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CO₂-cycle in cement and concrete
Part 7: Models for CO₂-absorption.

A new model for CO₂-absorption of concrete structures

Lars-Olof Nilsson

Funded by the Swedish Consortium for Financing Basic Research in the Concrete Field

Report TVBM-3158 Lund 2011
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1 Background

In a previous project, funded by the Nordic Innovation Centre, CO$_2$-uptake during the concrete life-cycle was studied, with project participants from Denmark, Iceland, Norway and Sweden. The objective was to provide documentation of concrete carbonation during the service life and recycling and re-use. The project resulted in a number of reports.

Preliminary results indicated significant effects: “In countries with the most favourable recycling practice it is realistic to assume that 86 % of the concrete is carbonated after 100 years, taking up approximately 57 % of the CO$_2$ emitted during the calcining process. Examples of impact of CO$_2$ uptake in life cycle screenings show that 70-80 % of the potential CO$_2$ uptake has been absorbed within 100 years lifetime after demolition and crushing”, Glavind (2006).

The project had a limited budget and the analysis was based on a number of assumptions and uncertainties. The need for further work was identified.

One weakness in the previous project was the simple square-root of time models used for carbonation and the lack of models for carbonation of crushed concrete. One part of the present project, part 7, concerns the development of new models for CO$_2$-uptake.

In a previous report, Nilsson & Fridh (2009), we summarized the present state of knowledge on carbonation models and pointed out how those models could be used to predict the absorption of CO$_2$. We also identified some of the knowledge that is missing. Now, special attention has to be given to the moisture dependency of carbonation, to be able to treat absorption of CO$_2$ in indoor structures.

After further laboratory and field studies we have now used the previous and new findings to create a model for CO$_2$-absorption of concrete structures. Some knowledge is still missing and some results are still not fully understood. In those cases we have done our best to base the model on reasonable assumptions, from what could be expected from some new literature that has been studied.

The project is funded by the Swedish Consortium for Financing Basic Research in the Concrete Field. The consortium members are: Cementa, Färdig Betong, Abetong, Swerock, Betongindustri and Strängbetong.
# Assumptions and definitions

## 2.1 Assumptions/simplifications

In a previous report current models for carbonation were described. For the choice of model(s) for predicting CO$_2$-uptake by carbonation, these assumptions/simplifications seemed to be reasonable

1. Carbonation is modelled as a sharp carbonation front moving inwards, cf. figure 1.
2. The movement of the carbonation front is due to the combined effect of CO$_2$ diffusing through the already carbonated layer and CO$_2$ being bound by the carbonation reaction at the depth of the front.
3. The amount of CO$_2$ bound per volume of concrete must be quantified in the model.
4. The amount of CO$_2$ bound per weight of binder may not be a constant for a particular binder; possible effects of degree of hydration and moisture level should be part of a model and separately quantified.
5. The concrete is modelled as being homogenous, i.e. the higher binder content close to the exposed surface is neglected. This assumption is on the "safe" side.
6. All CaO is assumed to be fixed in position, i.e. no movement of Ca$^{2+}$ and OH$^{-}$ ions is considered.
7. The CO$_2$-binding capacity of carbonated concrete is set to be equal to 0, i.e. the diffusion of CO$_2$ could be modelled as a steady-state transport process.
8. The diffusion coefficient for CO$_2$ for a particular concrete should be modelled as a function of degree of hydration and moisture level.
9. The liberation of water from the carbonation reaction is neglected, i.e. does not increase the moisture content and humidity.
10. The effect of a surface treatment, paint, wall paper etc., must be considered, at least as an additional resistance to diffusion of carbon dioxide.

These assumptions/simplifications should be agreed upon, within the project. The assumptions are quantified in the next sections.

The consequences of these assumptions are several:

a) The CO$_2$-uptake is quantified by models that include assumptions that are well known and frequently used in models for carbonation.

b) The historic moisture profiles of the concrete structure must be estimated, not only the equilibrium conditions; the effect of outdoor humidity variations and rain periods, and the early drying of excess moisture, should be considered to some extent. This is, however, neglected due to lack of information.

c) The profiles of degree of hydration, originating from the time required for the early drying of the surface regions of the concrete structure, should be considered to some extent. This is, however, neglected due to lack of information.

d) The significance of these effects should be smaller and smaller with time, i.e. with larger depths of carbonation.
2.2 Definition of material properties

The model includes a few important and decisive material properties. They are defined here in detail.

2.2.1 Diffusion coefficient for CO₂ in concrete

The diffusion coefficient for CO₂-diffusion in concrete is defined by the flux of CO₂-gas through an infinitesimal layer of concrete

\[
J_{CO_2} = -D_{CO_2} \frac{\partial c}{\partial x} \left[ \frac{kgCO_2}{m_e^2 s} = \frac{m_{air}^3}{m_e s} \frac{kgCO_2}{m_e^2} = \frac{m^2}{s} \frac{kgCO_2}{m_{air} m_e} \right]
\]  

(1)

where \(m_e\) refers to the depth and cross-section of concrete. The concentration \(c\) of CO₂ is given as an amount in a unit volume of air. The amount could of course be given in moles as well.

The diffusion coefficient \(D_{CO_2}\) has, as usual, the dimension \(m^2/s\), but when using literature data it is important to clarify how the concentration of CO₂ is defined.

