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F. Lavergne\textsuperscript{a}, K. Sab\textsuperscript{a,∗}, J. Sanahuja\textsuperscript{b}, M. Bornert\textsuperscript{a}, C. Toulemonde\textsuperscript{b}

\textsuperscript{a}Université Paris-Est, Laboratoire Navier (ENPC, IFSTTAR, CNRS)
77455 Marne-la-Vallée Cedex, France

\textsuperscript{b}Département Mécanique des Matériaux et des Composants, EDF R&D, Site des Renardières, Avenue des Renardières, 77818 Moret-Sur-Loing Cedex, France

Abstract

Prestress losses due to creep of concrete is a matter of interest for long term operations of nuclear power plants containment buildings. Experimental studies by Granger (1995) have shown that concretes with similar formulations have different creep behaviors. The aim of this paper is to numerically investigate the effect of size distribution and shape of elastic inclusions on the long term creep of concrete. Several microstructures with prescribed size distribution and spherical or polyhedral shape of inclusions are generated. By using the 3D numerical homogenization procedure for viscoelastic microstructures proposed by Šmilauer and Bažant (2010), it is shown that the size distribution and shape of inclusions have no measurable influence on the overall creep behavior. Moreover, a mean-field estimate provides close predictions. An Interfacial Transition Zone was introduced according to the model of Nadeau (2003). It is shown that this feature of concrete’s microstructure can explain differences between creep behaviors.

Keywords: Modeling, Creep, Particle Size Distribution, Microstructure, Interfacial Transition Zone

Introduction

Concrete structures evolves due to time-dependent phenomena such as shrinkage and creep. Predicting basic creep is necessary to estimate the time-dependent prestress losses in thick structures such as nuclear containment buildings. For this purpose, experimental investigations were conducted by Granger [1] in order to assess the basic creep of six concretes with various formulations.

*Corresponding author. Tel: +33 1 64 15 37 49
Email address: karam.sab@enpc.fr (K. Sab)

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According to this study, concretes with apparently similar formulation may exhibit different long term behaviors. For instance, the concrete from Flamanville and the one from Paluel have very similar formulation regarding cement composition, water to cement ratio and grain size distribution as shown in table [Tab. 1]. Fillers were added to the concrete from Paluel so that the total amount of fine aggregates (smaller than 100 µm) was almost equal in these two concretes. However, the major difference between the two concretes is the nature of the aggregates: crushed granite (granodiorite) for Flamanville and semi-crushed sand-lime river gravel for Paluel. Actually, the time-dependent strains of these two concretes are very different as shown in figure [Fig. 1]: The average strain measured on concrete samples from Flamanville after three years of the same compressive loading was twice as much as the one measured on samples from Paluel. Since tests revealed comparable mechanical properties for aggregates, Granger [1] also measured the basic creep strains of the cementitious matrices without finding noticeable deviation. He concluded that the differences of overall creep properties were due to the aggregates (shape, mineralogy or interface with the cement paste).

<table>
<thead>
<tr>
<th></th>
<th>Flamanville</th>
<th>Paluel</th>
</tr>
</thead>
<tbody>
<tr>
<td>(12.5/25mm)</td>
<td>700kg</td>
<td>700kg</td>
</tr>
<tr>
<td>(5/12.5mm)</td>
<td>340kg</td>
<td>348kg</td>
</tr>
<tr>
<td>(0/0.8mm)</td>
<td>170kg</td>
<td></td>
</tr>
<tr>
<td>(0.1/0.5mm)</td>
<td>485kg</td>
<td></td>
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<tr>
<td>(0/5mm)</td>
<td>140kg</td>
<td>722kg</td>
</tr>
<tr>
<td>Filler Piketty</td>
<td>50kg</td>
<td></td>
</tr>
<tr>
<td>Cement (St Vigor)</td>
<td>375kg</td>
<td>375kg</td>
</tr>
<tr>
<td>(CPA HP PM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>180kg</td>
<td>180kg</td>
</tr>
<tr>
<td>BV40 plasticizer</td>
<td>1.5kg</td>
<td>1.56kg</td>
</tr>
<tr>
<td>Young Modulus of aggregates</td>
<td>51-65GPa</td>
<td>65GPa</td>
</tr>
<tr>
<td>Young Modulus of concrete at 28 days</td>
<td>31.9GPa</td>
<td>38.4GPa</td>
</tr>
<tr>
<td>Compressive strength $f_c$ at 28 days</td>
<td>50MPa-53.4MPa</td>
<td>43.0MPa-52MPa</td>
</tr>
</tbody>
</table>

Table 1: Formulation of two concretes of Granger [1]

Because experimental results are difficult to obtain, a natural way to perform parametric studies is to rely on models. The purpose of this paper is to numerically investigate the effect of size distribution and shape of inclusions on the long term creep of a matrix-inclusion material where the viscoelastic phase is the matrix. The matrix has the same behavior as a cement paste, the inclusions are perfectly bonded to the matrix and they have the elastic properties of aggregates.

One could use X-ray tomography images as inputs of 3D FFT-based compu-
Figure 1: Results of uniaxial compressive creep tests reported by L. Granger [1]. Although the formulations of concretes from Flamanville and Paluel were comparable, the basic creep strain \(10^{-6}\text{MPa}^{-1}\) vs days proved different. The compressive stress was 12 MPa and the concrete was loaded at \(t_0 = 28\) days.

This procedure has two drawbacks: the image must be segmented in order to differentiate the phases, and boundary effects may occur since the sample is not the unit cell of a periodic material. Filtering strategies have recently been defined to mitigate the effect of such boundary effects and get a macroscopically consistent estimate of the overall behaviour [3]. This study is focused on numerically generated periodic microstructures so as to tightly control the volume fraction of inclusions, their sizes and their shapes. The generation of several microstructures with prescribed grain size distribution and spherical or polyhedral shape of inclusions is made by dense sphere packing or Random Sequential Adsorption algorithm. The 3D numerical homogenization procedure for viscoelastic microstructures proposed by Šmilauer and Bažant [4] is used to compute the time-dependent response to a constant applied load. The Interface Transition Zone is described, the model of Nadeau [5] is recalled and the assumption to extend it to viscoelasticity are exposed. Then, the obtained results are compared to the predictions of mean-field homogenization.

In the first part of the paper, the microstructure generator is described, the 3D numerical homogenization method is introduced, some numerical tests are performed to validate and assess the performances of the simulations, the case of an Interfacial Transition Zone is exposed and its effect on the creep strain is numerically estimated. In the second part, the micromechanics methods are
recalled and their predictions are compared to the numerical results. The paper ends with a discussion and a conclusion.

1. Simulation of viscoelastic matrix-inclusion microstructures

1.1. Generation of matrix-inclusion microstructures

In this section, the microstructure generator and the 3D numerical method for computing the overall viscoelastic response are presented. A matrix-inclusion microstructure is generated according to given volume fraction, sieve curve and shape of inclusions.

1.1.1. Scope statement

Some features are to be enforced in order to obtain a valuable matrix-inclusion microstructure generator:

- Inclusions are placed in a cubic cell and they should not overlap.
- The microstructure should be periodic.
- A tight control of the volume fraction, the size distribution and the shape of inclusions are required.
- High volume fractions of inclusions (> 50%) are needed to represent realistic microstructures of concrete.
- The simulated material should be isotropic.
- The generator should run as fast as possible.

