Slag Rim Formation of two Mould Powders used for Casting of Construction Steel

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Abstract: Two different mould powders are used for casting peritectic steel grades at the voestalpine Stahl Linz GmbH slab casting machine. Powder A forms significantly larger slag rims than Powder B, although their chemical compositions are very similar. For the clarification of the different performance during service, the melting and solidification behaviour as well as the viscosity of the original mould powders have been investigated. Slag rim samples were taken during tail out and cross-sections were analysed by microscopic means. The findings show that the mould powder suppliers use two different raw material concepts. Consequently, the powders do not exhibit identical melting behaviour, which turns out to be the main influencing factor for the slag rim formation.

Keywords: Mould powder, Slag rim, Melting, Solidification

Schlackenkranzbildung zweier Gießpulver beim Vergießen peritektischer Stahlgüten

Zusammenfassung: Für das Vergießen peritektischer Stahlgüten werden bei voestalpine Stahl Linz GmbH zwei verschiedene Stahlgüten eingesetzt. Hierbei bildet Pulver A signifikant größere Schlackenkränze als Pulver B, obwohl ihre chemische Zusammensetzung nahezu ident ist. Zur Klärung dieses unterschiedlichen Verhaltens wurden das Schmelz- und Erstarrungsverhalten sowie die Viskosität der Originalpulver untersucht. Schlackenkränzproben wurden während des Ausfahrums gezogen und mikroskopisch untersucht.

Die Untersuchungen ergaben, dass die Gießpulverlieferanten zwei verschiedene Rohstoffkonzepte verwenden. Aus diesem Grund zeigen diese Pulver ein unterschiedliches Aufschmelzverhalten, welches maßgeblich für die Ausbildung von Schlackenkränzen verantwortlich ist.

Schlüsselwörter: Gießpulver, Schlackenkrantz, Aufschmelzen, Erstarren

1. Introduction

In steel casting operations employing mould powders, usually a crust on the mould in the meniscus region, called slag rim, is formed. This slag rim is attached to the copper mould and moves downwards with the mould oscillation [1]. Its thickness has a strong impact on the heat flux and a large influence on the initial shell formation. In order to achieve a steady state condition of the solidification process and the desired surface quality of the strand, the size of the slag rim should be stable and robust during the whole casting [2, 3]. Rims have been found to show a layered structure of mould slag, small intermediate phases, and original mould powder. They also include carbon black particles [4, 5]. It is mentioned that, during the upward mould motion, the bottom part of the slag rim is coated by liquid and solid powder. During the downward mould motion, this new powder layer melts when coming into contact with the hot steel meniscus. Thus, the infiltrated slag layer and the bottom part of the rim are renewed [6, 7]. According to Takawa [8], the temperature at the interface slag rim–slag lies around 1080 °C. In a previous study, it turned out that the chemical compositions of the mould powders show a deep impact on the slag rim size [9]. Even though the samples investigated in the present studies are very similar in their chemical com-
TABLE 1

| Powder | F  | Na₂O | MgO | Al₂O₃ | SiO₂ | CaO | MnO | C   | CaO/SiO₂ |
|--------|----|------|-----|-------|------|-----|-----|-----|----------|
| A      | 8.2| 8.0  | 2.0 | 3.4   | 30.9 | 38.6| 0.0 | 6.0 | 1.25     |
| B      | 8.1| 6.8  | 1.2 | 4.7   | 31.8 | 38.1| 1.0 | 5.9 | 1.20     |

Fig. 1: Slag rim samples after service

positions, Powder A tends to form larger slag rims during operation than Powder B.

2. Samples

Two mould powders from different suppliers but with almost similar chemical compositions (Table 1) are used for casting peritectic steel grades at the voestalpine Stahl Linz GmbH slab casting machine. Slag rim samples collected during tail out (Fig. 1) and samples from the corresponding mould powders have been analysed.

3. Experimental Procedure

The mineralogical compositions of the mould powders have been analysed by X-ray diffraction (XRD), and polished sections have been investigated with reflected light microscopy and scanning electron microscopy (SEM). For the investigation of the melting behaviour, Simultaneous Thermal Analyses (STA) have been performed. 100 mg of mould powder as delivered were heated to 1350 °C at a rate of 5 °C ⋅ min⁻¹ within a platinum crucible using synthetic air for purging. For the analysis of the microstructure during heating, the mould powders were annealed within steel crucibles (30 x 30 x 40 mm³) at 1000 °C, 1100 °C, and 1200 °C using a preheated high temperature chamber furnace. After a dwell time of 15 min, the samples were quenched within the crucibles to room temperature. Then the specimens were analysed by XRD and microscopic means.

