Mechanical Properties and Pore Structure Characteristics of Magnesium Oxychloride Cement Modified by Highland Barley Straw Ash

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Abstract: Highland barley straw ash contains a large amount of silica, and the ash calcined and ground under certain conditions has a higher pozzolanic effect. In order to study the effect of HBSA added into magnesium oxychloride cement mortar (MOCM) on the mechanical properties and pore structure, the activity of highland barley straw ash was studied firstly through the macroscopic mechanical properties test. Nuclear magnetic resonance (NMR) and Brunner–Emmet–Teller (BET) were used to test the distribution of full pore and micropore for MOCM respectively. The microstructure of MOCM was characterized by scanning electron microscope (SEM). The results illustrate that the highest activity of HBSA was obtained by calcining at 600°C for 2h and grinding for 2h. The addition of HBSA has a significant effect on the mechanical properties and pore diameter distribution of MOCM. A large amount of M-S-H gel was generated in MOCM added with 10% HBSA content, and had a consequence of decreased proportion of larger pores and the increased proportion of micropores as well as the better mechanical properties and pore structure.

Keywords: Magnesium oxychloride cement; Highland barley straw ash; Mechanical properties; Pore structure; Microstructure

1 Introduction

Magnesium oxychloride cement is a new type of ecological building material, which is mixed by MgO, MgCl2 and H2O in a certain proportion[1-3]. There is a wide application prospect in the engineering field due to the advantages as high early strength, good brine corrosion resistance, and good wear resistance, etc.[4,5]. However, the characteristics of poor water resistance and corroded steel bar limit its further application[6,7]. The main reason of poor water resistance is that
the 5 and 3 phases of magnesium oxychloride cement products are unstable phases, which could be converted into bischofite under certain conditions and make the structure become loose and porous[8-10].

It has been found that rice husk ash[11-13], fly ash[14-16] and wheat straw ash[17,18], etc. can improve the pore structure and mechanical properties of magnesium oxychloride cement. It has been studied that adding rice husk ash into MOC might be able to obtain higher density and better mechanical properties, and reduce water absorption and improve durability[19]. According to Guo et al., the incorporation of 30% fly ash in MOC could optimize the pore structure, make the structure more compact, and significantly improve the mechanical properties and water resistance[20]. Recent research has suggested that the cement-based material mixed with 15% wheat straw ash has smaller porosity and higher mechanical properties[21]. Wang et al. hold the view that nano silica is beneficial to improve the early compressive strength of cement, and the addition of fly ash is beneficial to improve the late compressive strength. The synergistic effect of the nanomaterials and the active mineral admixture can optimize the volcanic ash effect, reduce the porosity and improve the pore structure[22]. Tan et al. found that silica fume could significantly improve the pore structure of basic magnesium sulfate cement and the proportion of gel pores and small pores might be increased by adding silica fume. M-S-H gel with talc structure was formed due to the secondary hydration reaction between silica fume and magnesium hydroxide, which facilitates the formation of interconnected and dense microstructure[23].

From the above researches, the findings from these studies suggest that the main reason why the admixtures above could be able to improve the pore structure and mechanical performance is that they all contain active silica. The reactive silica reacts with the hydration products of the cement-based material to form new products that densifies its pore structure[24-27]. Similar to rice husk ash, fly ash and wheat straw ash, highland barley straw contains a large amount of ash, and the content of silica in the ash is high[28], which makes it possible to be used as cement-based auxiliary cementitious materials. However, these results would seem to suggest that there are few reports on the activity of highland barley straw ash and the effect of its incorporation into magnesium oxychloride cement on the mechanical properties and pore structure.
Therefore, there is great interest to study the effect of HBSA on the mechanical properties and pore structure of MOCM.

In this paper, the activity of highland barley straw ash (HBSA) was analyzed by mechanics performance testing. The HBSA was added into magnesium oxychloride cement mortar (MOCM) in different proportions, the effect on mechanical properties was studied. Its pore structure was analyzed and compared by NMR and BET. The microstructure of the MOCM was characterized and the influence mechanism was further revealed.

