Dissolution behaviour of ULC steel in carbon saturated hot metal

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In addition to hot metal, the Linz-Donawitz oxygen steelmaking process (LD) uses scrap as an iron source. Aside from this fact, scrap acts as a coolant for the exothermic reactions inside the LD converter and will be dissolved in the hot metal. The optimization of the LD process is also focused on the thermodynamic and kinetic modelling where literature-based dissolution equations are used. In laboratory scale experiments the dissolution behaviour of ULC steel scrap in hot metal with two different carbon concentrations was investigated. For the evaluation of the experiments, a literature model for diffusive melting of scrap in hot metal was examined. Based on the measured ablation rate of cylindrical scrap samples submerged in hot metal, the mass transfer coefficient for the dissolution of ULC steel was determined.

KEYWORDS: BASIC OXYGEN FURNACE — STEELMAKING — SCRAP DISSOLUTION — THERMODYNAMICS — PROCESS MODELLING

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INTRODUCTION

Oxygen steelmaking in an LD converter was developed in the early 1950s in Linz and Donawitz (LD) and it gradually became the most dominant method of crude steel production. Large quantities of scrap are used for an ordinary blowing process in an LD converter besides hot metal, which is the main charging material. Scrap is mainly used as a coolant for the process due to heat generation from the oxidation reactions of carbon, silicon, manganese and phosphorus. The dissolution and melting behaviour of scrap influences the whole process cycle in the converter. Many articles concerning the scrap melting process have been published in the past, whereby only a few describe the kinetics of a special steel scrap like ULC in hot metal with changing carbon content. (1 - 12) In this work, an experimental investigation of the melting behaviour of ULC (ultra-low carbon) steel scrap in hot metal with different carbon content was executed. The experiments are based on thermodynamic and kinetic calculations through a MatLab® coded single-zone LD model. Detailed descriptions of the model are published in (13 - 16). In previous calculations it was shown that low alloyed scrap types tend toward a stagnation of dissolution and melting during the oxygen blowing process using literature-based analytical equations (15). Based on these results, laboratory scale experiments were done for validation.
DERIVATIONS OF THE OBSERVED DATA

A collection of measurement data from LD-converters by various authors on the carbon concentration of the molten steel and the melt temperature during the blowing process is presented in (3). It was found that the change of the carbon content is a function of the melt temperature and its trend is parallel to the liquidus line (austenite-liquid melt) in the Fe-Fe₃C phase diagram. Most of the data points are in the liquid phase but some are in the two-phase area of austenite and liquid melt. A similar behaviour is also modelled with the LD model used and was partly published in (15). To describe the scrap melting and dissolution behaviour following literature-based considerations were applied.

\[- \frac{\partial r}{\partial t} = k_{\text{met}} \ln \left( \frac{\% C_{\text{Scrap}} - \% C_{\text{HM}}}{\% C_{\text{Scrap}} - \% C_{\text{liq}}} \right) \]  

The literature describes the melting of scrap in different phases where the diffusion process of carbon controls the scrap dissolution as long as the metal phase temperature is below the melting point of the scrap. If the temperature exceeds the melting point of the scrap a model for forced scrap melting is used. The scrap melting point is defined by the liquidus temperature in the phase diagram for the given chemical composition of the scrap.

The rate of diffusive melting of scrap is determined by the diffusive mass transport of carbon between the liquid metal and the charged scrap. Equation [1] is a mathematical model, proposed by Zhang L. and F. Oeters (6), where the mass transfer coefficient \( k_{\text{met}} \) in [m s⁻¹] is a decisive factor.

