Characterization and Modeling of Hydration and Microstructure Formation for Blast Furnace Slag Cement

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ABSTRACT

Degree of hydration of cement minerals and BFS, amount of hydration products of pastes were determined by XRD/Rietveld method. The hydration mechanism of slag was also investigated, considering the role of calcium hydroxide as activator and the Ca/Si ratio of C-S-H. Assumed that for low Ca/Si ratio, the C-S-H inner product resists strongly against ion diffusion and thus the hydration of BSF retards, an enhanced model for slag hydration was proposed. The validity of enhanced model is verified by both hydration degree and heat generation tests. The capillary porosity of BFS blended cement pastes is larger at lower hydration degree, and remarkably decreases as its hydration progresses. Gel water in BFS paste increase nonlinearly with the hydration degree of BSF. This implies that C-S-H produced from BFS hydration at later age has larger amount of gel water and becomes lower density, can fills in large pores more efficiently. Enhanced modeling of pore structure taking into account of these new findings could reproduce the experimental results of porosity and higher long-term strength development of BSF paste even though the hydration degree of BSF was lower than OPC.

INTRODUCTION

Realizing of sustainable society is a major driver for future innovations and prospects for development of construction industry. The main problem for the construction industry is that the global average CO₂ emission per ton of cement manufactured, which is mainly due to the decarbonation of limestone, is estimated to be about 0.83 tons [Gielen and Tanaka 2007]. It is actually quite low compared to most other manufactured products. For example, compare it to steel, the global average CO₂ emission per ton of steel manufactured rises up to about 3 tons of CO₂, or for aluminum, it rises up to about 15 ton/ton of product [Gaetner 2011]. However, global cement production is estimated around 3.6 billion tons of cement were produced worldwide in 2011 [Gielen and Tanaka 2007], per year, and continues to increase due to mega-urbanization and subsequent infrastructure development all over the world, predicted to rise to 5.8 billion tons by 2050 [Gartner 2009]. Thus, simply because of the enormous volumes of concrete produced globally every year, it is evident that any positive action that the cement manufacturing industry could take to reduce its specific CO₂ emissions would contribute to reducing overall global Green House Gas
emissions. If we are to tackle this perceived problem, energy efficient ways must be found to produce hydraulic binders that emit significantly less fossil CO₂ during manufacture.

The most promising way to meet this rising demand is to continue increasing the use of supplementary cementitious materials (SCMs) such as fly ash, ground granulated blast furnace slag, silica fume and natural pozzolan. The practice of using SCMs is increasing, with the world average percent clinker in cement having decreased from 85% in 2003 to 77% in 2010, and it is projected to further decrease to 71% in the future [Gielen and Tanaka 2007]. While the replacement Portland cement clinker with SCMs is often driven by the pressure that comes from the global environmental issues to reduce CO₂ emissions from concrete, the use of SCMs in concrete remains the relatively small amounts (5–20% replacement of clinker) by economics and improvements in the long-term mechanical properties and durability of concrete, since the high volume clinker replacements result in losses in performance at early ages, and larger drying and autogenous shrinkage that tends to induce cracking and decrease durability. The use of SCMs in concrete. On the other hand, many past studies shows that concrete made with slag blended cement has many advantages, including better chloride resistance, lower permeability and higher strength at later age. From the viewpoint of circular economy and sustainable development, blast furnace slag is a promising mineral admixture in the concrete industry.