2 Materials and Methods

2.1 Materials

The raw materials of MOCM are composed largely of light burning magnesium oxide (MgO), industrial magnesium chloride (MgCl₂·6H₂O), sand, water, a water reducing agent, and water reducer. Light burning magnesium oxide is produced by the Magnesium Chloride Plant of Qarhan Salt Lake in Golmud City, Qinghai Province. The amount of MgO is accounted for 98% of light burning magnesium oxide and the amount of active MgO is 62.4%. The chemical composition is shown in Table 1. Industrial magnesium chloride is produced by Qarhan Salt Lake Magnesium Chloride Factory in Golmud City, Qinghai Province. 96% of the industrial magnesium chloride are MgCl₂·6H₂O, and its chemical composition is shown in Table 2. Sand with a particle size less than 4.75mm was obtained from Guide River in Qinghai Province, it is graded well and adopts tap water mixing, which meets the requirements of the concrete mixing water standard. The water reducer adopts a UNF series high efficiency water reducer, with an efficiency of 21%. The water repellent is phosphoric acid, which is produced by Shuangshuang Chemical Co., Ltd., Yantai City, Shandong Province. The proportion of H₃PO₄ is at least 85% in the water repellent, and the chromaticity Hc unit is at most 25% of that. The highland barley straw was obtained from the Nanmenxia area of Huzhu County, Qinghai Province. After weeds and other impurities were removed, it was burned to ash in an outdoor natural environment. The burning point is approximately 300°C, and the burning time was approximately 3h. After soil, sand, gravel, and other impurities were removed, the ash from the burnt highland barley straw, the integrated SX2-12-10A intelligent box muffle furnace was used for secondary calcination under laboratory conditions. The calcination times were 2, 5, and 8h at temperatures of 500, 600, and 700°C, respectively. The second-calcined HBSA was ground in a roller ball mill for 0.5, 1, and 2h, respectively. HBSA was calcined at 500°C, 600°C and 700°C for 5h, respectively, and the chemical composition was measured by X-ray fluorescence method, as shown in Table 1.

Table 1 Chemical composition of HBSA at different calcination time of 600°C (wt%)
According to the previous research results [29], the mixture ratio of magnesium oxychloride cement mortar is shown in Table 2. To study the HBSA’s activity, it was prepared under the following conditions: calcination temperatures (500, 600, 700°C); calcination time (2, 5, 8h), and grinding time (0.5, 1, 2h). According to the mass ratio of magnesium oxide, 15% HBSA was added directly into the MOC mortar mixture. When the mixture was stirred evenly, it was loaded into a 40mm×40mm×160mm cast iron mold for casting, vibration, and compaction. Then, it was cured in the air and demolded after 24 hours, and continued to cure in the air until 28 days. Each group consisted of 3 pieces, with a total of 28 groups and 84 pieces. One group of specimens without HBSA were included. The specimens numbered HBSA-0-0-0 refer to MOCM specimens without incorporation of HBSA, and the specimen numbered HBSA-500-2-0.5 refers to the MOCM specimen containing HBSA after calcination for 2h at 500°C and grinding for 0.5h, etc.

### Table 2 The mixture ratio of MOCM added with HBSA (kg/m³)

| MgO   | MgCl₂ | Sand | Water reducer | Water Repellent | Water | HBSA |
|-------|-------|------|---------------|-----------------|-------|------|
| 135.59| 147.81| 625  | 16.02         | 4.58            | 0%    | 0    |
| 141.14| 16.02 | 4.58 |
| 146.69| 19.45 |
| 152.24| 38.90 |
| 157.79| 58.34 |
| 168.79| 77.79 |
| 168.79| 116.69|

In order to study the effect of HBSA incorporation into MOCM on water resistance, HBSA which prepared under the highest activity conditions was added into MOCM admixture with the mass ratio of 0%, 5%, 10%, 15%, 20% and 30% of MgO respectively. HBSA was added into MOCM directly rather than the equal proportional replacement of MgO. MOCM specimens were prepared to test water resistance according to the ratio of raw materials in the mixture ratio in Table 1. The prism test blocks of 40mm×40mm×160mm were also poured, with 3 pieces in each group, a total of 12 groups and 36 pieces. After curing under indoor natural conditions for 24h, the mold is removed and the natural curing is continued for
28d. Among them, 6 groups of specimens were cured for 28d and then soaked in water for 28d to make them fully hydrated.

