

Influence of finely ground limestone on cement hydration

Jean Péra, Sophie Husson, Bernard Guilhot

▶ To cite this version:

Jean Péra, Sophie Husson, Bernard Guilhot. Influence of finely ground limestone on cement hydration. Cement and Concrete Composites, Elsevier, 1999, 21 (2), pp.99-105. 10.1016/S0958-9465(98)00020-1. hal-00555525

HAL Id: hal-00555525 https://hal.archives-ouvertes.fr/hal-00555525

Submitted on 13 Jan 2011

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Influence of finely ground limestone on cement hydration

JEAN PERA(1)*, SOPHIE HUSSON(1,2), BERNARD GUILHOT(2)

- (1) Institut National des Sciences Appliquées de Lyon, Unité de Recherche Génie Civil-Matériaux, 20, Boulevard Albert Einstein, Villeurbanne Cedex, France
- (2) Ecole Nationale Supérieure des Mines de Saint Etienne ; Centre SPIN ; Département PMMC ; LPMG -UMR CNRS 5148 ; 158 Cours Fauriel, 42023 Saint-Étienne Cedex 2, France

Abstract

Some work has been carried out on the effect of calcium carbonate on cement paste, but there is no general agreement on the relative effects of different amounts of calcium carbonate on cement paste properties. The objective of the present work is to assess the effect of various amounts of calcium carbonate on the hydration of tricalcium silicate in order to explain the physico-chemical changes occurring during Portland cement hydration. It is shown that calcium carbonate has an accelerating effect on C_3S and cement hydration and leads to the precipitation of some calcium carbosilicate hydrate.

Keywords:

Calcium carbonate; Calorimetry; Carbosilicate; Cement; Hydration; Infrared spectroscopy; Microstructure; Strength

I. Introduction

For a long time, ground limestone has been considered as an inert filler. However, recent studies carried out in the 1980s have pointed out the following phenomena:

- calcium carboaluminate hydrates precipitate during the hydration of cements containing ground limestone [1-7],
- during the formation of ettringite, sulfate ions can be replaced by carbonate ions without modifying the sequences of the reaction [8-11],
- there is an interaction between calcium silicate (alite) and calcium carbonate; the latter accelerates the hydration of C3S and modifies the Ca/Si ratio of C-S-H [12-14].

As the interactions of ground limestone and C_3A are well documented, we have, in the present paper, focused our attention on the reactions which occur during the hydration of C_3S in the presence of calcium carbonate. The behaviour of blended Portland cements containing up to 50% ground limestone has also been investigated.

_

^{*} Corresponding author: jean.pera@insa-lyon.fr

II. Experimental

II.1. Materials

II.1.1. Calcium carbonate (CaCO₃)

The calcite studied was derived from marble and was very pure (98.6% CaCO₃). Its specific gravity was 2.75.

It was ground to get the following characteristics:

- * 100% particle below 20 μm,
- average diameter $D_sO = 2.5 \mu m$,
- * 35% particles below 1 μm.

For this product, the Blaine specific surface area was $680m^2kg^{\text{-}1}$ and the BET specific surface area was $3600\ m^2kg^{\text{-}1}$

Calcium silicate (C_3S)

Calcium silicate was synthesized by reacting a well mixed stoichiometric blend mixture of pure calcium carbonate and reactive silica. The blend was wetted and pellets prepared. The temperature program was as follows:

- ❖ 20-700°C: 20°C min⁻¹
- ❖ 700-1000°C: 15°C min⁻¹
- **▲** temperature maintained at 1000°C for 40 min.
 - **❖** 1000-1450°C: 8°C min⁻¹
 - **❖** 1450-1600°C: 5°C min⁻¹

▲ temperature maintained at 1600°C for 120 min.

The residual lime (CaO) was measured by hydrochloric acid titration of a solution containing 250 mg C_3S and 100 mL ethyleneglycol, shaken at 70°C for 30 min. The calcination was considered good as the free CaO content was less than 2%. A well-crystallized triclinic C_3S was then obtained and ground in a laboratory ball-mill. Its particle size distribution was characterized by:

- ❖ 100% particles below 80 µm
- ♦ $D_{50} = 15 \mu m$
- ❖ 30% particles below 10 µm

II.1.2. Ordinary Portland Cement (OPC)

The cement used in the present study was a CPA CEM I 52.5 according to the European prestandard ENV 197-1 for common cements. Its Bogue potential composition was $C_3S = 67.9\%$; $C_2S = 4.8\%$; $C_3A = 10.4\%$; $C_4AF = 9.3\%$. Its Blaine and BET specific surface areas were 364 m²kg¹¹ and 1400 m²kg¹¹, respectively. The average value of the particle size distribution was $D_{50} = 16 \ \mu m$.