For the investigations of the solidification behaviour and the viscosity for decarburisation, the specimens were annealed at 700 °C for 12 h. Afterwards the samples were melted in covered platinum crucibles at 1400 °C for 15 min, poured onto a steel plate at room temperature, and sprinkled with cold water to obtain glassy pieces. Finally, the samples were ground into powder and split up for the consecutive tests.

The solidification behaviour has been determined by the Furnace Crystallisation Test (FCT). This test is a big scale Differential Thermal Analysis performed in a high temperature chamber furnace. Sample weights of 27 g were put into one platinum crucible and the equal mass of alumina was used as a reference material in a second crucible. The crucibles were heated to 1350 °C, held for 15 min, and cooled to room temperature at a cooling rate of 10 °C/min. The temperatures of the mould powder and the alumina were measured with type S thermocouples situated within the samples and recorded via a data logger at intervals of 30 s. After this procedure the solidified specimen was cut close to the platinum wire into two pieces, analysed by XRD, polished, and investigated by microscopic means.

The dynamic viscosities and the break temperatures of the Fluxes A and B were measured with a commercial high temperature rotational viscometer. The crucible as well as the bob were made of platinum. 27 g of the sample were heated within 30 min to 1450 °C, and the measurement was carried out at a cooling rate of 10 °C ⋅ min⁻¹ with a rotational speed of 150 rpm. During the experiment, the furnace chamber was purged with argon.

From the slag rim samples, polished sections have been prepared and analysed with respect to their mineralogical composition in dependence on the position within the slag rim.

4. Results and Discussion

4.1 Characterisation of the Mould Powders

4.1.1 Raw Materials

Even though the chemical compositions of both mould powders are almost similar, the composition of the raw ma-
terials differs. Powder A is based on a crystallised fluorine containing slag consisting of wollastonite (CaSiO$_3$), cuspidine (Ca$_3$Si$_4$O$_7$F$_2$), and small amounts of oldhamite (CaS) besides a residual glassy phase. Quartz (SiO$_2$), fluorite (CaF$_2$), spodumene (LiAl$_2$O$_3$), and several carbon carriers are added secondarily. The granules contain some amount of gaylussite (Na$_2$Ca(CO$_3$)$_2$·5H$_2$O). Sporadic particles of fly-ash, alumina (Al$_2$O$_3$), natural wollastonite, diopside (CaMgSi$_2$O$_6$), and calcite (CaCO$_3$) have been observed, too.

By contrast, the main components of Powder B are natural wollastonite, fluorite, and fly ash. Calcite, quartz, and several carbon carriers are used in minor amounts. From the results of the STA, it can be concluded that the carbon carriers of Powder B and their amounts differ from those of Powder A. Diopside, pyrolusite (MnO$_2$), alumina, and hematite (Fe$_2$O$_3$) have been detected occasionally.

### 4.1.2 Melting Behaviour

From the STA results, it follows that the diverse mould powder recipes lead to a different melting behaviour. Up to 580 °C, the combustion of carbon is the predominant reaction, but, different to mould Powder A, Powder B contains another carbon carrier with an exothermal peak at 717 °C. The dissociation of the carbonates takes place for Powder A between 581 and 720 °C, but for Powder B this reaction finishes around 802 °C. At 772 °C a peak originating from the reduction of the pyrolusite is visible in the curve of Powder B. The curves exhibit large differences in the formation of liquid. Due to the diffusion processes of sodium into the glassy phase of the slag of Sample A, the first liquid phase is formed at 740 °C, whereas the main melting process starts at higher temperatures (990 °C). At 1062 °C and 1223 °C, two peak maxima are visible until the powder is completely liquefied at 1299 °C. By contrast, no such first liquid formation is visible in the DTA curve of Powder B. Its melting process starts at 1073 °C and ends at 1224 °C. The temperature of the total liquefaction of Sample B corresponds to the same temperature related to the largest melting peak of Powder A.

This different melting behaviour of the mould powders is also confirmed by the annealing experiments (Fig. 2). While for Powder B granules are still visible after annealing at 1000 °C, Powder A forms a compact cube at 1000 °C. From 1000 °C on, cuspidine is the main mineral phase for both powders, but its crystal size is always larger for Powder A (Fig. 3). After annealing at 1000 °C, residues of alumina are present. In case of Powder A, this alumina is surrounded by lazurite (Na$_2$Al$_6$Si$_6$O$_{24}$S$_3$) due to its higher sulphur content, whereas for Powder B nepheline (K$_{0.25}$Na$_{1.24}$Al$_{6.24}$Si$_{9.76}$O$_{32}$) is present in contact with alumina. However, fluorite segregated at 1200 °C from the liquid phase only for Powder B.

