Immobilization of Fluorine in Aqueous Solution by Calcium Aluminum Ferrite and the Mixture of Calcium Aluminate and Gypsum

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The immobilization of fluorine by calcium aluminum ferrites has been studied using 20 mass ppm F-containing aqueous solution. It is found that the rate of the F immobilization by Ca$_2$(Al,Fe)$_2$O$_5$ is initially slower than that by 3CaO·Al$_2$O$_3$ at the solid/liquid ratio of 1/200, but a similar immobilization degree is obtained at later stage. The hydration product contributed to the F immobilization is the Ca$_3$(Al,Fe)$_2$O$_5$·(OH,F)$_{12}$ phase, which is confirmed by the combination of XRD and electron microprobe analysis. For a ferrite phase of Ca$_2$(Al$_{0.68}$Fe$_{0.32}$)$_2$O$_5$, the hydration product has a chemical composition of Ca$_3$Al$_{1.54}$Fe$_{0.46}$(OH)$_{11}$F$_{0.0}$–0.13. The F immobilization by 2CaO·Fe$_2$O$_3$ does not occur to a significant extent, but the F immobilization by 12CaO·7Al$_2$O$_3$ is accelerated with the addition of 2CaO·Fe$_2$O$_3$ due to the formation of the Ca$_3$(Al,Fe)$_2$O$_5$·(OH,F)$_{12}$ phase. It is also observed that F can be immobilized by the formation of ettringite (3CaO·Al$_2$O$_3$·3CaSO$_4$·32H$_2$O, 3CaO·Fe$_2$O$_3$·3CaSO$_4$·32H$_2$O) through the substitution of F ion for the SO$_4^{2-}$ site.

KEY WORDS: immobilization; fluorine; calcium aluminum ferrite; ettringite.

1. Introduction

Fluorine has become a health and environmental concern ever since the discovery that fluoride is the cause of mottled enamel (dental fluorosis) and skeletal fluorosis. Human exposure to fluorides was primarily caused by the natural sources such as naturally fluoridated drinking water and fluoride uptake in fruits and vegetables from the soil and seafood. In recent years, however, F-containing waste has become a major concern on the F contamination.

The major waste form generated from ironmaking and steelmaking industry is slags with complicated compositions and these slags usually carry hazardous elements which sabotage the recycle and disposal of slags. Fluorine is one of the most commonly concerned hazardous elements in the steelmaking slags due to the usage of CaF$_2$ in refining process, particularly for the production of high quality steels. Fluorine contamination to the groundwater can occur when steelmaking slags are reused as road construction materials or subjected to landfill. The natural processes such as the weathering and leaching by rainfall can bring more fluorine into the groundwater. Therefore, a research has been carried out to immobilize fluorine in hot metal dephosphorization slags by using secondary refining slags whose main components are CaO, Al$_2$O$_3$, and SiO$_2$. A satisfactory immobilization degree has been attained, indicating that this remediation strategy is effective. However, the immobilization mechanisms involved are not clear yet due to the complexity of the composition of these slags.

The mechanisms of the F immobilization by calcium aluminates (3CaO·Al$_2$O$_3$, 12CaO·7Al$_2$O$_3$, 12CaO·7(Al,Fe)$_2$O$_3$) and/or calcium silicates (3CaO·SiO$_2$, 2CaO·SiO$_2$) which are major mineral phases in these slags have been investigated in the previous study. It was found that the immobilization of F ion could be established by the formation of the Ca$_3$(Al,Fe)$_2$O$_5$·(OH,F)$_{12}$ phase. For a ferrite phase of Ca$_2$(Al$_{0.68}$Fe$_{0.32}$)$_2$O$_5$, the hydration product has a chemical composition of Ca$_3$Al$_{1.54}$Fe$_{0.46}$(OH)$_{11}$F$_{0.0}$–0.13. The F immobilization by 2CaO·Fe$_2$O$_3$ does not occur to a significant extent, but the F immobilization by 12CaO·7Al$_2$O$_3$ is accelerated with the addition of 2CaO·Fe$_2$O$_3$ due to the formation of the Ca$_3$(Al,Fe)$_2$O$_5$·(OH,F)$_{12}$ phase. It is also observed that F can be immobilized by the formation of ettringite (3CaO·Al$_2$O$_3$·3CaSO$_4$·32H$_2$O, 3CaO·Fe$_2$O$_3$·3CaSO$_4$·32H$_2$O) through the substitution of F ion for the SO$_4^{2-}$ site.

