A thermodynamically self-consistent damage equation for grainsize evolution during dynamic recrystallization.

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SUMMARY

We employ basic non-equilibrium thermodynamics to propose a general equation for the mean grainsize evolution in a deforming medium, under the assumption that the whole grainsize distribution remains self-similar. We show that the grainsize reduction is controlled by the rate of mechanical dissipation in agreement with recent findings. Our formalism is self consistent with mass and energy conservation laws and allows a mixed rheology. As an example, we consider the case where the grainsize distribution is lognormal, as is often experimentally observed. This distribution can be used to compute both the kinetics of diffusion between grains and of dynamic recrystallization. The experimentally deduced kinetics of grainsize coarsening indicates that large grains grow faster than what is assumed in classical normal grain growth theory. We discuss the implications of this model for a mineral that can be deformed under both dislocation creep and grainsize sensitive diffusion creep using experimental data of olivine. Our predictions of the piezometric equilibrium in the dislocation-creep regime are in very good agreement with the observations for this major mantle-forming mineral. We show that grainsize reduction occurs even when the average grainsize is in diffusion creep, because the largest grains of the grainsize distribution can still undergo recrystallization. The resulting rheology that we predict for olivine is time-dependent and more non-linear than in dislocation creep. As the deformation rate remains an increasing function of the deviatoric stress, this rheology is not localizing.

Key words: Grainsize – Distribution – Rheology – Olivine

1 INTRODUCTION

The localization of deformation in narrow shear bands is necessary for plate tectonics to occur (see e.g., Bercovici et al. 2000). Weak faults can be formed during deformation but their weakness can persist even after a reorganization of the large scale stress pattern (e.g., Gurnis et al. 2000). This indicates that the rheology is not only controlled by the instantaneous stress field but has memory and healing.

Localization occurs by a feed-back between the rheological law and the deformation wherein a faster deformation can be obtained with a lower stress. In simple shear experiments, this happens when the derivative of deviatoric stress τ with respect to strain-rate $\dot{\epsilon}$ is negative (Bercovici 1993; Montési & Zuber 2002). The fact that the rheology of silicates is often expressed by a nonlinear expression with $\epsilon \propto \tau^n$ and $n \ge 1$ (Ranalli 1995) does not lead to strike slip localization as τ remains a monotonically increasing function of $\dot{\epsilon}$ (Bercovici 1995). Non-linear rheologies with large positive exponents tend, however, to narrow the zones of deformation (Weinstein & Olson 1992; Landuyt & Bercovici 2009).

Shear heating has often been proposed as a source of localization that provides some long term memory to the rheology (the thermal diffusion time) (Fleitout & Froidevaux 1980; Leloup et al. 1999; Kameyama et al. 1997). Unfortunately, although shear heating is necessarily associated with localization, it does not seem to explain either the narrowness of plate boundaries, or their geometries (e.g., Bercovici & Karato 2003). A local increase of porosity/microcraks occurring during deformation has also been invoked for localizing the deformation (Bercovici 1998; Ogawa 2003) but this process is not efficient in 3D simulations at generating toroidal motions, e.g. shearing between plates and plate rotations (Bercovici & Ricard 2005). Anisotropic mechanical behaviour due to an inherited preferred orientation of crystals could also control or favor the localization (e.g., Tommasi et al. 2009; Bystricky et al. 2000). Once the localization is effective, minor mineralogical phases like serpentine can also lubricate the motion (Hilairet et al. 2007).

Grainsize reduction seems the most attractive physical process for explaining the initial localization of the deformation. Cataclastic fracturing and recrystallisation are well known mecanisms of grainsize reduction and the ductile deformation in the diffusive regime is facilitated in the presence of small grains (Kelemen & Hirth 2007). However, models of localization by grainsize reduction are not self-consistent. Recrystallisation is observed in the dislocation regime (i.e., when the microscospic deformation proceeds by coherent motion of crystal dislocations rather than by the individual diffusion of atoms and vacancies). Grainsize reduction and localization by grainsize sensitive rheology occur therefore in somewhat exclusive regimes (Karato et al. 1980; Derby & Ashby 1987). Various models have however discussed the possible interactions between large scale deformation and grainsize evolution (Kameyama et al. 1997; Braun et al. 1999; Montési & Hirth 2003; Bercovici & Ricard 2005).

Up to now, most attempts to model the evolution of grainsizes have been derived from phenomenologic laws involving only a mean grainsize. A few attempts have been made to describe in a very general way the evolution of an assemblage of grains under deformation (e.g., Slotemaker 2006; Ricard & Bercovici 2009). In these approaches, one has to consider the complete distribution of grains $\nu(\mathcal{R}, \mathbf{X}, t)$ (which is the number of grains per unit volume near the position \mathbf{X} and at time t, having a size between \mathcal{R} and $\mathcal{R} + d\mathcal{R}$). Although a general and physically consistent theory has been proposed (Ricard & Bercovici 2009), the mathematical formalism remains cumbersome and a general implementation in a 3D and time dependent geodynamic simulation seems implausible.

The approach followed by Ricard & Bercovici (2009) shows that the grainsize reduction cannot be related to the stress or to the strain-rate tensor, alone, but necessarily to their scalar product $\underline{\tau} : \underline{\dot{\epsilon}}$ (where $\underline{\tau} : \underline{\dot{\epsilon}} = \sum_{ij} \tau_{ij} \dot{\epsilon}_{ij}$). This result is based on the second law of thermodynamics that requires the positivity of the entropy sources. This theoretical requirement already used in Bercovici & Ricard (2005), has been confirmed empirically by Austin & Evans (2007) that concluded that the grainsize is a "paleowattmeter" (i.e., a measure of the rate of dissipation) rather than a "paleo piezometer" (i.e., a measure of the deviatoric stress).

At a microscopic level and laboratory scale there are a large number of observations regarding the evolution of silicate rheology with grainsize, pressure, temperature and stress (but also, water content, oxygen fugacity, porosity...). At the same time, models of mantle convection simulate the existence of plates with ad hoc rheologies (Tackley 2000a,b; Stein et al. 2004). Up to now, these models are not based on experiments but often assume that the lithosphere has a linear viscosity with a plasticity threshold. The goal of this paper is to provide a theory based on laboratory experiments that could be used in large scale geodynamic modeling.

