

1 **Ultrastrong and ductile NiTi-based composite with large recoverable
2 strain mediated by a compositionally complex phase**

3 Jiayi Geng^{1,4}, Yunzhu Shi^{1,*}, Pere Barriobero-Vila^{2,3}, Meiyuan Jiao⁵, Yihuan Cao⁵, Yu
4 Tang⁶, Jingzhi He⁶, Chao Ma¹, Yan Ma⁷, Zhifeng Lei^{1,*}, Zhaoping Lu^{5,*}

5 ¹College of Materials Science and Engineering, Hunan University, Changsha, 410082,
6 China

7 ²Department of Materials Science and Engineering, Technical University of Catalonia
8 (UPC), Barcelona 08019, Spain

9 ³CIM, Technical University of Catalonia (UPC), Barcelona 08028, Spain

10 ⁴State Key Laboratory of High-Performance Ceramics and Superfine Microstructure,
11 Shanghai Institute of Ceramics, Chinese Academy of Sciences, Shanghai, 200050,
12 China

13 ⁵Beijing Advanced Innovation Center for Materials Genome Engineering, State Key
14 Laboratory for Advanced Metals and Materials, University of Science and Technology
15 Beijing, Beijing 100083, China

16 ⁶College of Aerospace Science and Engineering, National University of Defense
17 Technology, Changsha 410073, China

18 ⁷Max-Planck-Institut für Eisenforschung, Max-Planck-Straße 1, 40237 Düsseldorf,
19 Germany

20 *Corresponding authors. Email: yzshi@hnu.edu.cn (Yunzhu Shi); zflei@hnu.edu.cn
21 (Zhifeng Lei); luzp@ustb.edu.cn (Zhaoping Lu).

22

23 **Abstract**

24 NiTi-based composites possess great potential for concurrently improving both
25 mechanical and functional properties. However, relying on traditional alloy design
26 principles limits the design space and greatly hinders the advancement of high-
27 performance NiTi-based composites. The concept of high-entropy alloys has expanded
28 the compositional landscape, unveiling unique structural characteristics for alloy design
29 and providing new prospects for addressing these limitations. Here, we report a
30 compositionally complex NiTi-based composite that exhibits exceptional strength and
31 ductility, along with remarkable recoverable strain. The composite,
32 $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ (at.%), comprises a 78.0% B2 NiTi matrix, a 19.2% Nb-Mo-Ta-
33 W-Ti-Ni compositionally complex body-centered cubic (BCC) phase, and a small
34 amount of Ti_2Ni . Notably, this composite achieves a compressive strength of 3274 MPa,
35 a fracture strain of 44.2%, and a recoverable strain of 7.3%. These outstanding
36 mechanical properties arose from the interplay between the compositionally complex
37 phase and lattice strain matching facilitated by phase transformations. The substantial
38 recoverable strain was obtained through the reversible $\text{B}2 \rightleftharpoons \text{R} \rightleftharpoons \text{B}19'$ phase transition.
39 This work not only innovates a new category of high-performance NiTi-based
40 composites but also extends the applicability of the high-entropy concept.

41 **Keywords:** NiTi-based composites; High-entropy alloys; Mechanical properties;
42 Recoverable strain; Phase transformation.

43

44 **Introduction**

45 Shape memory alloys fall within the category of smart materials, owing to their
46 remarkable capability to undergo substantial reversible phase transformations that
47 define their functional properties^[1–3]. NiTi alloys with a simple CsCl (B2) structure lead
48 in shape memory alloys due to their outstanding attributes: exceptional shape memory
49 effect, pseudoelasticity, strong corrosion resistance, biocompatibility, and good
50 processability^[4–6]. To cope with the increasingly challenging and intricate service
51 environment, expanding industrial application of NiTi alloys primarily involves
52 improving their mechanical properties and functional capabilities. To meet this demand,
53 NiTi-based composites, such as NiTi-metal and NiTi-intermetallic compound
54 composites, have become widely adopted solutions. Examples of these composites
55 include NiTi-Nb^[1], NiTi-W^[7], NiTi-V^[8], NiTi-NbTi^[9], NiTi-Ti₃Sn^[5], NiTi-Ti₅Si₃^[10]
56 and NiTi-Nb₃Sn^[11]. Of particular note is the NiTi-Nb composite, which has a
57 substantial quasi-linear elastic strain of over 6% and an impressive yield strength of
58 1.65 GPa^[1]. However, conventional design principles restrict the composition range of
59 NiTi-based metallic composites, hindering their further development and performance
60 optimization. Moreover, traditional NiTi-based composites are usually soft due to
61 occurrence of martensitic transformation. Intermetallic compounds, characterized by
62 their rigidity and brittleness, hinder efforts to improve ductility and recoverable strain.
63 Therefore, developing innovative NiTi-based composites is imperative to tackle these
64 challenges.

65 The alloy design concept of high-entropy alloys (HEAs) composed of multiple
66 principal components is a promising solution to address these challenges. The strong
67 configuration entropy effect inherent in HEAs allows for a considerably wide range of
68 compositions in alloy design^[12–14]. This compositional diversity endows HEAs with
69 great potential to integrate various performance attributes^[15]. Moreover, the variations
70 in atomic size and chemistry among the multiple constituents lead to large lattice
71 distortion. This phenomenon serves to impede dislocation movement, thereby

72 substantially enhancing the strength of HEAs^[16–20]. In addition, the complex short-
73 range ordering within these solid solutions enriches their plastic deformation
74 mechanisms, thereby improving their ductility^[21–26]. Therefore, utilizing the concept of
75 HEAs reveals captivating opportunities for advancing NiTi-based composites.

