

Review of Mechanisms of Nanograin Generation

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Abstract. A brief review of the theoretical models which describe specific plastic deformation mechanisms of nanograin generation in nanocrystalline and ultrafine-grained materials is presented. In the framework of the models, formation of new nanograins and nanograin chains occur through stress-driven splitting and migration of grain boundaries near crack tips and grain boundary structures containing disclination dipoles and quadrupoles. Stress-driven migration of mobile grain boundaries serves as a specific plastic deformation mode and results in formation of new nanograins and nanograin chains.

1. INTRODUCTION

Ultrafine-grained (UFG) and nanocrystalline materials (NC) characterized by superior strength represent the subject of rapidly growing research efforts motivated by a wide range of their current and potential applications; see, e.g., [1-4]. The UFG structure in bulk metallic materials is typically formed due to severe plastic deformation that causes grain refinement in initially coarse-grained structures. In order to control the final UFG structure in severely deformed materials, it is highly interesting to understand and describe both the nature and micromechanisms for deformation-induced grain refinement. Despite of the progress in this area [5,6], micromechanisms for deformation-induced grain refinement are still the subject of intense debates. In these circumstances, of particular interest is the stress-driven grain boundary migration which by definition represents a plastic deformation mode carried by migrating grain boundaries [7,8]. It is conventionally treated that the stress-driven grain boundary migration carries both nanoscale plastic flow and grain growth [7,8]. In particular, this role of the stress-driven grain boundary migration is indirectly supported by experimental observations [9,10] of the athermal grain growth in the vicinities of cracks in NC materials. Recently, it has been

suggested that the stress-driven grain boundary migration plays an alternative role as a process carrying nucleation of new grains in NC and UFG materials [11-14]. In doing so, the grain nucleation is considered as a process occurring through splitting and migration of grain boundaries, in parallel with splitting and movement of grain boundary disclinations (rotational defects) [11-14]. In papers [11-14], the two types of the initial grain boundary structure (before the grain nucleation) were considered: a highly stressed, disclination-free region near a crack tip of a NC solid [13], and a grain boundary structure containing a disclination dipole in the vicinity of a triple junction of grain boundaries [11,12,14]. Whereas, in paper [15], rather similar situation of nanograin chain formation was considered. In this situation, the initial grain boundary structure contains a quadrupole of disclinations located at grain boundary junctions of a previously formed nanograin in a NC or UFG specimen. The final structure contains two neighboring nanograins which are elongated along the same direction and may serve as nuclei for experimentally observed [16-18] nanograin agglomerates (groups of nanograins with low-angle and/or special boundaries within comparatively large grains having high-angle boundaries) in NC and UFG materials.

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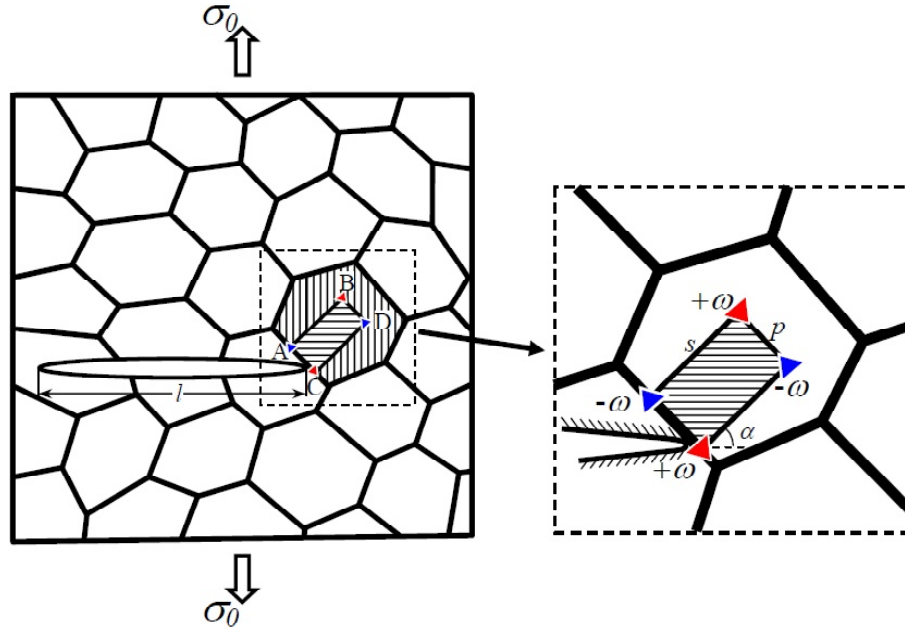


Fig. 1. Generation of a new nanograin and associated disclination quadrupole near a crack tip in deformed NC material.