2 EVOLUTION OF GRAINSIZE DISTRIBUTION

Within a volume of material, there is a continuous distribution of grainsizes, $\nu(\mathcal{R}, \mathbf{X}, t)$, so that the number of grains dn with sizes between \mathcal{R} and $\mathcal{R} + d\mathcal{R}$, per unit volume dV, at position \mathbf{X} and time t is

$$dn(\mathcal{R}, \mathbf{X}, t) = \nu(\mathcal{R}, \mathbf{X}, t) \ d\mathcal{R}dV.$$
(1)

The distribution ν has units of m^{-4} . From this distribution, average quantities like the mean grainsize can be computed. The equations for grainsize evolution that have been proposed are empirical and consider the mean grainsize only. Here, we also derive an equation for the mean grainsize but from theoretical thermodynamic considerations, starting explicitly from the existence of the grainsize distribution.

The grainsize distribution evolves through two different processes. First, mass transfer between grains can occur continuously through grain boundary migration or diffusion. This involves a change in the number of grains dn in the bin of size \mathcal{R} (i.e., a change in the number of grains of sizes between \mathcal{R} and $\mathcal{R} + d\mathcal{R}$) by coarsening of smaller grains (or continuous reduction of larger grains). This can be described by introducing the rate of movement of the grain in size space $\dot{\mathcal{R}} = \partial \mathcal{R}(\mathbf{X}, t)/\partial t$ (Lifshitz & Slyozov 1961). Mathematically, this process occurs at constant total number of grains per unit volume (experimentally, smallest grains shrink below observability). Second, the number of grains of a given size can be populated by a discontinuous transfer from remote population bins. For example, large grains can be subdivided by the formation of sub-grain boundaries that nucleate new small grains (Hobbs 1968), or can be broken by cataclasis. We call $\Gamma(\mathcal{R})$ the rate at which grains are added to or removed from the bin of size \mathcal{R} by discontinuous process. This discontinuous process changes the total number of grains per unit volume. The balance of grain population implies a continuity equation for the grainsize distribution itself (Atkinson 1988; Hillert 1965; Ricard & Bercovici 2009)

$$\frac{\partial\nu}{\partial t} + \frac{\partial\dot{\mathcal{R}}\nu}{\partial\mathcal{R}} = \Gamma, \qquad (2)$$

where t is time. For simplicity we omit the space variable X and the advection term $\nabla \cdot (\mathbf{v}\nu)$ where $\mathbf{v}(\mathbf{X}, t)$ is the macroscopic velocity of the grained medium (Ricard & Bercovici 2009).

Most attempts of modeling grain coarsening or damage have assumed that the whole grainsize distribution can be obtained by the knowledge of its mean grainsize (c.f., De Bresser et al. 2001; Ricard & Bercovici 2009). Representing the whole grainsize distribution by only a single size scale (e.g., some average) means mathematically that the distribution is self-similar, i.e. that

$$\nu(\mathcal{R}, t) = A(\mathcal{R}_0)H(u),\tag{3}$$

where \mathcal{R}_0 is some average grainsize function of t that will be defined later, $u = \mathcal{R}/\mathcal{R}_0$ is the self-similarity variable, and A is an amplitude. This self-similarity assumption is supported experimentally (e.g., Slotemaker 2006) although exceptions exist such as abnormal grain growth (Hillert 1965). Self-similarity has also a mathematical justification. If one solves the equation (2) starting from any arbitrary distribution, and for quite general assumptions for $\hat{\mathcal{R}}$ or Γ , the distribution evolves toward a self-similar solution like (3), after sufficiently long time. This mathematical result has been used in various classical studies of grain coarsening (Lifshitz & Slyozov 1961; Wagner 1961; Hillert 1965), grain fragmentation or aggregation (Collet 2004) and is valid under more general assumptions (Ricard & Bercovici 2009). Our basic start is therefore that although the self-similarity may not be strictly valid, the system remains close enough to a self-similar state than a mean size theory remains useful and is a reasonable compromise between the complexity of a general grainsize distribution theory and what can be constrained by the available observations. Future observations of time evolution of grainsize variance, skewness and other higher moments of the grainsize distribution will be needed to adjust the parameters of a more general theory.

The mass conservation equation, assuming a constant density of grains, implies that the unit volume is just the sum of the volumes of grains, which leads to the normalization

$$\int_{0}^{\infty} \breve{\mathsf{v}}\nu(\mathcal{R}) \, d\mathcal{R} = 1,\tag{4}$$

where

$$\breve{\mathsf{v}} = \frac{4}{3}\pi \mathcal{R}^3,\tag{5}$$

is the grain volume. The notations follow Ricard & Bercovici (2009) and the breve accent (e.g., $\check{}$) represents variables at the

grainsize level. The grains are assumed to be spherical, but more complex shapes could be handled easily by replacing π by a comparable factor accounting for polyhedral grain shapes (see Ricard & Bercovici 2009). In the normalization (4), we omit the space and time variables, and we integrate the distribution over all grainsizes, potentially from $\mathcal{R} = 0$ to $\mathcal{R} = +\infty$ (we assume that *H* is zero after some finite value or is well-behaved enough that all the integrals containing *H* converge at $u = +\infty$).

Using the self similar expression (3), this normalization condition implies

$$A(\mathcal{R}_0) = \frac{3}{4\pi\lambda_3} \frac{1}{\mathcal{R}_0^4} \tag{6}$$

where we define, as in Ricard & Bercovici (2009),

$$\lambda_n = \int_0^\infty u^n H(u) du. \tag{7}$$

The amplitude $A(\mathcal{R}_0)$ is therefore related to the grainsize distribution H.

