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Update of the HTSI method : application to the characterization of mechanical properties of CaF₂ from RT to 800 °C

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Abstract

The High-Temperature Scanning Indentation (HTSI) method [1] allows for the characterization of material mechanical properties quasi-continuously over a large temperature range in 1-day experiments, based on a cycle with a constant maximum load applied regardless of the temperature. For materials exhibiting an Indentation Size Effect (ISE), the variations in hardness with temperature can stem from both temperature and ISE. It is challenging to differentiate their impact on the mechanical properties. So, a new 1-second indentation cycle was implemented, with an adjustment of the applied maximum load to control the maximum achieved depth across temperatures. It allows for the determination of mechanical properties at a given maximum depth over a wide temperature range.

This methodology has been applied to CaF₂ single-crystal from RT to 800 °C. It enables the characterization of this material at 1000 nm depth over the entire temperature range.. The obtained results are consistent with conventional indentation results.

Keyword nano-indentation; Ca; F; crystal; extreme environment; hardness; in situ; nanoscale

Highlights

- A new 1-second cycle allowing maximum-depth control throughout thermal cycling for the HTSI method has been implemented
- This methodology was applied on CaF₂, known to present a ISE in temperature
- Mechanical and creep properties of CaF₂ were determined from RT to 800 °C

I Introduction

Calcium fluoride is an ionic metal used in various industries for its properties in temperature (lubrication [2–5], optics [6–9], etc.). It has been characterized at high temperature [10] but since it is quite brittle at room temperature [11], characterizing its mechanical properties from room temperature to 800 °C is quite complicated.

Nanoindentation testing allows the characterization of small samples at room temperature [12, 13]. The classical Constant Stiffness Measurement method (CSM) provides properties along the tested depth at a given strain rate [14]. Combining these tests with long-term indentation creep tests [15, 16] as well as relaxation tests [16, 17] now gives access to mechanical properties over 8 orders of magnitude in strain rate.

With the recent development of high-temperature nanoindentation devices [18, 19], nanoindentation testing can be carried out up to 1000 °C. Creep properties can then be determined at various temperatures using the methods implemented for room temperature testing. However, classical CSM, creep, and relaxation indentation tests require precise control of thermal equilibrium during the tests

[17]. Carrying out such tests at high temperatures remains a challenge to overcome, and is still time-consuming.

The High-Temperature Scanning Indentation (HTSI) method [1] allows performing 1-second indentation tests while heating the sample, providing access to mechanical properties over the entire studied temperature range. The idea is that performing 1-second indentation tests during slow heating (a few °C/min) greatly reduces the impact of thermal drift. The indentation cycle is programmed in load-control, meaning the same maximum load is applied regardless of the temperature. Moreover, because of the short duration of the indentation cycle, CSM cannot be easily implemented during loading [20–22]. Thus, mechanical properties are only determined at the maximum depth point. As hardness is expected to decrease with temperature, the tested volume increases significantly with temperature. However, to better understand the source of variations observed during testing, it is preferred to test the same volume of material. Moreover, some materials exhibit an Indentation Size Effect (ISE), where their mechanical properties change with depth [23]. Changing both temperature and testing depth does not allow distinguishing the impact of each phenomenon on changes in behavior. So is there a way to perform indentation tests to characterize CaF₂ mechanical properties from RT to 800 °C without impact of the ISE?

In the present work, we propose an updated version of the indentation cycle used in [1] to address some of the previously identified problems. A new strategy is presented, allowing for a change from maximum load-controlled to maximum depth-controlled indentation cycle. Then, this strategy is applied to characterize the mechanical (creep and hardness) properties of CaF₂ single crystals. Finally, discussion on the proposed method and the results is carried out to identify their limitations.

II Updates of the High-Temperature Scanning Indentation Technique

II.1 Preparation of the thermal cycle

As already presented in [1, 24], the HTSI method consists of applying multiple short indentation tests along a specific thermal cycle. To ensure thermal equilibrium throughout the tests, calibration steps are required. First, it is assumed that since the tip is quite small (1 mm to 2 mm long) and usually made of conductive materials [18], the temperature of the contact point is the same as the one measured at the tip backside (see Figure S1). So one should start by defining the thermal cycle to apply to the tip. Then, temperature calibration steps are performed following the method proposed by Minnert et al. [19]. For a given temperature on the tip side, one should determine the temperature to apply at the back of the sample to achieve thermal equilibrium at the contact point. We recommend performing such calibration steps at least at two temperatures in the studied temperature range. From experiments, if the maximum studied temperature is lower than approximately 700 °C, the difference between the settings is believed to evolve quasi linearly, as radiation is not expected to play a major role in the heating. Once the settings have been correctly determined, the thermal cycle to be applied on the sample side can be constructed. It should be noted that the heating rate on the tip and sample may be different since the maximum temperatures to be achieved are usually different. The main difficulty here is to have extremely precise control of the temperatures during the experiments. A schematic representation of the preparation of the thermal cycle is presented in Figure S1 in the Supplementary Materials.

II.2 Updates of the indentation cycle

II.2.a From quarter-sinus to half-sinus loading

The indentation cycle proposed by Tiphéne et al. [1] allows performing an indentation test in 0.8 s but induces a burst effect at the beginning of the loading step on all tested materials. Some may believe that it is a "pop-in" effect [24]. However, in this case, it is believed to be an experimental artifact since it was also observed on fused silica (see Figure S2). It is probably due to the infinite value of the derivative of the load at the starting point of the test. To overcome this effect, the quarter sinus loading function used in [1] has been replaced by a half-sinus loading function.

$$P(t) = P_{max} \sin\left(\frac{\pi t}{2t_{load}}\right) \Rightarrow P(t) = \frac{P_{max}}{2} \left(1 - \cos\left(\frac{\pi t}{t_{load}}\right)\right) \quad (1)$$

With P_{max} the maximum load and t_{load} the loading time. The corresponding loading cycles have been plotted in Figure S2(a). In Figure S2(b), the application on fused silica shows the disappearance of this artifact at the beginning of loading.

65 II.2.b From maximum load control to maximum depth control

66 As hardness usually decreases with increasing temperature, the tested volume increases when perform-
 67 ing maximum load-controlled HTSI experiments [1]. However, some materials presents an Indentation
 68 Size Effect (ISE) at room temperature : the mechanical properties depend on the tested depth. Then, the
 69 hardness reduction observed during testing is due to both the temperature and the increased maximum
 70 depth. As, the ISE may depend on temperature, it is not possible to predict its changes in tempera-
 71 ture and so to correct its effect based on room temperature measurements. To reduce the impact of
 72 the ISE and better control the tested volume, a cycle controlled in displacement would be much bet-
 73 ter. As it is quite complicated to implement a 1-second depth-controlled indentation cycle on the used
 74 force-controlled nanoindentation device, a different approach is used. The idea is to adapt the applied
 75 maximum load between each test using the result of the previous test. Under the hypothesis that the
 76 hardness does not change much between two tests, it is then possible to predict the load to target to
 77 always perform tests at a given maximum depth. Such a strategy requires a sample where hardness
 78 does not vary in space nor is expected to vary abruptly (due to microstructure changes, for instance
 79 [24]) during the thermal cycle. So the maximum load in equation 1 for test i would be calculated using
 80 the results of test $i - 1$:

$$P_{max}^i = P_{max}^{i-1} \left(\frac{h_{target}}{h_{max}^{i-1}} \right)^2 \quad (2)$$

81 With h_{target} the maximum depth that one wants to reach at all temperatures, and P_{max}^{i-1} and h_{max}^{i-1} repre-
 82 senting the maximum load and depth reached in test $i - 1$, respectively.

