The hierarchical energy landscape of screw dislocation motion in refractory high-entropy alloys

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1. Introduction

Dislocations, the line defects in a crystal lattice, dictate the strength and deformation behaviors of materials [1]. Their glide that must overcome an energy barrier due to lattice resistance can be facilitated by applied mechanical stress, thermal activation, or both of them. At zero temperature, a minimum stress (Peierls stress) is required to vanish the energy barrier (Peierls barrier) and move a straight dislocation in the absence of thermal activation [2,3]. As the temperature is raised, dislocation motion can occur via kink-pair mechanism at a stress level below the threshold value [4]. In pure metals, the three (constant) energy barriers associated with Peierls mechanism, kink-pair nucleation, and kink gliding render a smooth potential energy landscape (PEL) that regulates the dislocation motion. Because of the easy glide of edge dislocation in body-centered cubic (bcc) metals, the sluggish motion of screw resulting from the kink-pair process (considerably high energy barrier) is the essential origin of high strength [5]. The promotion of kink-pair nucleation and increased concentration of thermal kinks, with increasing temperature, would smear out the rate-limiting effects of the kink-pair mechanism, which is closely correlated to the apparent temperature dependence of strength of bcc metals [6,7]. In dilute bcc alloys, the addition of solute atoms that lowers the double kink nucleation barrier and increases the barrier for kink lateral migration [8,9] would roughen and tilt the PEL, in which kink-pair activation process is favored and kink propagation is impeded.

In contrast to bcc pure metals and dilute alloys, the refractory high entropy alloys (HEAs) [10] such as MoNbTaW [11] and MoNbTi [12], exhibit a gradual and slow decrease in strength with increasing temperature to half of their melting points. The high strength and its weak temperature dependence are ascribed to dislocations that operate in concentrated solid solutions. The locally high chemical fluctuations in HEAs inevitably influence the element mechanisms of dislocation motion and their corresponding energy barriers, and hence the structure of PEL. It is conceivable that a high PEL ruggedness and large variation of energy barriers can be induced by the composition fluctuations, and impact the dominance of individual mechanisms governing dislocation motion. For example, the ubiquitous chemical inhomogeneity creates many local trapping sites over the dislocation migration path, giving rise to a rugged energy landscape for dislocation movement [13]. The wide distribution of dislocation core energy indicating the

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High-entropy alloys (HEAs) with concentrated solid solutions are conceived to possess a rugged atomic and energy landscape in which dislocation motion necessarily proceeds to accommodate mechanical deformation. Fundamental questions remain as to how rough the energy landscape is and to what extent it can be influenced by the local ordering of their constituent elements. Here, we construct and report the potential energy landscape (PEL) governing screw dislocation motion in refractory HEAs that reveals a hierarchical and multilevel structure with a collection of small basins nested in large metabasin. This striking feature pertaining to HEAs exerts a trapping force and back stress on saddle point activations, retarding dislocation movement. By introducing chemical short-range order, the energy landscape is smoothed but skewed to different degrees that shifts the rate-limiting process from kink-gliding to kink-pair nucleation. The chemical disorder-roughened PEL in HEAs, analogous to structural disorder induced in metallic glasses, signifies the role of various barrier-hopping processes underlying the extraordinary mechanical behaviors of the two distinct groups of materials.
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high variation of Peierls valley level also reflects the rugged energy landscape in HEAs [14]. Indeed, the local chemical fluctuations and their interaction with the dislocation can be so strong to give rise to changes of the local dislocation core structure [15]. Since the dislocation energy merely presents local minima in the PEL, important questions remain as to how rough the energy landscape is and to what extent the energy barrier varies in HEAs.

