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Crystal viscoplastic modeling of UO$_2$ single crystal

Abstract

The viscoplastic behavior of uranium dioxide (UO$_2$) single crystal is of great interest to perform predictive multiscale modeling of the nuclear fuel. Here, a viscoplastic model is built considering dislocation glide in $\frac{1}{2}<110>{100}$ and $\frac{1}{2}<110>{110}$ slip systems. The constitutive law parameters are determined adjusting the temperature dependency of the experimental critical resolved shear stress for both principal slip modes. Crystal plasticity finite element simulations of single crystal compression tests show a reasonable agreement with experimental viscoplastic anisotropy of UO$_2$. However, for specific orientations where $\frac{1}{2}<110>{111}$ slip is observed experimentally, significant differences remain between experimental and computed compression stresses. Therefore, the role of $\frac{1}{2}<110>{111}$ slip is investigated based on a parametric study that provides new insights on UO$_2$ plastic deformation. Several parameterizations of $\frac{1}{2}<110>{111}$ slip are tested highlighting the complexity of UO$_2$ viscoplastic behavior. Significant improvements are still required to explain all simulation-experiment gaps.

1. INTRODUCTION

The primary fuel material in nuclear power plant is made of uranium dioxide (UO$_2$) integrated in a zirconium alloy cladding tube. While the viscoplastic behavior of the material plays a key role during nominal and accidental reactor operations, it is a complex process due to the heterogeneous fuel microstructure and its evolution under irradiation. In this context, the link between fuel pellet mechanical properties and the cladding tube integrity has been established [1-3] but complex multiphysics simulation tools are required to provide the scale transition between involved elementary mechanisms and the macroscopic behavior. The development of a constitutive model that captures the underlying physics, without irradiation effect at first, is important for polycrystalline behavior itself. In addition, fission products releases are known to strongly depend on the local mechanical state [4]. Thus, the improvement on the knowledge of the mechanical state at the lower grain scale is of great importance for the global multiphysics scheme. As a consequence, single crystal modeling appears as the first fundamental step to build a multi-scale and multiphysics modeling approach able to predict the global polycrystalline mechanical properties and also the consequences of fission products and mechanical behavior interactions.

UO$_2$ millimeter size specimens exhibit a plastic behavior under compression at temperatures larger than 800 K. UO$_2$ has the fluorite structure ($a_0=5.47$ Å) that is made of a face-centered cubic lattice of uranium atoms that entangles a cubic sublattice of oxygens located in tetrahedral interstitial positions. The three different dislocation slip modes of UO$_2$ usually referenced in the literature are $\frac{1}{2}<110>{100}$, $\frac{1}{2}<110>{110}$ and $\frac{1}{2}<110>{111}$ [5]-[14]. Those systems are also observed in other fluorite materials as e.g. CaF$_2$ [15]-[18], BaF$_2$ [19] and ThO$_2$: [20],[21]. In UO$_2$, critical Resolved Shear Stress (CRSS) for {100} and {110} modes show a strong dependency to the temperature, what underlines the role of thermally-activated deformation processes. Lefebvre [11] and Byron [6] emphasize the decrease of
experimental CRSS in \( \frac{1}{2}<110>{\{100}\} \), the most favorable slip mode, down to 20 MPa increasing the temperature up to 1900 K. Although Lefebvre data show a clear changeover to an athermal regime beyond a transition temperature \( T_c=1400 \text{ K} \), no transition is noticed from Byron’s dataset. In the 1400 K – 2000 K temperature range, \( \frac{1}{2}<110>{\{110}\} \) slip systems are characterized by larger CRSS values (when compared to \{100\}) that vary significantly with temperature [7], [9]. \( \frac{1}{2}<110>{\{111}\} \) slip is also observed but always in association with \( \frac{1}{2}<110>{\{100}\} \) or \( \frac{1}{2}<110>{\{110}\} \) slip systems. Dislocation glide and cross-slip from \{110\} and \{100\} slip systems are both observed in \{111\} slip planes. Those systems have also been reported in ThO\(_2\) [21]. For single mode orientations, \{110\} shows CRSS almost twice larger than \{100\} and crystal orientation dependency is roughly consistent with the Schmid law [9]. As \( \frac{1}{2}<110>{\{111}\} \) slip systems are not observed under single mode conditions, no CRSS can be easily derived from experimental single crystal tests.

