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### Thermodynamic analysis of zinc ferrite (ZnFe<sub>2</sub>O<sub>4</sub>) formation inside the HIsarna off-gas system

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#### ABSTRACT

HIsarna reactor is characterized by a high raw materials versatility and is therefore attractive for processing secondary iron sources. Among the materials that can be recycled through HIsarna, zinc-bearing material has drawn a special attention. Based on the plant data, once dust-containing Zinc was injected into the main reactor, a final collected dust with a zinc content of 16% was achieved which opened up possibilities of higher enrichment for direct reuse in Zn smelting as a secondary source and an alternative for Zn ore (the primary source). However Zn vapor can react with iron oxide to form zinc ferrite (ZnFe<sub>2</sub>O<sub>4</sub>), which is an undesired product. Hence, the main focus of this study is to minimize the formation of ZnFe<sub>2</sub>O using thermodynamic (FactSage) and computational fluid dynamic tools. After detecting regions with high potential of ZnFe<sub>2</sub>O<sub>4</sub> formation, proper geometrical and operational modifications of the off-gas system is proposed to minimize the formation of zinc ferrite.

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Thermodynamic analysis; equilibrium graph; HIsarna; CO-H<sub>2</sub> mixture combustion; zinc ferrite formation; dog leg; frog leg

#### Nomenclature

- absorption coefficient а
- С coefficients are constant
- Ε total energy [J kg<sup>-1</sup>]
- $\vec{F}_D$ drag force [N]
- g G gravitational acceleration [m s<sup>-2</sup>]
- production terms
- L spectral radiation intensity
- $\overline{J_i}$ diffusion flux of species
- effective conductivity of fluid [W m<sup>-1</sup> K<sup>-1</sup>] k<sub>eff</sub>
- k turbulent kinetic energy  $[m^{-2} s^{-2}]$ ,
- particle mass [kg]  $m_p$
- n spectral index of refraction of the medium
- р pressure [Pa]
- ŕ position vector [m]
- Ri net rate of chemical reaction
- ŝ direction vector
- Sh heat of chemical reaction
- S source term
- Т temperature
- t time
- $\vec{u_p}$ fluid phase velocity [m s<sup>-1</sup>]
- йp particle velocity [m s<sup>-1</sup>]
- u' fluctuating velocity [m s<sup>-1</sup>]
- ū mean velocity
- destruction terms

#### **Greek letters**

- Stefan-Boltzmann constant σ
- $\sigma_{s}$ scattering coefficient
- $\Omega'$ solid angle
- Г effective diffusivities [kg m<sup>-1</sup> s<sup>-1</sup>]
- dissipation frequency [1 s<sup>-1</sup>] З
- turbulent viscosity [m<sup>-2</sup> s<sup>-1</sup>]  $\mu_{t}$
- molecular viscosity [kg m<sup>-1</sup> s<sup>-1</sup>] μ
- density of fluid [kg m<sup>-3</sup>] ρ
- density of the particle [kg m<sup>-3</sup>]  $\rho_p$

particle relaxation time  $\tau_r$ 

- turbulent Prandtl number for kinetic energy = 1  $\sigma_k$
- turbulent Prandtl number for energy dissipation = 1.2  $\sigma_{\varepsilon}$

#### Introduction

Zinc (Zn) industry demands Zn ore at an acceptable price, but is also very interested in secondary resources as well to offset the need for prime ores. The availability of prime Zn ore and concentrates is declining and therefore driving up prices. Thus, there is an increasing interest in suitable secondary resources of Zn concentrates. One feasible way is to extract Zn from by-products and waste streams and reusing it for Zn production. This can contribute to reducing the need for landfilling, or even opens the possibility to process the contents of existing landfills and tailings, and ultimately lessening their environmental impacts and contributing to circular economy.

The steel industry produces large volumes of Zn-coated (galvanized) products which can be easily recycled via the electric arc furnace (EAF) route. In this route, scrap is charged into the furnace and is heated by means of an electric arc. The charged materials are melted to form slag and metal layers while Zn is vaporized and along with slag particles forming a fume (also known as dust) that escapes from the top space [1]. The EAF dust is rich in Zn, which is generally in the order of 7% to 40%, depending on the zinc content of the utilized scrap [2-6]. Havlik et al. [7] have reported an average of 22.14% Zn content based on a review of 19 different experimental resources with minimum and maximum of 13% and 39%, respectively.

During the EAF process, the Zn element can speciate into either zinc oxide (ZnO) or zinc ferrite (ZnFe<sub>2</sub>O<sub>4</sub>) compound. If

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the amount of ZnO in the dust is high enough, then it can be directly used in the Zn smelting process to produce pure metallic Zn. However, the presence of ZnFe<sub>2</sub>O<sub>4</sub> and also iron oxide compounds (on average 31% [7]) in the dust does require additional processing steps (pyro and hydrometallurgical routes) to enrich ZnO content before it is suitable for smelting application [1,2,8]. The most common routes are pyrometallurgical routes such as the Waelz kiln process which are characterized by high Zn recovery efficiency and also high energy consumption and pollutant emission. On the other hand, hydrometallurgical routes are less energy intensive and more eco-friendly, but complete recovery of Zn has proved to be very difficult due to the stability of ZnFe<sub>2</sub>O<sub>4</sub> [7,9]. If Zn content in the initial dust is mainly in the form of ZnO, high recovery efficiency is to be expected for any recovery routes [9].