3 AVERAGE GRAIN COARSENING AND DAMAGE

The rules of differentiation applied to distribution (3) taking into account the time dependence of the amplitude (6) lead to

$$\frac{\partial\nu}{\partial t} = -\frac{3}{4\pi\lambda_3} \frac{1}{R_0^5} \frac{d\mathcal{R}_0}{dt} \frac{1}{u^3} \frac{\partial u^4 H}{\partial u}.$$
(8)

This proves that $\partial \nu / \partial t$ is also a self similar function as it can be written as a function of \mathcal{R}_0 times a function of u. As $\partial \nu / \partial t$ appears in the evolution equation (2), $\partial \dot{\mathcal{R}} \nu / \partial \mathcal{R}$ (and therefore $\dot{\mathcal{R}}$) and Γ must also be self similar (or have non self similar contributions that cancel each other and are irrelevant to the grainsize evolution). Γ is therefore the product of a time-dependent (or \mathcal{R}_0 -dependent) amplitude $C(\mathcal{R}_0)$ times a shape functions of $u = \mathcal{R}/\mathcal{R}_0$ such that

$$\Gamma = \frac{3}{4\pi\lambda_3 \mathcal{R}_0^5} C(\mathcal{R}_0) \frac{1}{u^3} \frac{\partial u^4 H}{\partial u},\tag{9}$$

(the coefficient $3/(4\pi\lambda_3\mathcal{R}_0^5)$ is included for subsequent simplification). Similarly, given (2) and (8), $\dot{\mathcal{R}}$ can be written as

$$\dot{\mathcal{R}} = B(\mathcal{R}_0) \frac{1}{H} \left(b + \int_0^u \frac{1}{v^3} \frac{\partial v^4 H}{\partial v} dv \right)$$
$$= B(\mathcal{R}_0) \left(\frac{b}{H} + u + \frac{3}{H} \int_0^u H dv \right), \tag{10}$$

and where $B(\mathcal{R}_0)$ is an amplitude factor. The integration constant b can be obtained from the condition of mass conservation (see Ricard & Bercovici (2009))

$$\int_{0}^{\infty} \breve{v} \Gamma(\mathcal{R}) d\mathcal{R} = 0 \text{ and } \int_{0}^{\infty} \frac{d\breve{v}}{dt} \nu(\mathcal{R}) d\mathcal{R} = 0, \quad (11)$$

where \breve{v} is the volume of a grain (see (5)). This assumes that mass is conserved either during the fusion or fission of grains (first equality) or during continuous mass transfer between grains (second equality). According to (3), (9) and (15), these two equations become

$$\int_{0}^{\infty} \frac{du^{4}H}{du} du = 0 \text{ and } \int_{0}^{\infty} u^{2}(b+uH+3\int_{0}^{u}H\,dv)du = 0.$$
(12)

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The first condition is already verified provided that $u^4 H$ is zero for u = 0 and $u = +\infty$. Then, by using an integration by parts,

$$\int_{0}^{X} \left(3u^{2} \int_{0}^{u} H dv \right) du = \left[u^{3} \int H du \right]_{0}^{X} - \int_{0}^{X} u^{3} H du$$
(13)

in the limit of $X \to +\infty$, we therefore prove that mass conservation implies

$$b = -3 \int_0^{+\infty} H \, du. \tag{14}$$

which allows us to write

$$\dot{\mathcal{R}} = B(\mathcal{R}_0) \left(u - \frac{3}{H} \int_u^{+\infty} H dv \right).$$
(15)

As the *u*-shapes of Γ and $\hat{\mathcal{R}}$ are derived from (2), which is itself another expression of the condition of mass conservation, it is not surprising that mass conservation is naturally satisfied.

When the self similar expressions (9) and (15) are introduced back into (2), the shape function can be eliminated and only a differential equation for the average size \mathcal{R}_0 remains

$$\frac{d\mathcal{R}_0}{dt} = B(\mathcal{R}_0) - C(\mathcal{R}_0).$$
(16)

Up to now the signs of $B(\mathcal{R}_0)$ and $C(\mathcal{R}_0)$ are not known. The goal is therefore to constrain these quantities which control the kinetics of continuous and discontinuous grain processes, respectively from observations guided by the necessary condition of positivity of entropy production.

4 ENERGY CONSIDERATIONS

During deformation, part of the input energy is simply dissipated (entropy) and part is stored reversibly (work). The irreversible energy is dissipated as heat although some minor entropy component (here neglected) can be associated with interfaces when the surface energy is temperature dependent (Bailyn 1994; Bercovici et al. 2001a). Three reversible terms can be considered: the macroscopic elastic energy, the energy of dislocations (Karato 2008) and the surface energy of grains, γ (in J m⁻²). The elastic energy stored around dislocations is always negligible compared to the surface energy of grains (see Shimizu 2008). When a material is deformed, some of the input energy is initally stored elastically, then some is used to increase the density of dislocations (see, Kohlstedt & Weathers 1980; Karato & Jung 2003). At a given point, the dislocations will rearrange to form subgrain boundaries and later, grain boundaries. The elastic energy around dislocations is therefore mostly a temporary buffer converting some of the deformational work into surface energy. Our assumption is that, in the long term, the energy balance is between the deformational work, the viscous dissipation and the change of grain surfaces. Understanding the grainsize evolution is therefore understanding how the macroscopic energy partitions between dissipation (through heat production) and storage (by creation of new grain surface through a temporary elastic stage).

Ricard & Bercovici (2009) show that the energy conservation and the requirement of entropy positivity lead to

$$\int_{0}^{\infty} \left(-\frac{2\gamma}{\mathcal{R}} \frac{d\breve{\mathsf{v}}}{dt} \nu(\mathcal{R}) - \frac{3}{2} \frac{2\gamma}{\mathcal{R}} \breve{\mathsf{v}} \Gamma(\mathcal{R}) + \breve{\underline{\tau}} : \underline{\dot{e}} \ \breve{\mathsf{v}} \nu(\mathcal{R}) \right) d\mathcal{R} \ge 0.$$
(17)

Although complex, the meaning of this equation is straightforward. The first term represents the continuous change of surface energy of grains and the second term the surface energy stored or removed by the fusion or fission of grains. The third term represents the energy deposited in each grain by the deformation ($\underline{\check{\tau}}$ and $\underline{\check{e}}$ are the stress and strain-rate in each grain). We retain the term $2\gamma/\mathcal{R}$ as it appears in the Laplace expression for the excess pressure in a sphere due to surface tension γ (γ can either be interpreted as a surface energy (in J m⁻²) or as the surface tension (in N m⁻¹) (see Bailyn 1994).

