Investigation into the dephosphorization of ferromanganese alloys for the production of advanced high-strength steel

M.P. Maphutha¹,², J.D. Steenkamp¹, and P.C. Pistorius²,³

Synopsis
Advanced high-strength steels (AHSS) are sophisticated materials being developed by the steel industry to mitigate challenges related to the performance of motor vehicles. To meet the requirements of AHSS, the ferromanganese alloys (FeMn) utilized in the production of the steel are required to contain acceptable levels of unwanted impurities, i.e., P, S, N, H, and C. The focus of the current study was to investigate dephosphorization of ferromanganese to produce a low-P alloy that could be effectively utilized in the production of AHSS. The study involved conducting laboratory-scale testwork to study the efficiency of CaO-based slag systems to dephosphorize FeMn alloys. The addition of Na₂O, CaF₂, and BaO to MnO-CaO-SiO₂ slag was considered. The test work was carried out in a 25 kW induction furnace at temperatures of 1350°C, 1400°C, and 1450°C. The P partition coefficient \( L_p \) remained small at <1, which is an indication that dephosphorization had not been achieved. The baseline slag, comprising 40%CaO-40%SiO₂-20%MnO, reported higher \( L_p \) values. Addition of Na₂O and CaF₂ did not show any further benefit. Substituting half of the CaO by BaO, resulted in similar \( L_p \) values to those of the baseline slag under conditions of 1350°C and 1450°C at 30 minutes. In summary, based on the \( L_p \) values obtained, the conditions investigated with the CaO-based slags appeared to have been unfavourable for dephosphorization of FeMn alloys, as most of this impurity element remained in the alloy.

Keywords
ferromanganese, dephosphorization, advanced high-strength steel.

Introduction
In the automotive industry the drive towards lightweight, high-strength steel grades to mitigate the challenges around the escalating energy crisis and environmental problems is a priority. It has been estimated that a 10% weight reduction in automobiles would reduce fuel consumption by between 3% and 7% (Demiri, 2013). This has led to the development of advanced high-strength steels (AHSS) (Baluch, Udin, and Abdullah, 2014).

AHSS steels are a group of special steel grades which offer excellent strength, allowing the use of thinner gauges to reduce the weight of vehicles. As the manganese (Mn) content in AHSS can be up to 25%, these steels could potentially be a significant market for manganese ferroalloys (Safarian and Kolbeinsen, 2013). The challenge lies in the maximum allowable phosphorus (P) content of <0.01% in the steel grades (Bernhard et al., 2019).

P has a detrimental effect on the strength, brittleness, ductility, and fracture toughness of steel. As a result, the quality of the steel could become compromised, especially in the case of AHSS where the Mn content can be up to 25%. As a consequence, FeMn alloys utilized in the production of AHSS are required to contain low P levels (You, Lee, and Pak, 1999; Chaudhary and Roy, 2001).

During carbonaceous reduction of Mn ores to produce FeMn alloys, almost all the P is reduced to the alloy phase. The P originates mainly from the Mn ores, and is intimately associated with iron (Fe) and Mn, making it difficult to remove by mineral processing routes (Chaudhary and Goel, 2007). South African Mn ores are, however, generally low in P compared to ores from other countries such as China. The P content in the typical South African ores, namely Gloria ore and Nchwaning ores, varies between

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0.02% and 0.05% (Visser et al., 2013; Olsen et al., 2007). When using low-P ore, the P present in the slag and the alloy originates mainly from the carbonaceous reductants used (Olsen et al., 2007). To produce FeMn alloys containing P at levels acceptable to AHSS producers, methods have to be developed to lower the P levels in the alloy.

Background

Mechanisms of dephosphorization

Dephosphorization of the molten alloy under oxidizing conditions can be described by the following ionic reaction (Chaudary Minji, and Goel, 2007; Nasaralla, Fruhehan, and Min, 1991; Simeonov and Sano, 1985):

\[
\frac{3}{2}(O^{2-}) + [P] + \frac{5}{2}[O] = (PO_4^{3-})
\]

where:

[X]: species dissolved in the alloy

(Y): species dissolved in the slag.

The equilibrium constant can be expressed as follows:

\[
K_1 = \frac{a_{(PO_4^{3-})}}{a_{[P]} a_{[O]^\frac{5}{2}}}
\]

Based on the above expression, it can be deduced that the removal of P from the alloy can be aided by the following (Chaudary and Goel, 1994):

- Higher oxygen activity in the alloy
- High activity of basic oxide (\(\alpha_{O^2-}\)) in the slag
- Low activity coefficient of phosphate (\(\gamma_{PO_4^{3-}}\)) in the slag
- High \(K\) values, which can be achieved at low temperatures since the reaction is of exothermic nature.