At least two methods may fulfill these requirements. The Random Sequential Adsorption (RSA) algorithm [6] is the simplest method to design and implement. Its ability to handle convex polyhedra makes it useful to assess the effect of shape of inclusions. A high volume fraction of inclusions (up to 68%) may be reached for wide sieve curves. On the other hand, this method fails to simulate monodisperse grain size distribution for volume fraction greater than 30%.

The other algorithm is the one of Lubashevky and Stillinger [7] which is dedicated to pack efficiently sets of ellipsoidal inclusions. In the case of monodisperse spheres, volume fraction up to 63% can be reached. To the authors’ knowledge, there is no extension of this method to polyhedral inclusions.
1.1.2. Determination of the number of inclusions and their volumes

Given the sieve curve (volume fraction vs particle diameter), the volume fraction of inclusions $f_g$ and the dimension $l$ of the cubic cell, the starting point, common to both methods, is the generation of the random number of the inclusions in the cubic cell and their volumes. The sieve curve is first approximated by a piecewise linear function of the equivalent diameters on twenty classes, each class representing 5% of the total volume of inclusions. The equivalent diameter of an inclusion, $d$, is given by $d = \sqrt[3]{\frac{\pi}{6}v}$, where $v$ is the volume of the inclusion.

Each class must provide a volume $V_{5\%} = 0.05 \times l^3 \times f_g$ of inclusions. On each class $i \in \{0, 1, ..., 19\}$, the distribution of diameters is such that the approximated sieve curve $s(d)$ is an affine function of $d$ on the interval $[d_{i+1}, d_i]$ with $D_{\text{max}} = d_0 > d_1 > ... > d_{20} = d_{\text{min}}$. [Fig. 2]. The classes are filled one after another, starting from the class 0 of largest inclusions $[d_1, d_0]$ to the class 19 of the smallest inclusions $[d_{20}, d_{19}]$. While the volume of the generated inclusions in class $i$ is less than $V_{5\%}$, a new inclusion of diameter $d_{\text{picked}}$ is repeatedly generated in this class as explained hereafter.

![Volume fraction vs particle diameter]

Figure 2: The sieve curve : volume fraction vs particle diameter. This curve is approximated by a piecewise linear function on twenty classes, each class representing 5% of the total volume of inclusions.

Let $F_i(d)$ be the probability to pick in the class $i$ an inclusion of diameter lower than $d$. Since the sieve function is piecewise linear, $F'_i(d) \frac{\pi}{6}d^3$ must be uniform. Taking into account $F_i(d_{i+1}) = 0$ and $F_i(d_i) = 1$, we find that $F_i$ is given by
\[ F_i(d) = \frac{d_i^2 - d_{i+1}^2}{d_i^2 - d_{i+1}^2} \]

Hence, \( d_{\text{picked}} \) is generated by:

\[ d_{\text{picked}} = F_i^{-1}(R()) = (d_i^2 + (d_i^2 - d_{i+1}^2)R())^{-\frac{1}{2}} \]

where \( F_i^{-1} \) is the inverse function of \( F_i \) and \( R() \) is the generator of the uniform random law on \([0, 1]\).

### 1.1.3. Convex polyhedra and the RSA algorithm

A convex polyhedron is generated aiming at controlling its shape [Fig.3,4]. A center is generated and a number of points is picked according to a Poisson law (parameter \( \lambda > 0 \)). Then, directions are selected on the unit sphere according to the uniform random law, and distances to the center are generated according to a log-normal law (parameters \( \mu_l \in \mathbb{R}, \sigma_l > 0 \)). The polyhedron is defined as the convex hull of those points which is computed through the C++ interface of the Qhull software [8,9].

Each convex polyhedron is generated according to the previously described procedure and is placed into the cubic cell avoiding overlapping with already generated polyhedra: its position is randomly picked in the cubic cell until non-overlapping is achieved. By putting the bigger inclusions first, high volume fraction may be reached. This method has already been used to generate concrete and mortar microstructures with spherical [10] or polyhedral inclusions [11]. Unlike [11], the computation of overlapping between polyhedra is performed analytically in the proposed generator. Indeed, overlapping between two polyhedra is computed thanks to the Gilbert-Johnson-Keerthi distance algorithm [12] as implemented in the Bullet physics library [13]. This procedure suppresses the exclusion-zone of one voxel adopted in [11] and it speeds up the computations. In addition, in order to limit the potential pair of polyhedra to be tested, the cubic cell is divided into sectors: two polyhedra can potentially overlap if they share the same sector. Moreover, each polyhedron is embedded in a rectangular bounding box because testing overlapping of boxes is much easier than testing overlapping of polyhedra. A polyhedron may be present in different sectors of the cubic cell in order to ensure periodicity.

The maximum reachable volume fraction depends on the size distribution and on inclusions’ shape. A wide range of inclusions’ sizes allows larger volume fractions. Simulating spherical inclusions is faster than simulating unaligned elongated inclusions with sharp edges. Artificial concrete microstructures with high volume fraction (60 – 68\%) and polyhedral inclusions can be built with the RSA method for some size distributions [Fig. 3]. In particular, the ratio between the largest inclusions’ equivalent diameter to the smallest one must be less than 20. If this ratio is too large, the number of inclusions would become prohibitive.
If this ratio is too small, reaching the given volume fraction would take too long or could not be achieved.

Figure 3: A section of a 4 cm-wide sample of a concrete made of coarse sand \((d > 0.5 \text{ mm})\) and small gravel \((d < 13.5 \text{ mm})\) built by the RSA algorithm is shown. There are 28760 polyhedra in the cubic cell and the volume fraction is 68%.

Figure 4: On the left, polyhedra are rather round whereas polyhedra feature sharp edges in the middle. The RSA algorithm has been used. The sieve curve of the corresponding mortar is presented on the right.

1.1.4. Lubashevsky and Stillinger algorithm for spherical inclusions

The RSA algorithm described above cannot achieve high volume fraction for monodisperse particles. For this reason, we use the Lubashevsky and Stillinger algorithm for spherical inclusions.
inger algorithm for generating matrix-inclusion microstructures with monodisperse spherical inclusions.

In our implementation, we use the algorithm described in [7] and [14] for simulating polydisperse packing of spherical inclusions. This dynamic, event-driven algorithm enables very high packing ratio of spheres. Seeds are placed in the cubic cell and their growth rate are chosen to comply with the size distribution. Each radius increases proportionally to time $t$ and each center moves with a constant velocity. As spheres collide, the algorithm changes their velocities to avoid overlapping. Thanks to uniform speeds and growth rates, collisions, or events, can be computed analytically. There is no need for time steps since nothing happen until the next event. Therefore the procedure jumps from one event to another and the program runs tremendously fast. This method has been extended to ellipsoids [15, 16], but this implementation is limited to spherical inclusions or sets of spheres. This algorithm has already been used to model composite materials in the range of elasticity [17].

1.2. Numerical determination of the effective viscoelastic properties of heterogeneous materials

1.2.1. The unit cell problem

The aim of this section is to numerically determine the overall viscoelastic behavior of a matrix-inclusion microstructure where the inclusions are elastic and the matrix viscoelastic.

