The shrinkage of hardening cement paste and mortar

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THE SHRINKAGE OF HARDENING CEMENT PASTE AND MORTAR

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E.M. Theissing, M. van Wallendael

(Rec'd. Sept. 26, 1973; in final form April 9, 1975)
(Refereed)

ABSTRACT

This paper is an abstract from the report of the commission B10: "The influence of the shrinkage of cement on the shrinkage of concrete", of the Netherlands Committee for Concrete Research.
Measurements of pulse velocity, volume shrinkage and heat of hydration on hardening portland cement support the idea that the formation of ettringite is an important link in the mechanism of shrinkage in the plastic stage of cement paste and mortar.
Mechanical tests on prisms of $4 \times 4 \times 16 \text{ cm}^3$ gave some information about the difference in sensitivity to surface corrosion of different types of cement.

Auch mechanische Prüfungen an erstarrten Zement- und Mörtelprismen wurden durchgeführt zur Erklärung des Schwindprozesses.

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Introduction

The purpose of this article is to introduce to a greater public the findings of the commission B10 of the Netherlands Committee for Concrete Research (CUR) which studied the influence of the shrinkage of cement on the shrinkage of concrete.

In this article only the main points of the study will be discussed.

The shrinkage of cement paste during hardening is caused by chemical reactions of the clinker minerals with water during the first hours. At a later stage it is caused by changes in the water content of the hardened paste.

The purpose of the study was to obtain more knowledge about the changes in volume in the first stage. To do this the following properties were measured during the hardening process:

a) Heat evolution rate (with a conduction calorimeter)
b) Velocity of ultrasonic pulses
c) Volume shrinkage

Next to these measurements also the strength and the shrinkage of the hardened prisms, made with different types of cement, were measured.

Experimental

The heat evolution rate calculated per gram cement was measured using Lerch's method (1) by means of an isothermal conduction calorimeter as described by Stein (2). The first peak was not measured because the mixing procedure of 2 minutes stirring, 3 minutes rest, 2 minutes stirring was done outside the calorimeter.

The velocity of pulses in hardening cement pastes and mortars was measured with a Cawkell UCT 2 apparatus at 40 kHz. The specimen was placed in a thin rubber container between the piezo
electric measuring crystals of the instrument. The crystals were mounted on a fixed distance; the specimen filled exactly the space between them.

Changes in volume of a hardening paste or mortar due to chemical transformations were measured on a specimen of ± 600 grams placed in a water-tight rubber cover to exclude the influence of drying or water-uptake. This sealed specimen was placed in a vessel completely filled with water of 20 ± 0.1 °C. The vessel was connected with a tube filled with water to a beaker on the scale of a Mettler balance. When changes of volume occurred the rubber cover followed these changes and water was sucked out or pressed into the beaker. The changes in weight of the filled beaker were recorded and from these the changes in volume of the specimen were calculated. In these experiments the same batch of portland cement class A was used.

The chemical and calculated mineralogical composition of this portland cement is given in table 1.

**Results and Discussion**

**Heat Evolution Rate**

The influence of increasing amounts of the various clinker minerals, of water, of the addition of sand and of the time of aerating the Paste or Mortar.

**TABLE I**

<table>
<thead>
<tr>
<th>Chemical Analysis</th>
<th>in % by weight</th>
<th>Mineralogical Composition</th>
<th>in % by weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>L.O.I. 1000 °C</td>
<td>1.0</td>
<td>L.O.I. 1000 °C</td>
<td>1.0</td>
</tr>
<tr>
<td>SiO₂</td>
<td>22.5</td>
<td>C₂S</td>
<td>30.8</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>5.5</td>
<td>C₃S</td>
<td>44.8</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>2.6</td>
<td>C₃A</td>
<td>10.2</td>
</tr>
<tr>
<td>CaO total</td>
<td>64.5</td>
<td>C₄AF</td>
<td>7.9</td>
</tr>
<tr>
<td>MgO</td>
<td>1.2</td>
<td>CaO free</td>
<td></td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.1</td>
<td>MgO</td>
<td>1.2</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.5</td>
<td>Na₂O</td>
<td>0.1</td>
</tr>
<tr>
<td>SO₃</td>
<td>2.1</td>
<td>K₂O</td>
<td>0.5</td>
</tr>
<tr>
<td>CaSO₄</td>
<td></td>
<td>CaSO₄</td>
<td>3.6</td>
</tr>
<tr>
<td>Total</td>
<td>100.0</td>
<td>Total</td>
<td>100.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Specific Surface cm²/g</td>
<td>2900</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Specific Weight</td>
<td>3.11</td>
</tr>
</tbody>
</table>
cement with air containing carbon dioxide and water vapour on the flow of heat was studied.

