The recovery of platinum group metals from low-grade concentrates to an iron alloy using silicon carbide as reductant

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Introduction

South Africa is the world’s leading supplier of platinum group metals (PGMs). Most of the PGMs are contained in the Merensky and UG2 reefs of the Bushveld Complex, where they are associated with nickel- and copper-bearing minerals. In Merensky ore, the major base-metal sulphide mineral is pyrrhotite, with pentlandite, chalcopyrite, pyrite, and minor amounts of other sulphides also present. In UG2 (Upper Group 2) ore the major base-metal sulphide is pentlandite. Pyrrhotite is found in moderate amounts, and millerite and pyrite in minor amounts. Because of lower mining costs, platinum mining is becoming more UG2-based, and the resulting concentrates contain high levels of chromite unless blended with Merensky ore. It is therefore critical that new and improved extraction methods be developed and exploited. The methods used in the recovery of the PGMs from these ores consist of physical concentration techniques, pyrometallurgical processing, and hydrometallurgical extraction of the base metals followed by the PGMs (Jones and Kotze, 2004; Nell, 2004).

During pyrometallurgical processing, the nickel-copper concentrates from the mill-float concentration step are smelted to bring about physical and chemical changes that enable recovery of base metals, PGMs, and other valuable metals in crude form. In general, the idea is to melt the concentrate in a furnace to produce a matte, which contains all the sulphides, below a carefully maintained slag layer. This matte, which contains large amounts of iron and sulphur, is oxidized in a Peirce-Smith converter to lower the iron and sulphur levels, while at the same time increasing the PGM grade. Conventional PGM matte smelting essentially requires the presence of a certain quantity of base metal sulphides in order to collect the PGMs in a molten sulphidic phase in the smelting furnace. However, the quantity of chrome oxide in the feed materials (particularly UG2) need to be strictly controlled to avoid the build-up of high-melting chromite spinels (Jones and Kotze, 2004; Nell., 2004).

Currently, the matte-based collection process is most widely used for PGM recovery, but because PGM-containing concentrates are becoming more enriched with UG2 concentrates, it is expected to be integrated with or replaced by an alternative process. Mintek has developed an alternative process for smelting PGM-containing oxide feed materials that contain low sulphur levels, and often high levels of chromium oxide, known as the Conroast process. The process involves the smelting of dead-roasted sulphide concentrates...
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in order to generate a small amount of an iron alloy in which PGMs and base metals are collected. A slag is also produced, which contains mostly unwanted materials and very low levels of residual PGMs. The desired degree of reduction is controlled by adjusting the carbon addition. Highly reducing conditions promote high recoveries of the valuable metals to the alloy, but high iron collection dilutes the PGM grade of the alloy, which affects the PGM recoveries in the converting and refining processes (Jones and Kotze, 2004; Nell J., 2004).

In this study, low-grade (LG) concentrates from Lonmin’s Rowland and Easterns operations were smelted with SiC as reductant. The effect of process parameters such as reductant to concentrate ratio, temperature, and different reductants on metal fall, alloy composition, slag composition, and gas composition were investigated. FactSage modelling was also used to simulate the reduction process. Detailed chemical and mineralogical characterization of the feed, alloy, and slag was conducted by X-ray diffraction (XRD), X-ray fluorescence (XRF), scanning electron microscopy (SEM), and inductively coupled plasma (ICP) analysis.

The primary objectives of this study include understanding the reducing conditions to produce the minimum amount of alloy while maintaining a high as possible recovery of PGMs; quantifying the deportment of the various elements to the alloy and slag phases; and establishing the factors affecting the recovery of PGMs; and comparing the gas emissions from different reductants in terms of environmental impact. The benefits of the new process would include no constrain on the minimum quantity of base metals required in the feed material, as the PGMs are collected in an iron-based alloy; ability to integrate the relatively large quantities of the alloy product into an existing smelter complex;可能性 of hydrometallurgical refining of the alloy; chromium tolerance; efficient PGM collection; fewer sulphur emissions compared to current matte-smelting processes; and possibly an economic incentive for treating other waste materials in a similar manner.