Slag basicity is associated with the degree of polarization of oxygen in the slag. Highly basic slag contains more free oxygen ions (\(O^{2-}\)) and less bridging and terminal oxygen. Slag basicity can therefore be defined by the activity of oxygen ions in slags (Liu et al., 1998). As indicated by Equation [1], high concentrations of \(O^{2-}\), and therefore basic slags, are efficient for dephosphorization (Wagner, 1975).

Dephosphorization of FeMn can also be carried out with basic slag under reducing conditions to form a phosphide species. The reaction is shown below (Chaudary and Goel, 2007):

\[
[P] + \frac{5}{2}(O)^{2-} = P^{3-} + \frac{3}{4}O_2
\]

Based on Equation [3], it can be deduced that the removal of P from the alloy under these conditions can be aided by a high activity of basic oxide (\(\alpha_{O^2-}\)) in the slag. A relatively lower oxygen partial pressure is required for reducing conditions, and CO gas is typically used. Metallurgies such as Ca, typically added as calcium silicide or CaC\(_2\), would dissolve into the metal and react with P to form phosphide. The reducing reaction occurs through the reaction between CaO in the flux and Si in the melt (for SiMn alloy), and also through the transfer of Ca from the calcium silicide introduced into the melt. An example is shown below (Karbowiczek et al., 2014):

\[
[Si] + 2(CaO) = 2[Ca] + (SiO_2)
\]

\[
3[Ca] + 2[P] = (Ca_3P_2)
\]

The reaction product from Equation [5] is phosphide, which in the presence of atmospheric moisture produces toxic phosphene. Dephosphorization of FeMn alloy under reducing conditions is therefore not desirable.

P removal in vapour form, as phosphine gas, by the reaction of P with hydrogen dissolved in the metal can occur (Chaudary and Goel, 1994):

\[
[P] + 3[\text{H}] = \text{PH}_3 \hspace{1cm} \text{(g)}
\]

The reaction product (phosphine) would not be environmentally tolerable when released into the atmosphere (Chaudary and Goel, 1994). Based on Equation [6], it can be deduced that the removal of P from the alloy under these conditions can be aided by higher hydrogen activity in the alloy. Dephosphorization under gaseous conditions may also result in loss of Mn, which has a higher vapour pressure than P. Application of the technique is deemed undesirable for dephosphorization of FeMn alloys (Chaudary and Goel, 1994).

Considering the abovementioned reaction mechanisms, dephosphorization of liquid FeMn alloy by basic slags for selective removal of P over Mn under oxidizing conditions is deemed the most viable route (Chaudary and Goel, 1994). Based on Equation [6], it can be deduced that the removal of P from the alloy under these conditions can be aided by higher hydrogen activity in the alloy. Dephosphorization under gaseous conditions may also result in loss of Mn, which has a higher vapour pressure than P. Application of the technique is deemed undesirable for dephosphorization of FeMn alloys (Chaudary and Goel, 1994).

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The study involved conducting preliminary thermochemical FactSage calculations followed by experimental laboratory tests to investigate the dephosphorization of South African FeMn alloys by different synthetic CaO-based slag systems, similar to the systems applied in dephosphorization of iron – for the dephosphorization of FeMn alloys commercially produced in South Africa.

Method

The study involved conducting preliminary thermochemical FactSage calculations followed by experimental laboratory tests to investigate the dephosphorization of South African FeMn alloys by different synthetic CaO-based slag systems, similar to the systems applied in dephosphorization of iron.

FactSage calculations

FactSage™ version 7.3 was utilized. The package is a fully integrated thermodynamic database computing system developed in 1976 by a joint research project between McGill University and École Polytechnique de Montréal (Canada). FactSage provides access to several databases, including pure substances, oxides, solutions, and alloy databases. The tool is widely applied to study thermochemistry for different chemical and metallurgical processes (Bale et al., 2016).