The likely most common case of grainsize evolution is when the grain growth is dominated exclusively by the diffusive mass exchange in keeping with Lifshitz & Slyozov (1961); Wagner (1961); Feltham (1957), but where grain reduction is driven by the deformation populating the material with small grains issued from "breaking" large grains. This suggests we look for mechanisms that independently obey to

$$\int_{0}^{\infty} -\frac{2\gamma}{\mathcal{R}} \frac{d\breve{v}}{dt} \nu(\mathcal{R}) \, d\mathcal{R} \ge 0, \tag{18}$$

and

$$\int_{0}^{\infty} \left(-\frac{3}{2} \frac{2\gamma}{\mathcal{R}} \breve{\mathsf{v}} \Gamma(\mathcal{R}) + \breve{\underline{\tau}} : \dot{\underline{e}} \breve{\mathsf{v}} \nu(\mathcal{R}) \right) d\mathcal{R} \ge 0.$$
(19)

According to (3), (9) and (15) these two conditions can be readily expressed. The first one, (18), just implies

$$-B(\mathcal{R}_0)\int_0^\infty u\left(uH-3\int_u^{+\infty}Hdv\right)\,du=\frac{1}{2}B(\mathcal{R}_0)\lambda_2\ge 0,$$
(20)

where the equality is obtained by integration by parts (similar to (13) with n = 2) and where λ_2 is the positive integral defined in (7). This inequality means that $B(\mathcal{R}_0) \ge 0$, and therefore that continuous transport must always lead to grain coarsening. The second inequality imposed by the positivity of entropy sources (19), becomes

$$\frac{3\gamma}{\mathcal{R}_0^2} \frac{\lambda_2}{\lambda_3} C(\mathcal{R}_0) \le \int_0^\infty \underline{\breve{\tau}} : \underline{\dot{e}} \, \breve{\nu}(\mathcal{R}) \, d\mathcal{R}.$$
(21)

Notice that while the continuous coarsening of grains is necessarily an entropy source (i.e., $B(R_0)$ must be positive), the processes of fragmentation/coagulation of grains can be either a source (when $C(R_0) \leq 0$) or a sink (when $C(R_0) \geq 0$) of entropy. The dissipation term itself $\underline{\check{\tau}} : \underline{\check{\epsilon}}$ is always positive.

The process reducing the grain size is an entropy sink $C(R_0) \ge 0$. This term is bounded by the mechanical dissipation expressed by the previous inequality. As advocated in a series of papers (Bercovici et al. 2001a,b; Bercovici & Ricard 2005; Ricard & Bercovici 2009), a reasonable and pragmatic choice is to introduce a partitioning function $\check{f}(\mathcal{R}, \tau)$, with $0 \le \check{f}(\mathcal{R}, \tau) \le 1$ so that

$$C(\mathcal{R}_0) = \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \int_0^\infty \breve{f} \, \breve{\underline{\tau}} : \dot{\underline{e}} \, \breve{\nu}(\mathcal{R}) \, d\mathcal{R}$$
(22)

The general expression of the grainsize evolution (16) becomes therefore

$$\frac{d\mathcal{R}_0}{dt} = B(\mathcal{R}_0) - \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \int_0^\infty \breve{f}\, \breve{\underline{t}} : \dot{\underline{\breve{e}}}\,\breve{\breve{v}}\nu(\mathcal{R})\,d\mathcal{R} \qquad (23)$$

The coarsening term is often given on the form $B(\mathcal{R}_0) = G/\mathcal{R}_0^{p-1}$ (Hillert 1965; Atkinson 1988; Karato 2008) where p is of order 2 (corresponding to a coarsening law where $\mathcal{R}_0 \propto t^{1/2}$). The kinetic term G is temperature dependent, G =

 $k_0 \exp(-E_g/(RT))$ but we are not aware of an experimentally observed dependence on stress (cf. Table 1). We therefore use

$$\frac{d\mathcal{R}_0}{dt} = \frac{G}{p\mathcal{R}_0^{p-1}} - \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \int_0^\infty \check{f}\, \check{\underline{\tau}}: \dot{\underline{\acute{e}}}\, \check{\mathbf{v}}\nu(\mathcal{R})\, d\mathcal{R} \qquad (24)$$

5 THE CASE OF LOGNORMAL DISTRIBUTIONS

5.1 Experimental distributions

Various experimental studies have indicated that the grainsize distribution is lognormal (e.g., Feltham 1957; Slotemaker 2006; Faul & Scott 2006). The distribution of grainsizes (1) is in this case, a Gaussian when plotted as a function of $\ln R$, i.e.,

$$dn \propto \exp\left(-\frac{(\ln(\mathcal{R}/\mathcal{R}_0)^2)}{2\sigma^2}\right) d\ln \mathcal{R} dV$$

$$= \frac{1}{\mathcal{R}} \exp\left(-\frac{(\ln(\mathcal{R}/\mathcal{R}_0)^2)}{2\sigma^2}\right) d\mathcal{R} dV.$$
(25)

In this expression, σ is the dimensionless variance of the distribution of $\ln(\mathcal{R}/\mathcal{R}_0)$ and \mathcal{R}_0 is the "mean" grainsize.

The lognormality of observed grainsize distributions is only approximate. In a recent compilation of quartz grainsizes, Stipp et al. (2010) argue that the distribution is not lognormal but shows modes indicative of the various processes occuring during dynamic recrystallisation. However, to first order, the distribution observed by Stipp et al. (2010) is, in fact, not far from lognormal when plotted in appropriate logarithmic coordinates. We therefore introduce the lognormal shape function

$$H(u) = \frac{1}{\sqrt{2\pi\sigma u}} \exp\left(-\frac{(\ln u)^2}{2\sigma^2}\right),\tag{26}$$

(while experimentalists often use the common (base 10) logarithm, we use the natural logarithm which makes the computations of derivatives and integrals less confusing).

