Shear-coupled grain boundary migration assisted by unusual atomic shuffling

Liang-Liang Niu1,2, Ying Zhang1, Xiaolin Shu1, Fei Gao2, Shuo Jin1, Hong-Bo Zhou1 & Guang-Hong Lu1

Shear-coupled grain boundary (GB) migration can be an efficacious mechanism to accommodate plastic deformation when the grain size of polycrystalline materials goes small. Nevertheless, how this kind of GB motion comes into play at the atomic level has not been fully revealed. Here, we have investigated the shear-coupled migration (SCM) of typical [100] group symmetrical tilt GBs in bcc W using atomistic simulations. Depending on GB character, the SCM is found to proceed via dislocation slipping in the ⟨100⟩ or ⟨110⟩ mode with striking shear strength difference between them. We demonstrate that there exists an unusual atomic shuffling along the tilt axis, which greatly assists SCM to operate in the easier ⟨110⟩ mode instead of the ⟨100⟩ one. The present results highlight the significant role of GB character in the atomistic SCM process and contribute to the future design and fabrication of high-performance materials in GB engineering.
The results are in excellent agreement with those found in bcc Fe37.

Figure 1. Grain boundary energy as a function of misorientation angle for the [100] STGBs in bcc W.

STGBs in W are revealed utilizing atomistic simulations based on MD methods. The implications of this work are discussed.

Results and Discussion
Grain boundary energy and structure. The energy and structure of GBs in bcc, fcc and hcp metals have been under extensive investigation for several decades37–42. Here we studied 36 [100] STGBs with misorientation angle θ varying between 5.5° and 84.5°. Almost all distinct GB structures have been covered within this interval for this particular GB group. Figure 1 shows the extra energies of these GBs as a function of misorientation angle, among which the GB energies of low-angle GBs (with θ → 0° and θ → 90°) are much lower than the high-angle ones. Notably, $\Sigma 5(01\bar{3})$ and $\Sigma 5(01\bar{2})$ GBs shown as the minor cusps stand out among the high-angle category. The results are in excellent agreement with those found in bcc Fe37.

Typical equilibrium GB structures are illustrated in Fig. 2. At the lower end of the misorientation angle, the low-angle GBs present a discrete array of (100) dislocations with a kite-shaped core. Actually, these GBs can also be described by the type 2 tilt wall model39,40 proposed previously. At the higher end of the range, the low-angle GBs consists of an array of revolved $\frac{1}{2} (110)$ dislocations (see Fig. 1l–p), which well fit the type 1 tilt wall model39,40. As the effective misorientation angle increases, the dislocation cores approach each other and begin to overlap. The GB structures thus formed can be readily characterized by the structural unit model43.

Previous studies have demonstrated that GBs with noticeable energy cusps are favorable. In hcp metals, these favorable GBs correspond to coherent STGBs which serve as the base boundary structure for the STGBs neighboring them39,40, whereas in bcc and fcc metals, these favorable GBs are consisted of some basic structural units and a combination of these favored structural units forms other STGBs with complexity. As shown in Fig. 1, the $\Sigma 5(01\bar{3})$ and $\Sigma 5(01\bar{2})$ STGBs are clearly favored for this GB group. We have identified five basic structural units as labelled in Fig. 2. Specifically, structural units A and A' corresponds to the (100) and (110) components of the perfect lattice, B and C are the basic units of the $\Sigma 5(01\bar{3})$ and $\Sigma 5(01\bar{2})$ STGBs, while B' is a filled version of B due to the added row of atoms to minimize the system potential energy. The STGBs can be determined by a combination of structural units from the favored boundaries neighboring them. The STGBs in Fig. 1a–e are formed by the structural units of the $\Sigma 5(01\bar{3})$ and that of perfect lattice, STGBs in Fig. 1f–j come from a combination of structural units of $\Sigma 5(01\bar{3})$ and $\Sigma 5(01\bar{2})$, while the STGB in Fig. 1k is composed of structural units of $\Sigma 5(01\bar{2})$ and that of the perfect lattice. For example, $\Sigma 85(01\bar{1}3) = 10A + 2B$, $\Sigma 13(01\bar{5}) = 2A + 2B$, $\Sigma 97(05\bar{1}3) = 2B + 4B' + 2C$ and $\Sigma 53(05\bar{9}) = 2A' + 4C$.