DESCRIPTION OF EXPERIMENTS

Dissolution tests

For the experimental procedure, an alumina crucible was charged with 320 g to 345 g of hot metal and positioned in a high temperature vertical tube furnace. The heating rate was 300 K/min to reach the starting temperatures, which are listed in Tab. 1. Before the first and between each further dissolution experiment, a holding time of 30 min was set. To prevent oxidation of the hot metal, the vertical tube furnace was flushed with nitrogen during the heating and melting process. The scrap geometry was cylindrical with a diameter of 12 mm and a length of 30 mm. The whole specimen, including the sample holding, is shown in Fig. 1. The cylinder was submerged into the hot metal with a depth of 20 mm, whereby the axial heat flow to the specimen holding was diminished through a notch. The starting temperature of the specimen was 25 °C. The axial movement of the cylinder to the melt was carried out with a vertical pneumatic controlled cylinder. No stirring by rotation of the cylinder or crucible was performed during the dissolution experiment (i.e. static conditions).
Before each dissolution experiment, the mass of the cylinder was determined. After a defined dissolution time, the cylinder was extracted from the melt and immediately quenched with water to inhibit further carbon diffusion in the sample and weight change by oxidation with air. Through the mass difference (Δm), the ablation rate of the radius Δr/Δt was evaluated. For this evaluation, the density of the cylinder, defined by equation [2], was used and assumed to be equal at the equilibrium temperature. After the experiments, it was observed that only a melting in radial direction occurred. Local density differences can explain this according to the temperature gradient in the boundary layer of the hot metal. The denser liquid will move downwards along the cylinder surface and inhibit a melting on the bottom surface of the cylinder, which is why the length of the cylinder remains nearly constant in all experiments.

The density of the scrap $\rho_{scr}$ is defined by equation [2] published by Miettinen in (19) and is dependent on the temperature (in °C) and the scrap composition. The scrap is assumed to be ferritic and the density for a multicomponent system (composition $C_{i,scr}$ in wt.-%) is, according to the considered elements $i$ of the scrap, explained with equation [2]. (19)

$$
\rho_{scr} = 7875.96 - 0.2977T - 5.62 \times 10^{-5}T^2 + \left(-206.35 + 0.007787T + 1.472 \times 10^{-6}T^2\right) \times C_{C,scr} + 36.86 \times C_{Mn,scr}
$$

The compositions of the hot metal and scrap samples used in the experiments are listed in Tab. 1 and Tab. 2, respectively. The dissolution tests were executed with ULC scrap in carbon saturated hot metal (hot metal 1) at three different temperatures (experiment numbers 1 to 3). Experiment number 4 was executed with hot metal 2 (carbon content of 1 wt.-%), which has a composition close to the liquidus line of the ULC at 1443 °C. The starting temperature of the experiment was assumed to be 1550 °C according to the calculation of the equilibrium temperature, which will be explained subsequently. According to Zhang and Oeters’ model, a stagnation of the melting behaviour should occur when the driving concentration difference (%$C_{s,scrap}$ - %$C_{s,liq}$) in equation [1] should become zero.
Pre-Tests for Verification of Starting Conditions

Before the dissolution experiments, pre-tests were executed to verify the starting conditions for the individual dissolution tests. The starting temperature of the melt was measured with a thermocouple type B, which was directly submerged into the melt. This temperature is shown in Tab. 1 for experiments 1 to 4.

In a preceding test series of this research work, the melting and dissolution behaviour of S235JR scrap was investigated. In the course of this investigation, S235JR specimens (see Tab. 2) were used for verification of the equilibrium temperature between the liquid metal and the submerged sample, which results from the heat exchange between hot melt and cold scrap. For this purpose, a thermocouple type S was located in a bore with a diameter of 1.7 mm in the cylinder centre and at a distance of 10 mm from the cylinder tip. The cold cylinder was submerged into the liquid hot metal and the temperature was monitored until the equilibrium between the melt and the cylinder was reached. This temperature was always below the starting temperature of the hot metal, according to Tab. 1. The S235JR has a very similar enthalpy to ULC steel as a function of temperature. Fig. 2 shows these functions for ULC steel and S235JR, calculated with the FactSageTM FSstel and FactPS database. Accordingly, it was assumed that the equilibrium temperature measured for S235JR should be the same as for ULC steel.

