Study on structure characteristics of B$_2$O$_3$ and TiO$_2$-bearing F-free mold flux by Raman spectroscopy

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Abstract: Structure characteristics of fluoride-Free mold flux containing simultaneously B$_2$O$_3$ and TiO$_2$ have been investigated by Raman spectroscopy in this work. Raman spectra for glass samples with different basicities, different contents of TiO$_2$ and B$_2$O$_3$ were recorded during the experiments. According to the experiments results, increase of TiO$_2$ content leads to the appearance of [TiO$_4$] and [TiO$_6$] structure groups, and [TiO$_4$] becomes the main structure unit in the system. TiO$_2$ produces a certain destructive effect on Si-O-Si network structure as well as large borate group and conducive to the formation of some other complex structure groups, such as (Si,Ti) coupling in sheet unit. It can be concluded that, with the increase of B$_2$O$_3$, the ratio of mixing of the Q$^0$ structure unit and [TiO$_4$] structural group decrease and the ratio of sheet structure unit increase, and there forms large borate group. Existence of B$_2$O$_3$ increases polymerization degree of the slag system. In addition, increasing basicity causes to the decrease of Q$^2$ and sheet structure unit and increase of mixing of the Q$^0$ structure unit and [TiO$_4$] structural group, and weakening the large borate group. It could be concluded that the increase of basicity reduces the degree of polymerization of the system.

Keywords: Fluoride-Free mold flux, B$_2$O$_3$, TiO$_2$, structure, Raman spectroscopy

1. Introduction

As critical materials in continuous casting process, mold flux plays an irreplaceable role in metallurgical industry. However, the fluorides within traditional mold flux produce harmful influence both on the natural environment and on equipment seriously, such as, volatilization of fluorides leading to erosion to continuous caster, pollution of environment, acidification of the cooling water, human health hazard, and so on [1-4]. Thus, the development and application of fluorine free mould flux is of great significance and has become a research area of interest [2-5]. It is reported that slags bearing B$_2$O$_3$ and/or TiO$_2$ become most promising substitute for traditional mould fluxes [2-8]. According to some previous studies [2, 4, 7, 8], the existence of TiO$_2$ in mould fluxes can easily lead to the formation of some crystals (eg. CaTiO$_3$ or CaO·SiO$_2$·TiO$_2$) with high melting point, which is potential to replace the cuspidine generated by fluorine in mould fluxes to ensure the crystallizability of slag to achieve good heat transfer performance. Besides, B$_2$O$_3$, as an effective fluxing agent to lower the melting point of mould flux, has been taken into consideration for being added to mould fluxes to adjust the viscosity or melting properties of slag [3, 5, 6]. So it is absolutely essential to investigate the properties of mould fluxes simultaneously containing B$_2$O$_3$ and/or TiO$_2$ to develop the optimal fluorine free mould fluxes.
The performance of mold flux, such as heat transfer and lubricity, serves an important function to ensure the process stability of the continuous casting process and surface quality of products. And benign physio-chemical properties, which contribute the excellent performance of mold fluxes [1-9], are closely related to structure characteristic of slag. Knowledge of structure of molten slags is essential to understand the properties of slags. Consequently, it is practical importance of investigating the structure of mold slag to explore the appropriate physio-chemical properties of mold flux so as to guarantee the good performance, and thus achieve the purpose of controlling metallurgical processes.

For a better understanding of the structure characteristic of molten slag, a lot of analytical techniques have been developed and widely used, such as IR spectrum analysis, nuclear magnetic resonance (NMR), electronic probe rays microscopic analysis (EPMA), high temperature X-ray diffraction analysis, Raman spectroscopy, and so on. In comparison to other methods, Raman spectroscopy is with a lot of advantages in terms of microanalysis, high analysis speed, high precision and accuracy, application to high-temperature condition and no-destruction [10-14]. Therefore, Raman spectroscopy has been successfully applied to detect the vibration mode of molecular or micro-structure units and has become a very popular technique to acquire the information of constitutes and structure of materials [10-12].

It is effective to study the structure of high temperature melts through analyzing the Raman spectra of the corresponding glasses at room temperature in the fact that the spectra of glass and melts are similar based on the comparison of the infrared and Raman spectra of various silicate glasses and that of the corresponding melts by many researchers [13, 14]. Thus, the structure characteristic of B_2O_3 and TiO_2-bearing fluoride-free mold flux has been investigated by analyzing the room-temperature Raman spectra of the glass systems in this work. Structural information of glass systems with variable TiO_2 and B_2O_3 content and different basicities (R=w(CaO)/w(SiO_2)) were obtained by Raman spectra to examine the influences of TiO_2, B_2O_3 content and basicities on the structure of B_2O_3 and TiO_2-bearing fluoride-free mold flux.