76 Stimulated by the field of HEAs, here, we developed a NiTi-based composite,
77 achieving remarkable strength and ductility, coupled with a significant recoverable
78 strain. The Ni₄₀Ti₄₀(NbMoTaW)₂₀ (at.%) composite exhibits a compressive strength of
79 3274 MPa, a fracture strain of 44.2%, and a recoverable strain of 7.3%. This composite
80 comprises an Nb-Mo-Ta-W-Ti-Ni compositionally complex body-centered cubic (BCC)
81 phase (19.2% area fraction) embedded in the B2 NiTi matrix (78.0% area fraction),
82 along with a dispersion of a small amount of Ti₂Ni phase. The compositionally complex
83 phase and the phase transformation-assisted lattice strain matching contribute to the
84 exceptional mechanical properties. The large recoverable strain originates from the
85 reversible B2 \rightleftharpoons R \rightleftharpoons B19' phase transformation. The current findings not only introduce a
86 new class of NiTi-based composites for engineering applications, but also offer insights
87 into the design of innovative multifunctional metallic materials using the HEA concept.

88

89 **Results**90 **Mechanical and superelastic properties**

91 **Figure 1a** illustrates the compressive engineering and true stress-strain curves of
92 the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, revealing a prominent double-yielding
93 phenomenon. Stress-induced martensitic transformation of the B2 NiTi phase within
94 the composite contributes to the first yielding^[27]. The first yield stress σ_{FY} and strain
95 ε_{FY} are 469 ± 24 MPa and $2.87\% \pm 0.16\%$, respectively, whilst the second yield stress
96 σ_{SY} and strain ε_{SY} are 1089 ± 30 MPa and $11.34 \pm 0.55\%$, respectively. The ultimate
97 strength σ_{US} and fracture strain ε_{FS} are significantly increased to 3274 ± 56 MPa and
98 $44.2\% \pm 0.2\%$, respectively. **Supplementary Figure 1** presents the fracture morphology
99 of the composite, showing pronounced ductile tearing and a large number of dimples
100 in $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$, indicative of its superior ductility. **Figure 1b** shows a plot of
101 the ultimate strength against fracture strain for the current $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$, and
102 previously reported NiTi-based alloys and composites. For comparison, compressive
103 mechanical properties of the $\text{Ni}_{40}\text{Ti}_{40}\text{Nb}_{20}$ composite were also examined
104 (**Supplementary Figure 2**). The $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite exhibits high
105 strength and large ductility. In addition, the composite features a low Young's modulus
106 of 17.9 ± 0.9 GPa, enabling significant elastic strain under low stress conditions and
107 potentially enhancing its superelastic performance.

108 **Figure 1c** depicts the compressive cyclic stress-strain curve of
109 $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$. Notably, during cyclic loading, the ultimate strength and
110 fracture strain of the composite are consistent with those under continuous loading
111 (**Figure 1a**). By analyzing the cyclic stress-strain curve, we calculated the superelastic
112 strain ε_{SE} and recoverable strain ε_{RS} of the composite (**Supplementary Figure 3**).
113 Initially, the superelastic strain ε_{SE} , induced by the reversed martensite to austenite
114 phase transformation, increases during the first three cycles ($\varepsilon < 9\%$) and stabilizes at
115 approximately 1.5% (see the lower part of **Supplementary Figure 3b**). In contrast, the

recoverable strain ε_{RS} of the composite increases with cyclic loading. The recoverable strain climbs from 2.1% to 7.3% under cyclic loading (see the upper part of **S3b**). Thus, $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ exhibits not only superb mechanical properties but also attractive superelastic characteristics. **Figure 1d** compares the superelastic properties of $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ with the $\text{Ni}_{40}\text{Ti}_{40}\text{Nb}_{20}$ composite (**Supplementary Figure 4**) and other NiTi-based alloys/composites^[28,29]. As shown, $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ shows a maximum recoverable strain comparable to NiTi-Nb and NiTi-Ag composites, underscoring its superior superelastic features.

Microstructure characterization

To unveil the intrinsic mechanisms governing the mechanical and superelastic properties of this material, we meticulously examined its phase constitutions. **Figure 2a** displays the high-energy X-ray diffraction (XRD) pattern of $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$. The composite primarily consists of B2 and BCC phases, accompanied by a limited quantity of Ti_2Ni and R phases. **Figure 2b** captures a scanning electron microscope (SEM) micrograph of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, unveiling a microstructure characterized by large white particles dispersed throughout the matrix. After careful examination, the matrix and white strips form a mesh-like eutectic structure, with a small number of black strips located at the junctions of the mesh-like white strips (**Figure 2c**). The corresponding energy-dispersive X-ray spectroscopy (EDS) results reveal that Ni and Ti elements are enriched in the matrix, Nb, Mo, Ta, and W elements are enriched in the large white particles and white strips, and Ni and Ti elements are enriched in the black strips (**Supplementary Figure 5**). The detailed compositions for each phase are provided in **Table S1**. In the matrix, the atomic ratio of Ni and Ti elements is close to 1:1, while in the black strips, the ratio tends to be 1:2. Combining the high-energy XRD and SEM results, it was found that the matrix corresponds to the B2 NiTi phase, the black strips is the Ti_2Ni phase, and the white particles and strips are the BCC phase. Specifically, for the BCC phase in $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$, the concentration of Ni, Ti, Nb, Mo, Ta and W were measured to be 10.05 ± 2.50 at.%,

144 19.95 ± 3.84 at.%, 13.96 ± 1.77 at.%, 13.51 ± 0.93 at.%, 18.23 ± 1.36 at.% and 24.29 ± 6.82
145 at.%, respectively. The BCC phase exhibits an ideal configurational mixing entropy of
146 approximately $14.56 \text{ J} \cdot \text{mol}^{-1} \cdot \text{K}^{-1}$, well within the range established for HEAs by
147 Yeh et al^[30]. Moreover, the area fractions of the B2, BCC, and Ti_2Ni phases were
148 determined to be $78.0\% \pm 4.1\%$, $19.2\% \pm 5.3\%$, and $2.8\% \pm 1.6\%$, respectively. Therefore,
149 we can classify the current BCC phase as a compositionally complex phase.