In the following of the paper, we refer to \mathcal{R}_0 as a mean grainsize, in agreement with experimental literature. However the real mean grainsize is defined as

$$<\mathcal{R}>=\int_{0}^{\infty}\mathcal{R}\nu(\mathcal{R})d\mathcal{R}/\int_{0}^{\infty}\nu(\mathcal{R})d\mathcal{R}=\mathcal{R}_{0}\exp\left(\frac{\sigma^{2}}{2}\right)$$
 (27)

while the true definition of \mathcal{R}_0 is $\exp(<\ln \mathcal{R}>)$ as using (3),

$$<\ln(\mathcal{R})>=\int_{0}^{\infty}\ln(\mathcal{R})\nu(\mathcal{R})d\mathcal{R}/\int_{0}^{\infty}\nu(\mathcal{R})d\mathcal{R}=\ln(\mathcal{R}_{0})$$
 (28)

The "mean" grainsize reported in most publications is indeed the mean of the grainsize distribution plotted as a function of $\ln \mathcal{R}$ rather than $\langle \mathcal{R} \rangle$. The difference between \mathcal{R}_0 and $\langle \mathcal{R} \rangle$ decreases with the decreasing variance of the distribution σ , see (27). For a monodisperse distribution of grainsizes (a Dirac distribution) the two quantities become identical. The choice of a lognormal distribution allows us to compute explicitly the integrals that appear in various equations (e.g. (27) or (28)). A similar exercise could have been done with any other distribution.

The lognormal distribution means that when the maximum of the grainsize distribution is at \mathcal{R}_0 , the grains of sizes

$$\mathcal{R}_{\sigma} = \mathcal{R}_0 \exp(\pm \sigma \sqrt{2 \ln 2}), \tag{29}$$

are two times less frequent. From observations on olivine (Slotemaker 2006; Karato 2008), σ seems to be between 0.5 and 1. This indicates that grains, 3 times larger or 3 times smaller than

the average grainsize are half less probable than grains of the grainsize (this estimate uses $\sigma = 0.93$). In other words, the center of the distribution covers already about one order of magnitude in sizes (3²) and the tails of the distribution cover another order of magnitude.

With this choice of distribution law, we get

$$\int u^n H(u) du = -\frac{1}{2} \exp\left(\frac{n^2 \sigma^2}{2}\right) \operatorname{erf}\left(\frac{n\sigma^2 - \ln(u)}{\sqrt{2}\sigma}\right),\tag{30}$$

which implies

$$\lambda_n = \int_0^\infty u^n H(u) du = \exp\left(\frac{n^2 \sigma^2}{2}\right),\tag{31}$$

$$\dot{\mathcal{R}} = B(\mathcal{R}_0) \left[u - \frac{3}{2H(u)} \operatorname{erfc}\left(\frac{\ln u}{\sqrt{2\sigma}}\right) \right], \quad (32)$$

where erfc is the complementary error function, $(1 - \operatorname{erf})$, and finally,

$$\Gamma = \frac{3C(\mathcal{R}_0)}{4\pi\lambda_3\mathcal{R}_0^5}H(u)\left(3-\frac{\ln u}{\sigma^2}\right).$$
(33)

These distributions are plotted in Figure 1. The function H is shaded and the $\dot{\mathcal{R}}$ and Γu^3 normalized functions (i.e., $\dot{\mathcal{R}}/B(\mathcal{R}_0)$) and $4\pi\lambda_3\mathcal{R}_0^5\Gamma u^3/(3C(\mathcal{R}_0)))$ are depicted in blue dot-dashed and red dashed. We display a curve proportional to Γu^3 instead of Γ to emphasize the negative lobe of the damage term. These two curves change sign roughly at the average grainsize around u = 3. As expected the function Γ , accounting for the discontinuous formation of grains, implies the breaking of large grains ($\Gamma < 0$ for large \mathcal{R}), and thus the formation of small grains ($\Gamma > 0$ for large \mathcal{R}). Although the $\dot{\mathcal{R}}$ function (32) seems very complex, it is a simple monotonically increasing function.

Notice that Feltham (1957) in a well-known paper, using an approach that is, in principle, equivalent, writes that a lognormal distribution corresponds approximatively to $\dot{\mathcal{R}} = C'(\mathcal{R}_0) \ln u/u$. His result is not in agreement with our findings. In his demonstration, Feltham (1957) identifies wrongly the grainsize distribution with the size distribution of grain sections. The probability to have a grain section of radius r is actually the convolution of the grainsize distribution by the probability to cut any grain of radius larger than r, at the specific grainsize section of radius r.

The grain growth law, $\dot{\mathcal{R}}$, is not much different from the functions previously used (see Figure 2), which are based on the inference that the kinetics of intergranular exchange is related to $2\gamma/\overline{\mathcal{R}} - 2\gamma/\mathcal{R} = 2\gamma/\mathcal{R}_0(1/\overline{u} - 1/u)$, i.e., the difference between the pressure in a given grain due to its surface tension, $\propto 2\gamma/\mathcal{R}$, and some average grain pressure written as $\propto 2\gamma/\overline{\mathcal{R}}$ (where $\overline{\mathcal{R}}$ is related to the grainsize distribution through mass conservation, see (11)). When u is large the shape of $\dot{\mathcal{R}}$ looks like $u(1/\overline{u}-1/u)$, while the shape used by Lifshitz & Slyozov (1961) was $1/u(1/\overline{u} - 1/u)$, and the shape used by Hillert (1965) was $(1/\overline{u} - 1/u)$ (see Figure 2). The stronger dependence of $\dot{\mathcal{R}}$ with \mathcal{R} at large grainsize, suggests that grains significantly larger than the average grainsize are more favorable than expected by Lifshitz & Slyozov (1961) or Hillert (1965). These other authors' assumptions yield narrow grainsize distributions of finite extent (see Ricard & Bercovici (2009)) where the maximum grainsize is only twice the average grainsize, while observations show distributions with much longer tails (Faul & Scott 2006).



Figure 1. The lognormal distribution H is shaded. The normalized function Γu^3 is depicted with a red dashed line, the normalized grain growth $\dot{\mathcal{R}}$ by a blue dot-dashed line. The value $\sigma = 0.6$ has been used.

