Modelling and simulation of concrete carbonation: competition of several carbonation reactions

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Abstract
Concrete carbonation, i.e. the reaction of alkaline species (inside the concrete) with atmospheric carbon dioxide, is one of the major physicochemical processes compromising the service life of concrete structures. While the main carbonation reaction is that of calcium hydroxide, other constituents such as calcium silicates or calcium-silicate hydrates in the concrete can also carbonate. Many authors neglect the carbonation of these additional constituents competing with calcium hydroxide for carbon dioxide when formulating prediction models. This paper is concerned with the theoretical and numerical investigation of this competition. In particular, the effect on the simulated carbonation depth, i.e. the depth how far the carbonation layer has advanced into the concrete at any given time, is investigated. For this purpose, the concrete-carbonation process is modelled by a semi-linear coupled system of reaction-diffusion equations. For this system, a dimensional analysis is carried out and it is solved by numerical techniques. Experimental data is used for reference.

Key words: Reaction-diffusion systems, concrete carbonation, fast reactions, Thiele modulus, internal layers

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1 Introduction

Steel bars in reinforced concrete are protected from corrosion by the highly alkaline environment inside the concrete ($\text{pH} \approx 13$). If the pH decreases in the environment of the reinforcements, this protection ceases and the steel bars can corrode. Consequently, the rusting of the reinforcement usually leads to a severe reduction of the durability of the structure. The main process that destroys the protection by alkalinity is concrete carbonation, i.e. the reaction of alkaline species (in the concrete) with atmospheric carbon dioxide to produce calcium carbonate amongst other species. Detailed surveys and literature accounts on the carbonation problem and related aspects concerning the durability of concrete can be found, for instance, in [Bie88, Kro95, Cha99, MIK03, Sis04] and references therein.

The carbonation process is associated with several reaction and transport mechanisms. The main carbonation reaction is that of calcium hydroxide which may be described by

\[
\text{CO}_2(g \rightarrow aq) + \text{Ca(OH)}_2(s \rightarrow aq) \rightarrow \text{CaCO}_3(aq \rightarrow s) + \text{H}_2\text{O}. \tag{1.1}
\]

Other constituents in the concrete can also carbonate, particularly calcium-silicate hydrate (CSH) and unhydrated calcium silicates (CS), but are commonly neglected when simulating concrete carbonation. The main focus of this paper is the investigation of the influence of these (usually neglected) species and the effects of their competition with calcium hydroxide for carbon dioxide in concretes with ordinary portland cement (OPC).

A short summary of the carbonation scenario that we focus on here can be given as follows: The atmospheric carbon dioxide diffuses through the unsaturated concrete matrix, dissolves in the pore water via a Henry-like transfer mechanism, and then reacts in the presence of water with available carbonatable constituents. These are available in the pore solution by dissolution from the solid matrix. At the same time, CS hydrate (i.e. they react with water) to calcium hydroxide and CSH. Once building up, carbonates precipitate quickly to the solid matrix.

Experimental evidence suggests that the bulk of the carbonation reaction (1.1) is located on an internal reaction layer of a priori unknown width which is formed initially and progresses into the material afterwards [cf. PVF89, e.g.]. Therefore, our main interest is in the carbonation penetration depth, which determines how far this reaction layer has advanced into the concrete sample at a given time. It was also shown experimentally that the CSH starts carbonating as soon as the available calcium hydroxide has been depleted by via (1.1) [cf. Cha97, e.g.]. However, it is not clear when the carbonation of CS starts.
The carbonation reactions of the other carbonatable species are not as fast as that of calcium hydroxide but are still faster than the hydration reactions (in the late stage of hydration considered here). However, all of these reactions compete for carbon dioxide. It is the aim of this research to investigate the competition of carbonation and hydration as well as the competition between several parallel carbonation reactions. This is particularly interesting since many authors neglect all of these aspects when modelling and simulating concrete carbonation which may lead to an over-estimation of the real carbonation depth. We investigate the formation and the evolution of possible additional internal layers associated with the respective reactions of the other carbonatable constituents. This is particularly studied with respect to the layer formed due to the carbonation of calcium hydroxide. The numerical studies are based on a proper nondimensionalisation of the governing system of equations allowing easy comparisons and identification of characteristic time scales.

Cf. [Ram01, Tay97], e.g., there are various problems associated with the determination of water chemically associated with CSH. For instance, it is difficult to really distinguish this (gel) water from the free water which is present in the pores. In our model, we explicitly distinguish between the moisture (pore water and moist air) and the gel water and only consider the evolution of the moisture while assuming a constant (sufficient) amount of gel water to be present in the porous matrix. However, more research has to be initiated in order to translate the present considerations in a double-porosity model [cf. Hor97, e.g.] which might be more appropriate to describe such scenarios. Such a model allows for variations in the gel porosities as well. Evidence on concrete materials presenting a double-porosity structure has been drawn by Houst and Wittmann [cf. Hou96, HW02]. We do not strictly follow this direction here and rather try to see some of the effects due to the carbonation of CSH and unhydrated constituents on the penetration curve which is the main output of the model. This problem is also related to the various morphologies and multiple choice of stoichiometry for the CSH compound. In this work, the precise morphology and stoichiometry of CSH plays no role. For definiteness, we only consider one particular stoichiometry for CSH.

We consider the hydration process in its final period (namely, much time after the 26 hours after the mixing referred to in table 1 in [MS01], e.g.). In an accelerated test, it is usually assumed that the hydration is complete [cf. Sis04, Ste00]. Nevertheless, we account for an almost complete hydration reaction and study a special competition between hydration and carbonation of unhydrated constituents. We do not go into detail in what the modelling of the hydration is concerned but rather refer the reader to [Tay97, Ram01, PVF89] for fairly detailed descriptions of the physicochemical processes, or to [PBK01, SBK03] for a presentation of some mathematical issues concerning some of the existing hydration models.

The paper is organised as follows: In section 2, the problem under consideration is specified in detail and we particularly discuss the chemical processes associated with the concrete-carbonation problem. In section 3, these results are used to formulate a complete system of reaction-diffusion equations including initial and boundary conditions for all species under consideration. In order to investigate numerically the course of the carbonation process we introduce a weak formulation of the problem and perform its nondimensionalisation in section 4. The largest part of this paper is section 5. It contains a significant part of our numerical experiments. Basic results (penetration curves and concentration profiles) obtained when simulating an accelerated carbonation test are given. We particularly study the competition of carbonation and hydration and that between the different carbonation reactions as well as their effect on the penetration curves. Moreover, we investigate the effect of changes in the rapidness of the carbonation and hydration reactions on the penetration curves. Finally, we summarise the simulation results and conclusions in 6. The values of all parameters used in this paper can be found in the appendix.
2 Main setting

In this section we shortly describe the setting under consideration and discuss some chemical issues. Moreover, the basic geometry and notation is introduced.

2.1 Basic geometry and porosity

We focus on a part of a concrete member which is exposed to ingress of gaseous CO$_2$ and humidity from the environment. Fig. 1a shows a typical control volume (box A) in such a structure. We denote by $\Omega$ the part of the concrete sample contained in box A, for which we model the carbonation process. The dark area points out a zone $\Omega_s(t)$ or a very thin front $\Gamma(t)$ of steep change in pH dependent on the time variable $t$. This indicates that the carbonation reaction is in its fast regime: namely the characteristic reaction time is much faster than the characteristic diffusion time of the fastest active species (here CO$_2$(g)). $\Omega_s(t)$ denotes the uncarbonated zone, $\Omega_t(t)$ is the carbonated zone, and $\Gamma^N$ and $\Gamma^R$ are the interior and exterior boundaries, respectively (not shown in the figure). For ease of notation, we also define $\Gamma := \Gamma^N \cup \Gamma^R$. Our considerations are also valid for cylindrical concrete structures as long as the carbonation-reaction zone has not advanced too far into the concrete. Then, what we have in mind is depicted in Fig. 1b and $\Omega$ is the concrete part of box B. Since we are only interested in the forming and propagation of the carbonation layers within $\Omega$, we do not consider situations where the layers get close to the unexposed boundary $\Gamma^N$. We prescribe homogeneous Neumann boundary conditions at these parts (cf. section 3). Therefore, the real width of the concrete sample is of no relevance in our considerations as long as $\Omega$ is chosen wide enough. Unlike the case of moving-interface formulations [BKM03b, e.g.], we cannot say anything a priori about the widths of the carbonated and uncarbonated regions.