The HIsarna process is a new concept based on the smelting reduction technology for producing liquid carbonated iron directly from iron ore and coal. Compared to the blast furnace route, coking and iron ore agglomeration (sintering and pelletizing) processes are eliminated, which inherently leads to at least 20% reduction in  $CO_2$  emission. Further reduction up to 80% can be achieved by incorporating carbon capture and storage technologies.

Based on the plant data and analysis which will be discussed in the coming sections, the process is characterized by high raw material versatility and is therefore attractive for processing secondary iron sources similar to the EAF process. Among the materials that can be recycled through HIsarna, galvanized steel scrap, basic oxygen furnace dust and zinc-bearing briquettes worth mentioning.

Based on the laboratory analysis of the collected dust in upstream, considerable amount of Zn in the form of ZnFe<sub>2</sub>O<sub>4</sub> and ZnO was detected. This finding led HIsarna technologists and researchers to believe that by injecting zincbearing material into the process, the Zn content is evaporated and concentrated in the off-gas dust (similar to the EAF process). However, it is desired to obtain a dust with the minimum content of ZnFe<sub>2</sub>O<sub>4</sub> (rich in ZnO) from which it is more convenient to recover Zn element.

In this study, the HIsarna process is described and then using thermodynamic (FactSage) and computational fluid dynamic (CFD) tools, the regions with a high risk of ferrite formation are detected inside the main reactor and off-gas system. According to findings, with current operation and configuration, the formation of ZnFe<sub>2</sub>O<sub>4</sub> is inevitable and can only be minimized through proper geometrical and operational modifications. The study is mainly based on thermodynamic equilibrium calculations and the effect of kinetics is not considered. Since the current system is a mixture of complex surface and volumetric reactions, thermodynamic calculations will be useful to have qualitative analysis on the behaviour of Zn vapour. For detailed quantitative analysis, incorporating kinetics is required which is the topic of another future study by the same authors.

#### HIsarna process

A pilot scale process flowsheet is shown in Figure 1(A). The process is currently at the development stage at ljmuiden site of Tata Steel in the Netherlands and since 2010 is fully operating with the capacity of 8 tones per hour of hot

metal production. The reactor is divided into two different regions namely cyclone converter furnace (CCF) and smelting reduction vessel (SRV) as depicted in Figure 1(B).

Fine iron ore and oxygen are fed into the CCF through sets of injectors. The iron ore particles are pre-reduced and melted during the fly time inside the CCF and are flung against the wall of the furnace to create a thin liquid film which drips and falls into the molten bath of SRV. The necessary heat for melting and pre-reducing fine ore is coming from partial combustion of the flue gas (containing CO–H<sub>2</sub> mixture) coming from SRV, with injected oxygen.

Inside the SRV, pulverized coal is injected into the molten bath through two sets of coal injectors which partially penetrate into the metal bath to carburize the molten iron. Coal particles reduce the pre-reduced iron oxide (FeO<sub>x</sub>) droplets fall into the SRV bath from CCF. Carbon monoxide (CO) and Hydrogen (H<sub>2</sub>) are produced during the reduction processes in the form of bubbles that are ultimately released from the bath to enter the top space of SRV. Pure oxygen is injected through two sets of injectors and partially combusts the gas mixture to provide the necessary heat inside the SRV.

Above CCF, the off-gas system is placed, which consists of four parts namely 'Reflux Chamber', 'Air Quench', 'Up Leg' and 'Down Leg' as depicted in Figure 1(C).

The reflux chamber is a slightly angled horizontal pipe with two bends to treat the flue gas coming from CCF. This configuration of the reflux chamber is also called 'Dog Leg'. It operates at high temperatures to fulfil two important tasks: post-combustion of the remaining CO, H<sub>2</sub> and carbon particles from CCF via oxygen injection and also capturing fine molten pre-reduced ore particles escaping the CCF using the described bent geometry. Above the chamber, there is an air quench system, which consists of square channels to inject air to serve two main purposes: cooling the flue gas to reach an appropriate temperature for flue gas cleaning and freezing the molten pre-reduced iron ore particles (escaped from the CCF and reflux chamber) to avoid molten particle accretion on the walls. Further cooling is achieved via nitrogen and water spray injection (evaporative cooling) in the down leg and ultimately flue gas enters the gas cooler (point D) to reach a proper temperature for the bag house and sulphur removal unit.

#### Zinc recovery from the HIsarna process

Zinc-bearing material can be fed into the process from different sections. For example, materials in the dust form are injected through nozzles into the CCF. On the other hand, granular materials are fed into the SRV through an inclined chute as illustrated in Figure 1(B). The zinc content of the injected material is rapidly evaporated, travels upward and ultimately will end up in the off-gas dust collected by cyclone and bag houses.

During the fourth HIsarna campaign in 2014, a test was performed where dust from the BOS (basic oxygen steelmaking) plant gas cleaning system was injected into the CCF together with iron ore. The dust was mainly made up of iron oxides, but also contained up to 5 wt% Zn. During the tests, it was demonstrated that it is possible to enrich Zn up to 16% into the HIsarna dust, as shown Figure 2.

However, during this test, it was only possible to inject a small amount of this dust for a short period of time. This meant that the achieved Zn enrichment was only limited,



Figure 1. (A) Schematic representation of HIsarna main components; (B) main reactor components; (C) off-gas system with plant measurement points (point A: reflux chamber outlet, Point B: end of up leg, Point C: 3 m above water quench atomizers, Point D: exit to gas cooler).

but it offered the prospect to achieve much higher levels of Zn in the Hlsarna off-gas dust without creating a zinc cycle (like in a blast furnace) and detrimental zinc accumulation within the process.

