese(II). For a partially neutralised PLS, the amount of precipitated manganese(II) increased at a molar ratio greater than the stoichiometric molar ratio (>0.50) before reaching a relatively constant value close to 100%. As for a post-partially neutralised PLS, the optimum value obtained was at a stoichiometric ratio less than or equal to 105% of the stoichiometric molar ratio (≤ 0.70) where the amount of precipitated manganese(II) was close to 100%. However, in the case of either the partially neutralised or postpartially neutralised PLS, inevitable losses in nickel(II) and cobalt(II) were observed in the results. Hence, the molar ratio applied on the oxidative precipitation of manganese(II) from either an actual partially neutralised or post-partially neutralised was limited to stoichiometric molar ratio and 105% of stoichiometric molar ratio. The results obtained are discussed in Section 4.3.

4.3 Application of Optimised Values on Oxidative Precipitation of Manganese(II) from Actual PLS

The oxidative precipitation experiments using actual partially neutralised and postpartially neutralised PLS were carried out using the same experimental procedure outlined in Section 3.6. As determined from the optimisation experiments using synthetic partially neutralised PLS, the optimum value obtained for pH was 3 (Section 4.2.1) while for molar ratio, it was at stoichiometric molar ratio (0.50) (Section 4.2.2). However, since the amount of precipitated manganese(II) at 0.50 was only 85%, experiments using 105% of the stoichiometric molar ratio (0.53) were also carried out even at the expense of losses in nickel(II) and cobalt(II) to test whether a higher amount of manganese(II) would precipitate from an actual partially neutralised PLS. Application of the technique using molar ratios of 0.50 and 0.53, both maintained at pH 3, were investigated in separate sets of experiments. In the case of the post-partially neutralised PLS, it was determined from the optimisation experiments that the optimum value for molar ratio should be less than or equal to 0.70 (Section 4.5.1) to avoid the overdose of permanganate ion, which can lead to the generation of additional manganese(II). Hence, the values chosen were stoichiometric molar ratio (0.67) and 105% of the stoichiometric molar ratio (0.70) since it was at these values that the amount of precipitated manganese(II) was relatively constant based on the OVAT experiment. The application of the technique using molar ratios of 0.67 and 0.70 was also investigated in separate sets of experiments. In either the actual partially neutralised or post-partially neutralised PLS, each set of experiments was conducted twice to test the repeatability of the results when applied to the actual PLS.

Repeatability of the results was examined by constructing a bar chart of the mean of precipitated metal ions (%) against the metal ions in the actual PLS. Error bars for each metal ion were also included to represent the standard deviation (SD) of the mean of precipitated metal ions (%). Data that are narrowly clustered around the mean will have a small SD while data that are broadly dispersed around the mean will have a large SD. In the case of the present study, a SD of less than or equal to 10% (SD \leq 10%) was considered to be practically acceptable for the application of the technique using an actual PLS. The acceptability of the SDs, however, may vary depending on the willingness of the nickel laterite processing operations to tolerate any variation in the results on the basis of their desired final product as well as the variability in the composition of the actual PLS they use. Moreover, the number of results taken for the repeatability test was based on the practicality of the procedure and the availability of an appropriate measurement system at the time the present study was conducted.

4.3.1 Actual Partially Neutralised PLS as Feed Solution for the Application of Optimised Values on Oxidative Precipitation of Manganese(II)

Table 13 shows a summary of the standard deviation (SD) of the mean of precipitated metal ions (%) obtained from the application of oxidative precipitation of manganese(II) using actual partially neutralised PLS. It is apparent from Table 13 that the SD for all of the metal ions was less than 10% in either a molar ratio of 0.50 or 0.53. This means that given the tolerance used for the variance in the results of the present study was less than or equal to 10% (SD \leq 10%), the results obtained from the application of the technique using actual partially neutralised were repeatable for every metal ion.

| | Molar Ratio | |
|------------|-----------------------------|------|
| Metal lons | 0.50 | 0.53 |
| | Standard Deviation (SD) (%) | |
| Al | 2.59 | 3.62 |
| Са | 0.78 | 2.29 |
| Со | 3.07 | 7.32 |
| Cr | 4.11 | 0.04 |
| Cu | 9.83 | 6.91 |
| Fe | 0.31 | 0.19 |
| Mg | 1.23 | 3.38 |
| Mn | 0.20 | 1.77 |
| Ni | 1.75 | 2.29 |
| Zn | 0.73 | 2.70 |

Table 13. Standard deviation (SD) of the mean of precipitated metal ions (%) inthe actual partially neutralised PLS.