Stick-slip dynamics of SCM. Figure 3 presents the variation of shear stress and GB displacement as a function of time at 0.1 K employing a velocity of 0.2 m/s. Note that altering the velocity by one order of magnitude has practically no influence on the elementary mechanisms of SCM, whereas the low temperature can dramatically suppress the effect of thermal noises which might jeopardize the examination of SCM. It can be seen that the yz component of the shear stress ($S_{yz}$) exhibits a sawtooth behavior characteristic of stick-slip dynamics and the GB migrates in a stop-and-go manner. The stress-induced GB migration can be either positive or negative depending on GB character. According to Cahn37, two branches of GB geometry-dependent coupling factor $\beta$, characterized by a linear relation, $\beta = T/N$, between normal GB displacement $N$ and parallel GB translation $T$, can be determined based on the slip direction of GB dislocations. The (100) branch, $\beta_{1(100)} = 2 \tan(\frac{\pi}{4})$, whereas $\theta$ is the misorientation angle, corresponds to the positive GB migration, whereas the (110) branch, $\beta_{1(110)} = -2 \tan(\frac{\pi}{4} - \frac{\theta}{2})$ dictates the negative one.

We found that for STGBs migrating in the (110) mode, there exists an inherent stress component along $xz$ (see the methods section for the crystallographic directions) in the ground state GB structure. As shown in Fig. 3e,k,l, this stress component changes periodically during the SCM process. Accompanying each step of GB displacement, $S_{xz}$ exhibits a precipitous drop accompanied by the sign change of $S_{xz}$ and then $S_{xz}$ may change sign or
maintain that sign until the next GB displacement. Interestingly, no noticeable $S_{xz}$ values have been recorded in GBs moving in the $\langle 100 \rangle$ mode and we thus did not show them. The atomistic origins of these phenomena are illustrated in the following sections.

**Dimensionality of atomic displacement associated with SCM.** Next, we provide further atomic insights into the SCM regarding the dimensionality of the atomic displacements. Figure 4 shows four typical atomic displacement maps at the end of the MD simulation run, among which STGBs in (a) and (b) migrate in the $\langle 100 \rangle$ mode and those in (c) and (d) migrate in the $\langle 110 \rangle$ one. Interestingly, for STGBs moving in the $\langle 110 \rangle$ mode (see Fig. 4c,d), the $x$-axial displacement maps demonstrate that concomitant with each step of the SCM, atoms locating at opposite sides of the GB dislocation glide plane are found to undergo transient displacements along the tilt axis towards opposite directions. Moreover, atoms in the regions, which are both traversed by the GB migration and sandwiched between neighboring dislocation glide planes, experience the equal total $x$-axial displacements. These effects give reasonable explanations as to why $S_{xz}$ changes sign (Fig. 3e,k,l). Note that the $y$-axial displacement maps indicate the rotation of dislocation core structures, while the $z$-axial displacement maps imply the existence of shear deformation of the region swept by the GB. In short, the atomic displacement maps demonstrate that atomic shuffling along the tilt axis in the SCM process contributes to the sign change of the shear stress component along the tilt axis, while the same shear stress component has not been observed when there is no such shuffling. The important role played by this kind of atomic shuffling will be subsequently discussed.

**SCM assisted by the tilt axis-oriented atomic shuffling.** In previous sections, we have pointed out that there exists an unusual tilt axis-oriented atomic shuffling associated with an intrinsic shear stress component along the tilt axis during the SCM process of the $\langle 110 \rangle$ branch GB. Here we try to understand how this kind of atomic shuffling correlates with the shear strength, which constitutes the central results of this work. We plot in Fig. 5 the critical stress, SCM mode, and whether tilt axis-oriented atomic shuffling exists as a function of

