Drying creep of concrete: constitutive model and new experiments separating its mechanisms

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A new experimental method which allows the direct separation of the components of drying creep due to microcracking and stress-induced shrinkage is developed, demonstrated and validated. The basic idea is to compare the curvature creep of beams subjected to the same bending moment but very different axial forces. The results confirm that drying creep has two different sources: microcracking and stress-induced shrinkage. The latter increases continuously, whereas the former first increases and then decreases. The test results are fitted using a finite element model. The results validate the present model for drying creep. The microcracking is described by an established model, and the free (unrestrained) shrinkage of a material element is shown to depend approximately linearly on the humidity drop.

1. INTRODUCTION

The Pickett effect [1], also called drying creep, is the excess of creep at drying over the sum of shrinkage and basic creep, as shown in Fig. 1. Over the last 50 years, several hypotheses have been presented to explain the mechanisms of this effect [1–8, 32].

Most of the models have tried to explain the excess deformation by the so-called microcracking effect. Because of the non-uniformity of moisture distribution in a shrinkage specimen (specimen No. 1, Fig. 1), the surface layer of the specimen dries and shrinks first, while the inner layer remains wet and does not shrink. As a result, the surface layer is in tension while the inner layer is in compression at the initial stage of the drying process. The tensile stress causes local microcracking or tensile strain-softening in the surface layer, which has been investigated by many authors [2, 9–13]. Due to the nonlinear inelastic behaviour and unrecoverable creep of concrete caused by the tensile stress, the microcracks cannot close fully when the moisture distribution finally approaches a uniform state. As a result, the measured shrinkage of the specimen is always smaller than the true shrinkage of the specimen. It is this irreversible deformation that reduces the total shrinkage of the specimen. On the other hand, for drying creep specimens (specimen No. 3, Fig. 1), the whole cross-section is in compression, and thus there is no microcracking on the planes normal to the direction of compression. So the observed specimen shrinkage does not get reduced by microcracking and the related irreversible deformations. This means that the drying creep must be larger than the sum of the separately observed basic creep and shrinkage.

Partly differing from the others, Bažant and Chern [6, 7] pointed out that microcracking (or tensile strain-softening) and other irreversible strains form only a part of the Pickett effect. Another mechanism exists: stress-induced shrinkage. This mechanism may be explained from a different point of view. Basically, two different types of moisture diffusion process can be distinguished: macrodiffusion (drying and wetting) and microdiffusion. Microdiffusion transports water locally between the capillary pores (macropores) and gel pores (micropores), and this transport affects the deformation rate of the solid framework of cement gel. The movement of water molecules through the gel pores promotes the breakage of bonds which are the source of creep, and thus influences creep [14, 15].

However, there are no experimental data currently in the literature that distinguish clearly among the proposed mechanisms. The occurrence of the latter mechanism has been shown previously rather indirectly by a complete computer analysis of various results of tests [6, 7], and was implied indirectly in the thermodynamic analysis by Bažant [14, 15]. The conventional creep test with centric loading and the bending creep test can show only the
Pickett effect itself, but cannot detect the different mechanisms involved. The first objective of this paper is to present a drying creep experiment which directly distinguishes the aforementioned mechanisms and provides a clear experimental basis for establishing the correct theoretical model. The basic idea of this experiment was presented in 1989 [16], and preliminary reports on the test results were made at two recent conferences [17, 18].

Many mathematical models with diverse mechanism concepts have been proposed for creep, shrinkage, and drying creep of concrete (for review see [19] chap. 2). Among them, Bažant and Chern's model [6-8] has been the only one that attempts to describe both of the aforementioned mechanisms of drying creep. At the time of those studies, however, some aspects were not understood as well as now: for example, the solidification aspect of basic creep and certain consequences of the diffusion theory for shrinkage. Furthermore, the parameters of the model could not have been calibrated accurately due to the absence of experiments that clearly separate the effect of each mechanism.

The second objective of this paper is to establish an experimentally well supported theoretical model for drying creep. For basic creep, the recently developed continuous retardation spectrum will be used to characterize the non-ageing creep component [20], and then the solidification concept [21, 22] will be applied to introduce the ageing aspect of creep. For shrinkage, a new local constitutive relation for relative humidity and shrinkage strain will be formulated, based on recent test results. For stress-induced shrinkage, the microporosity concept of Bažant and Chern [6] will be retained, but the mathematical formulation will be refined and modified. For the microcracking effect, a model that has been used for other similar types of softening behaviour of concrete will be employed. Finally, the parameters of the model will be calibrated by new test results.

2. NEW EXPERIMENTS TO SEPARATE VARIOUS MECHANISMS

2.1 Basic idea

The basic idea is to measure the curvature creep of specimens subjected to the same bending moment but very different axial forces. The advantage of measuring curvature creep is that shrinkage does not contaminate the results because it causes no curvature in a symmetric and symmetrically drying specimen (it is assumed that the stresses and microcracking do not affect the water diffusion significantly).