The influence of the specific surface and the amount of alkali of the cement, which are thought to be large, have not been investigated.

In general three periods of increasing and decreasing heat evolution (peaks) can be observed when the hardening of portland cement is studied with an isothermal conduction calorimeter. On the decreasing branch of the second peak after ± 18 hours a small hump or shoulder can sometimes be seen (Figure 1). The cause of this hump is not clear; it is frequently attributed to both β-C₂S and C₄AF.

The magnitude of the peaks, and the time at which they occur, in the first place depends on the mineralogical composition and fineness of the cement. In table 2 the changes in time and height of the second and third peak are summarized. The exact values of time shift and peak height are not given because they depend too much on the mineralogical composition of the clinker used.

The influence of C₃S added (Figure 2) on the second peak is rather clear: a larger surface of C₃S is exposed to water, so more reaction heat can be evolved.

![Figure 1](image-url)
The Influence on Time of Occurrence and Height of the Peaks in the Heat Flow Curve by varying the Composition

<table>
<thead>
<tr>
<th>Increasing Amounts of</th>
<th>Second Peak</th>
<th>Third Peak</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Height</td>
<td>Time</td>
</tr>
<tr>
<td>$C_3A$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$C_3S$</td>
<td>higher</td>
<td>-</td>
</tr>
<tr>
<td>$CaSO_4$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>water</td>
<td>lower</td>
<td>later</td>
</tr>
<tr>
<td>quartz sand</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>aerating time</td>
<td>lower</td>
<td>-</td>
</tr>
</tbody>
</table>

- = no distinct influence

Addition of more $C_3A$ (see Figure 3) gives a larger sulphate consumption during the first peak, the grains become covered with ettringite crystals through which the reaction rate becomes low (appr. at 2 hours in Figure 3). The ettringite layer is less thick with more $C_3A$ because the same amount of gypsum is available. Therefore the layer is more permeable for water.
This results, either according to the "scaling off" mechanism (2), (3) or by chemical destruction (4), (5), in a faster consumption of sulphate until the stage is reached at which all sulphate is consumed. This stage will be reached in shorter time and a new C₃A hydration peak (the third peak) is formed sooner.

The influence of the addition of gypsum can be described by the same mechanism with the opposite result; the third peak is developed later.

The influence of the water cement ratio on the heat of hydration of portland cement per gram of cement is given in Figure 1. The second peak is developed between 2 and 34 hours after mixing. The heat developed during this peak is high when the water cement ratio is low.

When interpreting these data, it should be remembered that part of the heat developed by a hydrating paste is used for heating the paste (there must exist a temperature difference between paste and constant temperature heat sink in order to make a
From calibration experiments, the heat flow recorded can be related to the temperature gradient existing over the heat flow meter; the temperature differences are found to be small, e.g. at the top of the second peak in Figure 1: 0.11 °C in the experiment with w/c ratio = 0.2, and 0.08 °C in the experiment with w/c ratio = 0.6. Consequently, only a small fraction (about 2%) of the total heat developed is employed for heating the hydration paste; thus, differences in heat capacities of paste + container caused by varying w/c ratios do not noticeably influence the accuracy of the heat evolution rate measurement. Moreover, the heat capacities do not differ much since they are determined largely by that of the containers.

Therefore, it must be concluded from the results mentioned in Figure 1, that the hydration reaction connected with the second heat evolution peak (C₃S hydration) is faster in pastes of lower w/c ratio. This stands in contrast with data reported by Kantro and Copeland (6) but agrees with experiments reported by Locher (7).

A slower C₃S hydration with increasing w/c ratio can be attributed to the influence of C₃A hydration. When a high w/c ratio is used, more C₃A reacts with water in the initial stages before the C₃A is covered with ettringite, and more material dissolves, so more aluminium is taken up by the calcium-silicate hydrates formed on the C₃S grains, shielding them from water. At low w/c ratio the C₃A will react less because the layer of ettringite is built up sooner around these grains since the water is super-saturated more quickly.

