Corrosion of MgO-C with Magnesium Aluminate Spinel Addition in A Steel Casting Simulator

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Abstract: For more than 20 years, the sidewalls and bottom of steel ladles have been lined with carbon-bonded magnesia (MgO-C) and magnesia-alumina bricks (MAC). The alumina raw materials react with magnesia forming a spinel, which decreases open porosity and slag infiltration. The amount, grain size, and chemistry of the added spinel impact the properties of spinel-containing MgO-C. Corrosion tests have been performed in a steel casting simulator at 1580 °C using 18CrNiMo7-6 steel and Fe-rich slag as corrosion medium. Digital light microscopy and SEM/EDS (scanning electron microscope with energy dispersive spectroscopy) were used to evaluate the corrosion mechanisms. The metal casting simulator test showed that the addition of CaO-MgO-Al₂O₃ aggregates results in the highest corrosion resistance against molten steel and synthetic basic slag compared to alumina-rich spinel aggregates.

Keywords: refractory ceramics; lifetime; corrosion

1. Introduction

MgO-C exhibits excellent slag penetration resistance due to the non-wetting property of carbon. The combination of this good corrosion resistance with the high thermal shock resistance, low thermal expansion, high thermal conductivity, and high toughness makes it a superior refractory material for the slag line of steel ladles [1]. But poor high-temperature strength and the oxidation of carbon at operation temperature limit the service life of MgO-C refractories. Lee and Zhang summarized the corrosion mechanisms of MgO-C [2]. The first decomposition step is the oxidation of the carbon component by surrounding oxygen and/or FeO (from the slag). Consequently, slag penetration and dissolution of the MgO grain proceed. Also, the reduction of MgO by carbon takes place at high temperatures. However, the subsequent reaction of MgO with oxygen, which leads to the formation of a dense secondary MgO-layer at the slag–refractory interface, will hinder slag penetration in turn. Additionally, the CO gas formed by carbon oxidation creates an excess pressure at the slag–refractory interface, which will hamper corrosion as well. The need of refractories with enhanced thermomechanical properties by steel producers led to the development of carbon-containing spinel-forming or spinel-containing bricks of type Al₂O₃-MgAl₂O₄-C [3]. Compared to conventional high alumina and doloma refractories, Al₂O₃-MgAl₂O₄-C bricks show better performance in steel ladle [4]. Al₂O₃ containing MgO-C (AMC-bricks), where periclase and alumina form spinel (MgAl₂O₄), exhibit excellent properties like high thermal shock and corrosion resistance as well as high thermal conductivity [5]. Similar to Al₂O₃-MgAl₂O₄-C, the improved performance of AMC ladle bricks compared to conventional high
alumina and doloma refractories have been proven [4]. Spinel contributes with its high refractoriness, high resistance to slag penetration and corrosion, and high thermal shock resistance [6]. In MgO-C, slag penetration decreases with increasing alumina content due to (a) spinel formation, (b) formation of aluminum nitride phases and aluminum carbide (Al$_4$C$_3$), which reduce oxidation of carbon, and (c) an increase of slag viscosity by reaction of Al$_2$O$_3$ with slag compounds like SiO$_2$ and CaO [7]. The addition of pre-reacted spinel (instead of Al$_2$O$_3$) to MgO-C led to the development of MgO-MgAl$_2$O$_4$-C bricks, a new type of carbon-bonded spinel-containing refractories. Thereby, the material composition (spinel and carbon content) and the manufacturing process influence the properties of MgO-MgAl$_2$O$_4$-C refractories. Choi and Jun presented a way of producing MgO-C-MgAl$_2$O$_4$ by adding thermite reaction products of MgO and Al to MgO-C leading to superior corrosion resistance against CaO/SiO$_2$ slag (ratio 3.3) to MgO-C and MgO-C-Al [8]. Ultra-low-carbon-containing MgO-MgAl$_2$O$_4$-C bricks (C <1 wt.%) exhibit improved oxidation resistance, volume stability, HMXR, and slag resistance. By applying these bricks in the slag line of a 200 t converter, an average service life of 75 heats could be obtained [9]. Ganesh et al. demonstrated that 20 wt.% of spinel addition improves the slag erosion resistance of MgO-C by 48 % when the MgAl$_2$O$_4$ was prepared by reaction of a stoichiometric mixture of caustic MgO (particle size of 5 µm) with Al(OH)$_3$ (particle size of 85 µm) [10]. Compared to MgO-C with commercially available micron-sized spinel, the addition of 0.5 and 1 wt.% nano-sized MgAl$_2$O$_4$ spinel prepared by citrate-nitrate gel method leads to improved resistance against oxidation, thermal spalling, and corrosion [11]. Nano-spinel affects the pore size distribution and hence limits the transportation of oxygen and slag into the matrix. This will protect graphite from being decarburized. Also, the chemistry of the spinel raw material impacts the properties of MgO-C. Wu et al. found that CaAl$_2$O$_7$-MgAl$_2$O$_4$ improves the thermal shock resistance of a MgO-C brick (4 wt.% C) and recommended the addition of 6 wt.% [12]. However, the differences in physical, mechanical, and thermomechanical properties of MgO-MgAl$_2$O$_4$-C with various spinel raw materials have been found to be only marginal [13–15].