2.2.2 Resistance to CO₂-diffusion of a surface cover

The resistance \(R_{CO_2}\) to CO₂-diffusion of a thin layer, i.e. a surface cover, is defined by the flux of CO₂-gas through the layer

\[
J_{CO_2} = \frac{\Delta c}{R_{CO_2}} \left[ \frac{kgCO_2}{m_e^2 s} = \frac{kgCO_2}{m_{air}^3 m_e s} = \frac{m^2}{s} \frac{kgCO_2}{m_{air} m_e} \right] \left[ \frac{m}{s} \right]
\]  

(2)

The resistance \(R_{CO_2}\) has the dimension \(s/m\).
2.2.3 Amount of CO\(_2\) bound in a unit volume of concrete

A model for carbonation, and for CO\(_2\)-absorption, should be a solution to a mass-balance equation for CO\(_2\). A general mass-balance equation is

$$\frac{\partial c_c}{\partial t} = - \frac{\partial J_{CO_2}}{\partial x} - \frac{\partial c_b}{\partial t} \left[ \frac{kgCO_2}{m^3 \cdot s} = \frac{kgCO_2}{m^2 \cdot s \cdot m_x} \right]$$

where \(c_c\) is the concentration of CO\(_2\) in a unit volume of concrete and \(c_b\) is the amount of bound CO\(_2\), also in a unit volume of concrete. In this mass-balance equation the amount of bound CO\(_2\) \(c_b\) will later be called \(a\), the amount of CO\(_2\) bound in a unit volume of concrete.
3 Model for the depth of carbonation

With these assumptions, and definitions, a carbonation model may look like the one in figure 2.

The flux of diffusing CO\textsubscript{2} is based on Equation (1)

\[ J_{CO_2} = -D_{CO_2} (RH, \alpha) \frac{dc}{dx} = \frac{c}{R_{CO_2}} \]  

where the diffusion coefficient depends on humidity and degree of hydration. The concentration difference over the carbonated layer with thickness \(X_{CO_3}\) \(\Delta c = c-0 = c\). The resistance to diffusion of CO\textsubscript{2} of the carbonated layer is \(R_{CO_2}\), which is given by

\[ R_{CO_2} = \int_{x=0}^{x=X_{CO_3}} \frac{dx}{D_{CO_2} (RH (x), \alpha (x))} \]  

In the model in figure 2 the mass balance equation of CO\textsubscript{2} at the depth of carbonation \(X_{CO_3}\) will be

\[ J_{CO_2} \cdot dt = a(X_{CO_3}) \cdot dX_{CO_3} \text{ [kg CO}_2\text{/m}^2\text{]} \]  

where \(dt\) is a short time step \([s]\) during which the carbonation front advances \(dX_{CO_3}\) \([m]\). This can also be written

\[ \frac{dX_{CO_3}}{dt} = \frac{J_{CO_2}}{a(X_{CO_3})} = \frac{c}{R_{CO_2} \cdot a(X_{CO_3})} \]  

After integration, the depth of carbonation is

\[ X_{CO_3} = \int_{x=0}^{x=X_{CO_3}} dX_{CO_3} = \int_{t=0}^{t=t} \int_{x=0}^{x=X_{CO_3}} \frac{c}{a(X_{CO_3} (t)) \cdot \int_{x=0}^{x=X_{CO_3}} \frac{dx}{D_{CO_2} (RH (x), \alpha (x))}} \]
The “problem” with this model is that the $X_{CO_3}$ appears on both sides of the equality sign! Consequently, the equation must be solved numerically, by dividing the carbonated layer into small slices with a thickness of $dx$ and by taking small time-steps $dt$. During each time-step, the moisture distribution $RH(x)$ must be known.

If further assumptions are made, the model will of course become much more simple. If the humidity $RH$, the degree of hydration $\alpha$ and the amount $a$ of $CO_2$ bound in a unit volume of concrete are assumed to be constant throughout the carbonated layer, the depth of carbonation will be

$$X_{co_3} = \sqrt{\frac{2D_{co_2} (RH) \cdot c}{a_{co_3} (RH)}} \cdot \sqrt{t} \left( \frac{t}{t_n} \right)^n$$

(9)

where the exponent $n$ depends on the humidity conditions and can be quantified from exposure data or from a more elaborated model. That was done by CEB (2000). A value of $n = 0.17$ was found for surfaces exposed to rain and $n \approx 0$ for surfaces protected from rain, Nilsson & Rodhe (1997a).

The parameters in this model are, theoretically, possible to quantify from laboratory and field measurements but very good data is required. This is further discussed in sections 4, 5 and 7.

3.1 Effect of a surface coating

A coating at the concrete surface will add a resistance to diffusion of carbon dioxide but will not change the carbonation reaction.

Performing the same kind of derivation as in equations (4)-(9), but adding a resistance $R_{CO_2}$ to the resistance against diffusion of $CO_2$ of the carbonated layer with a thickness of $X_{CO_3}$, will give a slightly different equation for the depth of carbonation

$$X_{co_3} = \sqrt{\left( D_{co_2} (RH) \cdot R_{co_3} \right)^2 + \frac{2D_{co_2} (RH) \cdot c \cdot t}{a_{co_3} (RH)}} - D_{co_2} (RH) \cdot R_{co_3}$$

(10)

if we set $n=0$, which is OK for not too wet structures or structures that are not exposed to rain, CEB (2000).

The consequence of a surface coating is that the time-dependency of the carbonation process no longer follows the square-root law. An example is shown in figure 3.

An example of the effect of the resistance of the surface coating on 50 years of carbonation is shown in figure 4. Surface resistances that are changing the rate of carbonation between 10 and 90 % have a magnitude of somewhere between $R=100000$ s/m and $R=2000000$ s/m.
Fig. 3 An example of the carbonation process for a concrete structure with a surface coating. Indoor concrete with w/c=0.8 and a surface resistance R=1 \cdot 10^6 \text{s/m}.