II.2. Methods used for investigation

Isothermal calorimetry was used to study the interactions between C_3S (or OPC) and C_3C_3S (or OPC) was mixed with C_3C_3S in the weight ratio 1:1. The mixture (300 mg) was then hydrated in presence of the same quantity of water (300 mg). The total heat developed during the reaction was recorded. The length of the experiment was 15 h.

The morphology of these hardened pastes was investigated by means of scanning electron microscopy (SEM), and the hydrates formed were identified, using infrared spectrometry (IRS), X-ray diffraction (XRD) and differential thermal analysis (DTA).

Pastes of pure C₃S (or OPC) and CaCO₃ were prepared at equivalent consistency. The water to solid ratios are shown in Table 1.

From Table 1, it is clear that the presence of CaCO₃ has a plasticizing effect on the paste: the higher the CaCO₃ content, the lower the amount of mixing water. Minicylinders of paste were cast ($\phi = 20$ mm, h = 40 mm), demoulded after 2 days of hydration, and kept at 20°C and relative humidity 95% until mechanical testing, which occurred after 7, 28 and 60 days of age.

III. Results and discussion

III.1. Reactions between C₃S and CaCO₃

III.1.1. Isothermal calorimetry

The isothermal calorimetry curves showing the rate of heat development of 300 mg C_3S and (150 mg C_3S+150 mg $CaCO_3$) during hydration, up to 15 h, are given in Figure 1. The values recorded for the blend ($C_3S+CaCO_3$) were always higher than those of hydrated C_3S .

The total heat resulting from pure C_3S hydration was 145 joules, whilst that of the mixture $(50\% \ C_3S+50\% \ CaCO_3)$ reached 260 joules. These results are in good agreement with those obtained by Ramachandran [14]. They indicate that $CaCO_3$ cannot be considered as an inert addition towards C_3S hydration.

III.1.2. SEM examination

As shown in Figure 2, the pastes presented about the same texture after 7 days of hydration; namely platlets of calcium hydroxide and fibrils of C-S-H. After 60 days of hydration, the morphology of the pastes is different, as presented in Figure 3. In pure C_3S , platlets of calcium hydroxide and C-S-H type II are present according to Taylor [15]. In $^{\rm L}C_3S+CaCO_3$ paste, granules of C-S-H are formed.

III.1.2. Infrared spectrometry

The IR spectra of hydrated C₃S at different ages are shown in Figure 4. Those of the 'C₃S+CaCO₃' paste are shown in Figure 5. The wavenumbers present in each product hydrated either for one day or 28 days are reported in Table 2. The vibrations associated with each wavenumber are also presented in Table 2.

In hydrated C₃S, all the Si-O-Si stretching bands were reorganized between 1 and 28 days. The structure of tobermorite gel was reached [16], as shown by the emboldened wavenumbers.

In the blend ' $C_3S+CaCO_3$,' the characteristic bands of the CO_3 ion appeared. Some SiO_4 bands were shifted after one day of hydration, demonstrating the accelerating effect of $CaCO_3$ on C_3S hydration. Those results were also obtained using the diffuse reflectance method, as shown in Figure 6.

The results of XRD investigations are given in Figure 7, after 60 days of hydration. The following conclusions can be drawn from these analyses:

- calcium hydroxide is present in all compositions;
- **❖** the intensity of the C₃S peak decreases when the amount of CaCO₃ increases such that in the blend containing 50% CaCO₃, these peaks have almost disappeared;
- ***** the intensity of the peak at $2\theta = 14.9^{\circ}$ increases with the CaCO₃ content and is due to the formation of some hydrated carbosilicate.

The consumption of $CaCO_3$ was studied by means of DTA. The area of the decomposition peak of $CaCO_3$ decreased with the hydration time and the temperature corresponding to the maximum of this peak shifted towards smaller values as shown in Figure 8. Such results were previously observed by Henning [6] in the case of precipitation of carbonate hydrates.

III.1.4. Compressive strength of pastes

The compressive strength of the different pastes is presented in Figure 9. The presence of $CaCO_3$ at levels higher than 30% has a beneficial effect on the strength, whatever the hydration time may be.

