Numerical study of crystalline plasticity : measurements of the heterogeneities due to grain boundaries under small strains

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Among all the methods developed to characterize the local behaviour in polycrystalline materials, the experimental determination of the fields inside grains requires to cut-off slices of the material to enable three-dimensional measurements. That is why such measurements are often made on multicrystals with one or two grains along the shortest direction of the piece of material. Numerical representation of the local behaviour inside an aggregate is possible, through homogenization or finite elements computations (12, 26, 19). The heterogeneities displayed with these approaches have been confirmed by comparison to experimental measurements. From the numerous studies concerning the intragranular description of the mechanisms of deformation, we know that the behaviour inside a grain is governed by dislocation arrangements that may cause greatly varying responses from point to point. Dislocation cells and walls form intragranular structures with sizes-spacing depending upon the distance to a grain boundary, a so-called geometrically necessary boundary (4) or even a free surface (17). Such features have been introduced in many non-local modellings with more or less accordance to physics : strain gradient plasticity (27, 1), Cosserat plasticity (14), explicit construction of dislocation structures (24) ... As well as experimental characterizations, these models, as long as they intend to represent intragranular heterogeneities, are restricted to structures containing few crystals.

With the use of FE parallel computations and with a classical approach to crystalline plasticity under small strains, it is possible to describe the intragranular heterogeneity in a model of three-dimensional polycrystals (25, 6, 7). The results thus deal with a huge amount of points inside each grain – largely enough to observe variations of gradients inside grains – where the interactions / slip patterns are three-dimensional. This is a fundamental requirement to justify performing statistical considerations : the number of individuals – points – is large enough. Another requirement is that the medium be isotropic so that the analysis be valid in any direction of space. This condition is respected as it has been checked in previous works with similar number of grains and the same material (5, 6). Consequently, all the points can be regarded as equivalent in any direction of

A finite element 3D polycrystalline aggregate made of 216 grains is subjected to a mean axial deformation up to 1 %. The heterogeneities inside the polycrystal are due to the crystallographic orientations of the grains – this appears at the scale of the mean responses per grain – and to the local interaction between neighbouring grains – this appears at the intragranular scale. The intragranular heterogeneity is measured as a function of the distance to the grain boundary, inside a single grain and for all the points of the microstructure. It is characterized by dispersions of the local responses due to grain boundaries. These measurements correspond to dispersions – variances – ; the statistical errors have also been determined.
Étude numérique de la plasticité cristalline : mesures des hétérogénéités dues aux joints de grains en petits déformations

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Cette étude présente une analyse par éléments finis de la plasticité polycristalline en petites déformations, avec une attention particulière portée sur les hétérogénéités de contraintes et déformations, tant intragranulaires que de grain à grain, dans le but de caractériser les effets mécaniques des joints de grains.

L’analyse par éléments finis offre l’avantage, par rapport à une démarche expérimentale, de fournir des informations sur les champs de contraintes et déformations sur un volume d’étude véritablement tridimensionnel et contenant suffisamment de grains pour constituer un polycristal.

Les précédents travaux menés avec cette approche numérique nous ont permis de déterminer une configuration minimale (en termes de taille de maillage, de nombre d’éléments par grains...) pour l’étude des hétérogénéités intragranulaires, configuration qui a été retenue pour cette étude : nous utilisons un maillage de 36*36*36 éléments finis de type briques à 20 noeuds et à intégration complète.

Dans ce maillage nous faisons figurer 216 grains, représentés par des polyèdres de Voronoï. À chaque grain est attribuée une orientation cristallographique dont il sera tenu compte dans le modèle de comportement de chaque grain pour la détermination des réponses locales en contraintes et déformations : des lois de comportement viscoplastiques avec écrouissage isotope et cinématique sont attribuées à chaque système de glissement d’un cristal, et la déformation du cristal est déduite via la loi de Schmid (équations [1] à [5]). Le polycristal est soumis à une traction à 1 % de déformation axiale.

Les hétérogénéités des champs mécaniques (visibles fig. 1) sont donc la conséquence de la différence de comportement des grains (ceci apparaît notamment à l’échelle de la réponse moyenne par grain) combinée aux interactions locales entre grains voisins. Les grains étant nombreux et ayant des géométries différentes, caractéristiques d’un milieu désordonné, le nombre de points d’intégration par grain étant très élevé (de l’ordre de 6 000), nous avons choisi de classer tous les points d’intégration du maillage en fonction de leur distance par rapport aux joints de grains, ceci afin de représenter :

- la réponse moyenne – en contrainte par exemple,
- la dispersion des variables mécaniques (équations [6] et [7]),
- l’erreur relative (équation [8]) en fonction de la distance aux joints de grains.

