Atomic-scale study of dislocation glide in a model solid solution

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Based on atomic-scale simulation techniques, we study the dislocation pinning mechanism in a dilute Ni(Al) model solid solution. For a solute concentration between 1 and 10 at.%, we found that the pinning of the dislocation on obstacles made of Al pairs is an interaction that operates significantly. The statistics of the dislocation motion is then modified accordingly to the nature of the obstacles and follows modified Mott–Nabarro statistics. Finally, a method to address thermal activation is proposed and exemplified on a periodic row of solute pairs.

1. Introduction

Computer modelling has addressed all aspects of physical metallurgy [1, 2]. While dislocations are extended defects for which mesoscale modelling seems to be the most promising tool [3], atomic-scale modelling proved to be very fruitful for some specific issues, where the elastic theory reaches its limit (e.g. when core interactions are important) [4]. Most studies are concerned with pure metals [5, 6] while a number of classical issues in physical metallurgy concerns alloys [7–10]. Recently, we addressed the question of dislocation glide in concentrated solid solutions, with Ni(Al) as a prototype of such problems [11] (C_{Al} ∼ 5 at.%). One conclusion was that the hardening mechanism cannot reduce to the classical solute dislocation interaction but could rather be attributed to solute–solute pairs, in course of dislocation crossing. Although solid solutions have been studied experimentally for many decades [12], they are still a matter of debate, especially because it is difficult to discriminate between theories due to the limited range of solubilities for alloying elements which have a strong interaction with the dislocations [13]. The classical scheme for describing solid solution hardening is inherited from that

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of dilute solution- and precipitate-hardening. In the Mott–Nabarro model [7–9, 14], the increase in yield stress originates from the long-range elastic interaction between the edge part of the dislocations and the solutes. It leads to a hardening which varies as a power law of the number of solutes per unit area of the glide plane, $C$. A first estimation was made in the late 1950s [7] that showed a linear behaviour, which was later corrected [8, 14, 15] to account for dislocation waviness, leading to $C^{2/3}$. An alternative approach, proposed by Friedel [10, 16], considers the short-range interaction between the dislocation and the pinning centres located in its glide plane. The yield stress then varies as $C^{1/2}$. It is noteworthy that a transition in $C$ has been predicted between both aforementioned statistics [15, 17]: in diluted solid solutions Friedel statistics should apply while in concentrated solid solutions, Mott–Nabarro statistics should become more relevant [18, 19] because in the latter case the dislocations are quasi-straight between the closely spaced obstacles. Here we note that the nature of the obstacles does not change with concentration, i.e. the solutes are assumed to remain isolated in the glide plane. In addition, since the dislocation–solute interaction was thought to be the dominant hardening mechanism, the classical theories model the dislocation glide in solid solution as occurring in the potential of interaction with a ‘frozen’ population of solutes (i.e. solute atoms the positions of which are independent of the dislocation location). In fact, in the case of a concentrated Ni(Al) solid solution ($C = C_{Al} \approx 5$ at.%) we showed that a source of hardening was due to solute–solute short-range repulsion, or more specifically, was due to pairs of solute atoms brought into positions of strong mutual repulsion in the course of dislocation glide. According to that finding, the nature of the obstacles thus changes with concentration since the concentration of these pinning pairs varies as $C_{Al}^2$. Since this $C_{Al}^2$ is much smaller than $C_{Al}$, Friedel statistics should be relevant for the Al pairs and their contribution to hardening should be linear in $C_{Al}$ (F.R. Nabarro is gratefully thanked for this remark made to one of us, GM). From a general point of view, it is noteworthy that, if hardening is due to pairs of Al atoms that are brought closer by dislocation sweeping, edge and screw dislocations should play similar roles by way of contrast with the earlier models where hardening originates from the hydrostatic interaction between the solute atoms and only the edge part of the dislocations. Note also that, due to the limited solubility, experimental data on solid solution hardening can equally well be fitted with $C^{1/2}$ as well as $C^{2/3}$ or even $C$ [20]. In spite of these difficulties, some experimental studies tentatively examined the role of small solute clusters in solid solution hardening and worked out exponents that differ from those shown in the standard statistics, for instance an exponent of 0.57 was found by Wille et al. [13] in Cu(Mn) and Cu(Ge) solid solutions. This contradicts the previous theoretical power laws and thus the debate still remains open.

Molecular dynamics (MD) simulations provide opportunities to work out interesting new elements about the hardening of solid solutions. Indeed the simulations allow us to investigate ideal cases such as a true random solid solution and permit us to test the theories in a more reliable way. This primarily motivates our present investigations. From our previous attempt [11] to simulate dislocation glide in solid solutions with concentrations between 1 and 8 at.%, at finite temperature $T = 300$ K, we identified a linear statistics, i.e. a resolved shear stress $\tau$, proportional to $C_{Al}$, which came as confirmation of the argument about the validity of Friedel
statistics for Al pairs. In the present paper, we work out the details of the pinning effect of Al solutes, i.e. single solutes and pairs in different geometries. In order to do so, a well-controlled geometry of regularly spaced obstacles is generated and its pinning effect on a moving dislocation is investigated. This allows us to clarify the different contributions to hardening (section 3). Then our study is extended to a larger range of concentrations, ranging from dilute to concentrated. The strategy is there to evaluate the density of a given type of obstacle and to weight it by its pinning strength in order to estimate an overall pinning force (section 4). The effect of a finite temperature is studied (section 5) for a regular array of obstacles. Our notation is summarized in Appendix A.

2. Simulation method and simulation cell

To model the interatomic potential, we employ an embedded atom method (EAM) which provides a good compromise between the realism of dislocation simulation and the computational load necessary to handle a system large enough to keep at least part of the complexity of the real alloy. For pure nickel we chose the potential developed by Angelo et al. which has proved to be well adapted to the atomic-scale simulation of dislocations [21, 22]. In particular, the dissociation width of the edge dislocation is quantitatively well reproduced by this potential. In our simulations, the dissociation width equals 3.3 nm, which agrees with the observations of Carter and Holmes [23] who estimate this width to be between 1.8 and 3.6 nm. For pure aluminium we chose the potential proposed by Voter and Chen [24], which has the drawback of yielding a low stacking fault energy (compared, for example, to the potential of Ercolessi et al. [25]), but has the advantage of being written in the very same functional form as that for Ni. The interaction potential between Ni and Al has been developed to account for the physics of alloys NiAl and Ni₃Al. The form and the optimization of this potential are described elsewhere [11]. The reader interested in more detailed information about the simulation technique and physical properties of the NiAl solid solution model we use is referred to [11].

The simulation cell is schematically represented in figure 1. It is oriented so that the horizontal Z-planes are (111) planes while the X-direction is a [110] Burgers vector direction. An edge dislocation is introduced between the two (111) central mid-planes of the cell with a line oriented along the [112] Y-direction. The simulation box size along the different directions, \( i = \text{X, Y, Z} \) is denoted by \( L_i \) and the Ni lattice parameter by \( a_0 \). The dislocation core is dissociated in two partials and the atoms that take part in the partials are recognized by their default configurations in first neighbour positions. Periodic boundary conditions are applied both parallel and perpendicular to the dislocation line. In figure 1, we show an Al pair (filled squares), with its first periodic images (open squares). Due to the periodic boundary conditions imposed along the dislocation line, i.e. in the [112] direction, a chain of obstacles is formed with a spacing \( L \) which equals \( L_y \), the size of the simulation box in the Y-direction (see figure 1).

