



Article Effect of MgO and K₂O on High-Al Silicon– Manganese Alloy Slag Viscosity and Structure

Xiangdong Xing ^{1,2}, Zhuogang Pang ^{1,2}, Jianlu Zheng ¹, Yueli Du ¹ and Shan Ren ^{3,*} and Jiantao Ju ^{1,2,*}

- ¹ School of Metallurgical Engineering, Xi'an University of Architecture and Technology, Xi'an 710055, China; xaxxd@xauat.edu.cn (X.X.); hjpangzhuogang@163.com (Z.P.); zhengjianlud@163.com (J.Z.); 18821795206@163.com (Y.D.)
- ² Metallurgical Engineering Technology Research Center of Shaanxi Province, Xi'an 710055, China
- ³ College of Materials Science and Engineering, Chongqing University, Chongqing 400044, China
- * Correspondence: shan.ren@cqu.edu.cn (S.R.); jiguangheng@xauat.edu.cn (J.J.)

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Abstract: The viscosity, melting proprieties, and molten structure of the high-Al silicon–manganese slag of SiO₂–CaO–25 mass% Al₂O₃–MgO–MnO–K₂O system with a varying MgO and K₂O content were studied. The results show that with the increase in MgO content from 4 to 10 mass%, the measured viscosity and flow activation energy decreases, but K₂O has an effect on increasing those of slags. However, the melting temperature increases due to the formation of high-melting-point phase spinel. Meanwhile, Fourier transform infrared (FTIR) and X-ray photoelectron spectra (XPS) were conducted to understand the variation of slag structure. The O^{2–} dissociates from MgO can interact with the O⁰ within Si–O or Al–O network structures, corresponding to the decrease in the trough depth of [SiO₄] tetrahedral and [AlO₄] tetrahedral. However, when K₂O is added into the molten slag, the K⁺ can accelerate the formation of [AlO₄] tetrahedra, resulting in the increase in O⁰ and O[–] and the polymerization of the structure.

Keywords: viscosity; structure; silicon-manganese slag; melting properties

1. Introduction

Nowadays, the silicomanganese (SiMn) alloy, a common deoxidant and desulfurizer, has been widely used in most steels containing silicon and manganese [1]. Calcium-based silicomanganese slag is primarily adopted in the production of SiMn because it easily controls the furnace conditions and alloy composition, but a large amount of slag will be generated and reduce the recovery of Si and Mn. In this case, many plants have begun to produce silicomanganese alloys using high-alumina slag (\geq 15 mass% Al₂O₃) to improve the recovery rate of Si and Mn elements and reduce the slag production as well as energy consumption. Meanwhile, as the consumption of high-quality manganese ores, the cost of raw materials for manganese ferroalloys producers gradually rises, which increases the utilization of low-grade, low-cost materials, but it also leads to the increase in the content of Al₂O₃ and alkaline oxides such as K₂O in silicomanganese alloys slags.

In the production of SiMn, the viscous behavior of molten slag is one of the most crucial factors that affects not only the smelting process, but also the reaction kinetics and the mass/heat transfer [2]. For instance, when viscosity increases significantly, it will delay the transfer of silicon and manganese between slag and metal, as well as the final slag removal. Controlling the viscosity of the slag within an appropriate range can ensure that slags have good fluidity and increase the amount of MnO and SiO₂ reduced from the slag to the alloy. However, the increment of Al₂O₃ and K₂O content in the slag will significantly increase viscosity, resulting in many operating and product quality problems. MgO is the main compound in slag, and an appropriate increase in its content can improve the fluidity of

the slag. Thus, in the present study, we aimed to adjust the MgO content of slag to alleviate the increment in the viscosity of slag with high aluminum and potassium.