With this mechanism, a large influence of mixing conditions and cement composition is comprehensible, which explains the contradictory findings reported concerning the influence of w/c ratio on the hydration rate of C₃S.
The third peak is higher for higher w/c ratio and there is a small shift to shorter times. This difference can be explained by a difference in morphology of the ettringite formed.

At low w/c ratio (<0.30) a closely packed ettringite layer is formed. The layer formed at higher w/c ratio consists of ettringite with a more needle-like structure, which is less dense. This layer is less able to cut off the C₃A grains from the supply of water. Thus at high w/c ratio the process of water transport through the ettringite layer elapses faster; sulphate is consumed sooner and an unhindered C₃A reaction can start.

The resulting third peak will be higher at higher w/c ratio because more water is available and less hydrosilicates have been formed (as evidenced by the lower second peak), and the silicate hydrates formed may be less dense, so the water transport to the grains is less hindered. This third peak is not seen at very low w/c ratios because the low amount of water is completely blocked from the C₃A by the higher amount of CSH-gel formed during the second peak. The hump on the second peak at these w/c ratios can be attributed to the reaction of C₃A at a stage where the ettringite layer is repaired after bursting through crystallization pressure (see also shrinkage measurements).

At the start of the investigations it was found that measurements on the same cement with the same w/c ratio in various tests at different times gave different heights of the peaks and different times of occurrence. This was thought to be caused by the influence of humid air during storage. To prove this, a batch of portland cement was exposed to air with different amounts of CO₂ and water during different periods. The moisture from the air reacts with C₃A and C₃S, the product reacts with CO₂. This was shown by infra-red spectroscopy, where the band of the calcium-sulphate decreases with the band of anhydrous calcium-aluminate
The disappearance of anhydrous calcium-aluminate by exposure to CO₂ and humid air at 19.3 μm, indicating that calcium-sulpho-aluminates and/or calcium-carbo-aluminates are formed (see Figure 4).

The total heat development of a paste from aerated cement (Figure 5) is decreasing with increasing aerating time especially at the more humid conditions (>44 % r.h.). The second peak decreases and disappears at longer aerating times. The third peak is formed sooner. These peaks are very sharp.

In view of these effects, care was taken that aerating of the cement used in the other investigations could not take place. The strength of hardened paste or mortar made with aerated cement was very low.

The influence of quartz sand on the flow of heat of portland cement is given in the Figures 6 and 7 for w/c ratios of 0.50 and 0.30. The addition of sand influences predominantly the third peak. The peak becomes smaller and occurs sooner when the amount of sand is increased.
Flow of heat of portland cement with different amounts of quartz sand (w/c ratio 0.50) (cement-aggregate ratio by weight 1:0; 1:1; 1:3 and 1:6)

The first minimum comes later, indicating that the first peak lasts longer, so less $C_3A$ is available to produce the third peak. The sulphate consumption must be faster too, which results in a shift to shorter times for this peak.

It is not clear how the sand influences the early $C_3A$ reaction. The following explanations are possible:
a) Sand gives nuclei for crystallization

b) Sand absorbs the sulphate

ad a) The products of the C₃A reaction (C₂AH₈, C₄AH₁₄ and ettringite) can be formed in the vicinity of the sand when the crystallization of these products form faster by nuclei given by the sand. By this the C₃A can react longer and faster, so sulphate is consumed more quickly. The outcome of this will be that the third peak is developed sooner and will be smaller. As C₃A grains react longer, the C₃S grains will be coated with more hydro-aluminates too, so the second peak will be retarded.

The fact that C₃A reacts longer under the influence of sand can be proved by integrating the first peak in the flow of heat curve which was not possible with the method used.

ad b) Sulphate absorption is not likely because experiments on mortars with a cement/aggregate weight ratio of 1:1 with different gradings (ranging from 16-200 cm²/g) of the sand did not show an influence of the surface of the sand.

**Velocity of Ultrasonic Pulses**

Up to the time at which the top of the second peak in the flow of heat curve took place, the pulse velocity was higher for higher w/c ratios (Figure 8 and 9). After this period the pulse velocities were higher for lower w/c ratios. When the velocity is thought to be correlated to the compressive strength, the early strength of a paste or mortar made with high w/c ratio must be higher than one made with a low w/c ratio. During the first hours the velocities, however, were very low, even lower than in water, so the velocity was at that stage not directly related to the strength.