Fundamental investigations of the corrosion mechanism of MgO-MgAl$_2$O$_4$-C refractories for application in steel ladles are not available so far. For the properties of MgO-MgAl$_2$O$_4$-C refractories, the chemical composition of the spinel raw material is essential. Hence, the aim of this paper is to describe the influence of two common coarse-grained spinel raw materials on the corrosion mechanism of MgO-C. Laboratory corrosion test with molten steel and a basic synthetic ladle slag at 1580 °C under fully controlled conditions has been performed. The corrosion resistance and slag penetration were evaluated by analyzing the slag-refractory interface at macroscopic as well as microscopic scales in SEM.

2. Materials and Methods

As reference material, MgO with a maximum grain size of 4 mm and approx. 3 wt.% carbon content was produced (sample MgO-C). As carbon sources, carbon black and graphite were added in equal parts. CMA 72 clinker (Imerys Aluminates, Paris La Defense, France) and alumina-rich spinel AR78 with fraction 1–3 mm (Almatis GmbH, Frankfurt, Germany) serve as spinel raw material, respectively (mineral composition listed in Table 1). They were sieved to a grain size below 2 mm and replaced parts of the MgO 1–2 mm fraction. Additionally, 1.5 wt.% liquid and 1.5 wt.% powder resin and 0.3 wt.% hardener (hexamethylenetetramine) were added. The composition of the sample bodies can be found in Table 2.

| Phase (wt.%) | AR78 | CMA 72 |
|-------------|------|--------|
| CaO·Al$_2$O$_3$ | - | 18–22 |
| CaO·2Al$_2$O$_3$ | - | 8–12 |
| MgAl$_2$O$_4$ | 100 | 68–72 |
| Ca$_2$Al [AlSiO$_7$] | - | <1 |

Table 1. Mineral composition of fine-grained AR78 (−90 µm) and CMA 72 (cement).
In a first step, the spinel raw materials were mixed with the coarse MgO fractions for 3 min. Afterwards, tempered (70 °C) liquid resin was added and mixed for another 2 min to coat the coarse grains. Finally, the fine components (residual MgO, graphite, carbon black, powder resin, and hardener) were added followed by another 5 min mixing step. Afterwards, bar-shaped sample bodies with a dimension of 25 × 25 × 150 mm³ were prepared using a uniaxial press (RUCKS engine building GmbH, Glauchau, Germany) and applied pressure of 120 MPa. After pressing, the samples were hardened by applying a three-step schedule up to a maximum temperature of 180 °C. The sample bodies were placed in a box filled with petrol coke and hence, thermal treatment (carbonization) was applied for 5 h at 1400 °C in a reducing atmosphere. After carbonization, the residual carbon content of 3 wt.% was calculated resulting from carbon black and graphite content, and the carbon yield of the resin binders.