The strain tensor $\varepsilon(t)$ in a viscoelastic material depends on the history of stress tensor $\sigma(t)$. If the constitutive law is linear, the Boltzmann superposition principle states that the material properties are defined by a compliance function (fourth order tensor), $J(t, t')$, such that:

$$\varepsilon(t) = \int_0^t J(t, t') \frac{d\sigma}{dt}(t') dt'$$

If the elapsed time since loading is the only relevant parameter, the material is non-aging:

$$J(t, t') = J(t - t')$$

The determination of the overall viscoelastic behavior of a periodic microstructure can be obtained by solving the following auxiliary problem on the periodic unit cell $V$.

$$\begin{align*}
\text{div } \sigma(x, t) &= 0 & x \in V \\
\varepsilon(x, t) &= \int_0^t J(x, t, t') \frac{d\sigma}{dt}(x, t') dt' & x \in V \\
\varepsilon(x, t) &= E(t) + \nabla^s u(x, t) & x \in V \\
u(x, t) &= \text{periodic} & x \in \partial V \\
\sigma(x, t) \cdot n(x) &= \text{skew-periodic} & x \in \partial V
\end{align*}$$

Here $E(t)$ is the time-dependent overall strain, $u(t, x)$ is the displacement field in $V$, $\nabla^s u(x, t)$ is its symmetric gradient, $\partial V$ is the boundary of $V$ and $n(x)$ is
the outer normal to $\partial V$. Actually, $E(t)$ is the volume average of $\varepsilon(x,t)$ and we denote by $\Sigma(t)$ the volume average of $\sigma(x,t)$.

1.2.2. Numerical discretizations

We use the computational method of Šmilauer and Bažant in [4] developed for cementitious materials. This method which relies on the exponential algorithm [18, 19, 20] is a time-iteration procedure to solve the viscoelastic problem for the case of steady loads. It features an integration of the constitutive equations on each time step assuming a constant stress rate. It enables the time step to grow exponentially when performing a relaxation (or creep) simulation. This method has recently made use of the FFT algorithm as the solver for the unit cell tangent problem [4, 21].

Let $\Delta \varepsilon(x) = \varepsilon(x, t_{i+1}) - \varepsilon(x, t_i)$, $\Delta \sigma(x) = \sigma(x, t_{i+1}) - \sigma(x, t_i)$ and $\Delta u(x) = u(x, t_{i+1}) - u(x, t_i)$ be respectively the strain, stress and displacement increment between time $t_i$ and $t_{i+1}$. These fields are solution of the following unit cell tangent problem:

\begin{align*}
\text{div} \Delta \sigma(x) &= 0 & x \in V \\
\Delta \varepsilon(x) &= \int_{t_i}^{t_{i+1}} J(x, t_{i+1}, t') \frac{d\sigma}{dt}(x, t') dt' + \varepsilon_0(x) & x \in V \\
\Delta \sigma(x) &= \Delta E + \nabla \cdot \Delta u(x) & x \in V \\
\Delta \sigma(x) \cdot n(x) &= \text{skew} - \text{periodic} & x \in \partial V \\
\Delta \varepsilon(x) &= \Delta E + \nabla \cdot \Delta u(x) & x \in \partial V
\end{align*}

where

$$\varepsilon_0(x) = \int_{t_i}^{t_{i+1}} \left[ J(x, t_{i+1}, t') - J(x, t_i, t') \right] \frac{d\sigma}{dt}(x, t') dt'$$

and

$$\Delta E = E(t_{i+1}) - E(t_i).$$

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{rheological_model.png}
\caption{The rheological model is made of a series of Kelvin chains}
\end{figure}

It is assumed that the stress rate $\frac{d\sigma}{dt}$ in the interval $[t_i : t_{i+1}]$ is constant and
that the compliance corresponds to a series of Kelvin chains [Fig. 5]:

\[ J(t - t') = \int_{t'}^{t} \sum_{k=1}^{n} \tau_k e^{-\frac{t - t'}{\tau_k}} C_k^{-1} d\tau + C_0^{-1} \]

where \( C_k, k = 0, 1, \ldots, n \) are \( n + 1 \) fourth-order elasticity tensors and \( \tau_k, k = 1, 2, \ldots, n \) are \( n \) characteristic times. The integration of the constitutive equations on the time step gives the following incremental constitutive equation:

\[ \Delta \sigma(x) = C_i(x)(\Delta \varepsilon(x) - \varepsilon^i(x)) \]

where \( C_i(x) \) and \( \varepsilon^i(x) \) are explicitly given by:

\[ (C_i)^{-1} = C_0^{-1} + \sum_{k=1}^{n} \left( 1 - \frac{1}{\tau_k} e^{-\frac{t_{i+1} - t_i}{\tau_k}} \right) C_k^{-1} \]

\[ \varepsilon^i = \sum_{k=1}^{n} \tau_k (1 - e^{-\frac{t_{i+1} - t_i}{\tau_k}}) \gamma_k^i \]

where \( \gamma_k^i \) are \( n \) symmetric second-order strains corresponding to internal variables of Kelvin chains [Fig. 5]. They are defined incrementally as \( \gamma_k^0 = 0 \) and for \( i \geq 0 \):

\[ \gamma_{k+1}^i = 1 - e^{-\frac{t_{i+1} - t_i}{\tau_k}} C^{-1} \Delta \sigma + e^{-\frac{t_{i+1} - t_i}{\tau_k}} \gamma_k^i \]

Notice that the choice of a series of Kelvin chains combined with the above described exponential algorithm allows for the introduction of these internal variables. Hence, storing these few internal variables enables large computations (FEM or FFT).

To solve the unit cell tangent problem at each time step, a method is required. The FFT algorithm of H. Moulinec and P. Suquet has proved a powerful tool to compute the elastic [22] and viscoelastic [23] response of a periodic microstructure. It takes digital images of materials and prestress as inputs. Both the basic [Alg.1][22] and the accelerated [Alg.2][24, 25] versions are written for this study. An isotropic reference material \( C_0^* \) is to be introduced and its Green operator \( \Gamma_0^* \) is applied in the Fourier space. The loading can be either the prescribed macroscopic strain or the prescribed macroscopic stress as explained in [22].

Regarding implementation details, the MPI extension of the FFTW library [26,27] and the PETSc library [28,29,30] are combined. The software routinely performs its computation on several processors on shared-memory machines as well as on clusters (Ivanhoe at EDF R&D). Using less than twenty nodes and about 100-200 processes on this cluster, the size of the problem may be increased up to \( 700 \times 700 \times 700 \) for elasticity and \( 400 \times 400 \times 400 \) for viscoelasticity with 8 Kelvin chains and our computations last less than two hours (180 time steps).
1.2.3. Constituents’ properties

Long term behavior of concretes were experimentally identified by Granger in [1]. Characteristic times $\tau_k$ and stiffness $C_k$ of Kelvin chains correspond to a creep test of a concrete (Penly) loaded at 28 days. Eight characteristic times were introduced, one for each decade, from 0.002 days to 20000 days. Properties of the cement paste are assumed to be similar. To model a viscoelastic cementitious matrix, the elastic and viscoelastic stiffnesses of a concrete were divided by three [Tab. 2]. Its Young modulus is $E_{\text{matrix}} = 12$ GPa, which is comparable to the measured ones of reconstituted cementitious matrices [1]. A reverse identification of these values could have been performed. All Poisson’s ratios are assumed to be 0.2. The behavior of inclusions is elastic isotropic (Young Modulus $E = 60$ GPa and Poisson’s ratio 0.2 ).