Only few TEM studies focusing on dislocations in UO\(_2\) are available in the literature. Yust and McHargue [8] characterized the dislocation microstructures for the \{100\} slip mode in the 1050 K – 1700 K temperature range (i.e. within the thermally-activated regime). At low strains, they observed many edge dipoles, as well as long mixed dislocations with many direction changes leading to zig-zag shaped configurations. These straight dislocations are entangled at larger strains. Such observations have been confirmed by few studies for \{100\} and \{110\} slip modes [10], [12] that emphasize the relevance of lattice friction in UO\(_2\) at low and intermediate temperatures. However, no detail analysis about elementary dislocation processes (e.g. kink-pair mechanism, dislocation vs. point-defect interactions) can be found. Sawbridge and Sykes [9] pointed out the surprising higher mobility of screw dislocations in \( \frac{1}{2}<110>{\{100}\} \) as already observed in other fluorite materials [15],[18],[19]. These screw dislocations can cross-slip in \{110\} or \{111\} slip plans leading to the characteristic zig-zag shape. In addition, Keller and collaborators show a slip system transition toward \( \frac{1}{2}<110>{\{111}\} \) at 873 K in hyper stoichiometric UO\(_{2+x}\) [13], [14]. Besides the temperature, the U/O ratio also influences the slip mode activity [10],[13],[14] but no quantitative dependency has been shown up to now.

Atomistic simulation using semi-empirical interatomic potentials have been performed to characterize the Peierls barriers, the 0 K Peierls stress and CRSS vs. temperature profiles at very high strain rates typical of molecular dynamics (MD) simulations for the three slip modes of UO\(_2\) [24-26]. While simulation results show that \( \frac{1}{2}<110>{\{100}\} \) is the softer slip mode (as in the experiment), \( \frac{1}{2}<110>{\{110}\} \) is harder than \( \frac{1}{2}<110>{\{111}\} \). In addition, most of the interatomic parameterizations show that the \{100\} edge dislocation has the lowest Peierls stress. This last result is in contradiction with most of experimental observations indicating that edge dislocation is the rate limiting character in \( \frac{1}{2}<110>{\{100}\} \) slip systems. In addition, Lunev [27] and Soulié [28] show that \{100\} edge dislocation glide due to a kink-pair process.

To model the viscoplastic behavior of UO\(_2\) based on dislocation mechanisms, Sauter et al. [29] propose to include physics-based equations for grains orientation and temperature dependency as well as dislocation strain hardening for polycrystal. Nevertheless, the authors used a simplified polycrystalline approach considering slip on \( \frac{1}{2}<110>{\{100}\} \) only without
any prior parameter identification on the single crystal. On the other hand, Sawbridge and Sykes \[9\] adjust UO\(_2\) plastic anisotropy considering that the measured stress \(\sigma\) relies on single slip conditions only, with the CRSS \(\tau_0\) and the maximum Schmid factor \(S^\alpha:\sigma = \tau_0^\alpha / S^\alpha\). However, this approach is not consistent with the strain compatibility conditions needed to accommodate the total strain under uniaxial compression.

In summary, while \(\frac{1}{2}<110>|100\) and \(\frac{1}{2}<110>|110\) are generally reported as the two principal slip modes in the thermally-activated deformation regime of UO\(_2\), no clear conclusions can be made upon \(\frac{1}{2}<110>|111\) slip and its thermal activation, especially from the experiments. Furthermore, the same uncertainty remains for the dislocation character identification. Despite this lack of knowledge, a single crystal viscoplastic model is necessary to build a predictive polycrystalline model for UO\(_2\) mechanical properties in the thermally-activated regime of deformation. For this purpose, we propose to reprocess experimental flow stresses using a reversed engineering approach to assess a Crystal Plasticity Finite Element Model (CPFEM) that account for the different slip modes of UO\(_2\). At this stage, the stoichiometry dependence will not be integrated. This model will be used to compute plastic anisotropy properties and the contribution of each slip modes to lattice rotation will be discussed. A particular attention will be paid to the \(\frac{1}{2}<110>|111\) slip mode which is currently under debate in the literature.

2. METHODS

2.1. Crystal plasticity model

In the following, \(C_0\) denotes the reference configuration and \(C\) the current configuration. Those configurations are linked by the deformation gradient tensor \(\bar{F}\).

The crystal plasticity model is based on the classical multiplicative Kröner-Lee decomposition \([30-32]\) of the deformation gradient tensor \(\bar{F}\):

\[
\bar{F} = \bar{F}^e, \bar{F}^p
\]

where \(\bar{F}^p\) the plastic deformation gradient and \(\bar{F}^e\) the elastic deformation gradient. According to Mandel, the plastic deformation gradient is associated in this paper with the unique isoclinic intermediate configuration \(G_i\).

Considering equation (1), the velocity gradient \(\bar{L}\) can be expressed as :

\[
\bar{L} = \hat{\bar{F}}.\bar{F}^{-1} = \bar{L}^e + \bar{L}^e.\bar{L}^p, \bar{L}^e^{-1}
\]

With \(\bar{L}^e = \hat{\bar{F}}^e.\bar{F}^e^{-1}\) the elastic part of the velocity gradient in the \(C\) and \(\bar{L}^p = \hat{\bar{F}}^p.\bar{F}^p^{-1}\) the plastic part in \(G_i\). Equation (2) shows that the mechanical power is decomposed into a reversible part (linear in \(\bar{L}^e\)) and an irreversible part (linear in \(\bar{L}^p\)).
The elastic part of the behavior is based on the hyperelastic Saint-Venant Kirchhoff law in the intermediate configuration which relates the second Piola-Kirchhoff stress tensor to the Green-Lagrange strain $\bar{\varepsilon}_{GL}^e = \frac{1}{2} (\bar{F}^e \bar{F}^e - \bar{I})$ as follows:

$$\bar{\pi} = \bar{C} : \bar{E}_{GL}^e$$  \hspace{1cm} (3)

where $\bar{C}$ is the elastic stiffness tensor.