Materials and methods

A Carbolite STF 1800 tube furnace with a programmable Carbolite controller was used for the experiments. The alumina tube was heated by five surrounding lanthanum elements. A set point of 1600°C was set on the controller and the measured temperature recorded by a thermocouple located in the furnace. The initial plan was to attach a crucible to a wire and then suspend it from the top of the furnace at the tube outlet and lower it to the marked hot spot. However, in order to facilitate the operation, the furnace was removed from its original supporting brackets and fitted onto a fork-lifter. After the adjustment, the furnace could easily be moved up and down in vertical direction. The movement was carefully controlled by a hydraulic lever or foot pedal (Figure 1). After the furnace had reached the set temperature, the MgO crucible containing the sample mixture was placed into the tube furnace. Before the sample was placed into the hot zone, high-purity argon was allowed to purge into the Al work tube at a flow rate of 5 l/min for 20 minutes. The sample was then raised to the hot zone, and the Ar flow rate was reduced to about 600 ml/min at the start of the experiment. After a predetermined time of reduction, the

furnace was raised and the sample was allowed to cool rapidly under an Ar flow rate of 5 l/min for about 10 minutes.

The screen analysis showed that FeO values for Easterns concentrate, Rowland concentrate, and SiC reductant were 320 µm, 180 µm, and 380 µm respectively. The chemical compositions of the LG concentrates are given in Tables I and II.

Results and discussion

FactSage simulations

Initial FactSage simulations were conducted prior to laboratory-scale experiments in order to establish a range of optimum conditions. Thermodynamic equilibrium results are given in 100 kg concentrate per mass of reductant. The

| Constituent | Rowland LG concentrate (wt %) | Easterns LG concentrate (wt %) |
|-------------|-------------------------------|-------------------------------|
| Al₂O₃       | 5.41                          | 5.00                          |
| CaO         | 2.72                          | 3.03                          |
| Cr₂O₃       | 2.94                          | 4.31                          |
| Fe₂O₃       | 12.73                         | 14.24                         |
| K₂O         | 0.10                          | 0.10                          |
| MgO         | 22.79                         | 23.05                         |
| MnO         | 0.15                          | 0.18                          |
| Na₂O        | 0.38                          | 0.32                          |
| P₂O₅        | 0.02                          | 0.02                          |
| SiO₂        | 49.40                         | 47.81                         |
| TiO₂        | 0.31                          | 0.32                          |
| LOI *       | 3.08                          | 2.32                          |
| Total       | 100.05                        | 100.86                        |

* LOI includes the total volatiles content of the rock (including the water combined in the lattice of silicate minerals) and the gain on ignition related to the oxidation of the rock (mostly due to Fe₃O₄ converted to Fe₂O₃).
The stoichiometric amount of SiC required per 100 kg of LG concentrate is 0.27 kg for Rowland and 0.38 kg for Easterns. These were determined as the minimum amounts of SiC required to produce a molten metal phase, primarily being Fe (> 98%), given the starting oxide concentration and ease of reduction. Experimental results are then compared with FactSage results and discussed. Without any reductant in the system, MgCr$_2$O$_4$(s) spinel accumulates, therefore a reductant mass of 3 kg SiC per 100 kg LG concentrate was chosen as a base case. During the prediction of optimum temperature range, the requirement was that Fe concentrations should be high, while Cr and Si should remain low. The respective liquidus temperatures during SiC reduction are 1420°C for Rowland and 1410°C for Easterns concentrate. The results indicated that Fe and Cr concentrations in the alloy should not be significantly affected when the temperature increased from 1400°C to 1650°C, while Si concentrations would increase. The solubility of C in the alloy decreases with increasing temperature. The metal fall is not significantly affected, however, and about 1600°C should yield a good metal fall and avoid formation of solid phases like MgSiO$_3$(s) proto-enstatite. Therefore 1600°C was selected as the temperature at which to investigate the SiC reduction of the LG concentrates.