<table>
<thead>
<tr>
<th>$\tau_k$ (days)</th>
<th>Concrete, L. Granger</th>
<th>cement paste</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elastic</td>
<td>36.02</td>
<td>12</td>
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<tr>
<td>0.002</td>
<td>1901.73</td>
<td>634</td>
</tr>
<tr>
<td>0.02</td>
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<tr>
<td>0.2</td>
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<td>2</td>
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<tr>
<td>20</td>
<td>112.68</td>
<td>37.6</td>
</tr>
<tr>
<td>200</td>
<td>54.79</td>
<td>18.3</td>
</tr>
<tr>
<td>2000</td>
<td>36.5</td>
<td>12.2</td>
</tr>
<tr>
<td>20000</td>
<td>27.04</td>
<td>9.01</td>
</tr>
</tbody>
</table>

Table 2: Mechanical behavior of a concrete identified on a creep test [1] and the behavior of the defined cement paste. All Poisson ratios are assumed to be 0.2.

1.2.4. Studied microstructures

To explore the effect of grain size distribution and shape, the volume fraction was set to 63% and three different microstructures were defined:

- Unimodal: all inclusions are spherical and their diameters are equal. There are 150 inclusions for a volume of 8 mm$^3$ and their radius is about 0.2 mm.

- Bimodal: all inclusions are spherical. Two diameters coexist: about 0.2 mm and 1 mm.

- Concrete: inclusions are polyhedra and their grain size distribution is described on [Fig. 3]. Regarding their shape, crushed inclusions correspond to convex hulls of about $\lambda = 12$ points whose distances to center are picked according to a log-normal law ($\mu_l = 0.08, \sigma_l = 0.02$) before scaling.
With these parameters, the polyhedral inclusions are similar to inclusions used in concrete mixes. Indicators describing the inclusions’ shape such as elongation ratio and flakiness ratio have been defined and measured on real samples of aggregates [31, 32, 33, 34]. For an ellipsoidal inclusion with principal axes of lengths $a > b > c$, the elongation ratio is $a/b$ and the flakiness ratio is $c/b$. The average elongation ratio of the polyhedral inclusions used in our simulations is 1.08 and their average flakiness ratio is 0.8. The elongation index and flakiness index correspond to the volume fraction of inclusions featuring respectively an elongation ratio greater than 1.8 and a flakiness ratio lower than 0.6. For normal mix design, the combined (flakiness + elongation) index for coarse aggregates must be limited to 25%, for workability reasons [35]. In our simulations, the elongation index is less than 1% and the flakiness index is 11%.

Examples of these microstructures are displayed on [Fig. 9].

1.2.5. Validation of the numerical procedure

To ensure the reliability of the method and its implementation, some basic tests were performed.

**Time Discretization**

Is the response of a homogeneous microstructure precisely computed? The integration of the constitutive law on the time step may induce a numerical error. In the case of Kelvin chain, the result of a relaxation test is estimated by the Laplace-Carson transform and it serves as a reference to compute the relative numerical error for the cement paste (Fig. 6). It should be emphasized that the long term stress of a series of Kelvin chain under a constant strain is steady. Therefore the numerical error due to time discretization vanishes. Different values of the rate $b = \frac{t_{i+2} - t_{i+1}}{t_{i+1} - t_i}$ of the geometric growth of the time steps were tested. In all our simulations, we chose the geometric growth $b = 1.12$. Therefore, 180 unit cell tangent problems have to be solved to cover the range between $t - t_0 = 5 \times 10^{-6}$ days and $t - t_0 = 21900$ days where $t_0$ is the time of the initial loading.

**Space discretization**

In the case of heterogeneous microstructures, the discretization of the FFT grid also triggers numerical errors:

- The stress and strain field are stored per voxel which induces numerical errors. See [36] for further discussions about this issue.

- The heterogeneous microstructure is projected on a grid. Therefore, there is a loss of information about the local behavior of the material. A critical comparison of numerical methods working on conforming meshes or Cartesian grids was performed in the range of thermal conduction and linear elasticity at high phase contrast [37]. FFT solvers lead to very good
approximations of overall properties since extremely fine meshes can be considered.

In order to lower the error due to the projection on a grid, for each voxel, a local volume fraction based on 64 sensing points is computed and a Reuss-like constitutive law is computed and assigned to each voxel. Therefore, the microstructure is made of black voxels (pure matrix), white voxels (pure inclusions) and gray voxels (composite). To estimate the error due to discretization, the response of the unimodal microstructure with a volume fraction of 55% is computed at various resolutions [Fig. 78]. The use of a low contrast between the Young modulus of the phases (<20), the ability of parallel FFT to deal with large grids and the choice of the Reuss bound for gray voxel lead to a mesh-independent overall response.

1.2.6. Representative Volume Element

It is well-known that the asymptotic overall response should not depend on the generated sample neither on the size of the unit cell which must be large enough to be representative of the microstructure ([38, 39] among others). For each microstructure defined in section 1.2.4, different size of the unit cell and many samples were generated. Numerical hydrostatic and shear creep tests are performed.

Two criteria were defined to ensure that the unit cell is a representative volume element :

- For a hydrostatic creep test, the unit-cell hydrostatic compliance at time \( t \) is defined as \( \frac{1}{3K(t)} = \frac{tr(\varepsilon(t))}{tr(\sigma)} \) and for a shear creep test, the unit-cell shear
compliance is $G(t) = \frac{\varepsilon(t)}{\sigma}$ The corresponding relative standard deviation of the compliance is defined as the ratio of the standard deviation to the average of the unit-cell compliance over the samples that were generated. It represents the statistical dispersion of the estimated overall time-dependent strains and it is expected to decrease as the size of the unit cell increases.

- An anisotropic indicator $a$ is defined to check that the unit-cell compliance is isotropic. For a hydrostatic creep test, it is defined as
  $$a = \sqrt{\frac{(\varepsilon_{xx}^{dev})^2 + (\varepsilon_{yy}^{dev})^2 + (\varepsilon_{zz}^{dev})^2 + 2\varepsilon_{xz}^2 + 2\varepsilon_{yz}^2 + 2\varepsilon_{xy}^2}{(tr(\varepsilon)/3)^2}}$$
  where $\varepsilon^{dev}$ is the deviatoric part of the overall strain. For a shear creep test, $\sigma_{xy}$ being the direction of loading, it writes :
  $$a = \sqrt{\frac{\varepsilon_{xx}^2 + \varepsilon_{yy}^2 + \varepsilon_{zz}^2 + 2\varepsilon_{xz}^2 + 2\varepsilon_{yz}^2}{\varepsilon_{xy}^2}}$$
  $a$ is positive and $a$ is null if and only if the overall strain corresponds to the response of an isotropic material.

Figure 7: Top: One microstructure of concrete of size 40 mm made of 63% of inclusions at discretization $100^3$, $200^3$ and $400^3$. Bottom: The delayed responses to hydrostatic (left) and shear (right) creep tests are computed.
The results of these tests are described in [Tab. 3]. The relative standard deviation of the estimated macroscopic compliances are very low and the overall response is isotropic. These features are obtained for cubic unit cells of concrete of 4 cm × 4 cm × 4 cm size, the largest inclusion diameter being less than 1.25 cm and the constituents’ properties being those described in section 1.2.3. In the case of bimodal microstructures, a 2mm unit cell, twice as large as the largest aggregates, was sufficient to retrieve a precise estimate of the overall strain. The frame of periodic homogenization allows precise results on small unit cells in the range of elasticity [40, 41]. Gal and Kryvoruck have performed 3D elastic computations on 50mm large samples of concrete, the maximum diameter of aggregates being 15mm [42]. Our simulations indicate that the size of the RVE remains small in the range of linear viscoelasticity, as long as the elastic contrast between phases is limited, while it may enlarge for phenomena like plasticity [43] or fatigue [44].