Previous works by the present authors [3] found that MgO decreased the viscosity and break temperature of BaO-bearing blast furnace (BF) slag. The results of Sun et al. [4] indicated that MgO provided oxygen ions to the molten slag which caused the depolymerization of the complex structure into simple units, resulting in decreasing the viscosity of SiO₂–CaO–17 mass% Al₂O₃–MgO slag, while the viscosity increased when MgO content exceeded 10 mass%. Yao et al. [5] investigated the structure of high alumina blast furnace slag, and found that Si–O tetrahedral linked together to form the network structure, and basic oxides (MgO) could destroy the complex polymerization. Otherwise, Liu et al. [6] studied the effect of K₂O on the viscosity and structure of CaO–SiO₂–8 mass% MgO–17 mass% Al₂O₃–2.5 mass% BaO–K₂O slag and revealed that K₂O increased the polymerization degree of aluminosilicate anions, which increased slag viscosity. Zhang et al. [7] suggested that the K⁺ compensated Al³⁺ to form a bridging oxygen bond O_{ALK}, which is stronger than O_{ALCa}, increasing the slag viscosity, and reached a maximum at K₂O/Al₂O₃ > 1. Although research works related to the influence of MgO and K₂O on the viscosity and structure of blast furnace slag have been implemented, the data of that on the unique SiO₂–CaO–Al₂O₃–MgO–MnO–K₂O silicomanganese alloys slag with a content of 25 mass% Al₂O is still lacking.

In this study, the effect of MgO and K₂O content on the viscosity of silicomanganese alloy slag with 25 mass% Al₂O₃ at a fixed CaO/SiO₂ mass ratio of 0.65 was investigated using the rotating spindle method. In addition, the melting properties of the slag were also analyzed. To correlate the effect of MgO and K₂O on the viscosity and structure, Fourier transform infrared (FTIR) spectroscopy and X-ray photoelectron spectroscopy (XPS) of the slag were performed to understand the structural rule of the slags.

2. Experimental

2.1. Materials Preparation

In this study, the chemical composition of Al₂O₃-based SiMn alloy slag was first analyzed by X-ray fluoroscopy (XRF, S4 Explorer; Bruker AXS, Karlsruhe, Germany), and the result was shown in Table 1. Based on the information of Table 1, the experimental slag samples were synthesized using reagent grade chemicals of SiO₂, CaO, Al₂O₃, MgO, MnO, and K₂CO₃. The target compositions of the experimental samples are listed in Table 2, and the basicity of all synthesized samples was fixed at 0.65. The raw materials precisely weighed 180 g to form the given compositions and were then mixed thoroughly in a mortar. Thereafter, the mixture was placed into a Mo crucible and pre-melted at 1600 °C under Ar (99.99%, 0.25 L/min) atmosphere. After keeping 1600 °C for 180 min, to obtain uniform melt, it was poured into water. Then, the quenched sample was collected and dried.

Composition	SiO ₂	CaO	Al ₂ O ₃	MnO	MgO	K ₂ O	Other
Mass%	38.54	26.21	22.87	4.24	3.97	0.32	3.85

Table 2. The intended composition of the synthesized sample.

Table 1. Chemical composition of high-Al silicon-manganese alloy slag.

Sample	SiO ₂	CaO	Al ₂ O ₃	MnO	MgO	K ₂ O	Basicity
	(mass%)	(mass%)	(mass%)	(mass%)	(mass%)	(mass%)	(CaO/SiO ₂)
1	38.79	25.21	25	5	4	2	0.65
2	37.58	24.42	25	5	6	2	0.65
3	36.36	23.64	25	5	8	2	0.65
4	35.15	22.85	25	5	10	2	0.65
5	36.97	24.03	25	5	8	1	0.65
6	35.76	23.24	25	5	8	3	0.65
7	35.15	22.85	25	5	8	4	0.65
	Sample 1 2 3 4 5 6 7	SiO2 (mass%) 1 38.79 2 37.58 3 36.36 4 35.15 5 36.97 6 35.76 7 35.15	SiO2CaO(mass%)(mass%)138.79237.5824.42336.36435.15536.9724.03635.7623.64735.1522.85	SiO2CaOAl2O3(mass%)(mass%)(mass%)138.7925.2125237.5824.4225336.3623.6425435.1522.8525536.9724.0325635.7623.2425735.1522.8525	$\begin{array}{c c c c c c c c c } SiO_2 & CaO & Al_2O_3 & MnO \\\hline & (mass\%) & (mass\%) & (mass\%) & (mass\%) \\\hline 1 & 38.79 & 25.21 & 25 & 5 \\ 2 & 37.58 & 24.42 & 25 & 5 \\ 3 & 36.36 & 23.64 & 25 & 5 \\ 3 & 36.36 & 23.64 & 25 & 5 \\ 4 & 35.15 & 22.85 & 25 & 5 \\ 5 & 36.97 & 24.03 & 25 & 5 \\ 6 & 35.76 & 23.24 & 25 & 5 \\ 7 & 35.15 & 22.85 & 25 & 5 \\ \end{array}$	SampleSiO2CaOAl2O3MnOMgO(mass%)(mass%)(mass%)(mass%)(mass%)138.7925.212554237.5824.422556336.3623.642558435.1522.8525510536.9724.032558635.7623.242558735.1522.852558	SampleSiO2CaOAl2O3MnOMgOK2O(mass%)(mass%)(mass%)(mass%)(mass%)(mass%)(mass%)138.7925.2125542237.5824.4225562336.3623.6425582435.1522.85255102536.9724.0325581635.7623.2425583735.1522.8525584