X-ray diffraction (XRD) analysis (PHILIPS diffractometer with CuKα-radiation, K-alpha1 = 1.54060 [° Å]) was used for phase identification. Therefore, bars of the three compositions were crushed and milled to achieve a maximum grain size of 20 µm.

The corrosion resistance of the different compositions against molten steel and slag were determined in a laboratory melting device (metal casting simulator, SYSTEC, Karlstadt, Germany), the structure and functionality of which have been described by Dudczig et al. [16]. The samples were fixed to an equipment and attached to a crucible (Figure 1). Afterwards, the crucible was filled with approximately 30 kg of steel grade 18CrNiMo7-6 (Deutsche Edelstahlwerke GmbH, Witten, Germany).

![Figure 1. Equipment for corrosion test of samples.](image)

A basic slag was synthesized as corrosion medium by mixing 19 wt.% Ciment Fondu, 52 wt.% LDSF RG, 6 wt.% CMA 72 (all Imerys Aluminates, Paris La Defense, France), 3 wt.% MS 971 (ELKEM AS, Oslo, Norway), 3.5 wt.% nedMag 99 (Nedmag B.V., Veendam, The Netherlands), and 30.5 wt.% CaCO₃ (Carl Roth GmbH, Karlsruhe, Germany). Table 3 lists the chemical composition of the synthesized slag (analyzed by X-ray fluorescence analysis). After mixing, the slag was granulated, calcined at 1000 °C,
and sieved to grain size in the range of 100–500 µm. This will prevent dust formation during addition to the steel melt as well as prevent degassing reactions during the test run. The steel was melted at approximately 1580 °C under a fully controlled argon atmosphere. Afterwards, the slag was stepwise added to the top of the steel melt via a feed unit. Investigations in a hot stage microscope in air revealed a melting temperature (hemisphere temperature) of the synthetic slag of 1304 °C. Due to metal melt flow caused by induction field heating, the slag melts and first accumulates at the sidewall of the crucible, where the samples are located. After adding more slag, the whole steel melt gets covered by the synthetic slag having a thickness of 5 to 10 mm (see Figure 2). These conditions were kept constant for 30 min, followed by shutting down the metal casting simulator.

Table 3. Chemical composition of the synthesized ladle slag.

| Compound | Synthetic Slag (wt.%) |
|----------|-----------------------|
| Al₂O₃    | 42.4                  |
| CaO      | 40.0                  |
| SiO₂     | 6.0                   |
| MgO      | 8.3                   |
| Fe₂O₃    | 0.1                   |
| TiO₂     | 0.9                   |
| P₂O₅     | 0.002                 |
| S        | 0.07                  |
| Traces   | 2.228                 |

Figure 2. Formation of slag layer on top of molten steel in the crucible inside the metal casting simulator (exemplary without sample equipment).

After laboratory testing, the corrosion was evaluated at a macroscopic scale, with aid of digital light microscopy and with SEM/EDS (scanning electron microscope with energy dispersive spectroscopy) (ESEM XL30 FEG, FEI, Eindhoven, The Netherlands).

3. Results and Discussion

3.1. Phase Composition—XRD

Since the CA-phases containing in spinel-binder CMA 72 are able to interact with MgO in a different way compared to AR78 (slight alumina-excess, see Table 1), it is expected that thermal treatment will lead to different phases depending on the added spinel raw material [17]. The spinel portion of CMA 72 is almost stoichiometric, and it is not expected that they will interact with the other constituents of the sample body during carbonization. But the CA-phases (CA and CA₂) may react...
with MgO forming additional spinel if the carbonization temperature is high enough. In contrast, AR78 contains free alumina and may react with MgO to form spinel, even at a lower temperature.