83 II.3 Nanoindentation analysis

84 Analysis of the HTSI results is performed using a Python code. The classical definitions of hardness,
 85 Young's modulus, etc., used in nanoindentation, are implemented. To confirm those results, CSM at
 86 constant strain rate (CSR), creep, and relaxation tests are also performed at different temperatures. The
 87 equations for the analysis are reminded in the following section. HTSI values are determined at the
 88 maximum depth point, while the CSM loading gives the mechanical properties all along the depth.
 89 Depending on the contact, pile-up or sink-in is observed: one should then prefer Loubet's model [13] or
 90 Oliver and Pharr's model [12].

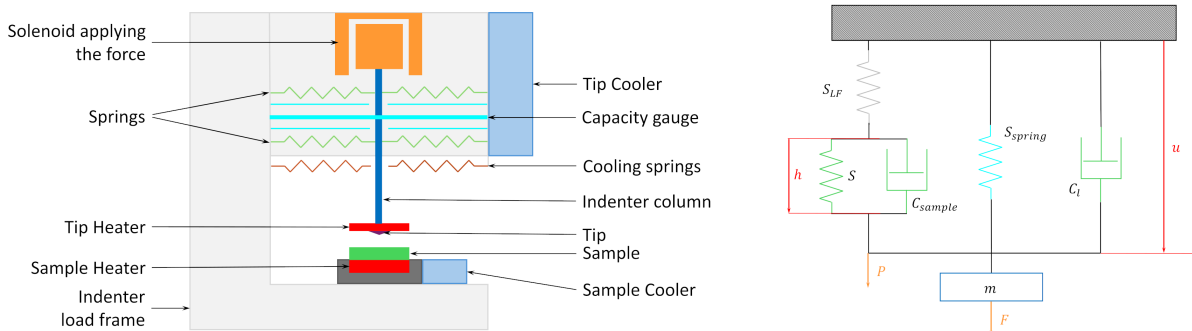


Figure 1: Schematic representation of the indenter and corresponding rheological model. We apply the raw force F and measure the raw displacement u and the raw stiffness K . To determine the mechanical properties, the load P , depth h and contact stiffness S are required. Adapted from [25].

91 II.3.a Determination of modulus and hardness

92 We measure the raw force F and the raw displacement u through the solenoid and capacity gauge of
 93 Figure 1. The total stiffness K is then:

$$K = \frac{dF}{du} \quad (3)$$

94 The load is defined as:

$$P = F - S_{spring}u \quad (4)$$

and the depth:

$$h = u - \frac{P}{S_{LF}} = u \left(1 + \frac{S_{spring}}{S_{LF}} \right) - \frac{F}{S_{LF}} \quad (5)$$

The contact stiffness S is related to K though:

$$\frac{1}{S} = \frac{1}{K - S_{spring}} - \frac{1}{S_{LF}} \quad (6)$$

with $S_{S_{LF}}$ denoting the load frame stiffness, calibrated on fused silica. It is supposed to stay constant in temperature [19]. In the following, the spring stiffness term is neglect in regard of its value (≈ 700 N/m) compared to the total stiffness ($\approx 5 \times 10^5$ N/m).

For HTSI tests, the total stiffness K is calculated using the unloading part of the indentation cycle [1], assuming a linear fit.

$$F = K(u - u_0) \quad (7)$$

with u_0 the raw displacement when the force is null.

In the present work, SEM observation confirmed that no pile-up phenomenon was observed on post-mortem indents on CaF_2 . Oliver and Pharr's [12] model was then used to determine the contact depth h_c in this case. According to this model:

$$h_c = h - 0.75 \frac{P}{S} \quad (8)$$

with h , the measured depth, P the applied load and S the contact stiffness.

The contact area is:

$$A_c = \sum_0^n C_i h_c^{2/2^i} \quad (9)$$

with the C_i been calibrated on fused silica. For a perfect Berkovich tip, $C_0 = 24.5$ and the others C_i are null.

The hardness H is defined as:

$$H = \frac{P}{A_c} \quad (10)$$

Through Sneddon work's [12, 13, 26], the stiffness can be related to the reduced modulus:

$$E_r = \frac{S}{2} \sqrt{\frac{\pi}{A_c}} \quad (11)$$

The indentation modulus M [27–29] is then defined as:

$$\frac{1}{M} = \frac{1}{E_r} - \frac{1 - \nu_i^2}{E_i} \quad (12)$$

with E_i and ν_i the tip Young's modulus and Poisson ratio respectively. ν_i was considered constant in temperature. The tip modulus changes in temperature were implemented, following the formula in [18].

In case of homogeneous isotropic materials, the Young's modulus E is then computed as:

$$E = \frac{M}{(1 - \nu^2)} \quad (13)$$

With ν the Poisson ratio of the sample. We are dealing here with single-crystal samples, whose behaviors are anisotropic. However, we will still consider that equation 13 holds true in the following. For a more precise analysis, the work of [27–30] should be consulted.

II.3.b Creep properties

To characterize the creep properties of the sample, the representative stress σ_r is first computed using the method described in [31]:

$$\sigma_r = \frac{H}{\gamma} = \frac{\xi_3 \cot \theta}{\xi_1 \cot \theta - (1 - \xi_2) \frac{H}{E}} H \quad (14)$$

123 with θ the equivalent conical angle (70.32° for a Berkovich tip) and ξ_1 , ξ_2 and ξ_3 characteristic tip's
 124 parameters. For HTSI tests, the hardness is calculated along the creep segment using the unloading
 125 contact stiffness to compute the contact depth (see Equation 8).

126 During the long-term creep tests, the load is maintained constant, and the contact stiffness is mea-
 127 sured continuously throughout the holding period. The strain rate is then defined as described in [16]:

$$\dot{\epsilon}_{creep} = \frac{\dot{S}}{S} \quad (15)$$

128 In the case of the relaxation tests, the stiffness is held constant [17]. Therefore, the strain rate can no
 129 longer be defined through the stiffness variations. Following the analysis of Baral et al. [16], the strain
 130 rate is defined using the variations in representative stress:

$$\dot{\epsilon}_{relaxation} = -\frac{1}{E} \frac{d\sigma_r}{dt} \quad (16)$$

131 Finally, as the HTSI tests are quite quick, stiffness is not recorded during the test. Therefore, the
 132 strain rate is then defined using the classical definition [32, 33]:

$$\dot{\epsilon}_{HTSI} = \frac{\dot{h}}{h} \quad (17)$$

133 An example of the calculation of the creep properties in the case of HTSI tests is plotted in Figure S3 in
 134 the Supplementary Material.

135 Supposing that the material follows a Norton-Hoff creep law:

$$\dot{\epsilon} = \alpha \sigma_r^{1/m} \exp\left(-\frac{Q}{RT}\right) \quad (18)$$

136 with α a pre-exponential factor, m the strain-rate sensitivity, Q the creep activation energy, R the gas
 137 constant and T the absolute temperature. At a given temperature, one can determine the strain rate
 138 sensitivity through [15, 16]:

$$m = \frac{d \ln(\sigma_r)}{d \ln(\dot{\epsilon})} \quad (19)$$

139 One can then calculate the activation volume of creep v_{ac} using the relation described in [15, 16]:

$$v_{ac} = \frac{\sqrt{3} k_B T}{m \sigma_r} \quad (20)$$

140 with k_B , Boltzmann's constant.

141 In nanoindentation, one should exercise caution when comparing creep and relaxation tests [16].
 142 Strain rate sensitivity values obtained here were lower than 0.1, so the representative strain rate defini-
 143 tion of [16] was not used.