The effect of reductant to concentrate ratio was investigated from 0.5 kg to 4 kg SiC per 100 kg LG concentrate. It was found that an increase in reductant mass will cause more oxides, particularly FeO, to reduce, thereby increasing metal fall. For both concentrates a molten metal phase starts to form when 0.5 kg of reductant is present per 100 kg concentrate, emphasizing the ease of reduction of both concentrates, and almost all Cr$_2$O$_3$ is reduced to CrO, thereby increasing the solubility of Cr in the slag phase. The metal fall continues to rise and it therefore becomes easier to recover PGMs. Better metal fall for reductive smelting of Easterns LG concentrate is expected at higher reductant to concentrate ratios, due to the marginally higher FeO concentration. The molten metal phase consists primarily of Fe and other easily reduced base metals; however, some Cr and Si deport to the metal phase once the SiC to concentrate ratio is increased. Si and Cr start to dissolve in the molten metal at significant concentrations when the reductant mass exceeds 3 kg SiC per 100 kg LG concentrate. Concentrations in the alloy also become more significant at a higher SiC to concentrate ratio. Cr solubility in the slag phase is greatly affected by the oxidation state of Cr (Nell, 2004; Jones, 2009). Chrome in its trivalent state (spinel) has a very low solubility in the slag phase – divalent chrome, in contrast, is highly miscible in the slag phase (Nell, 2004). The FactSage simulation in Figure 2 shows how Cr is reduced from Cr(III) to Cr(II) as function of SiC additions to Easterns LG concentrate charge.

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| Table II | ICP-MS results for PGM values (wt %) in the LG concentrates |
|----------|------------------------------------------------------------|
|          | Au % | Ir % | Pd % | Pt % | Rh % | Ru % |
| Rowland LG concentrate | <0.005 | 0.057 | 0.10 | 0.059 | <0.005 | <0.005 |
| Easterns LG concentrate | <0.005 | 0.064 | 0.11 | 0.083 | <0.005 | <0.005 |

Figure 2 – The distribution of chromium between the phases as a function of SiC additions to Easterns LG concentrate.
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Experimental results
An alloy from each experiment was separated from the slag phase and weighed. The percentage metal fall for each experiment was determined as the amount of alloy formed per amount of initial concentrate. Some thermogravimetric analysis (TGA) experiments were also conducted to investigate the reaction mechanisms and kinetics. The alumina pedestal with the attached crucible containing the LG concentrate charge and SiC reductant were placed on a scale, which was connected to a computer with a VGA cable to acquire the change in mass of the concentrate charge. It was found that the mass of the crucible, refractory support material, and alumina pedestal remained constant throughout the experiment. Therefore the only change in mass was that of the concentrate and reductant charge in a gas-tight furnace. The mass changes on the scale were noted every 0.5 seconds on the computer, and typical graphs of mass loss were constructed. The results showed that no significant mass changes occurred after about 10 minutes of reduction time, indicating that equilibrium was reached due to the reduction of iron oxides, given their relatively high ratio of reduction time, indicating that equilibrium was reached.

Table III
Alloy composition (by ICP-MS) from SiC reduction of Rowland LG concentrate as a function of reductant addition and reduction time (wt %)

| SiC Concentrate | ratio (x10^2) | Reduction time (min) | Fe   | Cr   | Si   | Cu | Ni | S  | Ir | Pt | Pd |
|-----------------|---------------|----------------------|------|------|------|----|----|----|----|----|----|
| 2.5             | 60            | 81.2                 | 0.24 | <0.01 | 3.67 | <0.01 | <0.01 | 0.05 | 0.02 | 0.08 |
| 3.0             | 35            | 67.6                 | 8.81 | <0.01 | 2.65 | <0.01 | 0.01  | 0.04 | 0.01 | 0.08 |
| 3.0             | 45            | 71.7                 | 9.36 | <0.01 | 2.36 | <0.01 | <0.01 | 0.06 | 0.01 | 0.06 |
| 3.0             | 60            | 81.4                 | 2.72 | <0.01 | 2.08 | <0.01 | <0.01 | 0.06 | 0.01 | 0.06 |
| 3.0             | 75            | 76.4                 | 5.9  | <0.01 | 2.25 | <0.01 | <0.01 | 0.06 | 0.01 | 0.06 |
| 3.0             | 90            | 75.1                 | 6.12 | <0.01 | 2.67 | 4.42 | 1.11  | 0.04 | 0.07 | 0.04 |
| 3.0             | 180           | 74.9                 | 6.35 | <0.01 | 2.71 | 4.51 | 1.13  | 0.08 | 0.06 | 0.06 |
| 3.5             | 60            | 66.1                 | 9.59 | 0.01  | 2.02 | <0.01 | 0.01  | 0.04 | <0.01 | 0.06 |