The plastic deformation gradient rate $\bar{L}_p$ is computed as the sum of each slip system shear rate contribution as defined in equation (4). The latter one uses the usual Schmid tensor definition $\bar{S}_\alpha = \bar{m}_\alpha \otimes \bar{n}_\alpha$, where $\bar{n}_\alpha$ and $\bar{m}_\alpha$ are respectively the system plane normal and the slip direction of the slip system $\alpha$. $N_s$ is the number of slip systems.

$$\bar{L}_p = \bar{F}_p \bar{F}_p^{-1} = \sum_{\alpha=1}^{N_s} \dot{\gamma}_\alpha \bar{S}_\alpha = \sum_{\alpha=1}^{6} \dot{\gamma}_\alpha \bar{S}_\alpha + \sum_{\alpha=7}^{12} \dot{\gamma}_\alpha \bar{S}_\alpha + \sum_{\alpha=13}^{24} \dot{\gamma}_\alpha \bar{S}_\alpha$$  \hspace{1cm} (4)

Considering constant strain rate under steady state loading, the shear rate $\dot{\gamma}_\alpha$ is derived from the Orowan’s law (equation (5)) that relies on the dislocation velocity $v_\alpha$ and the mobile dislocation density $\rho_m^\alpha$ for the system $\alpha$.

$$\dot{\gamma}_\alpha = \rho_m^\alpha b v_\alpha$$  \hspace{1cm} (5)

In the thermally activated regime, the dislocation velocity depends on both the temperature and the resolved shear stress $\tau_\alpha$ defined hereafter.

To express the stress dependency of the dislocation velocity and ensure the energetic consistency of the proposed model, the work conjugate $\bar{M}$ of the plastic velocity gradient $\bar{L}_p$, known as the Mandel stress tensor, is introduced as follows:

$$\bar{M} = \det(\bar{F}^e) \bar{F}_e \cdot \bar{S} \cdot \bar{F}_e^T = \bar{F}_e^T \cdot \bar{F}_e \cdot \bar{S}$$  \hspace{1cm} (6)

The resolved shear stress $\tau_\alpha$ can now be expressed the Schmid tensor $\bar{S}_\alpha$:

$$\tau_\alpha = \bar{M} : \bar{S}_\alpha$$  \hspace{1cm} (7)

Due to the lack of information on physics-based deformation processes in UO$_2$, we used an empirical Arrhenius equation for dislocation mobility that depends on both shear stress and temperature as derived in [33], equation (8). More information about the mobility equation is provided Appendix 1.

$$v^\alpha = v_0 b \exp \left( - \frac{\Delta H_0}{k_b T} \right) \left( \cosh \left( \frac{\tau_\alpha}{\tau_0} \right) - 1 \right)$$  \hspace{1cm} (8)
Where \( v_D = 4.94 \times 10^{13} \text{s}^{-1} \) is the Debye frequency, \( b \) is the Burgers magnitude, \( k_B \) the boltzman constant, \( T \) the temperature, \( \Delta H_i^\alpha \) is the dislocation glide activation energy and \( \tau_0^i \) a critical shear stress. Here, \( i \) refers respectively to \{100\}, \{110\} or \{111\} modes. For \( i=\{100\}, \alpha \in [1;6], \) for \( i=\{110\} \alpha \in [7;12] \) and for \( i=\{111\} \alpha \in [13;24]. \) \( v_D \) is deduced from the Debye temperature \( \Theta_D=377 \text{K} \) [31-33], where \( v_D = \Theta_D /(h. k_B) \) with \( h \) the Plank constant.

2.2. Identification of the shear rate equation parameters

Parameters of equations (5) and (8) are defined using a reversed engineering process based on equation (9). Parameters \( \rho_m^\alpha, \Delta H_i^\alpha \) and \( \tau_0^i \) are adjusted to reproduce experimental CRSS measured during single crystal compression tests. Numerical CRSS \( \tau^\alpha \) are computed using the least square method and refer to a given temperature \( T \) and a resolved shear rate \( \dot{\gamma}^\alpha. \)

\[
\tau^\alpha = \tau_0^i \cdot \cosh^{-1}(1 + \frac{\dot{\gamma}}{0.5 \times \rho_m^\alpha b^2 e^{-\Delta H_i^\alpha / k_B T \cdot \Theta_D}})
\]

(9)

Assuming single slip and low strain conditions, \( \dot{\gamma}^\alpha \) of equation (5) is computed assuming the macroscopic strain rate \( \dot{\varepsilon} = 1 \times 10^{-4} \text{s}^{-1} \) with \( S = 0.5. \)

The identification process is performed using several sets of experimental CRSS that refer to \( \frac{1}{2}<110>\{100\} \) and \( \frac{1}{2}<100>\{110\} \) slip modes [6], [7], [9], [11]. See supplementary information for a detailed discussion about experimental data.