144 III Results

145 III.1 Interest in the maximum depth-controlled indentation cycle

146 This strategy has been applied to a CaF_2 single crystal. Figure 2 presents the changes in the maximum
 147 load and depth achieved during the HTSI tests. As observed, the proposed changes in maximum load
 148 between each test allow for good control of the maximum depth at various temperatures. Experimen-
 149 tally, false surface detection leads to a false hardness value near 40°C . This results in a significant
 150 decrease in the applied load. However, the system corrects itself quickly (within 2 to 4 indents), and the
 151 target depth value is once again reached. The error remains quite low throughout the entire tempera-
 152 ture range (see Figure 2c). CaF_2 is known to exhibit an Indentation Size Effect (ISE) on hardness [23,
 153 34]. As observed in Figure 2, hardness decreases more rapidly when using a maximum load-controlled
 154 cycle instead of a maximum depth-controlled cycle due to this effect.

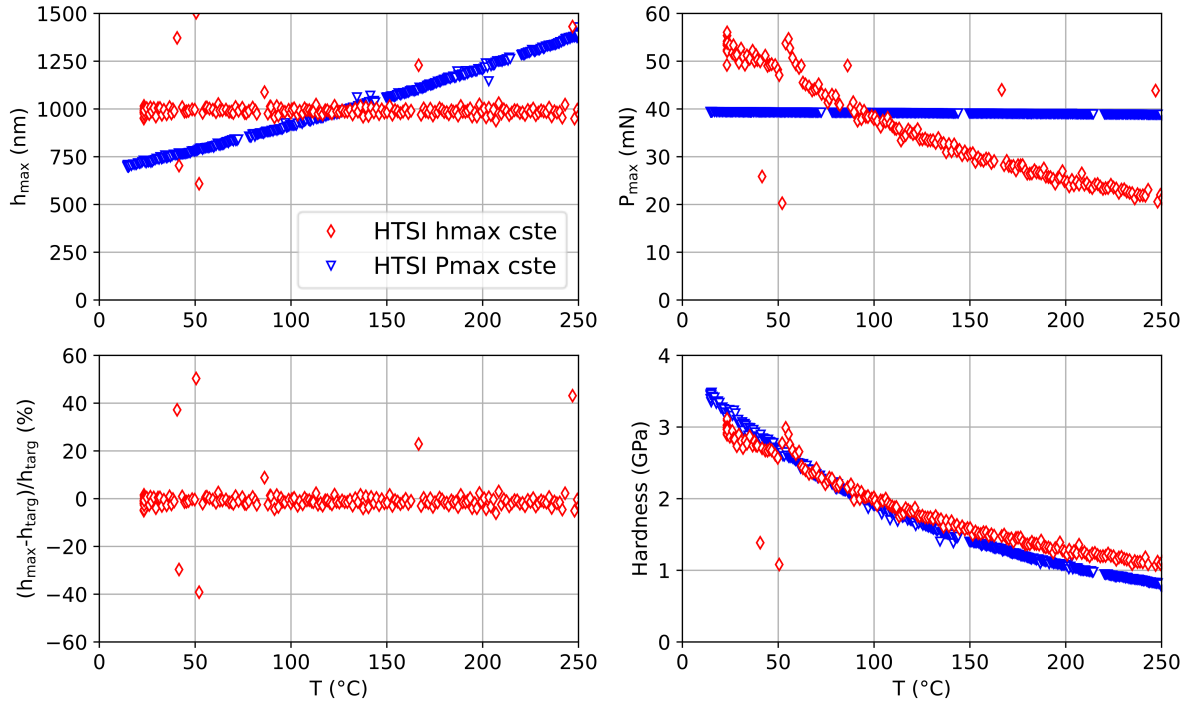


Figure 2: (Top) Maximum depth (left) and load (right) versus temperature measured on CaF_2 during heating. (Purple) Indentation cycle with a target load of 40 mN at all temperatures. (Cyan) The adapted indentation cycle is used with a 1000 nm target depth at all temperatures. (Bottom left) Percentage of error on achieved maximum depth compared to target depth for the maximum depth-controlled HTSI tests. Except for the points related to false surface detection, the error stays within $\pm 5\%$. (Bottom right) Corresponding hardness versus temperature curves on CaF_2 . The impact of increasing testing depth can clearly be observed on the hardness values here.

155 III.2 CaF_2 at room temperature

156 CaF_2 was first characterized at room temperature. Figure 3 presents the hardness and Young's modulus
 157 determined by CSM at various loading rates. Mean values at 900 nm are recap in Table ?? . Values are
 158 consistent with literature [23, 34]. As it can be seen, there is no clear impact of the strain rate on the
 159 modulus. However, on hardness, there is both an impact of loading rate [35] and a size effect [23]. For
 160 future analysis, when comparing results, they are taken at a depth of 1000 nm.