Another salient and emergent feature pertaining to HEAs that makes them distinct from traditional alloys is chemical short-range order (SRO) [16]. The actual materials, which are processed, homogenized or annealed at temperatures below their melting points, unavoidably accommodate SRO, because of the attractive/repulsive interactions among the constituent elements [17]. Especially when the mixing enthalpy becomes the predominant term in the Gibbs free energy, the diffusion kinetics leading to a lower free-energy state renders the formation of SRO inevitable [18,19]. It has been shown that, in fcc alloys, SRO can have a considerable impact on dislocation motion and modifies its structure from wavy to planar [20], impacting the mechanical behaviors, e.g., slip resistance [21] and hardness [18]. In the context of bcc HEAs, the role of SRO on individual mechanisms underpinning dislocation motion and the PEL structure has yet to be elucidated [22].

This study samples and reconstructs the energy landscape formed by concentrated solid solutions and reveals the saddle point activation events along minimum energy pathways that trigger individual mechanisms (Peierls mechanism, kink-pair formation, and lateral kink glide) underlying screw dislocation motion. Considering a model bcc MoNbTaW alloy using a state-of-the-art machine learning interatomic potential [23] that captures solute interactions and ordering, the potential energy landscape demonstrates hierarchical structures with large metastable basins encompassing a bundle of small local basins - a striking feature resulting from random HEA solid solutions. The energy landscape, sensitive to local chemical order, can be altered by introducing SRO (Appendix A), which transfers the predominance of individual mechanisms as the rate-limiting step for screw dislocation motion. The detailed analysis of the PEL in the random solid solution (RSS) alloy shows that, in contrast with both the very recent understanding [24,25] of screw dislocation glide, kink-pair nucleation can be a rate-limiting mechanism that competes with kink pair propagation in concentrated solid solutions. The PEL of the SRO system highlights the various roles of breaking chemical order and consequent diffuse anti-phase boundary (APB) generation in modifying the energy barriers of elementary gliding processes.

2. Phenomenology of screw dislocation motion in RSS and SRO

We first report screw dislocation gliding behaviors in the two different chemical environments, i.e., RSS and SRO. The length of the modeled dislocation is 60 b (b is the Burgers vector magnitude), which is large enough to enable the three elemental mechanisms (Appendix B), Fig. 1a and b shows the typical sequences of a screw dislocation gliding in RSS and SRO, respectively, at T=300 K and 800 MPa applied shear stress. The evolution of dislocation morphologies shows that the dislocation line in RSS is more rugged, and many kinks persist as the motion progresses. Since the screw dislocation line vector and the Burgers vector are parallel, a glide plane is not defined, and the screw dislocation can move in any of the three [110] planes that contain the Burgers vector. In Fig. 1a and b, we use various colors to indicate dislocation slip on different planes.

Examining the detailed dislocation motion (Fig. 1a), we can see two kink-pairs are nucleated within the first 4 ps from the initially straight dislocation, moving part of the dislocation line two Peierls valleys away and on a different plane than the maximum resolved shear stress (MRSS) plane. Lateral kink migration of the two kink-pairs stops at 7.5 ps, suggesting that a large energy barrier has been encountered. The further advancement of the dislocation proceeds with nucleation of a third kink-pair in another region. The new kink pair expands rapidly and one of its kinks recombines with the immobile kink, overcoming the local barrier which was retarding the dislocation motion. At 10.5 ps, the entire dislocation has moved away from its initial Peierls valley after further lateral kink glide and annihilation of opposite-sign kinks. At 11.25 ps, three more kink-pairs nucleate, one of which is on a different (red) plane. Two kinks belonging to different planes collide and generate a cross-kink, leading to screw dislocation self-pinning (12 ps). The dislocation unpinns from cross-kinks requiring a long-waiting time implies a strong self-pinning effect of the cross-kink (Supplementary Fig. S1). In contrast to RSS, dislocation motion in the presence of SRO reveals an overall less rugged structure in the displaced configurations, involving fewer kink-pairs (Fig. 1b). At 5.5 ps, a kink-pair forms on the MRSS plane, and the kinks migrate along the dislocation line (7.75 ps) to eventually recombine, as one would expect to occur in an elemental bcc metal. After the entire dislocation has moved to the next Peierls valley (10 ps), the same processes (kink-pair nucleation and kink propagation) repeat, yet involve kink-pair nucleation on a different plane (18.75 ps). Cross-kink formation is not observed for this 60 b long dislocation after simulating glide for 500 ps.