#### Zinc oxide and zinc ferrite formation

In the HIsarna process, similar to EAF, the Zn vapour content in the flue gas can end up into two main zinc-bearing compounds. It could react directly with oxidants such as  $O_2$ ,  $CO_2$  or  $H_2O$  to form ZnO or react with oxygen and prereduced ore (mainly Fe<sub>3</sub>O<sub>4</sub>) to form ZnFe<sub>2</sub>O<sub>4</sub> [10–13]. In this context, ZnO is a favourable by-product since Zn recovery from ZnFe<sub>2</sub>O<sub>4</sub>, due to its high stability, requires further treatment [2]. There have been numerous studies on the  $ZnFe_2O_4$  and ZnO formation in iron making industry especially during the EAF process.

Pickles [14,15] have investigated the effect of different parameters on the formation of zinc–manganese ferrite in EAF dust using thermodynamic analysis for the Zn–Fe– Mn–O–H–C system. He has reported that gas composition, mainly high ratio of CO/CO<sub>2</sub> and H<sub>2</sub>/H<sub>2</sub>O has a significant impact on limiting the formation of both ZnO and ZnFe<sub>2</sub>O<sub>4</sub>. Temperature reduction is reported to be a favourable condition for the formation of ferrite. Increasing the amount of Zn and manganese content in the gas can reduce the formation of ferrite, however, an increase in the ferrite formation is reported at high concentration of iron oxide.



Figure 2. Result of Zn enrichment of the HIsarna flue dust after injection of Zn-containing dust into CCF (data taken from HIsarna plant).

Sureerat et al. [16] have investigated the reduction properties of reducing gases (H<sub>2</sub>-CO mixture) to eliminate and decompose the  $ZnFe_2O_4$  compound in EAF off-gas over a temperature range of 400–1600°C. Both experimental and thermodynamic analyses were performed. They have reported that ferrite reduction occurs at relatively high temperature and strongly depends on the reducing gas composition. For example,  $ZnFe_2O_4$  reduction by hydrogen is feasible at a temperature above 1100°C but temperatures above 1500°C are required for reduction by carbon monoxide.

Suetens et al. [10] have performed a detailed experimental study on the formation of  $ZnFe_2O_4$  in EAF dust to investigate the possibility of in-process separation of Zn from the dust. They have concluded that the  $ZnFe_2O_4$  formation is thermodynamically favourable in oxygen-rich atmosphere and low temperature same as previously mentioned studies.

Nedar [13] has proposed a gas–solid reaction mechanism for the ferrite formation where the oxygen is reacting with zinc vapour in the presence of liquid iron. The liquid iron is oxidized to form FeO and Fe<sub>3</sub>O<sub>4</sub> and depending on the oxidized form, two different routes are possible. Their proposed mechanism is confirmed by experimental analysis and again points out the importance of oxygen and temperature for the ZnFe<sub>2</sub>O<sub>4</sub> formation. The same mechanism has been investigated by other researchers [10–12].

Learning from the mentioned researches, one can find many similarities between the dust formation in the EAF process and occurring phenomena inside the HIsarna reactor when zinc-bearing material is injected. As will be discussed in coming sections, the off-gas system provides the most favourable environment for the  $ZnFe_2O_4$  formation. This is due to a high concentration of molten pre-reduced iron oxide particles (Fe<sub>3</sub>O<sub>4</sub>), low temperature and high partial pressure of oxygen (or equivalently low partial pressure of carbon monoxide) [10–17]. From the literatures and for the condition inside the HIsarna process, there are few main reactions for  $ZnFe_2O_4$  and ZnO formation and reduction as listed below:

$$Zn + CO_2 \rightarrow ZnO(s) + CO$$
 (1)

$$Zn + H_2O \rightarrow ZnO(s) + H_2$$
 (2)

$$Zn + O_2 \rightarrow ZnO(s)$$
 (3)

$$Zn + Fe_3O_4(s) + O_2 \rightarrow ZnFe_2O_4(s)$$
 (4)

$$ZnO(s) + CO \rightarrow Zn + CO_2$$
 (5)

$$ZnFe_2O_4(s) + CO \rightarrow ZnO(s) + 2FeO(s) + CO_2$$
 (6)

$$3 ZnFe_2O_4(s) + CO \rightarrow 3 ZnO(s) + 2 Fe_3O_4(s) + CO_2$$
 (7)

$$ZnO(s) + Fe_2O_3(s) \rightarrow ZnFe_2O_4(s)$$
(8)

$$Zn + CO_2 + Fe_2O_3(s) \rightarrow ZnFe_2O_4(s) + CO$$
(9)

Figure 3 shows Gibbs energy change versus temperature for reactions (1) to (9).

Reaction (4) which is main in the ZnFe<sub>2</sub>O<sub>4</sub> forming reaction [8,9] is highly stable (negative  $\Delta G^{\circ}$ ) for the defined temperature range, especially at lower temperature. From the same analysis, it can also be seen that the ZnFe<sub>2</sub>O<sub>4</sub> formation reactions are more stable than both ZnFe<sub>2</sub>O<sub>4</sub> reduction and ZnO formation which makes the prevention of the ZnFe<sub>2</sub>O<sub>4</sub> formation quite challenging in the HIsarna process.