![Figure 1: a) Typical corner of a concrete structure. Box A is the region which our model refers to. b) Cross section of a cylindrical concrete sample. Box B is the region which our model refers to.](image)

We introduce some concepts usually needed to describe reactive processes taking place in porous media. $\Omega$ is composed of the solid matrix $\Omega^s$ and of the totality of pore voids $\Omega^p$. Furthermore, since the pore space is unsaturated and carbonation is a heterogeneous process, $\Omega^p$ splits into $\Omega^a$ (the parts filled with dry air and water vapors) and $\Omega^w$ (the parts filled with liquid water). We denote by the volumetric ratio $\phi := |\Omega^p|/|\Omega|$ the concrete porosity and by $\phi^j := |\Omega^j|/|\Omega^p|$ the air and water fractions, where $j \in \{a, w\}$. Like other authors [cf. SSV95, SV04, Ste00, e.g.], we assume a constant concrete porosity, i.e. the porosity does not change during the course of carbonation and last stage of hydration. Cf. [PVF89], we calculate it as

\[
\phi := \frac{R_w/\rho_w}{R_w/\rho_w + R_a/\rho_a + 1},
\]  

(2.1)
where $R_{w/c}$ and $R_{a/c}$ represent the water-to-cement and aggregate-to-cement ratios, while $\rho_a$, $\rho_w$ and $\rho_c$ are aggregate, water and concrete densities, respectively. Throughout the paper, the space variable is denoted by $x \in \Omega$.

### 2.2 Chemistry

It is well-known [cf. PVF89, e.g.] that if the hydration reactions are not complete, then some of the unhydrated constituents may carbonate. Consequently, we expect that a special competition between hydration and carbonation reactions may take place in the concrete structure. Such a competition does not seem to be too significant in the case of accelerated testing, mainly due to the high hydration degree which is reached by the sample during the period before the accelerated test starts. On the other hand, since much more CO$_2$ is available for reaction, the effect might be stronger yet again. For natural carbonation, these competitive effects might be stronger because of the lower (initial) degree of hydration of the sample [cf. Ste00, PVF89]. On the other hand, for a natural carbonation setting the state of hydration at a given time is generally unknown. For this reason, we focus on the simulation of accelerated tests for which the state of hydration is fairly certain at the beginning of the test. Nevertheless, the model derived in the sequel also applies to natural carbonation. The carbonation of other alkaline species like KOH, NaOH and Mg(OH)$_2$, e.g., and of the aluminate phases is neglected in this first approach. See [Ste00, e.g.] for more details on this subject.

#### 2.2.1 Hydration reactions

The main species to be hydrated are 2CaO·SiO$_2$ (dicalcium silicate, C$_2$S) and 3CaO·SiO$_2$ (tricalcium silicate, C$_3$S). The products of their hydration are 3CaO·2SiO$_2$·3H$_2$O (calcium silicate hydrate, CSH) and Ca(OH)$_2$. The hydration reactions of C$_2$S and C$_3$S are given by

\[
\begin{align*}
2(2CaO&-SiO_2) + 4H_2O &\rightarrow (3CaOcdot 2SiO_2cdot 3H_2O) + Ca(OH)_2, \quad (2.2) \\
2(3CaO&-SiO_2) + 6H_2O &\rightarrow (3CaOcdot 2SiO_2cdot 3H_2O) + 3Ca(OH)_2, \quad (2.3)
\end{align*}
\]

respectively [cf. Tay97, PVF89, e.g.]. In the hydration reactions (2.2) and (2.3), water is consumed. However, these reactions take place in the (porous) matrix space (as opposed to the pore space) at the late stage of hydration considered here. By matrix space we mean the concrete matrix which can be considered a porous medium (of low permeability) itself [see Hor97, for some details concerning the notion of double-porosity models]. We assume that there is a sufficient amount of water available in the matrix space for the hydration reactions to be completed.

#### 2.2.2 Carbonation reactions

The main carbonation reaction can be described by

\[
Ca(OH)_2 + CO_2 \rightarrow CaCO_3 + H_2O. \quad (2.4)
\]

Beside Ca(OH)$_2$, the other hydration product CSH and as well as unhydrated constituents such as C$_2$S and C$_3$S are susceptible to carbonation. Their carbonation reactions can be described via

\[
\begin{align*}
(3CaO&-2SiO_2cdot 3H_2O) + 3CO_2 &\rightarrow (3CaCO_3cdot 2SiO_2cdot 3H_2O), \quad (2.5) \\
(2CaO&-SiO_2) + 2CO_2 + \nu H_2O &\rightarrow (SiO_2cdot \nu H_2O) + 2CaCO_3, \quad (2.6) \\
(3CaO&-SiO_2) + 3CO_2 + \nu H_2O &\rightarrow (SiO_2cdot \nu H_2O) + 3CaCO_3, \quad (2.7)
\end{align*}
\]

respectively. The coefficient $\nu$ is an arbitrary positive number of moles of water that is conserved through the reactions (2.6) and (2.7). Note that these reactions, especially (2.6) and (2.7), take place in the matrix space and therefore, the water entering in these reactions is not assumed to be mobile. Moreover, also note that the carbonation reactions (2.6) and (2.7) do not directly affect
the alkalinity of the concrete. We may account for this scenario provided the concrete is sufficiently wet and all phases are available to CO$_2$.

In general, CSH has a variable stoichiometry. There are at least five different morphologies of CSH-phases which grow during the hydration process [cf. SD96, CTTJ04, DdNC05, e.g.]. In this approach, the exact morphology is not relevant and for definiteness, we restrict ourselves to the stoichiometric combination (3-2-3) (also cf. section 5). Amorphous silica gel and calcium carbonate are usually the end products of cement phases carbonation (see [KK01], [Sis04, p. 22] and references therein).

2.3 Active concentrations

We define the active concentrations (in grams per cm$^3$) by:

\[
\begin{align*}
    c_{\text{CO}_2(g)} & \quad \text{concentration of CO}_2 \text{ in pore air}, \\
    c_{\text{CO}_2} & \quad \text{concentration of CO}_2 \text{ in pore water}, \\
    c_{\text{Ca(OH)}_2} & \quad \text{concentration of Ca(OH)}_2, \\
    w & \quad \text{concentration of (mobile) moisture}, \\
    c_{\text{CaCO}_3} & \quad \text{concentration of CaCO}_3, \\
    c_{\text{CSH}} & \quad \text{concentration of CSH}, \\
    c_{\text{C}_2\text{S}} & \quad \text{concentration of C}_2\text{S}, \\
    c_{\text{C}_3\text{S}} & \quad \text{concentration of C}_3\text{S}.
\end{align*}
\]

We refer to $w$ as total humidity, or total moisture, or simply, moisture – in all cases this variable incorporates the pore water and the vapors from the air-filled parts of the pore. However, it does not include the immobile water stored in the porous matrix (like gel water, e.g.).

3 Mass balances

Before formulating the mass balances of the active species, we introduce the specific form of the production terms.

3.1 Production terms

The kinetics of the carbonation and hydration reactions are assumed to be of power-law type [cf. FB90, e.g.]. While we assume standard reaction-rate constants for the carbonation reaction (2.5)–(2.7) and the hydration reactions (2.2), (2.3), an improved reaction constant is used for the rate constant of the carbonation reaction of Ca(OH)$_2$. It incorporates the fact that the rapidness of this reaction depends on the moisture potential $w$. We only give a short summary here. For more details see [MPMB05]. Other production terms due to dissolution, precipitation and exchange with the environment are discussed afterwards.

3.1.1 Production by hydration

The production terms by hydration, $\eta_j^{\text{hydr}}$ for $j \in \{\text{C}_2\text{S}, \text{C}_3\text{S}\}$, are defined by means of power-law kinetics in the following way

\[
\eta_j^{\text{hydr}} := C_j^{\text{hydr}} (c_j^{0})^{p_j} \times (w)^{q_j}, \tag{3.1}
\]

where $C_j^{\text{hydr}} := k_j^{\text{hydr}} / (c_j^{0})^{p_j}$. Here, $c_j^{0}$ is the initial concentration of species $j$, while the factor $k_j^{\text{hydr}}$ is the reaction constant for the (hydration) reaction between constituent $j$ and water. The exponents $p_j$ are the partial reaction orders of species $j$ with respect to the hydration reaction. Following [PVF89], we choose $p_{\text{C}_2\text{S}} = 3.10$, $p_{\text{C}_3\text{S}} = 2.65$. Therefore, we define the production terms
due to hydration by
\[ f_{C_2S}^{\text{hydr}} := C_{C_2S}^{\text{hydr}} (c_{C_2S})^{p_{C_2S}} \quad \text{and} \quad f_{C_3S}^{\text{hydr}} := C_{C_3S}^{\text{hydr}} (c_{C_3S})^{p_{C_3S}}. \] (3.2)

### 3.1.2 Production by carbonation

We define the reaction rate of reaction (2.4) as
\[ \eta_j^{\text{reac}} := C_j^{\text{reac}} f_{\text{hum}}^{(w)} c_{CO_2}^{p_{CO_2}} c_{\text{Reac}(OH)}^{q_{\text{Reac}(OH)}}, \] (3.3)
where \( C_j^{\text{reac}} \) is the reaction constant. For the partial orders of reaction \( p, q \), we assume \( p, q \geq 1 \).