The collision of kink-pairs on different slip planes results in cross-kinks that act as self-pinning points. Depending on the applied stress and temperature, cross-kinks can be overcome by two distinct mechanisms: (1) stress-driven cross-kink unpinning and (2) thermally-activated and diffusion-mediated cross-kink annihilation. Fig. 2a elucidates the atomistic processes underlying cross-kink formation and breaking at low/moderate temperatures (300 K). The unpinning process resulting from stress-induced cross-kink breaking produces debris that is left behind the moving dislocation and that consists of vacancy and self-interstitial loops [26]. At high temperatures (900 K), the second type of mechanism occurs, as shown in Fig. 2b. The two cross-kinks diffuse
along the dislocation line (the interstitial-type kink drifts faster) and interact with each other. Eventually, the (cross-kink) vacancy and self-interstitial recombine, thus annihilating the cross-kinks. This thermally-activated disentanglement, which is enabled by the high-temperature diffusion, suggests that the self-pinning strengthening might become weakened with increasing temperature, calling for other strengthening mechanisms sustaining the strength at high temperatures [27].

The screw dislocation motion in RSS and SRO environments reveals that distinct mechanisms arise at different length- and time-scales. The spatial and temporal scales of dislocation motion mechanisms, regulated by their energy barriers and the PEL structure, are smaller and more frequent in the RSS case, possibly originating from the faster chemical and energy fluctuations compared with SRO. Next, we focus on the individual processes, i.e., the Peierls mechanism, kink-pair nucleation, and kink glide that mediate dislocation motion and reveal their underlying energy landscapes in SRO and RSS, with an attempt to isolate respective roles of concentrated solid solutions and local chemical order. It is noted that the current study focuses on (i) the energy landscape of three elementary mechanisms, Peierls barrier jumps, kink-formation, and lateral kink glide; (ii) the role of SRO on the energy landscape. There are other essential events, including kink-pair collision, cross-kink diffusion, and cross-kink breaking, for describing overall strengthening effects, and their energy landscape deserves future and careful study.

3. Peierls mechanism and its energy landscape

The Peierls mechanism, involving glide of a straight dislocation segment from one Peierls valley to the neighboring one, occurs via Peierls hill (barrier) crossing on PEL. To reveal its energy landscape and how SRO alters it, we consider a short screw dislocation with 4b length. For such a short-length segment, kink-pair nucleation is prevented, and the entire dislocation can glide over the Peierls. Fig. 3a shows the distance traveled by the screw dislocation as a function of time (glide distance-time curves) in both RSS and SRO cases, subjected to various shear stresses at 300 K. Irrespective of the applied stress, the dislocation glides faster in RSS than SRO. By tracking the dislocation core position upon gliding (Fig. 3b), at the same stress level, the dislocation in RSS experiences more cross slip events than SRO. This appears from the larger vertical displacement component (Y-axis) of the dislocation core, hence a larger slope in the X–Y plane (0.31 for RSS and 0.10 for SRO). These two findings suggest that the presence of SRO retards the Peierls mechanism and reduces the cross-slip frequency.

We now investigate the mechanistic origin of the reduced dislocation mobility by analyzing PEL and the Peierls barrier using minimum energy path calculation (Appendix C). Fig. 4a shows the constructed minimum energy pathway spanning 25 Peierls valleys for both SRO and RSS configurations. The potential energy (per Burgers vector) variation is plotted along with the dislocation move distance over 25a (Peierls valleys spacing a = 0.94b). It can be seen
that the hierarchical structure emerges from the energy landscape with a series of local basins (Peierls valleys) embedding in large metastasins, a feature resulting from concentrated random solutions (upper panel of Fig. 4a). Remarkably, the SRO energy landscape shows an increasing trend as the dislocation advances (middle panel of Fig. 4a). This trend reflects that the chemical order on the slip plane is destroyed when the screw dislocation glides through the SRO crystal. The breaking of SRO, also known as diffuse APB generation, incurs an energy cost, thus resulting in an overall energy increase. By subtracting the APB energy (Supplementary Section 1), we obtain the energy landscape of the Peierls mechanism associated with the chemistry fluctuations (green line in Fig. 4a). While the energy pathway for SRO after subtracting the APB energy shows a similar trend as in the RSS case, the dislocation core energies (local minima) have a more concentrated distribution in SRO than RSS (Supplementary Section 2). The SRO shrinking core energy (reduced variation of Peierls valley) is consistent with the recent study using density functional theory calculations [14].