---

**Figure 2.** Typical equilibrium GB structures with basic structural units constituting the (a) $\Sigma 85(01–13) = 8.8^\circ$, (b) $\Sigma 25(01–7) = 16.3^\circ$, (c) $\Sigma 13(01–5) = 22.6^\circ$, (d) $\Sigma 85(02–9) = 25.1^\circ$, (e) $\Sigma 5(01–3) = 36.9^\circ$, (f) $\Sigma 97(05–13) = 42.1^\circ$, (g) $\Sigma 29(02–5) = 43.6^\circ$, (h) $\Sigma 29(03–7) = 46.4^\circ$, (i) $\Sigma 97(04–9) = 47.9^\circ$, (j) $\Sigma 5(01–2) = 53.1^\circ$, (k) $\Sigma 3(05–9) = 58.1^\circ$, (l) $\Sigma 89(05–8) = 64.0^\circ$, (m) $\Sigma 13(02–3) = 67.4^\circ$, (n) $\Sigma 37(05–7) = 71.1^\circ$, (o) $\Sigma 65(07–9) = 75.7^\circ$ and (p) $\Sigma 41(04–5) = 77.3^\circ$. The left and right images are equilibrium GB structures projected onto the yz and xy planes, respectively. Atoms are colored according to their potential energies and a darker color indicates higher energy.
misorientation angle. It can be seen that the critical stress for SCM generally decreases with increasing misorientation angle accompanied by mode change from $\langle 100 \rangle$ to $\langle 110 \rangle$ at $\theta \in (31.9^\circ, 36.9^\circ)$. Clearly, a correlation between GB energy and critical stress cannot be established. Surprisingly, seven GBs, i.e., $\Sigma 73(03\text{--}8)$, $\Sigma 97(05\text{--}13)$, $\Sigma 29(02\text{--}5)$, $\Sigma 29(03\text{--}7)$, $\Sigma 97(04\text{--}9)$, $\Sigma 5(01\text{--}2)$ and $\Sigma 5(01\text{--}2)$, which are supposed to move in the $\langle 110 \rangle$ mode according to previously defined criterion, fall into the region of $\langle 100 \rangle$ mode with an extraordinarily high critical stress. 'Coincidentally', no atomic shuffling (no shear stress component along the tilt axis) can be observed for all these GBs moving in the $\langle 100 \rangle$ mode, whereas a relative atomic shuffling of $\approx a/2$ (see Fig. 4c,d) was detected for all GBs moving in the $\langle 110 \rangle$ mode. In Fig. 6 and Supplementary Videos, we present further evidence to support this finding by showing the SCM of the $\Sigma 53(05\text{--}9)$ STGB at 0.1 K and the $\Sigma 97(05\text{--}13)$ STGB at 300 K. Figure 6a and Supplementary Video S1 show that no atomic shuffling was observed at the initial stage of the shearing run and the stress accumulates to a value as high as $\approx 7$ GPa. We observed slight positive GB migration in the $\langle 100 \rangle$ mode. Surprisingly, at $\approx 7.2$ ns, we observed pronounced structural transformation accompanied by atomic shuffling along the tilt axis and the GB quickly moves down in the $\langle 110 \rangle$ mode with a dramatically reduced stress level. As far as the $\Sigma 97(05\text{--}9)$ STGB is concerned, it moves in the $\langle 100 \rangle$ mode at 0.1 K in the 12 ns simulation (see Fig. 3f and Supplementary Video S2). However, once the temperature is increased to 300 K, the change of SCM mode was observed once again (see Fig. 6b and Supplementary Video S3). Specifically, at $\approx 2.3$ ns, structural transformation accompanied by atomic shuffling along the tilt axis was observed and the GB quickly moves down in the $\langle 110 \rangle$ mode with a greatly reduced stress level. We thus conclude that the tilt axis-oriented atomic shuffling can effectively assist the STGBs to move in a more energy-efficient ($\langle 110 \rangle$) mode.