1.2.7. Influence of grain size distribution and shape

Responses to hydrostatic and shear creep tests are computed, the matrix being made of cement paste. The microstructures are those defined in section 1.2.4 on page 11. The volume fraction of inclusions is set at 63%.

The overall time dependent strains remain close, even for very different microstructures[Fig. 9]. The sieve curve and the shape of inclusions have little effect on the overall viscoelastic behavior of the matrix-inclusion composite. Therefore, the only relevant parameter is the volume fraction of inclusions.

Thanks to 3D computations, stress concentrations can be estimated. During
<table>
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<tr>
<th>microstructure</th>
<th>size of cubic cell (mm)</th>
<th>maximum of inclusion diameter (mm)</th>
<th>average compliance, in $10^{-6}$MPa$^{-1}$</th>
<th>compliance relative standard deviation</th>
<th>anisotropic indicator $a$</th>
<th>number of tests</th>
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<td></td>
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</tr>
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<td>0.01</td>
<td>10</td>
</tr>
<tr>
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<td>19.8</td>
<td>0.0005</td>
<td>0.003</td>
<td>5</td>
</tr>
<tr>
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<td>0.0005</td>
<td>0.013</td>
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<tr>
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<td>0.0003</td>
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<td>0.0005</td>
<td>0.012</td>
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<tr>
<td>$t - t_0 = 21900$ days</td>
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<td>39.3</td>
<td>0.002</td>
<td>0.006</td>
<td>10</td>
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<tr>
<td>$t - t_0 = 21900$ days</td>
<td></td>
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<td></td>
</tr>
<tr>
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<td>88.9</td>
<td>0.02</td>
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<tr>
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<td>0.4</td>
<td>88.3</td>
<td>0.005</td>
<td>0.005</td>
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<td>0.01</td>
<td>5</td>
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<td>92.7</td>
<td>0.006</td>
<td>0.02</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 3: For each kind of microstructure and different size of the unit cell, many samples are built and numerical hydrostatic and shear creep tests are performed. The relative standard deviation of the estimated macroscopic compliances are very low and the overall response is isotropic.
a shear creep test $\sigma_{xy} = 1\text{MPa}$, the instantaneous local shear stress is above 1.37MPa in 5% of the microstructure. After 60 years, the local shear stress is above 1.71MPa in 5% of the microstructure: for prescribed constant average stress, the stress distribution in the microstructure evolves with time. The matrix tends to relax while the stress increases in inclusions [Fig. 10]. The concrete and the bimodal microstructures cannot be distinguished based on the probability distributions of the stress component $\sigma_{xy}$ during a shear creep test [Fig. 10].

The limited heterogeneity between the inclusions and the matrix may be one of the reasons for the overall time dependent strains being so close. On the first time step, the elastic loading, the inclusions are 5 times stiffer than the matrix. On the last time step, 60 years after loading, the tangent behavior of the inclusions is 14 times stiffer than the one of the matrix.

As long as concrete is modeled as a matrix-inclusion microstructure, the discrepancies between creep strains of similar concretes cannot be attributed to the sieve curve or to the shape of inclusions. The description of concrete’s microstructure must be refined.

Considering the cementitious matrix as homogeneous seems arguable as there is no strong scale separation in concrete. Considering a 40 mm-wide sample of concrete and a $400^3$ discretization, each voxel is 100 $\mu$m wide. At this scale, SEM images [45, 46, 47] clearly show that the behavior of the matrix may differ from one voxel to another due to fine aggregates, bundled partly unhydrated cement grains or a variable porosity [Fig. 11]. The assumption of a matrix-inclusion material might also be questionable for concrete since there sometimes exists an Interfacial Transition Zone between the inclusions and the cement paste. This feature is studied in the next section.

1.2.8. Introducing an Interfacial Transition Zone

The Interfacial Transition Zone is a $\approx 20 \mu$m-thick layer of the cement paste that is more porous than the rest of the cement paste and has a different chemical composition [45]. It is due to the imperfect packing of cement grains (< 50 $\mu$m) on aggregates known as wall effect and it evolves during hydration. This feature may be observed on SEM images [45, 46, 47] and it might affect the mechanical properties [45, 48]. Garboczi and Berryman have successfully compared their analytical model to numerical elastic computations in the range of elasticity, the ITZ being introduced as effective layers around particles[49]. For the elastic 3D computations of Gal and Kryvoruck [42], this porous zone is merged with the aggregates. This procedure increases the volume fraction of inclusions, but it decreases their elastic stiffness, especially for small aggregates if the thickness of the ITZ is taken as uniform. For the 2D finite element computations of Grondin [50], an effective mixed interphase around each aggregate is formed by the ITZ and by a volume fraction of the bulk cement paste. Micromechanics methods may
Figure 9: Viscoelastic responses to hydrostatic or shear macroscopic loadings. Various matrix-inclusion microstructures with the same inclusions’ volume fraction lead to very close time-dependent strains. The unimodal microstructures seem to creep slightly less than the other microstructures.
Figure 10: Probability distributions of a stress component $\sigma_{xy}$ during a shear creep test for different microstructures. On the top and bottom left, the relaxation of the matrix and the increase of stress in aggregates are displayed. On the bottom right, there are some differences between unimodal and concrete microstructures after 60 years.

Figure 11: A blue 100 $\mu$m grid is put on top of a SEM image from Scrivener [45]. The red lines indicate the largest inclusions. The cement paste is clearly heterogeneous over a 10 0$\mu$m voxel.
also take account of the ITZ as an effective layer surrounding inclusions \[5, 49\] or as an imperfect bonding between inclusions and matrix \[51\].

Representing the ITZ would not be possible a concrete scale since the size of the voxel would be 100 µm. Multiscale approaches may deliver a consistent way to set the behavior of composite voxels \[3\], since the finer details of the microstructure cannot be accurately described at concrete scale. To perform realistic homogenization computation, a 4 mm-wide sample of a mortar was considered, so that 10µm details such as an ITZ can be pictured. The model of Nadeau \[5\] described hereafter was used to estimate the local behavior as a function of the distance to the nearest inclusion. Let \( \delta \) be the thickness of the ITZ.

- The local volume fraction of cement \( \alpha_c(d_i) \) depends on the distance to the nearest inclusion \( d_i \):
  \[
  \alpha_c(d_i) = \begin{cases} 
  \bar{\alpha}_c \left(1 - 0.5 \left( \frac{\delta - d_i}{\delta} \right)^2 \right) & d_i < \delta \\
  \bar{\alpha}_c & d_i > \delta
  \end{cases}
  \]
  where \( \bar{\alpha}_c \) is the volume fraction of cement far from the inclusions. As described below, it is computed for each microstructure so that the overall water to cement ratio \( w/c \) is equal to 0.48.