An explanation for these low velocities was given by Dr. E. Drescher
from Poland, when working in the Netherlands, by considering the cement paste as a visco-elastic material, using a Maxwell model. According to this, the velocity is depending on the frequency of the pulse (see Figure 10). At low frequencies (till 500 kHz) there is a large influence of the viscous part of the model by which the velocities are very low during the first 10 hours. For higher frequencies (2000-4000 kHz) the velocity is much nearer to the velocity of sound in water. At these frequencies the elastic part is the most important.

After 10 hours the differences between the velocities are smaller but the velocities are still higher for higher frequencies. At the same frequency the curves for the higher and lower w/c ratio cross each other, but the time at which they cross is not the same, more details have to be obtained however.

From this model the conclusion is drawn that the velocity in paste and mortar during the first hour is to a high degree dependant on
Pulse velocity in paste as a function of the time for different pulse frequencies (experimental).

Internal viscous forces. These forces are formed by the interaction of particles together and particles with fluid. The particles change in number, form and mass during the hardening process. The fluid changes in viscosity by the dissolving clinker minerals. So the difference in velocity between paste or mortar made with either low or high w/c ratio can probably be attributed to the difference in morphology of ettringite, the degree of hydration of C₃A and the degree of supersaturation of the water. Later this difference disappears because more particles are formed, during the second peak, giving a more solid gel with more elastic properties.

* A pulse can be seen as an external force working on the paste. At low w/c ratio a paste is called dilatant, the opposite of thixotrope, so an external force induces more internal force, that is a more viscous behaviour, resulting in a low velocity. At high w/c ratio the paste is thixotrope so an external force gives an abolition of the viscous forces resulting in higher velocities during the first hours.
This is particularly distinct in low w/c ratio pastes: here particle to particle interaction would be higher than in high w/c ratio pastes at the same degree of hydration because of the latters' higher porosity; moreover, according to the large heat evolution rate during the second peak more silicate hydrate is formed in low w/c ratio pastes. Mortars have always higher pulse velocities than pastes at the same w/c ratio because more "solid material" is found per unit of volume.

**Shrinkage**

In Figures 11 and 12 the shrinkage curves combined with the pulse velocity and flow of heat are given for paste and mortar with a cement : aggregate weight ratio 1 : 1. Different w/c ratios were used. During the first hours until the time at which the second peak is fully developed, the shrinkage is larger at lower w/c ratio. Later the shrinkage becomes lower for the lower w/c ratio so the curves cross each other during the second peak. This is ascribed to the following mechanism:

![Graph showing combined curves of flow of heat, shrinkage, and pulse velocity for pastes.](image-url)
Directly after mixing CaAl₂O₄ dissolves, giving a fast shrinkage (Figure 13). After some time the ettringite crystals are formed, CaAl₂O₄ dissolves more slowly and the velocity of shrinkage decreases.
The formation of ettringite compensates for the shrinkage because the volume of ettringite is only a little smaller than the total volume of the starting materials, while all other hydrates have a much smaller volume than the starting materials.

At low w/c ratios the ettringite crystals are formed close around the C₃A and the C₃S reaction starts early. After approximately 4 hours the unstable CSH-gel is formed causing an increase of the velocity of shrinkage. When the C₃S reaction is at its maximum value the velocity of shrinkage decreases while stable CSH-gel is formed.

When the hump on the second peak occurs the velocity of shrinkage gets a small decrease that is followed by an immediate increase after this shoulder. This effect can be explained by the formation of shrinkage resisting ettringite during the stage at which the bursted coating is sealed around the C₃A grain.

At high water cement ratios, shrinkage is less compared with low w/c ratios (Figure 11). This is supposed to be caused by the different morphology of the ettringite that has been formed. More C₃A has reacted but shrinkage is less so the ettringite formed must have more shrinkage resisting properties than the ettringite formed at low w/c ratio. When the crystals are longer and are more perpendicular to the grain they can keep the grains wider apart.

After 10 hours, shrinkage becomes higher at high w/c ratios than at low w/c ratios.

This can be attributed to differences in porosity of the CSH-gel: at the same degree of hydration the interaction between the calcium-silicate hydrate particles would be lower at higher w/c ratio than at lower w/c ratio, resulting in a weaker total system; in the pastes investigated here this is enhanced by a higher degree of C₃S hydration in low w/c ratio pastes as evidenced by the high heat evolution rate during the second peak.