150 To delve into the atomic-scale microstructure features of $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$,
151 we utilized transmission electron microscopy (TEM). **Figure 2d** shows a representative
152 bright-field TEM image. The selected area electron diffraction (SAED) pattern (**Figure**
153 **s. 2d1-d3**) clearly identifies the matrix as B2, white strips as BCC, and black strips as
154 Ti_2Ni . The B2 and BCC structures were further affirmed through a high-resolution TEM
155 image and the corresponding fast Fourier transformation (FFT) image (**Supplementary**
156 **Figure 6**). Both distinctly illustrate the ordered atomic arrangement of B2 versus the
157 disordered atomic arrangement of BCC. The corresponding TEM-EDS mapping
158 analysis further demonstrates the enrichment of Nb, Mo, Ta, and W elements in the
159 BCC phase, while Ni and Ti elements dominate the composition of the B2 matrix and
160 Ti_2Ni phase (**Figure 2d**). Remarkably, the Ni enrichment in the Ti_2Ni phase is lower
161 than that in the B2 matrix, whereas the Ti enrichment follows an opposite trend. The
162 Ti_2Ni strips exhibit a tendency to connect the BCC phase, forming a continuous mesh-
163 like structure. A similar structure, arising from the partial substitution of Ti atoms by
164 Nb atoms in the B2 phase, has also been observed in NiTi-Nb composites. This
165 phenomenon leads to the accumulation of expelled Ti atoms at grain boundaries, thus
166 resulting in the formation of the Ti_2Ni phase^[31]. It is worth noting that other refractory
167 elements may have similar effects.

168 Transformation behavior

169 To comprehend the phase transformation mechanisms in the present
170 $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, we conducted differential scanning calorimetry

171 (DSC) measurement (**Figure 3a**). In the heating and cooling cycle, the material
172 manifests two exothermic peaks and two endothermic peaks. Microstructure analysis
173 reveals the predominant presence of the B2 NiTi and BCC phases within the composite.
174 Specifically, when doped with additional elements, the B2 NiTi phase typically
175 experiences a two-step phase transition $B2 \rightleftharpoons R \rightleftharpoons B19'$. This dual transformation is
176 characterized by two exothermic peaks and two endothermic peaks^[32]. Thus, it can be
177 deduced that the $Ni_{40}Ti_{40}(NbMoTaW)_{20}$ composite undergoes a two-step $B2 \rightleftharpoons R \rightleftharpoons B19'$
178 phase transformation during both the cooling and heating cycles. In this case, both the
179 R and B19' phases undergo diffusionless transformations characterized by minimal
180 displacements. The R phase takes the form of a rhombic structure resulting from the
181 elongation of B2 along the [111] direction, while B19' adopts a monoclinic structure
182 formed by the displacement of a pair of orthogonal (110) crystal faces in B2 by a certain
183 distance^[6,33,34]. The specific reversible transformation process can be described as
184 follows: $(\bar{1}\bar{1}1)_{B2} \rightleftharpoons (100)_R$; $(\bar{1}21)_{B2} \rightleftharpoons (010)_R$; $(2\bar{1}1)_{B2} \rightleftharpoons (001)_R$; $(111)_R \rightleftharpoons (100)_{B19'}$;
185 $(\bar{2}11)_R \rightleftharpoons (010)_{B19'}$; and $(0\bar{1}1)_R \rightleftharpoons (001)_{B19'}$ (**Figure 3b**). The occurrence of this two-step
186 transition within the B2 phase in NiTi-based composites is associated with factors such
187 as the introduction of a third element and the ratio of Ni to Ti elements^[33,35]. This
188 implies that the inclusion of refractory elements may facilitate the transition from B2
189 to the R phase. The $B2 \rightarrow R$ transformation start temperature (R_s) and the $R \rightarrow B19'$
190 martensitic transformation start temperature (M_s) are measured at $35.4^\circ C$ and $-85.7^\circ C$,
191 respectively. The R_s value exceeds room temperature, suggesting the likely presence of
192 the R phase in the $Ni_{40}Ti_{40}(NbMoTaW)_{20}$ composite. This aligns with the high-energy
193 XRD result (**Figure 2a**). Also, the low M_s implies a significant difference in chemical
194 potential between B19' and B2 in $Ni_{40}Ti_{40}(NbMoTaW)_{20}$, potentially yielding a
195 substantial driving force for the reverse transition from B19' to B2. Moreover, the
196 $B19' \rightarrow R$ transition finish temperature (R'_f) and the $R \rightarrow B2$ austenitic transformation
197 finish temperature (A_f) are measured at $-46.5^\circ C$ and $71.7^\circ C$, respectively. The low R'_f
198 suggests a thermodynamically favored transition from the B19' phase to the R phase.

199 **Deformation mechanisms**

To unravel the deformation mechanism exhibited by $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite under both continuous and cyclic loading, we conducted *in-situ* synchrotron XRD experiments. **Figure 4a** depicts the *in-situ* synchrotron XRD measurement during continuous loading, revealing a four-stage deformation process. In stage I (macro-strain $\epsilon=0\text{-}1.8\%$), we observe the presence of only the BCC and B2 phases. Notably, their peak positions exhibit subtle leftward shifts, while their peak widths remain largely unaltered. Consequently, this initial stage is primarily characterized by the elastic deformation of the BCC and B2 phases. Moving on to stage II ($\epsilon=1.8\text{-}2.6\%$), we observe that the peak positions of the BCC phase continue to shift slightly leftward, indicating that the BCC phase remains in the elastic deformation period. Concurrently, the transition from the B2 phase to the R phase commences, and with further deformation, the emergence of the B19' phase follows suit, while the retained B2 phase continues its elastic deformation. As we transition into stage III ($\epsilon=2.6\text{-}8.6\%$), the B2 phase undergoes a significant transformation into the R phase and subsequently into the B19' phase. Thus, during this stage, the deformation of the composite is primarily driven by the $\text{B}2 \rightarrow \text{R} \rightarrow \text{B}19'$ phase transformation. Stage IV ($\epsilon>8.6\%$) witnesses the composite mainly comprising the BCC and B19' phases. Though the peak positions of both phases display minimal changes, the peak widths progressively increase with deformation, signifying that this stage is primarily characterized by the plastic deformation of both phases. **Figure 4b** illustrates the lattice strain of the BCC and B2 phases during continuous loading. In the elastic strain regime (stage I and II, $\epsilon=0\text{-}2.6\%$), the BCC phase demonstrates a maximum lattice strain of 0.45%, while the B2 phase undergoes elastic deformation at macro-strain ϵ of less than 2.6%. Remarkably, the BCC phase exhibits a substantial elastic strain ($\epsilon=2.7\%$) during loading, closely resembling the onset strain for the B19' transition (stage II, $\epsilon=1.8\text{-}2.6\%$) (**Figure 4c**). This phenomenon facilitates a favorable alignment between the BCC and B2 phases. In the course of the B19' phase transition (stage III, $\epsilon=2.6\text{-}8.6\%$), the d -spacing strain of the BCC phase increases (**Figure 4b**). The slope during this stage is higher than that in the plastic regime (stage IV, $\epsilon>8.6\%$), indicating the presence of lattice strain and