The desired loadings are axial loads with different eccentricities. Fig. 2 shows type 1 and type 2 specimens with small eccentricity, and Fig. 3 shows type 3 and type 4 specimens with large eccentricity. Type 1 and type 3 specimens are for basic creep, and are therefore sealed on all four sides, whereas type 2 and type 4 specimens are for drying creep and are sealed on two sides (and the ends) to make the moisture diffusion process one-dimensional.

The load magnitudes and eccentricities are chosen so that the bending moments for the two loading cases are exactly the same. Consequently, the deformations due to the two different mechanisms of drying creep can be separated by measuring curvatures.

First, when specimens with small eccentricity are loaded, the whole cross-section is in compression (Fig. 2), which eliminates the tensile microcracking effect. Thus, deformation of the type 1 specimen is due solely to basic creep. Therefore, if any excess deformation is observed on the type 2 specimen, it means that stress-induced shrinkage exists and contributes to the drying creep. Second, when the specimens with large eccentricity are loaded, there is both tension and compression in the cross-section. Deformation of the type 3 specimen (see Fig. 3) is from basic creep only. The irreversible deformation, particularly microcracking, causes excess deformation in the tensile region of the cross-section of the type 4 specimen, and thus extra curvature. Hence, any extra curvature in the type 4 specimen compared with the type 2 represents deformation due to microcracking. This is the basic idea of the present experimental approach.

Keeping the stresses within the service stress range, one can treat the creep as approximately linear. Thus, the bending curvatures observed can be characterized in terms of the compliance function \( J(t, \tau) \). This function can then be decomposed into the terms corresponding to the two different mechanisms.
2.2 Experimental procedure

The experimental setup, shown in Fig. 4, was the standard setup recommended by ASTM. The specimens were square prisms of length 16 in (40.64 cm) and side 4 in (10.16 cm). For eccentric loading, a one-way hinge at the top of the specimens and a two-way hinge at the bottom were used. In this way, the load axis could be kept at the centre of the cross-section in one horizontal direction and off-centre in the perpendicular horizontal direction.

Lateral deflections were measured on both sides during the test, and then averaged. Each time, before taking a reading, the load was adjusted to a constant load value. The experiments were carried out in a moisture room. The environmental relative humidity was controlled at 50%, and the temperature was held constant at 80°F (26.7°C).

Two sets of tests have been carried out. In set I, there were two different eccentricities, small and large, and two different moisture conditions, drying and sealed. For each loading and moisture condition, there were two identical specimens, with a total of eight specimens in set I. The lateral deflection measurement in set I was designed to be used to verify different mechanisms. Therefore, all specimens in set I were subjected to the same bending moment and were cast from the same batch of concrete.

Set II involved four specimens cast from another batch. There were two moisture conditions but only one eccentricity – small. For each loading and moisture condition, two identical specimens were cast. To check the drying creep mechanisms at different loading levels, the bending moment in set II was designed to be different from that in set I. Further information on the two sets is listed in Table 1.

All the specimens used in the two sets were cast identically, in a horizontal position, and cured identically. The mix ratios were water:cement:sand:gravel = 0.5:1.0:2.5:3.0 (by weight). Type I cement was used. The aggregate was crushed limestone with a maximum size of 3/4 in (1.9 cm). The forms were removed after one day of moist curing at room temperature, and the sides of each specimen were sealed with siliconized acrylic latex caulk.

2.3. Test results and discussions

Fig. 5 shows the averages of the lateral deflections for each pair of identical specimens. First, by comparing the deflections of the sealed and unsealed specimens with the small eccentricity (solid squares), it is noticeable that there is an additional deformation, added to basic creep. Since, with small eccentricities, the whole cross-section is in compression, microcracking cannot occur. So the additional deformation that is seen in Fig. 5 must come from the stress-induced shrinkage. This is a direct experimental proof of the existence of stress-induced shrinkage.

Furthermore, comparing the sealed and unsealed specimens with large eccentricity (the circles in Fig. 5), it is seen that there is also an additional deformation, added to basic creep. This extra deformation combines the contributions from the two mechanisms, microcracking and stress-induced shrinkage.

<table>
<thead>
<tr>
<th>Set I</th>
<th>Set II</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Small $e_0$</td>
</tr>
<tr>
<td>$e_0$ (in)</td>
<td>0.21</td>
</tr>
<tr>
<td>No. spec.</td>
<td>4</td>
</tr>
<tr>
<td>$f'_1$ (psi)</td>
<td>5619</td>
</tr>
<tr>
<td>Size (in)</td>
<td>$4 \times 4 \times 16$</td>
</tr>
<tr>
<td>$P$ (lb)</td>
<td>26971</td>
</tr>
<tr>
<td>$\sigma_{\text{max}}$ (psi)</td>
<td>2248</td>
</tr>
<tr>
<td>$\sigma_{\text{min}}$ (psi)</td>
<td>1124</td>
</tr>
</tbody>
</table>

Fig. 4 Experimental setup for eccentric loading.