Table IV
Alloy composition (by ICP-MS) from SiC reduction of Easterns LG concentrate as a function of reductant addition and reduction time (wt %)

| SiC Concentrate | ratio (x10^2) | Reduction time (min) | Fe   | Cr   | Si   | Cu | Ni | S  | Ir | Pt | Pd |
|-----------------|---------------|----------------------|------|------|------|----|----|----|----|----|----|
| 2.5             | 180           | 81.2                 | 0.24 | <0.01 | 3.67 | <0.01 | <0.01 | 0.03 | 0.02 | 0.08 |
| 3.0             | 180           | 84.5                 | 1.47 | 0.05  | 1.47 | 4.49 | 2.04  | 0.8  | 0.52 | 1.24 |
| 3.5             | 180           | 88.6                 | 1.33 | 0.15  | 1.08 | 2.81 | 1.52  | 0.8  | 0.66 | 1.26 |
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Table V
Recovery of Ir, Pd, and Pt from SiC reduction of Rowland LG concentrate

| SiC Concentrate ratio(x10^2) | Reduction time (min) | Ir recovery (%) | Pd recovery (%) | Pt recovery (%) |
|------------------------------|----------------------|-----------------|-----------------|-----------------|
| 2.5                          | 60                   | 2.410           | 2.198           | 0.931           |
| 3.0                          | 75                   | 4.449           | 3.233           | 0.86            |
| 3.0                          | 90                   | 3.957           | 2.637           | 6.497           |
| 3.0                          | 180                  | 3.816           | 3.516           | 7.914           |
| 3.5                          | 60                   | 4.082           | 2.905           | 0.889           |

Table VI
Recovery % of Ir, Pd, and Pt from SiC reduction of Easterns LG concentrate

| SiC Concentrate ratio(x10^2) | Reduction time (min) | Ir recovery (%) | Pd recovery (%) | Pt recovery (%) |
|------------------------------|----------------------|-----------------|-----------------|-----------------|
| 2.5                          | 180                  | 11.232          | 1.538           | 0.599           |
| 3.0                          | 180                  | 69.728          | 62.882          | 34.95           |
| 3.5                          | 180                  | 94.536          | 86.829          | 59.221          |

Table VII
Slag composition (by XRF) from SiC reduction of Rowland LG concentrate

| SiC Concentrate ratio(x10^2) | Reduction time (min) | Al2O3 | CaO | CrO* | FeO* | MgO | SiO2 | TiO2 | Total |
|------------------------------|----------------------|-------|-----|------|------|-----|------|------|-------|
| 2.5                          | 60                   | 5.2   | 2.67| 2.25 | 5.87 | 27.60| 52.17| 0.33 | 97.54 |
| 3.0                          | 75                   | 5.22  | 2.72| 2.43 | 3.35 | 28.35| 52.17| 0.32 | 96.72 |
| 3.5                          | 90                   | 5.23  | 2.70| 2.34 | 3.07 | 29.21| 52.11| 0.33 | 97.11 |
| 3.0                          | 180                  | 5.22  | 2.69| 2.32 | 3.03 | 28.11| 52.09| 0.33 | 96.91 |
| 3.5                          | 180                  | 5.22  | 2.69| 1.61 | 2.97 | 28.74| 54.01| 0.33 | 97.72 |

Table VIII
Slag composition (by XRF) from SiC reduction of Easterns LG concentrate

| SiC Concentrate ratio(x10^2) | Reduction time (min) | Al2O3 | CaO | CrO* | FeO* | MgO | SiO2 | TiO2 | Total |
|------------------------------|----------------------|-------|-----|------|------|-----|------|------|-------|
| 2.5                          | 180                  | 4.75  | 3.08| 3.11 | 6.51 | 29.11| 50.17| 0.33 | 97.56 |
| 3.0                          | 180                  | 4.71  | 3.1  | 3.09 | 5.84 | 30.06| 49   | 0.33 | 97.61 |
| 3.5                          | 180                  | 4.78  | 3.12| 3.15 | 3.12 | 29.78| 51.85| 0.33 | 97.10 |

Alloy are very favourable for downstream processing because of the difficulties in removing these elements during converting and refining. The Pt and Pd concentrations are higher in alloys from SiC reduction of Easterns LG concentrate. The better metal fall is the most likely factor to have contributed to the higher PGM concentrations in these alloys. PGM recoveries (Ir, Pd, and Pt) to the alloy phase for the respective concentrates are shown in Table V and Table VI.