Let \( f(d_i)d\delta \) be the volume fraction of the cementitious matrix located at a distance of the nearest inclusion between \( d_i \) and \( d_i + \delta \). \( f(d_i) \) is numerically estimated for each microstructure. The following equality stands:
  \[
  w/c = \frac{\int_{0}^{\infty} \alpha_w(x)f(x)dx}{\rho_c \int_{0}^{\infty} \alpha_c(x)f(x)dx}
  \]
  where \( \rho_c \approx 3.15 \) is the density of cement and \( \alpha_w(d_i) \) is the local volume fraction of water. If cement and water are the only constituents of the cement paste, \( \alpha_w(d_i) + \alpha_c(d_i) = 1 \) for all \( d_i \) and :
  \[
  \int_{0}^{\infty} \alpha_c(x)f(x)dx = \frac{1}{1 + \rho_c w/c}
  \]
  and finally :
  \[
  \bar{\alpha}_c = \left(1 + \rho_c w/c\right)\left(\int_{0}^{\delta} \left(1 - 0.5 \left( \frac{\delta - x}{\delta} \right)^2 \right) f(x)dx + \int_{\delta}^{\infty} f(x)dx \right)^{-1}
  \]

- The elastic behavior \( C_0 \) as a function of the local water to cement ratio, equal to \( \frac{\alpha_w}{\rho_c \bar{\alpha}_c} \), is provided by fit on experimental tests, as reported by Nadeau \[52\]. To set viscoelastic parameters, two assumptions were made.
  
  1. The viscoelastic Poisson’s ratio of each Kelvin chain is the same as the elastic one.
2. The effect of water to cement ratio on the elastic and viscoelastic parameter are identical: \( C_k = A_k C_0 \left( \frac{\alpha_w}{\rho_c \alpha_c} \right) \) where \( A_k \) is a constant. For instance, if increasing the water to cement ratio from 0.48 to 0.55 decreases the elastic Young Modulus \( (C_0) \) by 30%, then the drop of all viscoelastic modulus of Kelvin chains \( C_k, k \geq 1 \) is also 30%.

The assumption \( C_k = A_k C_0 \left( \frac{\alpha_w}{\rho_c \alpha_c} \right) \) holds true if the cement paste is considered as a porous material which solid phase is a non aging linear viscoelastic material with a uniform Poisson’s ratio. Indeed, in this case, at the scale of the porous material, the local compliance writes:

\[
J(t, x) = J(0, x)f(t)
\]

where \( f(t) \) is a scalar function of time \( t \). The determination of the overall response of the cement paste as a function of porosity necessitates the resolution of a homogenization problem on the unit cell representing the porous material.

In the Laplace-Carson space (Appendix B), the constitutive equation for this unit cell problem becomes:

\[
J^*(p, x) = J(0, x) f^*(p)
\]

Let \( \varepsilon(x) \) be the elastic solution of the unit cell problem for prescribed overall strain \( E \). Then, \( \varepsilon(x) \) is also the solution of the unit problem with prescribed overall strain \( E \) for all \( p \) in the Laplace Carson space. Therefore:

\[
J_{\text{hom},*}(p) = J_{\text{hom}}(0) f^*(p)
\]

Inverting the Laplace Carson transform gives:

\[
J_{\text{hom}}(t) = J_{\text{hom}}(0) f(t)
\]

This demonstrates that if the cement paste is a two-phase porous material, the local compliance having a uniform in time Poisson’s ratio, then the cement paste has a uniform in time Poisson’s ratio and the effect of time and microstructure are uncoupled.

If the effect of changing the water to cement ratio were limited to a change of porosity in a two-phase porous material, the consequence on the viscoelastic behavior would be similar to the one described above. But it is known that the microstructure of the cement paste includes porosity, elastic unhydrated cement, elastic or viscoelastic products of hydration and other additives [53]. The higher the water to cement ratio is, the higher the hydration degree becomes [54] and the hydrated phases are assumed to be responsible for the viscoelastic behavior of cement paste [4]. Therefore, the assumption \( C_k = A_k C_0 \left( \frac{\alpha_w}{\rho_c \alpha_c} \right) \) probably underestimates the long term effect of an ITZ.

Numerical tests were performed on a 4 mm wide cubic cell depicting the microstructure of a mortar [Fig. 12]. At this scale, a voxel is approximately 10 \( \mu \)m wide and the ITZ can be introduced. Various thickness \( \delta \) between 0 \( \mu \)m (no ITZ) and 50 \( \mu \)m were tested. The larger the ITZ is, the lower the elastic and viscoelastic stiffnesses are. The relative decreases of identified viscoelastic
Lamé coefficient $\lambda$ (GPa)
at elastic time step, $t - t_0 = 0$,
for $\delta = 0 \mu m$, 20$\mu m$ and 50$\mu m$.

Figure 12: Top : an ITZ can be pictured on the microstructure of a mortar, depending on its thickness $\delta$. Bottom left : the local water to cement ratio is displayed as a function of the distance to the inclusion. The corresponding elastic Young modulus and the Young modulus of a 10 $\mu m$ voxel overlapping with inclusions are also displayed. Bottom right : the numerical solution to uniaxial creep test depends on the thickness of the ITZ. Both the elastic response and the long term strain are modified by the introduction of an ITZ.

stiffnesses are twice larger than the one of the elastic stiffness [Tab. 4]. Moreover, according to the previous paragraph, it is likely that this effect is underestimated by our computations.

Therefore, the presence of an ITZ is a plausible explanation of discrepancies between creep strains of various concretes.

2. Comparison of numerical and micromechanics methods

2.1. Semi-analytic methods

2.1.1. Linear viscoelastic homogenization

Micromechanics methods remain practical to upscale the overall viscoelastic properties. In the frame of non-aging creep, the Laplace-Carson transform turns the time-depend problem into several elastic problems thanks to the correspondence principle [55, 56]. In this space, the Hashin-Shtrikman bounds [57], the Mori-Tanaka model [58, 59], the three-phase model of Christensen & Lo [60] or the $n + 1$-phase spherical model of Hervé & Zaoui [61] deliver estimates of the macroscopic response. Coming back to the real space requires inverting the Laplace-Carson transform with numerical tools such as the Gaver-Stehfest algorithm [62, 63] or the collocation method [64, 65]. Semi-analytical elastic homog-
Table 4: Effect of the ITZ on the identified macroscopic Young modulus of $C_k$. The effect of the interface is more important on the long term creep behavior than on the elastic behavior as shown by the relative difference to the no-ITZ case.

enization models may also provide the time-dependent response through time stepping procedures [66, 67, 68]. The analytic method combining the Hashin-Shtrikman bound and the Laplace Carson transform is described in Appendix B.

2.1.2. The tri-sphere model: a scalar parameter to account for the morphology of the microstructure

Compared to the Hashin-Shtrikman model, the tri-sphere model of de Larrard & Le Roy [69, 70] introduces an additional parameter $g^*$ to account for the morphology of the microstructure. The cementitious matrix is divided in two parts. A minimal part fills the gaps between the aggregates at maximum packing density $g^*$. The rest of the matrix ensures good rheological properties.

The corresponding homogenization procedure has two steps. First, the minimal part is merged with the inclusions using the upper Hashin-Shtrikman bound (volume fraction $g^*$). Then, the rest of the matrix is introduced thanks to a lower Hashin-Shtrikman bound [Fig. 13].

The close packing volume fraction $g^*$ arises from the compressible packing model of de Larrard [69], which extends the linear packing model of Stovall [71]. It depends on the close packing density $\Phi^*_i$ and volume fraction of solid material $y_i$ of each class $i$ of the sieve curve.