#### **Tools and methods**

In order to have a detailed analysis on the formation of  $ZnFe_2O_4$  inside the off-gas system, profiles of temperature, gas composition and molten pre-reduced ore flow rate inside the reflux chamber and the rest of the off-gas system are needed. The temperature values are only available and measured at points A, B and D and the flue gas compositions



Figure 3. Gibbs energy change versus temperature for possible reactions involving Zn.

are only measured at points A and D as shown in Figure 1(C). The molten particle flow rate measurements are quite complex and almost impossible due to a very high temperature and liquid state of particles inside the reflux chamber. These limited measurement points do not provide enough information to have a thorough analysis at every point of the off-gas system.

CFD is a very useful tool to predict the behaviour of the flow in such systems to obtain necessary information regarding gaseous composition, temperature, particle flow rate, etc. A CFD model of the HIsarna off-gas system is setup and validated using plant data in another study by the same authors [18]. Using the developed model, it is possible to obtain all of the profiles, and measure all necessary parameters of the system at any desired points. FactSage thermo-chemical software [19] is used to plot equilibrium diagrams by varying temperature and partial pressure of oxygen to calculate all formed phases. Using the calculated temperature and composition profiles from the validated CFD model, discrete operating points are mapped on the equilibrium graphs to see possible formed phases and compounds inside the off-gas system.

#### **CFD** calculation

Detailed discussion on the CFD model setup, mesh, boundary conditions, grid independency and governing equations can be found in the study of Hosseini et al. [18]. In this study, a detailed discussion on the CFD model development is omitted for brevity, however, a quick overview and summary is presented in the following.

Table 1 shows the composition and operating condition of the inlet flue gas entering the reflux chamber. These data are obtained at normal plant operation where there is no zincbearing material injection into the process. The inlet flue gas contains  $CO-H_2$  mixture, carbon particles and molten pre-reduced ore particles which mainly contain Fe<sub>3</sub>O<sub>4</sub>.

 $CO-H_2$  and carbon particles are burnt by injecting oxygen inside the reflux chamber and slag particles are captured using the reflux chamber bent forming a flowing liquid layer which runs back into the CCF.

Iron ore is injected with 8 tons  $h^{-1}$  (2.223 kg s<sup>-1</sup>) into the CCF at the current pilot plant and according to mass balance calculations, on average 10% of the injected iron ore escapes CCF and enters the reflux chamber (0.223 kg s<sup>-1</sup>). Particle density is considered to be 5000 kg m<sup>-3</sup> with a heat capacity of 922 j kg<sup>-1</sup> K<sup>-1</sup> which are properties of molten Fe<sub>3</sub>O<sub>4</sub> that form majority of the particles. Rosin–Rammler distribution is used to set particle

size distribution for the particles with fixed minimum particle size, maximum particle size, mean diameter, spread parameter and number of intervals as 30, 1870, 166  $\mu$ m, 0.805 and 22, respectively.

To include chemistry/turbulence interaction and calculating reaction rates, the eddy dissipation concept model [20] is considered. The detailed kinetic mechanism proposed by Cuoci et al. [21] is used for  $CO-H_2$  mixture combustion.

For carbon particle combustion, field char oxidation model is used which is a simplification of unreacted shrinking core modelling. Kinetic data and expressions proposed by Wen et al. [22] are used for carbon combustion. Water droplet evaporation is also included using the convection/diffusion controlled sub-model.

Particles behaviour (molten ore and carbon) are modelled using the Lagrangian discrete phase method. The trajectory of a discrete phase particles is predicted by integrating the force balance on the particles, which is written in a Lagrangian reference frame. The dispersion of particles due to turbulence in the fluid phase is included using the stochastic tracking model (random walk).

Reflux chamber walls are made of steel tubes and inner side is covered with a refractory of 4 cm. Above the reflux chamber, the walls are only made of steel tubes (OD: 0.038, thickness: 0.005 m). Water flows through the pipes and cools the wall in counter-current flow. All walls are modelled according to the shell conduction approach to consider different material layers. It is assumed that molten particles are at the liquid phase and trapped once they hit the reflux chamber walls (trapped wall boundary condition), but reflected anywhere above the reflux chamber section as they tend to solidify.

#### CFD results

Figure 4(A) shows the temperature and composition profile along the off-gas system (shown in Figure 1(C)). The length axis refers to the length of a line passing through the middle of the off-gas geometry. The calculated temperature and compositions are averaged on a cross-section sweeping along the length axis. As can be seen, the model predictions (solid lines) are in good agreement with plant measured values (symbols).

Figure 4(B) illustrates the calculated flow rate of the ore particles across the reflux chamber. The particles are captured by the walls and the flow rate is reduced and after point 8 (reflux chamber outlet) the flue gas and remaining particles

 Table 1. Inlet boundary conditions for flue gas, oxygen port and air quench.

	Reflux chamber	Air quench	Oxygen port	Nitrogen ports	Water Spray
Inlet temperature [°C]	1813 (2086 K)	20 (293 K)	20 (293 K)	20 (293 K)	20 (293 K)
Inlet mass flow rate [kg s <sup>-1</sup> ]	4.33	3.69	0.27	0.205	0.45
Inlet Carbon Particles flow rate [kg s <sup>-1</sup> ]					
diameter = 120 $\mu$	0.0282	_	-	-	-
Inlet molten pre-reduced ore particle [kg s <sup>-1</sup> ]	0.223	-	-	-	-
Composition – average mole fraction at inlet					
CO	0.0261	0	0	0	0
CO <sub>2</sub>	0.61	0.0003	0	0	0
H <sub>2</sub>	0.002	0	0	0	0
02	0	0.21	0.995	0	0
N <sub>2</sub>	0.166	0.78	0.005	1	0
$H_2^{-}O$	0.2	0.012	0	0	1
Post combustion ratio	96.63%	-	-	-	-



Figure 4. (A) Calculated gaseous composition and temperature profile and (B) the flow rate of particles across the reflux chamber (dog leg geometry) along the off-gas system.

enter the air quench section where particles are frozen and no more capturing by walls occurs.