A key issue in materials design based on materials science is to establish quantitative relationships between processing, structure, and properties. It is broadly agree that for ordinary Portland cement, higher strength usually derives from a higher degree of hydration. However, when we switch to blast furnace blended cement (BSF) from Portland cement (PC), it seems insufficient to attribute strength development only to the hydration process. For instance, it is well known that under room temperature when PC is replaced partially by BFS, strength decreases at an early age while higher strength can be achieved at a later age. On the other hand, researchers [Luke 1987, Lumley 1996, Demoullian1980, Battagin 1992, Sakai 2004] indicate that the hydration degree of slag ranges from 30% to 70% after long time curing such as 1–2 years, which is lower than that of PC. Obviously BFS gains higher strength than PC at a later age, even if the hydration degree is lower than that of PC. Microscopic observation [Regourd 1987, Escalant-Garcia 2001, Tan 1998] by Scanning Electron Microscopy (SEM) indicates that the BFS matrix with full curing has a denser structure than PC, and porosity analysis such as Mercury Intrusion Porosimetry (MIP) [Feldman 1981, Feng 1989, Aldea 2000, Kwak 2004] also shows that at a later age the pore distribution curve moves towards a much finer diameter. Therefore, dense pore-structure can be considered to contribute more than hydration for the strength of BFS at later ages. Actually, other properties of BFS that differ from PC may be also related to the pore-structure. The moisture loss of BFS under drying conditions is reported to be less than that of PC [Dan 2009, Hogan 1981], which can be explained by finer pores and stronger retention of water. It is also reported that drying shrinkage of BFS is larger than PC, especially for those cases with sufficient curing [Brooks 1992, Li 20011, Chern 1989]. This larger shrinkage deformation can be attributed to the finer pore structure, because higher capillary tension force may be induced in the paste. Beside, BFS concrete is more sensitive to the curing conditions. Insufficient water supply during curing may retard the hydration process and cause a significant decrease in the long-time properties. Therefore, for the purpose of promoting the efficient application of BSF, it is of great necessity proper modeling and simulation of its hydration and subsequent microstructure
formation are necessary in order to evaluate or predict the macro-properties of BFS concrete based on micro-information.

This paper summarizes the advances in our studies (Sagawa 2006a, 2006b, 2010, Ishida 2011, Luan 2012) on characterization and modeling of hydration and pore structure formation achieved in the last decade to improve the material design for BFS concrete. The hydration of BSF and pore structure of the BFS matrix can be properly simulated. Furthermore, based on the correct information on pore-structure, prediction of the macro properties of BFS concrete such as thermal temperature rise and strength development could also be improved.

CHARACTERIZATION FOR HYDRATION AND MICROSTRUCTURE FOR BSF

The accurate measurement of the degree of reaction of SCMs in hydrating blended cements is necessary to unravel the parameters governing SCM hydration. Several different methods have been proposed and applied, but all previously proposed methods have more or less serious limitations. The selective dissolution approaches are unreliable as they show large, unrealistic discrepancies with other methods and problems with inter-laboratory reproducibility [Ohsawa 1985, Luke 1987, Ben Haha 2010, Kocaba 2012]. Application of image analysis to electron microscopy images and element maps has shown promising results for SCMs [Kocaba 2012, Weerdt 2011, Deschner 2013]. However, the proposed approaches are time-demanding and cannot resolve the reaction of particles smaller than about 2 μm, rendering the method unsuitable for fine materials such as silica fume and metakaolin. A more generally applicable method makes use of $^{29}$Si and $^{27}$Al solid-state NMR spectroscopy. The method has been successfully applied to SCMs [Saout 2011,Hjorth 1988, Lothenbach 2012]. However, widespread use will remain limited because access to solid-state NMR equipment is restricted. $^{29}$Si NMR data acquisition is time-consuming (1–2 days per measurement) and measurements are restricted to systems with low levels of paramagnetic ions. XRD is the most prominent analytical techniques in the characterization of Portland cement based materials because of ease and speed of measurement and its accuracy. Full XRD profile fitting methods such as the Rietveld method can realize the more quantitative analysis of crystalline phases in unhydrated and hydrated cement has resolved crystal structures and can be quantified. It is well-known that the glass content of slag can be determined by thermal analysis: the first reversible exothermic peak in the temperature range 700–800 °C corresponds to the glass-transition temperature, the next two exothermic peaks with well-defined maxima in the range 925–1040 °C are attributed to the devitrification process. Using X-ray diffraction, these peaks were respectively assigned to merwinite (metastable phase) and melilite (solid solution gehlenite to akermanite) [Regourd 1980, Fredericci 2005]. It has been suggested that quantification of unhydrated BFS in hardened BFS blended cement can be determined by Rietveld method to measure the crystallized BFS.