Fig. 5 Averaged deflections.
The deflection magnitude depends on many parameters, such as the prism length, cross-section size, and eccentricity. To compare test results, the compliance function can be employed. The compliance function $J$ can be calculated easily from the measured mid-span lateral deflection $y_{\text{max}}$. According to the theory of elastic buckling [23], chap. 1,

$$J = \frac{1}{E} = \frac{1}{P} \left[ \frac{2}{l} \arccos \left( \frac{e_0}{y_{\text{max}} + e_0} \right) \right]^2,$$

where $l$ is the centroidal inertia moment of the cross-section, $P$ is the axial load, $l$ is the length of the column, and $e_0$ is the initial eccentricity. For small $P$ and small $y_{\text{max}}$, Equation 1 may be approximated as

$$J = \frac{1}{E} \approx \frac{8l}{l^2 P e_0} y_{\text{max}}.$$

This eliminates from the creep observations the effects of $P$, $e_0$, $l$, and $I$. Therefore, the test results for different loading moments, column lengths, and cross-sections can be compared with each other in terms of $J$. This is exactly the same as in the analysis of centric loading tests. In that case, the axial strain $e_t$ is measured during the test. Instead of $e_t$, the compliance function $J = e_t A / P$ ($A$ is the area of the cross-section) is analysed to get rid of the effects of $A$ and $P$.

The averaged compliance function, computed from Equation 1, is shown in Fig. 6. One can see that, for identical specimens, loading and moisture conditions, the analysis of compliance function is equivalent to the analysis of lateral deflection and will give the same conclusions as in Fig. 5.

The different mechanisms shown in Fig. 6 can be decomposed as shown in Fig. 7. It is evident that the stress-induced shrinkage increases with time. The microcracking effect can be decomposed further, as shown in Fig. 8. It appears that the microcracking effect increases from the very beginning, reaches its maximum (for this cross-section size) in about 10 days, and then decreases. This is because moisture distribution gradually approaches a uniform state. However, even after a long time, the microcracks that have developed in the early drying process close only partly, which must be explained by irreversibility of the crack openings.

Fig. 9 shows the averaged compliance function obtained from the results of set II. Comparison with Fig. 6 shows clearly that there is also another deformation, added to the basic creep. Since the whole cross-section is also in compression for set II, there is no microcracking effect. The observed additional deformation must come from the mechanism of stress-induced shrinkage. Noting that the bending moments and axial loads in sets I and I are different, we further conclude that the drying creep mechanisms, distinguished from each other by the experiment in set I, truly occur for various loading levels.
3. CONSTITUTIVE MODEL

3.1 Review of basic creep model with ageing

The ageing aspect of creep is handled by solidification theory. The key feature of the theory (Fig. 10) is that the ageing aspect of concrete creep is considered to be caused by the growth of the volume fraction \( \eta(t) \) of the effective load-bearing portion of solidified matter (i.e., hydrated cement), representing an increase of both the volume fraction of the hydrated cement and the load-bearing solid fraction, which is growing due to the formation of further bonds (or polymerization of calcium silicate hydrates). The principal advantage is that, in this theory, the properties of all the constituents, including the load-bearing matter, are age-independent (as required by thermodynamics). This makes it possible to apply the conventional theory of non-ageing viscoelasticity.

The creep strain rate \( \dot{\varepsilon}_c \), corresponding to the solid part, can be expressed as the product of the age-independent strain rate of solid \( \dot{\gamma} \), and the increase of the volume fraction \( \eta(t) \) of the solid [21, 22]:

\[
\dot{\varepsilon}_c(t) = \frac{F[\sigma(t)]}{\eta(t)} \dot{\gamma}(t),
\]

where function \( F[\sigma(t)] \) is introduced to reflect nonlinear behaviour at high stress (at low stress, \( F[\sigma(t)] = 1 \)).

In this formulation, all of the procedures developed for non-ageing basic creep are applicable to the viscoelastic microstrain \( \dot{\gamma} \). Applying the well-known Kelvin chain model with \( N \) Kelvin units, we then have, for a constant uniaxial stress \( \sigma \) [20],

\[
\dot{\gamma} = \left[ \sum_{n=1}^{N} A_n (1 - e^{-\frac{t}{\tau_n}}) \right] \sigma
\]

where \( \xi \) is the load duration. Analogously to Equation 3, we have for the flow term:

\[
\dot{\varepsilon}_f(t) = \dot{\varepsilon}_c(t) = \frac{F[\sigma(t)]}{\eta(t)} \sigma(t),
\]

where \( q_3 \) is an empirical coefficient which depends on the composition of concrete.