We further recognize that the discovered correlation between atomic shuffling, the SCM mode and critical stress can be traced back to the equilibrium GB structure (see Fig. 2). The $\langle 100 \rangle$ branch STGBs are highly mirror-symmetric to the GB interface, and it takes tremendous time to break the mirror symmetry at low temperatures due to the suppressed thermal fluctuations, thus they are more likely operate in the difficult $\langle 100 \rangle$ mode. As a matter of fact, we observed more frequent SCM mode changes from $\langle 100 \rangle$ to $\langle 110 \rangle$ induced by structural transformation with increasing temperature (e.g. Fig. 6b and Supplementary Video S3) due to the fact that high temperature increases the probability of symmetry breaking. In contrast, this kind of mirror symmetry is broken for the $\langle 110 \rangle$ branch STGBs at the beginning due to the shift along the tilt axis, they can, therefore, readily

Figure 3. Shear stress and GB displacement as a function of time for representative STGBs of (a) $\Sigma 85(01\text{--}13) = 8.8^\circ$, (b) $\Sigma 25(01\text{--}7) = 16.3^\circ$, (c) $\Sigma 13(01\text{--}5) = 22.6^\circ$, (d) $\Sigma 85(02\text{--}9) = 25.1^\circ$, (e) $\Sigma 5(01\text{--}3) = 36.9^\circ$, (f) $\Sigma 97(05\text{--}13) = 42.1^\circ$, (g) $\Sigma 29(02\text{--}5) = 43.6^\circ$, (h) $\Sigma 29(03\text{--}7) = 46.4^\circ$, (i) $\Sigma 97(04\text{--}9) = 47.9^\circ$, (j) $\Sigma 5(01\text{--}2) = 53.1^\circ$, (k) $\Sigma 89(05\text{--}8) = 64.0^\circ$ and (l) $\Sigma 13(02\text{--}3) = 67.4^\circ$ at 0.1 K. The $xz$ component of the shear stress is shown only in the $\langle 110 \rangle$ branch STGBs because it is zero for the $\langle 100 \rangle$ branch.
operate in the easier (110) mode. The present finding pinpoints the important effect of inherent GB character on the atomistic mechanism and critical stress of SCM. We remark that the atomistic mechanism revealed here is expected to be generally applicable to the SCM process of all STGBs in bcc metals and how the present mechanism depends on temperature, velocity or defects will be a subject of future work.

Conclusions
In this work, we have conducted extensive atomistic simulations to investigate the SCM behavior of [100] STGBs in bcc W at low temperature. Analysis of the equilibrium GB structures shows that the STGBs can be adequately described by the previously proposed structural unit model and dislocation model. We show that the SCM proceeds through two modes of GB dislocation slipping (along (100) or (110) directions) depending on GB character. The shear strength of the (100) branch GBs is much higher than the (110) ones. Surprisingly, we reveal that

Figure 4. Atomic x-, y- and z-axial displacement maps of the final snapshot for (a) $\Sigma 221(01\overline{2}1) = 5.5^\circ$, (b) $\Sigma 13(01\overline{5}) = 22.6^\circ$, (c) $\Sigma 5(01\overline{3}) = 36.9^\circ$ and $\Sigma 145(08\overline{9}) = 83.3^\circ$ at 0.1 K. The atomic x-axial displacement map is shown only in the (110) branch STGBs because it is zero for the (100) branch.
there exists an unusual tilt axis-oriented atomic shuffling that can effectively assist SCM to operate in the easier \langle 110 \rangle mode. This kind of atomic shuffling is directly responsible for the sign change of the shear stress component along the tilt axis. We attribute the observed atomistic mechanism to the inherent GB equilibrium structure. Transformation of the GB structure may also simultaneously change the SCM mode. The present work points out the strong dependence of the atomic-scale SCM process on GB character. The atomistic mechanism presented here suggests another factor be taken into consideration when designing and fabricating novel GB-containing materials in GB engineering.

Figure 5. Critical stress, SCM mode and whether tilt axis-oriented atomic shuffling exists for the SCM process of the \{100\} group STGBs at 0.1 K. The tilt line separates the two modes of SCM. The transition from \langle 100 \rangle mode to \langle 110 \rangle one with increasing misorientation angle was traditionally thought to occur at \theta \in (31.9°, 36.9°). The seven STGBs surrounded by the oval box that are normally expected to move in the \langle 110 \rangle mode actually move in the \langle 100 \rangle one due to lack of atomic shuffling along the tilt axis.