2.2. Measurement of Viscosity

The slag viscosity measurements were conducted using the rotating cylinder method. The schematic figure of the viscosity measurement equipment can be seen in the previous work [3]. The apparatus mainly consists of five U-shaped MoSi₂ heating components, a Pt–6% Rh and Pt–30% Rh thermocouple (the error < ± 2 °C), a Mo rotor, and a differential transformer. The quenched samples were melted at 1600 °C and kept for 30 min to obtain a homogeneous liquid phase. The spindle was then immersed into the liquid slag. Thereafter, the furnace was cooled down slowly from 1600 °C at the rate of 5 °C/min. The viscosity measurements were performed at intervals of 20 °C during the cooling cycle. The measurements were conducted until the viscosity value of melt > 5 Pa·s. The viscosity–time curve was recorded by the software. There was very little Mo (<0.05 mass%) infiltrated into the slag, which has an ignorable effect on the viscosity of molten slags. The viscosity measurement equipment was calibrated by castor oil before each experiment.

2.3. Measurement of Melting Properties

The melting properties of slags were determined by a melting-point and melting-rate comprehensive measurement system. (MTLQ-RD-1600, Chongqing University of Science and Technology, Chongqing, China), as illustrated in Figure 1. The system mainly included a high-temperature furnace, a charge coupled device (CCD) camera, a cooling water tank, and an analysis computer. The height of the cylindrical slag sample (Φ 3 × 3 mm²) was reduced during the heating process (the heating rate was 15 °C/min). The analysis computer recorded the temperature at which the height of the sample reduced to 75%, 50%, and 25% of the initial height as the softening, hemispherical, and flow temperature, respectively. Each sample was tested three times and we calculated the average values of the experiment.



Figure 1. The schematic figure of melting-point and melting-rate comprehensive measurement system.

2.4. Measurement of Structure

All the samples were pre-melted at 1600 °C and quenched in water to maintain the slag's molten structural state. The quenched powders (–200 mesh) were conducted by X-ray diffraction (XRD, D8 Advance A25; Bruker AXS, Karlsruhe, Germany) for ensuring the state of the slag. The XRD used a Cu K α radiation at a scan rate of 3°/min over the degree of 20–90°, and the XRD patterns are presented in Figure 2. There are no characteristic peaks that can detected in Figure 2, suggesting that the samples were amorphous. The quenched samples were used to analyze the slag structure through FTIR (Nicolet iS5; Thermo Fisher Scientific, Waltham, MA, USA) and XPS (EscaLab Xi+, Thermo Fisher Scientific, Waltham, MA, USA) and XPS (EscaLab Xi+, Thermo Fisher Scientific, Waltham, MA, USA) spectroscopy. During the FTIR measurement, 2.0 mg sample (–300 mesh) and 300 mg KBr were mixed, and then the mixture was pressed into a film. The sample was recorded at a scope of 400–4000 cm⁻¹ with a resolution of 4 cm⁻¹. Each sample was scanned for 32 s. In addition, XPS was carried out at a vacuum chamber with 8 × 10⁻¹⁰ Pa and used an Al K α monochromatic source (1486.6 eV). The O_{1s} peak was deconvoluted and peak fitted using XPS Peak (Version 4.1) and Origin software (Version 2019). The peaks were appropriately calibrated by the C_{1s} of externally doped carbon as the reference peak, and the binding energy of C_{1s} is 284.80 eV.