For the molecular statics (MS) simulations (at 0 K), we use a gradient algorithm, which is a minimization of the total potential energy, achieved through a gradient procedure under a fixed applied shear stress. The applied stress \( \tau_{xy} \) is produced
by adding extra forces to the atoms in the upper and lower Z-surfaces. This external applied stress is incremented by 0.75 MPa and for each increment the gradient procedure is performed till either it converges to a required precision or the dislocation starts to glide in the cell, in which case we consider that the yield stress is reached. In figure 2, the average position of a dislocation has been plotted versus the increasing applied stress. The obstacles were positioned at \( x = 6 \text{ nm} \) in the simulation cell. Apart from the initial stage where the Peierls stress has

Figure 1. Schematic view of the simulation cell containing an edge dislocation and an Al pair (filled squares) along with its first periodic images (open squares).

Figure 2. A plot of the average position of the dislocation core versus the external shear stress during molecular statics simulations at 0 K, for different types of obstacle (see table 1). The labels (A, B and C) are described in the text.
to be overcome to force the dislocation to glide towards the obstacle, the dislocation was found to stop at two different positions (labels A and B in figure 2) because of its dissociation in two partials that interact differently with the obstacle. When the external stress is sufficient to unpin the trailing partial (label C in figure 2), the dislocation glides freely in the simulation box.

For the MD simulations (at 300 K), we integrate the dynamics under a fixed $\tau_c$. It is worth noticing that in MD simulations, the time scale is typically one nanosecond which is far from the time required to model diffusion processes in solid solutions. As a consequence, we consider ideal, purely random, solid solutions and even though this solid solution is supersaturated, precipitation cannot take place.

3. Case of periodic chain of obstacles: strengthening effect of different pinning configurations

3.1. Geometry of significant obstacles

Beside the classical situation of the interaction between an isolated Al atom (see figure 3) and a dislocation, several other configurations involving pairs of Al atoms have been tested (see figure 4). Regular chains of obstacles of given types were generated, with a distance $L$ between obstacles. The shear stress necessary to glide through the array of obstacles is $\tau_c = \alpha \mu b / L$, where $\alpha$, the interaction coefficient, characterizes the strength of the obstacle, $b$ is the Burgers vector and $\mu$ is the shear modulus of pure Ni. The $1/L$ term gives the number of obstacles per unit length of dislocation and $\alpha \mu b^2$ is the maximum force exerted by each obstacle. Among the different configurations, we selected those with highest yield stress. The dislocation may interact with Al atoms in three ways. Either it interacts with isolated Al atoms (figure 3) or it interacts with pre-existing Al–Al pairs (figure 4a and b), or it generates such pairs by its passage (figure 4c–g). The investigated pinning configurations are shown in figures 3 and 4, which represents a portion of the two central $(111)$ planes contiguous to the dislocation glide plane. Atoms in the lower $(111)$ plane are shown as full circles, whereas those in the upper plane are represented by

Figure 3. Portion of the two nearest $(111)$ planes that bound the dislocation glide plane showing the configurations of isolated Al atoms (type I obstacles in table 1) that have the strongest pinning effect. The square symbol is the Al atom whereas the circles are Ni atoms. Open symbols represent atoms situated above the glide plane, the full symbols the ones below the glide plane. The arrows indicate the motion of the Al atom in the course of the shearing process.
empty circles. The Al atoms are squares with larger sizes, in agreement with the size ratio between Al and Ni atoms (14\%). Plotting the normalized yield stress $\tau_c/\mu$ as a function of the normalized obstacle density $b/L$ as done in figure 5 gives a direct measure of $\alpha$.

For isolated Al atoms (hereafter called type I), the strongest configuration is such that the Al atom occupies a site in the (111) plane just above the glide plane (see figure 3a), so as to visit the compression zone of the core in the course of dislocation crossing. The interactions between an isolated Al and the two Shockley partials do not have the same magnitude. The controlling reaction is

Figure 4. Same as figure 3 for the configurations of Al pairs that have the strongest pinning effect (see text for details).
the unpinning of the trailing partial. For a solute atom in the plane below the glide plane (see figure 3b), the pinning coefficient of the trailing partial is weaker (see figure 5a) than when the Al atom is in the plane above, and the pinning strength on the leading partial is negligible since it is of the same order as the Peierls stress of Ni. The strength for single Al atoms in \(\{110\}\) planes that do not bind the dislocation glide plane has been found to be also of the order of the Peierls Ni stress.

The pair configurations shown in figure 4(a) and (b) are first neighbour Al atoms (called type II) placed in the plane above the glide plane. As for isolated Al, the dominant interaction is the one between the trailing partial and the Al pair

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**Figure 5.** The yield stress \(\tau_c\) versus the inverse of \(L\), the distance between pinning centres along the \(Y\)-axis. The computation has been carried out by molecular statics at 0 K in various samples, containing different types of obstacles, detailed in the text. The stress has been normalized by the Ni shear modulus and the distance \(L\) by the Ni Burgers vector. For obstacles in the \(\{111\}\) plane just above the glide plane, the yield stress of the trailing partial is represented as a continuous line (circle symbols); the dashed line (square symbols) corresponds to the leading partial. For obstacles in the \(\{111\}\) plane just below the glide plane, the yield stress of the trailing partial is plotted in (a) and (b) as a dot-dashed line (triangle symbols).
with a strength $\alpha$ which is approximately twice that for single Al. The strength on the leading partial depends on the direction of the dimer bond (e.g. [112] shown in figure 4a, [011] or [101] shown in figure 4b). The same pair configurations in the plane below the glide plane have strength of the same order as obstacles of type I (see the dot-dashed line in figure 5b).

In configurations shown in figures 4(c) and (f), before the passage of the dislocation, the Al atoms are third and second neighbours (called type III), respectively. One Al atom is in the lower ($\langle 1 \bar{1} 1 \rangle$) plane and the other Al atom in the upper plane. As schematically depicted in figures 4(c) and (f), if the edge dislocation glides over this pair, the two solute atoms become first neighbours. This configuration is highly energetic as seen from the fact that the ordered phase in equilibrium with the solid solution of this alloy is Ni$_3$Al, which is L1$_2$ with the Al atoms in second neighbour positions. This energy excess is at the origin of the pinning effect on the dislocation. The glide process of the dislocation over the third neighbour Al pair comprises two stages corresponding to the passage of two Schockley partial

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**Figure 5.** Continued.
dislocations: when the first partial glides over the Al pair, the latter is shifted by the \( a/\sqrt{12} \) Burgers vector of the leading partial and adopts an intermediate configuration; when the second partial glides over the pair, the Al atoms are brought into first neighbour positions which are the most energetic configurations. More precisely, the configuration of figure 4(c) (resp. figure 4f) becomes that of figure 4(d) (resp. figure 4g). Again, the limiting reaction is the unpinning of the second partial from the Al pair.