The bar chart (Figure 26) shows the mean of precipitated metal ions (%) against the different metal ions present in the actual partially neutralised PLS. The trend for the mean of precipitated metal ions (%) for the metal ions found in actual partially neutralised PLS was relatively comparable for either a molar ratio of 0.50 or 0.53.



Figure 26. Mean of precipitated metal ions (%) from an actual partially neutralised PLS. Conditions: pH = 3.00; ambient T = 23.3–26.9 °C; agitation speed = 50 rpm; molar ratio = 0.50 (stoichiometric) and 0.53 (105% of stoichiometric); t = 1 h.

This finding suggested that nearly the same amounts of metal ions were precipitated from the actual partially neutralised PLS whether the molar ratio is either 0.50 or 0.53. By using a molar ratio of 0.50, however, it will be more economical for the application of the technique since lesser amount of potassium permanganate will be consumed to achieve a comparable result when a molar ratio of 0.53 was used. Further analysis of Figure 26 showed that manganese(II) and iron(II) had the highest mean of precipitated metal ions in comparison with all the other metal ions in the actual partially neutralised PLS. The result is consistent with the trend observed from the OVAT approach using synthetic partially neutralised PLS where manganese(II) and iron(II) also attained the highest amount precipitated (Section 4.2.1 and Section 4.2.2), suggesting that the optimum values used for the application of the technique were suitable for the oxidative precipitation of manganese(II) and iron(II) from the actual partially neutralised PLS. It can be seen from Figure 26, however, that there were certain losses in nickel(II) and cobalt(II), which were approximately 27% and 50%, respectively. This reinforced the finding in Section 4.2.1 and Section 4.2.2 that losses in nickel(II) and cobalt(II) were inevitable during the oxidative precipitation of manganese(II) by KMnO₄. The results in Figure 26, however, were not compared with standard precipitation curves as a function of pH for the reason it was only limited to metal hydroxides, sulfides, arsenates, and phosphates and does not include the possible compounds that can form during the oxidative precipitation by potassium permanganate. Characterisation of the precipitate still needs to be carried out in future studies before the results can be compared with standard precipitation curves.

4.3.2 Actual Post-Partially Neutralised PLS as Feed Solution for the Application of Optimised Values on Oxidative Precipitation of Manganese(II)

Table 14 shows the summary of the standard deviation (SD) of the mean of precipitated metal ions (%) obtained from the application of oxidative precipitation of manganese(II) using actual post-partially neutralised PLS. It is apparent from Table 14 that majority of the metal ions had a SD of less than 10% in either a molar ratio of 0.67 or 0.70. Table 14, however, shows that the SD of chromium(III) slightly exceeded 10% at a molar ratio of 0.67 while copper(II) had a large SD at both a molar ratio of 0.67 and 0.70. A possible reason for this may be due to the very low concentrations of chromium(III) (0.004 g/L) and copper(II) (0.01 g/L) in the actual post-partially neutralised PLS where co-precipitation occurred, which resulted in variable amounts of precipitated chromium(III) and copper(II).

| | Molar Ratio | |
|------------|-----------------------------|-------|
| Metal lons | 0.67 | 0.70 |
| | Standard Deviation (SD) (%) | |
| Al | 2.56 | 2.46 |
| Са | 5.23 | 3.50 |
| Со | 7.41 | 0.77 |
| Cr | 10.13 | 2.33 |
| Cu | 26.08 | 45.82 |
| Fe | 4.87 | 0.20 |
| Mg | 2.79 | 1.67 |
| Mn | 2.28 | 0.00 |
| Ni | 4.69 | 0.43 |
| Zn | 9.02 | 2.92 |

Table 14. Standard deviation (SD) of the mean of precipitated metal ions (%) inthe actual post-partially neutralised PLS

The bar chart (Figure 27) shows the mean of precipitated metal ions (%) against the different metal ions present in the actual post-partially neutralised PLS. The trend for the mean of precipitated metal ions (%) for the metal ions found in the actual post-partially neutralised PLS was relatively similar for either a molar ratio of 0.67 or 0.70.