Les figures 1a et 1b apportent des informations sur les distances maximales aux joints de grains et sur le nombre de représentants (i.e. le nombre de points d’intégration) dans chaque intervalle de distances aux joints de grains où nous choisissons de recoller les variables mécaniques. Nous vérifions ainsi les domaines où il est permis d’effectuer des mesures à caractère statistique.

Les figures 3, 4 et 5 portent sur les réponses en termes de contrainte équivalente et de déformation équivalente dans un seul grain du polycristal ; elles présentent respectivement la réponse moyenne en fonction de la distance, la dispersion relative et l’erreur relative due au nombre de représentants par intervalle. Les figures 6, 7 et 8 présentent le même type de diagrammes que les figures 3, 4 et 5, pour des mesures prises sur tous les points du polycristal, indépendamment de leur orientation cristallographique.

Il apparaît qu’il n’y a pas d’effet de la distance aux joints de grains sur la réponse moyenne en contrainte ou en déformation équivalente : même si on observe une légère variation de ces quantités en fonction de la distance au joint de grain, elle n’est pas significative ; on ne peut donc pas dire que les contraintes sont plus élevées ou plus basses à proximité d’un joint de grain par rapport au cœur du grain. Par contre, la dispersion des variables apparaît nettement affectée par la proximité à un joint de grain, tant pour la contrainte équivalente (de l’ordre de 17 %) que pour la déformation (de l’ordre de 30 %). Retenons que cette dispersion, du fait du cadre de notre modélisation, n’est due qu’à l’anisotropie de la plasticité locale, combinée aux hétérogénéités structurelles du polycristal.
space. In these works, we have also checked that the loa-
ding boundary conditions of the structure had negligible
effect on the polycrystal effective response and on the
mean response per grain. Only a small layer of one-grain-
depth is affected by the boundary conditions (for a same
tensile test, homogeneous strain boundary conditions were
used, as well as mixed boundary conditions where the mini-
num necessary displacements were imposed). There thus
might not be any distinction to make between points having
different positions toward the boundary of the structure – at
least, once all the points are mixed together through avera-
ging operations, the slight effect due to structure bounda-
ries vanishes –. Furthermore, we can choose not to make
any distinction between points of different crystallographic
orientation and classify the points regarding their minimum
distance to a grain boundary. This distance thus becomes
the only parameter considered for the analysis of
stress/strain heterogeneities.

In the following, tools for the modelling of the behaviour
of polycrystalline aggregates and an example of computa-
tion (section “Polycrystal modelling”) are briefly presented.
Section “Measurements of heterogeneities” deals with the
measurements taken from a simulated 216-grain-polycrys-
tal subjected to mean tensile loading: first the definitions
of the measurements are given and then the resulting mea-
surements are provided for the case of our polycrystal.
They consist in:

– the mean response in sets/classes of points being at a
same distance to a grain boundary,
– the dispersion in these sets of points,
– the estimated error associated to the number of repre-
sentative points in each set.

Results are then discussed in section “Results and observations”.

POLYCRYSTAL MODELLING

Single crystal

It is assumed that slip is the predominant deformation
mechanism and that Schmid’s law is valid. The resolved
shear stress can then be used as a critical variable to eva-
luate the inelastic flow. A viscoplastic framework is chosen,
in order to avoid the problems related to the determination of
the active slip systems in plastic models. A threshold is intro-
duced both in positive and negative directions on each slip
system: twelve octahedral slip systems will be used for FCC
materials. Two variables are defined for each slip system s,
r^s and x^s, corresponding respectively to isotropic hardening
(expansion of the elastic domain), and kinematic hardening
(translation of the elastic domain (11). A system will be
active provided its resolved shear stress r^s is greater than x^s
+ r or less than x^s - r, and the slip rate will be known as
long as stress and the hardening variables are known. The
state variables used to define the evolution of r^s and x^s are
the accumulated slip v^s for isotropic hardening and the
variable v^s for kinematic hardening. Knowing the stress ten-
sor \sigma applied to the grain s, the resolved shear stress for
system s can be classically written according to [1], v^s and
v^s being respectively, for the system s, the normal to the slip
plane and the slip direction in this plane. The hardening
variables x^s and r^s can then be expressed as a function of v^s
and v^s following [2], their current values allowing us to com-
pute the viscoplastic slip rate \gamma^s; the viscoplastic strain rate
tensor \dot{\varepsilon}^s [3], and the hardening rules ([2.1] and [5]). The
present formulation gives a saturation of the hardening in
both monotonic and cyclic loading, and takes into account
the interactions between the slip systems, through matrix
h_{rs}, as in (18). Nine material-dependent coefficients are
involved in the model (E, v, K, n, c, d, R_0, Q, b).