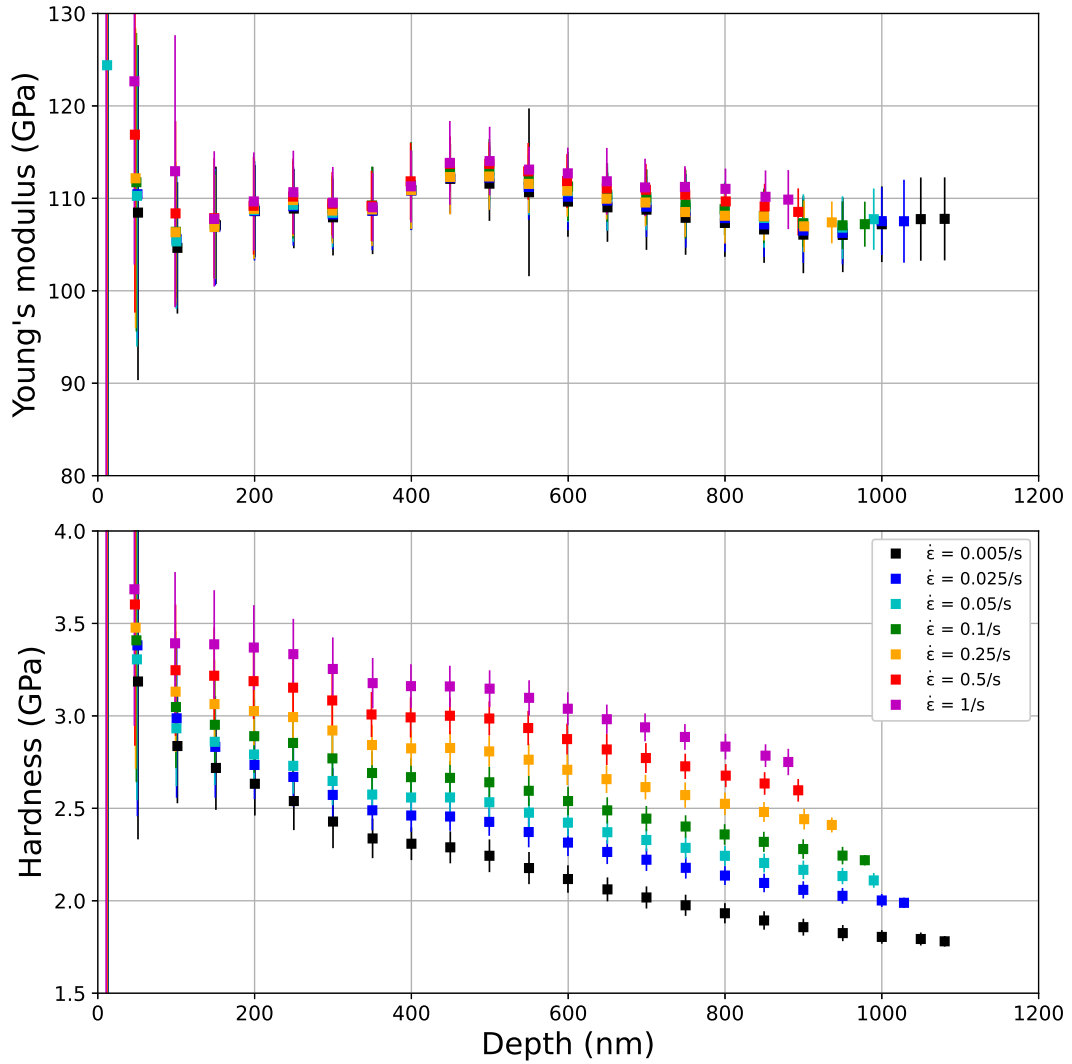


Figure 3: Impact of strain rate on hardness and Young's modulus at RT on CaF_2 samples. The given values are the mean \pm 3std values. 5 to 10 indents were carried out at each strain rate. Contrary to modulus, hardness is sensitive to strain rate and exhibits a size effect. Unfortunately, the tip used to get those results has been damage by HT indentation tests, which explain the oscillations that can be observed at low depths.

161 To gather more information on the creep behaviors of CaF_2 at room temperature, indentation creep
 162 and relaxation tests, as well as constant high-strain rate tests, were conducted. The results of strain rate
 163 versus modulus-compensated stress are plotted in Figure 4. The evolution is consistent across the tests.
 164 However, the variations are not linear at all, and the sample appears more sensitive to strain rate as it
 165 increases. This behavior is unexpected for FCC structures but quite common in BCC structures [36].

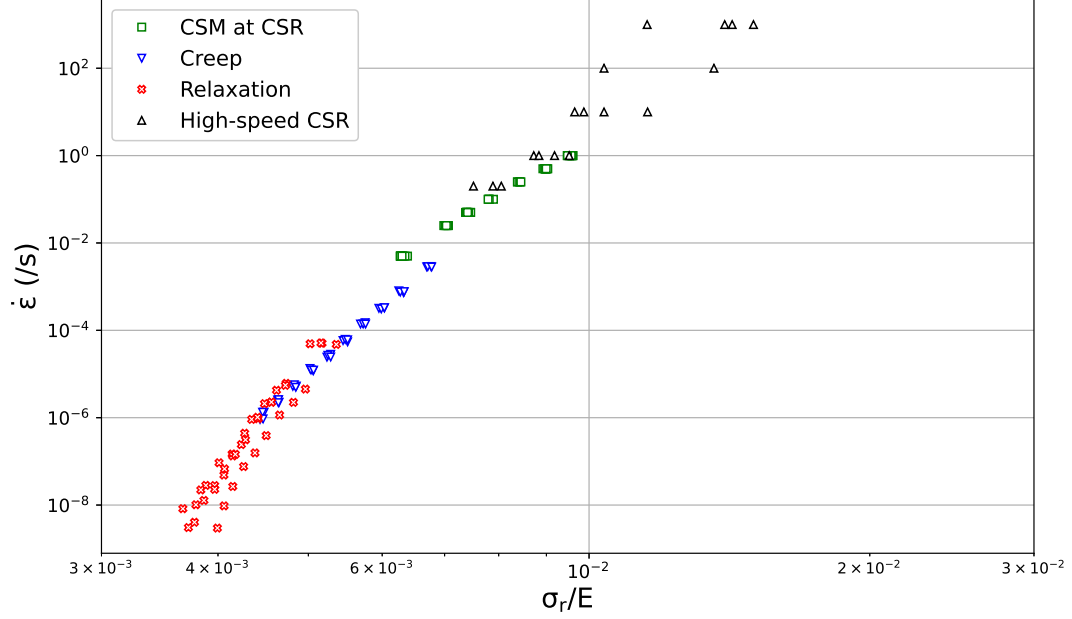


Figure 4: Stress versus strain rate curve on CaF_2 single-crystals at room temperature, obtained using various indentation tests. Non-linearity indicates different creep behaviors, depending on strain rate and temperature.

166 III.3 CaF_2 at high temperatures

167 III.3.a Mechanical properties in temperature

168 Young's modulus and hardness changes with temperature are plotted in Figure 5. The black curve
 169 corresponds to the expected changes in Young's modulus for CaF_2 along the studied orientation [35,
 170 37]. As expected, the measured values decrease with temperature. However, the absolute values vary
 171 around the expected value at a given temperature. In the HTSI tests at temperatures higher than 750°C ,
 172 the variations in Young's modulus are quite unreliable. Due to issues with the automatic testing, the
 173 tests at these high temperatures were conducted manually, and thermal equilibrium is not expected to
 174 be achieved. It can be observed that all CSM tests at 200°C yield very small values.

175 As previously observed, the hardness value at room temperature depends on the strain rate. A
 176 mean strain rate of 2s^{-1} is estimated during loading in the HTSI tests, which is consistent with the
 177 higher hardness value obtained at room temperature. When increasing the temperature, hardness starts
 178 decreasing rapidly until 200°C . At higher temperatures, it still decreases but less rapidly. A slight drop
 179 in hardness is observed around 500°C before the decrease in hardness smoothens again until 750°C
 180 approximately. Strangely, an increase in hardness is observed near 800°C .

181 Hardness variations with temperature are consistent with those measured by Deadmore [11], al-
 182 though the absolute values differ. This discrepancy is expected to be due to differences in tested depth
 183 since Vickers hardness yields results at larger depths compared to nanoindentation tests. Finally, the
 184 impact of strain rate on hardness seems to decrease with increasing temperature.