2.3 Test Methods

2.3.1 Mechanical Properties

To study the HBSA’s activity, 84 pieces of MOCM specimens added HBSA which prepared in 28 kinds of conditions were tested for mechanical properties. After curing naturally for 28d, the flexural and compressive strength tests were carried out on the specimens by using the Hualong cement mortar flexural and compressive testing machine. The prismatic specimen was tested for flexural strength firstly, and then the compressive strength was tested after breaking. Each group of test blocks’ mean value was taken as the strength test result.

The MOCM was mixed with HBSA under the optimal preparation conditions. At each dosage, the flexural strength and compressive strength of one group of test blocks were tested after natural curing for 28d. The other one group of test blocks were soaked in water for 28 days and taken out. After wiping the water on the surface, the flexural strength and compressive strength were tested immediately. The softening coefficient φ is calculated according to the following formula [30]:

\[ \phi = \frac{f_w}{f_d} \]  

in which \( f_w \) is the compressive strength of MOCM specimen after soaked in water for 28 days, and \( f_d \) is the compressive strength of MOCM specimen after natural curing for 28d.

2.3.2 Aperture Test

NMR technology could measure the real pore size distribution in the sample more accurately because it does not destroy the pores during the testing process[31]. NMR technology are widely used to calculate the volume of pores in porous media by measuring the mass and density of water in the sample to obtain such parameters as porosity and pore size distribution. By testing the signal of an H atom in the sample, a greater number of H atoms indicates more water contained, i.e., the proportion of the pore is higher where water is under this signal[32]. A 40mm×40mm×40mm cube block was intercepted from the MOCM specimen for NMR test, and the effective aperture testing range was 2nm–1mm. The pore size distribution could be calculated by the transverse relaxation time T2 distribution as follows[33]:

\[ \frac{1}{T_2} = \rho_2 \frac{S}{V} \]  

in which \( T_2 \) is the medium’s transverse relaxation time, ms; \( S \) is the pore’s surface area, \( \text{um}^2 \); \( V \) is the pore’s volume, \( \text{um}^3 \); \( \rho_2 \) is \( T_2 \)’s surface relaxation ratio, \( \text{um/\text{ms}} \).

2.3.3 Micro Measurement
An automatic BET specific surface area analyzer was used to test the micro-pore size distribution of MOCM. The effective pore diameter test range was 2nm~100nm, and the adsorption and desorption medium were nitrogen. The test sample is taken from the crushed block after the compressive strength test, and the length of the side is not more than 5mm. Field emission scanning electron microscopy (SEM) was used to test the micromorphology of MOCM samples. The test samples were taken from the flakes crushed by the compressive strength test, and the microcosmic scanning was carried out after gold spraying. The BET and SEM test samples were soaked in alcohol for 24h before the test to terminate further hydration, and then taken out and dried before the test.

3 Results and discussion

3.1 HBSA Activity

From the chemical composition of HBSA, it can be seen that the content of silica is about 60%, which is a potential pozzolan materials. The compressive strength test results of MOCM added with HBSA which prepared under the above 28 conditions, as shown in Fig.1. The calculated results of activity index are shown in Table 3. It can be seen that the compressive strength of MOCM with HBSA calcined at 600℃ is higher than that of 500℃ and 700℃, and the compressive strength of MOCM at 700℃ is the lowest. The results indicated that the activity calcined at 600℃ was the highest, followed by that at 500℃, and the activity calcined at 700℃ was the lowest. When the calcination temperature is 500℃ and 600℃, the compressive strength of MOCM is the highest calcined at 2h, followed by that at 5h and the lowest calcined at 8h, indicating that the activity of HBSA is the highest calcined at 2h. While the calcination temperature is 700℃, the compressive strength is the highest calcined at 8h, followed by that at 5h, and the lowest calcined at 2h. This is because that a large amount of SiO₂ exists in the form of crystal calcined at 700℃, and the content of active SiO₂ is less. At this time, the particle filling effect of HBSA was mainly worked in MOCM, rather than the active effect. Therefore, when the calcination temperature is 700℃, the longer the calcination time is, the more crystalline SiO₂ content is, and the better the filling effect is. When the calcination temperature is 500℃, 600℃ and 700℃, and the calcination time is 2h, 5h, and 8h, the compressive strength increases with the growth of the grinding time. The strength under 2h of grinding time is the highest, followed by that for 1h and the lowest grinding for 0.5h. The results indicate that the incorporation of HBSA after grinding for 2 hours into MOCM could effectively improve its activity and filling effect. The compressive strength of MOCM without HBSA is 49.7MPa after curing for 28d, while the compressive strength of MOCM with 15% HBSA content
calcined at 600 °C for 2 h and ground for 2h is 51.2 MPa and the activity index reach to 1.03. According to the test results of macroscopic mechanical properties, the activity effect of prepared HBSA is the highest when the calcination temperature is 600°C, the calcination time is 2h, and the grinding time is 2h.