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Figure 1 – a) Polycrystalline aggregate made of 200 grains represented by 32^3 full integration elements; b) amount of plastic slip on the
contour of the polycrystal – INCO600 – loaded at 1.5 % axial strain with homogeneous boundary conditions.

Figure 1 – a) Agrégat polycristallin à 200 grains représenté par 32^3 éléments briques à intégration complète; b) quantité de glissement plastique
sur le contour du polycristal – INCO600 – sous chargement en déformation axiale à 1,5 %, avec des conditions homogènes sur le contour.
Such a formulation (10, 21) is an extension of the classical crystallographic approach for single crystal modelling in plasticity or in viscoplasticity (see for instance (28, 20, 3). It has been extensively used for single crystal modelling including Finite Elements simulations (22, 9). The material coefficients correspond to INCO600:

- isotropic elastic behaviour: \( E = 196,000 \text{ MPa}, \quad n = 0.3 \);
- viscous effect in (2.1) and (5): \( K = 10 \text{ MPa.s}^n, \quad n = 25 \);
- kinematic hardening in (2):
  - \( c = 1,600 \text{ MPa}, \quad d = 40 \);
- isotropic hardening in (2) and (3): \( R_0 = 111 \text{ MPa}, \quad Q = 35 \text{ MPa}, \quad b = 7 \).

Computation and as-given results

The constitutive laws for single crystals are applied to each integration point of a FE structure representing a polycrystal. Each grain is made of a set of Gauss points that represents a Voronoi polyhedron (see fig. 1a for illustration and (6) for the details about Voronoi polyhedra generation – contribution due to Decker and Jeulin, LMM-ENSMP). The dependence of our results at different scales on the mesh refinement has been determined in (8). At least, 4*4*4 20-node-full-integration-elements per grain must be used to expect a correct result at the intragranular scale. The results produced in the following analysis deal with a structure made of 36*36*36 elements (1,259,712 integration points). For such a big structure, we have resorted to parallel computing on 32 processors of the Linux PC Cluster of the “Centre des Matériaux (École des Mines de Paris)” : the structure was divided into 32 sub-domains computed separately and balanced through the FETI method at each increment of the load (13, 25). As an example of the results obtained from such computations, the contour of the amount of plastic slip after 1.5 % axial strain on a 200-grain-polycrystal is shown on figure 1b. It displays the large variation in the local values at the intragranular scale. But our purpose here is to quantify those variations; it is referred to (5, 7) for a detailed analysis of as-given results. In the following, the analysis focuses on results of an axial tensile test to 1 % total strain performed on a 216-grain-polycrystal.

**MEASUREMENTS OF HETEROGENEITIES**

Definitions

Each point of the structure is an individual which has a proper set of values, one for each mechanical variable \( e_{eq} \) and \( s_{eq} \) will be treated in the following). Collecting all the points of a grain or of the whole structure at a same distance to the grain boundary ensemble, we build classes of individuals.
one class per mechanical variable, that we consider to be
discrete random variables. Then we can try to find an esti-
mator for each variable, and the related dispersion – square
root of the variance – and error. The most direct estimator
suited to our case is the esperance, i.e. the normal avera-
ging over all the points at a same distance to a grain boun-
dary. With the assumption that this estimator is correct, we
compute the relative dispersion (7) around the estimation
and the relative error (8) that is made by choosing the aver-
age as estimator.