One important advantage of the solidification theory is that, when the age-independent creep strain \( \dot{\gamma} \) is modelled by a generalized Kelvin chain, \( A_n \) in Equation 4 can be determined [20] by the method of continuous retardation spectrum \( L(\tau_n) \). If \( \tau_n \) is known, then, approximately (but with very good accuracy)

\[
A_n = L(\tau_n) \ln(10) \Delta(\log \tau_n),
\]

where \( \Delta(\log \tau_n) \) is the time interval between the retardation times of two adjacent Kelvin units in the logarithmic time scale. Computational experience shows that the choice \( \Delta(\log \tau_n) = \log 10 = 1 \) gives a sufficiently smooth compliance function, while greater separations of \( \tau_n \) give a bumpy appearance to the creep curves. \( L(\tau_n) \) depends on the compliance function of the concrete. For the compliance function of the non-ageing constituent, we may assume the log-power creep law:

\[
J(\xi) = q_2 \ln \left[ 1 + \left( \frac{\xi}{\xi_0} \right)^n \right]
\]
in which \( \lambda_0 = 1 \text{ day} \); and \( q_2 \) is a parameter depending on the concrete composition.

The method for obtaining \( A_0 \), analogously to the method used in classical non-ageing viscoelasticity, has been presented by Bazant and Xi [20] and will now be outlined briefly. For Equation 7, it has been shown that

\[
L(\tau) = \left[ \frac{-2n^2(3\tau)^{2n-3}[n - 1 - (3\tau)^n]}{[1 + (3\tau)^n]^3} \right. \\
+ \left. \frac{n(n - 2)(3\tau)^{n-3}[n - 1 - (3\tau)^n] - n^2(3\tau)^{2n-3}}{[1 + (3\tau)^n]^3} \right] \\
\times q_2^*(3\tau)^3 \frac{1}{2}.
\]  

(8)

This is the approximate retardation spectrum of order 3. According to the fitting of test data by Bazant and Prasannan [21, 22], \( n \) is approximately a constant. In the case of small \( n \), the terms with \( n^3 \) in Equation 8 may be neglected without much loss of accuracy. Combining this with some other simplifications of Equation 8, the following expression for the retardation spectrum can be obtained:

\[
L(\tau) = q_2^*n(1 - n) \frac{(3\tau)^n}{1 + (3\tau)^n}.
\]  

(9)

Because of its simple form, Equation 9 is attractive as means of creep characterization that could be used in a design code. For computational analysis, such as finite element analysis, or when \( n \) is large (\( n > 0.45 \)), Equation 8 ought to be used. To analyse the present tests, eight units of Kelvin chain have been used. To be able to cover the test range by a minimum number of Kelvin units, \( \tau_\nu \) are spaced by decades in \( \log \tau_\nu \), i.e., \( \tau_1 = 10^{-3} \) and \( \tau_{\nu+1} = 10\tau_\nu \), \( \mu = 1, 2, \ldots \). \( A_0 \) depends on \( n \) as given by Table 2, and \( \tau_0 = 10^{-7} \); \( A_0 \) is given by Equation 6. The rate-type formulation used in finite element analysis is presented in detail by Bazant and Xi [20] and is in principle the same as given by Bazant and Prasannan [21, 22].

### 3.2 Shrinkage model: relation of local shrinkage to relative humidity

In the idealized case of free shrinkage, in which there is no restraint and no stress and the whole specimen shrinks uniformly, the rate of local shrinkage strain is [6–8]

\[
\dot{\varepsilon}_{sh} = k_{sh} \dot{H}
\]  

(10)

\[
k_{sh} = -\varepsilon_0^* g_s(t) \frac{d f_s(H)}{dH}
\]  

(11)

where \( k_{sh} \) is the shrinkage coefficient; \( \varepsilon_0^* \) is the ultimate shrinkage, \( g_s(t) = E(t_0)/(E(t)) \), and \( E(t) \) can be estimated by

\[
E(t) = E(28) \sqrt{\frac{t}{4 + 0.85t}}
\]

with

\[
E(28) = 33w^{1.5} \sqrt{f'_c}
\]  

(12)

where \( w \) is the unit weight of concrete in lb ft\(^{-3} \), and \( f'_c \) and \( E \) are in psi. For \( t \to \infty \), \( g_s(t) \to 1 \).

Now the important question is the meaning of \( f_s(H) \). This is a complicated problem because, until now, there have been no experimental results to identify clearly the function \( f_s(H) \). As an approximation, the following equation has been used [19]:

\[
f_s(H) = 1 - H^3
\]  

(13)

It was obtained by fitting a large amount of test data on shrinkage deformations averaged over the whole
cross-section for different ambient relative humidities [24], and was intended only for such deformations which include the effects of distributed cracking, and creep caused by shrinkage stresses. As a local constitutive relation for a material point, however, Equation 13 does not give best results.

Fig. 11 shows finite element results for the shrinkage strain according to Equation 13. The humidity profile (Fig. 11, upper right) indicates that the outside layer of the specimen has a larger humidity drop. However, calculations based on Equation 13 indicate that the inner layer could shrink more than the outside layer, which is not correct.

