Carbon Nanotubes Enabled Laser 3D Printing of High-Performance Titanium with Highly Concentrated Reinforcement

HIGHLIGHTS

- Take advantage of reaction between CNTs and Ti for performance breakthrough
- Use laser 3DP as a unique route to tailor in situ formation of novel nanostructure
- Increase strength and ductility simultaneously via multiple strengthening mechanisms
- Broaden structural applications of CNTs in load-bearing Ti-based engineering materials
Carbon Nanotubes Enabled Laser 3D Printing of High-Performance Titanium with Highly Concentrated Reinforcement

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SUMMARY
Zero- to two-dimensional nanomaterials have been incorporated into metal-matrices to improve the strength of metals, but challengingly, high-volume-fraction nanomaterials are difficult to disperse uniformly in metal matrices, severely degrading the ductility of conventionally processed metals. Here, a considerably dense uniform dispersion of in situ formed nanoscale lamellar TiC reinforcement (16.1 wt %) in Ti matrix is achieved through laser-tailored 3D printing and complete reaction of Ti powder with a small amount (1.0 wt %) of carbon nanotubes (CNTs). An enhanced tensile strength of 912 MPa and an outstanding fracture elongation of 16% are simultaneously achieved for laser-printed components, showing a maximum 350% improvement in “product of strength and elongation” compared with conventional Ti. In situ nanoscale TiC reinforcement favors the formation of ultrafine equiaxed Ti grains and metallurgically coherent interface with minimal lattice misfit between TiC lamellae and Ti matrix. Our approach hopefully provides a feasible way to broaden structural applications of CNTs in load-bearing Ti-based engineering components via laser-tailored reorganization with Ti.

INTRODUCTION
Modern aerospace, aircraft, and automobile industries have an urgent demand for the increased payload and fuel efficiency, promoting the development of better metallic materials with higher strength/weight ratios (Pollock, 2010). Since the strength-ductility trade-off has been a long-standing dilemma for materials (Wei et al., 2014), increasing the strength of metals without sacrificing ductility is critical for their competitiveness (Lu, 2010). An essential solution of this challenge relies on a radical change in the way that metals are reinforced, which in turn calls for revolutionary production technologies. However, conventional synthesis and processing methods have reached certain limits in further elevating the properties of metals including titanium (Ti) (Chen et al., 2015). Titanium may be reinforced by other materials in a controlled way to form titanium matrix composites (TMCs), which provides a possible route to enhance overall properties (Attar et al., 2018; Dadbakhsh et al., 2019; Tian et al., 2019). Performance enhancement of novel materials also urgently calls for even advanced processing technologies. Laser-based three-dimensional printing (3DP)/additive manufacturing (AM) of metallic materials is currently attracting worldwide research interest, owing to the immense application demands in recent years from high-end industries (Bourell et al., 2017; Debroy et al., 2019; Macdonald and Wicker, 2016). Laser-based 3DP has demonstrated a significant capability for creating high-performance/multi-functional components with complex geometries in a cost-effective manner (Martin et al., 2017; Shukrun et al., 2018; Zhakeyev et al., 2017). More importantly, its highly non-equilibrium physical metallurgical nature, featured by superhigh heating/cooling rates (10³–10⁸ K s⁻¹) (Das et al., 2010), makes this technology more unique and valuable for the development of non-equilibrium nano-phases and fine-grained microstructures and attendant enhancement in final mechanical performance (Wang et al., 2018).

Ceramic nanoparticles including TiC (Gu et al., 2012b), TiB₂ (Attar et al., 2015), and BaC (Li et al., 2019), as a category of zero-dimensional nanomaterial, have been added into Ti matrix to produce Ti-based nano-composites by laser 3DP. Laser printing of TMCs based on external addition of ceramic reinforcing...
particles is essentially a powder metallurgy route, with the main cores of ceramic particles remaining unmelted. Nevertheless, it is rather difficult to incorporate nanoparticles into Ti since they tend to agglomerate into clusters within the matrix, especially for a high-weight-fraction ceramic addition, thus initiating cracks in composites rather than reinforcing them (Lin et al., 2019). A large difference in surface tensions of ceramics and Ti also causes poor wettability and weak interfacial coherence of reinforcing phase by molten metal (Ma et al., 2017).

