Material characteristics of prefabricated concrete elements from a late 19th century church in lower Austria

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Abstract

The late 19th century has preserved several important historic objects in the former Austro-Hungarian Empire made of Portland cement concrete. One of the most interesting examples of early concrete objects in Austria is the Parish church of Weissenbach an der Triesting near Vienna. The church was constructed between 1891 and 1893 using a novel double wall system: the inner wall was built from brick and the whole façade was covered with different prefabricated PC concrete elements. The present study introduces the material characteristics of some of the original concrete elements at the church. Thin and polished sections of the samples were analyzed in the optical and scanning electron microscope in order to determine the properties of the cement and aggregates. Façade elements, such as ledges, show a double-layered structure: a dense outer layer is followed by a porous tamped concrete layer to lighten the element. Composition and grains size of unhydrated clinker phases and the presence of solid fuel remnants suggest the use of a shaft kiln, inhomogeneous raw meal, firing conditions and slow cooling rates. Differences in the microporosity between the two layers refer to different w/c-ratios during the production of materials. SEM-EDX measurements and element ratio plots highlight the composition and change in the binder due to the interaction of cement hydrate phases, aggregates, and atmospheric pollutants. Additionally, dolomite aggregates exhibit the phenomenon of dedolomitization, manifesting in the transformation of CaMg(CO₃)₂ grains into calcite and brucite. The phenomenon, however, is not connected with the formation of alkali-silica gel, thus it is harmless and cannot be classified as deleterious ACR.

Keywords: Portland cement, historic concrete, clinker phases, optical microscopy

I. INTRODUCTION

Portland cement was first produced in 1856 as natural cement in Austria. The last decades of the 19th century, however, were dominated by the use of Roman cements as hydraulic binders (Zehetner, 2010). At the turn of the 20th century this trend rapidly changed and the success story of Portland cement has begun in the Austro-Hungarian Empire. Some of the most interesting applications of Portland cement in Eastern Austria were connected to the talented Austrian engineer Adolf Freiherr von Pittel (1838-1900). In the 1870s he established several cement plants near Vienna, where hydraulic lime and Roman cement were produced, but he promptly completed his array of products with prefabricated casts and artificial stones made from Portland cement (Heimel et al, 2010). In the 1870s he established several cement plants near Vienna, where hydraulic lime and Roman cement were produced, but he promptly completed his array of products with prefabricated casts and artificial stones made from Portland cement (Heimel et al, 2010). The present study is a part of an international cooperation aimed at studying the 19th century concrete buildings within the territory of the former Austro-Hungarian Empire.

Due to the abundance of large unhydrated cement grains and non-carbonated cement paste in the samples, the paper focuses on the chemical-mineralogical characterization of residual clinker phases and hydration products, attempting to determine the original composition, characteristics, and production techniques of the cement and concrete.
II. Sampling and analytical methods

Two samples, one from a slightly damaged ledge (WB-1) at the western façade (Figure 2) and another concrete fragment from the socle (WB-2) on the southern side, were taken for analyses. Thin and plane sections prepared from the samples were etched with Nital (1.5 ml of HNO$_3$ in 100 ml of isopropyl alcohol; Campbell 1999) for 7 to 9 seconds and analyzed in the optical microscope (Zeiss AXIOScope A1) using reflected and transmitted polarized light. Depending on the reactivity of the cement phases, alite normally turned blue to tan; belite was brown to blue and both phases showed details of internal structure (Campbell, 1999). Phases of the interstitial matrix (C$_3$A and ferrite) were light gray and white in color. Non-etched polished sections were coated with carbon and analyzed in the scanning electron microscope (SEM, Zeiss EVO MA 15), coupled with an energy dispersive X-ray spectrometer (EDS, Oxford DryCool, 15kV, 300 pA). A number of standardized point analyses were used to characterize the chemical composition of the binders.