Let us first call $X_{d_i}$ the discrete random variable defined by
the set of responses, either $\varepsilon_{eq}$ or $\sigma_{eq}$, at a given distance $d_i$
to the grain boundary ensemble :

$$X_{d_i} = \{x_k\}_{k=1,N(d_i)}$$

where $N(d_i)$ is the number of points at the distance $d_i \pm
(d_i - d_{i-1})/2$.

The absolute dispersion for this variable $X_{d_i}$ is the square
root of the variance :

$$D_{abs}(d_i) = \sqrt{\frac{\sum_{k=1}^{N(d_i)}(x_k - \langle X_{d_i} \rangle)^2}{N(d_i)}}$$

where

$$\langle X_{d_i} \rangle = \frac{\sum_{k=1}^{N(d_i)} x_k}{N(d_i)}$$

is the mean value of $X_{d_i}$.

The relative dispersion is the dispersion normalized by the
mean value $\langle X_{d_i} \rangle$ :

$$D_{rel}(d_i) = \frac{D_{abs}(d_i)}{\langle X_{d_i} \rangle}$$

Relative error associated to the number of representatives :

$$E_{rel} = \frac{D_{abs}}{\langle x_k \rangle_{d_i}} \cdot \frac{1}{\sqrt{N(d_i)}}$$

The measurements are made for all the points of a single
grain and then considering the points of the structure all-
together, without any care on their grain belonging. In the
second case, there are points of a same class (same dis-
tance to a grain boundary) that have different distances to
their respective grain centers. In other words, the maximum
distance to a grain boundary varies from grain to grain,
which is illustrated on figure 2a that gives the distribution
among grains of this maximum distance to a grain boundary.
But doing so, one makes estimations on larger sets of indi-
viduals than if a single grain were treated. The total number
of points of the structure in each class of distance to a grain
boundary is given on figure 2b. As for other graphs featuring
the distance to a grain boundary, the measurements are per-
formed in intervals of distances to the grain boundary having
two different sizes : 0.01 (the set of distances then corres-
ponds nearly to the set of every possible discrete distance)
and 0.1 which is less than one tenth of the minimum dis-
tance to a grain boundary. It is thus seen that measurements
for each possible discrete distance shall be considered
carefully if the distance to the grain boundary exceeds 2.5
whereas measurements in intervals sized 0.1 do not suffer
from lack of representatives until the distance 3 is attained.

The distance from an integration point to the grain boundary
is defined as the distance between the point concerned and
the closest integration point located along the grain boun-
dary inside the same grain. For this reason, for any integra-
tion point of the structure, the minimum distance to the grain
boundary is necessarily 0.

Results and observations

In a single grain

First, von Mises equivalent stress and strain inside a single
grain are considered on figure 3 where all the local responses
are given as functions of the distance to the grain boundary,
together with the normal average response per interval of
distance to the grain boundary. Apart from the increasing
variation of the responses as a grain boundary is approached, it is noticeable that the normal average made in intervals of distance does not vary much from grain center to grain boundary (about 10% of variation in strain and 5% in stress). If the normal average over all the points of a class is taken as the estimator, on figure 4 we see that there is an increasing dispersion from the grain center to the grain boundary for both stress and strain. But it is greater with strain (about 30%) than with stress (about 5%). The associated error (on fig. 5) shall be interpreted carefully: it does not vary although an increase could be expected around the grain center where number of representatives becomes very low: the largest errors are even reached on points far from grain center. So we actually just can deduce that the dispersions measured have a statistical validity, at least for distances less than 2.5 where stress and strain obviously appear to be largely dispersed among a large number of representatives (fig. 3). For longer distances, the number of representatives becomes very low and in the extreme limit, there is just a single point per class.

In the whole structure

The same plots of 1) the mean response per class of points, 2) the dispersion, 3) the error, determined from all the points of the structure, are given in figures 6, 7, 8. Figure 6 confirms the tendency featured by the single grain analysed: there is no significant variation of the mean behaviour with the distance to the grain boundary. There is just a slight 6% monotonous decrease in strain from the distance 0.5 to the distance 2.5. About stress, one can observe a tendency to increase in a monotonous manner in the interval [0.7-2.5], but it is too low an increase to be a significant effect (about 1.5% increase in stress...).

Moreover, from the analysis of another polycrystal of INCO600 containing 200 grains, it has been found varying signs of the variations of stress/strain from grain centers to grain boundaries (5). These variations were also very low so...
Figure 6 – a) $\varepsilon_{eq}$ et b) $\sigma_{eq}$ (MPa) en fonction de la distance au joint de grain, déterminés à partir des réponses en tout point de la structure.