- For unimodal spherical inclusions, $g^* = \Phi^* = \frac{\pi}{3\sqrt{2}} \approx 0.74$

- For bimodal spherical inclusions and if there exists no interaction between the two classes:

$$g^* = \min \left( \frac{\Phi^*}{1 - y_{small}}, \frac{\Phi^*}{1 - (1 - \Phi^*)(1 - y_{small})} \right)$$

At packing limits, if $y_{small} < 0.21$, large inclusions dominate and small inclusions fill in the gaps. On the opposite, if $y_{small} > 0.21$, large inclusions...
are scarce and small inclusions are tightly packed. The bimodal distribution used in our numerical simulations has a volume fraction of solid material of small inclusions of $y_{\text{small}} = 0.5$.

- A more precise knowledge of interactions between classes is required to address continuous inclusions’ size distributions and irregular shapes. Typical values of $g^*$ for concretes are between 80% and 90%.

2.1.3. $n+1$–phase spherical model to account for the Interfacial Transition Zone

The $n+1$–phase spherical model of Hervé & Zaoui [61][Fig. 16] extends the three-phase model of Christensen & Lo [60]. It has been used by Nadeau [5] to upscale the elastic properties of mortars with an ITZ. The ITZ is depicted as successive layers of cement paste having different mechanical properties. This method is combined with the Laplace-Carson transform to estimate the time dependent strain of such a material.

In our case, the inputs are the same as in the numerical method: the distribution of the cement paste as a function of the distance to the nearest inclusion $f(d_i)$ is provided by the microstructure. The link between $d_i$, the water to cement ratio and the viscoelastic properties is described in section 1.2.8. Each layer has a width of $1 \mu m$ and the corresponding volume fraction of the microstructure is estimated according to its discretized image. These volume fractions are then used as inputs for the $n+1$–phase spherical model in the Laplace-Carson space.

The sieve curve and the shape of inclusions change the output of this micromechanics method since they affect the distribution of the cement paste as a function of the distance to the nearest inclusion $f(d_i)$ and the local volume fraction of cement $\alpha_c(d_i)$.

Outputs of these models are to be compared to full field numerical results.

2.2. Comparison with numerical estimates

2.2.1. Influence of volume fraction

Concrete microstructures made of crushed inclusions are built thanks to the RSA algorithm. Three unit cells of each volume fraction of inclusions are built and
Figure 14: Viscoelastic responses to a shear loading for different volume fraction of inclusions. Results of crushed inclusions are close to the Hashin-Shtrikman bound in the Laplace Carson space.

Figure 15: Results of the tri-sphere model [69, 70] for unimodal (left), bimodal (center) and concrete microstructures are compared to numerical estimates. The volume fraction of inclusions is 63%.

discretized on 384³ grids [Fig. 4] to run viscoelastic computations. A numerical shear creep test is performed on these unit cells. The overall strains is close the one estimated by the Hashin-Shtrikman bound in the Laplace-Carson space [Fig. 14] at different volume fractions of inclusions.

The volume fraction of inclusions being set to 63%, a small deviation from the Hashin-Shtrikman bound in the Laplace-Carson space occurs at long term [Fig. 9]. The heterogeneity between the inclusions and the matrix may explain this feature. On the first time step, the elastic loading, the inclusions are 5 times stiffer than the matrix. On the last time step, 60 years after loading, the tangent behavior of the inclusions is 14 times stiffer than the one of the matrix.
2.2.2. Hashin-Shtrikman, tri-sphere model and numerical estimates

The creep strains estimated by the tri-sphere model are close to our numerical results for the bimodal and concrete microstructures. As $g^*$ is large, this model delivers estimates close to the Hashin-Shtrikman ones. But estimated time-dependent strains are different for the unimodal microstructure. Though the tri-sphere model anticipates a lower creep strain due to $g^* = 0.74$, the numerical results for unimodal microstructures remain close to the other ones [Fig. 15]. Since concrete materials exhibit large $g^*$, the three methods are relevant to produce estimates of the overall macroscopic strain.

2.2.3. Interfacial Transition Zone : $n + 1$ phase spherical model and numerical estimates

Numerical uniaxial creep tests are performed on the mortar with ITZ defined in section 1.2.8 on page 21. There is a difference between the estimated overall strain estimated by the numerical method and the one produced by the $n + 1$-phase spherical model: the later seems to slightly underestimate the effect of the ITZ [Fig. 16].

Inputs are similar up to the mechanical computations. Therefore, representing the ITZ as successive layers of coating around the inclusions is not precise for this mortar. At this scale, with this volume fraction of inclusions (50%) and the given sieve curve, a 10µm-wide ITZ is a percolated phase. Moreover, stress concentrations occur between close inclusions, that is in the ITZ. It could explain why the effect of the ITZ is larger than expected by the semi-analytical model.

A two-scale homogenization method was defined by merging the layers of the ITZ and the matrix first according to the Reuss bound. Then, the inclusions are introduced through a lower Hashin-Shtrikman bound. Even if this crude method performs slightly better than the $n + 1$-phase spherical model on this mortar, it still underestimates the time dependent strain. The fact that the numerical estimates are softer than the response of this crude model is a clue than the scale of the ITZ and the scale of the inclusions cannot be separated.

2.3. Discussions

There are several assumptions to question:

- Are the inclusions perfectly bonded to the matrix?
- Is viscoelasticity the right frame to model the basic creep of concrete?

The long-term discrepancies between these two concretes [Tab. 1] may also be partly due to the chemical reaction between calcium carbonate $\text{CaCO}_3$ from the limestone filler at Paluel and tricalcium aluminate $\text{C}_3\text{A}$ from the clinker. For the concrete of Paluel, using the model defined by de Larrard [69], the long term
effect of the additional binder on compressive strength is comparable to 30kg/m$^3$ of cement per cubic meter of concrete.

It is well known that the time dependent behavior of concrete goes beyond viscoelasticity. The time-depend strain may be separated in different components: endogenous shrinkage, drying shrinkage, basic creep and drying creep. This divide-and-conquer strategy is the frame of EU standards Eurocode 2 and models of Granger [11], Benboudjema [72], de Larrard [69] and B3 model of Bžant [73] among others. It eases the introduction of moisture content, structural effects and temperature effects. Rossi pledges instead for a unified strategy in which basic creep is a self-drying shrinkage under stress [74]. The assumption is that the creation of microcracks during a creep step (constant load level imposed during the creep test) generates water transfers which induces some additional self-drying shrinkage. This model is consistent with records of acoustic emissions during compressive creep tests [75].

The basic creep of concrete is an ageing phenomenon and the current numerical model could be improved by considering ageing linear viscoelasticity. Moreover, additional eigenstrains due to shrinkage (autogeneous or drying) could be introduced, in order to better estimate the local stresses in the matrix and assess the possible occurrence of damage. The introduction of microcracking, or damage, would also help to explain non-linear creep above $\approx 40\%$ of compressive strength. In [74], stress levels are above $\approx 50\%$ of compressive strength whereas the overall biaxial prestress of containment buildings is below 30% of compressive strength in each direction ($\sigma_{\theta\theta} \approx 12MPa$ and $\sigma_{zz} \approx 8MPa$).
Conclusion

Methods to build various artificial microstructures have been defined and a numerical frame to study the viscoelastic behavior of concrete as a matrix-inclusion material is presented. Numerical results do not differ largely from the Hashin-Shtrikman estimates as long as volume fractions remain below 63% and the aspect ratio of inclusions and the contrast between phases remain low. Either the sieve curve and the inclusion shape do not influence creep or the matrix-inclusion model is too simple to be accurate. Since concretes with close volume fraction of aggregates exhibit very different long term creep, considering such concretes as a two-phase matrix-inclusion materials to study creep is questionable. A heterogeneous cementitious matrix, interfacial zones or decohesion phenomena might explain such differences.