III. Results and discussion

General features of the concrete samples

Based on macroscopical observations on broken surfaces (Figure 3), samples exhibited compact structure; material loss due to weathering could not be observed. By the use of phenolphthalein on fresh fracture surfaces an intensive purple discoloration (Figure 3) at ca. 0.5 to 1 cm from the surface indicated the preserved alkalinity of the binder. This phenomenon suggests adequately low capillarity, probably due to a relatively low w/c ratio of the fresh concrete (Idorn & Thaulow, 1983). On the contrary to this, the more porous tamped concrete layer (Figure 3) in the sample WB-1 did not show any discoloration with effect from the phenolphthalein test, which suggests higher capillary porosity and the influence of carbonation and/or leaching on the cement paste, due to contact with air and humidity along e.g. fitting gaps on the façade.

Optical microscopy

Confirming the phenolphthalein test, the high birefringence of the matrix in the sub-surface zones corresponds to carbonated binder portions. Estimated water to cement ratios being 0.4-0.45 (outer concrete, Figure 4, WB-1) and ~0.50 for the sample WB-2 (Figure 5). The sample WB-1 contained many irregularly shaped compaction pores (Figure 4), while the sample WB-2 had very few entrapped air voids (1 to 2 vol.%, Figure 5). The cement paste of both samples were free of any cracks.
The most characteristic feature of both samples is a large amounts of very coarse residual cement particles, exhibiting a maximum of 0.9 and an average diameter of 0.2 to 0.3 mm (Figures 6 to 11), typical for historic PC mortars and concretes (Idorn & Thaulow, 1983; Blezard, 1984, Weber et al, 2012).

A significant portion of cement grains have remained unhydrated (Figures 6 to 11). Traces of hydration could mainly be observed on the margins of the residual phases. Due to the higher w/c ratio the inner concrete layer of the sample WB-1 exhibits higher degree of hydration.

The aggregate contained in the mortars at grain sizes of 0.1 to 3 mm (WB-1) and 0.1 to 4 (WB-2) is exclusively of crushed dolomite; the estimated binder to aggregate ratio being 1 to 2.5-3 (WB-1) and 1 to 4 (WB-2) with a definitely higher amount of fine fraction (0 to 0.05 mm) in the sample WB-2, which probably also had an impact on the higher water demand and thus higher capillary porosity of the mortar (cf. Figures 4 and 5).

**Characteristics of cement phases**

Residual cement grains that entirely or partly survived the hydration and later carbonation were used to determine the composition and properties of the original cement. In terms of composition and grain size of single crystals, cement grains exhibited very inhomogeneous features. Interpretation of the observations was based on Insley (1936), Gille et al (1965), Idorn & Thaulow (1983), Blezard (1984), and Campbell (1999).

**Alite**

Residual cement grains contain large amount of unhydrated euhedral to subhedral, partly distorted C3S (Figures 7, 8 and 10). The size of single crystals...
The high birefringence of the matrix in the sub-surface zones (0.5 to 1 cm from the surface) can be observed in transmitted light or after the structural etching in reflected light. The predominant types are rounded alite (tan) and rounded belite (blue) crystals. Interstitial phases are made up of aluminate (grey) and ferrite (white). WB-1, reflected light, Nital etching.

Alite mostly contains small residual belite nuclei (e.g., Figure 8). Several crystals show corroded surfaces and marginal decomposition into belite (Figure 10). This latter observation indicates slow rates of cooling, while the different sizes of alite can also be explained by an inhomogenous raw meal and/or inhomogenous firing conditions in the kiln.