The factor \( f_{\text{hum}}^{(w)} \) is defined as
\[ f_{\text{hum}}^{(w)} := g_{\text{hum}}^{(RH(w))}. \] (3.4)

RH is the relative humidity calculated from \( w \). The equilibrium with the relative humidity RH is implicitly assumed by using an sorption isotherm RH(w). For a range of RH \( \in [50\%, 80\%] \) the sorption isotherm can be well approximated by an affine function, namely
\[ RH(w) = a + b \cdot \phi \cdot w, \] (3.5)
where \( a \) and \( b \) are porosity-dependent fitting parameters from [Ste00], see table 1 in the appendix for their values. The clipping factor (3.4) describes the dependence of the carbonation kinetics on RH. According to [Ste00, SSV93], it can be written as
\[ g_{\text{hum}}^{(RH)} := \begin{cases} 0, & \text{RH} \leq 0.5, \\ 5/2(\text{RH} - 0.5), & 0.5 < \text{RH} < 0.9, \\ 1, & \text{RH} > 0.9. \end{cases} \] (3.6)

We are only aware of few references where possible values for \( C_j^{\text{reac}} \) are mentioned ([Ste00, IM01, Cha99], e.g., in case of a first-order kinetics w.r.t. CO\(_2\)).

To point out the structure of the carbonation-reaction rates of reactions (2.5)–(2.7), we follow the modelling ideas in [PVF89]. For \( j \in \{ \text{CSH}, C_2S, C_3S \} \), we define the production terms by carbonation by as
\[ \eta_j^{\text{reac}} := C_j^{\text{reac}} c_{CO_2}, \] (3.7)
where \( C_j^{\text{reac}} := k_j^{\text{reac}} a_j^a \) and \( k_j^{\text{reac}} \) is the reaction constant for the reaction between the constituent \( j \) and CO\(_2\)(aq). The factor \( a_j^a \) represents the liquid-exposed surface area of the constituent \( j \). There is some uncertainty about the ratio of the constants of reactions (2.4) and (2.5). For this, only a lower estimate can be obtained from the literature, namely \( \frac{k_{\text{Ca(OH)}_2}^{\text{reac}}}{k_{\text{Ca(OH)}_2}^{\text{reac}}} > 2.4 \cdot 10^{-3} \) [cf. PVF89].

For the simulations we select a value for \( k_j^{\text{reac}} \) such that the latter estimate is satisfied. Then, we point out the effect of varying \( k_j^{\text{reac}} \) on the penetration depth. Therefore, we define the local carbonation-reaction terms as
\[ f_{\text{Ca(OH)}_2}^{\text{reac}} := C_{\text{Ca(OH)}_2}^{\text{reac}} f_{\text{hum}}^{(w)} c_{CO_2}^{p_{CO_2}} c_{\text{Reac}(OH)}^{q_{\text{Reac}(OH)}}, \] (3.8a)
\[ f_{\text{CSH}}^{\text{reac}} := C_{\text{CSH}}^{\text{reac}} \Theta(c_{\text{CSH}}) c_{CO_2}, \] (3.8b)
\[ f_{C_2S}^{\text{reac}} := C_{C_2S}^{\text{reac}} \Theta(c_{C_2S}) c_{CO_2}, \] (3.8c)
\[ f_{C_3S}^{\text{reac}} := C_{C_3S}^{\text{reac}} \Theta(c_{C_3S}) c_{CO_2}, \] (3.8d)
where \( \Theta: R \rightarrow \{0, 1\} \) is the Heaviside function,
\[ \Theta(u) = \begin{cases} 1, & u > 0, \\ 0, & u \leq 0. \end{cases} \] (3.9)
3.1.3 Other production terms and boundary conditions

We model the interfacial mass transfer from pore air to pore water by means of Henry’s law. The corresponding production term has the form

\[ f_{\text{Henry}}^\text{ex} := C_{\text{Henry}}^\text{ex} (c_{\text{CO}_2(g)}^\text{Henry} - c_{\text{CO}_2}) , \]

where \( C_{\text{Henry}}^\text{ex} \) is the interfacial mass-transfer coefficient and \( c_{\text{CO}_2(g)}^\text{Henry} \) is the dimensionless Henry constant.

We implicitly assume instantaneous dissolution and precipitation of some of the carbonation reactants and products. Moreover, we assume homogeneous Neumann boundary conditions for all diffusing species at the interior boundary. At the exterior boundary, Robin boundary conditions are assumed for \( c_{\text{CO}_2(g)} \) and \( w \) while homogeneous Neumann boundary conditions are applied for all other diffusing species. The fluxes across the external boundary are expressed as

\[
\begin{align*}
\eta_{\text{CO}_2(g)}^\text{Rob} &:= C_{\text{CO}_2(g)}^\text{Rob} (c_{\text{CO}_2(g)} - c_{\text{CO}_2}^\text{ext}), \\
\eta_{\text{H}_2\text{O}}^\text{Rob} &:= C_{\text{H}_2\text{O}}^\text{Rob} (w - w^\text{ext}),
\end{align*}
\]

where the constants \( C_{\text{CO}_2(g)}^\text{Rob} \) and \( C_{\text{H}_2\text{O}}^\text{Rob} \) represent mass-transfer coefficients at the external boundary and \( c_{\text{CO}_2}^\text{ext} \) and \( w^\text{ext} \) are given exterior concentration of \( \text{CO}_2 \) and the moisture potential, respectively.

3.2 Mass balances of active species

We formulate the macroscopic mass balances for \( \text{CO}_2 \) in both phases (gaseous and liquid) and for \( \text{Ca(OH)}_2, \, \text{CaCO}_3, \, \text{CSH}, \, \text{C}_2\text{S} \) and \( \text{C}_3\text{S} \) in the liquid phase. The fact that the latter species should actually be associated with the solid matrix is taken care of via the structure of the reaction-rate constants. For moisture, we assume a local equilibrium of the gas and liquid phase [see Ste00, Arf98, Gru97, e.g.]. Moreover, we do not further distinguish between the products \( 3\text{CaCO}_3, \, 2\text{SiO}_2 \cdot 3\text{H}_2\text{O} \) and \( 3\text{CaCO}_3 \). A more detailed description on the general modelling aspects of the carbonation problem can be found in [BKM03b, BKM03a, Mun05].

We formulate the system of reaction-diffusion equations independent of the space dimension. The time interval of interest is denoted by \( S \). For numerical simulations, we only consider the case of one space dimension. In the formulation of the mass balances, we make use of some additional notation, namely \( m_j \) (the molar mass of species \( j \)) and \( D_j \) (the microscopic diffusivity of species \( j \)), \( \nu \) (the outer normal unit vector), and the superscript \( 0 \) which denotes initial concentrations. Based on the discussion in the previous subsections, the complete model can be formulated in the following manner:

**Mass balance for \( \text{CO}_2(g) \):**

\[
\begin{align*}
&\partial_t (\phi \phi^w c_{\text{CO}_2(g)}(x, t)) - \nabla \cdot (D_{\text{CO}_2(g)} \phi \phi^w \nabla c_{\text{CO}_2(g)}(x, t)) = -f_{\text{Henry}}^\text{ex}(x, t), \quad x \in \Omega, \ t \in S, \\
&- (D_{\text{CO}_2(g)} \phi \phi^w \nabla c_{\text{CO}_2(g)}(x, t)) \cdot \nu = 0, \quad x \in \Gamma^N, \ t \in S, \\
&- (D_{\text{CO}_2(g)} \phi \phi^w \nabla c_{\text{CO}_2(g)}(x, t)) \cdot \nu = C_{\text{CO}_2(g)}^\text{Rob} (c_{\text{CO}_2(g)}(x, t) - c_{\text{CO}_2}^\text{ext}), \quad x \in \Gamma^\text{R}, \ t \in S, \\
&c_{\text{CO}_2(g)}(x, 0) = c_{\text{CO}_2(g)}^0(x), \quad x \in \Omega.
\end{align*}
\]

**Mass balance for \( \text{CO}_2 \):**

\[
\begin{align*}
&\partial_t (\phi \phi^w c_{\text{CO}_2}(x, t)) - \nabla \cdot (D_{\text{CO}_2} \phi \phi^w \nabla c_{\text{CO}_2}(x, t)) = f_{\text{Henry}}^\text{ex}(x, t) \\
&+ m_{\text{CO}_2} \phi \phi^w \left( - f_{\text{Ca(OH)}_2}^\text{reac}(x, t) - 3 f_{\text{CSH}}^\text{reac}(x, t) - 2 f_{\text{C}_2\text{S}}^\text{reac}(x, t) - 3 f_{\text{C}_3\text{S}}^\text{reac}(x, t) \right), \quad x \in \Omega, \ t \in S, \\
&- (D_{\text{CO}_2} \phi \phi^w \nabla c_{\text{CO}_2}(x, t)) \cdot \nu = 0, \quad x \in \Gamma, \ t \in S, \\
&c_{\text{CO}_2}(x, 0) = c_{\text{CO}_2}^0(x), \quad x \in \Omega.
\end{align*}
\]
Mass balance for Ca(OH)$_2$:

\[
\begin{aligned}
\partial_t (\phi \phi^w c_{\text{Ca(OH)}_2}(x,t)) - \nabla \cdot \left( D_{\text{Ca(OH)}_2} \phi \phi^w \nabla c_{\text{Ca(OH)}_2}(x,t) \right) &= m_{\text{Ca(OH)}_2} \phi \phi^w \left( -f_{\text{Ca(OH)}_2}^{\text{reac}}(x,t) + \frac{1}{2} f_{\text{CSH}}^{\text{hydr}}(x,t) + \frac{3}{2} f_{\text{C}_3S}^{\text{hydr}}(x,t) \right), \\
-(D_{\text{Ca(OH)}_2} \phi \phi^w \nabla c_{\text{Ca(OH)}_2}(x,t)) \cdot \nu &= 0, \quad x \in \Gamma, \ t \in S, \\
s_{\text{Ca(OH)}_2}(x,0) &= c^0_{\text{Ca(OH)}_2}(x), \quad x \in \Omega.
\end{aligned}
\]  