The calculations were conducted in the Equilib module. In all calculations, the pure substance database (FactPS), which is suitable for pure solids, liquids, and gases, was utilized. As solution databases, FToxid was utilized to describe the slag system and FSstel the alloy system. FToxid includes a wide range of components such as \(\text{Al}_2\text{O}_3\), \(\text{As}_2\text{O}_3\), \(\text{B}_2\text{O}_3\), \(\text{CaO}\), \(\text{CoO}\), \(\text{CrO}\), \(\text{Cr}_2\text{O}_3\), \(\text{CuO}\), \(\text{FeO}\), \(\text{Fe}_2\text{O}_3\), \(\text{GeO}_2\), \(\text{K}_2\text{O}\), \(\text{MgO}\), \(\text{MnO}\), \(\text{Na}_2\text{O}\), \(\text{NiO}\), \(\text{PbO}\), \(\text{SiO}_2\), \(\text{SnO}\), \(\text{TiO}_2\), \(\text{Ti}_2\text{O}_3\), \(\text{ZnO}\), and \(\text{ZrO}_2\). Furthermore, FToxid contains many solid solution databases, which include those listed below (Bale et al., 2009):

- Wollastonite: \(\text{CaSiO}_3\), \((\text{FeSiO}_3\), \(\text{MgSiO}_3\), \(\text{MnSiO}_3\)
- Olivine: \((\text{Ca}_2\text{O}_2\), \(\text{CO}_2\), \(\text{Fe}_2\text{O}_2\), \(\text{Mg}_2\text{O}_2\), \(\text{Mn}_2\text{O}_2\), \(\text{Ni}_2\text{O}_2\), \(\text{Zn}_2\text{O}_2\)
- \(\text{Ca}_3\text{Si}_2\text{O}_7\): \(\text{Ca}_3\text{Si}_2\text{O}_7\), \((\text{Fe}_2\text{SiO}_4\), \(\text{Mg}_2\text{SiO}_4\), \(\text{Mn}_2\text{SiO}_4\)
- Monoxide: \(\text{CaO-MgO-MnO-CoO-NiO-FeO} + \text{Fe}_2\text{O}_3\), \(\text{Al}_2\text{O}_3\), \(\text{ZnO-Cr}_2\text{O}_3\)
- Corundum: \(\text{Al}_2\text{O}_3\), \(\text{Cr}_2\text{O}_3\), \(\text{Fe}_2\text{O}_3\)

FSstel is a steel database covering a wide range of compositions related to steelmaking processes (Bale et al., 2009).
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Although thermodynamic data exists for HCFeMn, it does not contain equilibrium data on P (Tang and Olsen, 2006). Fstel has been proven in the past to represent equilibrium conditions in SiMn production, and as it contained data on P, was selected for the calculations presented here (Steenkamp, Pistorius, and Tangstad, 2015).

As pure species, all liquids and solids were selected except the Mn phosphate phases. As pure ideal gas species, only O₂ and Ar were selected. For solution systems, all solution phases were selected, except in cases where only one option existed. Solution phases that were suppressed were SlagD, Slag? MeP?, cPyrB, and cPyr?. SlagA model was selected over Slag? model because SlagA provides much more reasonable L_p values for steelmaking than does Slag?, which overestimates P removal. Since MeOB, does not include BaO as a solute, this solution phase was not selected. MeP?, cPyrB, and cPyr? were not selected because more than one option existed.

The calculations were done to investigate the phosphorus partition coefficient (L_p) between slag and alloy. The expression for L_p is

\[ L_p = \frac{\text{Wt}\% \text{P}_{\text{FeOxid}} \text{SLAGA}}{\text{Wt}\% \text{P}_{\text{Fstel-Liq}}} \]  

[7]

The typical HCFeMn and MCFeMn compositions obtained from the literature (see Table I) were utilized for the dephosphorization thermochemical calculations.

The slag systems in Table II were utilized. To determine the L_p for the different slag-alloy systems, 100 g of the alloy was reacted with 10 g of slag, i.e. the slag-to-alloy mass ratio was maintained constant at 0.1 (but equilibrium is essentially independent of the relative masses, other than the effect of equilibration of Mn and Si between metal and slag). A small amount of Ar (0.01 g) was also added to allow for reactions to converge (Steenkamp, Pistorius, and Tangstad, 2015). The temperature range considered was 1300–1700°C at 50°C intervals.