The nearest neighbour pairs in figures 4(d) and (g) (type IV) proves to constitute also pinning centres with a force of the same order as other pairs. The glide process on type IV obstacle in figure 4(d) conserves the pair of nearest neighbours, but forces the direction of the pair to flip so as to form the configuration in figure 4(e). The glide over this latter configuration forces the dimer to pass through a second neighbour position which is favoured by the stability of the L12 phase. The passage of the trailing partial breaks that bond and thus it involves a pinning force. The different obstacles interact differently with both partials as summarized in table 1, which lists the nature of obstacles, the manner they interact with the dislocation, their interaction coefficients \( \alpha \) and the force range \( w \).

### 3.2. Interaction coefficients at 0 K: origin of the pinning strength

The pinning force of each type of obstacle is given by the \( \alpha \) coefficients in table 1. The physical origin of the pinning force can have various contributions. The classical interpretation considers the interaction between individual solute atoms and the dislocation. This is likely to apply to type I defects. According to elastic theory [27], the interaction potential between an edge dislocation and an impurity is:

\[
W(\vec{r}) = \beta_{cl} \frac{\sin \theta}{r}
\]

where \( \theta \) is the impurity azimuthal departure from the glide plane, \( r \) the distance between the impurity and the dislocation and

\[
\beta_{cl} = \frac{\mu b_p}{3\pi} \frac{1 + v}{1 - v}(v_s - v)
\]
with $\mu$, the shear modulus of Ni, $b_p$, the edge leading partial Burger vector ($b_p = 0.5b$), $\nu$, Poisson’s ratio and $v_s$ and $v$, the atomic volumes of, respectively, the impurity (the solute) and the matrix (the solvent). For Ni(Al) we have $\nu = 0.28$, $v_s = 16.6 \text{Å}^3$, $v = 11 \text{Å}^3$.

Let us consider, as in the simulations, a regular chain of isolated Al atoms at the closest position from the dislocation core, i.e. $z \approx a_0 \sqrt{3}/2 = b \sqrt{3}/8$. This corresponds to the interplane distance between planes that bound the dislocation glide plane. The $X$-axis force sustained by the dislocation and due to one impurity is given by $2\beta_d x z/(x^2 + z^2)^2$, the maximum of which is reached at $x = z/\sqrt{3}$. This force is rewritten as $a \mu b^2$ so as to obtain a pinning strength

$$\alpha = \frac{1 + \nu}{1 - \nu} (v_s - v)/(2\sqrt{3} \pi b^3)$$

which yields numerically in the present case of Ni(Al): $\alpha = 0.06$. This result is much larger than the one obtained from the simulations in figure 5 at zero Kelvin, i.e. $\alpha = 0.009$ for the trailing partial and $\alpha = 0.0062$ for the leading partial. As seen from figure 6(a), the short-range variations of the interaction potential do not vary monotonically with distance between the dislocation and the obstacle, as assumed by an elastic interaction model as in equation (1). On the other hand, for a single impurity in the second nearest plane above the glide plane (dot-dashed line in figure 6a), the long-range repulsive interaction, expected from elastic theory, is recovered. Indeed, for a distance between the average position of the dislocation core and the obstacle larger than 20 Å, the internal energy increases monotonically as the distance decreases. Below this cut-off distance, which corresponds to a distance of $3b$, between the leading partial and the obstacle, the internal energy decreases to reach a minimum when the obstacle is at the centre of the dislocation stacking fault ribbon. It is noteworthy that for all obstacle types, we found that the internal energy is minimum when the defect visits the dislocation stacking fault. This Suzuki effect [16] implies that the pinning of the trailing partial is stronger than that of the leading partial. As may be expected, this short-range interaction is not properly described in the standard elastic theory as in equation (1).

For an obstacle situated immediately above or below the glide plane, in the planes that bound the glide plane, the long-range interaction is found to be attractive whereas from elastic theory one would expect that these interactions would have opposite sign, because of $\sin \theta$ in equation (1). In fact, the position of the glide plane, situated between two atomic planes that are separated by $b \sqrt{3}/2$, is arbitrary. As this fixes the azimuthal angle $\theta$, we have no means to determine accurately this angle for atoms in the nearest atomic planes. This concerns the long-range interaction forces whose strengths are negligible compared to the short-range forces. This can be seen from figure 6(b) and figure 7. Moreover the long-range forces with different directions, i.e. either pulling or retaining the dislocation, are superposed linearly and thus compensate provided the solute concentration is large enough to give a quasi-uniform distribution in a shell that surrounds the dislocation core [9].

In the case of type II dimers, the interactions are expected to result from the size effect of the solute atoms. Indeed, we found that the type II pinning coefficients are approximately twice that of type I obstacles, in agreement with the elastic theory.
where interactions add linearly. The case of type III and IV dimers is different. If we consider third neighbour pairs of solute atoms across the dislocation glide plane, then according to equation (1), the dislocation is attracted by the Al atom in the lower plane and repelled by that in the upper plane. Both effects compensate at first order and thus, from elastic theory, the interaction between the edge dislocation and the dimer is expected to be much weaker than for a single atom. This prediction of the elastic theory is not verified, as shown in table 1. The pinning force for type III and IV is much larger than for single atoms. The physical reason is that in these configurations, the dominant pinning effect is not due to the solute–dislocation elastic interactions, but rather to the solute–solute interaction that is either repulsive or attractive.

Figure 6. (a) Internal energy of our simulation box versus the average position of the dislocation core. The simulation box contains a single obstacle of type I situated either in the plane above the glide plane (full line), in the plane below (dashed line) or in the second nearest plane above (dot-dashed line). (b) The derivative of the internal energy for a type I obstacle in the plane just above the glide plane.
In the plane below the glide plane, we noticed that the pinning forces of dimers are comparable or smaller than that of single Al atoms, depending on which partial is considered. Placing pairs in second nearest neighbouring \( \{1/2,1,1\} \) planes, above or below the glide plane does not lead to a yield stress larger than single Al atoms in the same planes. This will allow us to neglect the role of pairs in configurations that are placed far from the plane above the glide plane. In these cases, the interaction between the dislocation and solutes could be well approached by considering only type I obstacles.

### 3.3. Range of the interactions

In the previous section we quantified the strength of the individual obstacles. In the classical theory of collective pinning [14] another quantity of interest is the range of the interactions. To measure that range, it is sufficient to calculate the variation of the internal energy of a system under a prescribed stress containing a dislocation which adapts its position. The internal energy is obtained from the sum of all the atomic potential energies at the convergence of each step of the energy minimization procedure under increasing applied stress. This has been done in figure 6(a) for type I obstacle. The calculation of the first derivative of the internal energy with respect to the average position of the dislocation is plotted in figure 6(b). The positive part of that quantity corresponds to the internal force which is opposed to the dislocation displacement. The range of the interaction between an obstacle of a given type and the dislocation is estimated from the distance which separates the maximum of the force and the nearest position where the force falls to zero. The calculation has also been carried out for different pairs, which is exemplified in figure 7. The results for all types are reported in table 1. As expected from simple considerations on the size of the obstacles, the range of interaction is of order of the atomic spacing for type I obstacles, and 2–3 atomic spacings for type II, III and IV pairs.