When the third peak in the flow of heat curve appears, the velocity of shrinkage becomes faster again. This is caused by the hydration
of C$_3$A. Probably the shrinkage is even boosted up by the transformation of ettringite in the less voluminous monosulphate and aluminates. The tobermorite gel is not strong enough at this stage to withstand this shrinkage. Moreover, the new components may form in a less stress state or in low stress locales within the sample and therefore will not provide the support which the ettringite has previously provided.

The influence of quartz sand on the shrinkage is as follows: During the first hours a mortar with c/a weight ratio 1:1 shrinks more than a paste with the same w/c ratio. After ~8 hours the shrinkage of a mortar is less (the curve in Figure 12 is an extension to the rule). The shrinkage curves for low and high w/c ratios cross each other during the formation of the second peak. The explanation for this is the same as that given for paste. The higher value of the shrinkage of mortar than of paste during the first hours can be explained if it is admitted (see flow of heat) by the fact that the sand has a large influence on the morphology of the ettringite formed. At low w/c ratio the compact coating around the C$_3$A is not formed, C$_3$A reacts longer causing greater shrinkage. The shrinkage retaining influence of the ettringite at higher w/c ratio is partly maintained. Shrinkage is less in relation to mortar with low w/c ratio but there is more shrinkage compared with paste with the same w/c ratio, notwithstanding the larger amount of solid material per unit of volume.

As a consequence of the observed influence of the sand on shrinkage, a large dependance of the shrinkage on the specific surface of the sand was expected. But the measurements on mortar 1:1 with different gradings of the sand (from 16-200 cm$^2$/g) gave no clear relation between these quantities. So the explanation for the influence of sand needs more research as does the explanation for the influence of sand on heat evolution.
Conclusions

The measurements of the flow of heat, shrinkage and pulse velocity show that during the hardening a clear difference exists between pastes or mortars made with either high or low water/cement ratio. During the second peak in the heat evolution, a change takes place in the course of the shrinkage and pulse velocity. High starting values become low ones and low starting values become high. For these effects two explanations may be envisaged. One is as follows:

At low water cement ratios the ettringite formed during the first peak is thought to consist of a great number of small crystals that form a compact layer close around the C₃A grain hindering the further reaction of the C₃A. This gives high shrinkage and low pulse velocity. High shrinkage because all particles can come closer to each other and a low pulse velocity because the viscous forces in this compact system are larger.

At high water cement ratios the ettringite has the possibility to form more needle-like crystals that are able to keep the clinker particles wider apart thus retaining shrinkage and giving a more "elastic" network in which pulses can proceed faster.

Alternatively, the low pulse velocity in the initial stages in low water cement ratio pastes could be ascribed to formation of hydrate layers in the vicinity of the contact points between the solid particles.

Then the solid particles do not move independently under the action of an ultrasonic pulse, and damping may occur especially in the transmission of motion from one particle to its neighbour.

The low pulse velocity in the initial stages in low w/c ratio pastes should then be ascribed to a relative large extent of hydration (as evidenced by the heat evolution rate data), which causes a large amount of hydrate crystals (rather flexible in this state) to surround the contact points. In the later stages, the hydrate crystals
obtain so many mutual contact points that they obtain together a certain elastic stiffness; then the pulse velocity is increased by a higher extent of hydration. Future research is needed for settling this point.

During the second peak in the flow of heat curve, the CSH-gel is formed. At low w/c ratio a gel with good mechanical properties is formed, that resists the shrinkage better than the gel formed at high w/c ratio. This stronger gel is more elastic; the pulse velocity becomes higher.

The CSH-gel formed at high w/c ratios is also less able to resist the extra shrinkage that occurs during the third peak, where C₃A reacts and ettringite is transformed into other products with a smaller volume, such as monosulphate.

The influence of the w/c ratio gives the same trends for mortars as for pastes. The presence of quartz sand, however, influences the formation of ettringite, this results in deviations in the measured values in heat flow, shrinkage and pulse velocity compared with paste.

In the heat flow the third peak comes sooner and has a lower intensity, because the C₃A reacts longer in the first hours, perhaps through nucleation by the sand. This first C₃A reaction gives higher shrinkage in the early hours too.