dislocation slip in the BCC phase. Notably, this alignment between the BCC phase and the B19' phase is achieved primarily through lattice strain rather than dislocation slip, a principle well-documented in studies involving nanowire and shape memory alloy composites, known as the lattice strain matching approach^[1,36]. The lattice strain matching approach considerably diminishes the intensity of the elastic strain field at the phase boundary. As we enter stage IV ($\epsilon > 8.6\%$), the d -spacing strain of the BCC phase gradually increases, with the corresponding slope being lower than that in stage III ($\epsilon = 2.6\%-8.6\%$). This suggests that the BCC phase now predominantly undergoes dislocation slip. The B19' phase transition primarily occurs in stage III ($\epsilon = 2.6\%-8.6\%$) (**Figure 4c**). The fraction of the B19' phase exhibits a rapid increase with macro-strain, with approximately 75% of the B2 phase ultimately transforming to the B19' phase by the end of this stage. In the early period of stage IV ($\epsilon = 8.6\%-16.7\%$), the fraction of the B19' phase remains relatively constant, possibly owing to the orientation of the B2 phase, which does not favor martensitic phase transformation. However, in the later period ($\epsilon > 16.7\%$), the residual B2 phase initiates transformation into the B19' phase, suggesting that increased stress and strain trigger the reinitiation of martensitic phase transformation.

Figure 4d presents the *in-situ* synchrotron XRD measurement under cyclic loading, revealing a distinct reversible $B2 \rightleftharpoons R \rightleftharpoons B19'$ transition. In the initial loading-unloading cycle (O-A-B, $\epsilon = 0-3\%$), the BCC phase peak position shifts leftward during loading (O-A) and then rightward during unloading (A-B). Initially, the peak width remains constant until significant broadening close to point A, indicating elastic deformation in the BCC phase, followed by plastic deformation beyond its elastic limit, generating dislocations and peak broadening. The B2 phase transforms to the R phase during loading (O-A) and reverses to the B2 phase during unloading (A-B). The second cycle (B-C-D, $\epsilon = 1.7\%-6.7\%$) sees the BCC phase peak position shift leftward during loading (B-C) and rightward during unloading (C-D). Peak width remains initially constant, gradually broadening during loading (indicating elastic and plastic deformation) and remaining constant during unloading. The $B2 \rightarrow R \rightarrow B19'$ phase

transition occurs during loading (B-C), with the reverse transition during unloading (C-D). In the third cycle (D-E-F, $\varepsilon=4.2\%-18\%$), the BCC phase undergoes elastic and plastic deformation during loading (D-E) and primarily elastic deformation during unloading (E-F). Similar to the previous cycle, the $B2 \rightarrow R \rightarrow B19'$ phase transition occurs during loading (D-E), with the reversed transition during unloading (E-F). After unloading, the residual $B19'$ phase remains (point F), primarily due to plastic deformation of the BCC phase hindering the $B19'$ phase's reverse transformation.

Figure 4e depicts d -spacing strain versus macro-strain for the BCC and B2 phases during cyclic loading. Residual stress persists in both phases post-unloading. A negative d -spacing strain reflects tensile stress, whereas a positive value indicates compressive stress. Specifically, after unloading at different stages, we find that the BCC phase exhibits d -spacing strains of -0.037% (at point B), -0.131% (at point D), and -0.199% (at point F), respectively. In contrast, the B2 phase displays d -spacing strains of 0.035% (at point B), 0.059% (at point D), and 0.093% (at point F), respectively. These observations imply that tensile stress persists in the BCC phase after unloading, while the B2 phase maintains compressive stress. With an increasing number of cycles, residual stress becomes more pronounced. In the BCC phase, tensile stress partially counteracts the applied compressive stress during subsequent loading, enhancing the elastic strain limit, reaching 3.2% during the third unloading. Furthermore, for reversible phase transformations, the $B2 \rightleftharpoons R$ transformation primarily occurs at low strain (< 6.7%), while the $B2 \rightleftharpoons B19'$ transformation mainly takes place at high strain (> 6.7%) (**Figure 4f**). Plastic deformation induced by the BCC phase at elevated strain levels impedes the $B19' \rightarrow R \rightarrow B2$ phase transition during unloading, resulting in a substantial 60% residual presence of $B19'$ post-unloading.

282 Discussion

We present a schematic representation of the deformation mechanisms in $Ni_{40}Ti_{40}(NbMoTaW)_{20}$ during continuous and cyclic loading processes (**Figure 4g**). During the loading process, in stage I ($\varepsilon=0\%-1.8\%$), the material undergoes the $B2 \rightarrow R$

transformation within the B2 phase, concurrently experiencing elastic strain in both the untransformed B2 and BCC phases. In stage II ($\varepsilon=1.8\%-2.6\%$), R \rightarrow B19' and B2 \rightarrow B19' transitions take place. The untransformed B2 and BCC phases continue their elastic deformation. Progressing to stage III ($\varepsilon=2.6\%-8.6\%$), the B19' phase fraction increases with strain, with the B19' phase initially undergoing elastic deformation, followed by plastic deformation. The BCC phase and the residual B2 phase undergo plastic deformation during this stage. In stage IV ($\varepsilon=8.6\%-16.7\%$), plastic deformation occurs in both the B19' and BCC phases, culminating in the transformation of the residual B2 phase into the B19' phase. During unloading, the B19' \rightarrow B2 transformation takes place. After unloading, the BCC phase experiences microscopic tensile stress, while the B2 phase incurs microscopic compressive stress. Therefore, in Ni₄₀Ti₄₀(NbMoTaW)₂₀ composite, the B2 NiTi matrix seamlessly integrates with the BCC phase, ensuring exceptional mechanical performance and recoverable strain capacity.