Another function, however, is suggested by noting that shrinkage of small specimens has been found to be proportional to the water loss $\Delta \varepsilon$, and according to the desorption isotherms, $\Delta \varepsilon$ is often approximately proportional to the humidity drop. According to this argument, we will use

$$f_s(H) = 1 - H.$$  

(14)

Indeed, good finite element results for shrinkage have been obtained with this relation (see Fig. 12). By substituting Equation 14 into 11 and then into 10, we obtain

$$\Delta \varepsilon_{\text{sh}} = k_{\text{sh}} \Delta H, \quad k_{\text{sh}} = \varepsilon_0^s g_s(t)$$  

(15)

$\varepsilon_0^s$ in Equation 15 is a material constant representing the magnitude of the final shrinkage value. It is left as a free empirical constant in our model, because of the difficulties in measuring $\varepsilon_0^s$ directly.

### 3.3 Formulation for stress-induced shrinkage

The Kelvin and Maxwell chains can both approximate linear creep behaviour with any desired accuracy [19]. For the sake of a theoretical argument, it is now simpler to use (temporarily) the Maxwell chain to analyse the stress-induced shrinkage. The same result could be obtained using the Kelvin model, but the derivation would be less clear because the Kelvin units involve not only the strain rate but also the strain itself. The equation for a single Maxwell unit is

$$\frac{\dot{\sigma}}{E} + \frac{\sigma}{\eta} = \dot{\varepsilon} - \dot{\varepsilon}_{\text{sh}} - \dot{\varepsilon}_T$$  

(16)

where $\eta$ is the dashpot viscosity; $\dot{\varepsilon}_{\text{sh}}$ and $\dot{\varepsilon}_T$ are the shrinkage strain rate and thermal strain rate; and $\dot{\varepsilon}_T = a_T \dot{T}$, where $a_T$ = coefficient of thermal expansion.

Because microdiffusion of pore water may be expected to enhance the debonding and reboning process in cement gel that is the source of creep, the viscosity $\eta$ should be a function of the microdiffusion flux $\dot{H}$ of water which is driven by a difference in the chemical potential between the capillary water and the gel pore water. At variable temperature $T$, this difference, in turn, is proportional (as shown by Bažant and Chern [6, 7] to effective pore humidity

$$\dot{H} = a_1 \dot{H} + a_2 \dot{T}$$  

(17)

in which $a_1$ and $a_2$ are constants. Note that (1) the microdiffusion flux increases creep and thus decreases viscosity $\eta$, and (2) only the absolute value of $\dot{H}$ matters since the creep increases regardless of whether the microdiffusion of water goes into or out of the capillary pores. Thus, as a first-order approximation,

$$\frac{1}{\eta} = \frac{1}{\eta_0} + k' |\dot{H}|$$  

(18)

where $\eta$ is the viscosity with no microdiffusion, and $k'$ is a constant. Substituting Equations 17 and 18 into 16,

$$\frac{\dot{\sigma}}{E} + \frac{\sigma}{\eta} = \dot{\varepsilon} - k \dot{H} - \dot{\varepsilon}_T$$  

(19)

where

$$k = k_{\text{sh}}(1 + r \sigma \text{ sign}(\dot{H}))$$  

(20)

$$\dot{\varepsilon}_T = a_T(1 + \rho \sigma \text{ sign}(\dot{H}))$$  

(21)

where $r = k'a_1/k_{\text{sh}}$, $\rho = k'a_2/a_T$, and sign($\dot{H}$) is the sign of $\dot{H}$.
One can see that the total shrinkage coefficient $k$ (Equation 20) is stress dependent. That is why the associated deformation is called the stress-induced shrinkage. Furthermore, Equation 21 reveals that the total coefficient of thermal expansion $\tilde{z}$ is also stress dependent, as proposed already by Thelandersson [25, 26]. The corresponding increase of thermal deformation, termed the stress-induced thermal expansion, has been detected by some experiments [25, 26].

For three-dimensional generalization, one can apply the same derivation to the volumetric and deviatoric components separately, and then sum them up. One obtains

$$\Delta\epsilon_{\text{sh,ij}} = k_{ij} \Delta H, \quad \Delta\epsilon_{\text{T,ij}} = a_{ij} \Delta T \quad (22)$$

in which

$$k_{ij} = k_{\text{sh}} [\delta_{ij} + (\rho \sigma_{ij} + r' \sigma') \text{sign}(\dot{H})], \quad a_{ij} = a_{T} [\delta_{ij} + (\rho \sigma_{ij} + r' \sigma') \text{sign}(\dot{H})] \quad (23)$$

where $\sigma' = \sigma_{kk}/3$, $\delta_{ij}$ is the Kronecker delta, $r$ and $r'$ are constants for shrinkage, and $\rho$ and $\rho'$ are constants for thermal expansion.

Equations 22 and 23 for shrinkage and thermal expansion have been used in computations, along with $k_{\text{sh}}$ from Equation 15.