Figure 7 – Dispersion relative de $\varepsilon_{eq}$ a) et $\sigma_{eq}$ b) en fonction de la distance au joint de grain, déterminée à partir des réponses en tout point de la structure.

Figure 8 – Erreur relative sur $\varepsilon_{eq}$ a) et $\sigma_{eq}$ b) en fonction de la distance au joint de grain, déterminée à partir des réponses en tout point de la structure.
there is no conclusion to draw from the slight variations observed. Nevertheless, from a detailed analysis of the mechanisms of slip (number of slip systems having a slip (resp. resolved shear stress) superior to a fixed value, number of systems having slip (resp. resolved shear stress) superior to a certain proportion of the maximum slip (resp. resolved shear stress) ...), it has been observed qualitative variations with distance to a grain boundary consistent with the results of the analysis of free surface effect: a free surface, compared to the behaviour in the bulk, favours higher strains and lower stresses by allowing activated slip systems to reach a great amount of slip; by contrast, a grain boundary tends to activate more slip systems while limiting the amount of slip on these systems. The analysis can actually not be restricted to the simple consideration of stress/strain and number of activated slip systems: there could be no effect on the stress/strain while there is an effect on the mechanisms of plasticity; the analysis requires to clearly differentiate the ways a slip system can be defined to be active. So, as a conclusion to the previous observations, one just can say that there is no significant effect of the grain boundaries on the mean response of points at a same distance to the grain boundaries. Unlike the widespread opinion that grain boundaries act as hardening layers that reinforce the behavior of each grain and that are taken to be responsible for the hardening of a polycrystal by comparison to single crystals alone (16, 23, 2, 15), there is no hardening nor softening effect due to the grain boundaries in our results. Some of these characteristics have been observed in the experimental measurements of Raabe et al (26) on a quasi 2D polycrystalline layer of aluminium strained at 8%: a spectrum of -87.5% to +87.5% deviation from the average value has been measured; some regions near grain boundaries correspond to strain localization whereas some others do not.

The quantitative dispersion and the qualitative variation of the dispersion with the distance to a grain boundary (fig. 7) are also confirmed: 30% of dispersion in strain along the grain boundaries and 17% in stress i.e. about half the dispersion in strain). From figure 2b we know that there just remain about 60 grains (a quarter) having a maximum distance to their boundary superior to 2.5. Those 60 grains occupy the volume fraction 90/216 i.e. 42% of the whole structure. One reasonably may suppose that the behaviour inside these grains might not be so particular that their responses are not significant of a "normal" behaviour, especially since they are the biggest grains, those who offer the best conditions for statistical considerations. Accordingly, one can bear on the measurements made between the distance 0 to 2.5: the remaining part of distances is representative of the behaviour in the core of the biggest grains. So a linear variation of the dispersion of strain from 30% at the grain boundary to 20% at distance 2.5 is observed. In stress, this variation goes from 17 to 12.5%, nearly linearly as well.

The error arising from the number of discretization points confirms that one can lay on our measurements for the major part of intragranular regions, at least for distances from 0 to 2.5: it reaches about 2% in strain and 1% in stress for a distance 2.5. Above this distance, the phenomenon expected in the single grain is present: the error remarkably increases, especially when it is measured in intervals sized 0.01. Indeed, the individuals of a class made from all the points of the structure have a priori a more pronounced random character than the points of a class inside a single grain: the behaviour in a grain may be significantly different from the one of another grain (for 1% total axial strain on this polycrystal, the mean response per grain varies from 250 to 500 MPa in stress and from 0.06 to 0.14 in strain) whereas two points of a same grain located near the center have no reason to vary greatly around the response at the center.

**CONCLUSION**

It has been shown that, in a purely local mechanical modeling of the behaviour of polycrystal, where no variation of the constitutive behaviour with position towards a grain boundary or any type of interacting region is accounted for and where only the geometrical arrangement of crystals generates intragranular heterogeneities, the major effect of the grain boundaries is the large dispersion that they cause; only the anisotropy of plasticity combined to structural heterogeneity is responsible for this in our case. Grain boundaries may favour localization in some places but no localization appears by average: grain boundaries do not harden nor soften the behaviour in their surroundings.

**references**