From Table V it is seen that the recoveries of Ir, Pt, and Pd are very low. The highest Pt and Pd recoveries were achieved at a reductant to concentrate ratio of 3 kg SiC per 100 kg Rowland LG concentrate and residence time of 180 minutes. One factor that may contribute significantly to the low PGM recoveries is slag viscosity. The recoveries of Ir, Pt, and Pd from SiC reduction of Easterns LG concentrate are significantly better (Table VI). Ir, Pd, and Pt recoveries increase significantly when the quantity of reductant is increased. Overall, Ir has the highest recovery, followed by Pd and Pt. The very low Cr and Si concentrations in these alloys will allow for a further increase in reductant quantity and therefore it should be possible to recover more than 90% of PGMs by SiC reduction of Easterns LG concentrate. The slag compositions from the SiC reduction of Rowland and Easterns LG concentrates are shown in Tables VII and VIII respectively.

The results from Tables VII and VIII show a significant decrease in FeO concentration compared with the initial concentrates. The lowest FeO content corresponded to the experiments with the highest quantity of reductant. The composition of the slag has a major effect on the viscosity,
The recovery of platinum group metals from low-grade concentrates to an iron alloy and FeO, NiO, CuO, and CrO will all aid in lowering the viscosity. Once the SiC comes in contact with these oxides, it will reduce and settle to join the alloy phase, thereby removing them from the slag phase and increasing the viscosity. It can be seen that the reduction process lowered the FeO concentration compared with the initial concentrate (refer to Table I).

The viscosity module in FactSage was used to determine the viscosities at 1600°C. At a temperature of 1600°C it can be safely assumed that the slag is molten. The calculated viscosities of the slags from SiC reduction of Rowland and Easterns LG concentrate at the beginning of the melts are 3.74 and 3.06 poise, respectively. The significantly higher slag viscosity for Rowland LG concentrate would make it more difficult for small metal prills to settle and coalesce.

Table IX shows viscosities of the slags at the end of the melt. The viscosities were calculated in the FactSage viscosity module from the slag compositions in Table VII and Table VIII.

Table IX shows that the calculated viscosities of the slags from all the tests are higher than the calculated viscosities at the beginning of the melts. This is because the bulk of the Fe has partitioned into the alloy. Some Cr has also deported to the alloy phase. The viscosities of slags from SiC reduction of Rowland LG concentrate are significantly higher than slags from SiC reduction of Easterns LG concentrate. This could explain why the PGM recoveries from Rowland LG concentrate are significantly lower.

Viscosities of PGM smelting slags are reported in the literature (Eric, 2004; Eric and Hejja, 1995) as being in the range 1.5–4 poise, which could hinder the settling of particles 15 µm or smaller in size. The FeO/SiO₂ ratio of the slag seems to have a major influence on the viscosity. The Rowland LG concentrate under investigation has a lower FeO/SiO₂ ratio than Easterns LG concentrate. When FeO is reduced, slag viscosity will increase and valuable metal prills will not settle. The high initial slag viscosity will contribute to insufficient coalescence of metal prills and phase separation. The corresponding viscosities from these experiments conducted with Rowland LG concentrate fall mostly above the range of 1.5–4 poise. It therefore seems that high slag viscosities are the primary reason for the low PGM recoveries. Slag viscosities from SiC reduction of Easterns LG concentrate do, however, fall in the recommended range. The lower viscosities will aid in the coalescence of metal prills, enhance phase separation, and induce good PGM recovery.

Figures 3 and 4 are SEM images of the slag phases from SiC reduction of Rowland and Easterns LG concentrate.
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respectively, from experiments that had the same reductant to concentrate ratio and reduction time.

Figure 3 shows a SEM image of the slag phase from SiC reduction of Rowland LG concentrate. Metal prills are entrained in the slag. These metal prills are very small and are most likely trapped in the slag phase and will never settle or coalesce with other metal prills of similar size. Studies in the literature indicate that many small metal prills (< 2 µm) are expected never to settle under the force of gravity alone. This phenomenon agrees well with the work done by Fagurland and Jalkanen (1999) and Poggie et al. (1969), who also reported that very small droplets may either settle very slowly or may be trapped as ‘rafts’ of droplets floating on the top surface of the slag or as droplets suspended below small gas bubbles. However, the high viscosity of the slag is most likely restricting the coalescence and settling of metal prills. An EDX spot analysis on the metal prill indicated by the arrow detected Ir and Pd.