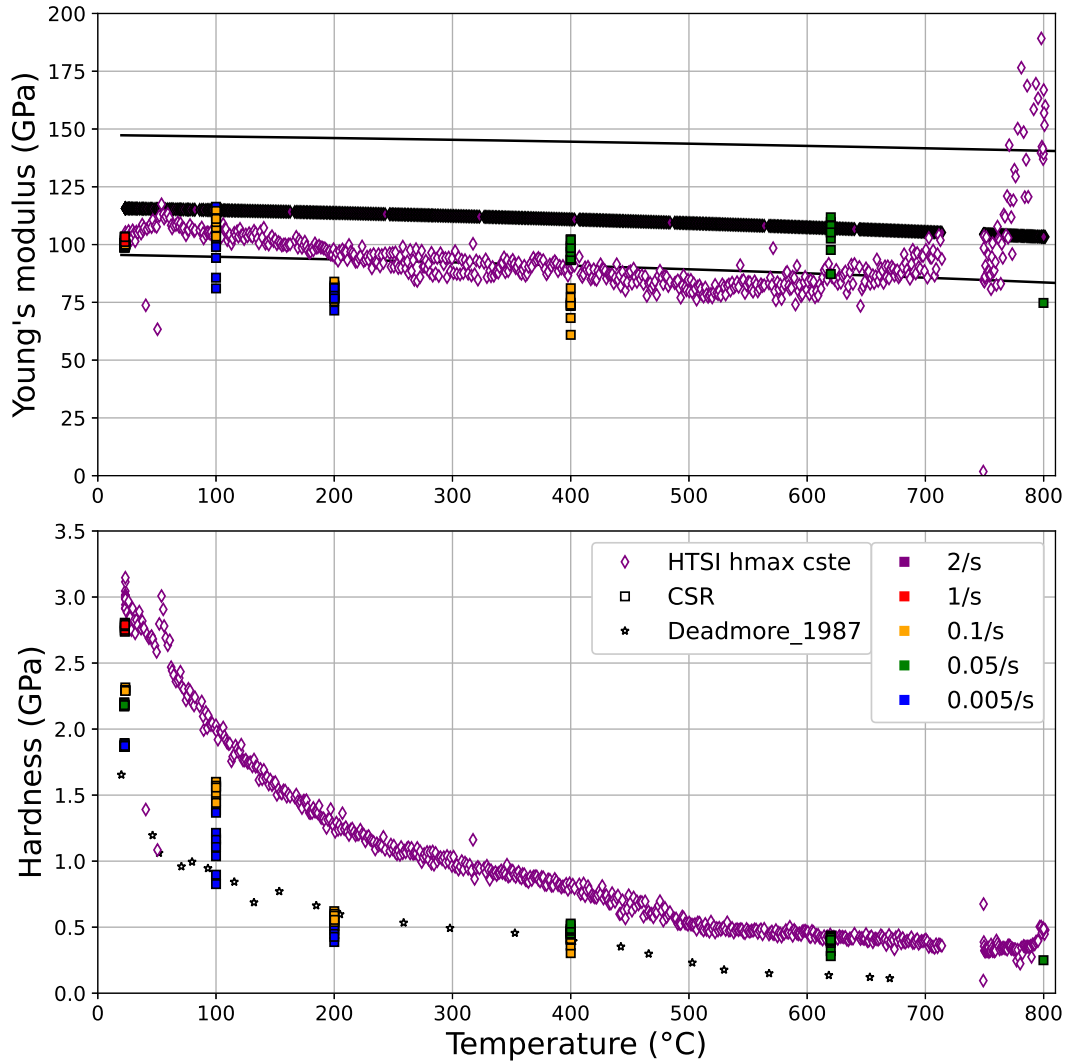


Figure 5: (Top) Young's modulus and (bottom) hardness changes with temperature for CaF_2 single-crystal. All data are measured at a depth of 1000 nm. The colors indicate the strain rate during loading (mean values for HTSI tests). Strain rate greatly impacts the hardness values at room temperature, but its impact decreases at high temperature. The hardness changes are consistent with literature data [11]. On the Young's modulus data, the black lines represent Voigt (top) and Reuss (bottom) bounds. The black diamonds show the expected Young's modulus changes, given the crystal orientation. The HTSI data appears consistent with these values up to 750 °C.

185 III.3.b Creep properties

186 To obtain the creep properties of CaF_2 at different temperatures, creep and relaxation tests were con-
 187 ducted. The strain rate versus modulus-compensated stress curves at all temperatures can be founded
 188 in Figure S4 in the supplementary material. Classical constant strain rate (CSR) and creep indentation
 189 tests are compared to relaxation tests and HTSI results.

190 Under the assumption that a single creep mechanism dominates the creep behavior of CaF_2 (Norton-
 191 Hoff creep law, Equation 18), the master curve has been plotted in Figure 6 for an activation energy
 192 of 100 kJ/mol. Such analysis give access to more than 20 orders of magnitude for the temperature-
 193 compensated strain rate.

194 The different tests overlap quite well, and the trend at high temperatures appears consistent with
 195 the results of Sadrabadi [10]. However, assuming a constant value of the activation energy over this
 196 wide temperature range is questionable since, according to the literature, we would expect it to increase
 197 with temperature: Oneill *et al.* [38] found an energy of 42 kJ/mol for temperatures below 300 °C, while

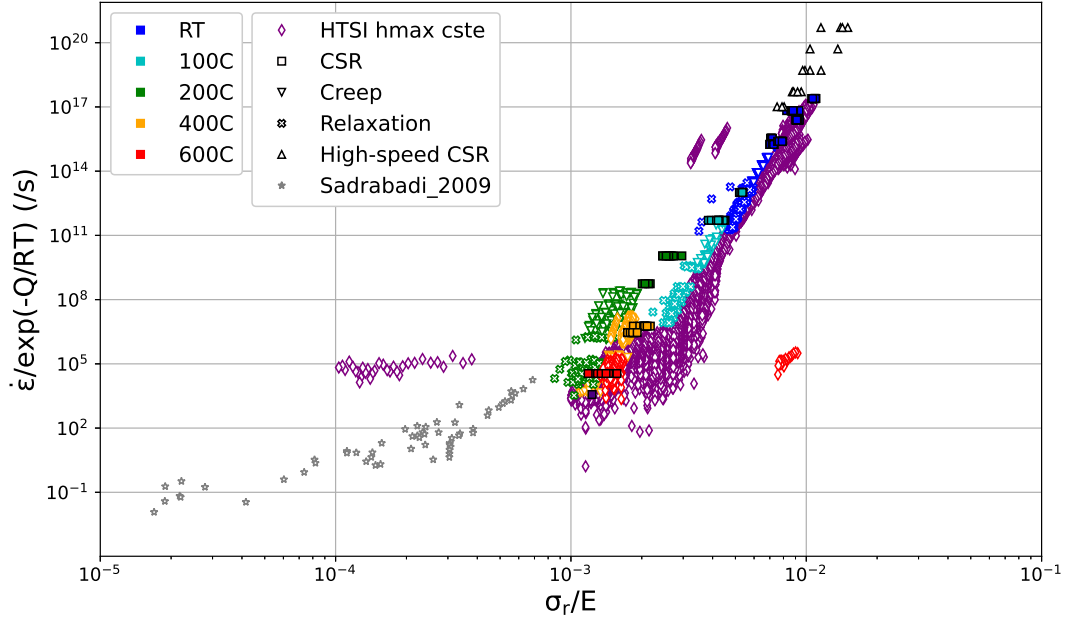


Figure 6: Master curve obtained for CaF₂ with an activation energy of 100 kJ/mol. The various indentation tests exhibit good agreement, and the trend is consistent with the high-temperature tests conducted by Sadrabadi [10].

198 Mekala [39] proposed an apparent activation energy (in J/mol) for creep following:

$$Q(T) = 8.31(20.8 T(K) + 7640) \quad (21)$$

199 when the temperature is in the range 400 °C to 927 °C (180 kJ/mol to 271 kJ/mol).

200 The strain rate sensitivity and activation volume have been plotted against the temperature-compensated
 201 strain rate in Figure 7. At room temperature, the activation volume is relatively low (ranging from 1 to
 202 $5b^3$ depending on the strain rate). An increase in temperature or a decrease in strain rate leads to an
 203 increase in this volume.

204 The activation volume appears to increase linearly up to 200 °C. Interestingly, the activation volume
 205 from the HTSI tests increases more rapidly than that from the creep and relaxation tests between 200 °C
 206 and 400 °C. However, at higher temperatures, the HTSI results align once again with those of the creep
 207 and relaxation tests. At high temperatures, the trend is consistent with the results of Sadrabadi [10].