### Table I

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Mn</th>
<th>C</th>
<th>Si</th>
<th>P</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCFeMn</td>
<td>78.00</td>
<td>7.50</td>
<td>0.30</td>
<td>0.18</td>
<td>14.00</td>
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<tr>
<td>MCFeMn</td>
<td>80.00</td>
<td>1.50</td>
<td>0.60</td>
<td>0.20</td>
<td>17.70</td>
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### Table II

<table>
<thead>
<tr>
<th>Slag label</th>
<th>Slag systems</th>
<th>Total</th>
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<tbody>
<tr>
<td></td>
<td>MnO</td>
<td>CaO</td>
</tr>
<tr>
<td>A</td>
<td>20.0</td>
<td>40.0</td>
</tr>
<tr>
<td>B</td>
<td>25.0</td>
<td>37.5</td>
</tr>
<tr>
<td>C</td>
<td>15.0</td>
<td>42.5</td>
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<tr>
<td>D</td>
<td>20.0</td>
<td>40.0</td>
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<tr>
<td>E</td>
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<td>20.0</td>
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<tr>
<td>F</td>
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</tr>
<tr>
<td>G</td>
<td>20.0</td>
<td>20.0</td>
</tr>
<tr>
<td>H</td>
<td>20.0</td>
<td>40.0</td>
</tr>
</tbody>
</table>

### Table III

<table>
<thead>
<tr>
<th>Target chemical compositions of different slag systems</th>
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</thead>
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<tr>
<td>Slag label</td>
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<tr>
<td>------------</td>
</tr>
<tr>
<td></td>
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<td>A</td>
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<td>B</td>
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<tr>
<td>C</td>
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<tr>
<td>D</td>
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<tr>
<td>E</td>
</tr>
<tr>
<td>F</td>
</tr>
<tr>
<td>G</td>
</tr>
<tr>
<td>H</td>
</tr>
</tbody>
</table>

### Table IV

<table>
<thead>
<tr>
<th>Average bulk chemical composition of the industrial MCFeMn alloy and HCFeMn master alloys utilized in the study (mass%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Element</td>
</tr>
<tr>
<td>-------------</td>
</tr>
<tr>
<td>Mn</td>
</tr>
<tr>
<td>Fe</td>
</tr>
<tr>
<td>Si</td>
</tr>
<tr>
<td>C</td>
</tr>
<tr>
<td>P</td>
</tr>
<tr>
<td>S</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

### Materials and equipment

Two FeMn alloys were utilized for the dephosphorization experiments, namely MCFeMn and HCFeMn alloys. The MCFeMn and HCFeMn alloys were sourced from an industrial producer in South Africa. HCFeMn was utilized in only a few cases to investigate experimentally which of the two alloys would be easier to dephosphorize. Table III summarizes the average chemical composition of the MCFeMn alloy and HCFeMn master alloy. Because a graphite crucible was used for the experiments, these alloys converged on a similar carbon concentration after the reaction.

The slag reagents utilized in the study were sourced from the commercial suppliers Sigma Aldrich and ACE chemicals. Analytical grade CaO, SiO₂, MnO₂, CaF₂, BaCO₃, Na₂CO₃, and BaF₂ were utilized. Different synthetic slags were prepared. The slag components, i.e. CaO, SiO₂, Na₂CO₃, and BaCO₃, were calcined separately at 1000°C for 2 hours in air to remove volatile matter. Decomposition of the carbonaceous matter for Na₂CO₃ and BaCO₃ was, however, not successful at 1000°C. The calcined materials were then weighed and blended in a ring mill to generate a homogeneous mixture and charged into a graphite crucible. The crucible charges were subsequently heated in an induction furnace at 10°C/min to 1600°C and maintained at the temperature for 1 hour under an inert atmosphere using Ar gas. The molten materials were allowed to cool to room temperature under an inert atmosphere. The crucibles were then retrieved from the crucibles, crushed, milled, and decarburized at 1200°C for 2 hours in air, using alumina crucibles to burn off the residual carbon. The decarburized slags were analysed to determine their elemental compositions.

The MnO₂ reagent was not added during the melting stage of the synthetic slags to avoid reaction with the graphite crucibles. The component was incorporated after the decarburizing stage. BaCO₃ was also added only to the decarburized slags. The inclusion of BaCO₃ during the melting stage resulted in the erosion of the graphite crucibles.

### Equipment

The dephosphorization tests were conducted in a 25 kW induction furnace Figure 1 presents the schematic diagram of the furnace. The furnace operates by radiation heating from a graphite susceptore heated by magnetic induction created by a water-cooled copper coil. The furnace power is switched on by using an on/off button on the control panel. The power was adjusted manually; the temperature was controlled by manually increasing the power setpoint at an interval of 30 minutes until the target temperature was reached. As it is generally difficult
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Recipe design

Baseline CaO-based slag (Slag A)

In the current study, the slag system comprising 20%MnO-40%CaO-40%SiO\textsubscript{2} was adopted as a baseline slag. The basic oxide CaO was selected to provide free oxygen (O\textsuperscript{2−}) when dissolved in the liquid slag (refer to Equation [1]). MnO was added to provide the oxidizing condition in the system. The acidic SiO\textsubscript{2} was selected as a flux to reduce the melting point of the high-melting, basic MnO-CaO binary system. Figure 2 presents the MnO-SiO\textsubscript{2}-CaO phase diagram, showing the fluxing strategy chosen and the expected change in slag liquidus temperature at 40% SiO\textsubscript{2} addition. As shown in Figure 2, the baseline slag has a liquidus temperature of 1320°C.