2. Experimental

2.1 Sample preparation

Analytical grade CaO, SiO_2, Al_2O_3, MgO, TiO_2, Na_2CO_3 and H_3BO_3 were taken as raw materials, with Na_2CO_3 and H_3BO_3 being substitutes for Na_2O and B_2O_3, respectively. Table 1 presents the compositions (wt. %) of the glass samples investigated.

Glass samples with different TiO_2, B_2O_3 content and different basicities were prepared by the conventional melting and quenching method. Raw materials were mixed, taken into a platinum crucible and then melted in high temperature furnace at approximately 1573K in air atmosphere. The samples were held at 1573K for nearly 3h to make sure complete melting and homogenization. After melting, the melts were quenched by water and then bulk glass samples are formed. These glass samples are proved to be amorphous by XRD (e.g. Figure 1).
Table 1 Chemical compositions of experimental samples (wt. %)

<table>
<thead>
<tr>
<th>Sample Number</th>
<th>basicity</th>
<th>CaO</th>
<th>SiO₂</th>
<th>Al₃O₃</th>
<th>MgO</th>
<th>Na₂O</th>
<th>TiO₂</th>
<th>B₂O₃</th>
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<tr>
<td>1</td>
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<td>11</td>
<td>R=0.8</td>
<td>31.6</td>
<td>39.4</td>
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<td>10</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

Fig. 1 A typical XRD pattern of the quenched sample

2.2 Raman spectroscopy measurement

Bulk glass samples were subjected to the Raman spectroscopy analysis. Raman spectra were acquired using a laser confocal micro-Raman spectrometer, JY-HR800, manufactured by Jobin Yvon of France. The experiments were carried out at room temperature using excitation wavelength of 532 nm and the light source was a semiconductor laser with power of 1mw. The frequency band measured in this work ranged from 100 to 2000 cm⁻¹.

3. Results and discussion

Figure 2, 3 and 4 present the room-temperature Raman spectra for glass samples with different contents of TiO₂, B₂O₃ and different basicities, respectively. All the backgrounds of the measured Raman signals have been subtracted. All of the measured Raman spectra of glass samples are deconvolved by Gaussian-Deconvolution method similar to method by Mysen et al. [15] with the minimum correlation coefficient $r^2 \geq 0.998$. The deconvoluted results have been shown in Figure 5, 7 and 9. The focus of attention in this work is the middle and high frequency of the Raman spectra. Assignments of Raman peaks that are to the interest of present work have been listed in Table 2.
Table 2 Assignments of Raman bands in spectra for the mold flux glass system

<table>
<thead>
<tr>
<th>Raman shift (cm(^{-1}))</th>
<th>Raman assignments</th>
</tr>
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<tbody>
<tr>
<td>677~680</td>
<td>O-Si-O deformation motion</td>
</tr>
<tr>
<td>690~700</td>
<td>Ti-O stretch vibrations of Ti in six-fold coordination ([TiO(_6)])</td>
</tr>
<tr>
<td>~870</td>
<td>(\text{SiO}^2_4) stretching in monomer structure unit (Q(<em>0)) or Q(</em>{4(4A)}) structure unit</td>
</tr>
<tr>
<td>~950</td>
<td>(\text{SiO}^2_4) stretching with two bridging oxygen in chain structure unit (Q(_2))</td>
</tr>
<tr>
<td>1030~1037</td>
<td>Si-O' antisymmetric stretching in any structural units that contain bridging oxygen</td>
</tr>
<tr>
<td>1192~1204</td>
<td>pyroborate group</td>
</tr>
<tr>
<td>1369<del>1379, 1435</del>1470</td>
<td>(\text{BO}_2)O(_3) triangles units attached to the large borate group</td>
</tr>
</tbody>
</table>

![Raman spectra for samples with different contents of TiO\(_2\) (No. 1, 2, 3, 4 and 5) at room temperature, after background subtraction](image1)

**Fig. 2** Raman spectra for samples with different contents of TiO\(_2\) (No. 1, 2, 3, 4 and 5) at room temperature, after background subtraction

![Raman spectra for samples with different contents of B\(_2\)O\(_3\) (No. 3, 6, 7, 8 and 9) at room temperature, after background subtraction](image2)

**Fig. 3** Raman spectra for samples with different contents of B\(_2\)O\(_3\) (No. 3, 6, 7, 8 and 9) at room temperature, after background subtraction
Figure 4 Raman spectra for samples with different basicities (No. 3, 10 and 11) at room temperature, after background subtraction