Fig. 1 shows the degree of hydration of BFS and C₃S in cement. Hydration degree of finer BFS with around Blaine fineness of 6000 cm²/g (representing B6 in Fig.1, N, M and L is ordinary, moderate heat and low heat Portland cement) was higher than coarse BFS with fineness of 4000 cm²/g. Hence it can be seen that reactivity of BFS is higher as the particle size of BFS is small. In addition, hydration reactivity of BFS was strongly affected by the amount of C₃S in base Portland cement, hydration of BFS tended to
be delayed at smaller amount of $C_3S$ in base Portland cement. On the other hand, it is confirmed that the hydration of $C_3S$ in blended cement accelerated by the presence of BFS. This implies that BFS plays the role of precipitation site of C-S-H which is the hydrate of $C_3S$.

Fig. 1 Degree of hydration of BFS and $C_3S$ in BSF blended cement.

It is well agreed that the principle hydrates of slag are C-S-H gels. Compared with Portland cement, researchers found that as the slag ratio increases, C-S-H gels produced by slag show a lower Ca/Si ratio (Harrisson et al. 1987; Shi et al. 1992; Richardson and Groves 1992; Gollop and Taylor 1992, 1996), as illustrated in Fig. 2. The physical properties of C-S-H may vary due to the reduced Ca/Si ratio.

Fig. 2 Ca/Si ratio of C-S-H in hydrated Portland and BSF blended cements.

Physical water was categorized into capillary water and water inside gel grains, i.e. gel and interlayer water. In experiments, it is impossible to divide capillary, gel and interlayer water precisely. Hence the author and coworkers define the capillary, gel and interlayer water: the weight loss by drying at 40 °C for specimens with saturated condition was considered as the total capillary water. The remaining physical water was regarded as the sum of the gel and interlayer water. Then the specimens were exposed at
105 °C to remove all the gel and interlayer water. The weight loss from 40 °C to 105 °C was measured as gel and interlayer water.

Fig. 3 shows experimentally calculated capillary pore, and gel and interlayers pores volume, as a function of hydration degree of BSF. One can find the experimentalt data agree with theoretical value predicted by the Powers’ model [Powes 1960]: the upper line and bottom line show the total pore volume and capillary pore volume, respectively. Finally, the specimens were heated up to 1000 °C to remove chemically combined water. The weight percentage is defined as the lost water to remaining paste weight after heating at 1000 °C. The definition of the gel and interlayer water needs to be clarified. As the definition in the test, the specimens were dried at 40 °C to remove capillary pore water and then exposed at 105 °C to measure the gel and interlayer water loss. Actually, after drying at 40 °C, a little water still remained in capillary pores while some of the gel water evaporated. Therefore, strictly speaking, the gel and interlayer water measured do not coincide completely. However, since only a small part of the gel water was lost during drying at 40 °C, it can be reasonably assumed that the test result is approximately comparable with the computed value.

Fig. 3 Relation between hydration degree of cement and pore volume.

Fig. 4 illustrates the relationship between the Ca/Si ratio and the H/Si ratio of C-S-H, the relationship between the ratio of Ca/Si ratio and gel water in C-S-H, respectively. The chemical combined water decreased with the decrease of the Ca/Si ratio of C-S-H, In contrast, the gel and interlayer water in BSF paste showed a reverse tendency. It increased with the decrease of the Ca/Si ratio of C-S-H, while for PC paste it was constant deposite the Ca/Si ratio of C-S-H. Therefore, when one consider a sum of chemical combined water and gel and interlayer water as water content, the H/Si ratio is gradually decreased with the decrease of the Ca/Si ratio of C-S-H for PC paste, while for BFS paste the H/Si ratio increased from around 3 to around 6 as the Ca/Si ratio decresing. Here, we focus only on the gel and interlayer water, calculated as the mass percentage of gel water in C-S-H, which were arranged in relation to the Ca/Si ratio is the right view of Fig. 4. For the PC paste, the sum of gel and interlayer water was substantially constant around 15% regardless of the Ca/Si ratio, whereas that in BFS paste significantly increased as
decreasing the Ca/Si ratio. Since Ca/Si ration of C-S-H decreased with age, these results implies that the gel and interlayer water in C-S-H for BSF paste increase with the age.