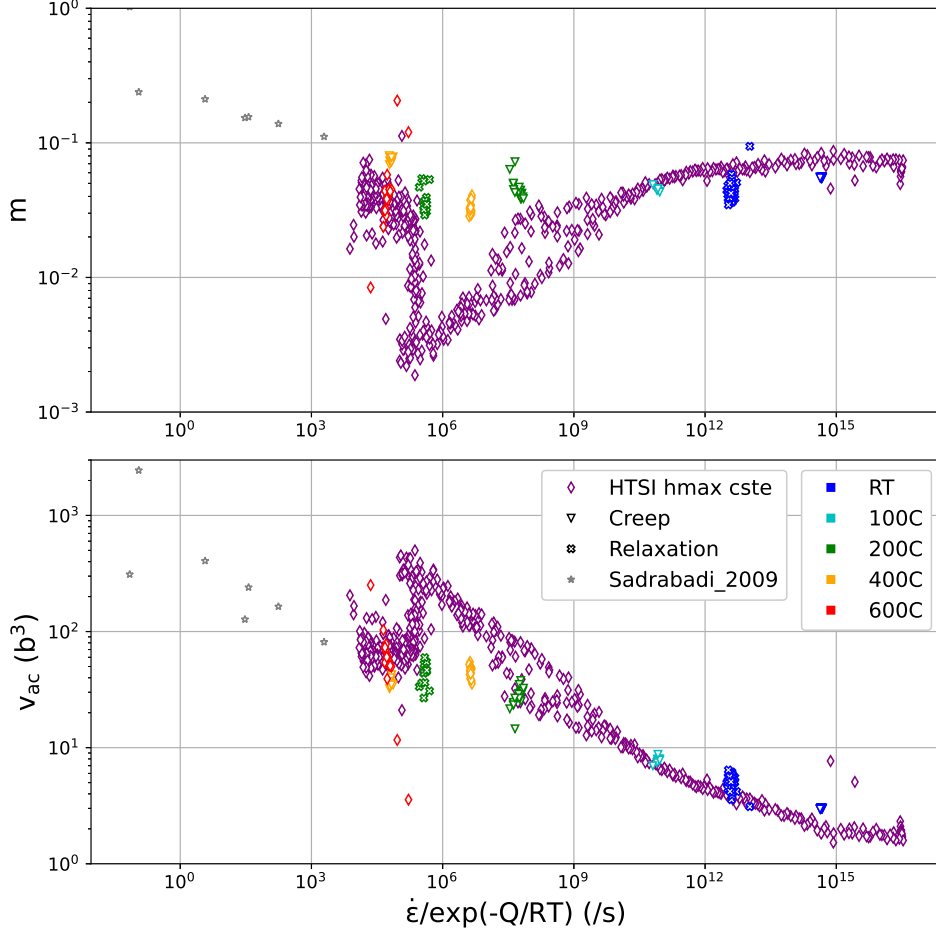


Figure 7: (Top) Strain rate sensitivity and (bottom) activation volume for creep versus temperature-compensated strain rate ($Q = 100$ kJ/mol). The results show consistency at both low and high temperatures (for $x > 1 \times 10^{10}$ or $x < 1 \times 10^6$). However, in the intermediate range, there appear to be two distinct behaviors, corresponding to the HTSI tests conducted between 200°C and 500°C .

IV Discussion

IV.1 On the HTSI method

IV.1.a Impact of temperature control on H and E

As observed previously (Figure 5), changes in hardness and Young's modulus with temperature appear consistent, at least up to 750°C for the modulus. In Figure 8, we plot the maximum depth, load, stiffness, contact depth and area, reduced modulus, Young's modulus, and hardness versus temperature determined by HTSI. It is worth mentioning that the control of the achieved maximum depth is satisfactory across the studied temperature range. Although there is some scattering, it results in a constant contact depth and contact area. Consequently, variations in hardness are primarily attributed to fluctuations in maximum load. Conversely, changes in reduced modulus (and consequently Young's modulus) are primarily influenced by variations in stiffness.

A detailed analysis of the measurements errors impact on the properties can be founded in the supplementary materials (Section S.III.). Considering Figure S5, it is evident that the error in stiffness will significantly impact Young's modulus but will have less impact on the contact depth, contact area, and hardness. For this reason, the authors have more confidence in the hardness data and attribute its variations to the material rather than the measurements. However, they caution that Young's modulus data should be analyzed with care. While the overall trend is usually consistent, the values should be interpreted cautiously.

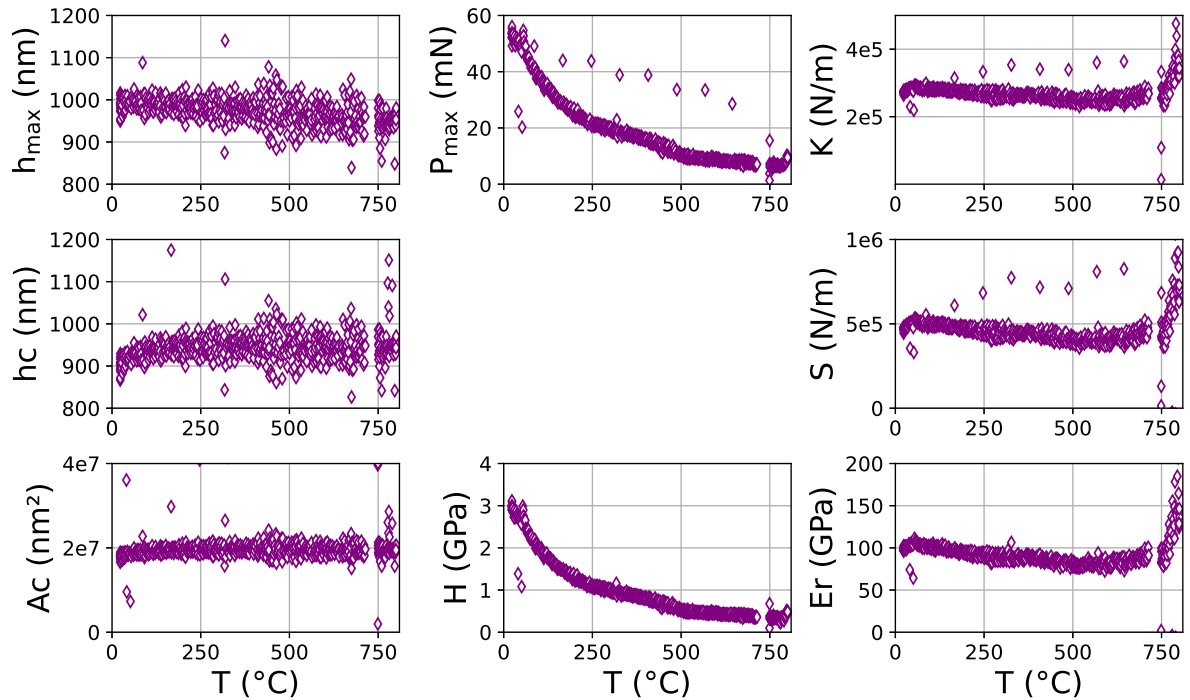


Figure 8: The evolution of maximum depth, load, contact stiffness, contact depth, reduced modulus, contact area, hardness, and Young's modulus with temperature for CaF_2 single-crystal, as determined through HTSI testing, is presented. It is evident that variations in hardness directly correspond to changes in load, while alterations in Young's modulus are associated with fluctuations in contact stiffness.

226 IV.1.b Control of the achieved maximum depth

227 As demonstrated previously, the maximum depth control methodology performs well on CaF_2 single-
 228 crystal samples. This approach was then applied to annealed copper samples with a grain size of around
 229 $50 \mu\text{m}$.

230 A series of 100 indents was conducted, with the first indent performed at a controlled maximum
 231 load. Subsequent ninety-nine indents were conducted with a target maximum depth of either 200 nm or
 232 1000 nm (see Figure 9). As observed, when targeting a maximum depth of 1000 nm, the proposed modi-
 233 fication leads to an achieved depth in the range of $\pm 20\%$ of this value. This is not that good. Moreover, if
 234 the targeted depth is decreased, higher errors are obtained. This technique results in significant discrep-
 235 ancies between the achieved maximum depth and the target depth on annealed copper. These errors are
 236 expected to arise from local variations in hardness related to the microstructure of the polycrystalline
 237 copper samples. An alternative methodology should be considered to conduct maximum depth control
 238 indentation tests on copper (and similar) samples. The assumption of low hardness variations in space
 239 is crucial for the successful implementation of this strategy. It should be notice that 20% difference in
 240 the maximum achieved depth at all temperatures may be better than multiplying the achieved depth by
 241 2 or 3 between RT and high temperature.