Figure 6. Shear stress and GB displacement as a function of time for the special case of (a) \sum_{53}(05–9) = 58.1° at 0.1 K and (b) \sum_{97}(05–13) = 42.1° at 300 K. The left and right inset images are the respective xy-plane projection of the equilibrium GB structures before and after the GB structural transformation.
Methods
A parallel MD package, LAMMPS \(^{44}\) is used to explore the SCM process of the [100] series STGBs constructed according to ref. 37. Firstly, based on coincident site lattice theory, two grains with different crystallographic orientations are brought together to obtain the bicrystal. Then, the upper and lower grains are translated with respect to each other in-plane by different magnitudes, followed by an atom deletion process to avoid close contact between atoms. Notably, due to the in-plane translation process, the mirror-symmetry of the STGBs is frequently observed to be broken, which has been reported in previous studies \(^{45,46}\). The size of the system varies depending on the specific GB type and typical bicrystal contains thousands of atoms. The interatomic potential for W-W interaction is described in previous works \(^{47,48}\). Periodic boundary conditions are applied parallel to the GB interface (x and z directions). For the convenience of imposing shear stress on neighbouring dynamic atoms, top and bottom slabs of ~12 Å thick (more than twice the cutoff radius) are frozen by setting the interatomic force to zero. An exemplary run of the SCM process and the crystallographic relation are illustrated in Fig. 7. A constant shearing velocity is subsequently applied to the upper slab along +z direction with the lower slab remaining fixed. Note that all shearing is conducted at 0.1 K with a velocity of 0.2 m/s unless specified otherwise. The position of the GB interface is tracked according to the maximum potential energy of atoms in the GB region. In addition, the standard viral stress tensor expression implemented in LAMMPS is employed to calculate the shear stress.

References
1. Li, C. H., Edwards, E. H., Washburn, J. & Parker, E. R. Stress-induced movement of crystal boundaries. Acta Metall 1, 223–229 (1953).
2. Legros, M., Gianola, D. S. & Hemker, K. J. In situ TEM observations of fast grain-boundary motion in stressed nanocrystalline aluminum films. Acta Mater 56, 3380–3393 (2008).
3. Zhang, K., Weertman, J. R. & Eastman, J. A. Rapid stress-driven grain coarsening in nanocrystalline Cu at ambient and cryogenic temperatures. Appl. Phys. Lett 87, 061921 (2005).
4. Mompiou, F., Legros, M. & Caillard, D. Direct observation and quantification of grain boundary shear-migration coupling in polycrystalline Al. J. Mater. Sci 46, 4308–4313 (2011).
5. Schäfer, J. & Albe, K. Competing deformation mechanisms in nanocrystalline metals and alloys: Coupled motion versus grain boundary sliding. Acta Mater 60, 6076–6085 (2012).