New carbonaceous nanomaterials, e.g., one-dimensional nanomaterial carbon nanotubes (CNTs) and two-dimensional nanomaterial graphene, owing to their extraordinary high specific strength (Bakshi et al., 2010), have also been introduced to strengthen Ti matrix, using both laser processes (Hu et al., 2019) and powder metallurgy routes (Li et al., 2013). Presently, the worldwide production capacity of CNTs exceeds several thousand tons per year (De Volder et al., 2013). For commercialization of CNTs products that find applications in load-bearing engineering structures, it is essential to integrate CNTs with metals to form nanocomposites (Kinloch et al., 2018). Besides the general processing difficulties of nanomaterials such as phase agglomeration and limited wettability, another long-term challenge arises from the possible chemical reaction between CNTs/graphene and Ti matrix during processing (Landry et al., 1998). A general principle of processing CNTs/graphene reinforced TMCs is to make CNTs/graphene “survive” during processing, thereby maintaining their initial nanostructural characteristics. However, owing to the extremely high atomic activity of CNTs/graphene, the in situ chemical reaction between C and Ti is normally inevitable during high-temperature processing. A close study of the microstructures of TMCs in previous work (Hu et al., 2019; Li et al., 2013) reveals a typical composite reinforcing structure, including the unreacted surviving CNTs/graphene and a fraction of in situ formed TiC during processing, hence increasing the microstructural heterogeneity.

In the present study, we propose a completely reverse processing philosophy to process CNTs/Ti TMCs, fully taking advantage of selective laser melting (SLM) 3DP to tailor an in situ reaction between CNTs and Ti to form novel nano-reinforcement. CNTs refer to a helical tubular structure grown with a single wall or multi-wall and typically have a diameter less than 100 nm (Bakshi et al., 2010). The high chemical activity of CNTs enables a complete in situ atomic-scale reaction between CNTs and Ti matrix during SLM 3DP. Thanks to the initial favorable nanostructure of CNTs, the reaction product is hopefully tailored in a nanoscale. More important, the unique physical metallurgical nature of laser 3DP, e.g., sufficiently high working temperature, superhigh temperature gradient, and superfast solidification rate, favors a homogeneous high-density precipitation of in situ nanophase, which is generally impossible for conventional powder metallurgy process for TMCs with limited thermodynamics. We demonstrate that a unique SLM process plus a class of new carbonaceous nanomaterial leads to a thorough reorganization of materials and attendant development of novel TMCs, yielding the significantly elevated mechanical performance of as-printed materials. We achieve a great balance between strength and ductility of TMCs via laser-tailored formation of nanoscale in situ reinforcement and regulation of interfacial strengthening mechanisms.

RESULTS AND DISCUSSION

Laser-Tailored Formation Mechanism of In Situ Nanoscale Reinforcement

Since CNTs powder demonstrates a high tendency toward agglomeration (Tjong, 2013), it is a long-standing challenge to homogenize CNTs dispersion while maintaining a high flowability of a nanocomposite powder, which plays a crucial role in achieving a smooth powder layer deposition and a sound SLM processability. We develop an approach for the scaled preparation of highly flowable and homogeneous Ti-based nanocomposite powder. Spherical CP-Ti powder is coated with CNTs powder using a “low-energy ball milling” method featured by a relatively low ball-to-powder ratio and a reduced rotational speed of grinding bowls. The original favorable spherical structure of Ti matrix powder is well maintained after ball milling (Figure 1A) to ensure a sufficiently high flowability of as-milled nanocomposite powder. CNTs are uniformly coated and distributed on the surface of Ti particles, with CNTs coating thickness being several tens of nanometers (Figures 1B–1D). Mass-produced CNTs/Ti nanocomposite powder was additively manufactured via the SLM process. We optimize an “island scanning” strategy by dividing the cross-section into islands to reduce the vector length (Figures 1E and 1F), which can greatly reduce residual stresses in SLM-processed parts. We also rotate the scan pattern by 37° in each island between neighboring layers; the directional anisotropy of current layer is compensated for by the following layers, thereby creating a more homogeneous stress and microstructure distribution (Thijs et al., 2013). SLM of our carefully prepared CNTs/Ti nanocomposite powder demonstrates a smooth laser-powder interaction...
Figure 1G) and a high capability to produce complex-structured integral turbine part with a dimension larger than 200 mm in a single print (Figure 1H).