Figure 5 shows the deconvoluted results of Raman spectra for glass systems with different TiO$_2$ content. It can be seen that in the TiO$_2$-free glass system (Figure 5 (a)), there is a dominant Raman peak at approximately 950 cm$^{-1}$ which is due to $SiO_3^{2-}$ stretching with NBO/Si=2 (non-bridging oxygen per tetrahedrally coordinated cation) and referred to as Q$^2$ (superscript refers to the number of bridging oxygen) species in a chain structure [16-21]. There are also three primary bands for sample without TiO$_2$, 679, ~870 and 1030 cm$^{-1}$. According to some previous researches [16-26], the band 679 cm$^{-1}$ could be assigned to O-Si-O deformation motion, the 870 cm$^{-1}$ band is assigned to $SiO_4^{4-}$ stretching with NBO/Si=4 and referred to as Q$^0$ species in monomer structure, or it is due to Q$^4$(4Al) structure unit, and the band at about 1030 cm$^{-1}$ is due to Si-O$^o$ (O$^o$-bridging O atoms) antisymmetric stretching in any structural units that contain bridging oxygen but does not need to be fully polymerized. It can be seen from Figure 5 that, as the addition of TiO$_2$ increases, the band at about 870 cm$^{-1}$ becomes the dominant peak and simultaneously shifts toward low frequencies gradually, as 856, 840 and 838 cm$^{-1}$, the 1030 cm$^{-1}$ band has shift toward low frequency, about 1012 and 997 cm$^{-1}$ respectively, and the ~950 cm$^{-1}$ peak shows a decrease in intensity. Based on the reports of Mysen et al [22, 27], for Ti-bearing silicate melts, the bands at about 830 and 810 cm$^{-1}$ could correspond to the vibration of Si-O$^2$ structural unit and Ti-O$^2$ vibrations in the form of [TiO$_4$], respectively, so it can be presumed that the emerging bands with increase of TiO$_2$ content, 856, 840 and 838 cm$^{-1}$ may indicate mixing of the Q$^0$ structure unit and [TiO$_4$] structural groups; the band in the range of 997–1030 cm$^{-1}$ probably corresponds to the mixing of (Si,Ti) coupling of stretch vibrations in a sheet structure with randomly distribution of Ti$^{4+}$ and Si$^{4+}$ [22] and Si-O$^o$ antisymmetric stretching in any structural units that contain bridging oxygen. Area ratios of the bands could reflect the abundance changes of different structure units, so the area fractions of bands in 600–1600 cm$^{-1}$ are taken into consideration to describe the changes of structure groups in this work. Figure 6 presents the area fractions of bands in 600–1600 cm$^{-1}$ as functions of TiO$_2$ contents. It can be seen that
with addition of TiO₂, the ratio of 838~875 cm⁻¹ bands increases, the ratio of 944~966 cm⁻¹ bands decreases and the ratio of 997~1030 cm⁻¹ bands shows minor increase. That is to say, TiO₂ exists in the system mainly in the form of [TiO₄] and the increase of TiO₂ leads to the increase of ratio of Q⁰ structure unit and [TiO₄] structural groups, reduction of Q² structure group, disappearance of the O-Si-O deformation. And simultaneously, some TiO₂ enters into Si-O-Si network to form (Si,Ti) coupling of stretch vibrations in a sheet structure.

Fig. 5 Deconvoluted results of Raman spectra for samples with different TiO₂ contents
Besides, with increase of TiO₂, there appear a new band, 690–695 cm⁻¹, which shifts to 712 cm⁻¹ as the content of TiO₂ increases to 10% in mass. According to Mysen et al [22], the band at approximately 690–695 cm⁻¹ could be attributed to Ti-O stretch vibrations of Ti in six-fold coordination, [TiO₆], and the band at 720 cm⁻¹ reflects the O-Ti-O deformation, so the band at 712 cm⁻¹ could be proposed to the mixing of the [TiO₆] structure unit and O-Ti-O deformation. There are also some changes for bands in high frequency with the increase of TiO₂. It can be seen from Figure 6 that, the ratio of Raman bands for small borate structure group, pyroborate (1192–1204 cm⁻¹), has somewhat decrease and the ratio of Raman bands for large borate structure group (1369–1379, 1435–1470 cm⁻¹) decreases to a certain extent.

![Graph](image-url)

**Fig. 6** Area fractions of bands in 600–1600 cm⁻¹ as functions of TiO₂ contents

In general, it can be concluded that increasing the content of TiO₂ leads to the appearance of [TiO₄] and [TiO₆] structure groups and [TiO₄] becomes the main structure unit in the system; TiO₂ produces a certain destructive effect on Si-O-Si network structure as well as large borate group and conducive to the formation of some other complex structure groups, such as (Si,Ti) coupling in sheet unit.