We harness the HEA concept to craft an inventive Ni₄₀Ti₄₀(NbMoTaW)₂₀ composite, uniting exceptional strength, ductility, and recoverable strain. Refractory HEAs inherently feature elevated intrinsic strength and pronounced lattice distortion^[19,37,38], both pivotal in achieving the high strength observed in the Ni₄₀Ti₄₀(NbMoTaW)₂₀ composite. Additionally, within our Ni₄₀Ti₄₀(NbMoTaW)₂₀ composite, the BCC phase exhibits a high ideal configurational entropy (**Supplementary Table 1**). The high configurational entropy fosters the formation of a stable BCC solid solution phase^[39], eliminating competing intermetallic compounds within the composite matrix. Thus, in Ni₄₀Ti₄₀(NbMoTaW)₂₀, the fraction of the brittle Ti₂Ni phase is minimized. This reduction significantly enhances the ductility of the Ni₄₀Ti₄₀(NbMoTaW)₂₀ composite (**Figure 1a**). Furthermore, during loading, the BCC phase exhibits a substantial elastic strain of 2.7% (**Figure 4b**), attributed to the effect of lattice strain matching between the transformation lattice distortion of the B2 NiTi matrix and the elastic strain of the BCC phase^[40]. These unique micromechanical characteristics further underpin the high strength and large ductility of Ni₄₀Ti₄₀(NbMoTaW)₂₀. Moreover, multiple-element doping induces a multi-step phase

315 transition process in $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, prompting the occurrence of the
316 $\text{B}2 \rightarrow \text{R} \rightarrow \text{B}19'$ phase transition during continuous loading and the $\text{B}2 \rightleftharpoons \text{R} \rightleftharpoons \text{B}19'$ phase
317 transition during cyclic loading. Consequently, the combination of lattice strain
318 matching and the reversible $\text{B}2 \rightleftharpoons \text{R} \rightleftharpoons \text{B}19'$ phase transition plays a pivotal role in
319 imparting $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ with its remarkable ductility and substantial
320 recoverable strain. Importantly, the extensive compositional flexibility of HEAs,
321 coupled with the manifold advantages stemming from the aforementioned structural
322 features, markedly expands the design landscape for NiTi-based composites. This
323 boundless potential facilitates the development of innovative NiTi-based composites
324 that seamlessly integrate a range of exceptional properties into a unified whole.

325 In summary, we pioneer the utilization of the HEA concept in the fabrication of
326 NiTi-based composites. The developed composite is primarily composed of a B2 NiTi
327 matrix and a BCC compositionally complex phase. Importantly, the composite exhibits
328 remarkable attributes, featuring exceptional strength, large ductility, and a superb
329 recoverable strain capacity. Notably, it showcases a compressive strength of 3274 MPa,
330 a fracture strain of 44.2%, and a recoverable strain of 7.3%, surpassing the overall
331 performance of the majority of NiTi-based composites. The extraordinary mechanical
332 performance is attributed to the unique structural characteristics of the compositionally
333 complex phase and the lattice strain matching induced by phase transitions.
334 Furthermore, multi-element microalloying within the B2 phase introduces a reversible
335 $\text{B}2 \rightleftharpoons \text{R} \rightleftharpoons \text{B}19'$ phase transition process, resulting in a significant recoverable strain. This
336 work not only introduces a novel category of NiTi-based composites but also offers
337 valuable insights into strategies for enhancing their performance through the HEA
338 concept.

339 **Methods**

340 **Material preparation.** Alloy ingots with a nominal composition of
341 Ni₄₀Ti₄₀(NbMoTaW)₂₀ (at.%) were prepared by arc melting a mixture of pure metals
342 (purity, >99.9 weight%) in a Ti-gettered high-purity argon atmosphere. The ingots were
343 remelted at least eight times to ensure chemical homogeneity, and then drop-cast into a
344 water-cooled copper mold with a dimension of 10×10×60 mm³.

345 **Mechanical measurements.** Compression and cyclic compression tests were
346 conducted at room temperature using an electric universal material testing machine
347 (AGS-X-50KN) with a loading/unloading rate of 1×10⁻³ s⁻¹. The sample size is Φ4 mm
348 × 6 mm. At least five samples were tested. In the loading-unloading tests, cyclic
349 compression strains increased in 3% increments, starting at 3%, and continued until
350 reaching failure.

351 **Microstructural characterization.** Microstructure and morphology were
352 characterized by MIRA3 LMU scanning electron microscope (SEM, TESCAN
353 company) operating at 20 kV. The energy-dispersive X-ray spectroscopy (EDS) was
354 utilized to analyze the elemental distribution. The area fraction of each phase was
355 calculated and measured by ImageJ software. The SEM sample was initially polished
356 with 2000-grit SiC paper. Then, the sample was mechanically polished with a diamond
357 plaster in a metallographic polishing machine (LMD-2C). Subsequently, the sample
358 was polished with a colloidal silica suspension in a vibratory polishing machine (Vibra
359 304). Transmission electron microscopy (TEM, FEI Talos F200X) operating at 200 kV.
360 The TEM sample was mechanically ground to a thickness of 50 µm and then twin-jet
361 electropolished using 10 % sulfuric acid + 90 % methanol (vol.%) solution under a
362 temperature of -20 °C, a voltage of 15 V. Phase transformation behavior of the
363 composite was studied through differential scanning calorimetry (DSC) using a TA
364 DSC25 unit at a cooling/heating rate of 10 K/min in a nitrogen atmosphere.