Fig. 1 Compressive strength of MOCM mixed with HBSA under different conditions (a) 500°C; (b) 600°C; (c) 700°C (fc is the compressive strength; t1 is Calcination time; t2 is grinding time)

Table 3 Activity index of MOCM mixed with HBSA under different conditions

| No. | Specimen number | Activity index | No. | Specimen number | Activity index |
|-----|----------------|----------------|-----|----------------|----------------|
| 1   | HBSA-0-0-0      | 1.00           | 15  | HBSA-600-5-1    | 0.88           |
| 2   | HBSA-500-2-0.5  | 0.79           | 16  | HBSA-600-5-2    | 0.92           |
| 3   | HBSA-500-2-1    | 0.89           | 17  | HBSA-600-8-0.5  | 0.72           |
| 4   | HBSA-500-2-2    | 0.92           | 18  | HBSA-600-8-1    | 0.83           |
| 5   | HBSA-500-5-0.5  | 0.74           | 19  | HBSA-600-8-2    | 0.86           |
| 6   | HBSA-500-5-1    | 0.73           | 20  | HBSA-700-2-0.5  | 0.67           |
The HBSA obtained by calcining at 600°C for 2h and grinding for 2h was added into MOCM with 0%, 5%, 10%, 15%, 20%, and 30% content, respectively. The specimens with natural curing for 28d and soaking in water for 28d were tested for flexural strength and compressive strength, respectively, and the results were shown in Fig.2. As can be seen from Fig.2(a), the trend of the compressive strength under dry and saturated conditions both increased firstly and then reduced with the increase of HBSA content. When the content of HBSA was 5%, the compressive strength was both the highest in dry and saturated conditions. The compressive strength in dry state increased by 26.56% and that in saturated state increased by 28.57% compared with that without HBSA. When the content of HBSA was less than 10%, the compressive strength was higher in dry and saturated conditions. When the HBSA content was more than 15%, the compressive strength under saturated condition was lower than that without HBSA. It can be seen from Fig.2(b), with the increase of HBSA content, the softening coefficient of MOCM increased firstly and then decreased. When the HBSA content was 10%, the softening coefficient was the largest, which rose by 6.80% compared with that without HBSA. Therefore, MOCM has the best water resistance with the 10% HBSA content. When the HBSA content was less than 10%, the water resistance was enhanced to a certain extent. When the content of HBSA was more than 15%, the softening coefficient decreased with the increase of the content of HBSA, and was all lower than that without HBSA. In conclusion, the addition of HBSA into MOCM could effectively improve the mechanical properties, but too much HBSA content is likely to decrease it. The optimum content of HBSA for improving the compressive strength and the water resistance might be 5% and 10%, respectively.

### 3.2 Mechanical Properties of MOCM

|    | HBSA-500-5-2 | 0.81 | 21 | HBSA-700-2-1 | 0.68 |
|----|--------------|------|----|--------------|------|
| 8  | HBSA-500-8-0.5 | 0.73 | 22 | HBSA-700-2-2 | 0.76 |
| 9  | HBSA-500-8-1  | 0.77 | 23 | HBSA-700-5-0.5 | 0.75 |
| 10 | HBSA-500-8-2  | 0.77 | 24 | HBSA-700-5-1  | 0.75 |
| 11 | HBSA-600-2-0.5 | 0.90 | 25 | HBSA-700-5-2  | 0.83 |
| 12 | HBSA-600-2-1  | 0.95 | 26 | HBSA-700-8-0.5 | 0.77 |
| 13 | HBSA-600-2-2  | 1.03 | 27 | HBSA-700-8-1  | 0.86 |
| 14 | HBSA-600-5-0.5 | 0.86 | 28 | HBSA-700-8-2  | 0.86 |
Fig.2 Mechanical properties of MOCM with different HBSA content (a) compressive strength; (b) softening coefficient