### 4.1.3 Viscosity and Solidification

The mean viscosities at 1300 °C are 0.042 ± 0.003 Pas for A and 0.062 ± 0.006 Pas for B. Due to their similar chemical compositions, the Break Temperatures of 1212 ± 4 °C for Sample A and 1213 ± 6 °C for Sample B are almost identical. The crystallisation temperatures detected with the FCT are in the same range: 1184 °C for Sample A and 1196 °C for Samples B. In both samples cuspidine is the main phase, similar to the melting process. The lower viscosity of Sample A supports the formation of bigger crystal compared to those of Slag B [10]. Furthermore, the matrix of both samples is different (see Fig. 4). Apart from cuspidine, crystallised NaAlF$_4$ and only small amounts of glassy phase have been observed for Sample A. For Sample B, by contrast, the amount of glassy phase was considerable higher and small amounts of fluorite and villiaumite (NaF) segregated out of the liquid.

### 4.2 Slag Rims

Depending on the mould powder, the shape of the slag rims taken during tale out differs. As can be seen from Fig. 5, Powder A forms large bulgy slag rims, whereas the slag rims of Powder B are considerably smaller (Fig. 5). For Slag Rim A, the significant increase of the CaO/SiO$_2$ ratio in the centre and the lower parts (marked boundaries in Fig. 5) is remarkable. The CaO/SiO$_2$ ratio detected by EDX in the samples annealed at 1200 °C is 1.09: In the middle of the slag rim bulge, it increases to 1.20, and a maximum of 1.37 can be observed at the lower parts. Simultaneously, the sodium content is decreased by 1–2 wt. Yet, the surrounding areas, marked in Fig. 6, are enriched in sodium and fluorine. From this it can be concluded that the liquidus temperature of the centre part is higher than that of the original mould powder and will not melt easily during service. By contrast, the chemical element distribution of Slag Rim B is homogeneous over the whole cross section.

The information about the formation of the slag rims is derived from the slag rims’ texture. In Rim A, areas with a high concentration of carbon surround the bulges. In Slag B these areas are marked. For Powder B by contrast, such areas can only be observed in contact with the mould. In both slag rims, residual carbon particles as well as pores evolving from gas formation due to the combustion of these carbon particles are entrapped in sintered areas. In Slag
Rim A, these pores are situated in the bulge, whereas for B the pores can be seen all over the cross section (Fig. 6). This gives strong evidence that these parts have been formed by sintered powder and not by liquid. For Powder B this is confirmed by the cuspidine crystal shape, which is similar to the samples fired at 1100 and 1200 °C (see Figs. 3 and 5). Areas corresponding to the quenched samples after a temperature treatment at 1000 °C can only be seen in contact with the mould and do not take part in the mechanism, since the amount of liquid is too low for sticking the granules together sufficiently strong. It can be concluded that most of Slag Rim B is made of molten powder sintered between 1100 and 1200 °C and that the painting mechanism [6] is responsible for the formation of the whole slag rim.
In the case of Powder A, the situation is more complex. The present carbon and spherical pores indicate that the bulge is mainly made of sintered powder, but the texture, especially in the centre of the bulge, looks like the crystallised samples. The cuspidine crystal size is extremely large, partially up to 2 mm. This is explained by the formation of considerable amounts of liquid below 1000 °C. This liquid acts as a binder and promotes slag rim formation in the upper region. Furthermore, the liquid facilitates the diffusion processes and crystal growth of the cuspidine particles already present in the raw material. Even during the negative strip time, the temperatures are too low for the liquefaction of these cuspidine crystals. Moreover, it looks as if the remaining sodium and fluorine rich liquid as well as the carbon particles are squeezed out of the structure and accumulate next to the mould. As a consequence, the C/S ratio and likewise the liquidus temperature in the centre part are increased. Due to this segregation, layers of Ca₂NaSiO₄F villiaumite (NaF), and carbon accumulation appear in the slag rim next to the mould. The lower part of the slag rim is probably made up of a mixture of liquid slag and powder sintered above 1100 °C. The low viscosity of Slag A supports the crystal growth rate. Segregation processes take place as well. As a consequence, even the lower parts do not liquefy easily during the negative strip time. For Slag Rim A, it can be concluded that the painting mechanism works, too. The slag rim, however, is not only built up by the attachment of liquid slag layers and powder sintered above 1100 °C molten powder, but also by granules held together by liquid phases formed from 740 °C upwards. The large melting range of Powder A is therefore responsible for the typical bulging shape of Slag Rim A.

5. Conclusion

According to Perrot [6], “slag rim coalescence” is the main mechanism of slag rim growth. Its grows is especially fast if slag and semi-molten granules are involved. During service the slag acts as a “binder” between the granules. The mechanism requires low viscosities and a special ratio of liquid to non-molten granules. For Powder B this ratio seems to be too low and the viscosity too high. Therefore, its slag rim growth is decreased in this case.
the melting behaviour of these mould powders, it can be concluded that the range of the melting temperature of the mould powder also influence the size of slag rims. If the melting range is small, smaller slag rims can be expected.

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