Fig. 4 An example of the effect of the resistance of a surface coating on carbonation after 50 years. Indoor concrete with w/c=0.6 and different surface resistances.
4 The amount of CO₂ bound in concrete

The aCO₂-value has already been shown to include only one uncertain parameter, the degree of carbonation that has to be measured for a set of typical binders and humidity conditions.

The amount of CO₂ bound in a unit volume of concrete is

\[
a_{\text{CO}_2}(x) = C \cdot \frac{\text{CaO}}{C} \cdot \frac{(\text{CaO})_{\text{CO}_3}}{\text{CaO}}(x) \cdot \frac{M_{\text{CO}_2}}{M_{\text{CaO}}} \text{ [kg CO}_2/\text{m}^3]}
\]

or

\[
a_{\text{CO}_2}(x) = C \cdot k_{a,C} \cdot k_{a,a} \cdot k_{a,RH} \cdot \frac{M_{\text{CO}_2}}{M_{\text{CaO}}} \text{ [kg CO}_2/\text{m}^3]}
\]

where \( C \) is the cement content, \( \text{CaO}/\text{C}_{\text{PC}} (=k_{a,C}) \) is the amount of CaO per weight of Portland cement, \((\text{CaO})_{\text{CO}_3}/\text{CaO} (=k_{a,a} \cdot k_{a,RH})\) is the degree of carbonation and \( M \) is the molar weight. The degree of carbonation consist of two factors \( k_{a,a} \cdot k_{a,RH} \) where the first one depends on the degree of hydration of the binder and the last one describes the moisture dependency of the carbonation reaction. The amount \( a(x) \) is marked as being dependent on the depth \( x \), since the degree of carbonation may be different at different depths.

The parameter \( \text{CaO}/\text{C}_{\text{PC}} (=k_{a,C}) \) follows from the binder composition, where the amount of CaO is only the portion that is possible to carbonate in a concrete. In a Portland cement this factor usually is in the order of 0.60-0.65. For a blended cement this factor must be multiplied with a factor \( \text{C}_{\text{PC}}/C \) that is the portion of Portland cement clinker in the cement.

The degree of carbonation \((\text{CaO})_{\text{CO}_3}/\text{CaO} (=k_{a,a} \cdot k_{a,RH})\) is the extent to which the available CaO actually is carbonated. The first factor is usually assumed to represent all of the CaO in the hydrated part of the cement. That factor is simply assumed equal to the degree of hydration

\[
k_{a,a} = \alpha
\]

The moisture dependency of the degree of carbonation is not well known, however. From a literature search some examples of actual measurements have been found. In the project some measurements have been done in indoor structures that could be used to estimate this parameter.

Goodbrake et al (1979) measured the degree of carbonation on paste samples of C₂S and C₃S after carbonation in various humidity, from 10 % RH. Their results are shown in figure 5. They concluded that the relationship between the degree of carbonation and RH is linear from 0 at 0 % RH up to 0.68-0.73 at 50 % RH and then constant in the interval 50-100 % RH.

They did not report the degree of hydration. It may very well be around 0.7, i.e. the moisture dependency factor would be 1.0 in the RH-interval 50-100 %.
Measurements by van Balen (2005) on lime showed a degree of carbonation of 0.80-0.85, with a mean value of 0.83. A few measurements of RH were made during the experiments, varying between 85 and 90 % RH. RH was said to increase because of the carbonation process.

El-Turki et al (2007) analyzed samples of hydraulic lime carbonated in 100 % CO$_2$ in two different climates. In 97 % RH they saw new Ca(OH)$_2$ and concluded that the carbonation was “complete”. Hydraulic lime should, however, contain some silica and, consequently, some CSH that also could be carbonated. In 65 % RH a small amount of Ca(OH)$_2$ was found; the carbonation was obviously not complete at that RH.

Fridh & Lagerblad (2010) measured degree of carbonation and RH in a few indoor structures. At an RH of 48 % she found the degree of carbonation to be around 0.42-0.48 in samples from an indoor beam and the bottom of a floor slab. The degree of hydration is not known for these samples; an estimation gives $\alpha=0.7-0.8$. With these numbers, the moisture dependency factor is $k_{a,RH} = 0.59-0.68$.

Data from measurements that Samuelsson (1977) presented could be used to evaluate the moisture dependency. He made carbonation experiments at young ages in climate rooms with a series of different relative humidity. After the specimens had carbonated for a couple of days or weeks they were exposed to water and the amount of CaO leached out into that water was measured. The amount of CaO that leached out is a measure of to what extent CaO was available, e.g. to what extent the CaO was not carbonated. Samuelsson’s results are summarized in figure 6.

Fig. 5 The moisture dependency of the carbonation reaction, Goodbrake et al (1979)
The data on the moisture dependency factor is summarized in figure 7.

More precise measurements should be done of the moisture dependency factor for carbonation. The data in figure 5 shows significant scatter. The full curve in figure 7 is the relationship used in the present model, but the uncertainty is of course significant.