The model for molten particles behaviour is at the initial stage of development and further improvement to consider the effect of particle temperature and state will be presented in another study. However, the predicted outlet flow rate of ore particles is in good agreement with the measured dust flow rate of 0.089 kg s<sup>-1</sup> in cyclone and bag houses of the pilot plant. Since the molten particles mainly contain Fe<sub>3</sub>O<sub>4</sub>, the reduction in the total particle flow rate means lesser available ingredients for the ZnFe<sub>2</sub>O<sub>4</sub> formation and a potential reduction of ZnFe<sub>2</sub>O<sub>4</sub> content in the collected dust.

#### FactSage calculation

The atmosphere inside the HIsarna reactor is a reducing environment with high CO and  $H_2$  content as shown in Figure 5(A). The mentioned ratio in the figure is from direct measurements and back calculations using the process flowsheet software. As discussed before, the presence of these reducing gases and also high flue gas temperature can significantly limit the formation of both ZnO and ZnFe<sub>2</sub>O<sub>4</sub> [10,14,15].

Figure 5(B) shows the effect of temperature on the formation of  $ZnFe_2O_4$  in a mixture of CO, CO<sub>2</sub>, H<sub>2</sub>O, O<sub>2</sub>, Zn and Fe<sub>3</sub>O<sub>4</sub>(s) for different CO/CO<sub>2</sub> ratios. The ratios of other gaseous and solid compounds are fixed with respect to CO<sub>2</sub>. Any Zn content ending up in the spinel phase is in the form of ZnFe<sub>2</sub>O<sub>4</sub> and as can be seen, increasing the ratio of CO/CO<sub>2</sub> can considerably reduce the formation of ZnFe<sub>2</sub>O<sub>4</sub>, especially at higher temperature. According to this thermodynamic analysis, the formation of ZnFe<sub>2</sub>O<sub>4</sub> at temperature and CO/CO<sub>2</sub> range of SRV and CCF is thermodynamically unfavourable. However, at the lower ratio of CO/ CO<sub>2</sub> which occurs inside the off-gas system, the formation of ferrite at any temperature range is highly appealing.

Furthermore, temperature inside the reflux chamber is quite high, but oxygen injection decreases the local temperature and increases the oxygen partial pressure. The formation of  $ZnFe_2O_4$  becomes even more favourable with further temperature reduction and oxygen injection above the reflux chamber and in the air quench section.

Altogether the current operating condition of the off-gas system makes this part of the process the region of interest for this study as it exhibits higher potential of the  $ZnFe_2O_4$  formation compared to the rest of the HIsarna reactor. From Figure 5(B), it can also be concluded that most of the Zn content will stay if the form of vapour at the inlet of the off-gas system.

'Phase Diagram Module' of FactSage software is used to plot the equilibrium graph (oxidation graph) using the listed compounds in Table 2. Figure 6 shows an example of a plotted graph.



Figure 5. (A) Gaseous composition and temperature at different outlet sections and (B) total zinc prediction in the spinel phase.

 Table 2. Components used to plot equilibrium graphs in FactSage software.

Component	Mole fraction/Partial pressure (atm)
CO <sub>2</sub>	0.547
H <sub>2</sub> O	0.173
N <sub>2</sub>	0.158
02	Variable from 0 to 0.12
Zn	Variable – arbitrary values
Fe <sub>3</sub> O <sub>4</sub>	Variable – arbitrary values

with a maximum value appearing above the air quench.

Equilibrium graphs are based on the temperature and gas', 'L  $log_{10}^{P_{O_2}}$  as y and x axes, respectively. Temperature is varied from 500 to 2000°C and  $P_{O_2}$  is varied from 0 to 0.12 atm is form

For all calculations, the molar ratio of Zn vapour to iron oxide  $\left(\frac{Zn_{(g)}}{Fe_3O_4 + Zn_{(g)}}\right)$  is varied from 0.01 to 0.9 and is fixed for each plot. For simplicity, this ratio will be called '*R*' from now on. The variation of '*R*' ratio depends on the zinc content of injected zinc-bearing material and also the amount of zinc vapour reaching the off-gas system inlet.

The equilibrium graphs are divided into different regions separated by solid lines. Each region might contain one or a combination of phases. Possible formed phases are 'Ideal gas', 'Liquid slag' (slag-liq), 'Spinel', 'Monoxide' (slag in solid form) and 'Fe<sub>2</sub>O<sub>3</sub>'. Another possible phase is 'Zincite' which is formed at a very high '*R*' ratio (when zinc vapour content is high in flue gas) as shown in later figures.



**Figure 6.** Equilibrium graph with mapped operating points for R = 0.01.

Table 3. Possible phases and components inside each phase.