3.3 Pore Structure Characteristics

3.3.1 NMR Analysis

Fig.3 shows the pore diameter distribution of MOCM specimens with HBSA content of 0%, 5% and 10% respectively by NMR test. The pore diameter distribution curves under the three content of HBSA are composed of three peaks. The first peak mainly represents the change of micro pores less than 0.2um, the second peak represents small pores of 0.2um ~ 10um, and the third peak represents large pores larger than 10um. The pore diameter distribution of MOCM is mainly concentrated in the micropores less than 0.2um, as well as some small pores and large pores. With the increase of HBSA content, the pore diameter distribution of micro pores changes obviously, and the first peak moved to the left. It shows that new micropores are formed with the increase of HBSA content, and the most probable aperture becomes smaller and the proportion is higher. When the HBSA content was 5% and 10%, the most probable aperture was smaller and the number of micropores less than 0.02um was more than that of without HBSA. It can be seen from the second peak that the pore diameter of MOCM specimens with 10% HBSA content are concentrated in the small pore range less than those with HBSA content of 0% and 5%. As shown from the third peak, the most probable aperture of MOCM specimen with HBSA content of 10% is less than that with HBSA content of 0% and 5%. When the HBSA content increases to 10%, the most probable aperture is the smallest, forming more and more new small-size pore, and the proportion of micropores was the highest.
The statistics of pore diameter distribution in each interval of MOCM specimens with different HBSA content are shown in Table 4. Wang et al. set up the division of pore diameter interval[22]. It can be seen from Table 3, the proportion of the harmless pores and the less harmful pores whose diameter is less than 0.05um increases gradually, while the proportion of the harmful pores and the more harmful pores whose diameter is more than 0.05um decrease by degrees with the increase of HBSA content. When the HBSA content was 10%, the proportion of harmful pores and more harmful pores decreased by 25.11% compared with that without HBSA, and the reduced pores were transformed into harmless pores and less harmful pores. The number of harmless pores less than 0.02um increased by 3.71 times, and the number of less harmful pores increased by 1.65 times. It indicates that HBSA added in MOCM could transform part of harmful and more harmful pores into harmless and less harmful pores, which make the pore structure more favorable.

Table 4 The pore diameter distribution statistics of MOCM with different HBSA content (%)

| HBSA content | <20nm | 20～50nm | 50～200nm | >200nm |
|--------------|-------|----------|-----------|--------|
| 0%           | 3.78  | 22.69    | 66.52     | 7.01   |
| 5%           | 6.06  | 29.46    | 58.84     | 5.64   |
| 10%          | 14.04 | 37.54    | 46.06     | 2.36   |

3.3.2 BET Analysis

According to the pore diameter distribution results of NMR test, most of the pore in MOCM are concentrated in the pores below 100 nm. For the furthering study of the pore diameter distribution within the range of 100nm, BET was used to test the mesopore structure of MOCM. The adsorption and desorption isotherms are the basis for analyzing the specific surface area and pore diameter distribution of MOCM[34,35]. MOCM’s adsorption and desorption
isotherms with different HBSA content were obtained by nitrogen adsorption method, as shown in Fig.4. The MOCM’s adsorption and desorption isotherms with the three HBSA content are basically the same, including initial, transition and end segment. At the initial segment, the slope of the curve gradually increases with the grow of HBSA content, which illustrates that the number of micropores in MOCM gradually increases[36]. Then, the adsorption and desorption isotherm are separated at transition segment. When the HBSA content increases to 10%, the relative pressure value corresponding to the separation point is the smallest, which demonstrates that the internal pore diameter is the smallest. The adsorption and desorption isotherms overlap again at the end segment. The curve slope of the adsorption and desorption isotherm of MOCM with 5% and 10% HBSA content was higher than that with 0% HBSA content at the end segment.