Figure 4 shows a SEM image of the slag phase from SiC reduction of Easterns LG concentrate. The slag phase is smoother, with no metal prills entrained. This is an indication that metal prills coalesced and settled well within the slag phase. From these findings it is very clear that the lower slag viscosity contributes significantly to PGM recovery from SiC reduction of Easterns LG concentrates. This corresponds to similar findings in the literature (Shahrokhi and Shaw, 2000; Saffman and Turner, 1956; Ammann et al., 1979).

From the results obtained experimentally, it was concluded that Easterns LG concentrate should form the bulk of the dry LG concentrate feed to a DC arc furnace, since it has a higher FeO/SiO\textsubscript{2} ratio, which is significant in reducing slag viscosity and increasing overall PGM recovery. The slag viscosities need to be kept in the region of approximately 3.2 poise (average viscosity determined from experiments with SiC reduction of Easterns LG concentrate) for acceptable PGM recoveries. Rowland and Easterns concentrates could be combined, with the main fraction being Easterns, or another source added that would lower slag viscosity. The slag from a Petrice-Smith converter has a high FeO/SiO\textsubscript{2} ratio and could be used to reduce the slag viscosity. FactSage was used to model such a scenario and an experiment was also conducted to back up the results. FactSage modelling was conducted at 3.5 kg SiC per100 kg Easterns LG concentrate at 1600°C with varying amounts of converter slag additions.

Figure 5 shows that addition of converter slag does decrease the slag viscosity. The converter slag is rich in FeO and thereby the FeO/SiO\textsubscript{2} ratio in the slag is increased, causing the slag viscosity to decrease. However, the converter slag contains about 30 wt% SiO\textsubscript{2}, which could act to increase the slag viscosity. Figure 5 shows that an increase in reductant will cause more FeO to be reduced, consequently a system with 5 kg and 10 kg converter slag additions will have a higher slag viscosity than a system with no converter slag if the reductant additions are increased past the point of intersection. Another critical point is that the conductivity and viscosity of the slag are inversely related. A decrease in viscosity will increase the conductivity of the slag; increasing iron oxide serves to depolymerize the slag and creates sites for electronic conduction (Nell, 2004; Eric, 2004). It is clear from the findings that around 10 kg of converter slag addition per 100 kg concentrate would decrease slag viscosity without increasing the total quantity of slag significantly. The alloy is also expected to have a good PGM grade. A test was therefore conducted to compare the results from experiments and FactSage modelling for similar conditions.

From Table X, it is apparent that FactSage predicts a slag with similar composition. The marginally lower FeO/SiO\textsubscript{2} ratio in the slag from FactSage modelling will cause the viscosity to be somewhat higher than the slag from the experiment. Table X also shows an improved PGM recovery with converter slag addition, which is likely to be a result of the decrease in slag viscosity. The Pd and Ir rd recoveries are nearly 100%, compared to Pd and Ir recoveries of 94.5 and 86.6% obtained experimentally with no slag addition. Pt recovery has also improved, from 59.2% to almost 70%. Pt recovery is expected to improve if the metal fall increases. More reductant could be added to increase metal fall and thereby increase the Pt recovery from the initial concentrate. Overall, this is a significant finding. A portion of converter slag can therefore be recycled and re-melted.
The recovery of platinum group metals from low-grade concentrates to an iron alloy together with any LG concentrate to increase metal fall and PGM recoveries.

FactSage modelling was used to predict an optimum ratio between Rowland and Easterns LG concentrates. The reductant to concentrate ratio was fixed at 3.5 kg SiC per 100 kg LG concentrate and the temperature at 1600°C. FactSage predicted that adding more than 30 kg converter slag per 100 kg LG concentrate decreases the alloy to slag ratio. Therefore a 0.7–0.8 fraction of Easterns LG concentrate in the feed and 20–30 kg converter slag per 100 kg LG concentrate should not significantly increase furnace slag quantities and the FeO/SiO$_2$ ratio should be adequate to sustain a slag viscosity close to 3.5 poise. At these operating conditions more than 90% of the PGMs should be recovered from a LG concentrate feed in an alloy with a high PGM grade. Cr and Si concentrations in the alloy are less than 1% in total, and C concentrations are less than 1%.