(3.14a)

(3.14b)

(3.14c)

Mass balance for (mobile) moisture:

\[
\begin{aligned}
\partial_t (\phi w(x,t)) - \nabla \cdot (D_{\text{H}_2\text{O}} \phi \nabla w(x,t)) &= m_{\text{H}_2\text{O}} \phi \phi^w f_{\text{Ca(OH)}_2}^{\text{reac}}(x,t), \quad x \in \Omega, \ t \in S, \\
-(D_{\text{H}_2\text{O}} \phi \nabla w(x,t)) \cdot \nu &= 0, \quad x \in \Gamma^N, \ t \in S, \\
-(D_{\text{H}_2\text{O}} \phi \nabla w(x,t)) \cdot \nu &= C^{\text{Rob}}_{\text{H}_2\text{O}} (w(x,t) - w^{\text{ext}}(x,t)), \quad x \in \Gamma^R, \ t \in S, \\
w(x,0) &= w^0(x), \quad x \in \Omega.
\end{aligned}
\]

(3.15a)

(3.15b)

(3.15c)

(3.15d)

Mass balance for CaCO$_3$:

\[
\begin{aligned}
\partial_t (\phi \phi^w c_{\text{CaCO}_3}(x,t)) &= m_{\text{CaCO}_3} \phi \phi^w \left( f_{\text{Ca(OH)}_2}^{\text{reac}}(x,t) + 3 f_{\text{CSH}}^{\text{reac}}(x,t) + 2 f_{\text{C}_2\text{SiO}_3}^{\text{reac}}(x,t) + 3 f_{\text{C}_3\text{SiO}_3}^{\text{reac}}(x,t) \right), \quad x \in \Omega, \ t \in S, \\
c_{\text{CaCO}_3}(x,0) &= c^0_{\text{CaCO}_3}(x), \quad x \in \Omega.
\end{aligned}
\]

(3.16a)

(3.16b)

Mass balance for CSH:

\[
\begin{aligned}
\partial_t (\phi \phi^w c_{\text{CSH}}(x,t)) &= m_{\text{CSH}} \phi \phi^w \left( -f_{\text{CSH}}^{\text{reac}}(x,t) + \frac{1}{2} f_{\text{C}_2\text{SiO}_3}^{\text{hydr}}(x,t) + \frac{1}{2} f_{\text{C}_2\text{SiO}_3}^{\text{hydr}}(x,t) \right), \quad x \in \Omega, \ t \in S, \\
c_{\text{CSH}}(x,0) &= c^0_{\text{CSH}}(x), \quad x \in \Omega.
\end{aligned}
\]

(3.17a)

(3.17b)

Mass balance for C$_2$S:

\[
\begin{aligned}
\partial_t (\phi \phi^w c_{\text{C}_2\text{S}}(x,t)) &= m_{\text{C}_2\text{S}} \phi \phi^w \left( -f_{\text{C}_2\text{SiO}_3}^{\text{reac}}(x,t) - f_{\text{C}_2\text{SiO}_3}^{\text{hydr}}(x,t) \right), \quad x \in \Omega, \ t \in S, \\
c_{\text{C}_2\text{S}}(x,0) &= c^0_{\text{C}_2\text{S}}(x), \quad x \in \Omega.
\end{aligned}
\]

(3.18a)

(3.18b)

Mass balance for C$_3$S:

\[
\begin{aligned}
\partial_t (\phi \phi^w c_{\text{C}_3\text{S}}(x,t)) &= m_{\text{C}_3\text{S}} \phi \phi^w \left( -f_{\text{C}_2\text{SiO}_3}^{\text{reac}}(x,t) - f_{\text{C}_2\text{SiO}_3}^{\text{hydr}}(x,t) \right), \quad x \in \Omega, \ t \in S, \\
c_{\text{C}_3\text{S}}(x,0) &= c^0_{\text{C}_3\text{S}}(x), \quad x \in \Omega.
\end{aligned}
\]

(3.19a)

(3.19b)

(3.19c)

For consistency, we also included the factors $\phi \phi^w$ in (3.17)–(3.19). Therefore, the formulation (3.12)–(3.19) is also true for porosities dependent on space or time.

4 Numerical implementation

In this section, we first present a weak formulation of the model (3.12)–(3.19). Afterwards, we perform a nondimensionalisation of all quantities which results in the final system of equations to be implemented using the finite element method [analogously as in MPMB05].
4.1 Weak formulation

We reformulate the system (3.12)–(3.19) in terms of macroscopic quantities. More precisely, we perform a transformation of the quantities from the previous section into volume-averaged concentrations of the form

\[ \tilde{c}_{CO_2}(g) := \phi \tilde{\rho}_{CO_2}(g), \quad \tilde{c}_{CO_2} := \phi \tilde{\rho}_{CO_2}, \quad \tilde{w} := \phi w, \quad \text{etc.} \tag{4.1} \]

We exclusively use the macroscopic quantities in the following, so for ease of notation we omit the tilde from now on. The main advantage of this procedure is that the porosities solely appear on the right-hand sides of the equations, i.e. in the production terms.

Define the spaces \( V \) and \( W \) as

\[ V = H^1(0, T; L^2(\Omega)), \tag{4.2} \]
\[ W = \{ v \in L^2(0, T; H^1(\Omega)) \mid \partial_v \in L^2(0, T; (H^1(\Omega))^2) \}, \tag{4.3} \]

and denote the \( L^2(\Omega) \)-scalar product by \((\cdot, \cdot)_\Omega\). Note that we have \( \partial \Omega = \Gamma^N \cup \Gamma^R \) (disjoint). See [DL92] for the definition of the function spaces in (4.2) and (4.3).

The weak formulation of problem (3.12)–(3.19) in terms of macroscopic concentrations is given by

\[ c_{CO_2}(g) \in W, \quad c_{CO_2}(0) = \phi \tilde{\rho}_{CO_2}(g) \quad \text{such that} \]
\[ (\partial_t c_{CO_2}(g) \mid v)_\Omega + D_{CO_2}(v c_{CO_2}(g) \mid \nabla v)_\Omega = -f_{Henry}(v)_{\Omega} - c_{Rob}(c_{CO_2}(g) - \phi \tilde{\rho}_{CO_2}(g)) v, \tag{4.4} \]

\[ c_{CO_2} \in W, \quad c_{CO_2}(0) = \phi \tilde{\rho}_{CO_2} \quad \text{such that} \]
\[ (\partial_t c_{CO_2} \mid v)_\Omega + D_{CO_2}(v c_{CO_2} \mid \nabla v)_\Omega = f_{Henry}(v)_\Omega + m_{CO_2}(-r_{\text{reac}_{Ca(OH)_2}} - 3f_{\text{reac}_{C_{CSH}}} - 2c_{S} - 3f_{\text{reac}_{C_{2S}}}) v, \tag{4.5} \]

\[ c_{Ca(OH)_2} \in W, \quad c_{Ca(OH)_2}(0) = \phi \tilde{\rho}_{Ca(OH)_2} \quad \text{such that} \]
\[ (\partial_t c_{Ca(OH)_2} \mid v)_\Omega + D_{Ca(OH)_2}(v c_{Ca(OH)_2} \mid \nabla v)_\Omega = m_{Ca(OH)_2}(-r_{\text{reac}_{Ca(OH)_2}} + \frac{1}{2}f_{\text{hydr}_{x,t}} + \frac{3}{2}f_{\text{hydr}_{x,t}}) v, \tag{4.6} \]

\[ w \in W, \quad w(0) = \phi w^0 \quad \text{such that} \]
\[ (\partial_t w \mid v)_\Omega + D_{H_2O}(v w) \mid \nabla v)_\Omega = m_{H_2O}(f_{\text{reac}_{Ca(OH)_2}}(v)_\Omega - c_{Rob}(w - \phi w^0) v, \tag{4.7} \]

where each equation needs to be satisfied for a.e. in \( S \) and all \( v \in W \) as well as