5.2 Microscopic rheology

At grainsize level, and for mantle conditions, the olivine rheology is generally found as a mixture of diffusion and dislocation creep (Ranalli 1995; Karato 2008; Hirth & Kohlstedt 1995a,b). At very high stress regimes, other deformation mechanisms might be present like grain boundary sliding (Kohlstedt & Wang 2001; Hirth & Kohlstedt 2003; Drury 2005; Langdon 2006; Kohlstedt 2007) or Peierls mechanisms (Evans & Goetze 1979; Frost & Ashby 1982; Raterron et al. 2004; Katayama & Karato 2008; Kohlstedt 2007). We only consider diffusion and dislocation creep laws for simplifi-



Figure 2. The shape of the function $\hat{\mathcal{R}}$, similar to Figure 1 but with a different scale, is depicted with a black line. Previous studies have assumed that $\hat{\mathcal{R}}$ was given by the red (dashed) (Ricard & Bercovici 2009), purple (dot-dashed) (Hillert 1965) or blue (double-dot-dashed) (Lifshitz & Slyozov 1961) lines. We arbitrary choose the amplitudes of these three later curves so they change sign with the same slopes.

cations, but other mechanisms could be added. We use

$$\underline{\breve{\varepsilon}} = (a\breve{\tau}^{n-1} + b\mathcal{R}^{-m})\underline{\breve{\tau}}$$
(34)

assuming that the mechanisms of deformation occur in parallel (and $\check{\tau}$ is the second invariant of $\check{\underline{\tau}}$). In (34), the two terms correspond to the dislocation stress-dependent mechanism and to the diffusion grainsize-dependent mechanism. Typically $n \sim 3$ and $m \sim 3$ (Hirth & Kohlstedt 2003; Kohlstedt 2007).

The macroscopic rheology $\tau = 2\eta \dot{\epsilon}$ is obtained by computing the volume average deformation rate and stress (see Ricard & Bercovici (2009) for more details). This can be done numerically but not analytically in the more general case although some variational estimates might be possible as in Hashin & Shtrikman (1963). As a simple case, we assume that the stress tensor at the grainsize level $\underline{\tau}$ is uniform and therefore equal to the macroscopic stress $\underline{\tau}$. This is akin to the Reuss averaging in elasticity (Reuss 1930). In this case,

$$\underline{\dot{\epsilon}} = \underline{\tau} \int_{0}^{+\infty} (a\tau^{n-1} + b\mathcal{R}^{-m}) \breve{\nu}\nu(\mathcal{R}) d\mathcal{R}$$
(35)

and the use of a selfsimilar distribution (3) allows us to express this relation as a function of average grainsize \mathcal{R}_0 and stress τ .

$$\underline{\dot{\epsilon}} = \left(a\tau^{n-1} + b\frac{\lambda_{3-m}}{\lambda_3}\mathcal{R}_0^{-m}\right)\underline{\tau} = a\tau^{n-1}\left(1 + \left(\frac{\mathcal{R}_c}{\mathcal{R}_0}\right)^m\right)\underline{\tau}$$
(36)

where

$$\mathcal{R}_c = \left(\frac{b}{a}\frac{\lambda_{3-m}}{\lambda_3}\tau^{1-n}\right)^{1/m} \tag{37}$$

is the average grainsize at the macroscopic transition between diffusion and dislocation creeps. The macroscopic rheology is thus a function of the mean grainsize \mathcal{R}_0 and the grainsize distribution through λ_{3-m}/λ_3 . Notice that when the average grainsize is \mathcal{R}_c , the macroscopic material is exactly at the transition between diffusion and dislocation, the individual grain with the same radius is still largely in the diffusion regime as $\lambda_{3-m}/\lambda_3 \sim 0.1$ according to (31). The experimentalist that works with a sample containing a distribution of grainsizes cannot measure directly the microscopic rheological factor of diffusion creep *b*, but only get the macroscopic coefficient $b(\lambda_{3-m}/\lambda_3)$. This coefficient together with the rheological factor *a* of dislocation creep, are both temperature dependent with an Arrhenius form and their own activation energies, i.e.,

$$a = A_1 \exp\left(-\frac{E_1}{RT}\right)$$
 and $b\frac{\lambda_{3-m}}{\lambda_3} = A_2 \exp\left(-\frac{E_2}{RT}\right)$, (38)

where R is the gas constant.

In order to provide a simpler expression for the grainsize evolution than the general expression (24), we need to know the value of the partitioning energy factor \tilde{f} . In the absence of direct observations, we can make two simple guesses:

(i) The simplest case is to assume a constant partitioning $\check{f} = f_0$.

The partitioning is the same irrespective of the deformation mechanism itself. In this case the general grainsize evolution would obey

$$\frac{d\mathcal{R}_0}{dt} = \frac{G}{p\mathcal{R}_0^{p-1}} - f_0 \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \underline{\tau} : \underline{\dot{\epsilon}}$$
(39)

(ii) The creation of new grains occurs by formation of subgrain boundaries (Twiss 1977; Derby 1990, 1991; Shimizu 1998a,b, 1999; Bresser et al. 1998; Bresser 2002). This requires the presence of dislocations density and it can be argued that the partition-



Figure 3. The normalized partitioning function f_1/f_0 (solid blue curve) for a mean grainsize varying between 10^{-1} and 10 times the critical grain size at the macroscopic boundary between diffusion and dislocation creep. The transition between diffusion and dislocation creeps occurs through a large range of grainsizes.

ing of energy only occurs during dislocation creep (in each grain, the strain-rate is determined by a mixture of diffusion and dislocation creep, but only the mechanical work done by dislocation creep can be used to form new grains). This means that

$$\frac{d\mathcal{R}_0}{dt} = \frac{G}{p\mathcal{R}_0^{p-1}} - f_0 \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \underline{\tau} : \underline{\dot{e}}_{disloc}$$

$$= \frac{G}{p\mathcal{R}_0^{p-1}} - f_1(\tau, \mathcal{R}_0) \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \underline{\tau} : \underline{\dot{e}},$$
(40)

where

$$f_1(\tau, \mathcal{R}_0) = f_0 \frac{1}{1 + (\mathcal{R}_c/\mathcal{R}_0)^m}.$$
(41)

which is depicted in Figure 3. The function goes from 0 (no grain reduction occurs when the mean grain is largely in the diffusion regime) to f_0 (all the grains are in the dislocation regime and they all shrink). The partitioning function is independent of σ , the spread of the grainsize distribution.

6 APPLICATION TO OLIVINE

The creep map of olivine has been described by various authors (Chopra & Paterson 1981; Karato et al. 1986; Karato & Wu 1993; Hirth & Kohlstedt 2003; Korenaga & Karato 2008). We compiled some reference values for exponents, prefactors and activation energies appropriate for dry olivine in Table 1. The exponents n and m are close to $n \simeq 3.5$ and $m \simeq 3$ for dry olivine although Korenaga & Karato (2008) suggest a larger n exponent, $n \simeq 4.94$ for dislocation creep.