![Plot of the internal force versus the dislocation average position. The obstacle is either a first neighbour pair (type II) (solid line) or a third neighbour pair (type III) (dashed line).](image)
4. Solute hardening in random solid solution at 0 K

4.1. Density of different obstacles

From the previous results, solid solution hardening has been shown to be mainly due to the solute obstacles contained in planes that bind the glide plane. It is interesting to count the number of significant interactions with each type of obstacle in the plane above the glide plane. We denote by $n_i$ the number of configuration per atomic site along the dislocation line, where the obstacle type is labelled by the subscript $i$. We saw in section 3.2 that a single Al above the glide plane could pin both partials, so we obtain simply $n_I = 2$.

In case of obstacles made uniquely of pairs, we have first to enumerate the different pinning configurations. The pair configurations of type II may have three different bond directions and can pin either the leading partial or the trailing one. This makes six different possibilities to place a type II obstacle per atomic site along the dislocation line, so that $n_{II} = 6$. By contrast one can consider that the interaction of type III obstacles and the leading partial is negligible. Depending on the relative location of Al atoms in the planes that bound the glide plane we found only two different possibilities for type III obstacles to pin the trailing partial, so $n_{III} = 2$. For type IV, this depends strongly on both the direction of the pair bond and the location of Al atoms. For the leading partial, we found two significant interactions while for the trailing partial there are three possibilities. The total of interactions thus rises to $n_{IV} = 5$. The total number of possibilities of interaction with Al–Al pairs is thus $n_{II} + n_{III} + n_{IV} = 13$.

4.2. Dislocation flexibility

For a chain of isolated Al atoms above the glide plane, the interaction coefficient $\alpha$ is $0.9 \times 10^{-2}$ (see figure 5). The bending of a dislocation reaches a maximum at the middle of the interval between obstacles, given roughly by $\alpha L/4$. To reach a value of $b$, $L$ must be larger than $L_b = 4b/\alpha = 556b$ (138 nm). According to figure 5(a), the corresponding resolved shear stress (around 2 MPa) is comparable to the intrinsic lattice friction of a few MPa. So the hardening due to a chain of isolated obstacles, with $L > L_b$, is negligible. For $L < L_b$ the bending can be neglected and the length of the dislocation segment between pinning points can be considered as equal to $L$. Similar considerations for pairs lead to $L_b = 236b$ and to a corresponding yield stress of 12 MPa. It is noteworthy that the maximum bending between two obstacles increases as the distance $L$. The number of sites that a single Al can occupy on a single partial is $N_y = L_y/(b\sqrt{3})$. For both partials, the proportion of Al at fixed $L_y$ is thus $1/(n_I N_y)$ for a regular chain. Assuming the same proportion in the solid solution would involve $C_{Al} = 1/(n_I N_y)$. In this solid solution, the mean distance between a single Al alongside the $Y$-axis is $L = L_y$. Replacing $L_y$ by $L_b$ in $N_y$ gives $C_{Al} = 0.3\%$. Since we choose to work on larger concentrations $C_{Al} > 1\%$, this indicates that in the present study, the dislocation cannot bend freely between two obstacles far apart as Friedel statistics would assume but is closer to the Mott–Nabarro assumption of a dislocation described by a sequence of quasi-straight segments.
To obtain a more precise idea of the dislocation behaviour in a random solid solution, we use the criterion of Labush as proposed in [9]. From our results in previous sections, we can estimate the Labush factor for the different types of obstacles $\beta = \alpha b^2 / (2c w^2)$ where $c$ is the density of obstacles along the dislocation line. For single solutes, taking $\alpha < 0.01$, $w = 1.8b$ and $C_{AI} = 1\%$ and considering that $c = n_1 C_{AI}$, we found $\beta < 0.08$. The factor $n_1$ in the atomic concentration of obstacles comes from the fact that both partials may be pinned with a strength of the same order. The value found for $\beta$ is rather close to the case studied by Mott and Nabarro [9], where $\beta \approx 0$ which means that the dislocation is nearly straight. The Friedel limit corresponds to $\beta = 1$ which means either a weak concentration or strong pinning obstacles. For the sake of simplicity, we assume that all pairs have the same strength and force range: $\alpha < 0.021$ and $w = 4b$. Then one can suppose that $c = (n_{II} + n_{III} + n_{IV}) C_{AI}^2$ which eventually gives an upper bound of $\beta$ that is $\beta < 0.55$. We thus work on a system that is far from solid solutions where Friedel statistics applies. This result emphasizes that it is not possible to determine the solid solution statistics without working out the details of individual obstacles. According to our present study, our first expectations in [11], made on the basis of a yield stress linear in solute concentration, proved to be hasty.

The typical picture of the dislocation moving inside the solid solution is shown in figures 8(a)–(c) for different Al concentrations. The three pictures figures 8(a), (b) and (c) correspond to an external shear stress just below the yield stress of the solid solutions, i.e. 70, 120 and 200 MPa, respectively. One notes that the bows formed along the partials are much less extended than $L_b$ introduced above, which confirms Mott–Nabarro statistics in contrast to Friedel statistics. The amplitude of static waves along the partials is smaller than the dissociation width (i.e. 13b in our simulations for pure Ni) but is still larger than b. This seems to scale as a few b and we do not notice a dependency on the external stress.

4.3. Simulations of solid solutions

To avoid size effects in the simulation of a random solid solution, we chose $L_x = 344b$ which allowed us to reduce the interaction between an obstacle and its periodic images so that this interaction could not generate a hardening larger than 4.8 MPa for pairs (see figure 5b–d). This is even smaller for type I obstacles. The yield stress observed in the simulations is larger than this value (see figure 9) so the interaction between periodic images can be neglected (for the details of size effects, see Appendix B).

We generated systems with concentrations from 2 to 10%. For each concentration, we generated different distributions of Al atoms. As done for the regular chains, the external shear stress is gradually increased and when the dislocation moves freely we let it glide for five passages in the simulation box in order to probe possible stronger pinning configurations. The simulation box along the $X$-axis was $L_x = 40b$, so we assumed that when the dislocation glided over $5 \times L_x$ we had reached the maximum of the applied shear stress for the considered random distribution of Al. For each concentration, the calculations of the yield stress from various distributions have been reported in figure 9. Each open symbol corresponds to a given random distribution.
4.4. A model for solid solution hardening at 0 K

In the approximation of a quasi-straight dislocation line (section 4.2) where the two partials are rigidly bound, we can now test different models for the hardening of solid solutions at 0 K. In the case of a quasi-straight dislocation (see equation 30 in [9]), the yield stress is given by the Mott–Nabarro statistics:

$$\tau_c = \frac{1}{b^2} \left( \frac{c^2 w f_m^4}{b \Gamma} \right)^{1/3}$$

with the same notation as in [9], so that $\Gamma = \mu b^2 / 2$, $f_m = \alpha \mu b^2$ and $c$ is the density of obstacles. In section 3.3, we calculated the parameters of this formula for different obstacles. However, as one has to account for forces on both partials from various configurations of different obstacles, the formula in equation (2) is in principle not directly applicable. We propose to extend the Mott–Nabarro statistics to account for obstacles of a different nature in interaction with a dissociated dislocation core.