Figure 2. The XRD pattern of quenched slags.

3. Results and Discussion

3.1. Viscosity of Slag

Figure 3 graphically shows the viscosity of the SiO₂–CaO–25 mass% Al₂O₃–MnO–MgO–K₂O system with a varying MgO content at a fixed basicity of 0.65. It can be clearly observed that the slag viscosity decreases with the addition of MgO content in the temperature range from 1370 °C to 1550 °C, and the decreasing rate of viscosity slows down when the MgO content exceeds 8 mass%. For example, the average decrement of viscosity is 0.284 Pa·s with the increment of MgO from 4 to 8 mass% at 1460 °C, whereas the viscosity just decreases by 0.168 Pa·s as the MgO content increases from 8 to 10 mass%. The viscous values of previous studies related to high Al₂O₃ containing slag are also included in Figure 3 [5,8,9]. Though the slag compositions are different, it exhibits a similar trend of slag viscosity to that of this study. However, it can be noted that the viscosity values of slag are lower than that of the present study, which is attributed to the high basicity (CaO/SiO₂ > 1.0). Work performed by Li et al. [10] suggested that MgO depolymerized the complex silicate and aluminate network structure, and then decreased viscosity. According to Kim et al. [11], MgO is classified as basic oxide and acted as a network modifier, and much of the network structure units were depolymerized into dimer or monomer units, as expressed by Equation (1):

$$O^{2-} + Si - O - Si = 2Si - O^{-}$$

The viscosity for samples 3, 5, 6, and 7 with varying K₂O contents is shown in Figure 4. On the contrary of MgO, K2O tends to increase the viscosity of slag. The effect of K2O is relatively weak when the temperature is higher than 1520 °C, which is because the excess thermal energy had sufficiently modified the intricate network structure of the slag, and large amounts of simpler structural units exist in molten slag. The same variation trends of slag viscosity with the addition of K₂O are found within the works of Liu et al. [6] and Chang et al. [12]. Generally, K₂O is classified as basic oxide which will introduce free oxygen (O^{2-}) to destroy Si–O bonds within an intricate silicate network, leading to the increment of non-bridging oxygens and eventually lowering slag viscosity. However, the result is in contradiction with the trend of viscosity of present slag, indicating that some other effects of K2O have more significant influence on slag viscosity. Based on the previous literature [7,12], it can be speculated that Al³⁺ prefers to form AlO₄⁵⁻ tetrahedrons and the process needs cations (such as K⁺ or Ca²⁺) to maintain electroneutrality, which is called charge compensation. However, K⁺ possesses higher priority than Ca²⁺ to compensate Al³⁺, leading to K⁺ gradually substituting Ca²⁺ and transforming O_{Al,Ca} to OALK. Since the length of chemical bonds between Al and Ca (α_{ALCa} = 4.996) are longer than those between Al and K ($\alpha_{Al,k}$ = 4.156), the chemical bonds around O_{Al,K} are stronger than those around O_{Al,Ca}. Thus, the charge compensation leads to an increase in the stability of slag structure and increases the

slag viscosity. Considering the increment of viscosity under the condition of K₂O, the effect of K⁺ seems to be more dominating. From the results of slag viscosity with different MgO and K₂O content, the addition of MgO is limited to the range of 6–8 mass%, and K₂O content should be controlled below 1 mass%, for the production of SiMn alloys.

In addition, the volume fractions of liquid and solid phases were calculated by Factsage thermodynamic software, and the results of calculation were presented in Tables S1–S7 and Figure S1 that were attached in the supplementary material. It should be mentioned that there are solid precipitates during the viscosity measurement, and the maximum amount of precipitation is 10%.



Figure 3. MgO content dependence on the viscosity of the slags at the temperature between 1370 °C and 1550 °C.



Figure 4. K₂O content dependence on the viscosity of the slags at the temperature between 1370 °C and 1550 °C.