Fig. 4b shows the energy barrier distributions in the RSS, SRO, and SRO-APB (SRO after removing diffuse APB energy). The SRO shows an overall higher energy barrier than RSS, implying an enhanced lattice resistance to dislocation glide in a chemically ordered environment. After removing the APB energy contribution (bottom panel of Fig. 4b), the barriers exhibit a similar distribution as in RSS: the mean value is similar, but the standard deviation is smaller (being 28.2 meV/b for SRO-APB and 41.8 meV/b for RSS case). This demonstrates the predominant role of breaking local chemical order and the generation of APB in increasing the Peierls barrier. It is worth noting that the average Peierls barrier in RSS and SRO-APB (103 meV/b) is close to the barrier estimated by averaging the elemental contributions (96 meV/b, see Fig. C12), while the diffuse APB energy in SRO raises the value to 127 meV/b.

The simulation results and energy pathway analysis signify that the local chemical order introduced into refractory HEAs retards the Peierls mechanism. The mobility reduction is associated with an increase in Peierls energy barrier connecting two energy valleys. As the dislocation jumps to the next valley, localized slip takes place and interrupts the chemical ordering on the slipped area, incurring an extra energy cost and hence raising the energy barrier. Concerning cross slip, the presence of SRO suppresses the frequency of cross slip and restrains the dislocation glide to one primary slip plane. The less frequent occurrence of cross-slip in the presence of SRO can be related to the smaller solute-dislocation interaction energy fluctuations [24] and to the energy cost of creating diffuse APBs, which might be larger in the cross-slipped case. In the RSS case, which encompasses more significant local chemistry (energy) fluctuations, the chance of finding an energetically more favorable environment on a different slip plane is also enhanced (originating from a wide distribution of dislocation core energy). Therefore, once the dislocation segment encounters on the MRSS plane a strongly unfavorable chemical environment that is hard to glide through, it prefers to change the glide plane towards a more favorable chemical environment. In contrast, the local environments in SRO are more uniform, and it is less likely to find a much lower energy environment on a slip plane different than the MRSS plane.

4. Kink-pair nucleation and its energy landscape

The kink-pair nucleation overcoming a sizable (constant) energy barrier in bcc pure metals is the rate-limiting step for screw dislocation motion. To inspect how the high solute concentration and SRO influence this kink-pair mechanism, we choose a screw dislocation with 20b length, a size that is long enough to prevent Peierls mechanism but appropriate to enable just one single kink-pair nucleation event (Appendix C). Fig. 5a and b illustrates the schematics of the kink-pair nucleation process in RSS and SRO cases, respectively. It is noted that the kink-pair nucleation and propagation in SRO would destroy the chemical order in the kinked region (dislocation swept area). This disruption of local order should be accounted for when considering kink-pair nucleation in SRO environment.

Fig. 4. Peierls mechanism and its energy landscape. (a) Minimum energy pathway and potential energy landscape spanning 25 Peierls valleys in RSS system (purple line). Middle panel shows the energy landscape (orange line) and diffuse APB energy due to dislocation glide-induced order breaking (blue dashed-line). Bottom panel presents the Peierls energy landscape after removing the APB energy. (b) The statistical distributions of energy barrier in RSS, SRO, and SRO with reduced APB energy. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
can be conceived as a reasonable approximation of the RSS energy cost due to kink-pair formation (depicted in Fig. 5a). In the presence of SRO, the diffuse APB contribution should also be accounted for the area swept by the kink pair (Fig. 5b), hence