Na\textsubscript{2}O-CaF\textsubscript{2} addition (Slag D, Slag E, and Slag F)

The addition of Na\textsubscript{2}O has proven to enhance the ability of CaO-based slags to dephosphorize liquid iron (van Niekerk and Dippenaar, 1998; Nassaralla and Fruehan, 1992). CaF\textsubscript{2}, on the other hand, is added during hot metal treatment to maintain the fluidity of lime-based slags. Both additives have been reported to influence the phosphate capacities of basic slags (Fujita et al., 1988) and were therefore investigated in the current study. CaO was partially replaced with Na\textsubscript{2}O and CaF\textsubscript{2}, while the MnO content was maintained constant.

BaO addition (Slag H)

Chaudhary and Roy (2001) found the BaO-BaF\textsubscript{2}-MnO slag system to be effective for the dephosphorization of FeMn alloys, as well as liquid iron when BaO was utilized. The effect of BaO addition was investigated in the current study by replacing a fraction of the CaO with BaO.

BaO-BaF\textsubscript{2} addition (Slag I)

Replacing both the lime and silica by BaO and BaF\textsubscript{2} flux was also investigated. This slag system has been reported to be effective for dephosphorization of high-carbon FeMn alloy (Chaudary, Goel, and Minz, 2008; (Fujita et al., 1988; (Dashevski et al., 1998). The use of BaO-based slag without CaO posed numerous challenges during the current test work. The slag was very wetting towards the graphite crucible and spilled from the crucible due to what is assumed to have been the Marangoni effect. This slag could therefore not be investigated further.

Basicity changes to baseline CaO-based slag

The effect of changes in basicity, %CaO/%SiO\textsubscript{2}, on the P partition coefficient was also investigated. The basicities investigated were 0.7, 0.9, and 1.3 using baseline slag A with the MnO content constant.

Dephosphorization test procedure

All tests used the same test preparation method, which entailed weighing the required amounts of the slag and alloy, followed by homogenization of the feed charges in a ring mill for about 30 seconds. About 10 g of both the starting alloy and slag was used for the experiments. The charge was then packed into the graphite crucible. Packing was done such that a dense, compact solid charge was formed to promote the interaction of solid particles in the crucible. Graphite crucibles were used to avoid attack of the crucible by the different slag systems. The crucibles were also selected with the intention of maintaining the slag chemistries during the tests without interference by the crucible.
The charged crucibles were weighed and placed in the furnace chamber. The furnace chamber was covered with high-temperature refractory blankets (Fiberfrax) to prevent air ingress and preserve heat in the chamber. An alumina tube piped argon gas into the furnace chamber to create an inert environment for the duration of each test. An alumina sheath encased the Type B control thermocouple. The same thermocouple monitored the sample temperature.

The furnace was heated gradually by incrementally increasing the power input until the target test temperature was reached. The typical heating rate was 8–10°C/min under an inert atmosphere. The argon flow rate was maintained at around 2 L/min for the duration of the test. The samples were maintained at the target temperature for retention times of 30 minutes and 60 minutes. A lower retention time of 5 minutes was also investigated, but the analysis of the samples could not be completed due to poor separation of the slag and alloy at 1350°C. Upon completion of each test, the furnace power was switched off and the refractory blankets slightly moved to allow removal of the graphite crucibles from the furnace chamber. The hot crucibles were removed from the furnace using steel tongs and immediately quenched in a water-and-ice bath to ensure rapid cooling of the molten slag and alloy and maintain the phase chemistry in both the slag and alloy at the specific test conditions. The quenched crucibles were then dried at 105°C overnight in a drying oven. The dried sample was weighed and broken to retrieve the slag and alloy.

The alloy and slag were separated manually, weighed, and pulverized for analysis. The bulk chemical compositions of the feed materials and products were determined by Inductively coupled plasma-optical emission spectroscopy (ICP-OES). C and S were determined by combustion (LECO). P was determined using wet chemical analysis by ICP.