3.2. Flow Activation Energy

In general, the variation of viscosity is related to the flow activation energy that represents the energy barrier to overcome the shear stress for liquid melt movement [13]. The viscosity of slag is affected by the temperature, and the Weymann–Frenkel equation is usually adopted to describe the relationship between the two, as expressed as Equation (2) [14]. This can be taken logarithmically and written as Equation (3):

$$\eta = \eta_0 + \operatorname{Exp}\left(\frac{E_f}{RT}\right)$$
⁽²⁾

$$\ln(\eta) = \ln(\eta_0) + \left(\frac{E_f}{RT}\right)$$
(3)

where η , η_0 , E_f , R, and T are the viscosity of the slag in Pa·s, the pre-exponential constant, the flow activation energy in J/mol, the ideal gas constant of 8.314 J·(mol·K)⁻¹, and the absolute temperature in K, respectively. The values of E_f are calculated on the basis of the slope of the fitting line depicted in Figure 5. Table 3 lists the results of the calculation.

As shown in Figure 5 and Table 3, the variation of E_f shows good agreement with that of viscosity. The flow activation energy increases with increasing K₂O content, while MgO has an effect on diminishing the values of E_f. This indicates that with the increase in MgO content, the complex structure transforms into simpler units which leads to a decline in the energy barrier for viscous flow, whereas K₂O increases the resistance of melt flows due to the polymerization of the structural units. The viscosity of silicate–aluminate melt is predominantly determined by its structure at high temperature, and many spectroscopic methods have been developed to identify the ionic structural units. Moreover, the FTIR spectroscopy and XPS were conducted in the present study.



Figure 5. The reciprocal of temperature dependence on the viscosity of the slags (**a**) with a varying MgO content; and (**b**) with a varying K₂O content.

MgO (mass%)	Ef (kJ/mol)	K ₂ O (mass%)	Ef (kJ/mol)
4	203.45 ± 12.95	1	183.06 ± 17.10
6	194.06 ± 8.29	2	190.14 ± 13.50
8	185.31 ± 9.11	3	203.29 ± 13.27
10	175.99 ± 3.86	4	218.47 ± 12.51

Table 3. The calculated results of apparent activation energy.

3.3. Structure Analysis of the Slag Using FTIR Spectroscopy

The viscosity of slag is closely related to its structure. Generally, the viscosity increases when the complexity of the structure increases, but it decreases when the simple structural units are formed. Figure 6 presents the FTIR transmittance spectra of glassy samples with different MgO content. The symmetric stretching bands of [SiO₄] tetrahedra are detected between 1200 and 750 cm⁻¹, and it is generally divided into four typical bands assign to Q⁰ ([SiO₄]⁴⁻, monomer, 880–850 cm⁻¹), Q¹ ([SiO₄]⁴⁻, dimers, 920–900 cm⁻¹), Q² ([Si₂O₇]⁶⁻, chains, 1000–950 cm⁻¹), and Q³ ([Si₃O₁₀]⁸⁻, sheets, 1100–1050 cm⁻¹), respectively [3,9,15,16]. It can be observed that as the MgO content is increased, the transmission trough between 1200–750 cm⁻¹ becomes regularly shallower. This suggests that the silicate structures are depolymerized under the action of MgO. In addition, the significant differences for both of the trough of [AlO₄] tetrahedra and Si–O–Al are also ascertained. The curves of the [AlO₄] tetrahedral band get smooth, indicating that MgO also plays a role of modifier on the aluminate network and decreases the relative fraction of the [AlO₄] tetrahedral units within the melt. According to the work of Qi et al., [15] the bridging oxygen in [AlO₄] tetrahedra can be transformed to non-bridging oxygen, which suggests the decrease in the degree of polymerization of slag, and the transformation is expressed by Equations (4) and (5). Meanwhile, the decline depth of the Si–O–Al trough further

illustrates the number of complex structures formed by the combination of [SiO₄] tetrahedra and [AlO₄] tetrahedra is reduced, that is, the complex network structure is depolymerized.

$$2AlO_2^- + O_2^- = Al_2O_5^{4-} \tag{4}$$

$$AlO_2^- + O_2^- = AlO_3^{3-}$$
 (5)



Figure 6. The effect of the MgO on the FTIR spectra of SiO₂–CaO–25 mass% Al₂O₃–MnO–MgO–2 mass% K₂O slag.