2. FORMATION OF NANOGRAINS NEAR CRACK TIPS

Paper [13] was concerned with nanograin nucleation in disclination-free region near crack tip. In the paper [13], a NC specimen with the mean grain size d contains a mode I crack and is subjected to one-axis tensile load σ_0 . Fig. 1 schematically shows a two-dimensional section of an NC solid with a flat crack of length l . Following the approach [11], the nanograin nucleation can serve as a special deformation mechanism occurring through splitting of a pre-existent GB AC into immobile boundary AC and mobile boundary BD , migration of GB BD , and formation of two new grain boundaries AB and CD (Fig. 1). The nucleation of the nanograin $ABCD$ is accompanied by formation of a quadrupole $ABCD$ of wedge disclinations with the strengths $\pm\omega$ near the crack tip [11] (Fig. 1). The new nanograin has a rectangular form with characteristic sizes s and p (serving also as distances between the disclinations located at junctions of the nanograin), and its orientation relative to the crack plane is characterized by angle α (Fig. 1).

The nanograin nucleation (Fig. 1) is characterized by the energy difference $\Delta W = W_2 - W_1$, where W_1 is the energy of the solid in its initial state containing the crack, and W_2 is the energy of the solid in its final state containing both the crack and new nanograin. The nanograin nucleation is energetically favorable, if $\Delta W < 0$. The energy difference ΔW (per unit length along the axis z) can be represented as the sum of the three terms [13]:

$$\Delta W = E^q + E^{q-\sigma} + E_{gb}, \quad (1)$$

where E^q is the proper energy of the disclination quadrupole, $E^{q-\sigma}$ is the energy that characterizes the interaction between the disclination quadrupole and the shear stress field σ_{ij}^c created by the external tensile load σ_0 in the solid containing the crack, and E_{gb} is the sum energy of the three new grain boundaries AB , CD , and BD .

In a first approximation (neglecting the effects of the crack on the disclination quadrupole stresses), the proper energy W^q of the disclination quadrupole is given by the following standard expression [19] for the disclination quadrupole energy in an infinite, elastically isotropic solid [13]:

$$E^q = \frac{D\omega^2 p^2}{2} \left((1+t^2) \ln[1+t^2] - t^2 \ln t^2 \right), \quad (2)$$

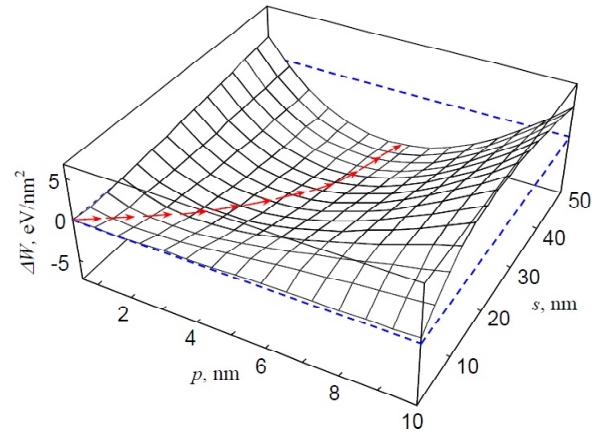


Fig. 2. Dependence of the energy difference ΔW on the nanograin sizes s and p .