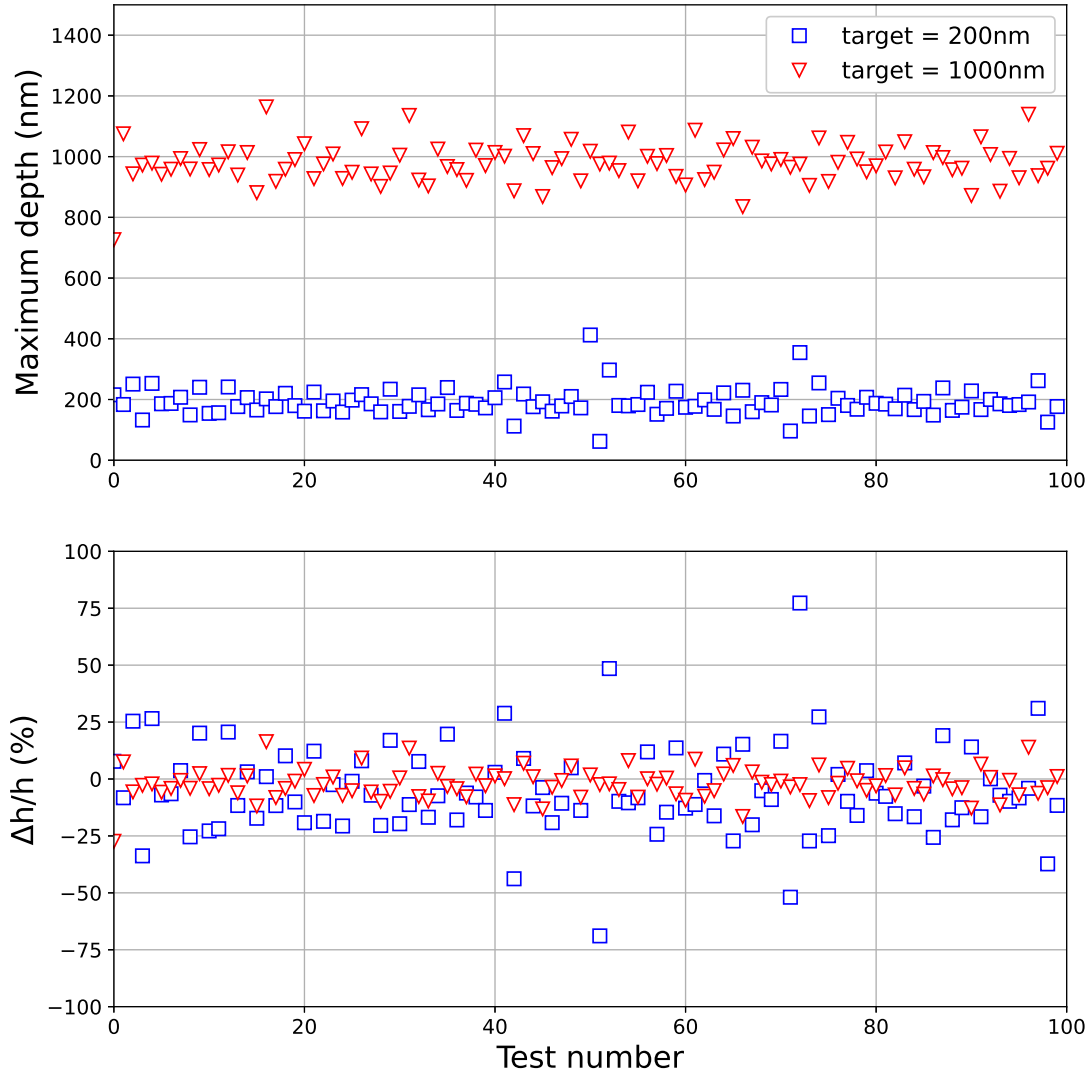


Figure 9: Updated indentation cycle applied to pure annealed copper. (Top) Maximum depth achieved in the test depending on the target depth. Blue squares represent a target depth of 200 nm, while red triangles represent a target depth of 1000 nm. (Bottom) Percentage of error in the achieved depth compared to the target depth.

242 IV.1.c Creep analysis

243 As it can be seen in Figure S3, depth increases during the creep segment during the HTSI tests. The
 244 same effect can be observed when performing classical creep test. As there is some ISE on the samples,
 245 it is expected that part of the hardness decrease comes from this effect. This is an error source in the
 246 estimation of the creep properties. However, in the case of HTSI tests, the authors have not yet found a
 247 way to correct it since stiffness is not recorded during the creep segment. Moreover, this effect may be
 248 negligible compared to the others hypotheses supporting the analysis.

249 For the classical creep tests is also around 100 nm at RT. As stiffness is measured during the creep
 250 segment, it should be possible to correct this effect by estimating the hardness decreases due to ISE
 251 before calculating the creep properties. However, the thermal drift may be a larger source of error so
 252 we do not correct this effect on the tests.

253 Finally, the ISE is temperature-dependent effect and is expected to be less and less pronounced as
 254 temperature increases. From CSM tests performed at various temperatures, the authors estimated that
 255 near 200 °C the hardness variation between 1000 and 1200 nm is 0.025 GPa ($H(200^{\circ}C) \approx 0.35$ GPa) while
 256 the decrease is estimated at 0.1 GPa at RT.

257 One should be careful when analyzing creep tests following the presented analysis as it may lead to
 258 inaccurate results if stationary creep is not reached in the timespan of the creep stage.

IV.2 CaF₂ behaviors

IV.2.a A BCC-like behaviors at RT

At room temperature (RT), the activation volume appears to be quite low (see Figure 7). This observation is intriguing because it suggests that a thermally activated mechanism such as climb could be the controlling mechanism at RT [40]. However, this proposition seems unexpected because climb mechanisms typically require thermal activation, as noted in previous studies [41].

Further insights from studies by Keig et al. [42] and Evans et al. [43] shed light on this phenomenon. These studies suggest that edge dislocations exhibit significantly lower speeds compared to screw dislocations at RT. Additionally, at low temperatures (below 700 °C), vacancies may arise due to the movement of interstitial fluorine ions, resulting in local charges [44]. Evans et al. [43] propose that near RT, a probable controlling mechanism could involve the interaction of edge dislocations with these local charges. In this scenario, edge dislocations would need to overcome electronic barriers to move within the crystal structure.

This behavior bears resemblance to the phenomenon of screw dislocations overcoming Peierls stress in body-centered cubic (BCC) structures at low temperatures [8, 41]. Further analysis would be required to better understand this behavior.

IV.2.b Changes of behaviors with high temperature

As observed previously, hardness exhibits significant scattering at low temperatures due to strain rate effects. This phenomenon can be attributed to the activation of a low number of gliding systems at low temperatures [38, 45, 46]. Such behavior resembles the characteristics of body-centered cubic (BCC) structures. As the temperature increases, more gliding systems become activated. Near temperatures of around 200 °C [38] and between 400-500 °C [46, 47], an increase in the number of active gliding systems is observed. This trend is consistent with the observed drop in hardness near 500 °C in Figure 5. Finally, at temperatures higher than 600 °C, plasticity is expected to become more isotropic [47].

As observed in Figure 5, an unexpected increase in hardness is observed near 800 °C. This could be attributed to a reaction of the sample with oxygen, resulting in material hardening [11]. After the high-temperature testing, the samples were no longer transparent, indicating a reaction had occurred. SEM observation of the samples (see Figure S6 and S7) revealed the presence of a structure on the sample surface that was not observed before high-temperature testing. EDX analysis (see Figure S8) of this structure showed a depletion of fluorine, replaced by oxygen. This suggests that the sample reacted with oxygen at high temperatures to form calcium oxide [48–50]. It appears that the vacuum level of 1×10^{-2} Pa was insufficient to prevent this reaction.