365 **In-situ synchrotron high-energy X-ray diffraction.** *In-situ* high-energy X-ray

366 diffraction (XRD) during compression was carried out at the beamline P07-HEMS of
367 PETRA III, Deutsches Elektronen-Synchrotron (DESY)^[41]. Uniaxial compression was
368 performed at room temperature (RT) using a modified dilatometer Bähr 805A/D
369 equipped with a deformation unit^[42]. Sequences of entire Debye-Scherrer rings were
370 recorded from the bulk material in transmission mode using a Perkin Elmer XRD 1621
371 image-plate detector. The incident X-ray beam (perpendicular to the loading direction)
372 was positioned at the center of the sample's length. A slit size of 0.8×0.8 mm² resulting
373 in a gage volume of $0.8 \times 0.8 \times 4$ mm³ was used. The sample-detector distance and
374 acquisition time were 1600 mm and 1 s, respectively. An illustrative diagram of the
375 experimental setup is presented in ref.[43]. The beam energy was 100 keV (wavelength
376 = 0.124 Å). The instrument parameters were calibrated using a LaB6 powder standard.
377 One-dimensional (1-D) high-energy XRD patterns were obtained by integrating across
378 all angles in the 2-D diffraction patterns under various applied strains. The lattice strain
379 under various applied strains was fitted by distorting two-dimensional (2-D) diffraction
380 patterns using GSAS-II software package^[44]. Positive for compression and negative for
381 tension in the direction of loading.

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394 **Author contributions**

395 Y.Z.S., Z.F.L., and Z.P.L. supervised and conceived this project. J.Y.G. fabricated the
396 materials and characterized the mechanical properties, phase transition behavior, and
397 microstructure of the materials. P.B.V. and Y.M. conducted the in-situ high-energy X-
398 ray diffraction experiments. Y.H.C. conducted the DSC experiments. C.M. and M.J.Y.
399 conducted and analyzed TEM experiments. Y.T. and J.Z.H. conducted the cyclic
400 compression experiments. J.Y.G., Y.Z.S., Z.F.L., and Z.P.L. wrote the manuscript. All
401 authors discussed the results and commented on the manuscript.

402 **Competing interests**

403 The authors declare no competing interests.

404 **Additional information**

405 **Supplementary information** The online version contains supplementary material
406 available at xxx.

407 **Correspondence** and requests for materials should be addressed to Yunzhu Shi,
408 Zhifeng Lei or Zhaoping Lu.

409 **Data availability**

410 The datasets generated and/or analyzed during the current study are available from the
411 corresponding authors on request.

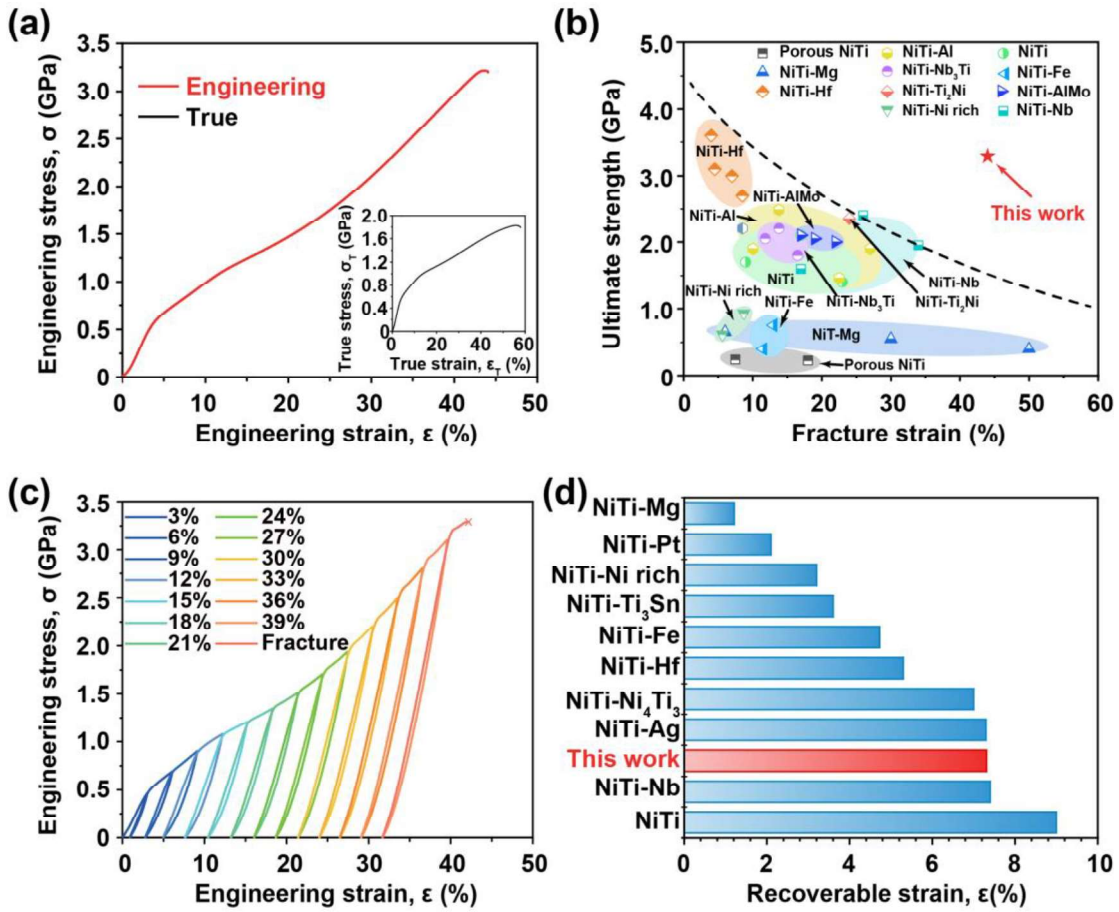
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- 483



484
485 **Figure 1. Mechanical and superelastic properties.** (a) Compressive engineering
486 stress-strain curve of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, with the inset displaying
487 the corresponding true stress-strain curve. (b) Ultimate strength versus fracture strain
488 of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite in comparison with NiTi and other NiTi-based
489 composites^[11,45–62]. (c) Engineering stress-strain curve during cyclic compression for
490 the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite. During the loading-unloading test, the cyclic
491 compression strains range from 3%, 6%, 9%, 12%, and so forth until failure occurs. (d)
492 Recoverable strain of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite in comparison with NiTi
493 and other NiTi-based composites^[1,29,45,47,52,53,56,59–68].