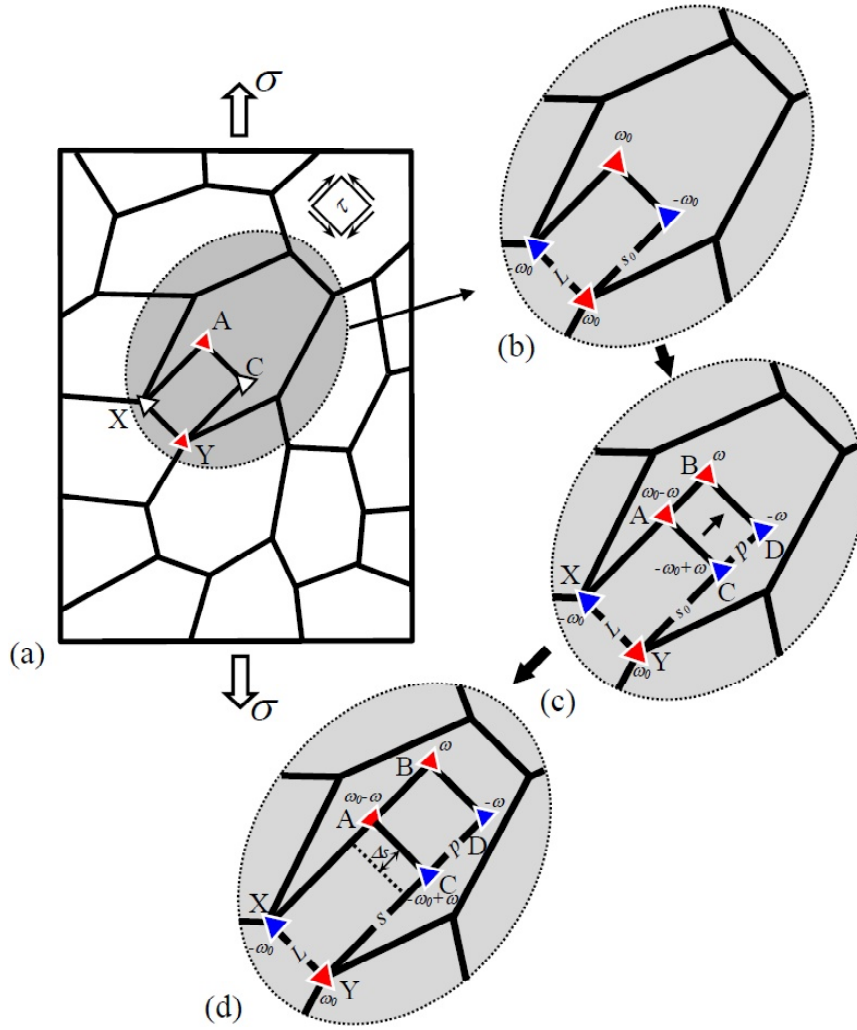


Fig. 3. Formation of a nanograin chain near disclinated nanograin in an UFG specimen. (a) A mechanically loaded specimen with the UFG structure. (b) Initial state with previously formed nanograin ACYX containing disclination quadrupole. (c) Formation a new nanograin ABDC through stress-driven splitting and migration of the grain boundary AC. (d) Formation of a nanograin chain due to growth (elongation) of grains ACYX and ABDC.

where $t = s/p$.

The energy $E^{q-\sigma}$ figuring in formula (2) is written as follows [13,20]:

$$E^{q-\sigma} = -\omega \int_S \sigma_{xy}^c dS', \quad (3)$$

where S is the area bounded by the disclination quadrupole, S' is the integration parameter, and σ_{xy}^c is a component of the stress field σ_{ij}^c (in the coordinate system (x,y) ; see Fig. 1) created by the external tensile load σ_0 in the solid containing the crack. The expression for σ_{xy}^c can be found in Ref. [21]. The energy E_{gb} of the new grain boundaries AB, CD, and BD is evidently given by formula [13]:

$$E_{gb} = \gamma_b (2s + p), \quad (4)$$

with γ_b being the grain boundary energy per unit area of a boundary plane.

Formulas (1)-(4) allow one to find the energy difference ΔW [13]. The dependence of the energy difference ΔW on the sizes s and p of the new nanograin in the example case of NC ceramic α -Al₂O₃ is presented in Fig. 2. In calculations, the following typical values of parameters of α -Al₂O₃ were used [22,23]: $G = 169$ GPa, $\nu = 0.23$, $\gamma = 1.5$ J/m², $\gamma_b = 0.5$ J/m². The external stress σ_0 is supposed to be equal to 1 GPa; this value is typical for mechanical tests of nanocrystalline ceramics. The crack length L is chosen as $L = 800$ nm. This value is close to (but slightly lower than) the Griffith crack length in α -Al₂O₃ at the considered stress level. The characteristic angle is chosen as $\alpha = 70^\circ$.

Arrows in Fig. 2 show evolution in the space (s,p) , corresponding to the most energetically favorable formation/growth of the new nanograin. As it follows from Fig. 2, the most energetically favorable growth of the new nanograin is specified by a simultaneous increase

in the nanograin sizes s and p . That is, for any value of the size s , the equilibrium rectangular shape (corresponding to the minimum energy difference $\Delta W(s,p)$ of the new nanograin is specified by equilibrium size p_{eq} .

Thus, the nanograin generation through stress-driven splitting and migration of grain boundaries near crack tips is energetically favorable in wide ranges of parameters of deformed NC materials. Formation of new nanoscale grains and associated disclination quadrupoles near crack tips relaxes stresses concentrated near these tips and enhances the fracture toughness in NC materials.