Phase	Possible component
Slag-liq Iron o	xide (major FeO, Fe <sub>2</sub> O <sub>3</sub> ) and ZnO (minor)
Spinel Fe <sub>3</sub> O <sub>4</sub>	and ZnFe <sub>2</sub> O <sub>4</sub>
Ideal Gas CO, CO	$D_2$ , Zn, $O_2$ , $H_2O$ , $H_2$ etc.
Zincite ZnO (i	major), iron oxide (minor)
Fe <sub>2</sub> O <sub>3</sub> Iron o	xide

Each phase contains certain compounds with calculated values. Table 3 summarizes the compounds of interest for this study in each phase. FactSage software allows to evaluate the composition at each point of the equilibrium graph which is not presented in this study for brevity.

Figure 6 also contains operating points (red dots) for different sections of the off-gas system. Those operating

points are discrete points along the reflux chamber and up leg (up to point B in Figure 1(C)) with calculated temperature and partial pressure from the discussed validated CFD model. For instance, point \* refers to the reflux chamber inlet with temperature of 1812°C and almost zero oxygen partial pressure ( $P_{O_2} \approx 10^{-4}$ . atm). The same is applied to all of the plotted operating points.

Using the equilibrium graphs with mapped operating pints, it is possible to observe where those operating points fall and which phases might form in the presence of a certain ratio of zinc vapour and iron oxide. For example in Figure 6, the operating points in the first section of the reflux chamber (before oxygen injection), fall into the 'Slagliq + Ideal gas' region where zinc elements speciate into the slag in the form of ZnO or remains in gaseous form due to



Figure 7. Equilibrium graph with mapped operating points for 'dog leg' geometry and for different 'R' values.



Figure 8. Laboratory analysis of HIsarna baghouse dust for Zincite (ZnO) and Zinc ferrite (ZnFe<sub>2</sub>O<sub>4</sub>) fraction.

high temperature and absence or existence of minor oxygen. In the second section, the partial pressure of oxygen is sharply increased and temperature drops locally. This is a favourable condition for the  $ZnFe_2O_4$  formation and as it can be seen most of the operating points fall into the 'Spinel' phase where the major component is  $Fe_3O_4$  and all of the zinc element speciate into  $ZnFe_2O_4$ . The same situation can be seen in the third section where the major phase is ' $Fe_2O_3$ ' and 'Spinel'.

#### **Results and discussion**

#### Current pilot scale - dog leg geometry

Figure 7 depicts equilibrium graph with mapped operating points for different '*R*' ratios. As can be seen, at the lowest investigated ratio where R = 0.01, the operating points before oxygen injection points fall in 'Slag-liq' and moved to 'Spinel' containing region after oxygen injection. As the ratio is increased (increasing zinc vapour content in flue gas), the

operating points in the first section fall into the 'Slag-lig + gas' region where most of Zn vapour remains in the gaseous phase. The 'Zincite' phase which contains almost pure ZnO content occurs around R = 0.4 to 0.5 and expands by increasing Zn to the iron oxide ratio. At the 'R' ratio of 0.8 and 0.9, most of the operating points fall into the 'Slag' and 'Slag + zincite' region where Zn element is in the form of ZnO. Nevertheless, at all ratios, the formation of spinel and ZnFe<sub>2</sub>O<sub>4</sub> is still inevitable especially in the second and third sections where local and global temperature drops fast and there is a sharp increase in oxygen partial pressure as shown in Figure 4(A). According to the calculations, increasing Zn vapour content in flue gas, even at low temperatures will promote the formation of ZnO which is in line with other researchers findings [2,11,12,14]. However, reaching such a high 'R' ratio is guite challenging. For example, during previous plant operations, 2 ton  $h^{-1}$  of scrap material containing 2%w zinc coating was injected into the SRV bath which accounts for 0.011 kg s<sup>-1</sup> of zinc vapour in total. If the whole vapour from the bath reaches the CCF outlet along with 0.223 kg  $s^{-1}$  of pre-reduced ore escaping CCF (which



Figure 9. Proposed modification to the reflux chamber, frog leg configuration.



Figure 10. (A) Predicted gaseous composition and temperature profile along frog leg and air quench (obtained from CFD calculations up to point C) and (B) predicted flow rate of particles across the frog leg geometry.

contains 85% of Fe<sub>3</sub>O<sub>4</sub>), the maximum ratio of R = 0.15 at inlet and 0.35 at outlet of the reflux chamber can be expected. At this ratio, and according to the calculated equilibrium graphs, most of the Zn element will be speciated into the ZnFe<sub>2</sub>O<sub>4</sub> component in the 'Spinel' phase. This is in agreement with laboratory analysis of collected dust in the baghouse (at the current pilot plant) shown in Figure 8. As can be seen, on average, around 2.5% of the total weight of the sampled dust contains zinc-bearing compounds and of that percentage, 92% is in the form of ZnFe<sub>2</sub>O<sub>4</sub> and only 8% of injected Zn ends up in the ZnO form. So it can be concluded that the ZnFe<sub>2</sub>O<sub>4</sub> formation is thermodynamically inevitable at the current operating condition and setup. Even in the absence of oxygen, prevention of ZnFe<sub>2</sub>O<sub>4</sub> formation is not a straightforward task, mainly due to high stability of the spinel at lower temperatures [10].

#### Reflux chamber geometrical modification

From previous discussion and calculations, the impact of temperature and oxygen partial pressure on the formation of ZnO and  $ZnFe_2O_4$  became evident. The main focus will be on geometrical and operational modification of the reflux chamber since, according to equilibrium graphs, this section of the off-gas system can still offer possibilities to minimize the formation of  $ZnFe_2O_4$ . To maximize the formation of ZnO-containing phases, high temperatures and lower partial pressures of oxygen (or high partial pressure of CO) are required inside the reflux chamber.