2.3. CPFE modeling of viscoplastic anisotropy

Single crystal compression tests of Sawbridge and Sykes [9] are used as reference to validate the proposed CPFE model and investigate the viscoplastic anisotropy. In both the experiment and the simulation, constant strain rate compression tests are performed at \( \dot{\varepsilon}=10^{-4}\text{ s}^{-1} \) and \( T=1600 \text{ K} \) for various orientations. In the following, the orientation of the compression axis (CA) in the crystal basis is defined using the two angles \( \phi \) and \( \Omega \) as proposed in the original study [9].

The constitutive model described in section 2.1 has been implemented in the \textit{MFront} integrator [37], [38] within the \textit{Cast3M} code [39] including an implicit quasi newton algorithm for the nonlinear mechanical equilibrium and a standard implicit newton scheme for the local nonlinear behavior. The fourth order elastic tensor (equation (3)) is derived from Hutching’s neutron scattering data that provide single crystal elastic constants for a large range of temperature [39]. The simulated sample has a parallelepiped shape (2.8 \( \times \) 2.8 \( \times \) 8 mm\(^3\)) and is completely meshed with 250 cubic (quadratic) elements with 20 integration points each to model the heterogeneous deformation of the sample. Uni-axial displacement loads in FE simulations are performed considering bottom and top surfaces sticked to the compression device. Free-boundary conditions are prescribed on the sample lateral surfaces in order to mimic experimental conditions. As the method for lattice rotation measures is not
detailed in Sawbridge and Sykes study, here we compute the average rotation of ten elements in the center of the sample. See supplementary information for more details on lattice rotation calculations.

3. RESULTS

3.1. CRSS

![Figure 1: CRSS versus temperature profiles for \( \frac{1}{2} <110> \{100\} \) and \( \frac{1}{2} <110> \{110\} \) slip modes. Experimental data are shown using symbols while blue and dashed red curves refer to equation (9) solutions respectively \{100\} and \{110\} slip modes.](image)

Table 1: Key parameters for the CRSS processing (equation (9)) of \( \frac{1}{2} <110> \{100\} \) and \( \frac{1}{2} <110> \{110\} \) slip modes under single slip conditions.

<table>
<thead>
<tr>
<th></th>
<th>( \frac{1}{2} &lt;110&gt; {100} )</th>
<th>( \frac{1}{2} &lt;110&gt; {110} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \rho^0 ) (m(^{-2}))</td>
<td>( 6.2 \times 10^{12} )</td>
<td>( 9.2 \times 10^{11} )</td>
</tr>
<tr>
<td>( \Delta H^0 ) (eV)</td>
<td>( 5.71 )</td>
<td>( 5.22 )</td>
</tr>
<tr>
<td>( \tau^0 ) (MPa)</td>
<td>( 1.35 )</td>
<td>( 4.78 )</td>
</tr>
</tbody>
</table>

Processing of equation (9) is shown in Figure 1 including CRSS for \( \frac{1}{2} <110> \{100\} \) and \( \frac{1}{2} <110> \{110\} \) slip systems. For \{100\}, Byron [6], Nadeau [7], Sawbridge [9], and Lefebvre [11] experimental data are used in the fitting procedure. At low temperature, \{100\} CRSS exhibit large discrepancies while they converge to an athermal stress of \( \tau^0 =18 \) MPa and a computed athermal transition temperature \( T_a =1750 \) K. For \{110\}, only Nadeau and Sawbridge and Sykes datasets are used to process the \{110\} mode leading to larger CRSS values when compared to \( \frac{1}{2} <110> \{100\} \). Assuming the same athermal stress for both modes, the athermal temperature is \( T_a =2250 \) K in \( \frac{1}{2} <110> \{110\} \). Both datasets exhibit extended thermally-activated regimes characterized by \( \Delta H^0 \{100\} =5.71 \) eV and \( \Delta H^0 \{110\} =5.22 \) eV. Derived mobile dislocation densities under single slip conditions are of \( 6.2 \times 10^{12} \) m\(^{-2}\) and \( 9.2 \times 10^{11} \) m\(^{-2}\) in \{100\} and \{110\} slip planes. These values are in relatively good agreement with usual measurements performed in the early stage of deformation. CRSS processed parameters are resumed Table 1.
3.2. CPFEM results

3.2.1. Mechanical behavior

CPFEM simulations are performed using rate equations (5) and (8) and Table 1 parameterization. Both \{100\} and \{110\} slip modes are considered and data are compared to Sawbridge and Sykes compression tests [9] in Figure 2. While some uncertainties exist for $\varepsilon \leq 1\%$, the stress-strain curves reproduce qualitatively the experimental profiles beyond 1\% strain. Sawbridge and Sykes [9] suggest that the low strain regime ($\varepsilon \leq 1\%$) refer to a so-called stage 0, that is commonly observed when the mobility of one dislocation character is higher than the others (see e.g. [41],[42]).