There are few studies and more uncertainty for the grain growth kinetics. As a reference case we use a value of p = 2 taken from Karato (1989) and in agreement with the theoretical model of Hillert (1965), and the form of $G = k_0 \exp(-E_g/RT)$ (see Table 1) is identical to that chosen in Kameyama et al. (1997). This choice based on Karato (1989) leads to a grain growth much faster than what is obtained in Evans et al. (2001) or Faul & Scott (2006) who use p = 4.3 and much larger than what is expected from observa-

tions of natural peridotites. Grain boundary pinning by impurities and minor phases may explain this discrepancy.

Equations (39) and (40) imply the existence of steady state regime, $d\mathcal{R}_0/dt = 0$ where a constant grainsize results from balancing grain coarsening by dynamic recrystallization. If we use (40), coarsening can only be driven by energy subtracted from the dissipation associated with dislocations. In this case, the steady state regime usually called piezometric equilibrium (Van der Wal et al. 1993; Bresser et al. 1998) satisfies

$$\mathcal{R}_0 = \left(\frac{3\gamma}{p}\frac{G}{fa}\right)^{1/(p+1)} \tau^{-(n+1)/(p+1)}.$$
 (42)

This equilibrium, where $\mathcal{R}_0 \propto \tau^{-1.5}$, occurs both in the dislocation and diffusion regimes because even in the diffusive regime, the larger grains in the grainsize distribution are still subject to dynamic recrystallization. If even the energy subtracted from dissipation in the diffusive regime contributes to grainsize reduction, as in (39), another equilibrium can be reached with

$$\mathcal{R}_0 = \left(\frac{3\gamma}{p} \frac{G\lambda_2}{fb\lambda_{3-m}}\right)^{1/(p+1-m)} \tau^{-2/(p+1-m)}.$$
 (43)

In the case p = 2 and m = 3, as p + 1 - m = 0, this equilibrium becomes independent of \mathcal{R}_0 and simply defines a plasticity threshold, $\tau_c = (3\gamma G\lambda_2/pfb\lambda_{3-m})^{1/2}$.

Figure 4, depicts the domains of dislocation dominant and diffusion dominant creeps for olivine at 1923 K. The dot-dashed black and solid blue curves indicate the equilibrium states $(\partial \mathcal{R}_0 / \partial t = 0)$ predicted by (42) and (43) when we choose $f_0 = 1.5 \ 10^{-6} \ (f_0 \ \text{con-}$ trols the intercepts of the theoretical piezometers, not their slopes). All grains below these curves grow until they reach the piezometric curve and reciprocally, above these curves the dynamic recrystallization dominates and the grainsize decreases. Purple squares represent experimental data from Karato et al. (1980) which fit our model for the chosen f_0 . In the diffusion creep domain, no experimental points are available. This is reasonable because if the hypothesis (39) is true, the plasticity threshold forbids any equilibrium in the diffusive regime. Even in the case where the recrystallization is only controlled by the dislocation, i.e., using (40) (Figure 4, blue curve), the equilibrium in the diffusive domain may be difficult to identify. Indeed, as the equilibrium grainsize is stress dependent, $\mathcal{R}_0 \propto \tau^{-1.5}$, the diffusive grainsize dependent rheology appears, at equilibrium, more stress dependent than in the dislocation domain since $\dot{\epsilon} \propto \mathcal{R}_0^{-3} \tau = \tau^{5.5}$. The experimentalist may interpret the data as an indication of grain boundary sliding rather than diffusion.

Figure 4 is computed at T = 1923 K. Other experimental piezometers have been proposed at different temperature and for olivine with different water contents (Post 1977; Ross et al. 1980; Karato et al. 1980; Van der Wal et al. 1993; Zhang et al. 2000; Jung & Karato 2001; Jung et al. 2006). For each set of experiments we can obtain an appropriate value for f_0 . These different values are depicted in Figure 5. We also infer an empirical fit $f_0 = \exp(-2.0(T/1000)^{2.9})$ that agrees with the requirement $0 \le f_0 \le 1$. The partitioning factor decreases significantly with temperature and does not seem to vary much with pressure or water content (f_0 might decrease slightly with water content and increase slightly with pressure). The non equilibrium thermodynamics imposes the form of the evolution equation and bounds the possible values of f_0 , but at this stage, we have not tried to derive this parameter from a microscopic kinetic model.

When the grainsize evolves with time and stress, the rheology



Figure 4. Equilibrium curves using our two approximations. We use the standard set of rheologies displayed in table 1, T = 1923K, P = 1 atm. The green shaded area corresponds to the zone where the rheology is predominantly in the diffusive regime. The equilibrium state (or piezometer) in the dislocative regime goes through the experimental points (Karato et al. 1980). A plasticity threshold is predicted for the case defined by (39) (label f_0).

becomes itself a time dependent function. We illustrate this point in Figure 6. We start with an olivine with very small grainsize (1 μ m) and impose a constant deviatoric stress at a temperature of 1100 K. In this case, the rheology is initially Newtonian, i.e., in the diffusion creep (dashed line). However, the grains coarsen or shrink (at very large stress) and the rheology becomes non-linear. For low deviatoric stresses (below 300 MPa), the equilibrium rheology is dislocative (stress exponent 3). For larger stresses, the grains deform under diffusion creep but as the grainsize is stress dependent, the equilibrium rheology is also non-linear (stress exponent 5.5).

Another illustration of the time dependence of the rheology is provided in Figure 7. We compute a stress profile within an oceanic lithosphere with age 50 Myrs (the temperature is an error function with standard parameters). A uniform strain-rate $\dot{\epsilon} = 10^{-15}$ s⁻¹ is imposed (Goetze & Evans 1979). We assume that the average grainsize is initially $\mathcal{R}_0 = 1 \ \mu$ m and, except in the shallower zone where the stress is limited by Byerlee failure criterion (Byer-



Figure 5. Various experimental piezometers can be fitted by choosing $f_0 = \exp(-2.0(T/1000)^{2.9})$. The fit does not seem to depend much on the pressure or the water content.