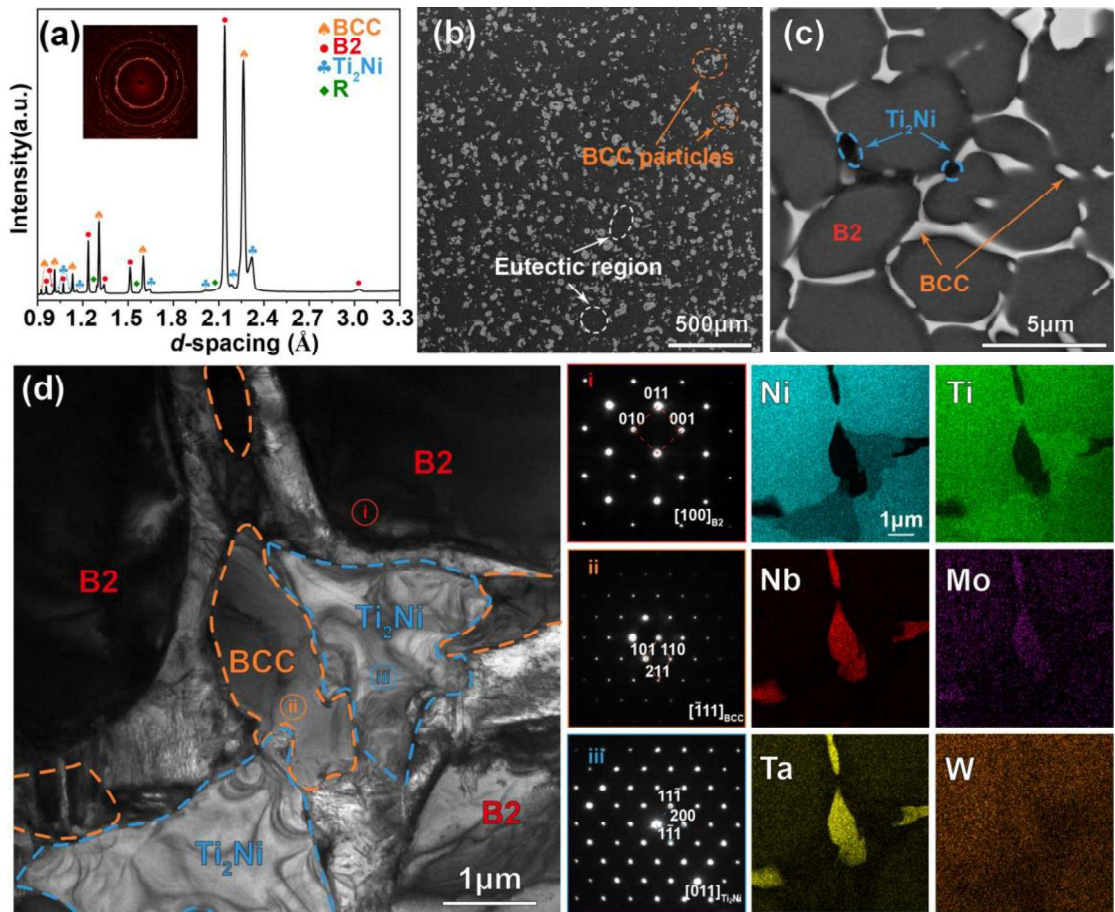
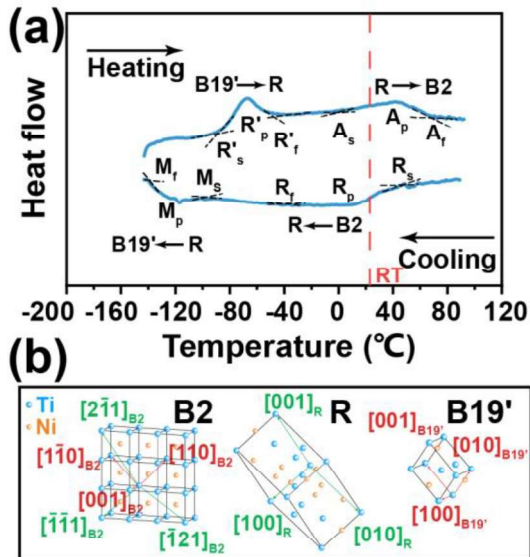