In the few measurements made of carbonation profiles, the degree of carbonation beyond the carbonation depth was never found to be zero! Möller (1994) found a degree of carbonation of around 0.1, cf. figure 1, and Fridh & Lagerblad (2010) found 0.15. This is yet to be explained. Obviously, some carbonate exists in the un-carbonated part of the concrete. It could originate from the binder or the aggregate. If the source is the binder, the “available” CaO for carbonation is somewhat overestimated. If, on the other hand, the source is the aggregate, the measured degree of carbonation is somewhat underestimated. Further research is needed to clarify this.
5 The diffusion coefficient for CO₂

The diffusion coefficient could be quantified together with the a-value by equation (7) in cases where the exponent \( n = 0 \). Alternatively, the diffusion coefficient could be estimated from an equation from literature. Numerous such equations do exist but the data behind them is rare. Here, the one derived for old German cements, Wierig (1964), has been used

\[
D_{\text{CO}_2} = B \cdot C \cdot \varepsilon^{4.5} \cdot (1 - RH \cdot (x)^{10})^5
\]

(14)

where \( C \) is the cement content, \( \varepsilon \) is the capillary porosity, \( B \) is a constant and the parenthesis gives the humidity dependency. The factor for the humidity dependency is shown in figure 8.

\[\text{Fig. 8 The moisture dependency of the diffusion coefficient for carbon dioxide according to equation (14)}\]

The capillary porosity is calculated as the capillary porosity of cement paste according to Power’s model

\[
\varepsilon = \frac{w/c - 0.39\alpha}{0.32 + w/c}
\]

(15)

where \( w/c \) is the water-cement ratio and \( \alpha \) is the degree of hydration. To obtain the correct capillary porosity the \( w/c \) must be based on the actual Portland cement content in the concrete, which could be different from the cement content, if the cement contains non-cementitious constituents such as limestone filler.

Now all the parameters, except one, in the carbonation model in equations (9) and (10) are quantified. The remaining parameter B is determined by calibration in section 7.
Model for CO$_2$-absorption

A model for absorption of carbon dioxide is now simple to derive, from the models for depth of carbonation in equations (9) or (10) and the amount $a_{CO_2}$ CO$_2$ bound in a unit volume of concrete. The CO$_2$-uptake of a unit area of concrete will be

$$m_{CO_2} = X_{CO_3} \cdot \int_{0}^{x_{CO_3}} a_{CO_2}(x) \cdot dx \quad [\text{kg CO}_2/\text{m}^2]$$

(16)

or, if $a$ is assumed to be constant throughout the carbonated layer, simply

$$m_{CO_2} = X_{CO_3} \cdot a_{CO_2} \quad [\text{kg CO}_2/\text{m}^2]$$

(17)

This model has been set up in an Excel sheet, including all the parameters mentioned above. The front sheet of this model is shown below.
7 Calibration/ verification

The parameter values in the model are dealt with in different ways. Most of them follow from the concrete composition, the environmental conditions and the assumptions given above. There is more or less only one parameter left to quantify and that is the parameter $B$ in the equation for the diffusion coefficient for $CO_2$.

It has to be quantified by calibrating the model against laboratory and/or field data. Such a calibration requires the other parameters to be known, especially the concrete composition, the degree of hydration and the climatic conditions. Good data where these parameters are known are rare. Carbonation experiments are usually not that well documented that all these three parameters are known. Instead, one or two of them has to be estimated. That will of course mean that the calibration is uncertain. For the time being we have to live with that uncertainty; future research may remove some of the uncertainties.

7.1 Calibration against the Swedish concrete cover standard

The Swedish concrete cover standard contains requirements for concrete covers and concrete w/c for different service-lives in the exposure classes of EN206. In some of the exposure classes XC, where carbonation initiated reinforcement corrosion is the dominant deterioration mechanism, the required cover must be regarded as the expected depth of carbonation in Swedish conditions and for Swedish Portland cement concretes. The carbonation depths are, however, the maximum expected depth of carbonation. That means that using those depths will give a certain overestimation of the average carbonation depth and, consequently, a certain overestimation of the absorbed amount of $CO_2$, which should be based on the average depth of carbonation. Using the larger covers, i.e. the longest service-lives, should decrease that over-estimation. Hergenröder (1992) found the difference between the maximum depth and the average depth of carbonation to be less than 5 mm at a carbonation depth of 20 mm.

To be able to calibrate the carbonation depth model, as many of the parameters in the model must be known. Therefore, the cover thicknesses in the two exposure classes that are of interest for carbonation initiated reinforcement corrosion, XC3 and XC4, have been chosen. In these two exposure classes the required covers are equal. To be relevant the most severe climatic conditions must be selected. As low RH as possible will give a maximum depth of carbonation. Only RH above the critical limit for corrosion, however, is relevant. This limit is some 85 % RH according to Tuutti (1982). Consequently, these exposure classes should have an RH of 85 % RH.

In the Swedish concrete cover standard SS137010 the required covers in these two exposure classes are given in table I.

Table I. Required covers in exposure classes XC3 och XC4 according to SS137010

<table>
<thead>
<tr>
<th>Exposure class</th>
<th>Max ((w/c)_{eq})</th>
<th>Required concrete cover (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>in L 100</td>
<td>in L 50</td>
</tr>
<tr>
<td>XC3, XC4</td>
<td>0.55</td>
<td>25</td>
</tr>
<tr>
<td></td>
<td>0.50</td>
<td>20</td>
</tr>
</tbody>
</table>

The cover depths are given as the nearest larger multiple of 5 mm which means that the shorter service-lives L20 and L50 should give a larger over-estimation of the required limit for the depth of carbonation than the longest service-life L100.
The concept "(w/c)_{eq}" must be seen as the water-binder ratio in a pure Portland cement concrete. When using other binders the \( (w/c)_{eq} \) will be larger which will affect the capillary porosity and the diffusion coefficient for CO\(_2\). 