It is possible to compute the K(t) value for each paste using Feret's formula:

 $R(t) = K(t)(V_s/V_t)^2$

where R(t) is the strength after t days of hydration, K(t) is the coefficient of reactivity of the binder, Vs, is the volume of binder, and Vt, the sum of the following three volumes: binder, water and entrained air. In the present study, the volume of air was measured at 1.5-1.6%. Table 3 shows the K(7) and K(60) values obtained for the different pastes. After 7 days hydration, all blended pastes develop higher K(7) values than pure C_3S which proves that $CaCO_3$ reacts chemically. After 60 days hydration, only pastes containing more than 30% $CaCO_3$, develop higher K(60) values.

III.2. Reactions between OPC and CaCO₃

III.2.1. Isothermal calorimetry

Figure 10 shows that the same type of curve is observed with OPC and pure C_3S . The heat produced by the reaction between 50% OPC and 50% $CaCO_3$ is about double that issued from plain OPC hydration.

III.2.2. Hydrates formed

As shown by XRD (Figure 11) carboaluminate and carbosilicate have been precipitated after 60 days of hydration.

The presence of calcium carboaluminate hydrate was also observed by infrared spectrometry (bands at 3670 and 3530 cm⁻¹). IR spectroscopy also pointed out differences in AF_m and AF_t phases (Figure 12). In the paste containing pure OPC, both monosulphate (bands at 100 and 1170 cm⁻¹) and ettringite (band at 1120 cm⁻¹) are present. In the paste containing 50% CaCO₃, monosulphate almost disappeared and the band at 1120 cm⁻¹ was reinforced, so some SO₄ ions in ettringite were probably replaced by CO₃ ions. The band assigned to the ν_3 SiO₄, vibration at 970 cm⁻¹ was also slightly modified and that of carbonate at 875 cm⁻¹ amplified.

III.2.3. Compressive strength of pastes

The compressive strength of pastes is presented in Figure 13. The strength is maintained or even increased in pastes containing 10% CaCO₃. Lower strengths are obtained with higher CaCO₃ levels. These results mean that the development of strength is not similar in C₃S and OPC pastes containing CaCO₃. In OPC, interactions occur between C₃A and CaCO₃, leading to the production of calcium carboaluminate hydrate and the modification of ettringite, whereas in blended C₃S pastes, only calcium carbosilicate hydrate is obtained.

Nevertheless, the level of strength remains acceptable for a 50% level of OPC substitution: it is 81% that of OPC alone. This result proves that chemical reactions occur, but in a smaller extent than in C_3S pastes and more $CaCO_3$, remains as an inert filler.

IV. Conclusion

As shown by isothermal calorimetry, more total heat is developed in C_3S or cement containing up to 50% calcium carbonate than in the absence of $CaCO_3$ under comparable conditions signifying the accelerating effect of $CaCO_3$. Hydration of C_3S in the presence of $CaCO_3$, results in the production of some calcium carbosilicate hydrate and good mechanical performance for amounts of $CaCO_3$ higher than 30%.

In cement paste, calcium carbonate modifies the AF_m and AF_t phases, and produces calcium carbosilicate and carboaluminate hydrates but does not lead to the same strength development as in C₃S paste.

References

- [1] A Bachiorrini, Interactions physico-chimiques entre l'aluminate monocalcique et différents carbonates au cours de l'hydratation. Thesis I, Université Claude Bernard, Lyon (1985), p. 215.
- [2] A Bachiorrini, AA Fournier, B Guilhot, M Murat, A Negro and M Soustelle, Influence de la calcite sur l'hydratation de l'aluminate monocalcique Corrélation entre les résultats expérimentaux et la thermodynamique du système. Proceedings of the 8th International Congress on the Chemistry of Cements, Rio de Janeiro, 22–27 September 1986 IV, FINEP, Rio de Janeiro (1986), pp. 376–380.