3D numerical simulations may be performed to assess the correctness of micromechanics models. For instance, it has been shown that numerical estimates of creep strains are different from the one obtained by the micromechanics $n + 1$ phase spherical model for modeling the Interfacial Transition Zone in a mortar. The $n + 1$ phase spherical model assumes that the ITZ is an isolated layer of around each inclusion whereas the numerical model accounts for the ITZ being a percolated phase in the mortar.

The gains of 3D numerical simulation are the ability to handle aging behaviors and, to a certain extent, the access to the magnitude of stress concentrations. Large scale computations are required to study concrete or other heterogeneous materials with a large representative volume element, especially if a non-linear phenomenon is introduced at the local scale.

Appendix A. FFT solvers

**Algorithm 1** Basic FFT scheme

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial strain field $\varepsilon^0(x)$ and prestress $\sigma_0(x)$ are provided</td>
<td></td>
</tr>
<tr>
<td>Initial strain field $\tilde{\varepsilon}^{-1}(\xi)$ is set</td>
<td></td>
</tr>
<tr>
<td><strong>while</strong> $\text{error}<em>{eq} &gt; 10^{-7} \times \text{error}</em>{eq,0}$ <strong>do</strong></td>
<td></td>
</tr>
<tr>
<td>$\sigma^i(x) = C(x)\varepsilon^i(x) + \sigma_0(x)$</td>
<td></td>
</tr>
<tr>
<td>$\tilde{\sigma}^i = \text{FFT}(\sigma)$</td>
<td></td>
</tr>
<tr>
<td>Compute error on equilibrium $\text{error}_{eq}$</td>
<td></td>
</tr>
<tr>
<td>$\tilde{\varepsilon}(\xi)^{i+1} = \tilde{\varepsilon}(\xi)^i - \Gamma^0(\xi) : \tilde{\sigma}^i(\xi)$ and $\tilde{\varepsilon}(0)^{i+1} = E$</td>
<td></td>
</tr>
<tr>
<td>$\varepsilon^{i+1} = \text{FFT}^{-1}(\tilde{\varepsilon}^{i+1})$</td>
<td></td>
</tr>
<tr>
<td><strong>end while</strong></td>
<td></td>
</tr>
</tbody>
</table>
**Algorithm 2** Accelerated FFT scheme

Initial strain field $\varepsilon^0(x)$ and prestress $\sigma_0(x)$ are provided

Initial strain field $\hat{\varepsilon}^{-1}(\xi)$ is set

while $error_{eq} > 10^{-7} \times error_{eq,0}$ or $error_{bc} > 10^{-7} \times error_{bc,0}$ or $error_{comp} > 10^{-7}$ do

$\sigma^i(x) = C(x)e^i(x) + \sigma_0(x)$

if $i \% 5 == 0$ then

$\hat{\sigma}^i = FFT(\sigma)$

Compute error on equilibrium $error_{eq}$

end if

$\tau(x) = (C(x) - C_0)e^i(x) + \sigma_0(x)$

$\hat{\tau} = FFT(\tau)$

$\dot{e}_b(\xi) = -2\Gamma_0(\xi) : \hat{\tau}(\xi)$ and $\dot{e}_b(0) = 2E$

$e_b = FFT^{-1}(\dot{e}_b)$

$e^{i+1}(x) = (C(x) + C_0)^{-1} : (\tau(x) + C_0 : e_b(x) - \sigma_0(x))$

Error on boundary conditions: $error_{bc} = \varepsilon^{i+1} - e_b$

Error on compatibility: $error_{comp} = \frac{||e^{i+1} - e_b||_2}{||e^{i+1}||_2}$

end while

---

**Appendix B. Hashin-Shtrikman bound and Laplace-Carson transform**

**Appendix B.1. The correspondence principle**

The non-aging linear viscoelastic problem corresponds to elastic problems thanks to the Laplace-Carson transform. The transform of a function $g(t)$ is $\hat{g}(p) = \int_0^\infty g(t)e^{-pt}dt$. The transform of its derivative $\dot{g}(t)$ is $\hat{\dot{g}}(p) = p\hat{g}(p) - g(0)$.

In our case, elastic inclusions (volume fraction $f_i$) are embedded in a viscoelastic matrix modeled by a single Kelvin chain. The relaxation problem reads:

\[
\begin{align*}
\text{div } \sigma(x,t) &= 0 & x \in V \\
\sigma(x,t) &= C_i \varepsilon(x,t) & x \in \text{inclusions} \\
\sigma(x,t) &= C_m \varepsilon(x,t) + \tau C_m \dot{\varepsilon}(x,t) & x \in \text{matrix} \\
\varepsilon(x,t) &= E(t) + \nabla u(x,t) & x \in V \\
u(x,t) &= \text{periodic} & x \in \partial V \\
\sigma(x,t) \cdot n(x) &= \text{skew} - \text{periodic} & x \in \partial V
\end{align*}
\]

In the Laplace-Carson space, for each $p$, this set of equation corresponds to the elastic problem:
The Hashin-Shtrikman bound is an analytical model which provides estimates of the macroscopic response of the elastic material.

\[ \langle \hat{\sigma} \rangle = \hat{C}_{HS}(p) \langle \hat{\varepsilon} \rangle = \hat{C}_{HS}(p) E \]

\[ K_{HS-}(p) = K_m(p) + \frac{1}{f_i} \left( \frac{3(1-f_i)}{3K_m(p)+4\mu_m(p)} \right) \]

\[ \mu_{HS-}(p) = \mu_m(p) + \frac{1}{f_i} \left( \frac{6(K_m(p)+2\mu_m(p))(1-f_i)}{5\mu_m(p)(3K_m(p)+4\mu_m(p))} \right) \]

The Laplace-Carson transform of the macroscopic stress \( \langle \hat{\sigma} \rangle \) is computed and the last stage is inverting this transform.

Appendix B.2. Inverting the Laplace-Carson transform

Lots of methods are available to invert the Laplace-Carson transform. In this study, the Gaver-Stehfest formula \[62\] has been used:

\[ g(t,M) = \sum_{k=1}^{2M} \frac{\xi_k}{K} \hat{g} \left( \frac{k\ln(2)}{t} \right) \]

and \( \xi_k = (-1)^{M+k} \sum_{j=E \left( \frac{M}{k} \right)}^{\min(k,M)} \frac{j^{M+1}}{M!} \binom{M}{j} \binom{2j}{j} \binom{j}{k-j} \)

Computing the binomial coefficients requires high precision and the long double type (IEEE 754, decimal on 128 bits) provided it. If \( M \) is too low, the formula lacks precision \[63\]. If \( M \) is too high, small errors on \( \hat{g} \left( \frac{k\ln(2)}{t} \right) \) may trigger large errors on the outcome. \( M \) is set to 7.

Therefore, to estimate the response at time \( t \), about 14 elastic computations are required. This formula does not seem to be practical for FEM since it lacks stability or precision. It is suitable as long as the numerical error in the Laplace-Carson space remains very low. This algorithm is useful when the Hashin-Shtrikman analytical formula or self-consistent estimate are computed in the Laplace-Carson space.


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