For the calibration, the other parameters must be estimated. A very important parameter is the degree of hydration, which appears both in the capillary porosity in the equation (9) for the diffusion coefficient for CO\(_2\), with an exponent of 4.5, and in the equation (11) for the degree of carbonation. The degree of hydration for different concretes and different curing can be estimated from the data by Norling-Mjörnell (1997). The curing could of course vary but since the most severe climatic conditions for carbonation-induced corrosion should be a climate with RH = 85 % and protection against rain, long-term curing in that climate should be relevant. Data by Norling-Mjörnell (1997) gives an \( \alpha = 0.79 \) and 0.75 for the two concretes in table I.

The predictions for these two concretes are compared with the depths of carbonation in table I. A B-value of \( B=5\cdot10^{-8} \) was used.

![Graph showing predicted depths of carbonation](image)

*Fig. 9 The predicted depths of carbonation from the model, with a B-value selected, compared with the data in table I for two concretes in XC3 and XC4.*

The comparison in figure 9 shows the predicted average depth of carbonation and the required covers in the exposure classes. The required covers should be somewhat larger, since they should consider the maximum depths of carbonation and not the average values. The difference could be a little too small. Maybe the B-value should be chosen somewhat smaller.

### 7.2 Calibration against Cementa’s field exposure data at Smöjen

Cementa has casted concrete specimens and exposed them for a long time at an exposure site in Smöjen on the island of Gotland. Recently, Ljungkrantz (2008) presented some of the results from these exposure tests. An attempt is made to calibrate or at least compare the prediction model with these data. Only the data from specimens that were sheltered from rain are used.

The data from Ljungkrantz (2008) is summarized in table II.
Table II Carbonation depths (mm) for samples stored outdoors at the island of Gotland, sheltered from rain, Ljungkrantz (2008).

<table>
<thead>
<tr>
<th>Cement, year, w/c</th>
<th>Years exposed</th>
<th>Assumed α</th>
<th>Phenolphthalein method</th>
<th>Thin section method</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>(Upper side)</td>
<td>(Upper side)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Bottom side</td>
<td>Bottom side</td>
</tr>
<tr>
<td>ByggC, 2000</td>
<td>8</td>
<td>0.80</td>
<td>(2.2)</td>
<td>5.3</td>
</tr>
<tr>
<td>W/C = 0.60</td>
<td></td>
<td></td>
<td></td>
<td>To be tested</td>
</tr>
<tr>
<td>StdP, 1982</td>
<td>26</td>
<td>0.80</td>
<td>(1.3)</td>
<td>12.3</td>
</tr>
<tr>
<td>W/C = 0.60</td>
<td></td>
<td></td>
<td></td>
<td>(10)</td>
</tr>
<tr>
<td>Std M, 1982</td>
<td>26</td>
<td>0.80</td>
<td>(7.5)</td>
<td>12.7</td>
</tr>
<tr>
<td>W/C = 0.60</td>
<td></td>
<td></td>
<td></td>
<td>(10)</td>
</tr>
<tr>
<td>StdP, 1985</td>
<td>23</td>
<td>0.85</td>
<td>(11.6)</td>
<td>23.2</td>
</tr>
<tr>
<td>W/C = 0.80</td>
<td></td>
<td></td>
<td></td>
<td>(5)</td>
</tr>
</tbody>
</table>

Designations according to EN 197-1: "ByggC" = CEM II/LL-42.5 R, "StdP" = CEM I 42,5 R, “StdM” = CEM II/B-V. These designations are used in figures 10-13.

RH was measured to 68-73 % in the upper and lower side of the specimens. RH=70 % was used here. It is uncertain whether this is representative for the annual climate but no better information is available. C_{PC}/C has been assumed to be 0.80 for the CEM II-cement.

No explanations have been found for the differences in carbonation depths between the two sides. The data from the bottom surfaces are used since occasional condensation at the upper surfaces could be possible.

The comparisons between the predictions with the model and the data in table II are shown in figure 10. The fly ash concrete is not included in this comparison since the model is not at all applicable for such a concrete. The effect of fly ash on the capillary porosity could be quantified but the effect on the other parameters is more uncertain. This is important to clarify since this type of cement may be use much more frequent in the future.

The comparison between measured depths of carbonation and the predicted ones show a large difference. The predictions gave much larger carbonation depths than the measurements.

The predictions are, however, based on a number of assumptions. If one instead assume that the measured RH=70 % is not representative, an RH=80 % would be much more likely for the climate at the exposure station, the comparison for the two last concretes will look like in figure 11a.
The difference between measured depths of carbonation and the predicted ones are especially large for the CEMII/A-cement concrete, cf. figure 10. In the predictions the limestone filler was regarded as completely inert and the amount of filler was assumed to be some 20%. This
may be quite unrealistic since the filler will have some reactivity which could decrease the capillary porosity to some extent and the filler content may be much lower. The degree of hydration may also be larger than 0.80. On the other hand, the field data in table II seems unrealistic; cf. the two top diagrams in figure 10. There is no major difference between the depths of carbonation in the two concrete with the same “w/c” = 0.60. This is strange and needs further analysis and research.

For comparison a prediction is done for the CEMII/A-binder with probably more realistic assumptions. The result is shown in figure 11b.

![Graph](image_url)

**Fig. 11b** The predicted depths of carbonation from the model (line) compared with the data in table II (point) for the CEMII/A concrete with w/C=0.60 exposed outdoor sheltered from rain, with new assumptions (filler content 10%, α=0.85, RH=80%).

Obviously, a possible explanation is that the filler content is small and, as a consequence, the equivalent w/c is lower than expected. Together with the higher RH the predicted depths of carbonation do not differ very much from the depths for a pure w/c 0.6 concrete.
7.3 Verification against field data in the project

Fridh & Lagerblad (2010) has measured carbonation depths, RH and degree of carbonation in a few indoor concrete structures at an age of 43 years. The degree of hydration in the samples is not known, which adds to the uncertainty. Not even the concrete compositions are known! Lagerblad (2010) made some estimates and said w/c to be around 0.55-0.60.