\[ W_{\text{KP}}^{\text{SRO}} = 2E_{\text{kink}} + E_{\text{int}} + \Delta E_{\text{seg}} + E_{\text{apb}} \]

and the APB contribution (disruption of local order) \( E_{\text{apb}} \) is always positive. Due to the local variation in solutes-dislocation interactions, it is expected that, on a long dislocation line, the nucleation site should be energetically favorable for segment advancement and hence \( \Delta E_{\text{seg}} \) is likely negative, in both RSS and SRO cases. However, in the SRO case, it is apparent from the data shown in Fig. 6 and the analysis above that segment advancement during kink-pair mechanism locally breaks the chemical order and incurs an extra energy cost because of the diffuse APB formation (area swept by the segment advancement). In the bottom panel of Fig. 6c, we present the kink-pair formation energy distribution after subtracting this APB energy from the SRO data. Interestingly, the reduced distribution still shows a slightly higher value but a more compact distribution compared with RSS. Since the average critical kink-kink separation is essentially unchanged between RSS and SRO (~5.9b in both cases), a plausible explanation for the difference is a larger kink-pair formation energy fluctuations in RSS compared with SRO, reflected by a larger standard deviation in the kink-kink separation for RSS (1.1b) compared with SRO (0.8b) (a rough energy landscape encountered by a kink is confirmed in the next Section). Another discrepancy between RSS and SRO-APB can originate from the typical energy changes \( \Delta E_{\text{seg}} \) experienced after segment hopping. In the RSS system, the screw dislocation is more likely to find a more favorable kink-pair nucleation site that is associated with a larger solute-dislocation interaction-energy decrease due to strong chemical fluctuations.

5. Kink glide and its energy landscape

In pure bcc metals, lateral kink migration along the screw dislocation line has a periodicity of b (magnitude of Burgers vector), and the activation energy for lateral glide is negligible so that kinks can glide easily, without thermal activation. The highly concentrated solid solute in HEAs is expected to appreciably change the kink-glide. Here, we construct a screw dislocation containing a single kink and carefully inspect the energy landscape over which the kink must travel (Appendix D). Fig. 8a shows the computed minimum energy pathways of a single kink glide spanning 615b glide distance in both RSS and SRO. Along the pathway, the intermittent kink glide events are interspersed with kink-pair formation events. When the kink is trapped by favorable local environments or faces unfavorable environments ahead of its propagation direction, new kink-pair nucleates somewhere along the dislocation line and the screw dislocation can continue to glide (illustrated in Fig. 7b).

In RSS (Fig. 8a, top panel), the minimum energy pathway and landscape consist of a series of energy minima and saddle points. The landscape is highly rugged and appears hierarchical, featuring small basins and larger meta-basins. Along the path, the two activation events, i.e., kink glide and kink-pair-nucleation, are denoted by filled and unfilled symbols, respectively. The migration of the existing kink will be the active mechanism (Fig. 8b) if its glide barrier is smaller than that of kink-pair nucleation in the dislocation. Once the kink experiences a large energy barrier to glide over (induced by its local environments), the dislocation emits a pair of kinks, and their propagation resumes the movement of the dislocation towards the next Peierls. As one of the newly formed kinks propagates to the pinned kink, eventually, the kinks annihilate with the aid of the elastic kink-kink attraction (configuration evolution and pathway shown in Fig. 8c). Thus, the moving screw dislocation in RSS exhibits both active kink glide and kink-pair nucleation. When

![Fig. 5. (a and b) Schematic illustrations of kink-pair nucleation in RSS and SRO, respectively.](image-url)
the existing kink is trapped and pinned, kink-pair nucleation takes over and produces mobile kinks that resume dislocation motion.