V Conclusions

This paper presents an update version of the High-Temperature Scanning Indentation technique [1]. It allows determining the mechanical properties of a material at a given depth on the whole studied temperature range. The main points of the article are as follows:

- The 1-second indentation cycle presented in [1] has been modified. HTSI tests are then carried out at a controlled maximum depth all along the thermal cycle. To do so, the applied maximum load is adjusted between each indent, using previous results.
- The old and update cycles have been applied to study the mechanical properties of CaF₂ single-crystals from room temperature up to 200 °C. CaF₂ is known to present a important ISE effect against depth. Thanks to the update methodology, the temperature effect has been separated for the ISE effect and has been clearly quantified here. The error on the maximum achieved depth stays within $\pm 5\%$ on the whole temperature range on CaF₂ single-crystals.
- Error analysis on the determination of hardness and Young's modulus shows that hardness values obtained by HTSI are quite reliable in temperature. On the other hand, stiffness and Young's modulus are greatly impacted by temperature mismatch: one should analyze them with caution.
- Hardness and creep properties of CaF₂ have been characterized from RT to 800 °C. The results are consistent with literature data.

- An equivalence between time and temperature was successfully plotted with an activation energy of 100 kJ/mol. Various indentations tests give consistent results with literature data. A focus on the activation energy values would be required to better understand the mechanism(s) controlling creep depending on temperature and strain rate for this material.
- Hardening of the sample near 800 °C has been related to the reaction of the sample with H₂O or O₂ present in the chambers through EDX characterization.

However, the proposed methodology is based on the assumption of low hardness variations in space. The error on the achieved maximum depth is quite high when applying the technique on annealed copper. It was expected but it means another methodology should be applied for this type of samples. Moreover, having the properties changes again depth at all temperature would be quite interesting. Finding a way to implement a CSM mode at high speed would be a way to go further on this topic.

VI Materials and methods

VI.1 Material

The CaF₂ single crystals were provided by Edmund Optics. These transparent crystals were uncoated flat samples of 2 mm-thickness and 12.5 mm-diameter. Parallelism is lower than 1 arcmin. The physical properties were taken from the literature: Burger's vector [51], lattice parameter [52], elastic constants [53], and Poisson ratio (0.26) [35].

The results on copper samples were obtained using the annealed samples previously used in [24].

VI.2 Experimental set-up

VI.2.a Microstructure characterization

All SEM characterizations were carried out using a Tescan SEM MIRA 3 equipped with an EDX Oxford Instruments XMax 80 mm² detector and a EBSD Oxford Instruments Nordlys camera. EBSD characterization was carried out to verify the orientation of the CaF₂ single-crystal. EDX analysis was conducted on CaF₂ samples after high-temperature nanoindentation to verify the surface composition. Post-processing was performed using the MTEX MATLAB ToolBox [54].

VI.2.b Nanoindentation testing of CaF₂

Load-controlled nanoindentation tests were conducted using the InSEM HT nanoindentation device (KLA Corporation), which was located inside a vacuum chamber (KLA, TN, USA) or inside a TESCAN SEM (LTDS, France) with a pressure of 1×10^{-2} Pa. A diamond Berkovich tip was used at room temperature, a sapphire tip was used up to 400 °C, and a WC tip was used up to 800 °C. Tip calibration was performed on a fused silica sample at ambient conditions prior to any experiments [13, 14]. Between high-temperature tests, the tip was regularly calibrated on fused silica. The specimens were mounted as indicated by Minnert et al. [19]. The minimal distance between each indent was 10 times the contact depth [55]. Temperature settings and controls were conducted in the same way as in [1, 24].

Table 1 presents the various indentation tests that were carried out on the CaF₂ samples. Classical CSM [12, 13] tests, as well as creep [15, 56] and relaxation [17] indentation tests, were conducted between room temperature (RT) and 800 °C after reaching thermal equilibrium. Due to difficulties in maintaining thermal equilibrium for long periods at high temperatures, relaxation tests were only conducted up to 200 °C and creep tests up to 400 °C.

HTSI [1] tests were then carried out to obtain the mechanical properties throughout the studied temperature range. The updated version of the indentation cycle was used to control the maximum achieved depth throughout this range. The loading is performed in 0.5 s which gives a mean strain rate during loading of 2 s^{-1} .

To complete these results, high-speed constant strain rate (HS-CSR) indentation tests at RT were performed on an Alemnis device (depth-controlled device) [57].

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ChatGPT 3.5 was used to correct the grammar and spelling of the present article.

Table 1: Indentation tests carried out on the CaF₂ samples.

type of test	Temperature	h_{max}	$\dot{\epsilon}_{loading}$	$t_{holding}$
CSM at constant strain rate (CSR)	RT	900 → 1000 nm	$1 \rightarrow 5 \times 10^{-3} \text{ s}^{-1}$	-
	100 °C	1000 nm	$0.1 \rightarrow 5 \times 10^{-3} \text{ s}^{-1}$	-
	200 °C	1000 nm	$0.1 \rightarrow 5 \times 10^{-3} \text{ s}^{-1}$	-
	400 °C	1000 nm	$0.1 \rightarrow 5 \times 10^{-2} \text{ s}^{-1}$	-
	600 °C	1000 nm	$5 \times 10^{-2} \text{ s}^{-1}$	-
	800 °C	1000 nm	$5 \times 10^{-2} \text{ s}^{-1}$	-
Creep	RT	1000 nm	0.1 s^{-1}	10 h
	100 °C	1000 nm	0.1 s^{-1}	1000 s
	200 °C	1000 nm	0.1 s^{-1}	1000 s
	400 °C	1000 nm	0.1 s^{-1}	1000 s
Relaxation	RT	1000 nm	0.1 s^{-1}	10 h
	100 °C	1000 nm	0.1 s^{-1}	1000 s
	200 °C	1000 nm	0.1 s^{-1}	1000 s
ine Half-Sinus Loading	400 °C	1000 nm	2 s^{-1} (mean)	1 s
	600 °C	1000 nm	2 s^{-1} (mean)	1 s
HTSI	RT → 800 °C	1000 nm	2 s^{-1}	1 s
	1.5 °C/min		(mean)	
HTSI	RT → 250 °C	750 → 1400 nm	2 s^{-1}	1 s
	2 °C/min	$P_{max} = 40 \text{ mN}$	(mean)	
HS-CSR	RT	1000 nm	$1 \rightarrow 1000 \text{ s}^{-1}$	-

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Conflict of interest

The authors declare that they have no conflict of interest.

Data Availability

The raw/processed data required to reproduce these findings cannot be shared at this time due to legal reasons.

Code availability

Not applicable

Authors' contributions

Gabrielle Tiphène: Methodology, Formal Analysis, Investigation, Data curation, Writing - Original draft preparation. **Benedicte Adogou:** Investigation, Writing - Review and Editing corrections **Gaylord Guillonneau:** Writing - Review and Editing corrections, Supervision. **Guillaume Kermouche:** Theoretical frame-work, Writing - Review and Editing corrections. **Jean-Michel Bergheau:** Writing - Review and Editing corrections, Supervision. **Warren C. Oliver:** Conceptualization of this study, Methodology, Software, Resources, Writing - Review and Editing corrections, Supervision. **Jean-Luc Loubet:** Conceptualization of this study, Resources, Writing - Review and Editing corrections, Supervision

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