Form the above-mentioned results, one can understand that the fraction of gel and interlayer water in C-S-H in PC paste is constant, while for BFS paste it was varied as hydration progressed. This can explain the fact that the capillary pore volume of BFS paste is larger than PC paste in an earlier age, but it decrease significantly as BSF hydration progresses, and at later age it became smaller than PC paste.

Fig. 4 Chemical combined H$_2$O to Si ratio (left), Chemical and gel and interlayer H$_2$O to Si ratio (middle) and gel and interlayer water content in C-S-H represented as a function of Ca/Si ratio of C-S-H.

ENHANCED PORE STRUCTURE MODEL FOR BSF AND ITS VERIFICATION

Otabe et al. (2005) is the strength development model which follows the concept proposed by Schiller (1958), in which it is assumed that the strength of concrete or mortar is principally related to capillary porosity development, because capillary pores are much larger than gel and interlayer pores and influence strength more significantly. As shown in Fig. 5, the model can predict the strength development of Portland cement (noted N in figure) accurately, while BSF (NB) had lower strength at an early age but reached higher strength at 91 d as hydration went on. This suggests that for BSF, more capillary pores should be filled by hydrates at later age.
From above discussion one can expect that when the degree of hydration is the same, the density of C-S-H produced due to BSF hydration may be smaller than that due to PC hydration. Fig. 6 shows a sum of the gel and interlayer water ratio as a function of the chemically combined water. The increase of chemically combined water reflects the hydration processes. For PC paste (noted N in figure), good agreement is obtained, but for BFS paste (NB), the analysis apparently underestimates gel and interlayer water development at a later age. Unlike the linear increase in N, all the BFS test data show a tendency toward nonlinear increase, with higher gel and interlayer water content gain at a later age. However, the analysis gives a linear increase for NB. Therefore it is rational to deduce that the current model is only applicable to PC rather than BFS due to inconsistent gel and interlayer pore water assessment. This inconsistency leads to underestimation of gel and interlayer porosity, and therefore overestimation of capillary porosity for BFS, which is also the cause of discrepancy of strength in Fig. 5. Hence, Ishida et al (20??) modified the pore-structure prediction model considering the impact of slag hydration on the gel and interlayer water. The results due to enhanced model are illustrated in Fig. 7. It is obvious that the enhanced model shows nonlinear increase and higher gel and interlayer water at a later age, which is consistent with the test results. Therefore, from the viewpoint of micro-physical properties of the matrix, the enhancement on the intrinsic porosity provides better simulation for BFS.

It is reported that C–S–H of BFS has a lower Ca/Si ratio [Talling 1989, Richardson 1992, Regourd 1983, Harrison 1986], which may contribute to gel grains and induce higher grain strength. Therefore, another modification for the ultimate strength of C-S-H is requested in order to estimate BSF matrix. The ultimate strength of C\textsubscript{3}S and C\textsubscript{2}S, slag are assumed to be 215 MPa, 250 MPa, and 260 MPa. As shown in Fig. 9, enhanced model agrees with the experimental data well: at the early age, the strength of BFS (NB) was less than PC (N), BFS tended to gain higher strength at later age as hydration progressed.
ENHANCED ING HYDRATION MODEL FOR BSF AND ITS VERIFICATION