![Fig.4 Adsorption and desorption isotherms of MOCM specimens with different HBSA content(a)0%;(b)5%;(c)10%](image)

The specific surface area of MOCM was obtained by BET multi-point test. The data points in the coordinate were linearly fitted to obtain the multi-point BET two-bit linear graph[37], as shown in Fig.5. The slope and intercept of the two-bit line were obtained by linear fitting. According to the BET test principle[38,39], the area of each milliliter of nitrogen molecules diffused into a monolayer is 4.354m$^2$. The specific surface area obtained through the BET method is shown in Table 5. It can be seen that the specific surface area of MOCM gradually increases with the rise of HBSA content, indicating that the number of micro pores inside MOCM increases correspondingly. When HBSA content was 10%, the specific surface area increased by 32.75% compared with that without HBSA, and the increased specific surface area was caused by the increase of internal micropores. Therefore, these results suggest that the addition of HBSA could form a part of micropores in MOCM, and the number of micropores is the largest when the HBSA content is 10%.
The distribution of pores diameter through the BET method is shown in Fig. 6. As can be seen from Figure 6(a) and (b), the variation rule of pore diameter distribution of MOCM with different HBSA content is basically the same, but the pore diameter proportion of MOCM with different HBSA content and the most probable aperture are obviously different. When the HBSA content is 10%, the most probable aperture is 3.730nm in the micropores range less than 20nm, and it is less than 3.831nm and 3.836nm when the HBSA content was 5% and 0%, respectively. The most probable aperture is 54.929nm in the mesoporous range of 20nm-100nm, which is less than 76.302nm and 77.459nm when the HBSA content is 5% and 0%, respectively. It can be seen that when the HBSA content is 5% and 10%, the growth rate of cumulative pore volume is significantly faster than that HBSA content is 0%. A possible explanation for these results may be that MOCM with the HBSA content of 5% and 10% generated more micropores.
The variation relationship of MOCM’s pore volume and surface area with different HBSA content is shown in Fig.7. It can be seen that MOCM’s micropores with 5% HBSA content have the largest cumulative volume, while MOCM’s micropores with 10% HBSA content have the largest cumulative specific surface area. However, the cumulative volume of MOCM’s micropores with HBSA content of 10% was less than that with HBSA content of 5%. Large surface area and small volume indicate a greater number of micropores and a smaller pore diameter. MOCM with HBSA content of 0% had the smallest proportion of micropores. In addition, it can be seen from the fig.3 that when the HBSA content was 5% and 10%, the cumulative pore volume growth rate was significantly faster than that when the HBSA content was 0%, indicating that MOCM with the HBSA content of 5% and 10% generated more micropores. The pore volume and surface area basically do not increase when the pore size is around 90nm, because the effective test pore size range of BET is within 100nm.
Fig. 7 MOCM with different HBSA content for (a) cumulative pore volume; (b) cumulative pore surface area

3.4 Microstructure

The microtopography of MOCM samples with HBSA content of 0%, 5% and 10% are shown in Fig. 8 (a), (b) and (c). When the HBSA content is 0%, the micromorphology of MOCM is mainly composed of 5-phase needle-like crystal and a small amount of M-S-H gel. When the HBSA content is 5% and 10%, the micromorphology shows that a lot of M-S-H gel is generated. The M-S-H gel generated in the MOCM with 10% HBSA content, which adhere uniformly to the 5-phase crystal and solid particle surface, filling a large number of crystal gaps and solid particle gaps, and eventually form in a more compact structure. When the HBSA content is 10%, the 5-phase crystal of needle-like is uniformly wrapped by M-S-H gel, forming the copolymer of needle-like crystal and gel. Some of the larger pores are transformed into micro pores due to the filling of the gel. This plays a positive and effective role in the pore diameter distribution of the pore structure, which is consistent with the results of NMR and BET test. Therefore, the fundamental reason for the change of pore structure is that the active silica in HBSA reacts with the hydration product from MOCM in a secondary hydration reaction to generate M-S-H gel. Some larger pores are filled with gel, forming a denser structure. In this work, HBSA was directly added into MCOM without replacing MgO, so that the molar ratio of MgO to MgCl2 remained unchanged at 6.25. Under the condition of magnesium surplus, hydration products are mainly 5-phase and magnesium hydroxide[40]. The basic process of hydration reaction is as follows:

The first hydration process: \[ 5\text{MgO} + \text{MgCl}_2 \cdot 6\text{H}_2\text{O} + 3\text{H}_2\text{O} \rightarrow 5\text{Mg(OH)}_2 \cdot \text{MgCl}_2 \cdot 8\text{H}_2\text{O} \]

\[ \text{MgO} + \text{H}_2\text{O} \rightarrow \text{Mg(OH)}_2 \]

The second hydration process: \[ 3\text{Mg(OH)}_2 + 2\text{SiO}_2 \rightarrow \text{Mg}_3\text{Si}_2\text{O}_5(\text{OH})_4 + \text{H}_2\text{O} \] (Serpentine)

\[ 3\text{Mg(OH)}_2 + 4\text{SiO}_2 \rightarrow \text{Mg}_5\text{Si}_4\text{O}_{10}(\text{OH})_2 + 2\text{H}_2\text{O} \] (Talcum)

\[ 8\text{Mg(OH)}_2 + 12\text{SiO}_2 + 8\text{H}_2\text{O} \rightarrow \text{Mg}_6(\text{H}_2\text{O})_4(\text{Si}_{16}\text{O}_{16})_2(\text{OH})_4 \cdot 8\text{H}_2\text{O} \] (Serpolite)

During the secondary hydration reaction, the values of x and y determine the structural form of M-S-H. The general M-S-H structure mainly includes sepiolite, talcum and serpentine[26,41]. When the content of HBSA is 10%, and the content of SiO2 in HBSA is about 60%, that is, the Mg/Si molar ratio is about 1.0. At this point, after the hydration reaction of 28 days, there was almost no residual silica and brucite in the hydration products, and all reacted to form M-S-H gel [42]. The main structural forms of M-S-H gel are talc and serpentine, forming layered silicate structure, which adheres to the surface of 5-phase crystal and solid particles. Therefore, the mass generation of M-S-H improves the mechanical properties of MOCM and optimizes the pore structure.
4 Conclusion

(1) The activity effect of HBSA obtained from highland barley straw is the highest after primary incineration of ash, secondary calcination at 600°C for 2h and grinding for 2h.

(2) The addition of HBSA into magnesium oxychloride cement mortar could significantly improve its mechanical properties. When MOCM is added with 5% HBSA, the compressive strength in dry state increases by 26.56% and that in saturated state increases by 28.57%. When the HBSA content increases to 10%, the water resistance of MOCM increases by 6.80%.

(3) The pore structure of MOCM is significantly different with different HBSA content, and the incorporation of HBSA make the proportion of micropores in MOCM higher and the pore diameter smaller. When the HBSA content is 10%, the number of harmful and more harmful pores decreases by 25.11% compared with that without HBSA. The number of harmless pores increases by 3.71 times, and the number of less harmful pores increases by 1.65 times. With the increase of HBSA content, the specific surface area of MOCM rises, and the most probable aperture decreases.

(4) The active silica in HBSA reacts with the hydration product of MOCM to form M-S-H gel, which fills part of the pores. Thus, some larger pores are transformed into smaller gel pores, and improve the pore structure. When the HBSA content is 10%, the most M-S-H is generated and the pore structure is the best.

Author Declarations
Ethics approval and consent to participate  Not applicable.

Consent for publication  Not applicable.

Availability of data and materials  All data generated or analyzed during this study are included in this published article.

Competing interests  The authors declare that they have no competing interests.

Funding  This work was financially supported by the National Natural Science Foundation of China (Nos. 51468039, 51868044). Authors appreciate the financial and equipment supports.

Authors' contributions  Feng Cao- Investigation, Methodology, Writing - original draft, Writing - review & editing. Hongxia Qiao- Supervision, Writing - review & editing. Penghui Wang- Writing - review & editing. Weijia Li- Writing - review & editing.

Acknowledgements  We appreciated Chongyan Qian and Jiaju Feng for their contribution in providing and collecting the raw materials of highland barley straw for this study.

Compliance with Ethical Standards

Disclosure of potential conflicts of interest  The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Research involving Human Participants and/or Animals  Not applicable.

Informed consent  Not applicable.

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