\[ c_{CaCO_3} \in V, \quad c_{CaCO_3}(0) = \phi \tilde{\rho}_{CaCO_3} \quad \text{such that} \]
\[ (\partial_t c_{CaCO_3} \mid v)_\Omega = m_{CaCO_3}(r_{\text{reac}_{CaCO_3}} + 3f_{\text{reac}_{C_{CSH}}} + 2f_{\text{reac}_{C_{2S}}} + 3f_{\text{reac}_{C_{2S}}}) v, \tag{4.8} \]

\[ c_{CSH} \in V, \quad c_{CSH}(0) = \phi \tilde{\rho}_{CSH} \quad \text{such that} \]
\[ (\partial_t c_{CSH} \mid v)_\Omega = m_{CSH}(c_{CSH} - c_{CO_2} - c_{CS_3}) v, \tag{4.9} \]

\[ c_{CS_3} \in V, \quad c_{CS_3}(0) = \phi \tilde{\rho}_{CS_3} \quad \text{such that} \]
\[ (\partial_t c_{CS_3} \mid v)_\Omega = m_{CS_3}(-r_{\text{reac}_{f_{\text{hydr}_{x,t}}}} - c_{CS_3} v, \tag{4.10} \]

where each equation needs to be satisfied a.e. in \( S \) for all \( v \in V \). Some of the production terms...
need to be re-defined due to the switch from microscopic to macroscopic quantities. These are

\[ f_{\text{Henry}} := C^{\text{ex}}(C^{\text{Henry}} (\phi^n)^{-1} c_{\text{CO}_2(g)} - (\phi^w)^{-1} c_{\text{CO}_2}), \]
\[ f_{\text{Ca(OH)_2}} := C^{\text{ex}}_{\text{Ca(OH)_2}} f_{\text{Henry}} (\phi^w)^{1-p-q} (c_{\text{Ca(OH)_2}})^q, \]
\[ f_{\text{hydr}}^{C_{2S}} := C_{\text{hydr}}^{C_{2S}} (\phi^w)^{1-p_{C_{2S}}} (c_{\text{Ca(OH)_2}})^q, \]
\[ f_{\text{hydr}}^{C_{3S}} := C_{\text{hydr}}^{C_{3S}} (\phi^w)^{1-p_{C_{3S}}} (c_{\text{Ca(OH)_2}})^q. \]

### 4.2 Nondimensionalisation

We introduce the following nondimensional quantities,

\[ u_1 := c_{\text{CO}_2(g)} / c^m_1, \quad u_2 := c_{\text{CO}_2} / c^m_2, \quad u_3 := c_{\text{Ca(OH)_2}} / c^m_3, \]
\[ u_4 := w / c^m_4, \quad u_5 := c_{\text{CaCO}_3} / c^m_5, \quad u_6 := c_{\text{CSH}} / c^m_6, \]
\[ u_7 := c_{\text{CSH}} / c^m_7, \quad u_8 := c_{\text{CaCO}_3} / c^m_8, \]

where \( c^m_j \), \( j = 1, \ldots, 8 \), are some maximal concentrations. In order to make a reasonable choice for the \( c^m_j \) and to simplify the model, we make the following assumptions:

1. \( c_{\text{CO}_2(g)}(0) = c_{\text{CO}_2}(0) = c_{\text{CaCO}_3}(0) = 0 \).
2. \( c^0_{\text{CO}_2(g)}, w^0, c^0_{\text{Ca(OH)_2}}, w^0, c^0_{\text{CaCO}_3}, c^0_{\text{CSH}}, c^0_{\text{CSH}}, c^0_{\text{CaCO}_3} \) are positive constants.
3. Diffusion of the species in water is sufficiently slow compared to diffusion in air.

Note that assumptions 1–3 are not inevitable but very much simplify making an appropriate choice for the \( c^m_j \). They lead to the following definitions of the maximal concentrations:

\[ c^m_1 := \phi^n c^0_{\text{CO}_2(g)}, \]
\[ c^m_2 := \phi^w C^{\text{Henry}} c^0_{\text{CO}_2(g)}, \]
\[ c^m_3 := \phi^n c^0_{\text{Ca(OH)_2}} + \frac{1}{2} m_{\text{CaCO}_3} \phi^w c^0_{\text{CaCO}_3} + \frac{3}{2} m_{\text{Ca(OH)_2}} \phi^w c^0_{\text{CaCO}_3}, \]
\[ c^m_4 := \max \left\{ \frac{m_{\text{H}_2O}}{m_{\text{Ca(OH)_2}}} c^m_3 + \phi w^0, \phi w^0 \right\}, \]
\[ c^m_5 := \frac{m_{\text{CaCO}_3}}{m_{\text{Ca(OH)_2}}} c^m_3 + \frac{3}{2} m_{\text{CaCO}_3} c^m_3 + \frac{2}{m_{\text{Ca(OH)_2}}} c^m_6 + \frac{3}{m_{\text{CaCO}_3}} c^m_8, \]
\[ c^m_6 := \phi^n c^0_{\text{CSH}} + \frac{1}{2} m_{\text{CaCO}_3} \phi^w c^0_{\text{CSH}} + \frac{1}{2} m_{\text{CaCO}_3} \phi^w c^0_{\text{CSH}}, \]
\[ c^m_7 := \phi^n c^0_{\text{CSH}}, \]
\[ c^m_8 := \phi^n c^0_{\text{CSH}}. \]

Define a characteristic diffusion time for the fastest species involved, \( \text{CO}_2(g) \), as

\[ T := L^2 / D_{\text{CO}_2(g)}, \]

and let \( \tilde{t} := t / T \) and \( \tilde{x} := x / L \) be the nondimensional time and space coordinates. An analogous nondimensionalisation was also used for the model considered in [MPMB05].
With definitions (4.16)–(4.18) we are led to the introduction of the following dimensionless combinations,

\[
\begin{align*}
\beta_2 & := \frac{c_2^m}{c_1^m}, & \beta_3 & := \frac{c_3^m m_{\text{CO}_2}}{c_1^m m_{\text{Ca(OH)}_2}}, & \beta_4 & := \frac{c_4^m m_{\text{CO}_2}}{c_1^m m_{\text{H}_2\text{O}}}, & \beta_5 & := \frac{c_5^m m_{\text{CO}_2}}{c_1^m m_{\text{CaCO}_3}}, \\
\beta_6 & := \frac{c_6^m m_{\text{CO}_2}}{c_1^m m_{\text{CSH}}}, & \beta_7 & := \frac{c_7^m m_{\text{CO}_2}}{c_1^m m_{\text{CS}_2}}, & \beta_8 & := \frac{c_8^m m_{\text{CO}_2}}{c_1^m m_{\text{CS}_3}}.
\end{align*}
\]

\[
\begin{align*}
\delta_2 & := \frac{D_{\text{CO}_2}}{D_{\text{CO}_2}(g)}, & \delta_3 & := \frac{D_{\text{Ca(OH)}_2}}{D_{\text{CO}_2}(g)}, & \delta_4 & := \frac{D_{\text{H}_2\text{O}}}{D_{\text{CO}_2}(g)}.
\end{align*}
\]

\[
W_{\text{H}_{\text{en}}} := \frac{L^2}{D_{\text{CO}_2}(g)} C_{\text{ex}}, \quad W_{\text{H}_{1}} := \frac{L}{D_{\text{CO}_2}(g)} C_{\text{Rob}}, \quad W_{\text{H}_{4}} := \frac{L}{D_{\text{CO}_2}(g)} C_{\text{Rob}},
\]

\(\Phi^2 := \frac{L^2 m_{\text{CO}_2} (c_2^m)^p (c_3^m)^q}{D^2_{\text{CO}_2}(g) c_1^{2m}} C_{\text{reac}}^\text{Ca(OH)$_2$}, \quad (\text{Thiele modulus})
\]

\[
\begin{align*}
R_6 & := \frac{L^2 m_{\text{CO}_2} c_6^m}{D_{\text{CO}_2}(g) c_1^m} C_{\text{reac}}^\text{CSH}, & R_7 & := \frac{L^2 m_{\text{CO}_2} c_7^m}{D_{\text{CO}_2}(g) c_1^m} C_{\text{reac}}^\text{CS}_2, & R_8 & := \frac{L^2 m_{\text{CO}_2} c_8^m}{D_{\text{CO}_2}(g) c_1^m} C_{\text{reac}}^\text{CS}_3,
\end{align*}
\]

\[
H_7 := \frac{L^2 m_{\text{CO}_2} (c_3^m)^p(c_4^m)^q}{D_{\text{CO}_2}(g) c_1^m} C_{\text{hydr}}^\text{CS}_2, & H_8 := \frac{L^2 m_{\text{CO}_2} (c_4^m)^p(c_5^m)^q}{D_{\text{CO}_2}(g) c_1^m} C_{\text{hydr}}^\text{CS}_3.
\]

The parameters \(\beta_j\) are usually called capacity factors, whereas the \(\delta_j\) are ratios comparing each diffusivity with that of \(\text{CO}_2(g)\), see [PVFS89, MPMB05]. The Thiele modulus \(\Phi^2\) as well as the factors \(R_j\) describe the rapidness of the carbonation reactions where as the \(H_j\) describe the rapidness of the hydration reactions. Furthermore, the factors \(W_{\text{H}_{\text{en}}}, W_{\text{H}_{1}}^{\text{Rob}}\) and \(W_{\text{H}_{4}}^{\text{Rob}}\) account for the rapidness of the different types of interfacial mass transfer.