Considering a fixed condition at the inlet of the chamber, the following solutions are available to practice:

- a) One solution is to increase the 'R' ratio as discussed in study of Pickles [14,15] and based on the thermodynamic calculations in section 'Reflux chamber geometrical modification'. This can be achieved through increasing zinc content inside the off-gas system by injecting more zinc-bearing material into the process. With this solution it is possible to increase the ratio to a certain extent as injecting very high amount of zinc-bearing material is complex and might cause technical issues. For example, injecting galvanized scrap particles in large quantities into the SRV molten liquid could cause local liquid freezing which is detrimental to the steady operation of the plant.
- b) It is also possible to increase the '*R*' ratio by increasing particle capturing inside the reflux chamber to limit the availability of  $Fe_3O_{4r}$  especially in those regions with high oxygen partial pressure and low temperature.
- c) It is also possible to modify the geometry of the reflux chamber to provide larger region with high temperature and low partial pressure of  $O_2$  or keeping CO partial pressure as high as possible before the post-combustion (creating a reducing environment). The presence of CO in the flue gas will limit the formation reaction of  $ZnFe_2O_4$ .

To fulfil these goals, a modified reflux chamber geometry shown in Figure 9 is proposed. This geometry will be called 'Frog Leg' from now on and as can be seen, the second



Figure 11. Equilibrium graph with mapped operating points for 'frog leg' geometry and for different R values.

bend is added to the original geometry (Dog Leg) and the oxygen port is shifted upward to delay the post-combustion.

The same discussed CFD model is used to determine temperature profile, compositional change and particle flow rate along frog leg geometry and results are illustrated in Figure 10.

In dog leg geometry, the temperature falls drastically at the length of 10 m, whereas for frog leg, this drastic reduction happens at length of 18 m, providing extra 8 m length for the flue gas to stay at high temperature. Also from the same figure, it can be seen that oxygen injection and CO combustion are delayed for 8 m which is again beneficial to prevent the  $ZnFe_2O_4$  formation and promoting the ZnO formation.

Once ZnO is formed inside the slag, it will be stable and in the form of small distinguished particles. This is confirmed by study of Suetens et al. [10] that reported the presence of ZnO particles inside the collected dust from EAF via EDS mapping (Energy Dispersive Spectroscopy).

The only way to destabilize and reduce the formed ZnO is through gaseous reduction or solid state reaction. The solid state reaction is quite slow and highly controlled by kinetics



Figure 12. (A) Calculated temperature profile for the off-gas system with frog leg geometry for different wall thicknesses (up to the point C) and (B) equilibrium graph with mapped operating points for frog leg geometry.

thus is unlikely to happen within the available short resident time inside the off-gas system. The possibility of ZnO reduction by reducing gases are also limited, because for the gas-solid reaction, the gaseous compound must diffuse into the solid surface which is again unlikely as the partial pressure of the CO-H<sub>2</sub> mixture is very low inside the off-gas system.

It also worth mentioning that according to thermodynamic calculations (Gibbs free energy) and study of Sureerat et al. [16], the reduction of both ZnO and  $ZnFe_2O_4$  can proceed only at high temperature and highly reducing environment. Thus, a stable ZnO content in hot zones and also cooling path (above reflux chamber) can be expected.

The other benefit of this modification is that the content of molten pre-reduced particles is constantly decreased as depicted in Figure 10(B). In dog leg geometry, the flow rate of particles entering the oxygen injection area is around 0.182 kg s<sup>-1</sup> however, for frog leg geometry the flow near oxygen injection is 0.089 kg s<sup>-1</sup>. This means an increase in '*R*' ratio from 0.2 to 0.4 in oxygen-rich region which can limit the formation of  $ZnFe_2O_4$  inside the reflux chamber.

These conclusions can be quantified with the same FactSage analysis to investigate possible  $ZnFe_2O_4$  forming spots inside the frog leg. The results are reported in Figure 11 and as can be seen, compared to the dog leg geometry, higher number of operating points in the first section of the frog leg falls into the liquid 'Slag' phase. However, the formation of spinel is still inevitable due to

the temperature reduction in oxygen and air injection regions. The operating points shift upward by increasing the '*R*' ratio and the formation of spinel (and  $ZnFe_2O_4$ ) will be limited in the same way discussed for the dog leg geometry.

The geometrical modification can also positively contribute to the combustion and removal of other undesired compounds such as carbon particles and  $CO-H_2$  mixture inside the off-gas system which has been investigated in another study by the same authors [23].

In conclusion, based on the analysis so far, for any reflux chamber configuration (dog leg or frog leg), the most efficient way to increase ZnO content of the collected dust would be increasing the '*R*' ratio by injecting higher amount of zinc-bearing material. Steel scrap injection into the molten slag would be an interesting option as it's a very efficient way to recover the iron content of the scrap and to reduce the consumption of iron ore. However, when it comes to zinc recovery, it has shortcomings. Achieving the high '*R*' ratio requires a considerable amount of steel scrap injection which can reduce the slag temperature and might cause the disturbance in temperature control and local freezing.

So it is suggested to inject zinc-bearing material not only in the form of the steel scrap injection into the molten slag, but also through other forms and from other section of the process such as zinc-bearing dust injection into the CCF or SRV upper section. This way the Zn content along the off-



Figure 13. Effect of CO content on temperature, oxygen profile and operating point for dog leg and frog leg geometry and for lowest and highest 'R' ratios.

gas system is evenly distributed and local disturbances are recovered faster.