KEY WORDS: immobilization; fluorine; calcium aluminum ferrite; ettringite.
and \( \text{Ca}_2\text{Al}_{1.36}\text{Fe}_{0.64}\text{O}_5 \) were synthesized by sintering mixtures of analytical grade \( \text{CaCO}_3, \text{Al}_2\text{O}_3 \), and \( \text{Fe}_2\text{O}_3 \) in an appropriate proportion at 1300°C for one week. The samples were ground to fine powder less than 100 μm in diameter. X-ray diffraction (XRD) analysis was used to confirm these compounds. Solid solution of \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) was prepared by heating a mixture of \( \text{C}_3\text{A} \) and analytical grade \( \text{Fe}_2\text{O}_3 \) at 1300°C for 2 weeks. The method for the preparation of \( \text{C}_3\text{A} \) has been already described elsewhere.\(^1\) The XRD analysis revealed that the \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) solid solution was formed and its X-ray diffraction pattern was almost the same as that of pure \( \text{C}_3\text{A} \).

The ettringite phase with fluorine was synthesized by precipitation in 20 mass ppm F-containing aqueous solution in which \( \text{C}_3\text{A} \) and \( \text{CaSO}_4 \cdot 2\text{H}_2\text{O} \) particles with the size less than 100 μm were added.

2.2. Shaking Test

Shaking test was made at room temperature. The test procedure which is basically followed by the Japanese standard test procedure (codified as Environment Agency Notice 46) has been already explained in previous article.\(^1\) The solid reactants were added in 20 mass ppm F-containing aqueous solution in 500 ml polyethylene bottle, and then the bottle was shaken horizontally at a speed of 200 times per minute. The solid/liquid ratio was 1/100 and 1/200, and the shaking time was up to 24 h. The F-containing aqueous solution was prepared by diluting hydrofluoric acid (47%) with distilled water. The solid phase was separated from the aqueous solution by filtration using the membrane filter with an open pore size of 0.1 μm after a certain period of shaking time. The solid phase was subjected to XRD analysis or microscopic observation. The concentration of F in an aqueous solution was determined by the selective ion-electrode method (JIS-K-0101), and those of Ca, Al and Fe were determined by inductively coupled plasma (ICP) atomic emission spectrometry.

2.3. Microscopic Observation

Microscopic observation was carried out by SEM equipped with electron microprobe analysis (energy dispersive or wave-length dispersive spectroscopy) to determine the contents of elements in F-containing minerals. The filtered residue after the shaking test and that in the experiments where large solid particles (0.84–1.0 mm) were reacted with 20 mass ppm F-containing aqueous solution were used for microscopic observation. These particles were rinsed by methanol, and then dried at 40°C for 24 h in CO₂-free environment.

3. Results and Discussion

3.1. Immobilization of F by Ferrite

The results for the F immobilization by \( \text{C}_2\text{FF} \), \( \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \) (\( \text{Ca}_3\text{Al}_{1.36}\text{Fe}_{0.64}\text{O}_5 \) and \( \text{Ca}_2\text{AF} \)) and \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) particles (≤0.1 mm) are shown in Fig. 1, in which the results\(^1\) for \( \text{C}_3\text{A} \) (≤0.1 mm) are also included for comparison. It can be seen that the F immobilization by \( \text{C}_2\text{FF} \) does not occur. However, \( \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \) demonstrates a good ability of removing F and the rate of F immobilization increases with an increase of Al/Fe ratio in the \( \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \) solid solution. This can be explained by the fact that the hydration rate of ferrites increases with increasing Al/Fe ratio. On the other hand, the \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) solid solution (\( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \), thereafter) demonstrates faster decrease of F content than the \( \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \) solid solution, especially at the beginning of the reaction. However, the F immobilization rate is obviously slower, compared to iron-free \( \text{C}_3\text{A} \).