3.4 Stress–strain relation for distributed microcracking

The microcracking that occurs in the tensile region of the cross-section under service loads has different characteristics from cracking and fracture under ultimate load. The microcrack density is quite low [9]. One has the so-called ‘dilute’ cracks, whose effect may be described in a continuum manner by a stress–strain relation. In the previous work, this was done by combining the uniaxial stress–strain relations for three principal directions in an isotropic model. The calculations for this paper have been performed with a different type of constitutive relation for dilute cracks, developed by Brockenbrough and Suresh [27], in which the microcracking damage is also assumed to preserve isotropy. Although this constitutive relation cannot be said to give distinctly more realistic results, it has a better theoretical foundation as it is to some extent based on the micromechanics of dilute microcracks. For the present problem, the stress–strain relations used by Bažant and Chern [6, 7] would nevertheless yield almost the same results.

As in some previous damage models, we will introduce the central idea that the elastic modulus of a microcracked material decreases gradually as the number of microcracks increases or the existing microcracks grow. For this purpose, different kinds of crack geometries and distributions have been analysed extensively, for example, randomly orientated penny-shaped cracks, elliptical cracks, slit cracks, aligned penny-shaped cracks, and aligned slit cracks [28, 29]. Based on those studies, the reduction of material stiffness can be expressed approximately as a function of a single parameter, called the crack density $f_\beta$: $E/E_0 = 1 - f_\beta(\beta)$, in which $E_0$ is the initial elastic modulus and $v_0$ is the initial Poisson’s ratio. A linear approximation is found acceptable for $f_\beta(\beta)$

$$\frac{E}{E_0} = \frac{v}{v_0} = 1 - \frac{16}{9} \beta \quad (24)$$

Anyhow, the exact form of $f(\beta)$ can hardly be determined because the crack system inside concrete does not involve a single crack geometry and the aggregates, acting as inclusions, also exert influence. We use

$$\beta = \begin{cases} \frac{A_\beta (\tilde{\sigma} - 1)^{n_\beta}}{\sigma_0} & |\tilde{\sigma}| > \sigma_0 \\ 0 & \text{elsewhere} \end{cases} \quad (25)$$

in which $\tilde{\sigma} = (\sigma_1^2 + \sigma_2^2 + \sigma_3^2)^{1/2}$; $A_\beta$, $n_\beta$, and $\sigma_0$ are constants; and $\sigma_\beta$ is a threshold stress for microcracking. A further parameter $\beta_\text{c}$, called the saturation crack density, has also been introduced in some models [27, 30].

To stimulate the irreversible deformation at unloading, an unloading parameter is introduced;

$$\lambda = 1 - \frac{\beta_\text{c}}{\beta_\text{max}} \quad (26)$$

$\beta_\text{c}$ is the closed crack density at unloading, and $\beta_\text{max}$ is the maximum crack density attained before unloading. It is required that $0 < \lambda < 1$. Then the unloading secant modulus $E_u$ is interpolated between $E_0$ and $E_\text{min}$ (the elastic modulus at the start of unloading), and the same is done for Poisson’s ratio $\nu_u = \nu_0 \lambda + \nu_\text{min} (1 - \lambda) \quad (27)$

The constitutive relations are

$$\sigma = C : \varepsilon \quad (28)$$

$$\tilde{\sigma} = \tilde{C} \tilde{\varepsilon} + \tilde{C_\text{e}} = \frac{\partial \tilde{C}}{\partial E} \frac{\partial E}{\partial \tilde{\varepsilon}} \tilde{\varepsilon} + \tilde{C_\text{e}} \quad (29)$$

From Equations 24 and 25,

$$\frac{\partial E}{\partial \beta} = -\frac{16}{9} E_0 \quad (30)$$

$$\frac{\partial \tilde{\beta}}{\partial \tilde{t}} = A_\beta n_\beta \left( \frac{\tilde{\sigma}}{\sigma_0} - 1 \right)^{n_\beta - 1} \frac{\sigma_0 - \tilde{\sigma}}{\sigma_0} \quad (31)$$

Then the incremental stress–strain relation for elastic deformations with cracking (but excluding creep and shrinkage) is

$$\Delta \sigma = C (\Delta \varepsilon - \Delta \varepsilon_\text{c}) \quad (32)$$

with

$$\Delta \varepsilon_\text{c} = -C^{-1} \frac{\partial C}{\partial \tilde{\varepsilon}} \frac{\partial E}{\partial \tilde{\varepsilon}} \frac{\partial \tilde{\beta}}{\partial \tilde{t}} \tilde{\varepsilon} \quad (33)$$

The unloading criterion is the sign of the increment of crack density $\Delta \beta$. $\Delta \beta < 0$ means that the crack density should decrease, which corresponds to unloading.
Equation 31 indicates that $\Delta \beta < 0$ is equivalent to $\sigma \cdot \Delta \sigma < 0$. At unloading, Equation 27 is used. For reloading ($\lambda > 0$ and $\Delta \beta > 0$), again Equation 27 is used.