Table 4 lists all main and trace phases of sample MgO-C, M-AR-C, and M-CMA-C after carbonization at 1400 °C. The MgO-C reference sample contains MgO and carbon, and Ca₃Mg(SiO₄)₂ as a trace phase, which is contained in the MgO raw material as a well-known impurity. Sample M-AR-C consist of MgO, MgAl₂O₄, C, and the trace phase Ca₃Mg(SiO₄)₂. By comparing the peak positions (main spinel peak) of sample M-AR-C and the raw material AR78, a slight shift from 31.35 °2Theta (AR78) to 31.45 °2Theta can be observed in the diffractogram (see Figure 3). This is an indication that AR78 incorporates some trace elements into the structure. In sample M-CMA-C, MgO, C, and MgAl₂O₄, but no CA-phases (e.g., CaAl₂O₄) can be detected. The absence of CA-phases indicates that (a) an additional spinel formation by reaction of MgO with Al₂O₃ of the CA-phases of the CMA raw material may take place, (b) the contained amount of CA-phases is below the detection limit of 1 wt.%, or (c) a glassy phase was formed by reaction of CA-phases with calcium silicate impurities in MgO. The detectable additional spinel formation by XRD method (peak high and width) is only marginal. Overall, a higher spinel peak can be detected in sample M-CMA-C compared to sample M-AR-C, as well as a peak shift from 31.3 to 31.45 °2Theta. Of course, based on the information given by XRD-diffractogram the total spinel content cannot be calculated.

Table 4. Main and trace phases after carbonization at 1400 °C.

| Sample      | Main Phase | Traces               |
|-------------|------------|----------------------|
| MgO-C       | MgO, C     | Ca₃Mg(SiO₄)₂         |
| M-AR-C      | MgO, MgAl₂O₄, C | Ca₃Mg(SiO₄)₂ |
| M-CMA-C     | MgO, MgAl₂O₄, C | -                  |

![Figure 3](image-url)  
Figure 3. XRD spectra variation in the vicinity of MgAl₂O₄ peak structure in sample MgO-C, M-AR-C, and M-CMA-C (extraction of diffractogram).

3.2. Laboratory Metal Casting Simulator Test

The slag and molten steel could not be poured into the tundish after the test run, as the samples were fixed to the crucible. Consequently, after shutting down the induction heating, the samples effectively froze in the steel and slag, but slag and steel could easily be separated. Hence, broken sample bits with contact to steel melt and slag are existent to perform investigations. Above the slag
Above the slag line, all three sample bars show a decarburization area. At the macroscopic scale, no specific differences are detectable. The edges are not sharp anymore, but no particular washout can be noticed. Light microscopy investigations on polished surfaces of the samples at 50-times magnification revealed much more differences in corrosion behavior (Figure 4). Compared to sample M-CMA-C and M-AR-C, the interface of sample MgO-C is irregular and presents numerous exposed grains. It is not possible to calculate the total slag infiltration depth as the initial sample edge cannot be discerned. But taking the highest and lowest part of the slag-refractory interface into account, an infiltration depth above 1.2 mm was calculated at sample MgO-C. The slag-refractory interface of sample M-AR-C with coarse AR78 has a wavy character, which indicates a washout of coarse and fine parts of the material. The slag infiltration depth was calculated being in the range of 800 µm. In contrast, the sample containing CMA 72 (M-CMA-C) shows a clear slag-refractory interface with average slag infiltration depth in a range of 100–200 µm, which indicates less interaction of the refractory material with slag.

Figure 4. Polished slag-refractory interface after corrosion test.