For notational purposes we finally define

\[
\begin{align*}
\phi_{\text{H}1} & := \phi_{\text{H}1}^\text{ext} \frac{c_2^m}{c_1^m}, & \phi_{\text{H}4} & := \phi_{\text{H}4}^\text{ext} \frac{c_6^m}{c_1^m}, & \phi_{\text{H}5} & := \phi_{\text{H}5}^\text{ext} \frac{c_6^m}{c_1^m}, & \phi_{\text{H}6} & := \phi_{\text{H}6} \frac{c_2^m}{c_3^m}.
\end{align*}
\]

Transformation of system (4.4)–(4.11) to the dimensionless quantities yields the system of equations to be solved numerically:

\[
\begin{align*}
u_1 & \in \mathcal{W}, & u_1(0) = 0 \text{ such that } (\partial_\nu u_1 | v)_\Omega + (\nabla u_1 | \nabla v)_\Omega = -(f_{\text{Henry}} | v)_\Omega - W_{\text{H}_{1}}^{\text{Rob}} (u_1 - u_1^{\text{ext}} | v)_\Gamma^n, \\
u_2 & \in \mathcal{W}, & u_2(0) = 0 \text{ such that } \beta_2 (\partial_\nu u_2 | v)_\Omega + \beta_2 \delta_2 (\nabla u_2 | \nabla v)_\Omega = +(f_{\text{Henry}} | v)_\Omega + (f_{\text{reac}}^3 - 3 f_{\text{reac}}^4 - 2 f_{\text{reac}}^5 - 3 f_{\text{reac}}^6 | v)_\Omega, \\
u_3 & \in \mathcal{W}, & u_3(0) = 1 \text{ such that } \beta_3 (\partial_\nu u_3 | v)_\Omega + \beta_3 \delta_3 (\nabla u_3 | \nabla v)_\Omega = (-f_{\text{reac}}^3 + \frac{1}{2} f_{\text{hydr}}^\text{CS}_2 + \frac{3}{2} f_{\text{hydr}}^\text{CS}_3 | v)_\Omega, \\
u_4 & \in \mathcal{W}, & u_4(0) = u_4^0 \text{ such that } \beta_4 (\partial_\nu u_4 | v)_\Omega + \beta_4 \delta_4 (\nabla u_4 | \nabla v)_\Omega = (f_{\text{reac}}^4 | v)_\Omega - W_{\text{H}_{4}}^{\text{Rob}} \beta_4 (u_4 - u_4^{\text{ext}} | v)_\Gamma^n.
\end{align*}
\]
4.3 Carbonation depth

where each equation has to be satisfied for a.e. $t \in S$ and for all $v \in W$ as well as

\[
\begin{align*}
 u_5 \in V, \quad u_5(0) &= 0 \
 & \text{such that } \beta_5(\partial_t u_5 | v) = (f_3^\text{reac} + 3 f_6^\text{reac} + 2 f_7^\text{reac} + 3 f_8^\text{reac} | v)\Omega 
\end{align*}
\]

\[(4.25)\]

\[
\begin{align*}
 u_6 \in V, \quad u_6(0) &= u_6^0 \
 & \text{such that } \beta_6(\partial_t u_6 | v) = (-f_6^\text{reac} + 1/2 f_7^\text{hydr} + 1/2 f_8^\text{hydr} | v)\Omega 
\end{align*}
\]

\[(4.26)\]

\[
\begin{align*}
 u_7 \in V, \quad u_7(0) &= u_7^0 \
 & \text{such that } \beta_7(\partial_t u_7 | v) = (-f_7^\text{reac} - f_7^\text{hydr} | v)\Omega 
\end{align*}
\]

\[(4.27)\]

\[
\begin{align*}
 u_8 \in V, \quad u_8(0) &= u_8^0 \
 & \text{such that } \beta_8(\partial_t u_8 | v) = (-f_7^\text{reac} - f_8^\text{hydr} | v)\Omega 
\end{align*}
\]

\[(4.28)\]

where each equation has to be satisfied for a.e. $t \in S$ and for all $v \in V$. The dimensionless production terms are

\[
\begin{align*}
 f^\text{Henry} &= W^\text{Hen}(C^\text{Hen}(\phi \sigma)^{-1} u_1 - (\phi \sigma)^{-1} \beta_2 u_2), \\
 f_3^\text{reac} &= \Phi^2(\phi \sigma)^{1-p-q} f_{\text{hum}}(u_4 \rho^m \phi \sigma^{-1}) u_2^p u_3^q, \\
 f_j^\text{reac} &= R_j \Theta(u_j), \quad j \in \{6, 7, 8\}, \\
 f_k^\text{hydr} &= H_k(\phi \sigma)^{1-p_k} u_k^p, \quad k \in \{7, 8\}. 
\end{align*}
\]

\[(4.29)\] \quad \[(4.30)\] \quad \[(4.31)\] \quad \[(4.32)\]

In the numerical implementation, we approximate the Heaviside function $\Theta$ by

\[
\Theta^\text{approx}(x) := \frac{x}{x + \gamma} 
\]

\[(4.33)\]

for $x \geq 0$ with $0 < \gamma \ll 1$ in order to avoid numerical problems.

4.3 Carbonation depth

As discussed earlier, we are interested in the prediction of the carbonation penetration depth, i.e. in how far the carbonation reaction of Ca(OH)$_2$ has advanced into the concrete sample at any given time. We define the penetration depth to be the isoline which corresponds to a (dimensionless) Ca(OH)$_2$-concentration equal to 0.1,

\[
s(t) := \{ x \in \Omega | u_3(x, t) = 0.1 \} \quad \text{for each } t \in S. 
\]

\[(4.34)\]

Analogous definitions of the carbonation front can be found in [SSV95, SSV93], e.g. We follow here the way indicated in [Ste00, SDA02].

4.4 Numerical solution

The equations (4.21)--(4.28) form a weakly-coupled system of semi-linear parabolic partial and ordinary differential equations. As in [MPMB05], we solve it numerically in one space-dimension by using a standard finite element discretisation method. More precisely, we accomplish a semi-discretisation in space on a uniform mesh of width $h = 1/(n - 1)$ by the Galerkin method. For the test and trial functions, first-order splines are used. In addition, we apply the standard mass-lumping-scheme, cf. [KA00], e.g. See [GM03] for a detailed description of a similar discretisation problem. The nonlinear terms $f_j^\text{reac}$ and $f_k^\text{hydr}$ are approximated by the trapezoidal rule.

The resulting stiff system of $8 \times n$ ODEs is numerically integrated using the MATLAB ODE stiff solver ode15s, which is a variable-order solver based on numerical-differentiation formulas (NDFs).

The examples in the following section are obtained by choosing $n = 80$.  

\[\text{1See www.mathworks.com for details and further references.}\]
5 Numerical simulation of an accelerated carbonation test

In figures 2 and 3, we show dimensionless concentration profiles of all involved species ($\text{CO}_2(g)$, $\text{CO}_2(aq)$, $\text{Ca(OH)}_2$, moisture, $\text{CaCO}_3$, CSH, $\text{C}_2\text{S}$ and $\text{C}_3\text{S}$) as well as the carbonation depth. For this, we use a set of parameters for the accelerated carbonation scenario based on data from [PVF89] (cf. the appendix for values of the parameters). We refer to this set of parameters as the standard set in the sequel. The experimental data we are plotting for the carbonation depth is also due to [PVF89].

The characteristic time scale of the carbonation reaction of $\text{Ca(OH)}_2$ is essentially faster than the characteristic time scales of the carbonation reactions of the other species and of the hydration reactions. For the standard set of parameters, we have

$$\Phi^2 \approx 10^3, \quad R_6 \approx 7, \quad R_7 \approx 5, \quad R_8 \approx 4, \quad H_7 \approx 5 \cdot 10^{-4}, \quad H_8 \approx 2 \cdot 10^{-4}.$$ 

Moreover, the characteristic diffusion time scale of $\text{CO}_2(g)$ is considerably faster than those of all other species (we have $\delta_1 = 1$ while the second largest is $\delta_2 \approx 8 \cdot 10^{-6}$).

The present formulation does not allow the diffusion of the additional phases CSH, $\text{C}_2\text{S}$ and $\text{C}_3\text{S}$. However, we also investigated what happens if they diffused with the diffusivity of ions in pore water. There is only a slight change in the quantitative results and none in the qualitative results. This is basically due to the fact that the ionic diffusivity is very small compared to that of $\text{CO}_2(g)$.