As a first step, we derive a model in the case of a quasi-straight undissociated dislocation which interacts with only one type of obstacle. This is the same situation as in [9, 15], but we proceed differently since we do not make any use of Labush’s method of distribution functions (see [14] for the original theory and [15] for its application to dislocation glide in solid solutions). In spite of the fact that our theory is less rigorous than Labush’s one, this will permit us to extend the calculation to a case more complex than the original one.

Let us consider a dislocation which can be separated into segments, the average length of which is $L_c$. Each segment corresponds to the wavelength of the static wave of our dislocation (see figures 8 and 10). The wavy shape of the dislocation
results from the coexistence of regions where the total force is pushing with regions where the total force is retaining the dislocation (see figure 10). So this wavelength $L_c$ characterizes the solute distribution in the neighbourhood of the dislocation. The difference of density between the pulling and retaining obstacles distributed along the dislocation segment $L_c$, in a ribbon $2w$ centred on the average core position, is given by the random statistics $\sqrt{2cwL_c/s}$. Here we denote by $s$ the atomic elementary surface in plane (11): $s = 3\sqrt{3}b^2/8$. Since we want to estimate the resistance to the external shear stress, we consider the case where the dislocation is stopped by an excess of retaining obstacles. The total force exerted by the obstacles on the dislocation is: $-(L_c/L_c)^{1/2}f_md\sqrt{2cwL_c/s}$. Submitted to an external shear stress $\tau$, the total variation of enthalpy due to a displacement $dx$ of the dislocation reads

$$dH = \tau b L_c dx - \frac{L_c}{L_c} f_md\sqrt{2cwL_c/s} + \delta E_t,$$

(3)

where $\delta E_t$ is the variation of the energy line tension due to the displacement. We assume that the displacement $dx$ is small compared to the force range $w$ and that the displacement does not involve a change in distribution forces around the dislocation core. To evaluate $\delta E_t$, we consider our dislocation as a sine-like string with line tension $\Gamma = \mu b^2/2$. By comparison of the three pictures in figure 8, we see that the amplitude of the dislocation bending, denoted by $A$, does not vary significantly with concentration nor with the external stress. On the other hand, one notes that the spatial extension along the dislocation line of the static waves clearly decreases as the concentration increases.

Figure 9. Variation of the yield stress versus the Al concentration, computed from the present simulations with different Al random distributions for each concentration (different symbols for each simulation) and different models: Mott–Nabarro statistics for isolated Al (dot-dashed line), for Al pairs (dashed line) and the modified Mott–Nabarro statistics to account for various types of obstacle (continuous line).
Since one assumes a sine-like shape for the dislocation segment $L_c$, the retaining and pushing regions have similar spatial extension. Provided that the dislocation remains pinned by the retaining obstacles, the only parts of the dislocation that can move significantly under the Peach–Kohler force are the regions where the sum of forces is pushing the dislocation. As depicted in figure 10, in the region where the obstacles retain the dislocation, the displacement is small compared with the region where the obstacles push the dislocation in the direction of the Peach–Kohler force. In a first-order approximation, we neglect the displacement in the region where the obstacles retain the dislocation. During a displacement $dx$ of the average position of the segment $L_c$, the dislocation displacement in the region where the obstacles push the dislocation is thus given by $2dx$, since the retaining and pushing regions have similar spatial extension. Thus, at the end of the displacement, we have $A = A_f \approx A + 2dx$. Since the string energy, $E_t$, is given by $(\Gamma/2)q^2 A^2$, where $q = 2\pi/L_c$ is the wavevector, we obtain:

$$
\delta E_t = L_y \frac{\Gamma(2\pi)^2}{2L_c^2} (A_f^2 - A^2) \approx L_y \frac{4\Gamma\pi^2}{L_c^2} A dx,
$$

(4)

Figure 10. Schematic view of the dislocation (thick line) interacting with a random distribution of obstacles whose forces are either pushing or retaining the dislocation. The small arrows indicate the different directions of forces. The notation is defined in the text. In the course of an average displacement $dx$, the dislocation is deformed because of the inhomogeneous force distribution. This leads to an increase of the curvature of roughly $2dx$ (see text above equation 5).
to first order in $dx$. The derivative of the enthalpy variation with respect to $dx$ is, thus to first order in $dx$:

$$\frac{dH}{dx} = L_y \left[ \tau b - f_m \sqrt{\frac{2cw}{sL_c}} + \frac{4\pi^2 A^2}{L_c^2} \right].$$

The position of the dislocation becomes unstable when $dH/dx > 0$, which defines the critical stress above which the dislocation glides:

$$\tau b = f_m \sqrt{\frac{2cw}{sL_c}} - \frac{4\pi^2 A^2}{L_c^2}.$$  \hfill (6)

The segment length $L_c$ depends on the distribution of Al encountered by the dislocation core. The yield stress corresponds to the maximum of $\tau b$ with respect to $L_c$, i.e. when all the dislocation segments of every length $L_c$ can be released under the stress $\tau$. This maximum value is reached for $L_c = L_c^*$:

$$L_c^* = \left( \frac{16\pi^2 \Gamma A}{f_m \sqrt{2cw/s}} \right)^{2/3}.$$ \hfill (7)

which decreases as the density of obstacles as expected from the simulations shown in figure 8. Combining equations (6) and (7), and assuming that $A \approx w$ leads to

$$\tau_c = \frac{3}{8b} \left( \frac{c^2 w f_m^4}{2\pi^4 \Gamma^2} \right)^{1/3},$$ \hfill (8)

which gives exactly the same power law as the formula derived by Nabarro in [9] (see equation 3), apart from a geometric factor. Equation (8) has also the same form as Mott’s result [8] when $A = w = b$. A first-order development of $A$ as a function of $w$ gives a linear function in which the constant term is zero since $A$ must go to zero as $w$. Indeed, for an external stress that is negligible, when the obstacles are punctual, i.e. $w = 0$, the dislocation is straight. According to the simulations (figure 8), it is reasonable to take $A = w$. With our notation, equation (8) may be rewritten as

$$\tau_c = \frac{3\mu b}{8} \left( \frac{c^2 w \alpha^4}{\pi^2 \delta^2} \right)^{1/3}.$$ \hfill (9)

We now extend the theory to different types of obstacles and to the dislocation which is dissociated into partials, considered as rigidly bounded. For the quantities $f_m$, $\alpha$ and $w$, we introduce the subscript $i$, which represents a single configuration of a given type in table 1, and a subscript $j = (1, 2)$, for the partial which is concerned. The concentration $c_j$ is either $C_{Al}$ or $C_{Al}^2$, since it does not depend on which partial is considered. Along the dislocation segment $L_c$, we consider the statistics of every
type of obstacle such that equation (5) becomes:

\[
dH = L_y dx \left[ \tau b - \sum_{i,j} f_{m,i,j} \sqrt{\frac{2c_j w_{i,j}}{s L_c}} + \frac{4\pi^2 W \Gamma}{L_c^2} \right],
\]