**Belite**

Belite shows round to angular, often atypical or distorted shapes. Its crystal size varies within a wide range of 5 to 100 µm, though single crystals around 20-30 µm are frequent (Figures 7 and 9). Up to 0.6 mm large nests of coarse, spherical C₂S with very little interstitial material suggest concentration and inhomogenities of quartz in the raw feed.
Segmented fingerlike sections (Figure 11) were observed both in thin and polished sections, referring to slow cooling and eventually to reducing conditions. Identical features were reported by Idorn & Thaulow (1983) and Blezard (1984) on a historic PC (Aspden’s cement) from 1850, burned in a shaft kiln. Very small belite droplets (Figure 11) could frequently be observed as inclusions within C₃A, due to slow cooling where excess silicates crystallized from the flux (Gille et al., 1965). The lamellar structure of the larger C₂S crystals can easily be observed in transmitted light or after the structural etching in reflected light. The predominant types are crystals with two or more sets of intersecting lamellae (Type I, Insley, 1936), as well as irregular grains with one set of parallel lamellae (Type II, Insley, 1936, Figure 9).
Aluminate and ferrite
Flux phases such as C₃A and Ca-Al-ferrite form very coarse, large and well-developed crystals (Figures 8, 9 and 11). These features are also consequence of a slow cooling rate.

Remnants of solid fuel and miscellaneous phases
Black, shard-like (Figure 12), in transmitted light opaque inclusions, in average of 50 to 500 μm of diameter, can be classified as solid fuel remnants. Observations in reflected cross polarized light proved the anisotropic nature of phases, suggesting the presence of graphite in the organic matter (Figure 13).

Apart from the PC cement clinker phases described above, the samples contain small quantities of other components indicating inhomogeneous firing conditions in the kiln. The most frequently observed phase is gehlenite (C₂AS) exhibiting platy shaped crystals of first order gray interference color (cf. Figure 7).

Dedolomitization of aggregates
Small amounts of the crushed dolomite sand used as aggregates shows dedolomitization in both concrete elements. The reaction products are mainly restricted to the outer rim of some of the grains, but decomposition along cracks was also observed (Figure 14). Evidence of dedolomitizations are small rims of brucite (Mg(OH)₂) and calcite (Figure 14) in the zones of interaction. The process itself has not produced expansion because no cracks originated from the dedolomitized aggregate into the cement paste were observed. The so-called deleterious alkali-carbonate-reaction (ACR) is made up of two reactions; the alkali-silica-reaction (ASR) occurs between the alkali-rich pore solution and micro- or cryptocrystalline SiO₂ as impurity in the dolomite stone and the above described dedolomitization (Katayama, 2010; Grattan-Bellew et al, 2010), which indicates the reaction between the high alkali pore solution and the CaMg(CO₃)₂. Former extensive research (Katayama, 2010) has shown that expansive forces do not happen upon dedolomitization. If expansion and formation of cracks, which reach the cement paste, are observable, the damage is always connected to ASR. Since neither microcrystalline SiO₂ in the dolomite aggregates, nor alkali-silica gel have been detected in the vicinity of the dedolomitized areas, the observed marginal decomposition of the aggregates cannot be classified as ACR.

Chemical composition of the binders
The elemental composition of hydration products, measured by EDS, according to the phenolphthalein
test, on the non-carbonated zone of the matrix, has been expressed in atomic ratio plots Ca, Si, Al and S.

\[ Si/\text{Ca} \text{ vs. } Al/\text{Ca} \text{ plots (Figures 16a and 17a) of both samples show the presence of different fine grained } \text{AFt} \text{ (ettringite) and } \text{AFm} \text{ phases. Furthermore, the } Si/\text{Ca} \text{ ratio of the inner C-S-H indicates, especially in the sample WB-2 (Figure 17b; yellow ellipse), higher values than those of the theoretical } Si/\text{Ca} \text{ ratio (0.58) given as a reference for pure synthesized } C_2\text{SH. This difference may be explained either by partly decalcified } C - S - H \text{ and/or by the influence of the clay fraction in the raw materials and subsequent clinker. The presence of portlandite confirms the alkaline environment in the binder.} \]

Values plotted in the Al/Ca vs. S/Ca plots (Figures 16b and 17b) confirm the nature of AFm phases. In the sample WB-1 (Figure 16b) the main AFm phase is monosulphate, coexisting probably also with some minor amount of monocarbonate, as suggested by the dots marked by the red ellipse. In the sample WB-2 monosulphate is completely replaced by monocarboaluminate, as characterized by the values 0.15 to 0.4 (Al/Ca) and 0.02 to 0.05 (S/Ca). The presence of monocarbonate can be explained by the high amounts of fine grained carbonate (dolomite) filler, which has partly reacted during hydration with the aluminate phases.