The equilibrium temperature was evaluated with a heat balance, based on equations [3] and [4]. Whereas mScrap and mhot metal are the masses of the scrap and the hot metal in [kg], Q is the heat flux in [W m⁻²], cp,hot metal is the specific heat capacity in [J kg⁻¹ K⁻¹], Tstart scrap/hot metal and T-equilibrium are the starting temperature of the melt or scrap and the equilibrium temperature in [K]. HScrap(T) is the temperature dependent enthalpy in [J kg⁻¹]. For the determination of T-equilibrium, the heat fluxes of the scrap and hot metal must be equal. (20)
\[ Q_{\text{Scrap}} = m_{\text{Scrap}} \times (H_{\text{Scrap}}(T_{\text{equilibrium}}) - H_{\text{Scrap}}(T_{\text{start \ scrap}})) \]

\[ Q_{\text{Hot metal}} = m_{\text{hot metal}} \times c_{p,\text{hot metal}} \times (T_{\text{equilibrium}} - T_{\text{start hot metal}}) \]

**DISCUSSION AND RESULTS**

The increase in the scrap core temperature based on the four investigated starting temperatures of the melt is shown in Fig. 3.

**Fig. 2** - Enthalpy of ULC steel and S235JR

**Fig. 3** - Measurement of the core temperature in the cylinder centre and equilibrium temperature
As seen in Fig. 3, the equilibrium temperature is reached within 10 seconds. The higher the starting temperature, the steeper the temperature gradient is. The measurement also shows that the equilibrium temperature is 65 °C to 80 °C below the starting temperature of the melt. In Tab. 3 the measured starting and equilibrium temperatures of the hot metal and the scrap are listed and compared with the calculated equilibrium temperatures from the heat balance obtained from equations [3] and [4] as well as the data from Fig. 3.

Tab. 3 – Starting and equilibrium temperatures of the hot metal and the scrap

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Starting temperature melt[°C]</th>
<th>Starting temperature scrap[°C]</th>
<th>Equilibrium temperature calculated[°C]</th>
<th>Equilibrium temperature measured [°C]</th>
<th>Temperature drop measured [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experiment 1</td>
<td>1305</td>
<td>25</td>
<td>1239</td>
<td>1230</td>
<td>75</td>
</tr>
<tr>
<td>Experiment 2</td>
<td>1370</td>
<td>25</td>
<td>1301</td>
<td>1300</td>
<td>70</td>
</tr>
<tr>
<td>Experiment 3</td>
<td>1450</td>
<td>25</td>
<td>1377</td>
<td>1385</td>
<td>65</td>
</tr>
<tr>
<td>Experiment 4</td>
<td>1550</td>
<td>25</td>
<td>1452</td>
<td>1470</td>
<td>80</td>
</tr>
</tbody>
</table>

Based on the well-fitting results of the thermocouple measurements, the determination of heat transfer coefficient $k_{net}$ according to equation [1] becomes possible. In the following phase diagram (Fig. 4) of the present ULC scrap, the measurement points according to the hot metal composition (magenta) and the temperature development of experiment 4 with a carbon content of 1 wt.-% are shown. The blue line indicates the way of the equilibrium temperature between 10 to 30 s., whereby the liquidus line is exceeded during this time. Such behaviour does not occur during the experiment with hot metal with 4.58 wt. % carbon. For each equilibrium temperature, the carbon concentration on the liquidus line (cliq) in the Fe-Fe$_3$C phase diagram was determined. The terms of the carbon differences from equation [1] for each experiment are presented in Tab. 4.

**Fig. 4** - Measured equilibrium temperature based on the present ULC scrap composition phase diagram
The experimental determination of the ablation rate of the radius (Δr/Δt) for experiments 1 to 3, executed in hot metal with a carbon content of 4.58 wt.-%, is presented in Fig. 5.

![Fig. 5 - Ablation rate of ULC steel scrap in carbon saturated hot metal](image)

At the beginning of the process, a positive ablation rate occurs. This phenomenon is attributed to a shell formation due to solidification of the liquid hot metal on the cold scrap surface. After a certain progressing time, the turning point is reached, the shell formation stops, and the melting of the shell starts. When the ablation rate turns negative, the melting of the mother scrap starts. The melting and dissolution is dependent on the equilibrium temperature, which is reached in this case, according to the measurements in Fig. 3, after 10 s. By using equation [1] the mass transfer coefficient for this system is calculated. According to the negative sign in the equation, a positive heat transfer is expected if the melting of the mother scrap is in progress. In Fig. 6 the calculated mass transfers of the three starting temperatures are plotted. As mentioned, the mass transfer is negative during the growth and melting of the shell layer at the beginning of the process and becomes positive after 25 s dissolution time at the latest. According to the progress, it could be established that the mass transfer coefficient will increase slightly with the temperature after the equilibrium temperature has been reached.