However, although the trend was similar for both of the molar ratios, there was a slight increase in the mean of precipitated metal ions (%) in most of the metal ions when a molar ratio of 0.70 was used. It is possible that the increase in the available permanganate ion (MnO_4^-) did not just result in the oxidative precipitation of manganese(II), but also the co-precipitation of other metal ions in the actual post-partially neutralised PLS. In addition, a comparison between the results from both the actual partially neutralised and post-partially neutralised PLS showed a dissimilar trend. A possible reason for this is due to the presence of considerable amount of iron(II) (0.28 g/L) in the actual post-partially neutralised PLS. The considerable amount of iron(II) in the actual partially neutralised PLS.

the results of the optimisation experiments (Section 4.2). In the case of nickel(II) and cobalt(II), it can be seen from Figure 27 that there were also certain losses in nickel(II) and cobalt(II), which was similar to the observation in Section 4.3.1 for actual partially neutralised PLS. This suggested that regardless of whether the actual PLS was partially neutralised or post-partially neutralised, losses in nickel(II) and cobalt(II) would be inevitable during the oxidative precipitation of manganese(II) by KMnO₄. With a similar reason as Figure 26, the results shown in Figure 27 were not compared with standard precipitation curves.

Despite the results showing slight increase in the mean of precipitated metal ions (%) for most of the metal species, using a molar ratio of 0.67 would still be more economical for the application of the technique since lesser amount of potassium permanganate will be consumed at only a marginal difference in the mean of precipitated manganese(II) between a molar ratio of 0.67 and 0.70. In addition, lower amounts of nickel(II) and cobalt(II) were lost in the precipitate when a molar ratio of 0.67 was used, which were approximately 3% and 43%, respectively.

Chapter 5

CONCLUSIONS AND RECOMMENDATIONS

The growing demand for nickel since the 1950s and the dwindling nickel sulfide ore resources have resulted in increasing interest in processing nickel laterite ores as an alternative source. Nickel laterite ores are processed via a hydrometallurgical route if the aim is to produce high purity nickel. There are three available leaching processes under the hydrometallurgical route, and these are the Caron process, pressure acid leaching (PAL), and atmospheric leaching (AL). Among the available three leaching processes, the most widely used for nickel laterite ores is PAL since it can yield high nickel and cobalt dissolutions (>95%). However, substantial co-dissolutions of other components of the ore occurs during PAL that results in a highly contaminated pregnant leach solution (PLS).

The PAL-generated PLS is partially neutralised to largely remove the remaining iron, aluminium, and chromium in the solution. Afterwards, the resulting solution will undergo further purification since large amounts of manganese, which can affect the efficiency of the downstream processing, still remain as the major impurity in the PAL-generated PLS even after the solution has been partially neutralised. Current purification techniques for PAL-generated PLS are carried out through either intermediate precipitation (MHP or MSP) or direct solvent extraction (DSX). These purification techniques, however, have proven to be unsuccessful in separating manganese from nickel and cobalt due to either co-precipitation as a hydroxide in intermediate precipitation or co-extraction in DSX where it can potentially oxidise during the electrowinning step and may degrade the organic when the spent electrolyte is recycled back to the solvent extraction. Moreover, the intermediate products (mixed hydroxides or mixed sulfides) coming from intermediate precipitation will require further processing using consecutive re-leaching and solvent extraction steps before nickel and cobalt can be recovered as final saleable products and therefore increasing the operational costs associated with processing nickel laterite ores.

The present study attempted to address the issue concerning manganese(II) by exploring the removal of manganese(II) from the PLS generated from PAL of nickel laterite ores by oxidative precipitation with potassium permanganate. As postulated by D.C. Ibana in a personal communication (Section 1.5), the removal of manganese(II) from PAL-generated PLS can be achieved through oxidative precipitation by potassium permanganate without introducing any new ions to the highly contaminated PLS. This study set out to determine the variables that significantly affect the oxidative precipitation of manganese(II) by potassium permanganate, optimise the determined variables by applying the technique using synthetic partially and post-partially neutralised PLS, and then apply the optimised variables using actual partially and post-partially neutralised PLS.