- [3] VS Ramachandran, Thermal analysis of cement components hydrated in the presence of calcium carbonate. Thermochimica Acta 127 (1988), pp. 385–394.
- [4] SM Bushnell-Watson and JM Sharp, The detection of the carboaluminate phase in hydrated high alumina cements by differential thermal analysis. Thermochimica Acta 93 (1985), pp. 613–616.
- [5] A Bachiorrini and L Cussino, Hydratation du ciment alumineux en présence d'agrégats calcaires. In: Proceedings of the 8th International Congress on the Chemistry of Cements, Rio de Janeiro, FINEP, Rio de Janeiro IV (22–27 September 1986), pp. 383–388.
- [6] O Henning and A Kudjakov, Einfluss von Calcit auf die Hydratation von Portland-Zement. Wissenchaftliche Zeitschrift de Hochschule für Architektur and Bauwesen Weimar 29 (1983), pp. 75–77.
- [7] O Henning and A Zkudjakov, Einfluss von Dolomit auf die Hydratation von Portland-Zement. Wissenchaftliche Zeitschrift de Hochschule für Architektur and Bauwesen Weimar 27 (1980), pp. 187–191.
- [8] C Vernet, Séquences cinétiques des réactions d'hydratation de l'aluminate tricalcique en présence de gypse, de chaux et de fillers calcaires. Proceedings of the 8th International Congress on the Chemistry of Cements, Rio de Janeiro, 22–27 September 1986 III, FINEP, Rio de Janeiro (1986), pp. 70–74.
- [9] A Negro, G Abbiati and L Cussino, Sur l'emploi du filler calcaire comme régulateur de prise. Proceedings of the 8th International Congress on the Chemistry of Cements, Rio de Janeiro, 22–27 September 1986 III, FINEP, Rio de Janeiro (1986), pp. 109–119.
- [10] J Bensted, Further hydration investigation involving Portland cement and the substitution of limestone for gypsum. World Cement 14 (1983), pp. 383–392.
- [11] A Bonnin and B Cariou, Système aluminate de calcium gypse-chaux et eau. 7e Congrès International de la Chimie des Ciments, Paris III (1980), pp. 158–163.
- [12] VS Ramachandran and Chun-Mei Zhang, Influenza del CaCO₃ sulla idratazione e sulle caracteristiche microstrutturali del silicato tricalcio. II Cemento 3 (July/September 1986), pp. 129–152.
- [13] VS Ramachandran and Zhang Chun-Mei, Dependence of the fineness of carbonate on the hydration behaviour of tricalcium silicate. Durability of Buildings Materials 4 (1986), pp. 45–66.
- [14] VS Ramachandran, Cement with calcium carbonate addition. Proceedings of the 8th International Congress on the Chemistry of Cements, Rio de Janeiro, 22–27 September 1986 III, FINEP, Rio de Janeiro (1986), pp. 109–119.
- [15] HFW Taylor, The hydration of tricalcium silicate. Matériaux et Constructions 17 102 (1984), pp. 457–468.
- [16]. VC Farmer, The Infrared Spectra of Minerals, Mineralogical Society, London (1974).

Figures

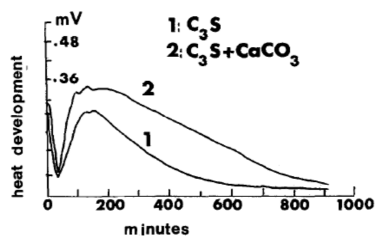


Fig. 1. Isothermal calorimetry data: C₃S and C₃S+CaCO₃.

Figure 1: Isothermal calorimetry data: C₃S and C₃S + CaCO₃.

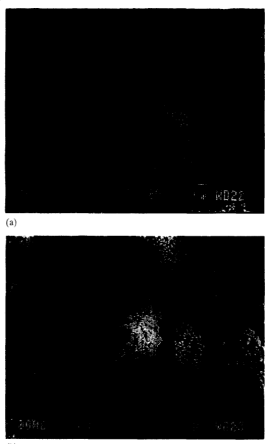


Figure 2: SEM micrographs after 7 days of hydration.

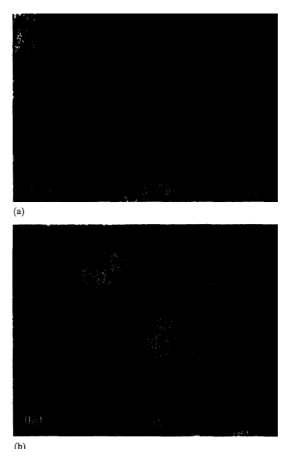


Figure 3: SEM micrographs after 60 days of hydration.

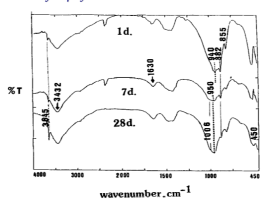


Figure 4: Infrared spectra of hydrated C₃S.