(10)

where we introduced the quantity \( W \) that is the average amplitude of the static wave of the whole core, that is taken as \( W = \sum_{i,j} c_i w_{i,j} / \sum_i c_i \). The line tension \( \Gamma \) is still given by \( \mu b^2 / 2 \), where \( b \) is the total Burgers vector since the partial dislocations are assumed to be rigidly bound. The coefficients \( w_{i,j} \) and \( \alpha_{i,j} \) are given in table 1. The same arguments as previously developed lead to the optimal wavelength:

\[
L_c^* = \left( \frac{16\pi^2 \Gamma W \sqrt{s}}{\sum_{i,j} f_{m,i,j} \sqrt{2c_j w_{i,j}}} \right)^{2/3}.
\]

(11)

The final formula for the yield stress of a random Al distribution with different types of obstacles is:

\[
\tau_c = \frac{3}{8b} \left( \frac{\sum_{i,j} f_{m,i,j} \sqrt{2c_j w_{i,j} / s}}{(2\pi^2 W \Gamma)^{1/3}} \right)^{4/3}.
\]

(12)

This is rewritten within our notation as follows:

\[
\tau_c = 3\mu b \left( \frac{\sum_{i,j} \alpha_{i,j} \sqrt{2c_j w_{i,j} / s}}{8(\pi^2 W)^{1/3}} \right)^{4/3}.
\]

(13)

The concentration of isolated obstacles is given by \( c_i = C_{\text{Al}} - 2C_{\text{Al}}^2 \) to avoid the double counting of pairs in the plane just above the glide plane. For pairs, we have simply \( c_i = C_{\text{Al}}^2 \). The corresponding yield stress is plotted in figure 9 as a continuous line and shows satisfactory agreement with our MS simulations. The length \( L_c^* \) (equation 11) varies between 164\( b \) for \( C_{\text{Al}} = 2\% \) and 7\( b \) for \( C_{\text{Al}} = 10\% \). Since \( L_y > L_c^* \), the simulation cell is large enough to reduce a possible size effect due to the fluctuations of the random statistics (Appendix B). Equation (13) represents an extension of the Mott–Nabarro statistics to the case of various types of obstacle. It is noteworthy that, not only is the upward curvature of the yield stress as a function of \( C_{\text{Al}} \) well approached by the present expression, but even the absolute value of the yield stress is calculated with good precision without any rescaling factor. Accounting for single solute pinning only would lead to underestimating the yield stress. The results for Mott–Nabarro statistics applied to pairs only is reported in figure 9 as a dashed line. The result is closer to our simulations than that for isolated Al. We are thus in a range of concentrations where the pair contribution dominates but both contributions have to be taken into account and properly summed to obtain a good quantitative result. According to Mott–Nabarro statistics applied to pairs only, an exponent 4/3 of the Al concentration can be worked out, replacing \( c \) by \( C_{\text{Al}} \) in equation (2) or (6). Since the main source of hardening comes from pair contribution, this exponent, larger than 1, involves the upward curvature of the overall yield stress, equation (13).
In the present development, we did not consider the possibility of Al triplets. In the range of concentrations where such a contribution could dominate that of pairs, an upward curvature should appear clearly since the exponent of $C_{Al}$ in equation (2) would be 2. According to the agreement between the simulations and the model presented above, we may reasonably expect that the triplet contribution is not significant in the range of concentrations studied here. At larger concentrations in Al, the dislocation pinning of triplets may thus be expected to operate. As we noted for pairs and isolated solutes, the contributions of various cluster size gather to enhance the hardening.

5. Temperature effect

5.1. Dislocation dynamics: stop and go motion

Thermal activation is often invoked to explain the unpinning of dislocations when $\tau < \tau_c$. This theory proves suitable when the activation energy $E_a$ for passing obstacles is much larger than the thermal fluctuations fixed by $k_B T$, i.e. the time spent at the top of the barrier $E_a$ is negligible in comparison to the occupation of the bottom. In the case of dislocations under an applied stress $\tau$, the activation energy $E_a$ is defined as $E - \tau V_{act}$, the difference between the internal energy barrier $E$ and the work of the Peach–Kohler force which involves the activation volume $V_{act}$. In the range of stress-temperature such that $E_a/k_B T \gg 1$, the process is thermally activated and it proves difficult to verify by MD whether the statistics of the activation law applies in such a case. In contrast, when $E_a/k_B T$ is of the order of unity, we are able to observe transient arrest periods, whose durations $\delta t$ we call waiting times (see figure 12). When the applied stress decreases below the critical stress, and below a value $\tau_c(T)$, $\delta t$ becomes larger than the duration of our simulations. $\tau_c(T)$ is defined as the finite-temperature yield stress of our system.

In his work on solution hardening [15], Nabarro derived a theory to work out the finite-temperature yield stress $\tau_c(T)$ from the zero-temperature yield stress $\tau_c$. As mentioned by Nabarro himself [9], the theory disagrees with experiments. For instance, the power law in concentration does not agree with results on silver alloys, obtained by Basinski et al. [29]. It would be worth carrying out MD simulations on the thermal activation of the glide process in solid solutions to attempt to improve the theory. This kind of study would require exceedingly long simulations of MD on large simulation cells, so we are not able to present a complete study on that precise point. However, as this proves to be fruitful for the case of zero temperature in section 3, our first step is to understand the consequences of thermal fluctuations on the crossing of simple regular chains with periodic condition along the $Y$-axis (see figure 1). In figure 11, we used MD to compute $\tau_c(T)$ as a function of the distance $L$ between obstacles, for a regular chain of type III obstacles (see section 3). The simulations were performed at 300 K. When the shear stress is large enough, the dislocation is unstopped. The external shear stress has been decreased till the dislocation remains pinned for the duration of our simulations, i.e. a few nanoseconds. Whether a dislocation will move or not depends on the distribution of initial atomic velocities. It is, however, possible to carry out
a statistical treatment over the initial atomic velocities. This results in a distribution of critical stresses rather than a well-defined yield stress. Typically, five independent simulations were performed for each of the seven values of $L$ depicted in figure 11. For a given $L$, these results are represented as error bars in figure 11. We can see that the temperature strongly lowers the resistance of type III Al pair. This decrease is analyzed thoroughly in section 5.2.

The atomistic processes involved are illustrated in figure 12. In our simulation box, we placed four pairs of type III (type III, figure 4c), aligned along the moving dislocation. The distance between the obstacles is fixed at $L = 14b$ which corresponds to a critical shear stress $\tau_c(300\,\text{K}) = 40\,\text{Mpa}$. The detailed atomistic mechanism is followed by measuring the length of the Al–Al bonds. For $\tau$ above $\tau_c(T)$, the dislocation passes so that the Al pairs follow the same trajectory and while crossing they are brought into first neighbour distance. The different stages of the Al–Al bonds length are labelled in figure 12(a). Initially the pairs are third neighbours (label A), they become second neighbours when the leading partial unpins (B) and then first neighbours (type IV, figure 4d) when the trailing partial crosses (C). The dislocation makes one loop in the simulation box between C and D, then the leading partial forces the Al atoms to a distance slightly smaller than first neighbours’ distance. After the second passage of the dislocation, the pairs are again first neighbours (type IV, figure 4e). Each further dislocation crossing (F) separates the pairs by one $b$.