3.5 Constitutive model for drying creep and comparison with test data

The experimental results shown in section 2 suggest that the deformations from each mechanism are additive. So the models for each mechanism may also be superposed to describe drying creep. Based upon such considerations, we use a series coupling model (Fig. 13). The basic creep is represented by a generalized Kelvin chain model, Equations 3–5. The free shrinkage and stress-induced shrinkage are from Equations 22 and 23; for a constant temperature field, the thermal strain vanishes; the microcracking model is from Equation 32. The total strain increment is

$$\Delta \varepsilon = \Delta \varepsilon_c + \Delta \varepsilon_s + \Delta \varepsilon_{sh} \quad (34)$$

where $\Delta \varepsilon_c$ is the strain increment representing the creep model, $\Delta \varepsilon_s$ is the strain increment due to microcracking, and $\Delta \varepsilon_{sh}$ is the strain increment representing shrinkage and stress-induced shrinkage:

$$\Delta \varepsilon_c = D_c^{-1} \Delta \sigma + \Delta \varepsilon_s^c, \quad \Delta \varepsilon_s = C^{-1} \Delta \sigma + \Delta \varepsilon_s^C \quad (35)$$

where $D_c$ is the material stiffness matrix and $\Delta \varepsilon_s^C$ are the inelastic strain increments [18]. By substituting Equations 35 and 22 into 34, we obtain the incremental constitutive relation for the total deformation:

$$\Delta \sigma = D(\Delta \varepsilon - \Delta \varepsilon^s) \quad (36)$$

in which

$$D = (D_c^{-1} + C^{-1})^{-1} \quad (37)$$

$$\Delta \varepsilon^s = \Delta \varepsilon_s^C + \Delta \varepsilon_s^C + \Delta \varepsilon_{sh} \quad (38)$$

Before calibrating all the parameters in the model, it is helpful to analyse the effect of each parameter in order to reduce the number of parameters involved to the fewest possible.

First, in the model of basic creep, there are four parameters to be calibrated by tests, $E_o$ (1/$E_o = q_1$), $n$, $q_2$, and $q_3$. Among these, $q_3$ can be neglected for the present purpose as long as the time ranges of the Kelvin units in the model cover the whole experimental duration; indeed, eight units are used in the present computations, with retardation half-times ranging from $10^{-4}$ to $10^3$. $E_o$ characterizes the asymptotic instantaneous deformation: Fig. 5 shows that the scatter of the instantaneous deformation is not large, and the computational results also indicate that the optimized $E_o$ will be slightly different for each deflection curve. Therefore, $E_o$ can be calculated by ACI approximate formula $E_o = 57000 \sqrt{f'_c}$ (with $E_o$ and $f'_c$ in psi, 1 psi $\approx 6895$ Pa), which gives quite good results. Thus, only $n$ and $q_2$ need to be calibrated.

Second, in the shrinkage model there are three parameters, $\varepsilon_0^n$, $r$, and $r'$. Computational experience shows that $r' = 0$, that is, the mean stress has no effect on the stress-induced shrinkage. So, only two parameters, $\varepsilon_0^n$ and $r$, remain.

Finally, in the microcracking model, there are five parameters, $A_p$, $n_p$, $\sigma_0$, $\beta$, and $\lambda$. From Equation 24 one can see that $\beta_p < 9/16$ (otherwise $E$ will decrease to 0); $\beta$ is taken as 0.4 (same as references [27] and [30]). The experimental results show that the unloading stiffness is lower than $E_o$ but higher than the stiffness just before unloading. So, $\lambda = 0.5$ is used, which implies that $E_c$ is the average of $E_0$ and $E_{min}$; $n_p$ is the degree of nonlinearity as indicated in Equation 25, and $n_p = 0$ corresponds to linear elastic behaviour. A large value of $n$, say 10, results in elastic-plastic behaviour, and a smooth transition can be obtained by $n_p = 1.5$ according to the one-dimensional analysis of the model. Consequently, only two parameters, $A_p$ and $\sigma_0$ are known. In total, 6 parameters remain to be found. Fortunately, they can be calibrated from the present test results sequentially.

The finite element mesh used is shown in Fig. 14. The finite elements were four-node quadrilaterals, 0.5 in by 0.5 in. The deformations for basic creep (sealed specimen, small eccentricity) are computed first and matched to the test results to identify the creep parameters $n$ and $q_3$ (see Fig. 15). Then, the test data on basic creep plus stress-induced shrinkage (drying specimens, small eccentricity) are analysed to determine the shrinkage parameters $\varepsilon_0^n$ and $r$; the result is shown in Fig. 16, where the values of $n$ and $q_3$ slightly differ from those in Fig. 15. Fig. 17 shows the deformed mesh of the drying specimen. Two-way spring elements are considered to be inserted in the interfaces between the end steel plate
elements and the concrete elements, to allow the concrete elements to slide. Finally, the test data exhibiting all the mechanisms (drying specimens, large eccentricity) are analysed to evaluate parameters $A_\beta$ and $\sigma_0^r$; the results are shown in Fig. 18.