Kishi (1994, 1997) and Maekawa et al. (2008) developed a multi-component hydration heat model based on the Arrhenius law of chemical reaction, in which parameters are temperature, available free water, CH content and slag fineness, the hydration heat rate under real condition is given by following the equation:

\[ H_{sg} = \min(\beta_{sg}, \lambda_{sg}) \cdot H_{sg,T_0}(Q_{sg}) \exp \left\{ -\frac{E_{sg}}{R} \left( \frac{1}{T} - \frac{1}{T_0} \right) \right\} \]  (1)

where \( H_{sg} \) is the hydration heat rate under real condition, \( H_{sg,T_0} \) is the reference heat rate at 293 K, \( Q_{sg} \) is the accumulated heat during hydration, \( E_{sg} \) is the activation energy of slag, \( R \) is the gas constant, \( T_0 \) is the reference temperature (293 K), and \( T \) is the real temperature. \( \beta_{sg} \) is the coefficient considering the effect of available free water, and \( \lambda_{sg} \) is the coefficient expressing hydration activation due to CH insufficiency. This model has been verified by adiabatic temperature rise tests for both Portland cement and slag blended cement concrete. On the other hand, verification of the hydration degree of slag has not been carried out, because of the scarcity of reliable data when the model was established. Fig. 6 shows the
comparison between experimental and analysis values of the hydration degree of BSF. It is obvious that extreme large differences between them exist.

Herein the hydration scheme of slag is discussed according to the basic information of activation (Tanaka et al. 1983; Taylor 1990). At the starting stage, due to the activation by Ca$^{2+}$, the protective film on the surface of the slag particle is destroyed. The internal structure, which is an anionic network composed of O, Si and other elements, starts to decompose, and silicate ions such as Si(OH)$_4$ are released into the solution. When silicate ions contact Ca$^{2+}$, a reaction occurs and hydrates start to precipitate. Like in the case of Portland cement, hydrates from slag can be divided into two categories, i.e. the inner product and the outer product. The inner product consists of hydrates generated inside the radius of the original particle and closely bond around an unhydrated core, while the outer product precipitates outside the range of the original particle and fills in capillary pores. In order to sustain hydration, it is necessary for silicate ions released from slag to diffuse outside the inner product, or alkali cations to diffuse inside. The inner product may show resistance against ion diffusion, i.e. as hydration goes on and the thickness of the inner product increases, the diffusivity of ions through the inner product becomes lower. Accordingly, the reaction comes slower or even stagnates. Therefore, the thickness of the inner product is a crucial factor that dominates the hydration rate of slag.

Furthermore, when we discuss the resistance effect of the inner product, not only its thickness but also the various other physical properties have to be taken into account. As discussed above, increasing the slag ratio varies the Ca/Si ratio of C-S-H, and accordingly one should consider the impact of physical properties of C-S-H on the hydration process from two aspects: First, since C-S-H with a lower Ca/Si ratio has a morphology that is foil-like rather than fibrillar, the tortuosity of gel pores becomes higher. This implies more twists and forks in pores, resulting in a much longer path for ions to diffuse through the inner product. The other factor may be the alternated electric charge on the surface of C-S-H. As indicated by Elakneswaran et al. (2009), pores with a diameter of less than 10 nm show the electrical double layer (EDL) effect. The inner product contains mainly gel pores and a large proportion of gel pores have a diameter smaller than 10 nm, and thus the EDL effect of the inner product is obvious. In the case of C-S-
H with a high Ca/Si ratio, the surface is positively charged. The repelling effect of the positive charge layer restrains the movement of Ca\(^{2+}\) towards the inner product. The principal ion movement is that of silicate ions that are released through the inner product. When the electric charges become less positive or even negative due to the lower Ca/Si ratio, the principal ion movement may change. Due to the absorbing effect, more Ca\(^{2+}\) ions move toward the inner product and react with silicate ions near it. The reverse ion movement implies that denser inner product having stronger resistance against ion diffusion is formed. Further ion diffusion may be blocked and the hydration rate afterwards is reduced significantly. The above discussion about the relationship between the hydration of the slag and inner product is illustrated in Fig. 5. Accordingly, the authors conclude that for a lower Ca/Si ratio due to both higher tortuosity and reversed electric charge, the inner product layer shows higher resistance to ion movement. This may be the main reason that the hydration degree decreases as the slag ratio increases.