Figure 5a displays the slag–refractory interface of reference sample MgO-C analyzed with SEM. It is obvious that the slag easily attacks the MgO coarse grains and matrix particles after the graphite and the carbon binder components were oxidized. The fine matrix particles were surrounded and dissolved by slag at an early stage of corrosion test. The coarse grains were infiltrated by slag as
well, which will cause a stepwise decomposition. Figure 5b presents an interspace between coarse MgO grains filled with slag located approx. 200 µm beyond the slag line. There are three different compositions detectable, which were analyzed by EDS. An overview of the composition at all EDS spots is given in Table 5. The main grey area contains 45 wt.% Al₂O₃, 30 wt.% CaO, 20 wt.% SiO₂, and 5 wt.% MgO (spot 1). According to the Al₂O₃-CaO-MgO-SiO₂ quaternary system with 5 wt.% MgO, the detected composition at spot 1 lies in the field of stability of spinel and nearby melilite (more precise akermanite) phase [18]. The grey area at spot 2 consists of about 38 wt.% Al₂O₃, 15 wt.% SiO₂, 7 wt.% MgO, 3 wt.% of sulphur-containing phase, and shows a depletion of CaO. This composition lies on the liquidus line of spinel and melilite (Ca₂MgSi₂O₇). In comparison to spot 2, the light grey spot 3 shows a low silica content and high amounts of TiO₂ and sulphur. Compared to initial synthetic slag composition (Table 3), an increase of silica and alumina, as well as depletion of CaO, can be detected in the slag between the MgO grains. The discrepancy of the composition of detected EDS-spots in comparison to synthetic slag implies the interaction of MgO-C reference material and its impurities especially with CaO of the slag. However, according to the MgO-CaO binary system, in range of 1600 °C MgO will not be dissolved by CaO but forms a solid solution. This was also proven by higher CaO content nearby slag-refractory interface compared to CaO content detected in slag far away from MgO-C material. The interaction of MgO-C with CaO or calcium silicates results in a eutectic slag with lower melting point and lower viscosity, which will easily penetrate the MgO-C reference material. Al₂O₃ and sulphur (originating from slag) seem to play only a minor role on the corrosion, and TiO₂ does not take part in the corrosion reaction. Furthermore, spinel, melilite, or anorthite are able to crystallize in the slag during cooling.

Figure 5c displays the slag-refractory interface of sample M-AR-C composed of a mixture of coarse grains and numerous fine particles surrounded by slag. Similar to sample MgO-C, slag with three different compositions is detectable inside the coarse grains of sample M-AR-C (Figure 5d). The dominating dark grey area (spot 4) consists of about 40 wt.% Al₂O₃, 30 wt.% CaO, 20 wt.% SiO₂, 5 wt.% MgO, and some sulphur. Like spot 2, these composition lies on the liquidus line of melilite and spinel. Composition at spot 5 is rich in CaO (50 wt.%) and SiO₂ (35 wt.%) and has a very low amount of Al₂O₃ (5 wt.%), MgO (2 wt.%), TiO₂, and sulphur. Light grey spot 6 correlates to a Ca- and S-rich phase. Compared to initial synthetic slag, CaO content is nearly constant, there is an increase in SiO₂ content and a depletion of Al₂O₃. Hence, in contrast to sample MgO-C, sample M-AR-C will react with CaO and additionally with Al₂O₃, which will intensify the basicity of the residual slag which results in higher melting point and hence higher viscosity, and consequently less slag penetration. EDS investigations of the slag far away from M-AR-C material interphase verify these findings, as lower Al₂O₃ content but higher CaO content was detected compared to slag far away from sample MgO-C. The slag inside sample M-AR-C has the potential of crystallization of rankinite as well as anorthite and CA. In addition, there is a crystallization of CaS.