The energy pathway of kink glide in SRO shows an increasing trend (the middle panel of Fig. 8a). As kink glides, the forward glide of the screw dislocation breaks the chemical order on the swept area (Fig. 7a), incurring an extra energy cost. Intriguingly, the fraction of kink-pair events during kink glide is markedly reduced in the presence of SRO, implying enhanced kink glide and hindered kink-pair mechanism. Since SRO increases the kink-pair energy barrier over glide barrier, it would raise the importance of kink-pair nucleation as the rate-limiting process. After removing the energy increase associated with diffuse APB creation (breaking SRO) along the migration path, we obtain the energy landscape of the kink glide along with the screw dislocation. The PEL shows a longer length scale (periodicity) between local energy minima of kinked configuration compared to RSS (14.7b for SRO, 11.6b for RSS, see Supplementary Section 3), and kink nucleation is less frequent in the SRO environment (large formation energy shown in Fig. 6c). We can then conclude SRO suppresses kink nucleation rate and favors kink glide as the mechanism facilitating dislocation motion.

6. Discussion and conclusion

The energy landscape structure that dictates dislocation motion in HEAs, revealed here by means of a minimum energy pathway sampling and a machine learning potential, exhibits a distinct structure from that of pure elements and traditional dilute alloys. Specifically, the trajectories of minimum energy migration pathway (Figs. 4a and 8a) suggest the HEAs possess a hierarchical energy landscape consisting of metabasins breaking into a collection of small sub-basins. The saddle point activations and elementary barrier hopping on the energy landscape, appearing as local pro-
processes such as dislocation segment advancement, kink-pair nucleation, and kink glide, regulate dislocation motion. When the system attempts to escape from one metabasin to an adjacent one, it goes through a series of local barrier activations and inevitably bears accelerated backward jumps during metabasin climbing (forward barrier larger than backward one). This emergent PEL feature, stemming from the multi-principal elements in high concentration, exerts a trapping force and back stress on saddle point activations, retarding dislocation movement. We regard the hierarchy of the energy landscape as the salient character of HEAs that makes them different from pure metals and dilute alloys. Interestingly, the hierarchical and fractal features are ubiquitous in glasses (the disordered materials that lack long-range structural order) [29] and are considered to be the root cause of many anomalous behaviors of glassy material, for instance, rheological behavior. In HEAs, however, the rugged PEL and its multilevel picture are induced by concentrated solid solutions and their chemical disorder. The resultant rugged PEL and its variability with chemical disorderliness act a vital role in dislocation motion and hence unusual strengthening in refractory HEAs.

When SRO is introduced into the system, the PEL of dislocation processes is tilted, because of the chemical order breaking and diffuse APB creation accompanied by individual dislocation advancement mechanisms. In the presence of SRO, the energy barriers for individual mechanisms (Peierls hopping, kink-pair, and kink glide) all increase but to different extents. Because of various SRO sensitivities, kink-pair mechanism is appreciably impeded, yet kink glide is comparatively promoted, leading to the dominant mechanism drift towards kink-pair nucleation. This is signified regarding kink propagation and new kink-pair formation [Fig. 8], where the SRO substantially reduces the occurrence of kink-pair nucleation (large energy barrier and slow process) and turns it into the predominant rate-limiting step for dislocation motion. The pure effects of SRO on dislocation core energy (Peierls valley), kink energy (local energy minimum), Peierls and kink gliding barriers (saddle points on PEL) are isolated by eliminating the energy penalty incurred by breaking local chemical order. The variances of local minimum and energy barrier distributions are shrunk, implying that SRO smoothes the PEL and reduces its ruggedness related to solute-dislocation interaction energy fluctuations.