3. FORMATION OF NANOGRAIN CHAINS NEAR DISCLINATED NANOGRAINS

In paper [15], a model situation of nanograin chains formation in the vicinities of previously nucleated nanograins containing disclination quadrupoles is considered (Figs. 3a and 3b). In the framework of the model [15], a new nanograin is nucleated in the vicinity of the pre-existent nanograin as follows. The grain boundary AC splits into the immobile grain boundary (also called AC) and the mobile grain boundary BD that migrates under the external shear stress τ (Fig. 3c). The migration causes plastic flow (within the area swept by the migrating boundary BD) and is accompanied by formation of new grain boundaries AB and CD (Fig. 3c). The junction points B and D of the mobile grain boundary contain the disclinations with strengths $\pm\omega$. These disclinations form a mobile dipole configuration and are resulted from splitting of the initial disclinations A and C with strengths $\pm\omega_0$ (Figs. 3b and 3c). As a result of the splitting transformations of the grain boundary AC and the disclinations, a new disclinated nanograin $ABCD$ is formed (Fig. 3c).

The distance between the grain boundaries AC and BD is denoted as p . In general, during the structural transformation, the external shear stress τ can cause migration of the grain boundary AC as well (Fig. 3d). In these circumstances, geometry of the structural transformation (Figs. 3b-3d) is specified by changes in both the sizes p and s . Finally, note that a series of structural transformations like that shown in Figs. 3b-3d can produce new nanograins. Nanograin chains (Fig. 3d) serve as nuclei of nanograin agglomerates (groups of nanograins with low-angle and/or special boundaries within comparatively large grains having high-angle boundaries) that typically form in NC and UFG materials under severe plastic deformation [16-18].

The splitting transformation (Figs. 3b-3d) is characterized by the energy change $\Delta W = W_2 - W_1$, where W_1 is the energy of the defect configuration in its initial state

with the ω_0 -disclination quadrupole (Fig. 3b), and W_2 is the energy of the defect configuration in its final state (Fig. 3d). The transformation is energetically favorable, if $\Delta W < 0$. In terms of the theory of defects in solids, the energy change ΔW is written as follows [15]:

$$\Delta W = E_s^{\omega_0} + E_s^{\omega} + E_s^{\omega_0-\omega} + E_{int}^{\omega_0-\omega} + E_{int}^{\omega_0-(\omega_0-\omega)} + E_{int}^{\omega-(\omega_0-\omega)} + E_\gamma - E_\tau - E_s^q. \quad (5)$$

Here $E_s^{\omega_0}$, E_s^{ω} , $E_s^{\omega_0-\omega}$ and E_s^q are the proper elastic energies of the dipole XY of $\pm\omega_0$ -disclinations, the dipole BD of $\pm\omega$ -disclinations, the dipole AC of $\pm(\omega_0-\omega)$ -disclinations and the initial quadrupole $XACY$ of $\pm\omega_0$ -disclinations, respectively; $E_{int}^{\omega_0-\omega}$ is the energy that characterizes the elastic interaction between the dipole XY of $\pm\omega_0$ -disclinations and the dipole BD of $\pm\omega$ -disclinations; $E_{int}^{\omega_0-(\omega_0-\omega)}$ is the energy that characterizes the elastic interaction between the dipole XY of $\pm\omega_0$ -disclinations and the dipole AC of $\pm(\omega_0-\omega)$ -disclinations; $E_{int}^{\omega-(\omega_0-\omega)}$ is the energy that characterizes the elastic interaction between the dipole BD of $\pm\omega$ -disclinations and the dipole AC of $\pm(\omega_0-\omega)$ -disclinations; E_γ denotes the sum of the specific energies (per unit area) of the new grain bounda-

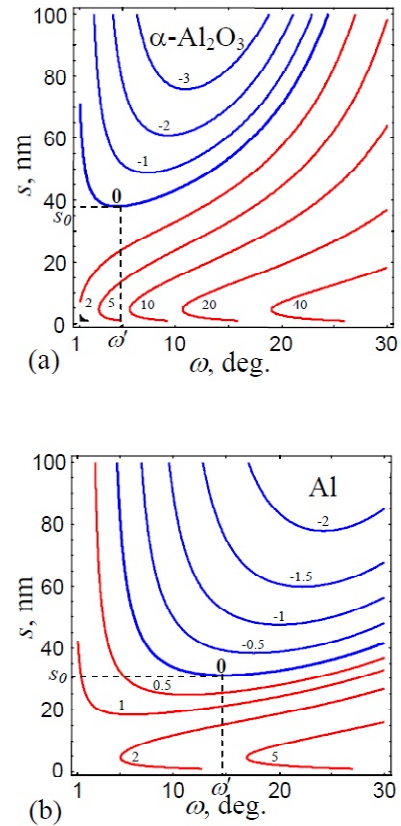


Fig. 4. The map of the energy difference ΔW in the coordinate plane of ω and s at the external stress value $\tau = 1$ GPa, for (a) NC ceramics $\alpha\text{-Al}_2\text{O}_3$ (sapphire) and (b) NC Al.