In simulating the creep tests, the load is first increased from zero to the creep test level and then is held constant for the rest of the computation. For the drying specimens, the relative humidity profile is computed at each time step using the nonlinear diffusion model of Bažant, Xi and Molina [31]. The length of the time steps used for the nonlinear diffusion process is increased such that $t_{i+1}/t_i = 1.003$. The exponential algorithm is used for the time steps in the drying creep analysis. At each step, after the given convergence criterion is met, all of the internal variables are evaluated and stored for all stress components, at all integration points and all output points. The iterations at structure level are carried out according to the Newton Raphson method. Thus, the strain increments accumulated during the iterations and the stress components in the previous time step need to be

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Fig. 14 Finite element mesh for analysis.

Fig. 15 Comparison with basic creep data.

Fig. 16 Comparison with test data of basic creep and stress induced shrinkage.

Fig. 17 Deformed mesh for drying creep.

Fig. 18 Comparison with drying creep test data.
stored to be available for the iterations at the element level, because the viscoelastic creep increments and shrinkage strain increments are computed between two time steps.

To obtain the optimum values of the model parameters, a nonlinear optimization program (based on the Levenberg–Marquardt algorithm) is used. The nonlinear finite element program is arranged as a subroutine for this optimization program.

Figs 15, 16, and 18 indicate that the present model simulates drying creep behaviour well. The optimized values of the parameters for the present test results are given in the figures. In terms of the order of magnitudes, \( n = 10^{-1}, \quad q_2 = 10^{-8}, \quad c_0 = 10^{-3}, \quad r = 10^{-5}, \quad A_p = 10^{-3}, \) and \( \sigma_0 = 10^4 \text{ psi}. \) The orders of magnitude of these parameters are of more general interest than their precise values because, for different cement type, w/c, g/c, and s/c, the values of these parameters may change but not the order of magnitude (for example, \( c_0 \) may vary from 0.0005 to 0.005).

4. CONCLUSIONS

1. A new experimental method, which allows direct separation of the components of drying creep due to microcracking and to stress-induced shrinkage, has been developed, demonstrated, and validated. The basic idea [16] is to compare the curvature creep of beams subjected to the same bending moment but very different axial forces. Such loading is achieved easily by applying axial forces of very different eccentricities and magnitudes.

2. Decomposition of the measured lateral deflections of eccentrically loaded prisms confirms the existence of the mechanism of stress-induced shrinkage. It follows that the deformations observed in drying creep test results from four mechanisms: free shrinkage, basic creep (with elastic strain), stress-induced shrinkage, and microcracking.

3. The test results for different load levels indicate that the stress-induced shrinkage contributes to the total deformation at all load levels.

4. The stress-induced shrinkage monotonically increases with time. The microcracking effect increases at the beginning of drying, reaches a maximum (for the present specimen thickness) at about \( t_m = 10 \text{ days} \), and then it starts to decrease (the value of \( t_m \) may be expected to be approximately proportional to thickness squared).

5. It is possible and advantageous to use for the non-ageing viscoelastic constituent of concrete a continuous retardation spectrum associated with the Kelvin chain model.

6. A linear local constitutive relation between the relative humidity and the free (unrestrained) shrinkage strain of a material element is proposed and validated.

7. The coefficient of stress-induced shrinkage is derived on the basis of the microdiffusion theory for pore water. This coefficient reflects the fact that the associated viscosity is decreased by the microdiffusion process regardless of whether the microdiffusion transports water into or out of the gel pores.

8. The microcracking model of Brockenhaupt and Suresh [27] is applied and is found to characterize nonlinear behaviour and irreversible deformation at unloading quite well.

9. To describe the drying creep tests, the models for basic creep, stress-induced shrinkage, and microcracking are coupled serially. The combined model characterizes each mechanism and the overall drying creep satisfactorily. Optimized model parameters are obtained, and the proper orders of magnitude of these parameters are given.

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REFERENCES


**RESUME**

**Flûage au séchage du béton: Modèle constitutif et nouveaux essais pour identifier ses mécanismes**

On a développé, démontré et validé une nouvelle méthode expérimentale permettant d’identifier directement les composantes du flûage au séchage dû à la microfissuration et à la contrainte de retrait. L’idée de base est de comparer le flûage en flexion de poutres soumises au même moment fléchissant mais de faire varier les différentes forces axiales. Les résultats confirment que le flûage au séchage a deux origines différentes: la microfissuration et le retrait sous contrainte. Celui-ci augmente de façon continue alors que celle-là augmente d’abord, puis diminue. On a utilisé un modèle d’élément fini pour ajuster les résultats d’essai qui valident le modèle actuel pour le flûage au séchage. On décrit la microfissuration par un modèle établi; quant au retrait libre (non empêché) d’un élément de matériau, on montre qu’il dépend de la chute d’humidité de façon approximativement linéaire.