For the two single mode orientations, $\phi = 2^\circ$ and $\phi = 44^\circ$, the Schmid factor of the activated slip systems is close to 0.5 respectively for four $\frac{1}{2}[110]\{110\}$ systems and for the [110](001) slip system (see Table 2). For $\phi = 2^\circ$, the computed flow stress shows a slight increase but remains in good agreement with the experiment. Considering experimental discrepancies (see the original article of Sawbridge and Sykes [9]), stresses are also comparable to Sawbridge and Sykes data for $\phi = 44^\circ$.

For $\phi = 54^\circ$, strain is accommodated in the simulation as in the experiment by dislocation glide in $\frac{1}{2}[110]\{100\}$. While yield stresses are comparable, the computed flow stress is slightly lower.

For other $\phi$ orientations of Figure 2, simulated strain is accommodated by both slip modes. In addition, orientation dependency on the stress is well captured despite a large overestimation for $\phi = 13^\circ$. Furthermore, one can note that the hardening tendency observed for $\phi = 36^\circ$ and $\phi = 25.5^\circ$ is well-captured although no hardening term is introduced in the crystal plasticity model. This effective hardening is induced by the coupling effect between lattice rotation and viscoplastic anisotropy.

![Figure 2: Stress-strain of 1600 K UO2 single crystal compression tests. CPFEM (dashed line) is compared to experimental compression tests (full line) [9]. Computed data are shown using the engineering stress definition.](image-url)

In the experiment, heterogeneous glide can be deduced from lattice rotations as e.g. in the case of Sawbridge and Sykes [9]. On Figure 3, both experimental and CPFEM results show that CA rotates toward [001] when $\phi > 16^\circ$. For these orientations, the $\frac{1}{2}[110](001)$ slip system contributes mainly to the total shear inducing significant deformation heterogeneities and lattice rotations. For orientations close to $\phi = 0^\circ$, CA misaligns from [001] in the experiment while simulated rotation is almost negligible. The numerical result is
due to homogeneous glide on four ½<110>{110} slip systems while only two slip systems are activated in the experiment.

Table 2: Slip system observations in the experiments and computed slip system activity. Experimental data refer to Sawbridge and Sykes study [9]. \( \phi \) and \( \Omega \) orientations are depicted figures 2 and 3 respectively.

<table>
<thead>
<tr>
<th>Orientations</th>
<th>Experiment [9]</th>
<th>Simulation results</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \phi = 2^\circ )</td>
<td>Two ½&lt;110&gt;{110} slip systems</td>
<td><a href="011">011</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.5 ) &lt;br&gt; <a href="011">011</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.5 ) &lt;br&gt; <a href="101">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.5 ) &lt;br&gt; <a href="101">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.5 )</td>
</tr>
<tr>
<td>( \phi = 13^\circ )</td>
<td><a href="001">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 38% ; \gamma \approx 0.22 ) &lt;br&gt; 4 other ½&lt;110&gt;{110} slip systems&lt;br&gt; ( \gamma / \gamma_{\text{tot}} \approx 14% ; \gamma \approx 0.42 )</td>
<td></td>
</tr>
<tr>
<td>( \phi = 25.5^\circ )</td>
<td>One ½&lt;110&gt;{100} slip system</td>
<td><a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 60% ; \gamma \approx 0.39 ) &lt;br&gt; 4 other ½&lt;110&gt;{110} slip systems&lt;br&gt; ( \gamma / \gamma_{\text{tot}} \approx 8% ; \gamma \approx 0.36 )</td>
</tr>
<tr>
<td>( \phi = 36^\circ )</td>
<td>One ½&lt;110&gt;{100} slip system</td>
<td><a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 83% ; \gamma \approx 0.48 ) &lt;br&gt; +2 other ½&lt;110&gt;{100} slip systems&lt;br&gt; ( \gamma / \gamma_{\text{tot}} \approx 2.12% ; \gamma \approx 0.36 ) &lt;br&gt; +2 other ½&lt;110&gt; {110} slip systems&lt;br&gt; ( \gamma / \gamma_{\text{tot}} \approx 1.11% ; \gamma \approx 0.12 ) &lt;br&gt; +4 other ½&lt;110&gt;{110} slip systems&lt;br&gt; ( \gamma / \gamma_{\text{tot}} \approx 2.67% ; \gamma \approx 0.24 )</td>
</tr>
<tr>
<td>( \phi = 44^\circ )</td>
<td>One ½&lt;110&gt;{100} slip system</td>
<td><a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 90% ; \gamma \approx 0.5 ) &lt;br&gt; <a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 5% ; \gamma \approx 0.42 ) &lt;br&gt; <a href="100">011</a>: ( \gamma / \gamma_{\text{tot}} \approx 5% ; \gamma \approx 0.42 )</td>
</tr>
<tr>
<td>( \phi = 54^\circ )</td>
<td>{100} slip mode</td>
<td><a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 38% ; \gamma \approx 0.48 ) &lt;br&gt; <a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 31% ; \gamma \approx 0.47 ) &lt;br&gt; <a href="100">011</a>: ( \gamma / \gamma_{\text{tot}} \approx 31% ; \gamma \approx 0.47 )</td>
</tr>
<tr>
<td>( \Omega = 0^\circ )</td>
<td>½&lt;110&gt;{100} and ½&lt;110&gt;{111} multi-slip</td>
<td><a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.35 ) &lt;br&gt; <a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.35 ) &lt;br&gt; <a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.35 ) &lt;br&gt; <a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 25% ; \gamma \approx 0.35 )</td>
</tr>
<tr>
<td>Intermediate ( \Omega ) orientations</td>
<td>Short traces on (001) and (111) planes which are mutual cross-slip planes for a [101] slip vector</td>
<td><a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 36% ; \gamma \approx 0.34 ) &lt;br&gt; <a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 29% ; \gamma \approx 0.32 ) &lt;br&gt; <a href="010">101</a>: ( \gamma / \gamma_{\text{tot}} \approx 17% ; \gamma \approx 0.28 ) &lt;br&gt; <a href="001">110</a>: ( \gamma / \gamma_{\text{tot}} \approx 24% ; \gamma \approx 0.27 )</td>
</tr>
</tbody>
</table>