When the external stress is slightly below the critical value $\tau_c(T)$, the dislocation stops (see figure 12b). As in figure 12(a), we labelled the different stages. Although the two first stages (A and B) are the same as in figure 12(a), the subsequent evolution differs. The pairs are tentatively put in nearest neighbour positions (C) but eventually return to a second neighbour configuration (D). The latter is preferred to the former because of the repulsive character of the Al–Al interaction in the core of the dislocation. It is noteworthy that the bond lengths of the four independent

![Figure 11. Critical shear stress of a regular chain of type III obstacles spaced by $L$ at 300 K (square symbols) and at 0 K (full line).](image-url)
pairs have been plotted in the same figures 12(a) and (b). The different plots are superposed so that they are hardly distinguishable which shows that the pairs flip simultaneously and that the dislocation crosses coherently the chain of obstacles.

For an applied stress between $\tau_c(T)$ and $\tau_c$, the dislocation may stop for a waiting time that decreases sharply as the shear stress increases and falls to an extremely low value when the shear stress is larger than $\tau_c$.

5.2. Models for temperature effect

We propose different models for the temperature effect on the crossing of simple regular chains. First, we denote us by $f(x)$ the force sustained by the dislocation due
to the presence of obstacles in its course. At equilibrium, \( \tau b L + f(x) = 0 \) where \( x \) is the distance between the obstacle and the partial which interacts significantly, and \( L \) is the distance that separates two neighbouring obstacles. The distance \( x \) is the same for all periodic images of the obstacle. We develop \( f(x) \) around its maximum \( (x_m) \):

\[
f(x) = f_m + \frac{1}{2} \frac{d^2 f}{dx^2}(x_m)(x - x_m)^2.
\]

At a distance corresponding to the range of the interaction \( w \), as estimated in section 3.3, we can estimate that \( f(x_m \pm w) \approx 0 \), thus \( d^2 f/dx^2(x_m) = -2f_m/w^2 \). The force then has a quadratic form. Furthermore the yield stress is such that \( f_m = -\tau_c bL \). Consequently, the equilibrium is given by:

\[
(\tau - \tau_c)bL + \frac{\tau_c bL}{w^2}(x_{eq} - x_m)^2 = 0.
\]

The stable equilibrium position is:

\[
x_{eq} = x_m - w\sqrt{(\tau_c - \tau)/\tau_c}.
\]

Integrating the total force \( \tau b L + f(x) \) between the equilibrium position \( x_{eq} \) and the limit of stability \( x_m \) gives the total work \( W \) one must provide to overcome the potential barrier:

\[
\delta W = -\frac{2b}{3} L w^{1/2}(\tau_c - \tau)^{3/2}.
\]

We assume that the whole dislocation moves coherently (which is confirmed by the simulations in section 5.1). From the bottom of this potential, i.e. \( x = x_{eq} \), to overcome the barrier, the kinetic energy of the dislocation core, \( E_c \), must rise by \( -\delta W \). The initial kinetic energy of each atom along the \( X \)-direction is, on the average, \( k_B T/2 \), which gives a thermal energy per unit length of dislocation \( k_B T/(2b\sqrt{3}) \), where \( b\sqrt{3} \) is the distance between two atoms along the dislocation line. Assuming that the thermal fluctuations are optimally distributed to pass the barrier, we obtain the yield stress at a finite temperature:

\[
\tau_c(T) = \tau_c - \left( \frac{3k_B T}{4\sqrt{3}b^2 w} \right)^{2/3} \tau_c^{1/3}.
\]

It is worth noting that this formula shows a \( 2/3 \) power of temperature, similar to the theory proposed by Nabarro in [9] for thermal effect on solid solution hardening. However, in our case of a regular obstacle chain, the difference \( \tau_c - \tau_c(T) \) is proportional to \( \tau_c^{1/3} \) whereas for solid solutions a \( \tau_c^{2/3} \) is expected [9]. Since the exponent of \( \tau_c \) is less than 1, there is a boundary value of \( \tau_c \) below which \( \tau_c(T) \) becomes negative and thus the predicted hardening becomes negligible. This boundary increases as the \( 2/3 \) power of temperature. Equation (18) is plotted versus \( 1/L \) in figure 11 (labelled by A, dot-dashed line). Clearly, this model fails to reproduce the numerical simulations. The discrepancy probably stems from the assumption that the thermal activation of the dislocation is associated with
a rigid body motion. One has to account for the finite line tension and the fluctuations in shape of the dislocation to describe more accurately the thermal activation process. In order to do so we assume that the free segments of the dislocation trailing partial can be approached by a string which oscillates around an equilibrium position as stationary waves of amplitude $A$ (see figure 13). The $X$-component of the total force exerted on the pinned point of the dislocation line is

$$F_p = \tau b L - \tau_c b L + N$$

where the first term is the Peach–Kohler force, the second term is the maximum of the obstacle force and the third one is due to fluctuations of the line tension. Assuming a sine form for the dislocation wave and denoting by $A$ the amplitude of oscillations and $q_n = n\pi/L$ the wave momentum with mode number $n$, then the maximum force due to the line tension projected on $X$ is $N = 2q_n A \Gamma$. According to the standard oscillating string theory, the vibration energy of the mode $n$ is $L q_n^2 A^2 \Gamma / 2$ which is assumed to reach $k_B T / \Gamma$ at thermodynamical equilibrium of the segment. The total force $F_p$ is zero when $\tau$ reaches $\tau_c(T)$ that is:

$$\tau_c(T) = \tau_c - 2/b \sqrt{k_B T \times \Gamma / L^3}.$$  \hspace{1cm} (19)

As the unpinning of the second partial is the controlling reaction, $\Gamma$ is here given by $\Gamma = \mu b_p^2 / 2$ where $b_p \approx b / 2$. Since $\tau_c = \alpha \mu b / L$, equation (19) can be rewritten as:

$$\tau_c(T) = \tau_c - \frac{2 \tau_c^{3/2}}{b} \sqrt{k_B T \times \Gamma / (\alpha \mu b)^3}.$$ \hspace{1cm} (21)