The FTIR spectra for glassy samples with varying K₂O content are shown in Figure 7. Though the band group of [SiO₄] tetrahedral stretching between 1200 and 750 cm⁻¹ deepens slightly with the addition of K₂O, the center of the trough declines from 961 to 938 cm⁻¹, which is presumably due to the formation of simpler Si–O structures, but it will cause the slag viscosity to decrease. In fact, the experimental viscosity of the slag is contrary to the aforementioned speculation. It can be noted that as the K₂O content increases, the shoulder at nearly 736 cm⁻¹ gradually moves to higher wavenumbers, broadening the [AlO₄] tetrahedral trough, and the depth increases from h₁ (0.95) to h₂ (2.8). In addition, the Si–O–Al bending troughs identified by the trough at nearly 480 cm⁻¹ also shifts to a higher wavenumber. The results indicate that K₂O has a different effect on [SiO₄] tetrahedral and [AlO₄] tetrahedral structures. According to Sohn et al. [17], K₂O can accelerate the formation of [AlO₄] tetrahedra that acts as a network former, which can be expressed by Equation (6). Combining the results of the viscosity experiment and the analysis of the FTIR spectra, it can be inferred that K₂O has more effect on aluminate structures to form network structure:

$$2AI^{3+} + 7O^{2-} + K_2O = 2AIO_4^{5-} + 2K^+$$
(6)



Figure 7. The effect of K₂O on the FTIR spectra of SiO₂–CaO–25 mass% Al₂O₃–8 mass% MgO–MnO–K₂O slag.

3.4. Structure Analysis of the Slag Using XPS

To elucidate the influence of K₂O and MgO on the structure of the SiO₂–CaO–Al₂O₃–MnO–MgO–K₂O system, the amorphous quenched samples were analyzed, as was the main O_{1s} peak, using XPS. According to the studies of Kim et al. [18] and Wu et al. [19], three different types of oxygens exists in silicate melts, which are bridging oxygen (BO, O⁰, connected to two Si⁴⁺), non-bridging oxygen (NBO, O⁻, connected to a Si⁴⁺), and free oxygen (FO, O²⁻, not connected with silicon cations), respectively. The fraction variation of different types of oxygen ions can provide a detailed bonding complexity of the melt. For instance, the higher the amount of bridging oxygen, the more the structural units (Si–O or Al–O) are connected, and the greater the degree of polymerization (DOP) of slag.

Figure 8 shows the deconvoluted result of the O_{1s} binding energy from the XPS, and the background is removed using Shirley's method. The binding energy, of nearly 530.3 eV, 531.3 eV, and 532.2 eV, assigned to FO, NBO, and BO, respectively, which is close to that observed by Wang et al. [20] and Kim and Shon. [18] From Figure 8a, the intensity peaks corresponding to different oxygen ion species change significantly with MgO additions. In particular, the O²⁻ always takes up a relatively large portion, and the fraction of O^0 gradually decreases with the addition of MgO. Additionally, it can be seen that the portion of O⁻ firstly enlarges with the content of MgO increasing from 4 to 6 mass%, and then decreases by further adding MgO. On the contrary, K2O plays a role in increasing the amount of O⁰ and O⁻, leading to the higher DOP of slag. By calculating the integrated areas of the deconvoluted peaks, the relative fractions of O⁰, O⁻, and O²⁻ can be observed visually, and the results of the calculation are depicted in Figure 9. According to Figure 9a, the fraction of O²⁻ increases from 41.25% to 49.35% with the MgO content up of to 10 mass%, while the average decrement of the fraction of O⁰, and O⁻ are 2.29% and 0.41%. It further suggests that MgO can dissociate free oxygen (O²⁻) and it interacts with the bridging oxygen (O⁰) to create the simpler non-bridging oxygen (O⁻), as expressed by Equation (7), which weakens the complexity of slag structure. It can note that by increasing the content of MgO from 8 to 10 mass%, the relative fraction of O⁰ did not obviously reduce, which leads us to speculate that the bridging oxygen has already been depolymerized by O²⁻, hence the reduction becomes less noticeable. Besides, the effect of K₂O on the fractions of oxygen species is presented in Figure 9b. The fraction of BO and NBO increases from 21.17% to 26.11% and from 33.42% to 34.29%, respectively. Based on the aforementioned analysis of FTIR spectra, the increment of BO and NBO is attributed to K₂O improving the formation of the [AlO₄] tetrahedral that acts as a network former. Thus, the slag structure becomes intricate, and the viscosity is higher, which is consistent with the results of the viscous measurement:

$$O^0 + O^2 = O^-$$
 (7)



Figure 8. The O_{1s} XPS spectra for SiO₂-CaO-25 mass% Al₂O₃-8 mass% MgO-MnO-K₂O slag as function of binding energy (**a**) with varying MgO content; (**b**) with varying K₂O content; BO is Bridging oxygen.