The pulse velocity is then higher, partly because the morphology of the ettringite is not the same as in paste, especially at low w/c ratios, or because contact points between quartz particles are less effected by hydrate formation than contact points between cement particles.

Later on, after the second peak, the sand serves as an inert aggregate and resists the shrinkage by the cement and makes possible higher pulse velocities.

The mechanism by which the sand influences the C₃A reaction during the first hours needs more detailed study.
The conclusion from these investigations is that good mechanical properties of pastes and mortars can be obtained when during the first five hours of hardening the shrinkage is relatively high, so that after this plastic stage one can expect low shrinkage and high pulse velocities (= high strength). This can be accomplished by promoting the growth of compact ettringite around the $C_3A$ grain or by suppressing ettringite formation by diminishing the $C_3A$ content. By this $C_3S$ can give more tobermorite and so strength will be built up sooner.

**Research on hardened Paste and Mortar**

Simultaneously with the preparation of the samples for the afore cited research, prisms of $4 \times 4 \times 16$ cm$^3$ were made. Prisms which were used for determination of flexural and compressive strength were stored during 28 days in water of $20 \, ^\circ C$. Prisms for shrinkage measurements were stored during 91 days at $20 \, ^\circ C$ and 65% relative humidity, after this period they were tested for strength. Strength and shrinkage of paste and mortar were related to the water cement ratio logarithmically (see Figure 14), and to the amount of gypsum (Figure 15) and of $C_3A$ (Figure 16) in portland cement.

At optimum gypsum content the strength is maximal and the shrinkage minimal (see Figure 15).

At other amounts of gypsum than the optimum amount, the water cement ratio of the sample has a larger influence on the strength and the shrinkage:

At higher w/c ratio ($> 0.40$) a high gypsum content gives relatively low strength. High shrinkage occurs when the amount of gypsum is low at high water cement ratio. When the gypsum content is nearly constant increasing amounts of $C_3A$ give higher strengths but above 10% $C_3A$ shrinkage increases excessively, independant of the w/c ratio (Figure 16). Measurements on different types of cement showed that aluminous cement shrinks the least, while normal blastfurnace
slag cement shrinks most at all water cement ratios.

The sequence in decreasing shrinkage of the types of cement becomes for pastes and mortars:

Portland blast-furnace slag cement class A
Portland cement class A
Portland blast-furnace slag cement class B
Portland cement class B
Portland cement class C
High-alumina cement

FIG. 14

Strength and shrinkage of cement-stone made from portland cement class B and C, portland blast-furnace slag cement class A, A winter quality and B and high-alumina cement
Strength and shrinkage of cement-stone made from portland cement clinker and 2, 5 and 8 % gypsum

When the aggregate cement ratio gets higher both strength and shrinkage become lower, so cement-stone has the highest strength and highest shrinkage.

In Figure 17 the ratio compressive strength to flexural strength
The ratio $\frac{\sigma_c}{\sigma_f}$ at dry conditions was always higher than the ratio for samples stored under water. The difference increased at higher compressive strength of the cement-stone.
For mortars with an aggregate cement ratio of 1:1 the type of cement when hardened under water had no influence on the ratio $\frac{\sigma_c}{\sigma_f}$ (Figures 17A and B). At dry conditions blast-furnace slag cement had approximately 30% higher values for $\frac{\sigma_c}{\sigma_f}$ than portland cement.

At the aggregate cement ratio of 3 there was no difference between the types of cement although the ratio again was higher under dry conditions than under water. At the aggregate cement ratio of 6 no difference could be found either between the types of cement or between the conditions. These systematics are interesting with respect to the outermost layer or skin of concrete where a transition in aggregate cement ratio from 6 or 7 to 0 occurs, going from the inside to the surface. At high flexural and compressive strengths of the bulk concrete the flexural strength of the skin of concrete can change considerably, especially under dry conditions, through a decrease of the aggregate cement ratio of the concrete + outermost layer, taking into account the large influence of the type of cement used.

Only very good curing and a low water cement ratio can make it possible that the flexural strength of the outermost layer of concrete made with blastfurnace cement is not lower than if made of portland cement (at the same compressive strength).

In case of both, surface corrosion and mechanical load (as in frost damage), the use of blastfurnace cement may lead to faster surface corrosion than the use of portland cement with the optimum amount of gypsum at equal compressive strength.

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