FactSage modelling was also undertaken to compare SiC reduction with carbon reduction. The optimum feed conditions, namely 75% Easterns LG concentrate, 25% Rowland LG concentrate, 25 kg converter slag per 100 kg LG concentrate, and 0–4 kg reductant per 100 kg LG concentrate at 1600°C were used in the comparison. With these conditions, it is assumed that slag viscosity will be low enough to ensure good phase separation and PGM recovery. The results indicated that C reduction of a LG concentrate charge results in a marginally higher metal fall. The alloy compositions were very similar and only small differences in Fe, Cr, and Si concentrations were noted. Gas emissions and energy requirements are higher for C reduction, arguably due to C reacting endothermically with FeO to produce Fe(l) and CO(g) in contrast to SiC reacting exothermically with FeO to produce Fe(l), SiO$_2$(l) and CO(g).

However, it must be borne in mind that SiC is not a naturally occurring compound and is manufactured by carbothermic reduction of silica at high temperatures, which entails its own high level of emissions which should be added to the emissions from SiC smelting. A preliminary FactSage analysis was conducted for the reaction to produce SiC:

$$\text{SiO}_2(s) + 3\text{C(s)} \rightarrow \text{SiC(s)} + 2\text{CO(g)} (+\text{O}_2 \rightarrow 2\text{CO}_2) \ [3]$$

The resulting estimates of energy requirements and CO emissions per ton of SiC product are given in Table XI.

It is seen from Table XI that not all of the SiO$_2$ and C has...
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reacted to form SiC. Thermodynamic modelling predicts that a small fraction of the SiO₂ will be converted to SiO₂(g), leaving some unreacted C(s) in the system. Overall more than 1.3 t of CO (g) is emitted to produce 1 t of SiC. Moreover, the energy required to produce the given SiC quantities will contribute to overall energy requirements for recovery of PGMs from an iron alloy. This needs to be taken into consideration from an environmental and economic point of view when considering a reductive smelting route. It was, however, determined that only approximately 35 kg SiC per ton of LG concentrate is required to recover most of the PGMs. It must also be taken into consideration that these PGMs need to be extracted from a more enriched LG concentrate feed, and reductive smelting is the only feasible option at the present time.

Conclusions

The feasibility of using SiC as reductant for Rowland and Easterns LG concentrates was investigated with respect to metal fall, PGM grade in the alloy, slag composition, and PGM recovery, using small-scale experiments in combination with FactSage modelling. The findings can be summarized as follows.

- SiC reduction of Easterns LG concentrate resulted in significantly better metal fall and PGM recoveries compared with Rowland concentrates. At a reductant to concentrate ratio of 3.5 kg SiC per 100 kg Easterns LG concentrate, more than 85% of Ir and Pd were recovered, and more than 60% of Pt. The higher FeO/SiO₂ ratio of Easterns LG concentrate and consequent lower slag viscosity is most likely to have contributed to the improvement of recoveries. SEM images from a slag from SiC reduction of Easterns LG concentrate showed no entrained metal prills, indicating that most metal prills had coalesced and settled. Cr and Si concentrations were below 2% in total in all alloys.

- In order to improve the PGM recoveries, the FeO content of the initial charge had to be increased. Peric-Smith converter slag from Lonmin was used as an addition to increase FeO/SiO₂ ratio. The experiments resulted in more than 95% recoveries for Ir and Pd, together with about 70% for Pt. FactSage modelling predicted that Easterns LG concentrate should make up 70-80% of the LG concentrate charge and 20-30 kg converter slag be added per 100 kg LG concentrate to attain relatively lower viscosities for phase separation. It is expected that PGM recoveries of more than 90% should be obtained from the LG concentrate, in conjunction with a good PGM grade in the alloy.

- Using the optimum LG concentrate charge, the effectiveness of SiC as a reductant was also compared to that of C through thermodynamic modelling. C reduction of a LG concentrate charge led to a marginally higher metal fall at the same reductant to concentration ratio compared with SiC reduction. The alloy composition was very similar and only small differences in Fe, Cr, and Si concentrations were noted. Gas emissions and energy requirements were higher for C reduction of a LG concentrate charge.

- From the evidence, it is clear that SiC reduction seems a reasonably attractive alternative to carbon-based reductants. Therefore, integrating this process into the present matte-based collection flow sheet could be considered as a future alternative for smelting LG concentrates. However, it must be borne in mind that SiC is not a natural compound and it should be manufactured by again carbothermic reduction of silica at high temperatures with its own level of emissions. This obviously will require further investigation.

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