Since the screw dislocation strength is the sum of the strengthening processes from segments/kinks gliding and cross-kinking [24,30], whether SRO will strengthen the alloy or not depends on specific compositions and their interaction with the dislocation. The presence of SRO brings about the back stress for local chemical order recovering but reduces variance in the distribution of solute-dislocation interaction energies compared to a purely random sample, including cross-kink separation, which controls the cross-kink strengthening [16]. The balance between these two effects gives rise to the net effects of SRO on strength, which highly depends on the alloy composition, as shown for instance in bcc NbTiZr alloys that exhibit SRO-softening [31]. In the MoNbTaW studied here, the SRO-breaking in individual mechanisms and the resulting energy penalty substantially raise the energy barriers and play a major role in dislocation glide. It is reasonable to speculate that SRO will contribute to the strengthening of this type of alloy at low or cryogenic temperatures. As temperature rises to intermediate range and cross-kinks take over as the dominant role in strengthening of refractory HEAs, the SRO can result in less strengthening because of infrequent kink-pair nucleation (enhanced barrier) and suppressed cross-slip (reduced probability of forming cross-kinks). At high temperatures, thermal diffusion mobilizes cross kinks and the subsequent cross-kink annihilation lessens the effectiveness of the strengthening, and therefore, the role of edge dislocations becomes predominant and controls the strength of bcc HEAs [27,32]. The importance of edge dislocation is closely related to its high energy barriers in the rough potential energy landscape, as carefully addressed for bcc high entropy alloys [33]. To assess the local strengthening behavior in solid solutions, an energy topology approach has been applied to study edge dislocation motion in fcc alloys [34,33].

Our study reveals and delineates the striking features of rugged energy landscape inherent in refractory HEAs, over which the dislocation motion necessarily operates and proceeds through cooperative elemental mechanisms along the minimum energy pathways. The unveiled PEL demonstrates a multilevel and hierarchical struc-

**Fig. 8.** Kink glide mechanism and its energy landscape. (a) The potential energy landscape of kink gliding over 615b distance in RSS (upper panel). The unfilled symbols indicate kink-pair events. Middle panel shows the energy landscape of kink gliding in SRO (orange curve) and the associated diffuse APB energy due to order breaking (blue dashed-line). The green curve in the bottom panel is the calculated energy landscape of kink glide in SRO after removing APB energy. (b-c) Typical minimum energy paths and structure evolution of (b) pure kink glide and (c) kink-pair nucleation with chemically pinned kink. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.).
ture, manifesting as a collection of small basins nested in a large metabasin, which is the notable feature making them distinct from their conventional counterparts. By tailoring local chemical distribution and introducing SRO, the energy landscape is appreciably smoothened (reduced variances in local minima and saddle point) yet skewed to different degrees that change the relative role of individual mechanisms as the rate-limiting process for dislocation motion. The energy landscape hierarchy and its high barrier variation, induced by chemical randomness in HEAs, highlight the similarity to that of structural disorder in metallic glasses [29] and render various barrier activation events and dislocation glide mechanisms underpinning the extraordinary behaviors of HEAs.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.actamat.2022.118022.

Appendix A. MC/MD simulations and chemical SRO parameter

We use a Monte Carlo (MC) swap of atoms coupled with MD simulations to prepare the NbMoTaW with lowered potential energy and increased SRO. The MC/MD simulation [36] is performed at 300 K with Nose-Hoover thermostat and involves 1,000,000 MD steps and calls MC trials every 1,000 timesteps to perform trials of exchanging each pair of elements. The atom swaps are accepted or rejected based on the Metropolis algorithm. It is worth noting that the temperature of 300 K is chosen to generate a well ‘annealed’ system with a high degree of SRO, which allows us to study the limiting effects of SRO on screw dislocation motion and the associated energy landscape. At high temperatures, it is possible that the degree of SRO and its contribution to strengthening will be lowered.

To measure chemical SRO, the non-proportional number is used to quantify the degree of chemical order [37]. The order parameter between any pair of atoms $i$ and $j$ is defined as $\delta_{ij}^k = N_{ij}^k - N_{ij}^0$, where $N_{ij}^k$ denotes the actual number of pairs in $k$th shell, and $N_{ij}^0$ the number of pairs for the pure random mixture. Following this definition, we can see a positive $\delta_{ij}^k$ indicates a favored and increased number of pairs, meaning element $i$ tends to bond with element $j$ in the $k$th shell, while a negative value represents an unfavored pairing. In the Fig. A9, we present the MC/MD simulation results, atomic configurations, and the calculated order parameters $\delta_{ij}$ for the first nearest neighboring shell. The order parameter values of all individual pairs in RSS system are zero, indicating the random nature. The MC/MD annealed structure exhibits strong ordering between Mo-Ta ($\delta_{Mo-Ta}^{k=1} = 2.044$), Nb-Mo ($\delta_{Nb-Mo}^{k=1} = 1.468$), and W-Ta ($\delta_{W-Ta}^{k=1} = 1.416$) pairs.