Figure 7. Exemplary SCM processes in the \(\Sigma 13(01–5) = 22.6^\circ\) STGB moving in the \(\langle 100 \rangle\) mode. The rest STGBs follow a similar crystallographic relationship and shearing procedure. Atoms are colored according to their potential energies and a darker color indicates higher energy. The violet marker is used to portray the shear deformation.
6. Rupert, T. J., Gianola, D. S., Gan, Y. & Hemker, K. J. Experimental observations of stress-driven grain boundary migration. Science 326, 1686–1690 (2009).
7. Zhang, Y., Sharon, J. A., Hu, G. L., Ramesh, K. T. & Hemker, K. J. Stress-driven grain growth in ultrafine grained Mg thin film. Scr. Mater. 68, 424–427 (2013).
8. Sharon, J. A., Su, P. C., Prinz, F. B. & Hemker, K. J. Stress-driven grain growth in nanocrystalline Pt thin films. Scr. Mater. 64, 25–28 (2011).
9. Mompiou, F., Caillard, D. & Legros, M. Grain boundary shear–migration coupling—I. In situ TEM straining experiments in Al polycrystals. Acta Mater. 57, 2198–2209 (2009).
10. Bobylev, S. V., Morozov, N. F. & Ovid’ko, I. A. Cooperative Grain Boundary Sliding and Migration Process in Nanocrystalline Solids. Phys. Rev. Lett 105, 055504 (2010).
11. Cahn, J. W., Mishin, Y. & Suzuki, A. Coupling grain boundary motion to shear deformation. Acta Mater. 54, 4953–4975 (2006).
12. Cahn, J. W. & Taylor, J. E. A unified approach to motion of grain boundaries, relative tangential translation along grain boundaries, and grain rotation. Acta Mater. 52, 4887–4898 (2004).
13. Velasco, M., Van Swygenhoven, H. & Brandl, C. Coupled grain boundary motion in a nanocrystalline grain boundary network. Scr. Mater. 65, 151–154 (2011).
14. Mishin, Y., Suzuki, A., Ubeuraqa, B. P. & Voter, A. F. Stick-slip behavior of grain boundaries studied by accelerated molecular dynamics. Phys. Rev. B 80, 224101 (2009).
15. Ivanov, V. A. & Mishin, Y. Dynamics of grain boundary motion coupled to shear deformation: An analytical model and its verification by molecular dynamics. Phys. Rev. B 78, 064106 (2008).
16. Frolow, T. Effect of interfacial structural phase transitions on the coupled motion of grain boundaries: A molecular dynamics study. Appl. Phys. Lett 104, 211905 (2014).
17. Yoshida, H., Yokoyama, K., Shibata, N., Ikuhara, Y. & Sakuma, T. High-temperature grain boundary sliding behavior and grain boundary energy in cubic zirconia bicrystals. Acta Mater. 52, 2349–2357 (2004).
18. Winning, M. In-situ observations of coupled grain boundary motion. Philos. Mag. 87, 5017–5031 (2007).
19. Gorkaya, T., Molodov, D. A. & Gottstein, G. Stress-driven migration of symmetrical (110) tilt grain boundaries in Al bicrystals. Acta Mater. 57, 5396–5405 (2009).
20. Gorkaya, T., Burlet, T., Molodov, D. A. & Gottstein, G. Experimental method for true in situ measurements of shear-coupled grain boundary migration. Scr. Mater. 63, 633–636 (2010).
21. Gorkaya, T., Molodov, K. D., Molodov, D. A. & Gottstein, G. Concurrent grain boundary motion and grain rotation under an applied stress. Acta Mater. 59, 5674–5680 (2011).
22. Caillard, D., Mompiou, F. & Legros, M. Grain-boundary shear-migration coupling. II. Geometrical model for general boundaries. Acta Mater. 57, 2390–2402 (2009).
23. Trautz, T. T., Adland, A., Karma, A. & Mishin, Y. Coupled motion of asymmetrical tilt grain boundaries: Molecular dynamics and phase field crystal simulations. Acta Mater. 60, 6528–6546 (2012).
24. Wang, J., Misra, A. & Hirth, J. P. Shear response of γ2,3 (112) twin boundaries in face-centered-cubic metals. Phys. Rev. B 83, 064106 (2011).
25. Honeker, E. R., Foiles, S. M., Holm, E. A. & Olmsted, D. L. Phenomenology of shear-coupled grain boundary motion in symmetric tilt and general grain boundaries. Acta Mater. 61, 1048–1060 (2013).
26. Frolow, T., Olmsted, D. L., Asta, M. & Mishin, Y. Structural phase transformations in metallic grain boundaries. Nat. Commun. 4, 1899 (2013).
27. Frolow, T., Divinski, S. V., Asta, M. & Mishin, Y. Effect of Interface Phase Transformations on Diffusion and Segregation in High-Angle Grain Boundaries. Phys. Rev. Lett. 110, 255502 (2013).
28. Hirth, J. P. & Pond, R. C. Steps, dislocations and disconnections as interface defects relating to structure and phase transformations. Acta Mater. 44, 4749–4763 (1996).