The calculations generally predict that dephosphorization of both alloys will not be possible with the CaO slags as the temperature is increased. The increase in the temperature was unexpected, as the dephosphorization reaction is exothermic (Chaudary and Goel, 1994; Simeonov and Sano, 1985). The addition of BaO in conjunction with CaO caused a detrimental effect, with lower predicted $L_p$ values obtained as indicated by slag H. Replacing both CaO and SiO$_2$ with BaO and BaF$_2$ flux in slag I resulted in higher $L_p$ values, above unity. The FactSage predictions concerning the BaO-BaF$_2$-MnO slag system agree with the observations made by various researchers (Fujita, et al., 1988; Chaudhary, Minj, and Goel, 2007; Dashevskii, et al., 1998). The effect of temperature on $L_p$...
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The results also show that at higher temperatures, generally lower $L_p$ values were obtained. Slag H, however, gave slightly higher values at 1450°C. This anomaly may be attributed to the analytical errors for samples from either the 1400°C or 1450°C experiments. Liu et al. (1995) similarly observed lower $L_p$ values as the temperature was increased during dephosphorization of FeMn alloy with BaO-containing slags.

Effect of changing MnO content of baseline slag

Figure 7 shows the $L_p$ values obtained at the three initial MnO slag contents at 1450°C, 30 minutes reaction time. Figure 8 represents the $L_p$ versus the final MnO slag contents, which increase in the final slags. The loss of Mn from the alloy is influenced by the oxygen partial pressure in the system. Factors that may have contributed to the loss of Mn include possible air ingress during the tests and the exposure of the alloys to oxygen during quenching, as Mn has a high affinity for oxygen. Liu et al. (1995) observed that the use of CaO, SiO$_2$, and CaF$_2$ to dephosphorize ferromanganese melts resulted in increased Mn losses to the slag. Watanabe et al. (1993) reported that MnO in basic slag acts as a diluent which may lessen the basic effect of the slag, reducing the capacity of the slag to remove P from the alloy. The presence of MnO in the slag is important to provide the required oxygen, although high MnO content can lead to low dephosphorization (Chaudhary and Roy, 2001).

The $L_p$ results show that there was an increase in $L_p$ from 15% to 20% MnO, followed by a reduction in the partition ratio. High amounts of MnO may affect the activity of the basic oxides such as CaO and BaO in the slag and can reduce the capacity of the slag to absorb P. Chaudhary, Minj, and Goel (2007) observed that an increase in the MnO content to above 20% (in BaO-containing slag) resulted in poor P removal. In the HCFeMn dephosphorization studies by Liu et al. (1995), it was observed times investigated. The results also show that at higher temperatures, generally lower $L_p$ values were obtained. Slag H, however, gave slightly higher values at 1450°C. This anomaly may be attributed to the analytical errors for samples from either the 1400°C or 1450°C experiments. Liu et al. (1995) similarly observed lower $L_p$ values as the temperature was increased during dephosphorization of FeMn alloy with BaO-containing slags.

Effect of changing MnO content of baseline slag

Figure 7 shows the $L_p$ values obtained at the three initial MnO slag contents at 1450°C, 30 minutes reaction time. Figure 8 represents the $L_p$ versus the final MnO slag contents, which increase in the final slags. The loss of Mn from the alloy is influenced by the oxygen partial pressure in the system. Factors that may have contributed to the loss of Mn include possible air ingress during the tests and the exposure of the alloys to oxygen during quenching, as Mn has a high affinity for oxygen. Liu et al. (1995) observed that the use of CaO, SiO$_2$, and CaF$_2$ to dephosphorize ferromanganese melts resulted in increased Mn losses to the slag. Watanabe et al. (1993) reported that MnO in basic slag acts as a diluent which may lessen the basic effect of the slag, reducing the capacity of the slag to remove P from the alloy. The presence of MnO in the slag is important to provide the required oxygen, although high MnO content can lead to low dephosphorization (Chaudhary and Roy, 2001).

The $L_p$ results show that there was an increase in $L_p$ from 15% to 20% MnO, followed by a reduction in the partition ratio. High amounts of MnO may affect the activity of the basic oxides such as CaO and BaO in the slag and can reduce the capacity of the slag to absorb P. Chaudhary, Minj, and Goel (2007) observed that an increase in the MnO content to above 20% (in BaO-containing slag) resulted in poor P removal. In the HCFeMn dephosphorization studies by Liu et al. (1995), it was observed...
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Figure 7—Effect of initial MnO content in the slag on \( L_p \) at 1450°C and 30 minutes retention time

Figure 8—\( L_p \) versus the final MnO in slags at 1450°C and 30 minutes retention time

that an increased concentration of MnO in slag had a negative influence on the phosphate capacity of BaO-MnO-BaCl\(_2\) slag. It was, therefore, concluded from the investigations that an MnO content above a certain value is detrimental to the capability of the slag to dephosphorize FeMn alloys. In the current study, the high MnO slag contents may have adversely affected the P capacity of the slags.