The deconvoluted results of Raman spectra for glass samples with different B₂O₃ content are presented in Figure 7. In the glass system without B₂O₃ (Figure 7 (a)), the main structure units are mixing of the Q⁰ structure unit and [TiO₄] structural groups, Q² in chain structure unit, [TiO₄] unit as well as mixing of (Si,Ti) coupling of stretch vibrations in a sheet structure and Si-O° antisymmetric stretching in any structural units that contain bridging oxygen and. Figure 8 shows the area fractions of bands in 600–1600 cm⁻¹ as functions of B₂O₃ contents. It can be observed that, as the content of B₂O₃ increases, the ratio of the dominant peak (in the vicinity of 866 cm⁻¹) decreases and the ratio of bands for sheet structure units (1010–1014 cm⁻¹) shows an increased tendency. It can be seen from Figure 7 and Figure 8 that as minor B₂O₃ is added into the system, such as 3% in mass, there firstly forms pyroborate structure unit, then, with the
increase of B$_2$O$_3$ content, BO$_2$O triangles units attached to the large borate group appear, and simultaneously the ratio of these borate structure groups increases to a certain extent. As the content of B$_2$O$_3$ reaches 10% in mass, the band at about 695 cm$^{-1}$ move to the vicinity of 712 cm$^{-1}$, that is, there forms some amount of O-Ti-O deformation due to the increase of B$_2$O$_3$. By and large, existence of B$_2$O$_3$ leads to enhancement of polymerization degree of the slag system.

**Fig. 7** Deconvoluted results of Raman spectra for glass samples with different B$_2$O$_3$ content
Fig. 8 Area fractions of bands in 600–1600 cm$^{-1}$ as functions of B$_2$O$_3$ contents

Fig. 9 Deconvoluted results of Raman spectra for glass samples with different basicities
Figure 9 exhibits the deconvoluted results of Raman spectra for glass systems with different basicities. It can be seen that, when the basicity is 0.8, the structure groups exist in the system mainly in the form of mixing of the Q⁰ structure unit and [TiO₄] structural groups (856–864 cm⁻¹), Q² in chain structure unit (945–968 cm⁻¹), mixing of (Si,Ti) coupling of stretch vibrations in a sheet structure and Si-O° antisymmetric stretching in any structural units that contain bridging oxygen (996–1037 cm⁻¹), and B₂O₃ exists in the system primarily in the form of BO₂O⁻ triangles units attached to the large borate group. Figure 10 presents the area fractions of bands in 600–1600 cm⁻¹ as functions of basicities. It can be seen that, as the basicity increases, both the ratio of Q² in chain structure unit (945–968 cm⁻¹) and sheet structure unit (996–1037 cm⁻¹) decrease, while the ratio of mixing of the Q⁰ structure unit and [TiO₄] structural groups (856–864 cm⁻¹) increases. It can be concluded that increasing the basicity produces a certain destructive effect on Si-O-Si network. Also, with the increase of basicity, and the band for BO₂O⁻ triangles units attached to the large borate group becomes too weak to be detected and along with the appearance of small borate group, pyroborate unit. That is, the increase of basicity causes simplification of borate network structure. Namely, the increase of basicity causes negative effects on the borate network and Si-O-Si structure containing bridging oxygen.

4. Conclusions

The Raman spectroscopy technique has provided a deep insight into the structure of the B₂O₃ and TiO₂-bearing fluoride-Free mold flux glass system. The increase of TiO₂ leads to the appearance of [TiO₄], [TiO₆] and O-Ti-O deformation structure groups. And TiO₂ exists in the slag mainly in the form of [TiO₄]. TiO₂ weakens the Raman signals of Q², O-Si-O deformation and BO₂O⁻ triangles units linked to the large borate group, increases the ratio of mixing of the Q⁰ structure unit and [TiO₄] structural groups. That is to say, TiO₂ produces a certain destructive effect on Si-O-Si network structure as well as large borate group and conducive to the formation of some other complex structure groups, such as (Si,Ti) coupling in sheet structure unit.
With the increase of B$_2$O$_3$, the ratio of mixing of the Q$^0$ structure unit and [TiO$_4$] structural group decreases and the ratio of sheet structure unit increase, and the peak for BO$_2$O$^-$ unit attached to the large borate group shows an increase in intensity. Generally, existence of B$_2$O$_3$ leads to the enhancement of polymerization degree of the slag system.

With the increase of basicity, the ratio of Q$^2$ and sheet structure unit decrease, and the ratio of mixing of the Q$^0$ structure unit and [TiO$_4$] structural group increase. And BO$_2$O$^-$ triangles units linked to the large borate group weaken, along with the appearance of pyroborate group. It suggests that the increase of basicity produces negative effects on the borate network and Si-O-Si structure containing bridging oxygen, and reduces the degree of polymerization of the system.

Acknowledgement

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References