The initial rates of releasing the Ca and Al contents in an aqueous solution suggest that the hydration ability of these minerals is in the increasing order of \( \text{C}_2\text{F} \ll \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \ll \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \ll \text{C}_3\text{A} \), which is in good agreement with the rate of F immobilization. Both \( \text{Ca}_3(\text{Al},\text{Fe})_2\text{O}_5 \) and \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) reduce the F content in solution to the same level as that by \( \text{C}_3\text{A} \) after about 12 h-shaking. This indicates that the major F-containing hydration product formed by them has a similar capability of holding F as that formed by \( \text{C}_3\text{A} \), which has been known as F-containing hydrogarnet phase (\( \text{Ca}_3\text{Al}(\text{OH,F})_2\text{F} \)).\(^1\) It is also noted that the concentration of Fe\(^{11+} \) ion in an aqueous solution is extremely low comparing with those of Ca and Al ions.

X-ray analysis revealed that no hydration products were identified for \( \text{C}_2\text{F} \) probably due to their small quantity. The main hydration product formed from \( \text{Ca}_3\text{Al}_{1.36}\text{Fe}_{0.64}\text{O}_5 \) and \( \text{Ca}_2\text{AF} \) has almost the same X-ray diffraction pattern to the hydrogarnet phase, \( \text{Ca}_3\text{Al}_2(\text{OH,F})_2\text{F} \). It was confirmed by electron microprobe analysis that this hydrate appears to be an iron solid solution of \( \text{Ca}_3\text{Al}_2(\text{OH,F})_2\text{F} \) phase containing F, that is, \( \text{Ca}_3(\text{Al},\text{Fe})_2(\text{OH,F})_2\text{F} \). In addition, the \( \text{C}_3\text{A}–8\text{H}_2\text{O} \) and \( \text{C}_3\text{A}–\text{Ca}(\text{OH})_2–18\text{H}_2\text{O} \) phases were also identified as a minor phase. The major hydration product for \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) was also \( \text{Ca}_3\text{Al}_2(\text{OH,F})_2\text{F} \) containing a small amount of iron (less than 1 mass%), as confirmed by electron microprobe analysis. The finding that the rate of F immobilization by \( \text{C}_3\text{A}–5\text{mass%Fe}_2\text{O}_3 \) is slower than that by \( \text{C}_3\text{A} \) may

![Fig. 1. Immobilization of F in aqueous solution by ferrites.](image-url)
be due to the fact that the retarding effect of a surface layer of reaction products may be greater when the surface layer contains ion, as observed by Boikova et al.  

The surface of a C$_2$F particle hydrated for 24 h in 20 mass ppm F-containing aqueous solution under shaking condition was examined by electron microprobe analysis. Hexagonal crystal C$_2$F·13H$_2$O and amorphous Fe(OH)$_3$ were identified and no significant amount of F was found in both phases.

The morphologies of a hydrated Ca$_3$Al$_{1.36}$Fe$_{0.64}$O$_5$ particle obtained at 2 and 24 h under static and shaking conditions are shown in Figs. 2 and 3, respectively. It can be seen from Fig. 2 that cubic crystals (C) along with two hexagonal crystals (A and B) are formed on the surface of a hydrated particle. Electron microprobe analysis revealed that the cubic hydrate corresponds to the iron- and F-containing hydrogarnet phase with the composition of Ca$_3$Al$_{1.54}$Fe$_{0.46}$[(OH)$_{1-x}$/H$_{1+0.02}$F$_x$]$_{12}$ (x=0–0.13). The hexagonal hydrates A and B correspond to the iron-containing Ca$_3$·8H$_2$O and Ca$_3$·Ca(OH)$_2$·18H$_2$O phases, respectively. The former contained no F, but the latter contained F up to 0.4 mass%. The surface layer consisted of mainly iron-containing hydrogarnet phase and its thickness increased with holding time.