#### Reflux chamber wall modification

In order to make sure that operating points inside the first section of the frog leg geometry stay away from the 'Spinel' region for minimum and maximum 'R' ratio, a further modification might be required. Considering the modified geometry, the only feasible way is to keep the temperature as high as possible inside the chamber. This can be achieved by either increasing refractory wall thickness to reduce the heat losses through the walls or increasing CO content at the inlet of the off-gas system to provide more fuel for post combustion. The validated CFD model is used to investigate both solutions. Figure 12(A) shows the temperature profile for different refractory wall thicknesses of the reflux chamber. For a fixed equilibrium graph, the operating points shift upward by increasing the flue gas temperature as depicted in Figure 12(B) for maximum and minimum considering the 'R' ratio. This shift will locate the operating points of the first section in more favourable equilibrium region where zinc element remains either in gaseous form or speciate into the slag in form of ZnO. It worth noting that even at high temperature, the formation of 'Spinel' and ZnFe<sub>2</sub>O<sub>4</sub> is still inevitable once the oxygen becomes available (upper region of frog leg). So practically this geometrical modification could minimize the ZnFe<sub>2</sub>O<sub>4</sub> to a certain extent and may not fully prevent the undesired product formation.

#### Increasing CO content inside the reflux chamber

Due to the harsh condition inside the reflux chamber, the refractory wall is exposed to thermo-chemical stresses which can cause wall material loss and thickness reduction. To make the wall thickness solution work fine, it is required to refresh the refractory wall on a regular basis which might not be feasible during long runs. Therefore, more feasible solution would be increasing the CO content of the flue gas at the inlet of the reflux chamber by reducing the CCF post-combustion ratio. Since the HIsarna process has a transient nature, in real plant operation the amount of CO at the inlet of the off-gas system can reach up to 10% (molar fraction).

Figure 13 shows the effect of CO content on flue gas temperature and oxygen composition along with the operating point position on the equilibrium graph for both dog and frog leg geometry. As can be seen, by increasing CO content at the inlet, most of the operating point will be shifted upward and into regions where Zn vapour speciates into ZnO even in the second section, where post combustion oxygen is injected in considerable amount.

Keeping CO partial pressure high inside the reflux chamber offers several benefits. Higher CO content means higher fuel for the post -combustion process which in turn signifies higher heat release and temperature profile to limit the evolution of  $ZnFe_2O_4$  similar to the effect of wall thickness. As mentioned earlier, with higher temperature, the operating points on the equilibrium graph are shifted upward to the regions with favourable form of Zn element. Moreover, high CO content consumes oxygen across the reflux chamber and lowers its partial pressure to create a reducing environment where  $CO/CO_2$  ratio is high. The

higher is the CO/CO<sub>2</sub> ratio, the lower would be the risk of  $ZnFe_2O_4$  formation (see Figure 5).

#### Conclusion

A thermodynamic analysis of  $ZnFe_2O_4$  and ZnO formation inside the off-gas system of the HIsarna process is performed. The aim is to avoid or minimize the formation of  $ZnFe_2O_4$ compound as it is an undesired by-product of Zn vapour reaction with iron oxide.

Since limited measure points from the real plant data were available, a CFD model was developed and validated to describe the composition and temperature profile variation inside the off-gas system.

The analysis was performed using equilibrium graphs and mapped operating point obtained from CFD calculations. It was found that the most favourable conditions for the ZnFe<sub>2</sub>O<sub>4</sub> formation are high oxygen partial pressure, high content of pre-reduced iron ore (high Fe<sub>3</sub>O<sub>4</sub> content) and low temperature which are in agreement with other experimental and numerical studies.

These conditions are met at the current state of the plant operation and according to thermodynamic analysis, the offgas system have higher  $ZnFe_2O_4$  formation potential compared to SRV and CCF sections. Inside the SRV and CCF, the formation of  $ZnFe_2O_4$  is limited due to highly reducing environment. On the other hand, in the off-gas system, the environment is highly oxidizing owing to the injection of post combustion oxygen and air. Since the oxidizer are injected at a very low temperature (20–30°C), it will also cause a rapid temperature drop across the off-gas system which again highly contributes to the formation of  $ZnFe_2O_4$ .

Among the proposed solutions, the easiest way was to increase the zinc to iron oxide molar ratio ('*R*' ratio) which will shift the mapped operating points into the phasic regions with ZnO compound as the final product of Zn vapour reactions. However, achieving a high '*R*' ratio is challenging and the formation of  $ZnFe_2O_4$ , even at the highest '*R*' ratio, is still predicted inside the off-gas system.

For further reduction of  $ZnFe_2O_4$  formation, a geometrical modification of the reflux chamber is proposed by adding another turn to the current design. The same analysis was performed for modified geometry and it was shown that the area with low oxygen content and high temperature is expanded which provides more residence time for Zn element to speciate into the desired phases with ZnO content.

The proposed modification can lead to a higher capturing efficiency of molten pre-reduced particles by the walls. This inherently can reduce the  $Fe_3O_4$  content of the flue gas and increase the '*R*' ratio which limits the formation of  $ZnFe_2O_4$ .

Nevertheless, the proposed modifications can only minimize the formation of undesired products to a certain extent and not fully eliminate it. The formation of  $ZnFe_2O_4$ is still inevitable as some of the operating point, especially those points near the oxygen and air injection still fall into the spinel-containing zones where Zn vapour will speciate into the  $ZnFe_2O_4$  compound in reaction with oxygen and iron oxide at low temperature.

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