For \( \phi \) close to 16°, CA misaligns with the \( \phi \) axis in the experiment while it rotates toward the [001] axis in the simulations. No experimental slip characterization is made for this orientation, but the computed lattice rotation is correlated to a significant activity of the ½ [110](001) slip system, as already seen before for \( \phi > 16^\circ \) (see Table 2). Therefore, it can be assumed that additional slip systems are involved in the experiment.

For orientations close to intermediate \( \Omega \), the CA rotates toward the [011] axis in the experiment while simulations show a rotation toward [001] with a significant activity of the ½[110](001) slip system comparable to the previous case. Finally, for CA close to [011], computed lattice rotations are negligible due to homogeneous glide on four {100} slip
systems (see Table 2). However, CA rotates toward [001] in the experiment. Here, experimental slip observations emphasize the activation of \( \frac{1}{2}<110>|\{111| \) slip systems that may contribute to lattice rotations for [011] and intermediate \( \Omega \) orientations.

Finally, when dislocation glide occurs on a single mode (typically for \( \phi \approx 0^\circ \) and \( \phi \approx 45^\circ \)), both the stress response and lattice rotation predictions are in good agreement with experimental measurements. For intermediate \( \phi \) orientations (\( \phi > 16^\circ \)), while the computed and experimental stress profiles show some discrepancies, lattice rotations are in good agreement due to a comparable slip mode activity.

To better understand the impact of the different slip modes on the strain accommodation, the following section will focus on the flow stress anisotropy and particularly based on additional simulations performed along \( \Omega \) orientations.

3.2.2. Viscoplastic anisotropy and shear contributions at 1600 K

Stresses (at 2% strain) versus lattice orientations \( \phi \) are shown on Figure 4(a) and numerical predictions are in good agreement with experimental data. The plastic shear distribution per slip mode is shown Figure 4(b) where the \{110\} mode accommodates the entire deformation for \( \phi = 0^\circ \). Up to \( \phi = 10^\circ \), the \{100\} mode handles less than 25% of the total strain. The latter mode becomes dominant beyond \( \phi = 13^\circ \) and accommodates all the strain between \( \phi = 40^\circ \) up to CA=[111]. Those results are consistent with experimental slip traces observations.

On the other hand, simulation results for \( \Omega \) orientations presented on Figure 4(c) and (d) are in contradiction with the reference tests. For \( \Omega < 25^\circ \), stress is underestimated (~50%) when compared to the experiment. The simulation is characterized by \{100\} multi-slip that does not agree with experimental observations which underlines \{100\}+\{111\} multi-slip [9]. For
the same orientations, two other experimental studies show multi-slip only on \{100\} at strains lower than 0.3% in addition to cross-slip from $\frac{1}{2}<110>$\{100\} to $\frac{1}{2}<110>$\{111\} slip systems [11],[12]. All these studies conclude that, plastic deformation occurs first in $\frac{1}{2}<110>$\{100\} slip systems and then, cross slip happens in \{111\}. For $\Omega \in [20;35]$, the computed stress increases up to $\sim 125$ MPa while measurements exhibit a steady-state of about 110 MPa. In the experiment, slip traces observations are non-crystallographic and follow the Maximum Resolved Shear Stress Plane (MRSSP) while simulations are characterized merely by shear in the \{100\} (\( \geq 75\% \)) and \{110\} (\( \leq 25\% \)) modes.

![Stress anisotropy](image1.png)

**Figure 4**: Stress anisotropy. (a)-(c) flow stress at 2% strain versus $\phi$ and $\Omega$ orientations. (b)-(d) Computed normalized plastic shear contribution for each slip mode ($\gamma^{(100)}/\gamma_{tot}$ in blue and $\gamma^{(110)}/\gamma_{tot}$ in red) versus $\phi$ and $\Omega$ orientations.