Under the shaking condition the amount of iron-containing hydrogarnet phase (C) increases compared with that under static condition, as shown in Fig. 3. The surface layer at 2 h is thinner and denser than that observed under static condition. The amount of iron-containing hexagonal crystals of C$_2$A·8H$_2$O (A) and C$_3$A·Ca(OH)$_2$·18H$_2$O (B) is less than that under static condition. This is due to the fact that these hexagonal crystals are metastable and change to iron-containing hydrogarnet phase, as observed by Collepardi et al. The formation of amorphous Fe(OH)$_3$ which did not contribute to the F immobilization was also confirmed by electron microprobe analysis.

The Ca$_3$(Al,Fe)$_2$(OH,F)$_{12}$ phase forms a complete solid solution in the Ca$_3$Al$_2$(OH)$_{12}$–Ca$_3$Fe$_2$(OH)$_{12}$ solid solution series. It was found impossible to synthesize Ca$_3$Fe$_2$(OH,F)$_{12}$ by either the direct hydration of anhydrous C$_3$F in the presence or absence of lime, or the reaction between ferric hydroxide gel and lime-containing aqueous solution. Even though the possibility that an unstable pure Ca$_3$Fe$_2$(OH)$_{12}$ phase may be formed exceptionally under special conditions still remains, it can be ruled out that this phase will play a significant role in the F immobilization.
3.2. Effect of C_2F/H_33525 on F Immobilization by C_3A and C_12A7

In the previous section, it was found that C_2F/H_33525 particles alone have little effect on F immobilization in comparison with ferrite, Ca_2(Al,Fe)O_5 which strongly contributes to the F immobilization. In this section, C_3A or C_12A7 particles (≤0.1 mm) together with C_2F/H_33525 particles (≤0.1 mm) were added to a F-containing aqueous solution and the shaking test was carried out.

The effect of the C_2F/H_33525 addition on the F immobilization by C_3A and C_12A7 is shown in the upper and the lower diagram of Fig. 4, respectively. The results for C_3A and C_12A7 (≤0.1 mm) are also included for comparison. The F immobilization by C_3A is not affected to any important degree by the addition of C_2F/H_33525, as shown in the upper diagram of Fig. 4. However, the addition of C_2F/H_33525 to C_12A7 improves the F immobilization, as shown in the lower diagram of Fig. 4. Especially, the F content decreases sharply in the case of C_12A7/H_11001 30%C_2F/H_33525.

XRD analysis was made on filtered residue obtained after shaking for 2 h. The results are summarized in Table 1 where the results for C_3A and C_12A7 are also given. It can be seen that in the case of C_3A the hydration products are the same in the presence or the absence of C_2F/H_33525 and the main phase is Ca_3Al_2(OH)_12. In the case of C_12A7, however, with the addition of 30% C_2F a small amount of Ca_3Al_2(OH)_12 phase is identified, which is effective to the F immobilization. These results of XRD analysis explains the phenomena observed in Fig. 4.

As confirmed by electron microprobe analysis, in the presence of C_2F, the Ca_3Al_2(OH)_12 phase formed for C_3A and C_12A7 contained F, as well as Fe. Therefore, it can be said that the addition of C_2F favors the formation of the Ca_3(Al,Fe)(OH,F)_12 phase and accelerates the F immobilization process of C_12A7.

3.3. Immobilization of F by the Formation of Ettringite

3.3.1. F Immobilization

The mixture of C_3A and gypsum (CaSO_4·2H_2O) was added to 20 mass ppm F-containing aqueous solution at C_3A/liquid ratio = 1/200 in weight. The results of the shaking experiments for C_3A/H_11001 40mass%CaSO_4 and C_3A/H_11001 66mass%CaSO_4 are shown in Fig. 5. It can be seen that the behavior of F content is influenced by the amount of CaSO_4. In the case of C_3A/H_11001 40mass%CaSO_4 the F content decreases rapidly at the beginning, reaches the minimum at about 2 h, and then increases up to around 3 mass ppm. However, in the case of C_3A/H_11001 66mass%CaSO_4 the F content decreases steadily and reaches a constant value approximately at 0.03 ppm after 12 h. Correspondingly, it is observed that the S content is nearly the same level for both cases in the beginning of 2 h, indicating that the solution is saturated with S by the addition of CaSO_4 over 40 mass%. Therefore, the S content decreases in the case of C_3A/H_11001 40mass%CaSO_4, while it remains nearly constant for C_3A/H_11001 66mass%CaSO_4. It is also noted that the Al content is extremely low in both cases.