Figure 6. Evolution of the rheology with increasing grain size (from 1µm). The shear stress varies initially with the strain-rate as τ (diffusive case), then as $\tau^{1/3.5}$ (dislocative case, for $\tau < 300$ MPa), or as $\tau^{1/5.5}$ (diffusive case with stress dependent grainsize).

lee 1978), diffusion creep prevails. The grainsize then evolves quite rapidly and the strength of the lithosphere increases with time. The lithosphere recovers in a few 10 kyrs. At equilibrium, the lithosphere is in the dislocation regime at large depth (above 50 km). From 20 to 50 km, the presence of small average grainsizes places the rheology in the diffusive regime. In this layer whose deeper limit is marked by a kink in the equilibrium grainsize (see Figure 7, bottom panel), the rheology is in fact more stress-dependent than in the underlying layer undergoing dislocation creep.

7 DISCUSSION AND CONCLUSION

In this paper, we started from simple thermodynamics requirements to derive a very general equation for grainsize evolution. Our method takes into account the spread of the grainsize distribution and considers that, at a given time and around a given position, not

Table 1. Chosen sets of parameters: $(^1$ from Duyster & Stockhert (2001), 2 from Hirth & Kohlstedt (2003), 3 from Kameyama et al. (1997).

Reference case			
Parameter	Value	Unit	
γ	1	$\mathrm{J}\mathrm{m}^{-2}$	Surface tension ¹
$ \begin{array}{c} E_1\\ A_1\\ V_1\\ n \end{array} $	$530 \\ 1.1 \ 10^5 \\ 2 \ 10^{-5} \\ 3.5$	$kJ mol^{-1}$ MPa ⁻ⁿ s ⁻¹ J mol ⁻¹ Pa ⁻¹	Act. Energy (disl.) ² Prefactor ² Act. volume ² disl. exponent ²
$ \begin{array}{c} E_2\\ A_2\\ V_2\\ m\end{array} $	$375 \\ 1.5 \ 10^9 \\ 5 \ 10^{-6} \\ 3$	$kJ \text{ mol}^{-1}$ $\mu \text{m}^m \text{ MPa}^{-1} \text{ s}^{-1}$ $J \text{ mol}^{-1} \text{ Pa}^{-1}$	Act. Energy $(diff.)^2$ Prefactor ² Act. volume ² diff. exponent ²
$E_g \\ k_0 \\ p$	$200 \\ 2.0 \ 10^4 \\ 2$	$kJ mol^{-1} \mu m^p s^{-1}$	Act. Energy (growth) ³ Kinetic factor ³ growth exponent ³



Figure 7. Time dependent stress profile (top) and time dependent grainsize profile (bottom). The initial grain size is $d = 1\mu$ m, a constant and uniform strain-rate is assumed, across a 50 Myrs oceanic lithosphere.

all grains deform with the same creep mechanism. Large grains can be submitted to recrystallization while the small grains can deform then coarsen in the diffusive regime. A corollary of our approach is that the kinetics of coarsening and recrystallisation can be estimated from the grainsize distribution itself. Experimental observations suggest that the growth of large grains is faster than what was considered in the classic theories of Lifshitz & Slyozov (1961) or Hillert (1965).

We infer a grainsize equation of the form

$$\frac{d\mathcal{R}_0}{dt} = \frac{G}{p\mathcal{R}_0^{p-1}} - f_0 \frac{1}{1 + (\mathcal{R}_c/\mathcal{R}_0)^m} \frac{\mathcal{R}_0^2}{3\gamma} \frac{\lambda_3}{\lambda_2} \underline{\tau} : \underline{\dot{\epsilon}}, \qquad (44)$$

where \mathcal{R}_c is the transitional grain radius between diffusion and dislocation. This equation assumes the self similarity of the grainsizes within a lognormal distribution. As already stated, self-similarity is approximately verified experimentally and is the consequence of the equation of grainsize distribution itself (2) whose solutions tend to self-similarity. If we reject this hypothesis, then we need to derive the expressions for $\dot{\mathcal{R}}$ and Γ directly from observations. The difficulty is then not so much with the mathematics than with the experiments: reporting the evolution of the average grainsize but not the evolution of the grainsize distribution itself is not sufficient to constrain the kinetics of grain diffusion and recrystallization.

The rheology of the grained material is naturally non-linear and time-dependent (it could also be anisotropic which is another complexity not accounted for in this paper (Tommasi et al. 2009)). This provides a memory to the mechanical behavior of the lithosphere, an ingredient that is missing to the models of convection with self-consistent plates (i.e. Tackley 2000a,b; Stein et al. 2004). When the grainsize-stress relation is taken into account, we predict that the diffusive rheology is more stress-dependent than the dislocative rheology, in contradiction with traditional rheological models.

At equilibrium, the strain-rate remains an increasing function of the stress and the rheology appears to be monotonic and simply non-linear with an exponent varying from 3.5 to 5.5. This is however, a very partial view of the rich dynamics implied by equation (44). In a realistic time dependent situation, the rheology maybe far from equilibrium and the effective stress–strain-rate relation can be much more complex than at equilibrium. The dynamics is also affected by the feedback between deformation and heat production by viscous dissipation as various parameters are temperature dependent. All these aspects are beyond the scope of the present paper but will be studied in the future.

The equation (44) can be compared to what is used in various publications (Braun et al. 1999; Kameyama et al. 1997; Bercovici & Karato 2003).

$$\frac{d\mathcal{R}_0}{dt} = s \frac{G}{p\mathcal{R}_0^{p-1}} - \frac{\dot{\epsilon}}{\dot{\epsilon}_T} (\mathcal{R}_0 - \mathcal{R}_P), \tag{45}$$

where *s* goes from 1 in the diffusion domain to 0 in the dislocation domain, $\dot{\epsilon}_T$ is an experimental parameter and \mathcal{R}_P the equilibrium grainsize function of stress.

Our formalism (44) differs from what has been used previously (45) by various points. We assume that coarsening occurs even in the dislocation regime while (45) considers that it only happens in the diffusion regime (although Bercovici & Karato (2003) use (45) with s = 1). Using non-equilibrium thermodynamics, we prove that grain reduction is related to the energy dissipated in the system, not to the strain-rate alone. In our model, the term representing recrystallization always reduces the grainsize while in (45), recrystallization facilitates grain coarsening when $\mathcal{R}_0 < \mathcal{R}_P$. At last, we predict the piezometric equilibrium while (45) must include an equilibrium radius \mathcal{R}_P which is not consistently deduced by the model itself.

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