There is a considerable amount of sulphate-bearing phases in both samples, which suggests the presence of primary Ca-sulphate in the cement. Historical data (Riepl, 2008) show that since 1889 the Association of Austrian Cement Plants allowed a maximum gypsum content of 2% in PC, which can be interpreted as a deliberately admixed additive. However, the use of a solid fuel containing natural SO3-bearing phases (Michaelis, 1869) and its effect on the clinker composition (e.g. larger amount of alkali-sulphates formed during firing) as well as on hydrate phases cannot be excluded.

The effect of excess sulphate is suggested by the atomic ratio plots. External SO3 has reacted with monosulphate to produce secondary ettringite, which could also be detected as secondary pore fillings or sometimes in the form of reactivated C3A around unhydrated residual cement. A crack pattern due to expansive forces cannot be seen.

IV. Conclusions

Using reflected and transmitted polarized light microscopy, completed with SEM-EDS measurements on polished and thin sections, the composition and material characteristics of two historic, prefabricated PC concrete elements from the Parish church in Weissenbach an der Triesting, dating back to 1893, were carried out. The low to moderate w/c ratio and the use of historical grinding techniques have induced the preservation of several unhydrated residual cement grains in the samples. These residual grains contained all four main PC phases; their relative amount, crystal size and shape suggested, however, different burning and cooling conditions and raw feed composition in comparison with modern PC clinkers. Features such as inhomogeneous crystal sizes of alite exhibiting frequently decomposition into belite on its margin, fingerlike sections of belite and coarse crystalline flux phases suggest slow cooling rates and eventually also reducing conditions during the calcination process. Due to the coexistence of the different belite types, uneven kiln temperatures and cooling rates, typical for shaft kilns, can be assumed. Beside the aforementioned observations, for modern PC atypical clinker phases containing C2AS and probably also other low temperature, non- or slightly hydraulic Ca-(aluminate)-silicates also suggest inhomogeneous heat distribution during firing. Small fragments of graphite-bearing inclusions indicate the use of a solid fuel.

Crushed dolomite sand used as aggregate in both concrete elements exhibited partly dedolomitization on marginal zones and along cracks. The reaction, however, is not connected with the formation of ASR, thus it is harmless and cannot be defined as alkali-carbonate reaction.

Binder portions with preserved alkalinity enabled to determine the nature of hydration products based on the chemical composition of the cement paste. Although the formation of secondary ettringite from monosulphate due to excess SO42− rather suggests the use of Ca-sulphate as setting retarder admixed to the cement, the effect of sulphur-rich fuel producing alkali-sulphates in the clinker phases during firing cannot be excluded either. One of the samples clearly indicated the effect of the fine-grained carbonate aggregate portion on the hydration products: monosulphate was replaced by monocarboaluminate. In the other sample monosulphate coexisted with some minor amount of monocarbonate.

Based on the extensive investigations of the samples it can be concluded, that the cement used for producing the concrete elements was burned in a shaft kiln above the sintering temperature, but still under inhomogeneous heat distribution, with probably long residence time, slow cooling rates and inhomogeneous raw feed. Chemical data of the cement paste suggests the use of artificially admixed Ca-sulphate as a setting retarder.

The factors such as the w/c-ratio, grading curve of the aggregates and the partial reaction of the sand particles with clinker phases during the hydration have played an important role in a remarkable performance of the concrete cast elements during the last 120 years.
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REFERENCES