Fridh & Lagerblad (2010) found depths of carbonation of 35 and 70 mm, respectively, in a beam and a floor slab made of Limhamn CEMI, a low-alkali cement. A comparison with predictions from the model is shown in figure 1.

In most of the points where Fridh & Lagerblad (2010) took samples, the indoors structures had a surface coating. These coatings had a significant effect on the depths of carbonation. Their data could be roughly summarized as follows:
- Surface coatings of wall paint etc. limited the depth of carbonation to around 50 % to what was expected after some 40 years.
- Surface coatings of ceramics, linoleum, parquet flooring limited the depth of carbonation to 0-10 % of what would be expected without a surface coating.

With these rough numbers the resistances to CO₂-diffusion can be estimated with the model. For a case with an indoor structure with the assumptions and conditions as in figure 12 the depth of carbonation has been predicted with different surface resistances. The results are shown in figure 13.

![Graph showing predicted vs. data for carbonation depths.](image-url)
Fig. 13  The predicted depths of carbonation from the model (line) for indoor structures, with the same conditions as in figure 12, with surface coatings with different resistances $R=0, 350000$ & $2000000$ $s/m$, from top to bottom.

### 7.4 Conclusions on the calibration

It is obvious that the calibration of the prediction model is very uncertain. Decisive parameters are not well known. Much better data is required. Reliable measurements of the climatic conditions and their variation with time must be better known. The degree of hydration is another decisive parameter that is rarely measured. It also varies with depth which cannot be reflected in the present model.

The calibration that was done is, however, what is possible today, with the data that was available. The uncertainty for CEMII/A-cement concrete is especially large.
8 CO₂-absorption in different types of concrete structures

The model gives the amount of carbon dioxide being absorbed as kgCO₂/m² concrete surface. This number can then be multiplied with the number of m² concrete surface of each type of structure and for relevant exposure in zones at these structures. Here the absorbed CO₂ per m² is calculated, with the model, for a number of structures.

The structures that are analyzed are

1. Indoor structures
   1.1 without surface coatings
   1.2 with somewhat permeable surface coatings (paint)
   1.3 with almost impermeable surface coatings (PVC, linoleum, parquet, floor paint)
2. Slab-on-grade
   2.1 on mineral wool; traditional and self-desiccating concrete
   2.2 on expanded polystyrene
   2.3 on coarse drainage
   2.4 on sand
3. Outdoor structures
   3.1 Sheltered from rain
   3.2 Exposed to rain
4. Bridges

For these structures estimations have been done on the type of cement, cement content, w/c, degree of hydration and relevant surface coatings. The surrounding climate has been estimated as an annual average RH in the carbonated part of the concrete.

The structures are briefly commented upon in the next sections. The detailed assumptions and the results of calculations are then presented in the Appendix as depths of carbonation and amount of absorbed CO₂ after different exposure times.

8.1 Indoor structures

The indoor climate in buildings varies a lot depending on the type of building, internal moisture sources, the surrounding climate and the rate of ventilation. The surrounding climate, especially the temperature and vapour content, is very different between the north and south parts of Sweden. Extremely dry conditions are expected in well ventilated buildings in the north during winter. For floor slabs with floor heating, RH could be as low as 10-20 % in wintertime. In summertime RH of the indoor air could be 50-60 % for longer periods of time.

Since the simple model uses a constant RH, a representative value of 40 % RH has been chosen.

Modern concrete slabs in buildings are made of self-desiccating concrete (SDC), with a low w/c, around 0.40. An additional calculation has been made for indoor structures with SDC in the various cases.

The surface resistances was assumed to be 0, 350000 and 2000000 s/m for structures without
surface coatings, with somewhat permeable surface coatings ("paint") and with almost impermeable surface coatings (PVC, linoleum, parquet, floor paint), respectively.

8.2 **Slab-on-grade**
Concrete slabs-on-grade actually have carbonation from beneath according to the field investigations, Fridh & Lagerblad (2010). Why this is possible has not been explained and the few cases that were investigated had mineral wool as underlying heat insulation. This insulation creates a temperature difference between the wet soil and the bottom surface of the concrete slab which lowers the RH to some 85-90 % RH.

Mineral wool is also open for diffusion of carbon dioxide. The surface resistance should be almost zero. A heat insulation of expanded polystyrene, however, must be assumed to have an infinite surface resistance.

The temperature difference between the bottom surface of the concrete slab and the wet soil depends on a number of parameters such as the type and thickness of the heat insulation, the type of soil, the size, shape and location of the building etc. For a 30 year old, heat insulated, small building, on clay in the south of Sweden, a temperature difference of some 3°C could be expected, corresponding to an RH = 85 %. For a building without heat insulation a much smaller (if any) temperature difference is expected. For buildings on a coarse drainage layer RH = 95 % was assumed; for buildings on sand 98 % RH.

The concrete quality in modern concrete floor slabs are frequently made of SDC, with a w/c of 0.40 compared to traditionally 0.65-0.70.

8.3 **Outdoor structures**
Carbonation of outdoor structures is fairly well documented. The absorption of CO₂ is, however, expected to be small due to the wet conditions. To demonstrate a comparison with indoor structures a few predictions have been made with the model. The model uses the RH of outdoor air and, to some extent the time of wetness at the surfaces, due to rain.

The age exponent n has been assumed to be 0.17 for structures exposed to rain and 0 for structures sheltered against rain. In both cases RH of the outdoor air was assumed to be 85 %.