4. DISCUSSIONS

The CPFE model reproduces correctly most of the CRSS anisotropy of UO$_2$ single crystal along $\phi$ and is also able to restitute qualitatively the stress-strain behavior (after stage 0) for most of the available experimental CA. However, three particular ranges of orientations remain under debate (i) $\phi \in [15;40]$ where the stress is overestimated, (ii) the region close to $[\bar{1}11]$ where stress is underestimated and, (iii) $\Omega < 25^\circ$ domain where the stress is underestimated. Theses discrepancies may have different origins as e.g. dislocation interactions, cross-slip and/or glide in $\frac{1}{2}<110>$\{111\} slip systems or composite slip. Indeed, the \{111\} slip hypothesis is supported by numerous evidences in the literature [5],[9],[11]-[14]. As a consequence, accounting for the \{111\} slip mode in the simulation seems to be of prime
importance. As pure $\frac{1}{2}<110>\{111\}$ slip is not referenced in the literature, the reverse engineering methodology developed section 2.2 cannot be applied in this case. Therefore, a parametric study focusing on $\frac{1}{2}<110>\{111\}$ is developed in what follows. Derived from the original model that accounts for $\frac{1}{2}<110>\{100\}$ and $\frac{1}{2}<110>\{110\}$ (labelled case 1), two additional cases that also include $\frac{1}{2}<110>\{111\}$ slip systems are set. They are labelled case 2 and case 3 in the following. Relative mode activation sensitivity is described in Table 3. The model case 2 accounts for $\{111\}$ slip using the same activation parameters than for $\{110\}$. This assumption should lead to a restricted activation of the $\{111\}$ mode. With a less severe hypothesis ($\{111\}$ slip slightly softer than $\{110\}$), Sawbridge and Sykes [9] underline that slip on $\frac{1}{2}<110>\{100\}$ slip system is more favorable than in $\frac{1}{2}<110>\{111\}$ for $\Omega$ orientations. Thus, case 3 is designed to check if a combination of $\{100\}$ and $\{111\}$ modes could improve the results, especially for $\Omega$ orientations. As a consequence, the activity of $\{100\}$ has been further reduced decreasing the mobile dislocation density on every $\frac{1}{2}<110>\{100\}$ slip systems down to $\rho_m^s = 1 \times 10^7$ m$^{-2}$ to promote $\{111\}$ glide. Finally, other $\{111\}$ activation parameters are equivalent to those of $\{110\}$ but with a $\tau_0^{\{111\}} = 3.50$ MPa (against 4.78 MPa for $\{110\}$ mode). This new parameter set influences the CRSS as shown in Table 3. Results for all cases are provided in Figure 5.

Considering $\{111\}$ as a hard mode (case 2) leads to a slight variation of the mechanical response (figure 5(a) and (c)) compare to case 1. In particular, $\{111\}$ accommodates up to 25% of the plastic shear for $0^\circ < \phi < 30^\circ$ and $25^\circ < \Omega < 40^\circ$ orientations. This result is consistent with several experimental observations where $\{111\}$ slip traces were observed for CA close to [001].

Table 3: Description of the model cases accounting for $\frac{1}{2}<110>\{111\}$ slip systems. CRSS are provided for each slip mode. They are computed at T=1600 K and $\dot{\varepsilon} = 10^{-4}$s$^{-1}$ using equation (9) and the maximum Schmid factor assumption.

<table>
<thead>
<tr>
<th>Model</th>
<th>Slip mode sensitivity</th>
<th>CRSS (MPa)</th>
<th>100</th>
<th>110</th>
<th>111</th>
</tr>
</thead>
<tbody>
<tr>
<td>case 1</td>
<td>soft ${100}$ + hard ${110}$</td>
<td>21.7</td>
<td>68.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>case 2</td>
<td>soft ${100}$ + hard ${110}$ + hard as ${111}$</td>
<td>21.7</td>
<td>68.4</td>
<td>68.4</td>
<td></td>
</tr>
<tr>
<td>case 3</td>
<td>harder ${100}$ + intermediate ${111}$ + hard ${110}$</td>
<td>39.8</td>
<td>68.4</td>
<td></td>
<td>50.1</td>
</tr>
</tbody>
</table>

For the case 3, strain is produced mainly by the $\{111\}$ and $\{100\}$ modes for $\Omega < 25^\circ$. This result is consistent with experimental slip traces observations made for CA close to [001]. Furthermore, this model improves lattice rotations when compared to the experiment (see Figure 3(b)) what confirms the role of $\frac{1}{2}<110>\{111\}$ slip systems in the mechanical behavior of UO$_2$, especially for orientations where CA is in the center of the standard triangle. Nevertheless, the activity of $\frac{1}{2}<110>\{111\}$ slip systems increases too much and the $\{110\}$ mode is nearly off, even close to CA=[001] (Figure 5(a)). Furthermore, CRSS become lower than expected for these orientations.
On the other hand, the \{100\} mode remains active especially close to CA=[\overline{1}11], but the stress level increases due to the changes of \{100\} activation parameters leading to significant discrepancies when compared to experimental data. Finally, along the \(\phi\) direction, the agreement with Sawbridge and Sykes [9] stress measurement is only reached for \(\phi=16^\circ\) i.e. at the intersection between \(\phi\) and \(\Omega\) directions. However, lattice rotations are still not consistent with experiments for this orientation where CA rotates toward [\overline{1}11] due to the larger activation of the \{111\} mode.