Compared to the simulation results of the simplified model [MPMB05], the concentration profiles shown in figures 2 and 3 differ only marginally, except that of $\text{CaCO}_3$ which now shows a bend in the decrease. This is due to the fact that in this model, there is more than one source producing $\text{CaCO}_3$. Moreover, the additional profiles of the concentrations of CSH, $\text{C}_2\text{S}$, and $\text{C}_3\text{S}$ show a similar behaviour like that of $\text{Ca(OH)}_2$. It can be observed that due to hydration reactions, the concentrations of $\text{C}_2\text{S}$ and $\text{C}_3\text{S}$ also decrease in the uncarbonated part of the concrete sample while those of $\text{CaCO}_3$ and CSH increase. However, these effects are comparably small for the chosen set of parameters.

Note that assuming a constant moisture profile instead of choosing a PDE for moisture does not change the qualitative results. Even the quantitative change is fairly small since the moisture content only enters in the improved reaction rate of the carbonation reaction of $\text{Ca(OH)}_2$ (cf. 3.1). Moreover, we always choose the mass-transfer coefficients at the external boundary of the concrete sample very high in order to account for Dirichlet boundary conditions. This seems to be a standard choice when simulating accelerated carbonation tests. For a detailed discussion of the effect of the moisture on the carbonation penetration we refer to [MPMB05].

The CSH-phases admit several stoichiometries [cf. SD96]. In our model, we only account for a particular stoichiometry for CSH (namely, 3-2-3). This does not influence the qualitative results although other stoichiometries may consume more $\text{CO}_2$ when carbonating and they may also have different (carbonation-) reaction constants (they also have different molar masses). We investigate the effects due to a change of the carbonation-reaction constant in section 5.2.1. This numerical test may be thought of as considering different CSH stoichiometries.

Since an extensive study of the effects due to variations of some of the relevant parameters was performed for the simplified model in [MPMB05] (neglecting CSH, $\text{C}_2\text{S}$, and $\text{C}_3\text{S}$), we concentrate on additional effects basically due to the inclusion of the other phases. We investigate the influence of each carbonation reaction, in particular the consumption of $\text{CO}_2$ by each reaction and the alternate neglect of CSH, $\text{C}_2\text{S}$, and $\text{C}_3\text{S}$ (cf. section 5.1) as well as the effects due to a variation of the carbonation- and hydration-rate constants (cf. section 5.2).
Figure 2: Concentration profiles of the involved species obtained with the standard set of parameters of the accelerated test.
Figure 3: Profiles of concentrations of $C_2S$ and $C_3S$ as well as carbonation depth obtained with the standard set of parameters of the accelerated test.
5.1 Influence of the competition of parallel carbonation reactions

We want to compare the influence of each carbonation reaction on the penetration depth. In figure 4, the dimensionless production terms

\[ n_j^{\text{reac}} := n_j^{\text{reac}} \frac{f_j^{\text{reac}}}{f_j} \quad j \in \{3, 6, 7, 8\}, \tag{5.1} \]

are plotted (with stoichiometric factors \( n_j^{\text{reac}} \) equal to 1, 3, 2 and 3 for \( j \) equal to 3, 6, 7 and 8, respectively).

It can be observed that the strongest carbonation reaction is that of Ca(OH)$_2$, followed by that of CSH. The reaction rates of C$_2$S and C$_3$S are essentially smaller. The reaction zone associated with Ca(OH)$_2$-depletion is the most narrow and most furthest advanced into the concrete sample at any given time. On the other hand, the reaction zone of CSH is the widest and also lagging behind the most. The reaction zone of C$_2$S is somewhere in between while that of C$_3$S is quite similar to that of Ca(OH)$_2$ (apart from the magnitude of the reaction rate). This last fact can be explained by the comparably small amount of C$_3$S available to carbonation.

Since we expect a strong competition for CO$_2$ by the three carbonation reactions, we are interested in how much CO$_2$ is consumed by each carbonation reaction. Moreover, since simpler models usually neglect CSH, C$_2$S, and C$_3$S, we are interested in how great their impact is on the carbonation of Ca(OH)$_2$. Therefore, we also consider two scenarios and compare them to the full scenario (cf. figs. 2 and 3): in the first, we neglect CSH, C$_2$S and C$_3$S; in the second, we neglect C$_2$S and C$_3$S.

Figure 4: Dimensionless production terms of the different carbonation reactions.
5.1.1 Consumption of $\text{CO}_2$ by each carbonation reaction

Since CSH, $\text{C}_2\text{S}$ and $\text{C}_3\text{S}$ cannot diffuse, we can calculate the amount of $\text{CO}_2$ which has been consumed due to each carbonation after any given time. We are particularly interested in the amount of $\text{CO}_2$ consumed by carbonation of each reactant. We denote this quantity with $P_j(x,t)$, $j \in \{3, 6, 7, 8\}$. Note that this quantity satisfies the ordinary differential equation

$$\frac{\partial}{\partial t} (\beta_2 P_j(x,t)) = n_j^\text{reac} f_j^\text{reac}, \quad x \in \Omega, \ t > 0, \ P_j(x,0) = 0. \quad (5.2)$$

The spatial profiles of $P_j(x,t)$ are plotted in figure 5.

It can be observed that all four species compete for $\text{CO}_2$. However, the amount of $\text{CO}_2$ consumed by the carbonation of $\text{Ca(OH)}_2$ is much greater than that of all other species. Namely, the amount consumed by carbonation of $\text{Ca(OH)}_2$ is four times as great as that of CSH, 16 times as great as that of $\text{C}_2\text{S}$ and roughly 36 times as great as that of $\text{C}_3\text{S}$. It is also worth noting that the reaction zone associated with $\text{Ca(OH)}_2$ is always ahead of the other reaction zones. This agrees with the experimental observations in [Cha97].

5.1.2 Alternate neglect of CSH, $\text{C}_2\text{S}$, and $\text{C}_3\text{S}$

In this section we investigate the influence of each carbonation process on the penetration depth. Recall that the carbonation depth is defined via an a priori fixed isoline of the concentration profile.
5.1 Influence of the competition of parallel carbonation reactions

of Ca(OH)$_2$ (cf. sec. 4.3). Therefore, we consider the amount of produced CaCO$_3$ as well as the carbonation depth in two cases:

- neglect of CSH, C$_2$S, and C$_3$S,
- neglect of C$_2$S and C$_3$S.

These two scenarios are illustrated in figure 6 (also compare with the full scenario, figs. 2e and 3c). Note that since C$_2$S and C$_3$S do not consume much CO$_2$ compared to CSH and Ca(OH)$_2$, it is reasonable to consider them together.

It can be observed that the carbonation of CSH affects the simulated carbonation depth as it consumes CO$_2$ which would otherwise be available for carbonation of Ca(OH)$_2$. On the other hand, the influence of C$_2$S and C$_3$S is almost negligible.

Figure 6: Profile of CaCO$_3$ and carbonation depth if CSH, C$_2$S and C$_3$S are neglected (a,b); and if C$_2$S and C$_3$S are neglected (c,d).
5.2 Variation of carbonation- and hydration-rate constants

As the carbonation- and hydration-rate constants are somewhat uncertain [cf. PVF89, SD96, e.g.] we want to illustrate the effects due to their variation. The effect of the variation of the Thiele modulus on the carbonation penetration depth was extensively investigated in [MPMB05]. Therefore, we concentrate on the reaction and hydration rates of CSH, C$_2$S, and C$_3$S. Note that the influence of the additional species is already quite small which is why we only investigate the effect of faster rates.

5.2.1 Variation of carbonation-rate constants

In figure 7, the dimensionless production terms $n^\text{reac}_j$ (also cf. (5.1)), the concentration profile of CaCO$_3$ as well as the carbonation depth are plotted for the standard setting, except that the reaction-rate constants of the carbonation of CSH, C$_2$S, and C$_3$S have been chosen ten times greater.

It can be observed that the carbonation depth is smaller than in the in the standard setting (figs. 2 and 3). This is due to the fact that more CO$_2$ is consumed by the carbonation of CSH, C$_2$S, and C$_3$S. Therefore, it is not available for carbonation of Ca(OH)$_2$. Moreover, the concentration profiles of CSH, C$_2$S and C$_3$S appear steeper and it can be seen that, in this setting, all reaction zones seem to coincide.

If the carbonation-rate constants of CSH, C$_2$S, and C$_3$S are chosen 100 times greater than in the standard setting it can be observed that the reaction zones of these additional phases overtake the one associated with Ca(OH)$_2$ (cf. fig. 8). It is not clear whether these observations reflect the reality. Nevertheless, it is due to the fact that there is not as much CSH, C$_2$S and C$_3$S available for carbonation as there is Ca(OH)$_2$. Therefore, these phases are used up more quickly and the reaction zone associated with them advances more quickly. As in the previous setting, the carbonation depth is only slightly affected, however. Regarding this last observation, recall again that the carbonation depth is defined via an isoline of the concentration profile of Ca(OH)$_2$ so that it does not recover this faster advancement of the reaction zones of the additional phases. It can also be observed that the maximum of the production term related to carbonation of Ca(OH)$_2$ increases (beside the maxima of the other production terms). This is due to the fact that, in this setting, the concentration of CO$_2$ at the position of the reaction zone is significantly higher (the profile of the CO$_2$-concentration changes only slightly compared to the standard setting).