X-ray patterns of hydrated mixtures of C_3A and CaSO_4 obtained after 2 and 24 h are shown in Fig. 6. It is clear that in the case of C_3A/H_11001 40mass%CaSO_4 the hydration products
are ettringite (C₃A·3CaSO₄·32H₂O) and monosulfate (C₃A·CaSO₄·12H₂O). At 2 h the amount of the former appears to be larger than that of the latter, as shown in the upper diagram. At 24 h the latter becomes the main one, as shown in the middle diagram. These results suggest that ettringite is formed at first and then converts to monosulfate. As will be pointed out later, ettringite phase is the one responsible for the F immobilization. The drastic decrease of the F concentration at initial stage shown in Fig. 5 for C₃A/HH₁₀₀₀¹₄₀mass%CaSO₄ can be explained by the formation of ettringite and the increase of F content after 2 h is attributed to the decomposition of ettringite. It is further noted from Fig. 5 that this decomposition is caused by the decrease of S content in an aqueous solution.

In the case of C₃A/HH₁₀₀₀¹₆₆mass%CaSO₄, ettringite is the only hydration product through the experimental period of time, as shown in the lower diagram in Fig. 6. This can explain the F immobilization behavior shown in Fig. 5. It is considered that since the formation of ettringite layer on a C₃A particle might be much faster in the presence of a large amount of CaSO₄, the process of F immobilization is retarded by this ettringite layer at beginning. It should be pointed out that the amount of CaSO₄ in this mixture corresponds to the stochiometric amount needed for the formation of ettringite.

Morphology of the surface of C₃A particles hydrated for 24 h in 20 mass ppm F-containing aqueous solution in which 40 and 66 mass% CaSO₄ was added, respectively, is shown in Fig. 7. The only hydration product observed in the case of C₃A/HH₁₀₀₀¹₆₆mass%CaSO₄ is needle-like ettringite (A) which contains up to 2 mass percent F. On the other hand, for C₃A/HH₁₀₀₀¹₄₀mass%CaSO₄, monosulfate (B) was identified together with ettringite (A). The former appears to be smaller acicular crystal, while the latter is much larger comparing with that formed for C₃A/HH₁₀₀₀¹₆₆mass%CaSO₄. This finding suggests that the ettringite is formed in the early stage of the reaction and is decomposed to monosulfate in the case of C₃A/HH₁₀₀₀¹₄₀mass%CaSO₄. No significant amount of F was identified in monosulfate phase. It can be seen that the microscopic observation corresponds well to the results of XRD analysis, as well as the results for the F immobilization behavior shown in Fig. 5.

The results for F immobilization by the mixture of ferrite (Ca₂Al₁.₃₆Fe₀.₆₄O₅) and CaSO₄ are shown in Fig. 8. The amount of CaSO₄ added in 20 mass ppm F-containing solution is the same as that shown in Fig. 5. It can be seen that the F immobilization does not occur within the experimental period of time and both Ca and S contents remain constant. As revealed by XRD, no hydration products were identified. This suggests that the hydration of Ca₂Al₁.₃₆Fe₀.₆₄O₅ is retarded by the presence of CaSO₄.

In order to prepare an iron-based ettringite, C₃F-
3CaSO\(_4\cdot32\)H\(_2\)O, a mixture of CaO and Fe\(_2\)SO\(_4\) in stoichiometric ratio for the formation of ettringite, was added into the aqueous solution containing 1 240 mass ppm F, followed by 30 h shaking. The SEM image of hydration product is shown in Fig. 9(a). Al-based ettringite was also synthesized in a similar manner by using CaO and Al\(_2\)SO\(_4\), and its morphology is shown in Fig. 9(b). It is clear that the morphologies of these two ettringite phases are different. The Al-based ettringite appears to be acicular crystals which are about 5 \(\mu\)m long, while the Fe-based ettringite is prismatic crystals which are much longer, about 9 \(\mu\)m. In the case of Fe-based ettringite, it was found that the value of F content determined by electron microprobe analysis depends on crystal plane. If the X-ray beam was focused on the plane perpendicular to the prism axis, the F content was about 2 mass%. However, the F content was 1 mass% on the plane parallel to the prism axis. In the case of Al-based ettringite, only the F content on the plane parallel to the prism axis was evaluated and the F uptake was about 1 mass%.