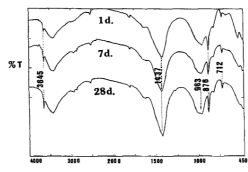


Figure 5: Infrared spectra of hydrated (C₃S+CaCO₃).

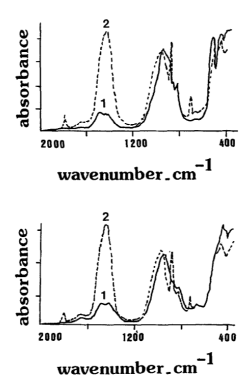


Figure 6: IR spectra obtained from the diffuse-reflectance method.

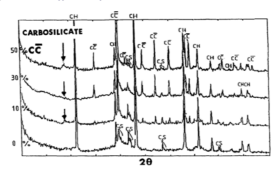


Figure 7: XRD patterns of the different pastes after 60 days of hydration.

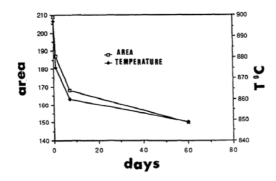


Figure 8: Evolution of the area and the temperature of $CaCO_3$ decarbonation.

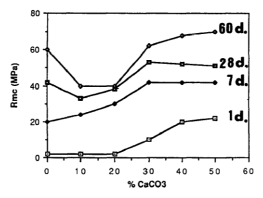


Figure 9: Influence of CaCO3 on the compressive strength of pastes.

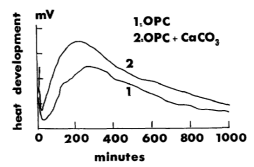


Figure 10: Isothermal calorimetry curves: OPC and OPC+CaCO₃.

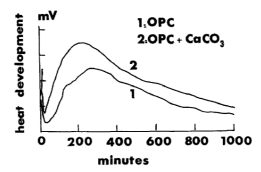


Figure 11: XRD spectrum of hydrated paste (OPC/CaCO₃ = 1).

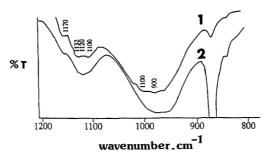


Figure 12: Modification of AF_m and Aft phases in the paste containing 50% OPC and 50% CaCO₃ after 60 days of hydration.

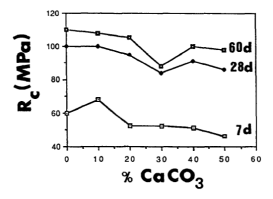


Figure 13: Compressive strength of cement pastes containing different amounts of CaCO₃.

Tables

Table 1: Water to solid ratios of pastes

C ₃ S (w _t %)	100	90	80	70	60	<i>50</i>
CaCO3 (wt%)	0	10	20	30	40	<i>50</i>
$W/SS = C_3S + CaCO_3$	0.35	0.34	0.33	0.32	0.31	0.3
OPC (wt%)	100	90	80	70	60	50
CaCO3 (wt%)	0	10	20	30	40	50
W/S S= OPC + CaCO3	0.28	0.28	0.27	0.26	0.25	0.24

Table 2: Infrared results of hydrated C₃S and (C₃S+CaCO₃)

	C_3S			C₃S + CaCO₃				
Wavenur	Wavenumber (cm ⁻¹)		Wavenun	nber (cm ⁻¹)	Vibration			
1 day	28 days		1 day	28 days				
1635	1630	v2 H2O	1635	1635	v2 H2O			
1480	1450	v3 CO3	1437	1437	v3 CO3			
938	960	v3 SiO4	1161	1161	v1 CO3			
905		v3 SiO4	954	963	v3 SiO4			
882	882	v3 SiO4	876	876	v2 CO3			
855	855	v3 SiO 4	849		v3 SiO 4			
660	665	v4 SiO4	712	712	v4 CO3			
	640	v4 SiO 4	660	660	v4 SiO 4			
518		v4 SiO4	506	486	v4 SiO4			
450	450	v2 SiO 4	452	462	v2 SiO4			

Cement and Concrete Composites, 1999, <u>21</u>(2), 99-105, <u>doi:10.1016/S0958-9465(98)00020-1</u>

Table 3: Reactivity of the different binders

C ₃ S	100	90	80	70	60	<i>50</i>
CaCO3 (Wt%)	0	10	20	30	40	<i>50</i>
K(7)	100	107	129	179	164	157
K(60)	225	125	130	300	375	410