8.4 **Bridges**
Concrete bridges in Sweden have been produced with "bridge concrete" for decades, having a w/c of 0.45. In recent decades the cement has been "Degerhamn Anläggning", a n SRPC.

The age exponent n was assumed to be 0.17 for parts of bridges exposed to rain and 0 for parts sheltered against rain. In both cases RH of the outdoor air was assumed to be 85 %.
8.5 Summary

The results from the Appendix are summarized in the table below, as the amount of absorbed carbon dioxide during 100 years.

Table III. Results of calculations for various concrete structures, cf. Appendix.

<table>
<thead>
<tr>
<th>Concrete structure</th>
<th>Absorbed amount of carbon dioxide (kgCO₂/m²) after 100 years</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. Indoor structures</strong></td>
<td></td>
</tr>
<tr>
<td>1.1 without surface coatings (SDC)</td>
<td>6.1 (1.7)</td>
</tr>
<tr>
<td>1.2 with somewhat permeable surface coatings (&quot;paint&quot;)</td>
<td>3.6</td>
</tr>
<tr>
<td>1.3 with almost impermeable surface coatings (PVC, linoleum, parquet, floor paint)</td>
<td>0.9</td>
</tr>
<tr>
<td><strong>2. Slab-on-grade (bottom surface)</strong></td>
<td></td>
</tr>
<tr>
<td>2.1 with mineral wool (SDC)</td>
<td>4.1 (1.4)</td>
</tr>
<tr>
<td>2.2 with EPS</td>
<td>0</td>
</tr>
<tr>
<td>2.3 with coarse drainage layer</td>
<td>0.7</td>
</tr>
<tr>
<td>2.4 with sand/gravel</td>
<td>0.1</td>
</tr>
<tr>
<td><strong>3. Outdoor structures</strong></td>
<td></td>
</tr>
<tr>
<td>3.1 exposed to rain</td>
<td>0.9</td>
</tr>
<tr>
<td>3.2 sheltered against rain</td>
<td>2.8</td>
</tr>
<tr>
<td><strong>4. Bridges</strong></td>
<td></td>
</tr>
<tr>
<td>4.1 parts exposed to rain</td>
<td>0.5</td>
</tr>
<tr>
<td>4.2 parts sheltered from rain</td>
<td>1.8</td>
</tr>
</tbody>
</table>
References


Fridh, K & Lagerblad, B. (2011) *Carbonation of indoor concrete*. Internal draft report, CBI-Cement & Concrete Institute, Stockholm, and Lund University


Lagerblad, B. (2010) Private communication within the project.


APPENDIX – calculations for different structures

1 Indoor structures

Assumptions:
Type of cement: CEMI
Water-cement ratio, w/c=0.65
Cement content, C=277 kg/m³
Degree of hydration, α=0.70
RCO₂ = 0
Relative humidity, RH=40 %
Age exponent, n=0.

The surface resistances was assumed to be 0, 350000 and 2000000 s/m for structures without surface coatings, with somewhat permeable surface coatings ("paint") and with almost impermeable surface coatings (PVC, linoleum, parquet, floor paint), respectively.

1.1 without surface coatings;

1.1.1 traditional concrete

Results: (degree of carbonation = 0.35)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>12</td>
<td>17</td>
<td>27</td>
<td>39</td>
<td>55</td>
<td>87</td>
<td>123</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.6</td>
<td>0.9</td>
<td>1.4</td>
<td>1.9</td>
<td>2.7</td>
<td>4.3</td>
<td>6.1</td>
</tr>
</tbody>
</table>

1.1.2 self-desiccating concrete

Assumptions:
Type of cement: CEMI
Water-cement ratio, w/c=0.40
Cement content, C=450 kg/m³
Degree of hydration, α=0.70
RCO₂ = 0
Relative humidity, RH=40 %
Age exponent, n=0.

Results: (degree of carbonation = 0.35)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>2</td>
<td>3</td>
<td>5</td>
<td>7</td>
<td>9</td>
<td>15</td>
<td>21</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.2</td>
<td>0.2</td>
<td>0.4</td>
<td>0.5</td>
<td>0.7</td>
<td>1.2</td>
<td>1.7</td>
</tr>
</tbody>
</table>

1.2 with somewhat permeable surface coatings (paint)

Results: (degree of carbonation = 0.35)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>1</td>
<td>2</td>
<td>5</td>
<td>10</td>
<td>19</td>
<td>42</td>
<td>72</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.1</td>
<td>0.1</td>
<td>0.3</td>
<td>0.5</td>
<td>0.9</td>
<td>2.1</td>
<td>3.6</td>
</tr>
</tbody>
</table>
1.3 with almost impermeable surface coatings (PVC, linoleum, parquet, etc.)

Results: (degree of carbonation = 0.35)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>9</td>
<td>19</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.1</td>
<td>0.2</td>
<td>0.5</td>
<td>0.9</td>
</tr>
</tbody>
</table>

2 Slab-on-grade

2.1.1 Slab-on-grade on mineral wool; traditional concrete

Assumptions:
Type of cement: CEMI
Water-cement ratio, w/c=0.65
Cement content, C=277 kg/m³
Degree of hydration, α=0.80
R_CO₂ = 0
Relative humidity, RH=85 %
Age exponent, n=0.