Figure 5e,f displays two slag-refractory interfaces of sample M-CMA-C. Two different zones can be detected: (a) an infiltration zone with coarse and fine particles surrounded by slag, and (b) the initial refractory material. Infiltration and dissolution take place but less of the solved material will be transported into the slag compared to sample M-AR-C. The infiltration zone (spot 8) is composed of 48 wt.% Al₂O₃, 37 wt.% CaO, 10 wt.% SiO₂, and 4 wt.% MgO, which can be assigned to the spinel stability field. Sample M-CMA-C will additionally react with Al₂O₃, which will result in a higher melting point, higher viscosity, and reduced slag penetration. EDS investigations of the slag far away from sample M-CMA-C revealed nearly similar composition like those of sample M-AR-C. The residual penetrating slag inside one of the bigger grains is composed of 32.5 wt.% MgO, 43 wt.% SiO₂, and 24.5 wt.% CaO (EDS spot 7). This composition lies next to the field of stability of MgO. By interaction with MgO, forsterite (M₂S) or monticellite (CMS) will be formed until the stability field of MgO will be reached very fast. This will disrupt further corrosion.
dissolved by slag at an early stage of corrosion test. The coarse grains were infiltrated by slag as well, which will cause a stepwise decomposition. Figure 5b presents an interspace between coarse MgO grains filled with slag located approx. 200 µm beyond the slag line. There are three different compositions detectable, which were analyzed by EDS. An overview of the composition at all EDS spots is given in Table 5. The main grey area contains 45 wt.% Al$_2$O$_3$, 30 wt.% CaO, 20 wt.% SiO$_2$, and 5 wt.% MgO (spot 1). According to the Al$_2$O$_3$-CaO-MgO-SiO$_2$ quaternary system with 5 wt.% MgO, the detected composition at spot 1 lies in the field of stability of spinel and nearby melilite (more precise akermanite) [18]. The grey area at spot 2 consists of about 38 wt.% Al$_2$O$_3$, 15 wt.% SiO$_2$, 7 wt.% MgO, 3 wt.% of sulphur-containing phase, and shows a depletion of CaO. This composition lies on the liquidus line of spinel and melilite (Ca$_2$MgSi$_2$O$_7$). In comparison to spot 2, the light grey spot 3 shows a low silica content and high amounts of TiO$_2$ and sulphur. Compared to initial synthetic slag composition (Table 3), an increase of silica and alumina, as well as depletion of CaO, can be detected in the slag between the MgO grains. The discrepancy of the composition of detected EDS spots in comparison to synthetic slag implies the interaction of MgO-C reference material and its impurities especially with CaO of the slag. However, according to the MgO-C binary system, in range of 1600 °C MgO will not be dissolved by CaO but forms a solid solution. This was also proven by higher CaO content nearby slag-refractory interface compared to CaO content detected in slag far away from MgO-C material. The interaction of MgO-C with CaO or calcium silicates results in a eutectic slag with lower melting point and lower viscosity, which will easily penetrate the MgO-C reference material. Al$_2$O$_3$ and sulphur (originating from slag) seem to play only a minor role on the corrosion, and TiO$_2$ does not take part in the corrosion reaction. Furthermore, spinel, melilite, or anorthite are able to crystallize in the slag during cooling.

Table 5. Results of EDS spot analysis in SEM.

| Spot | Compound (wt.%) |
|------|----------------|
|      | Al$_2$O$_3$ | MgO | SiO$_2$ | CaO | TiO$_2$ | SO$_3$ |
| 1    | 46.3        | 4.7 | 18.9    | 29.9 | 0.3     | -      |
| 2    | 37.6        | 7.2 | 15.5    | 35.4 | 1.0     | 3.3    |
| 3    | 39.9        | 3.8 | 5.2     | 34.0 | 8.0     | 9.1    |
| 4    | 40.2        | 4.5 | 22.6    | 30.1 | -       | 2.6    |
| 5    | 4.9         | 1.8 | 33.8    | 52.7 | 1.0     | 5.9    |
| 6    | -           | -   | -       | 36.7 | -       | 63.3   |
| 7    | -           | 32.5| 43.1    | 24.4 | -       | -      |
| 8    | 47.5        | 3.6 | 10.4    | 37.2 | 1.3     | -      |

4. Conclusions

The aim of this work was to identify the influence of AR78 and CMA 72 on the corrosion mechanism of MgO-C, determined by a laboratory corrosion test in a metal casting simulator with
steel 18CrNiMo7-6 and synthetic slag at 1580 °C for 30 min. The reference sample MgO-C without spinel addition gets infiltrated by slag and shows intense dissolving especially by CaO. The slag resulting from this reaction still has a highly reactive character, which explains the high corrosion rate. MgO-C containing alumina-rich spinel AR78 also reacts with CaO but additionally with Al2O3 from the slag. This leads to a depletion of Al2O3 in the infiltrated slag, increases its basic character, and consequently reduces its reactivity (stability field of melilite and spinel), observable in reduced corrosion compared to reference MgO-C. The MgO-C sample containing CMA 72 also reacts with CaO and Al2O3 of the infiltrated slag, but the resulting slag composition lies in the stability field of spinel and next to MgO. Consequently, by further reaction with the infiltrated slag, the formation of forsterite (M2SiO4) or monticellite (CMS) is expected until the stability field of MgO will be reached very fast. This will disrupt further corrosion and explains the very low corrosion rate.

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