The literature review revealed that most studies on the oxidative precipitation of manganese(II) by potassium permanganate were for treatment of water and acid mine water, which was usually carried out at basic pH conditions. There is, however, a firm thermodynamic basis to test the application of oxidative precipitation of manganese(II) from PAL-generated PLS in highly acidic conditions, which will produce manganese dioxide (MnO₂), based on the potential-pH equilibrium (Pourbaix) diagram of manganese-water system (Figure 5). Aside from the influence of pH, another variable found in the literature that has an influence on the oxidative precipitation of manganese(II) is the amount of potassium permanganate. The amount of potassium permanganate affects the ratio between moles of permanganate ion and total moles of iron(II) and manganese(II) (denoted as mol MnO4⁻: total mol Fe(II) and Mn(II)) for partially neutralised PLS and between moles of permanganate ion and moles of manganese(II) (denoted as mol MnO4⁻:mol Mn(II)) for post-partially neutralised PLS. The amount must carefully be determined to ensure that sufficient MnO_4^- is available for the complete oxidative precipitation of manganese(II) from PAL-generated PLS and to avoid the generation of manganese(II) due to the reduction of MnO4⁻ when there is an overdose of potassium permanganate. Given these considerations, an experimental program involving bench-scale experimental work was developed and implemented to explore and evaluate the viability of the proposed oxidative precipitation technique.

The main findings of the research work may be summarised as follows:

- Screening experiments using unreplicated full factorial design revealed that molar ratio (B) significantly affected the oxidative precipitation of manganese(II) from either the partially neutralised or post-partially neutralised PLS. This was evident in the normal probability plot of the main and interaction effects for precipitated manganese(II) (%) since molar ratio had the largest standardized effect (SE) compared with all the other variables investigated. Additionally, pH was also considered for the oxidative precipitation of manganese(II) from the partially neutralised PLS due to the presence of iron(II) in the solution, which would oxidise before manganese(II). The screening experiments revealed that pH (C) significantly affected the oxidative precipitation of iron(II), as evident in the normal probability plot of the main and interaction effects for precipitated iron(II) (%).
- Optimisation experiments using OVAT for the effect of pH revealed that the oxidative precipitation of manganese(II) can be carried out in acidic conditions, which was at a pH less than or equal to 4.0 (pH ≤ 4). It was found that a pH of 3 was optimum since appreciable amounts of nickel(II) and cobalt(II) were lost in the precipitate at pH > 3. This was evident in the plot of amount of precipitated metal ions (%) versus pH. In the case of the optimisation of the effect of molar ratio, the amount of manganese(II) precipitation increases with an increase in molar ratio and it was evident with the plot of amount of precipitated metal ions (%) versus molar ratio.
- Results from optimisation experiments using partially neutralised PLS revealed that a pH of 3 and a stoichiometric molar ratio of 0.50 were found to be optimum, where the amount of precipitated manganese(II) was 85% with minimal losses for nickel(II) and cobalt(II) at 2% and 7%, respectively.
- Results from optimisation experiments using post-partially neutralised PLS revealed that when pH was maintained between pH 1.66 and pH 1.71 and molar ratio was less than or equal to 0.70, the amount of precipitated

manganese(II) was nearly 100% with losses of nickel(II) between 3% and 10% while for cobalt(II), losses were between 50% and 60%.

Application of the optimised pH and molar ratio for the oxidative precipitation of manganese(II) from actual partially neutralised and post-partially neutralised PLS resulted in nearly 100% of precipitated manganese(II). Nickel(II) losses were approximately 27% while for cobalt(II), they were approximately 50% for the actual partially neutralised PLS. As for the actual post-partially neutralised, nickel(II) losses were approximately between 3% and 10% while for cobalt(II), they were approximately between 43% and 54%. This was evident with the bar chart for the mean of precipitated metal ions (%) versus metal ions in the actual PLS.

The findings of this work have contributed significantly to understanding the oxidative precipitation of manganese(II) from PAL-generated PLS using potassium permanganate. However, opportunities for further research that were outside the scope of the present study might explore the following:

- Investigate the mechanism by which nickel(II) and cobalt(II) partake on the oxidative precipitation of manganese(II) by potassium permanganate from either a partially neutralised or post-partially neutralised PLS by conducting kinetic and chemical speciation experiments.
- Characterise the precipitate by analysing the filtrate products from either the partially neutralised or post-partially neutralised PLS by conducting chemical composition, morphology, and minerology analysis.

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