Results: (degree of carbonation = 0.80)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>4</td>
<td>5</td>
<td>8</td>
<td>12</td>
<td>16</td>
<td>26</td>
<td>37</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.4</td>
<td>0.6</td>
<td>0.9</td>
<td>1.3</td>
<td>1.8</td>
<td>2.9</td>
<td>4.1</td>
</tr>
</tbody>
</table>

2.1.2 Slab-on-grade on mineral wool; self-desiccating concrete (SDC)

Assumptions:
Type of cement: CEMI
Water-cement ratio, w/c=0.40
Cement content, C=450 kg/m³
Degree of hydration, α=0.70
R_CO₂ = 0
Relative humidity, RH=85 %
Age exponent, n=0.

Results: (degree of carbonation = 0.70)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.1</td>
<td>0.2</td>
<td>0.3</td>
<td>0.4</td>
<td>0.6</td>
<td>1.0</td>
<td>1.4</td>
</tr>
</tbody>
</table>

2.2 Slab-on-grade on expanded polystyrene (EPS)
The objective was to be able to estimate the resistance against diffusion of carbon dioxide of the surface materials R_CO₂ through "calibration" against field data within the project. These data were, however, too few and too uncertain for such a calibration to
be possible, Fridh & Lagerblad (2010). No structures with expanded polystyrene have been studied. Because of this the resistances of the surface materials are assumed to be infinite, i.e. the absorption of CO₂ is zero.

2.3 Slab-on-grade on coarse drainage

If the flooring material is a PVC-sheet we have no other information than its resistance against CO₂-diffusion is infinite. A certain carbonation could happen from below, but there the moisture level is very large, close to 100 % RH. Here an RH of 95 % is assumed, which is reasonably true if the temperature difference between the soil and the bottom of the slab is some 1°C.

Assumptions:

Type of cement: CEMI
Water-cement ratio, w/c=0.65
Cement content, C=277 kg/m³
Degree of hydration, α=0.85
R_{CO₂} = 0
Relative humidity, RH=95 %
Age exponent, n=0.

Results: (degree of carbonation = 0.85)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.2</td>
<td>0.3</td>
<td>0.5</td>
<td>0.7</td>
</tr>
</tbody>
</table>

2.4 Slab-on-grade on sand

If the flooring material is a PVC-sheet we have no other information than its resistance against CO₂-diffusion is infinite. A certain carbonation could happen from below, but there the moisture level is very large, close to 100 % RH. Here an RH of 98 % is assumed, which is reasonably true if the temperature difference between the soil and the bottom of the slab is some 0.4°C.

Assumptions:

Type of cement: CEMI
Water-cement ratio, w/c=0.65
Cement content, C=277 kg/m³
Degree of hydration, α=0.85
R_{CO₂} = 0
Relative humidity, RH=98 %
Age exponent, n=0.

Results: (degree of carbonation = 0.85)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.01</td>
<td>0.01</td>
<td>0.02</td>
<td>0.03</td>
<td>0.04</td>
<td>0.07</td>
<td>0.09</td>
</tr>
</tbody>
</table>
3  Outdoor structures

3.1  exposed to rain

Assumptions:
Type of cement: CEM I
Water-cement ratio, w/c=0.55
Cement content, C=327 kg/m³
Degree of hydration, α=0.90
R_{CO2} = 0
Relative humidity, RH=85 %
Age exponent, n=0.17.

Results: (degree of carbonation = 0.90)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.2</td>
<td>0.3</td>
<td>0.3</td>
<td>0.4</td>
<td>0.5</td>
<td>0.7</td>
<td>0.9</td>
</tr>
</tbody>
</table>

3.2  sheltered from rain

Assumptions:
Type of cement: CEM I
Water-cement ratio, w/c=0.55
Cement content, C=327 kg/m³
Degree of hydration, α=0.80
R_{CO2} = 0
Relative humidity, RH=85 %
Age exponent, n=0.0.

Results: (degree of carbonation = 0.80)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>2</td>
<td>3</td>
<td>5</td>
<td>7</td>
<td>10</td>
<td>15</td>
<td>21</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.3</td>
<td>0.4</td>
<td>0.6</td>
<td>0.9</td>
<td>1.3</td>
<td>2.0</td>
<td>2.8</td>
</tr>
</tbody>
</table>

4  Bridges

4.1  parts exposed to rain

Assumptions:
Type of cement: CEM I
Water-cement ratio, w/c=0.45
Cement content, C=400 kg/m³
Degree of hydration, α=0.85
R_{CO2} = 0
Relative humidity, RH=85 %
Age exponent, n=0.17.
Results: (degree of carbonation = 0.85)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>0.6</td>
<td>0.7</td>
<td>1.0</td>
<td>1.2</td>
<td>1.5</td>
<td>2.1</td>
<td>2.6</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.10</td>
<td>0.12</td>
<td>0.17</td>
<td>0.21</td>
<td>0.26</td>
<td>0.36</td>
<td>0.45</td>
</tr>
</tbody>
</table>

4.2 parts sheltered from rain

Assumptions:
Type of cement: Anl
Water-cement ratio, w/c=0.45
Cement content, C=400 kg/m³
Degree of hydration, α=0.75
R_CO₂ = 0
Relative humidity, RH=85 %
Age exponent, n=0.

Results: (degree of carbonation = 0.75)

<table>
<thead>
<tr>
<th>Time (years)</th>
<th>1</th>
<th>2</th>
<th>5</th>
<th>10</th>
<th>20</th>
<th>50</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td>1.1</td>
<td>1.6</td>
<td>2.6</td>
<td>3.6</td>
<td>5.1</td>
<td>8.1</td>
<td>11.4</td>
</tr>
<tr>
<td>Amount (kgCO₂/m²)</td>
<td>0.17</td>
<td>0.25</td>
<td>0.39</td>
<td>0.55</td>
<td>0.78</td>
<td>1.24</td>
<td>1.75</td>
</tr>
</tbody>
</table>