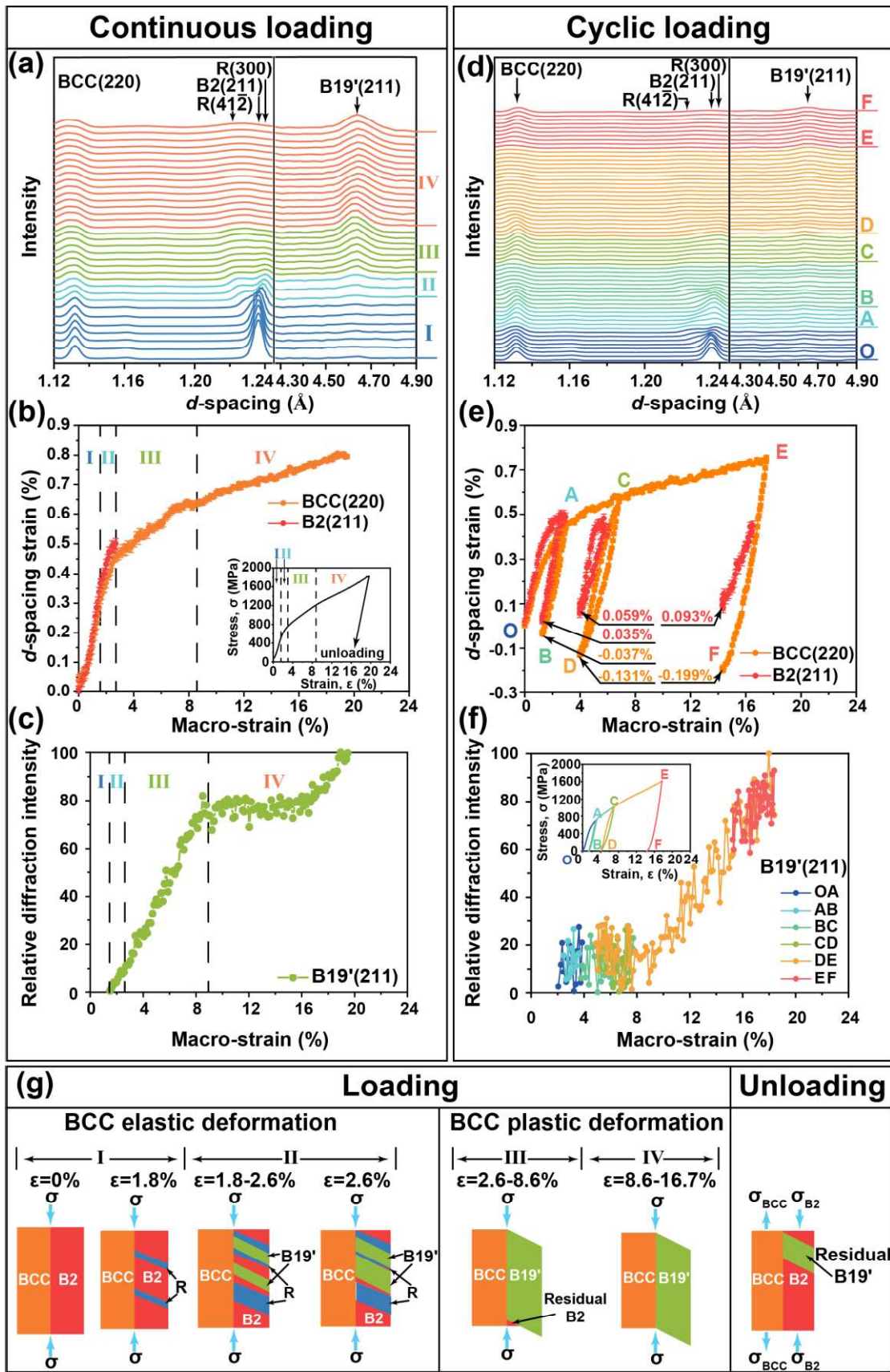
Figure 2. Microstructure characterization. (a) Synchrotron high-energy X-ray diffraction (XRD) pattern of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, with the inset revealing the Debye–Scherrer rings. (b) Scanning electron microscope (SEM) image of the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite. (c) Corresponding local enlargement of (b). The white arrows highlight the eutectic region, the orange arrows depict the body-centered cubic (BCC) particles and strips, and the blue arrows illustrate the Ti_2Ni strips. (d) Bright-field transmission electron microscopy (TEM) image offering a detailed view of the B2 NiTi matrix, BCC phases, and the Ti_2Ni precipitates. (i), (ii) and (iii) are the corresponding selected area electron diffraction pattern of the B2 NiTi, BCC and Ti_2Ni phases. The two columns on the right showcase the elemental distribution mapping, illustrating the spatial distribution of Ni, Ti, Nb, Mo, Ta, and W.



507

508 **Figure 3. Transformation behavior.** (a) Differential scanning calorimetry (DSC)
 509 curve for the $\text{Ni}_{40}\text{Ti}_{40}(\text{NbMoTaW})_{20}$ composite, providing insights into various
 510 transformation temperatures. Key points on the DSC curve are labeled as follows: RT
 511 (room temperature), R_s (start temperature of $\text{B}2 \rightarrow \text{R}$ transition), R_p (peak temperature
 512 of $\text{B}2 \rightarrow \text{R}$ transition), R_f (finish temperature of $\text{B}2 \rightarrow \text{R}$ transition), M_s (start temperature
 513 of $\text{R} \rightarrow \text{B}19'$ martensitic transformation), M_p (peak temperature of $\text{R} \rightarrow \text{B}19'$ martensitic
 514 transformation), M_f (finish temperature of $\text{R} \rightarrow \text{B}19'$ martensitic transformation), R'_s
 515 (start temperature of $\text{B}19' \rightarrow \text{R}$ transition), R'_p (peak temperature of $\text{B}19' \rightarrow \text{R}$ transition),
 516 R'_f (finish temperature of $\text{B}19' \rightarrow \text{R}$ transition), A_s (start temperature of $\text{R} \rightarrow \text{B}2$ austenitic
 517 transformation), A_p (peak temperature of $\text{R} \rightarrow \text{B}2$ austenitic transformation), and A_f
 518 (finish temperature of $\text{R} \rightarrow \text{B}2$ austenitic transformation). (b) Schematic diagram
 519 illustrating the interconversion between the $\text{B}2$, R , and $\text{B}19'$ phases.

520



521

522 **Figure 4. Deformation mechanisms.** (a) *In-situ* synchrotron high-energy X-ray

523 diffraction patterns captured during continuous loading, revealing the dynamic
524 evolution of peaks corresponding to BCC (220), B2 (211), R (41 $\bar{2}$), R (300), and B19'
525 (001). **(b)** The progression of d -spacing strain for BCC (220) and B2 (211) planes
526 during continuous loading, with analysis data obtained parallel to the loading direction.
527 An inset provides a schematic illustration of the continuous loading process. **(c)** The
528 evolution of the relative diffraction intensity of B19' (001) peaks during continuous
529 loading. **(d)** *In-situ* synchrotron high-energy X-ray diffraction patterns recorded during
530 cyclic loading. **(e)** The evolution of d -spacing strain for BCC (220) and B2 (211) planes
531 during cyclic loading, with analysis data acquired parallel to the loading direction. **(f)**
532 The evolution of the relative diffraction intensity of B19' (001) peaks during cyclic
533 loading. An inset offers a schematic depiction of the cyclic loading process. **(g)** A
534 schematic representation elucidating the elastic-plastic deformation within the BCC
535 phase and the phase transition in the B2 matrix. Here, σ represents the external stress;
536 σ_{BCC} symbolizes the microscopic stress on the BCC phase after unloading, and σ_{B2}
537 denotes the microscopic stress on the B2 matrix after unloading.