**Effect of adding \( \text{Na}_2\text{O} \) and \( \text{CaF}_2 \)**

\( \text{Na}_2\text{O} \) is a more basic oxide than CaO, so it was expected that dephosphorization would be improved by \( \text{Na}_2\text{O} \) additions. The results represented in Figures 5 and 6 generally show that the addition of \( \text{Na}_2\text{O} \), indicated by slag E, generally did not provide any added benefit as low \( L_p \) values were still obtained. The literature indicates that the addition of \( \text{Na}_2\text{O} \) to CaO-based slags improves the dephosphorization of liquid iron; however, \( \text{Na}_2\text{O} \) can be reduced by either C or Mn, producing CO/MnO. The instability of \( \text{Na}_2\text{O} \) can therefore have a detrimental effect on the dephosphorization of FeMn alloys due to loss of Mn (Tabuchi and Sano, 1985). In the current study, the loss of Mn from the alloy was observed even with the \( \text{Na}_2\text{O} \)-free slags, so it is not possible to establish the effect of \( \text{Na}_2\text{O} \) on Mn loss. Fujita et al. (1988) also observed that low degrees of dephosphorization were achieved when using \( \text{Na}_2\text{CO}_3 \) flux on FeMn alloy.

The fluoride ion (F\(^-\)) from \( \text{CaF}_2/\text{BaF}_2 \) is reported to stabilize the phosphate ion (PO\(_4^{3-}\)) during the ionic dephosphorization reaction in molten slag under oxidizing conditions, and thus contribute positively to dephosphorization (Liu et al., 1998). However, Ca\(^{2+}\) binds fluoride ions much more strongly than Ba\(^{2+}\) does, because of the difference in the ionic radius of the two cations. Therefore, fewer free fluoride ions would be available when using \( \text{CaF}_2 \) to enhance dephosphorization (Liu et al., 1998). Liu et al. observed that the addition of \( \text{CaF}_2 \) to MnO-BaO-BaF\(_2\) slag reduced \( L_p \) and increased the Mn capacity of the slag, i.e. Mn losses to the slag. In the current study, the addition of \( \text{CaF}_2 \) (slag F) showed no improvement in terms of \( L_p \).

**Effect of adding BaO**

The use of BaO in conjunction with CaO (slag H), also resulted in no improvement. Similar \( L_p \) values to those of slag A were obtained under conditions of 1350°C and 1450°C at 30 minutes. Slag A was, however, not outperformed by slag H under any of the other conditions. Nasaralla, Fruehan, and Min (1991) investigated the use of BaO as an additive in CaO-based flux for dephosphorizing Fe-C-P alloy. It was observed that amounts of BaO < 40% had no effect on the phosphate capacity of the slag. A larger increase in the phosphate capacity was only observed after increasing the BaO content of the CaO-CaF\(_2\) slag to above 40%. The researchers also later observed that the \( L_p \) increased significantly, by a factor of 4.6, when a high BaO content of 30% was utilized in the CaO-AI\(_2\)O\(_3\)-CaF\(_2\) slag system (Nasaralla and Fruehan, 1992). During the current study, a BaO content of 20% was utilized. The lower \( L_p \) values obtained could be attributed to the low concentration of the basic oxide and uncertainties in the analysis of P, as well as the dilution of the slag by MnO.

**Effect of slag basicity**

Increasing the slag basicity (%CaO/%SiO\(_2\)) to 1.3 caused the slag liquidus temperature to increase to about 1500°C. This led to difficulties in melting the slag and resulted in poor slag separation during the experiment at 1450°C, due to the alloy being entrained in the slag. This had a detrimental effect on the ability of the slag to remove P. As the reaction occurs in a liquid phase, a lower \( L_p \) was obtained by increasing the basicity to 1.3, as shown in Table VI. At the lower %CaO/%SiO\(_2\) ratio of 0.7, the P concentration in the slag was below the detection limit (<0.005%) and \( L_p \) could not be calculated. The results, therefore, show an initial increase in \( L_p \) value from a basicity of 0.7 to 1.0, followed by a slight reduction in \( L_p \).

**Effect of initial alloy composition**

Figures 9 and 10 show the \( L_p \) values obtained with HCFeMn at 1350°C and 1400°C respectively. Dephosphorization of HCFeMn by the respective slags was also not significant, as lower \( L_p \) values of <1 were generally obtained.