3.3.2. Immobilization Mechanism

It is known that the structure of ettringite is based on two distinct structural components\(^5\): columns of \(\{\text{Ca}_6([\text{Al,Fe})(OH)_6]_2 \cdot 24\text{H}_2\text{O}\}\)\(^{+\times}\) and channels of \(\{\text{SO}_4\}_6 \cdot 2\text{H}_2\text{O}\)^{6\text{+}} which contain four sites, of which three are occupied by sulfate and one by two water molecules. With this structure a modest deviation in composition can be tolerated by sulfate and one by two water molecules. With this structure a modest deviation in composition can be tolerated without causing a change in structure. The ions available for substitution in the ettringite structure are Ca\(^{2+}\), Al\(^{3+}\), SO\(^{4-}\) and OH\(^{-}\) and among them only OH\(^{-}\) and SO\(^{4-}\) are replaceable by F\(^{-}\). Comparing with OH\(^{-}\) ion site in columns, SO\(^{4-}\) site in channels is much more favorable for the ion substitution. A variety of this sulfate-substitution occurrences have been reported, like Cl\(^{-}\) and OH\(^{-}\) ions. Considering the similarities of F\(^{-}\) with Cl\(^{-}\), OH\(^{-}\), it seems reasonable to assume that the F immobilization occurs through the substitution of F\(^{-}\) ion for this SO\(^{4-}\) site.

This assumption is supported by the relation of the value of F content to the crystal plane, as mentioned above. The number of channels in the direction parallel to the prism axis in the unit cell of ettringite is twice as many as that in the direction perpendicular to the prism axis because a half of channels is shielded by columns. Therefore, if the F immobilization occurs through the substitution of F\(^{-}\) for SO\(^{4-}\) sites in channels, the value of F uptake obtained by electron microprobe analysis will depend on the crystal plane on which the X-ray beam is introduced.

The F immobilization mechanism can be represented by the following reaction:

\[
\text{C}_3\text{A}+3\text{CaSO}_4+6\text{xF}^-+32\text{H}_2\text{O} \rightarrow \text{C}_3\text{A} \cdot 3\text{Ca}[\text{SO}_4]_{1-x} \cdot \text{F}_x \cdot 32\text{H}_2\text{O}+3x\text{SO}_4^2^-,
\]

To balance charges, one SO\(^{4-}\) site is replaced by 2F\(^{-}\) ions, and on the basis of the aforementioned results obtained by electron microprobe analysis, the substitution degree of x is up to 0.2. Therefore, 0.2 mole of ion per ettringite molecular can be replaced by 0.4 mole of F\(^{-}\) ion at maximum.

Based on the present study, it is said that fluorine in waste water can be immobilized by using stabilizing agents. For this purpose, the stability of F-containing hydrates should be examined at various pH values.

4. Summary

The F immobilization by the formation of iron-containing hydrogarnet phase and ettringite phase has been studied. The effect of C\(_3\)F on the F immobilization by C\(_3\)A and C\(_{12}\)A\(_2\) is also discussed. Based on these results, the following conclusions are summarized:

1. The F immobilization by C\(_3\)F on the F immobilization by C\(_3\)A and C\(_{12}\)A\(_2\) occurs through the formation of C\(_3\)F\(_3\)(Al,Fe)\(_2\)O\(_{11}\) and demonstrates a similar ability to the F immobilization by 3CaO·Al\(_2\)O\(_3\).

2. C\(_3\)F does not immobilize F, but the addition of C\(_3\)F to C\(_{12}\)A\(_2\) gives a positive effect on the F immobilization by C\(_{12}\)A\(_2\), which is due to the formation of C\(_3\)F\(_3\)(Al,Fe)\(_2\)(OH,F)\(_{12}\).

3. Ettringite can immobilize F through the substitution of F\(^{-}\) for SO\(^{4-}\) site in channels.

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