Figure 9. The fractions of the different types of oxygens (**a**) with a varying MgO content; and (**b**) with a varying K₂O content.

3.5. Melting Properties of Slags

The composition of slag affects the melting properties of slags that are very important considerations in the SiMn production process. The influence of K₂O and MgO on the melting properties of SiO₂– CaO–Al₂O₃–MnO–MgO–K₂O slag are shown in Figure 10. It is clear that the softening, hemispherical,

and flowing temperatures of all slags rises with the increment of MgO and K₂O content. The flowing temperature (T_f) can be considered as the liquid temperature (totally melted), and it increases from 1372 °C to 1428 °C with the addition of MgO at 2 mass% K₂O. MgO has an effect on decreasing slag viscosity, but has an opposite effect on the liquid temperature. Besides, with the increase in K₂O content, the T_h rises from 1389 °C to 1425 °C.



Figure 10. The effect of MgO and K₂O on the melting properties of slags.

These phenomena can be explained from the phase diagram of the SiO₂–CaO–Al₂O₃–MnO–MgO–K₂O slag calculated by Factsage7.1 software, as shown in Figure 11. It was found from Figure 9a that the increase in MgO content leads to the transition from the liquid phase area to the spinel corresponding to the high melting point area, resulting in the increment of liquid temperature. The phase diagram in the condition of 8 mass% MgO varying with K₂O is shown in Figure 9b, from which the phase moves from the liquid phase to the spinel phase with increasing K₂O. Considering K₂O not only increases the liquid temperature but also increases the slag viscosity, thus, the decreasing K₂O content in the slag is beneficial to improve the melting speed and fluidity of the slag. Moreover, from the phase diagram, it is conducive to limit the content of MgO and K₂O to less than 8 mass% and 1 mass%, respectively, for the slag with a low liquid temperature.



Figure 11. Phase diagram of SiO₂–CaO–Al₂O₃–MnO–MgO–K₂O slag system at 1400 °C: (**a**) with a varying MgO content; (**b**) with a varying K₂O content.

4. Conclusions

The viscosity, structure, and melting properties of high-Al silicon–manganese alloy slag with a varying K₂O and MgO content were investigated in this study, and the main conclusions can be summarized as follows:

- (1) The viscosity of SiO₂-CaO-25 mass% Al₂O₃-MgO-MnO-K₂O slag decreases with the increment of MgO content, and the decreasing rate of viscosity slows down by further increasing the content of MgO from 8 to 10 mass%. However, the viscosity of slag increases with the increase in K₂O.
- (2) MgO acts as a network modifier that can dissociate O²⁻ and interact with the O⁰ within Si–O or Al–O network structures, leading to an increase in the simpler O⁻, and then depolymerizes the complex network structure. On the contrary, when K₂O is added into the molten slag, the K⁺ can accelerate the formation of [AlO₄] tetrahedra and connects aluminate–silicate structural units, which results in the increase in O⁰ and O⁻ and the polymerization of structure.
- (3) An increase in the MgO and K₂O content leads to the transformation from the feldspar and liquid phase to spinel, which increases the melting temperature. By combining with the results of the viscosity and melting properties experiment, it is suitable to limit the content of MgO and K₂O to less than 8 mass% and 1 mass%, respectively, for the production of silicon–manganese alloy.

Supplementary Materials: The following are available online at www.mdpi.com/2075-163X/10/9/810/s1, Figure S1: Relationship between the volume fraction of phases and calculated viscosity at different temperatures, Table S1: Calculation results of the liquid and solid phase fractions of sample 1#, Table S2: Calculation results of the liquid and solid phase fractions of sample 2#, Table S3: Calculation results of the liquid and solid phase fractions of sample 3#, Table S4: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractions of sample 5#, Table S6: Calculation results of the liquid and solid phase fractio

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