Comparison between the results shows that at 1350°C, relatively higher \( L_p \) values were obtained from MCFeMn, and lower values at 1400°C. As noted earlier, both the HCFeMn and the MCFeMn alloy would have reached carbon saturation during the experiments. The studies on the effect of initial C, Mn, and Si content on dephosphorization have shown that a high Mn in the alloy decreases the activity coefficient of P, and \( L_p \) increases.

<table>
<thead>
<tr>
<th>Table VI</th>
<th>Effect of %CaO/%SiO(_2) ratio on ( L_p ) at 1450°C and 30 minutes retention time</th>
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</thead>
<tbody>
<tr>
<td>%CaO/%SiO(_2)</td>
<td>( L_p )</td>
</tr>
<tr>
<td>0.7</td>
<td>1.3 0.13</td>
</tr>
<tr>
<td>0.90.26</td>
<td>1.30.13</td>
</tr>
</tbody>
</table>

- P slag analysis <0.005%
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![Graph showing phosphorus distribution](image1)

**Figure 9—Comparison of phosphorus distribution between MCFeMn and HCFeMn slag at 1350°C, 60 minutes reaction time**

![Graph showing phosphorus distribution](image2)

**Figure 10—Comparison of phosphorus distribution between MCFeMn and HCFeMn slag at 1400°C, 30 minutes reaction time**

| Table VII
<table>
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<tr>
<td><strong>Comparison between the results obtained in the current study and in the literature</strong></td>
</tr>
<tr>
<td><strong>Variables</strong></td>
</tr>
<tr>
<td>MnO content of baseline slag</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Na$_2$O and CaF$_2$</td>
</tr>
<tr>
<td>BaO additions</td>
</tr>
<tr>
<td>Slag basicity</td>
</tr>
<tr>
<td>Initial alloy composition</td>
</tr>
</tbody>
</table>

The results generally showed that $L_p$ remained small (<1): a significant proportion of P remained in the alloy, and thus dephosphorization was not favoured. The CaO-SiO$_2$-MnO (slag A) slag system yielded higher values of $L_p$. The addition of Na$_2$O (slag F) generally did not show any added benefit. Substituting half of the CaO by BaO (slag H) resulted in similar $L_p$ values to those of slag A under conditions of 1350°C and 1450°C at 30 minutes. Slag H did not outperform slag A under any of the other conditions. Increasing the temperature generally resulted in lower $L_p$ values. This may be attributed to the exothermic nature of the reaction which should be favourable at lower temperatures. Increasing the %CaO%/SiO$_2$ ratio of the starting slag gave an initial increase in $L_p$ value from a basicity of 0.7 to 1.0, followed by a slight reduction in the partition ratio. The latter results were not expected, as a higher basicity is expected.

with increasing initial C content until the C saturation point. High initial Si (>0.2%) is not favourable for dephosphorization as the Si is oxidized during dephosphorization, leading to increased SiO$_2$ in the slag, which consumes the high basic oxides and ultimately reduces the capacity of the slag to remove P (Chaudhary, Goel, and Minz, 2008; Bhardwaj, 2014). During the current study, the molten MCFeMn absorbed C from the crucible, forming HCFeMn, and the final average compositions of the alloys appeared to be similar after dephosphorization. The initial Si in both alloys was similar at about 0.4%, which is reported to be undesirable for dephosphorization of FeMn alloys.

**Conclusion**

The FactSage calculations indicated that none of the CaO-based slags would result in significant dephosphorization.
to improve the P capacity of the slag. Increasing the basicity increased the slag liquidus temperature, which negatively affected dephosphorization.

In summary, based on the $L_p$ values obtained, the conditions investigated are unfavourable for the removal of P from South African MCFeMn and HCFeMn industrial alloys, as significant amounts of P remained in the alloy. The experimental results are in line with FactSage predictions that the CaO slags are not suitable for dephosphorization of FeMn alloys.

Below is a summary of the shortcomings of the current study and recommended future work (Table VIII).

Due to the unsuccessful removal of P from the FeMn alloys by the CaO-based slag systems, it is recommended that further investigations be considered on the use of BaO-based slags without BaO addition. BaO slags are reported in the literature to be effective for dephosphorization of FeMn alloys. However, during the current study experimental challenges were encountered with the BaO-based slag. Further test work should be conducted with highly pure slag reagents that do not introduce P as an impurity in the master slags. It is also recommended that various other slags be investigated, with which higher basicity can be achieved at lower temperatures.

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References