**Tab. 4 – Carbon differences based on equation [1]**

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Equilibrium temperature [°C]</th>
<th>(%C_{Scrap} - %Clq) [wt.-%]</th>
<th>(%C_{Scrap} - %CHM) [wt.-%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experiment 1</td>
<td>1230</td>
<td>-3.4651</td>
<td>-4.5763</td>
</tr>
<tr>
<td>Experiment 2</td>
<td>1300</td>
<td>-2.7467</td>
<td>-4.5763</td>
</tr>
<tr>
<td>Experiment 3</td>
<td>1385</td>
<td>-1.7612</td>
<td>-4.5763</td>
</tr>
<tr>
<td>Experiment 4</td>
<td>1470</td>
<td>-0.5930</td>
<td>-0.9980</td>
</tr>
</tbody>
</table>
During the LD-process, the carbon content in the liquid melt decreases and the temperature increases in line with the exothermic reactions of oxidation. Therefore, the melting behaviour of ULC steel scrap was determined in modified hot metal with 1 % C. In Fig. 7 the mass transfer coefficient and the ablation rate of experiment 4 are presented. According to the heat balance calculation and measurement the specimen temperature increases to 1415 °C after 10 s and 1470 °C after 30 s, which is just above the liquidus line in the quasibinary phase diagram.
The smaller carbon content in the hot metal results in a strong shell formation at the beginning of the process with a higher negative mass transfer coefficient in comparison to experiments 1 to 3, with higher carbon contents in the hot metal. However, under real process conditions, shell formation will not occur. The scrap will be, due to heat conduction, almost at the equilibrium temperature when those amounts of carbon are reached. This will result in a higher expected negative mass transfer coefficient with an approach to 0. For that reason, a stagnation of the melting behaviour, described by equation [1], where the driving of (\%C_{scrap} - \%C_{liq}) would become zero or negative could be neglected. An explanatory argument is therefore that in this measurement just above the liquidus line, a dissolution of the ULC steel with high mass transfers occurs, which would not be stopped immediately when reaching the liquidus line or ends in an abrupt negative mass transfer coefficient when the temperature falls below the liquidus line.

CONCLUSION
This publication presents the results of laboratory scale experiments on the dissolution and melting behaviour of ULC steel scrap in hot metal with two different carbon concentrations. For the investigation, four different experiments with various hot metal temperatures were performed. For the evaluation of the hot metal mass transfer coefficient of the diffusive scrap melting process, a literature-based equation was used. To get the true process equilibrium temperature, a pre-test with a thermocouple in the core position of the specimen was executed, resulting in a temperature drop of more than 65 °C. Based on these measurements, the carbon concentrations on the liquidus line of the current available Fe-Fe_3C-Mn phase diagram were evaluated. With the measurements, the ablation rates of the radius were determinable and furthermore, the mass transfer coefficients were defined. What is mentionable is that the mass transfer coefficient is slightly dependent on the temperature. At the beginning of the process a shell formation occurs, which results in a negative mass transfer. If the carbon content in the melt decreases and the temperature increases, an increase in the mass transfer is observable. This results in a high melting rate of ULC scrap just above the liquidus line which would not turn into an abrupt negative mass transfer coefficient if the liquidus line were undercut. In summary, the outcomes of this work clearly indicate that the actual temperature and melt composition have a strong impact on the melting and dissolution behaviour of ULC steel scrap. However, the difference in the dissolution behaviour of hot metal with 4.58 and 1 wt.-% carbon is not fully explainable. More research work must be done to investigate and describe the melting and dissolution of scrap for the conditions in the LD process.

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