If the reaction-rate constants of CSH, C$_2$S, and C$_3$S are chosen significantly smaller than in the standard setting (multiplying them by a factor of 1/10 or 1/100, e.g.) the influence of the additional phases just decreases steadily.

5.2.2 Variation of hydration-rate constants

In figure 9, selected concentration profiles as well as the carbonation depth are shown for the standard setting but with hydration-rate constants which have been chosen ten times faster than in figures 2 and 3.

Recall that the overall hydration process is assumed in its final phase. Therefore, it is almost complete at the beginning of our simulations. However, some small competition effects can be observed. The concentration profiles of C$_2$S and C$_3$S show a considerable deviation. The influence on the profiles of interest (the carbonation depth, e.g.) is rather small. Note that even a multiplication of the hydration-rate constants by a factor of 1000 does not have a significant effect on the carbonation depth.
5.2 Variation of carbonation- and hydration-rate constants

Figure 7: Production rates, profile of CaCO$_3$ as well as carbonation depth obtained with carbonation-reaction rates ten times faster than in the standard setting.
Figure 8: Production rates, profile of CaCO$_3$ as well as carbonation depth obtained with carbonation-reaction rates one hundred times faster than in the standard setting.
5.2 Variation of carbonation- and hydration-rate constants

![Graphs of Ca(OH)$_2$, CaCO$_3$, CSH, C$_2$S, C$_3$S, and carbonation depth profiles](image)

Figure 9: Profiles of selected concentrations as well as carbonation depth obtained with *hydration-reaction rates* ten times faster than in the standard setting.
6 Summary and discussion

The carbonation of concrete was investigated taking into account several parallel carbonation and hydration reactions. A closed system of reaction-diffusion equations was used to model the mass balances of all active species under consideration. To allow easy comparisons of the importance of the involved carbonation and hydration reactions, a nondimensionalisation was carried out for this model. Using results from numerical simulations, the influence of each reaction on the penetration depth was investigated in detail. Experimental data from [PVF89] was recovered.

Compared to simplified models which neglect CSH, C$_2$S and C$_3$S, it can be observed that the two silicate phases only have a small influence on the total outcome in the late stage of hydration considered here. This was found for an accelerated carbonation scenario. Further research is required with respect to natural carbonation where the effect of CSH and unhydrated constituents on the penetration depth is not a priori clear.

In general, all species compete for carbon dioxide. This slows down the carbonation of calcium hydroxide since less carbon dioxide is available to carbonation. This effect, reducing the propagation speed of the carbonation layer, was of noticeable extent only for CSH. Under natural conditions, experiments suggest that the influence of the carbonation of CSH might become even smaller [cf. Cha97, e.g.] because of the worse accessibility of carbon dioxide and the possibly lesser amount of CSH available in the concrete (due to uncomplete hydration). On the other hand, for concretes with high CSH content (such as those containing blast furnace slag, fly ash, silica fume, etc.), this influence might be stronger again. Moreover, depending on the cement chemistry, several other alkaline species such as KOH, NaOH, Mg(OH)$_2$, and aluminate phases may also carbonate and, altogether, might have a noticeable effect on the simulated carbonation depth.

For a large range of parameters, different reaction layers are formed associated with the several carbonation reactions. On the other hand, the hydration reactions occur fairly uniformly as long as unhydrated constituents are available. The fact that the reaction layer associated with CSH lags behind that of calcium hydroxide, which was observed experimentally by [Cha97], is also recovered in our simulations. We are not aware of experimental information about the formation of reaction zones associated with the carbonation of C$_2$S or C$_3$S.

It is well-known that the effect of moisture on the carbonation process is of considerable importance. In the model considered here, a rather simple approach was chosen. The mobile moisture influencing the rapidness of the carbonation of calcium hydroxide was modelled by a reaction-diffusion equation weakly coupled with the production term associated with the carbonation of calcium hydroxide. The effect by immobile moisture stored in the porous concrete matrix was neglected. Further research is required in order to take into account possible effects of the mobile moisture (on the diffusivities, e.g.), the immobile moisture (on the reaction rates, e.g.) as well as couplings between them. A possible better approach could be the use of double-porosity-type models. More research is required to investigate this issue.

Acknowledgments

This work has been partially supported by the Deutsche Forschungsgemeinschaft (DFG) with a grant through the special priority program SPP1122 Prediction of the Course of Physicochemical Damage Processes Involving Mineral Materials. We would also like to thank J. Grunewald, J. Kropp, K. Sisomphon and L. Franke for fruitful discussions.
Appendix: Dimensional parameters

We list the *standard* sets of dimensional parameters of the reference setting which we use for the numerical simulations. It refers to ordinary portland cement. There are some parameters which are definite, for instance the molecular weights of the involved species. Others are generally unknown, for instance the carbonation-reaction constants, or they strongly depend on the setting considered. Therefore, we first list all parameters in table 1. We give exact values where the parameters are definite and ranges where they are uncertain. The particular values used for the numerical simulations are given in table 2, representing an accelerated carbonation setting. Note that due to some dependencies, changing a certain parameter may imply several other changes (e.g. changing $R_{w/c}$ implies a change in $\phi$ which, in turn, implies a different $u_1^{exc}$ etc.).

Most of the values of the dimensional parameters can be found in the standard literature, e.g. in [Lid02]. For some specifics, we refer to [BKM03b] and [Ste00], as well as [Arf98] for values of $D_{\text{H}_2\text{O}}$. 

<table>
<thead>
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<th>Parameter</th>
<th>Value</th>
<th>Unit</th>
<th>Description</th>
</tr>
</thead>
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<tr>
<td>$R_{wc}$</td>
<td>0.4 – 0.6</td>
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<td>water-to-cement ratio</td>
</tr>
<tr>
<td>φ</td>
<td>0.1 – 0.6</td>
<td>–</td>
<td>concrete porosity</td>
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<td>φ'</td>
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<td>–</td>
<td>volume fraction of air-filled pores</td>
</tr>
<tr>
<td>φw</td>
<td>0 – 1</td>
<td>–</td>
<td>volume fraction of water-filled pores</td>
</tr>
<tr>
<td>L</td>
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<td>cm</td>
<td>width under consideration</td>
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<table>
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<th>Value</th>
<th>Unit</th>
<th>Description</th>
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<td>cm²/day</td>
<td>diffusion constant for CO₂(g)</td>
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<tr>
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<td>cm²/day</td>
<td>diffusion constant for moisture</td>
</tr>
<tr>
<td>$C_{Henry}$</td>
<td>0.7 – 0.9</td>
<td>–</td>
<td>dimensionless Henry constant</td>
</tr>
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<td>$C_{ex}$</td>
<td>$10^0$ – $10^6$</td>
<td>1/day</td>
<td>mass transfer coefficient for absorption of CO₂(g)</td>
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<tr>
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</tr>
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<td>cm³/day</td>
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<td>exponent in hydration reaction rate of C₂S</td>
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<tr>
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<td>g/cm³</td>
<td>initial concentration of CO₂(aq)</td>
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<td>g/cm³</td>
<td>ambient concentration of CO₂(g)</td>
</tr>
<tr>
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<td>g/cm³</td>
<td>ambient concentration of moisture</td>
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<td>a</td>
<td>−0.11</td>
<td>–</td>
<td>fitting parameter in sorption isotherm</td>
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<tr>
<td>b</td>
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<td>cm³/g</td>
<td>fitting parameter in sorption isotherm</td>
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Table 1: Values and ranges of dimensional parameters.
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<td>( \phi )</td>
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<td>( \phi_w^w )</td>
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<td></td>
</tr>
<tr>
<td>( D_{CO_2} )</td>
<td>( 1 \times 10^{-4} ) cm(^2)/day</td>
<td></td>
</tr>
<tr>
<td>( D_{Ca(OH)_2} )</td>
<td>( 1 \times 10^{-7} ) cm(^2)/day</td>
<td></td>
</tr>
<tr>
<td>( D_{H_2O} )</td>
<td>( 1 \times 10^{-2} ) cm(^2)/day</td>
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<tr>
<td>( C_{Henry} )</td>
<td>0.85 g/cm(^3)/day</td>
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<td>( 1 \times 10^2 ) cm/day</td>
<td></td>
</tr>
<tr>
<td>( C_{Rob} )</td>
<td>( 1 \times 10^5 ) cm/day</td>
<td></td>
</tr>
<tr>
<td>( C_{Rob} )</td>
<td>( 1 \times 10^7 ) cm/day</td>
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</tr>
<tr>
<td>( C_{reac} )</td>
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<td>0.25 g/cm(^3)/day</td>
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<td>( C_{SH} )</td>
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<td>( C_{C_3S} )</td>
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<td>( p_{C_3S} )</td>
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<td>0.0074 g/cm(^3)</td>
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<td>( \phi_{w,C_3S}^w )</td>
<td>0.0015 g/cm(^3)</td>
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Table 2: Specification of the values of table 1 used in the simulations.
References


REFERENCES