29. Wan, L. & Wang, S. Shear response of the Σ9<110> (221) symmetric tilt grain boundary in fcc metals studied by atomistic simulation methods. Phys. Rev. B 82, 214112 (2010).
30. Rajabzadeh, A., Mompiou, F., Legros, M. & Combe, N. Elementary Mechanisms of Shear-Coupled Grain Boundary Migration. Phys. Rev. Lett 110, 265507 (2013).
31. Rajabzadeh, A., Legros, M., Combe, N., Mompiou, F. & Molodov, D. A. Evidence of grain boundary dislocation step motion associated to shear-coupled grain boundary migration. Philos. Mag. 93, 1299–1316 (2013).
32. Rajabzadeh, A. et al. The role of disconnections in deformation-coupled grain boundary migration. Acta Mater. 77, 223–235 (2014).
33. Khater, H. A., Serra, A., Pond, R. C. & Hirth, J. P. The disconnection mechanism of coupled migration and shear at grain boundaries. Acta Mater. 60, 2007–2020 (2012).
34. Merkle, K. L., Thompson, L. J. & Phillipp, F. Collective Effects in Grain Boundary Migration. Phys. Rev. Lett 88, 225501 (2002).
35. Borovikov, V. et al. Coupled motion of grain boundaries in bcc tungsten as a possible radiation-damage healing mechanism under fusion reactor conditions. Nucl. Fusion 53, 063001 (2013).
36. Yu, K. Y. et al. Removal of stacking-fault tetrahedra by twin boundaries in nanotwinned metals. Nat. Commun. 4, 1377 (2013).
37. Tschopp, M. A. et al. Probing grain boundary sink strength at the nanoscale: Energetics and length scales of vacancy and interstitial absorption by grain boundaries in α-Fe. Phys. Rev. B 85, 064108 (2012).
38. Rittner, J. D. & Seidman, D. N. (110) symmetric tilt grain-boundary structures in fcc metals with low stacking-fault energies. Phys. Rev. B 54, 6999–7015 (1996).
39. Wang, J. & Beyerlein, I. J. Atomic structures of symmetric tilt grain boundaries in hexagonal close packed (hcp) crystals. Modell. Simul. Mater. Sci. Eng. 20, 024002 (2012).
40. Wang, J. & Beyerlein, I. J. Atomic Structures of [0-110]Symmetric Tilt Grain Boundaries in Hexagonal Close-Packed (hcp) Crystals. Metallurgical and Materials Transactions A 43, 3556–3569 (2012).
41. Wang, J., Li, N. & Misra, A. Structure and stability of Σ3 grain boundaries in face centered cubic metals. Philos. Mag. 93, 315–327 (2013).
42. Tschopp, M. A. & McDowell, D. L. Structures and energies of Σ3 asymmetric tilt grain boundaries in copper and aluminium. Philos. Mag. 87, 3147–3173 (2007).
43. Sutton, A. P. & Vitre, V. On the Structure of Tilt Grain Boundaries in Cubic Metals I. Symmetrical Tilt Boundaries. Philos. T. Roy. Soc. A 309, 1–36 (1983).
44. Plimpton, S. Fast Parallel Algorithms for Short-Range Molecular Dynamics. J. Comput. Phys. 117, 1–19 (1995).
45. Ochs, T., Beck, O., Eksasser, C. & Meyer, B. Symmetrical tilt grain boundaries in body-centred cubic transition metals: An ab initio local-density-functional study. Philos. Mag. A 80, 351–372 (2000).
46. Wojdyr, M., Khalil, S., Liu, Y. & Szlufarska, I. Energetics and structure of <001> tilt grain boundaries in SiC. Modell. Simul. Mater. Sci. Eng. 18, 075009 (2010).
47. Ackland, G. J., Tichy, G., Vitre, V. & Finnis, M. W. Simple N-body potentials for the noble metals and nickel. Philos. Mag. A 56, 733–756 (1987).
48. Juslin, N. & Wirth, B. D. Interatomic potentials for simulation of He bubble formation in W. J. Nucl. Mater. 432, 61–66 (2013).
Acknowledgements
This work is supported by the National Natural Science Foundation of China (NSFC) through Grant No. 11405006 and 51371019. G.H.Lu acknowledges support from China National Funds for Distinguished Young Scientists through Grant No. 51325103.

Author Contributions
L.L.N. performed the calculation and wrote the initial manuscript under the supervision of Y.Z. and X.S. F.G. and G.H.L. conceived the research. S.J., H.B.Z., F.G. and G.H.L. commented and revised the manuscript. G.H.L. directed the project. All authors contributed to the scientific discussion of the results and reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Niu, L.-L. et al. Shear-coupled grain boundary migration assisted by unusual atomic shuffling. Sci. Rep. 6, 23602; doi: 10.1038/srep23602 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/