Fig. A9. MC-MD simulations and chemical SRO parameters. (a) Potential energy change as a function of MC in the hybrid MC-MD simulation. (b) The variation of SRO parameter $\delta_{Mo-Ta}^{k=1}$ with MC swaps. (c) The calculated SRO parameters at the first nearest neighbor in random solid solution (RSS) and short-range order (SRO) systems. The MC/MD annealed structure exhibits strong ordering between Mo-Ta, Nb-Mo, and W-Ta. (d) The corresponding atomic configurations of RSS and SRO, respectively.
Appendix B. MD simulations of screw dislocation motion in RSS and SRO

To reveal screw dislocation motion behaviors in RSS and SRO, we perform MD simulations of a periodic array of dislocations subjected to constant shear stress. The crystal has dimensions of ~250 Å in [112], 240 Å in y [110] and 168 Å in z [111] directions, containing 580,000 atoms. As shown in Fig. B10, a screw dislocation with a length of 60b is introduced to the system along z by imposing a linear displacement on all atoms in the upper half of the cell. The system orientations give rise to the glide direction is along x and the glide plane normal direction y. Periodic boundary conditions are imposed along x and z, and zero tractions on the two free surfaces parallel to the glide plane. To drive the motion of screw dislocation, we apply the constant shear stress to the system by adding a constant force to two atomic layers of upper and lower surfaces. A machine learning potential [23] is adopted to model interatomic interactions in NbMoTaW, which has been verified for predicting the non-degenerate core structure of screw dislocation.

Appendix C. Peierls mechanism and kink-pair nucleation models

We use the free-end nudged elastic band (NEB) method [38] to construct the minimum energy pathway of dislocation glide over two neighboring Peierls valleys. By moving the dislocation to next valley and repeating the same procedure, the energy landscape of dislocation glide over a large distance is obtained. For Peierls mechanism, a short dislocation of 4b length is considered and inserted in the system (see Fig. C11). We minimize and connect the energy pathways spanning 25 Peierls valleys using NEB calculation. The NEB inter-replica spring constant is set to be k = 0.01 eV/Å² and force tolerance 0.001 eV/Å. The choice of this value that optimizes convergence of the calculations results in essentially the same energy barrier using smaller tolerance and large spring constant (Supplementary Fig. S2). For pure elements, the Peierls barrier is constant in a perfectly periodic energy landscape. Fig. C12 shows the Peierls energy barriers for the four constituent elements, Mo, Nb, Ta, and W, in which the average energy barrier is estimated to be 96 meV/b.

In order to explore kink pair nucleation energy barriers, a longer dislocation with 20b is adopted, which naturally leads to one kink-pair event along the dislocation line. The kink-pair nucleation mechanism and the minimum energy pathways are calculated using NEB by considering 30 different sites in RSS and SRO, respectively.

Appendix D. Kink glide model

To construct dislocation with one single kink, we first build a crystal with orientations of [9 20], [110], and [1 1 0 0 0 1] along x,
y, and z directions. A screw dislocation is inserted into the system with a line direction of z. The angle between [10 10 9] and [1 1 1] directions is 2.79°, and the distance between two Peierls valley is 0.94b (b is Burgers vector). The dislocation length for forming one single kink is 54.25 Å in the z direction (see Fig. D13). After energy minimization, one single kink is formed in the relaxed and kinked dislocation line locating in the Peierls valleys (i.e., along with the [1 1 1] direction). We then perform NEB calculation and construct the minimum energy pathway of kink-gliding connecting two neighboring energy minima. By sampling a large glide distance of 650b, the potential energy landscape governing kink propagation is obtained in RSS. As to SRO, the initial crystal is annealed through MC/MD simulation, followed by the same procedure, i.e., [10 10 9] orientated dislocation generation and minimum energy path calculation.

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