ries AB , CD , and BD ; and E_τ is the work spent by the external shear stress ω to displacements of the disclination dipoles AC and BD (Fig. 3d).

After some calculation one can find the expression for the energy change ΔW (see [15], for details):

$$\Delta W = \frac{D\omega^2}{2} \left(L^2 \ln \frac{L^2 + p^2}{L^2} + p^2 \ln \frac{L^2 + p^2}{p^2} \right) + \frac{D\omega_0\omega}{2} \left(L^2 \ln \frac{L^2}{L^2 + s^2} + L^2 \ln \frac{L^2 + (s+p)^2}{L^2 + p^2} \right. \\ \left. + (s+p)^2 \ln \frac{L^2 + (s+p)^2}{(s+p)^2} - s^2 \ln \frac{L^2 + s^2}{s^2} - p^2 \ln \frac{L^2 + p^2}{p^2} \right) - (\omega_0\Delta s + \omega p)L\tau + (2(p + \Delta s) + L)\gamma_{gb}. \quad (6)$$

The maps ΔW in coordinates of the arm s of the initial $\pm\omega_0$ -disclination quadrupole and the disclination strength ω of the mobile $\pm\omega$ -disclination dipole in the exemplary cases of NC ceramics α -Al₂O₃ (sapphire) and NC Al (aluminum) are presented in Fig. 4. In calculations, the following typical values of parameters were used [23,24]: $G = 169$ GPa and $\nu = 0.23$, for α -Al₂O₃; and $G = 26.5$ GPa and $\nu = 0.34$, for Al. The values γ_{gb} of the energy of special grain boundaries in α -Al₂O₃ and Al are as follows: $\gamma_{gb}^{Al_2O_3} = 0.05$ J/m² [25] and $\gamma_{gb}^{Al} = 0.1$ J/m² [26]. The arm and strength of the initial $\pm\omega_0$ -disclination dipole AC (Fig. 1b) are taken as $L = 10$ nm and $\omega = 30^\circ$, respectively.

As it follows from Fig. 4, the splitting process is energetically favorable when the initial distance between the grain boundaries XY and AC has value of $s_0 \approx 37$ nm at the disclination strength $\omega' \approx 5^\circ$, for α -Al₂O₃ (Fig. 4a); and $s_0 \approx 31$ nm at $\omega' \approx 14^\circ$, for Al (Fig. 4b). In both the cases under examination, when the distance s (with the proviso that $s > s_0$) increases, so does the interval of disclination strength values ω at which the splitting process is energetically favorable (Fig. 4).

Thus, the splitting transformations of grain boundaries of previously formed nanograins with disclination quadrupoles at their junctions (Figs. 3b-3d) are energetically favorable and can initiate nucleation of new nanograins in nanocrystalline α -Al₂O₃ and Al in wide ranges of their parameters. Nanograin chains (Fig. 3d) are resulted from these transformations.

4. CONCLUDING REMARKS

Thus, formation of the nanograins and nanograin chains is energetically favorable in wide ranges of parameters of deformed UFG and NC materials. It is shown that the generation of nanograins near crack tips causes a partial stress relaxation near cracks. Besides, the stress-driven nanograin formation in NC ceramics is expected to be more effective as a deformation and toughening mechanism, compared to the stress-driven grain growth, because migration of pre-existent grain boundaries in nanoceramics is often suppressed by the difference in the chemical composition between the grain interiors and grain boundaries. Also, it is shown that the nanograin chains may serve as nuclei for experimentally observed [16-18] nanograin agglomerates (groups of nanograins with low-angle and/or special boundaries within comparatively large grains having high-angle boundaries) in NC and UFG materials. The suggested theoretical representations on plastic flow occurring through generation of nanograins at grain boundaries in UFG materials are well consistent with the experimental observation [27] of nanograins generation at grain boundaries in cobalt.

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