![Diagram](image)

**Fig. 10 Influence of inner product on ion movement and hydration of slag**

From above discussion, it can be seen that the hydration process is closely associated with the inner C-S-H product because of its resistance to ion movement. This resistance depends on both the thickness and intrinsic properties of C-S-H. In the original model proposed by Kishi and Maekawa et al., although the thickness of the inner product is considered, the various intrinsic properties of C-S-H determined by the Ca/Si ratio are not taken into account. On the other hand, inefficiency coefficient of CH, \(\lambda_{sg}\), seems to be able to ignore since sufficient amount of CH exists in the paste even for high slag ratios. After all, all factors are integrated into coefficient \(\beta_{sg}\). Since the hydration degree is related to the fraction of free water \(\omega_{free}\), slag fineness \(s_{sg}\), and the effect of the inner product. The Ca/Si ratio of C-S-H shows a clear correlation with the slag ratio, and thus the slag ratio seems a proper index of the effect of the inner product. Further as the inner product becomes thicker, the ion diffusion is strongly resisted so the hydration rate decreases sharply rather than gradually. This fact request the exponential formula seems appropriate to describe the significant relationship between inner product thickness and the resistance effect. Therefore, the effect of the inner product is proposed by a new coefficient \(\chi\), defined as follows:
\[ \chi = \exp(-a \cdot \eta_{sg}^b) \quad (2) \]

where \( \eta_{sg} \) is the thickness of the inner product. \( a \) and \( b \) are intrinsic parameters that are the functions of the slag ratio. In fact, the value of parameter \( \chi \) depends on the intrinsic properties of C-S-H which depends on the Ca/Si ratio and its influence on ion movement. The value of parameter \( \chi \) decreases as the thickness of the inner product \( \eta_{sg} \) increases, which implicitly represents the resistance by the inner product to ion diffusion. Besides, for a higher slag ratio, the parameter \( \chi \) decreases more sharply from the beginning. This has clear physical meaning that the C-S-H inner product has a stronger retardation effect on ion movement the lower the Ca/Si ratio. Integrating \( \chi \) into \( \beta_{sg} \), it is applicable to the simulation of the development of the hydration degree for various slag ratios. Accordingly, in the enhanced model, the new expression for \( \beta_{sg} \) is given by:

\[
\beta_{sg} = \left\{ 1 - \exp \left\{ -r \left( \frac{\omega_{free}}{20.0} \right) \frac{1}{s_{sg}^2} \right\} \right\} \cdot \chi \quad (3)
\]

where \( r \) and \( s \) are the material constants.

The enhanced model is verified by experiments. As shown in Fig.11, the analytical results are much improved.

(a) Original model  
(b) Enhanced model

Fig. 11 Comparison of hydration degree based on enhanced model with original model: the water/binder ratios are 0.3, 0.4 and 0.5, slag replacement ratio is 50%, and slag powder with JIS fineness grade of 4000 is used.
CONCLUSIONS

The following general conclusions can be drawn from the study provided in the paper:

• Rietveld analysis for unhydrated and hydrated BFS sample crystallized BFS with heat treatment is available to determine the hydration degree of compounds in BSF.

• Enhanced pore structure model was proposed for the BFS matrix. As the degree of hydration increases, more gel and interlayer pores are gained and hydration product fills in the capillary pores more efficiently, resulting in higher strength in a later age. Verification by strength experiment showed good agreement.

• The hydration process of BSF in cement is investigated focusing on the intrinsic properties of the inner product. Since the Ca/Si ratio of C-S-H decreases as the slag ratio increases, resulting in both higher tortuosity and changed positive electric charge accumulation in C-S-H, it is assumed that the C-S-H inner product has a strong resistance to ions movement, and therefore the subsequent hydration tends to be retarded. Based on the above assumption, an enhanced model is proposed, taking into account the resistance effect. Finally, the enhanced model is applied to simulation of slag hydration, and the results indicate good agreement with the experiment data.

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REFERENCES