Overall, \(\frac{1}{2}<110>\{111\}\) slip systems improve stress predictions along \(\Omega\) but have an opposite influence along \(\phi\) orientations. Therefore, \{111\} slip appears as a key feature to obtained reliable lattice rotations and CRSS in the central part of the standard triangle only. This latter observation suggests that \(\text{UO}_2\) slip system activity might be more complex than expected. Among others, cross-slip responsible for probable composite glide between \{100\} and \{111\} or \{110\} and \{111\} systems (depending on the orientation) should be accounted in the future, as well as the strain hardening induced by dislocation interactions between the different slip systems. Indeed, forest hardening should be particularly effective in the cases of multi-slip orientations (e.g. \(\Omega = 0^\circ\) and \(\phi = 54^\circ\)) that include several \(\frac{1}{2}<110>\{100\}\) slip systems characterized by an athermal transition temperature \(T_a\) very close from Sawbridge and Sykes [9] effective temperature.

![Figure 5: Stress anisotropy for the three model cases accounting for \{111\} slip. (a)-(c) flow stress at 2\% strain versus \(\phi\) and \(\Omega\) orientations. (b)-(d) computed normalized shear contribution for each slip mode versus \(\phi\) and \(\Omega\) orientations (\(\gamma^{(100)}/\gamma_{\text{tot}}\) in blue, \(\gamma^{(110)}/\gamma_{\text{tot}}\) in red and \(\gamma^{(111)}/\gamma_{\text{tot}}\) in in yellow).](image-url)
5. CONCLUSIONS

The goal of this study was to build a viscoplastic model for UO$_2$ single crystal based on the dislocation glide mechanism to reproduce the plastic anisotropy of UO$_2$. The activation parameters for the ½<110>{100} and ½<110>{110} slip systems have been adjusted on experimental CRSS vs. temperature data and integrated into a CPFEM framework. Then, compression simulations have been performed and compared to Sawbridge and Sykes experimental work [9]. Accounting for the two slip modes, the plastic anisotropy is correctly assessed only in the case of single-mode orientations i.e. when only one of the two slip modes is activated. To improve the model, ½<110>{111} slip systems have been integrated based on several experimental observations. As a result, accounting for ½<110>{111} slip systems greatly improves stress predictions and lattice rotations in central part of the standard triangle but other orientations remain under debate. This confirms that ½<110>{111} slip systems may play an important role in UO$_2$ single crystal mechanical response and related lattice rotations, together with other slip modes. In particular, cross-slip, combined slip and dislocation interactions appear as key processes to investigate in the future in order to better understand UO$_2$ single crystal mechanical properties.

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APPENDIX 1

Empirical power laws (as e.g. equation (10) [43]) are commonly used to compute mean dislocation velocities and describe viscoplasticity at the grain scale. This kind of equation relies on dislocation glide in slip system $\alpha$ for $\tau^\alpha$ reaching a critical value $\tau_0^\alpha$. Thus, the viscosity is given by the stress exponent $n$, i.e. the strain rate sensitivity is reduced when increasing $n$, up to a purely plastic behavior for high $n$ values.

$$\dot{\gamma}^\alpha = \rho_m b v_0^\alpha \exp \left( - \frac{\Delta H_0^\alpha}{k_b T} \right) \left| \frac{\tau^\alpha}{\tau_0^\alpha} \right|^n \text{sign}(\tau^\alpha)$$  \hspace{1cm} (10)

Usually, $n$ is defined for a given stress and temperature range and the use of a single $n$ value for larger stress range is proscribed. To avoid the identification of stress-temperature dependent exponent, an exponential form such as a hyperbolic sine equation (11) can be used [44-46]. Both equations (10) and (11) are equivalent for a given stress domain assuming different values of $\tau_0^\alpha$.

$$\dot{\gamma}^\alpha = \rho_m b v_0^\alpha \exp \left( - \frac{\Delta H_0^\alpha}{k_b T} \right) \sinh \left( \frac{\tau^\alpha}{\tau_0^\alpha} \right) \text{sign}(\tau^\alpha)$$  \hspace{1cm} (11)
Finally, the hyperbolic sine law (11) can be replaced by equations (5) and (8) to avoid dislocation glide at low stress and high temperature, in the void diffusion creep regime. This formulation originally introduced by Soulacroix [33] cancels the linear stress dependency†, given by exponential or hyperbolic sinus law, for low stresses.

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† A first order Taylor’s development around 0 leads to a linear function for exp(x) and sinh(x) when it gives a null function for cosh(x)


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