We note that the temperature exponent in equation (20) is 1/2 whereas it was 2/3 in equation (18). The exponent of $\tau_c$ in the second term at the right-hand side in
equation (20) is now larger than 1, by contrast to equation (18). Consequently, the predicted $\tau_c(T)$ in equation (20) remains positive for small values of $\tau_c$. The variation of the theoretical critical stress predicted by equation (20) is plotted in figure 11 (labelled by B, dashed line) and proves to be in good agreement with the simulation results, indicating that the oscillating string model captures the physics of thermally activated unpinning of the dislocation in interaction with a regular array of obstacles. The agreement can be improved by adjusting the value of the partial line tension $\Gamma$ upon atomic simulations in order to take into account the effect of the dissociated core [30]. A rescaling of 80% of the line tension allows us to obtain the plot corresponding to label C in figure 11. For small values of $L$ in equation (19) (i.e. large $\tau_c$ in equation 20), $\tau_c(T)$ can be negative, which indicates a drawback of the model that is ascribed to the fact that the dislocation cannot be considered as an oscillating string when the obstacles are close to each other. The combination of the rigid string model (model A) and of the oscillating string model (model B) might allow us to improve our approach of the temperature effect. The nucleation of thermally activated double kinks to pass the Al barrier could also be a possibility. In the case of not too small spacings between obstacles, the agreement between the simulations and the oscillating string model shows that it is surely the fluctuations of the dislocation line which operates the unpinning. In this case, it is interesting that the stronger the obstacle chain the larger the softening due to the temperature. Hence the effect of Al dimers is expected to be more attenuated than that of isolated solutes. Such a softening of the strongest obstacles could involve a change in the dislocation glide statistics as a function of the temperature.

6. Discussion

Atomistic simulations shed new light on the problem of solute hardening [7] to which F.R.N. Nabarro has made major contributions. The complexity of developing reliable interatomic potentials for alloys postpone the direct application of these simulations to a variety of solid solutions. However, this approach can guide us in the understanding of the atomistic aspects which cannot be accounted for by the classical elastic theories. The main finding of our paper is that solute pairs, whether they cross the glide plane or not, play a dominant role in hardening concentrated solid solutions; this is reflected in the variation of the yield stress as a function of solute concentration. The pair contribution is enhanced by the various possibilities of the bond direction. We have also demonstrated how to take advantage of such simulations to address the still open problem of the thermal activation of dislocation motion in a solid solution. The computer simulations described here can be seen as sophisticated thought experiments which helped us to generalize the Mott–Nabarro model for solid solution hardening: the extension we propose accounts for the effect of atom pairs and attempts to incorporate thermal activation effects. Since the computer simulations depend obviously on the interatomic potentials, our results need to be confirmed in different situations, i.e. different potentials and different systems [31]. A comparison with experiments proves difficult because of the low solubility limit in the solid solutions similar
to Ni(Al). By way of contrast in the computer simulations, the diffusion process of the solute can be ignored because the simulation time scale is much smaller than the jump rate of defects. This allowed us to properly address the case of one dislocation in a single monocrystal of purely random solution. In the experiments, the presence of the grain boundaries, other impurities or other dislocations may impede the analysis of the solid solution hardening.

The phenomena identified in this paper pinpoint two main issues which should not be overlooked: the importance of chemical bonds in concentrated solid solutions, and the role of the detailed local crystallography. One can expect for instance that short-range order hardening could be explored along the same lines, and the case of solute hardening in bcc metals, still largely unsolved, should also benefit from these approaches in the near future [32].

Appendix A: Notation

Our notation for constants and variables is summarized in tables 2 and 3, respectively.

Appendix B: Size effects in simulation of a solid solution

The size effects in solid solution simulations are mainly due to our finite size along the dislocation line, i.e. in the $Y$-direction, $L_Y$. This involves interactions between

<table>
<thead>
<tr>
<th>Symbols</th>
<th>Meaning</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a_0$</td>
<td>Ni lattice parameter</td>
<td>3.52 Å</td>
</tr>
<tr>
<td>$b$</td>
<td>Norm of Burgers vector for a edge dislocation</td>
<td>$b = a_0/\sqrt{2}$</td>
</tr>
<tr>
<td>$b_p$</td>
<td>Edge part of partial Burger vectors</td>
<td>$b/2$</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Ni shear modulus in $(111)$</td>
<td>74 600 MPa</td>
</tr>
<tr>
<td>$v$</td>
<td>Ni atomic volume</td>
<td>11 Å$^3$</td>
</tr>
<tr>
<td>$v_s$</td>
<td>Al atomic volume</td>
<td>16.6 Å$^3$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Symbols</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$</td>
<td>Pinning strength normalized with respect to $\mu b^2$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Labush factor</td>
</tr>
<tr>
<td>$\beta_{el}$</td>
<td>Coefficient of the hydrostatic work</td>
</tr>
<tr>
<td>$f_m$</td>
<td>Obstacle force</td>
</tr>
<tr>
<td>$L$</td>
<td>Nearest neighbour distance between obstacles</td>
</tr>
<tr>
<td>$L_c$</td>
<td>Dislocation core wavelength in a distribution of obstacles</td>
</tr>
<tr>
<td>$L_c'$</td>
<td>Dislocation core wavelength in the strongest distribution of obstacles</td>
</tr>
<tr>
<td>$n_i$</td>
<td>Number of significant obstacles of type $i = (I, II, III, IV)$</td>
</tr>
<tr>
<td>$\tau$</td>
<td>Resolved external shear stress</td>
</tr>
<tr>
<td>$\tau_c$</td>
<td>Yield stress at 0 K</td>
</tr>
<tr>
<td>$\tau_c(T)$</td>
<td>Critical shear stress at temperature $T$</td>
</tr>
<tr>
<td>$w$</td>
<td>Range of obstacle force</td>
</tr>
</tbody>
</table>
the dislocation and the solutes and their periodic images along $Y$ which may artificially pin the dislocation. As seen in section 3.2, according to our calculations, the pinning effect of a regular chain becomes negligible compared to the yield stress of our solid solutions, whenever the obstacle spacing is larger than 100 times the Burgers vector. We could thus have fixed our box size along $Y$ to $100b$ but a more subtle size effect has a different origin than the interaction between periodic images.

The number of obstacles (isolated Al atoms or pairs of Al) is denoted by $N_o$. As we assume purely random disorder, the fluctuation of this number is $\Delta_o = \sqrt{N_o}$. The random statistics is satisfactorily represented to the condition that $\Delta_o \ll N_o$. If this condition is not satisfied, then our dislocation may encounter an abnormally high or low concentration of obstacles either an excess or a lake of \( \sqrt{N_o}/2 \). The size effect on the statistics is tightly related to the characteristics of the interactions between the obstacles and the dislocation. The number of partials concerned as well as the obstacle density and the range of the interaction should be involved in the estimation of the required size. In order to account for these features, a model is proposed in section 4.4. According to this model, the scale of the statistical fluctuations along the dislocation line, $L^x$, is smaller than the size of the simulation cell along the $Y$-direction. We can thus reasonably expect that the inequality $\Delta_o \ll N_o$ is satisfied.

To reach the large $Y$ sizes, we have reduced the dimensions in the other directions and particularly in the $X$-direction. The size $L_x$ is only 10 times larger than the separation width between partials which could involve elastic interactions between the core dislocation and its periodic images along $X$. However the distance between periodic images is still $100b$, which is larger than the distance between the dislocation core and the solute atoms in solid solutions. The interaction between the dislocation core and the solute atoms is thus expected to dominate the possible interactions between the core dislocation and its periodic images along the $X$-direction.

References