Scanning electron microscope (SEM) and transmission electron microscope (TEM) characterizations of as-built Ti-based samples using a laser power of 350 W demonstrate that SLM of CNTs/Ti powder yields a completely new composite structure, showing a considerably dense uniform dispersion of the significantly refined lamellar reinforcing phase throughout Ti matrix (Figure 2B). The selected area electron diffraction (SAED) pattern of a lamella can be indexed as TiC crystal along the [001] direction on the basis of a cubic unit cell (Figure 2C), implying the occurrence of in situ reaction between CNTs and Ti during SLM. We accordingly propose a reverse processing philosophy on CNTs-reinforced Ti-based nanocomposites through laser-tailored reorganization of materials, fully taking advantage of conventionally “unfavorable” in situ reaction between CNTs and Ti matrix. Interestingly, our energy-dispersive spectroscopy (EDS) line-scan analysis across a lamella reveals a gradient distribution of C element (Figure 2D), indicating the formation of substoichiometric titanium carbide referred to as TiC$_x$ (x = 0.1–0.9) (Hugosson et al., 2001). The substoichiometric TiC$_x$ carbides are stable over a large range of C contents. Our TEM analysis reveals that the TiC$_x$ lamellae generally have a standard nanostructure, with the average thickness below 80 nm (Figure 2C). We attribute the formation of nanoscale TiC$_x$ to the combined effect of its material nature and unique thermodynamic behavior of SLM process. The standard Gibbs free energy ($\Delta G$) of TiC formation by a reaction between titanium and carbon above the melting temperature of titanium is $-136.178$ kJ/mol (Barin, 1995), indicating a spontaneous generation of TiC$_x$ during SLM. SLM is featured by a superfast melting/solidification metallurgical process, and the operative temperature within the molten pool exceeds 3,000 K within 1.1 ms, yielding an extremely high peak cooling rate of $2.73 \times 10^7$ K s$^{-1}$ (Figure S1A). The in situ precipitated TiC$_x$ nuclei have insufficient time for crystal growth, well retaining the nanostructure. Meanwhile, the calculated temperature gradient within the molten pool is as high as $4.89 \times 10^7$ K m$^{-1}$ (Figure S1B), giving rise to surface tension gradient and resultant Marangoni flow (Leung et al., 2018). The formation of convective stream within the molten pool induces liquid capillary force, which in turn exerts on the precipitated TiC$_x$ and accelerates its rearrangement throughout the melt, favoring a homogeneous dispersion in the solidified matrix (Figure 2B).
We further study the grain morphology and size distribution of Ti matrix by electron backscatter diffraction (EBSD). The inverse-pole figure (IPF) map shows an almost general formation of fine equiaxed Ti grains at a laser power of 350 W (Figure 2A), yielding a relatively large fraction of low-angle grain boundaries (LAGBs, 2°–15°) of ~17% (Figure S2A). The estimated average grain size of Ti matrix, 3.24 μm, is considerably low (Figure S2B). A successive change of color in the IPF map reveals that the crystallographic orientation of Ti grains is highly random (Figure 2A). Here we demonstrate that the in situ formation of nanoscale TiCx reinforcement exerts a significant effect on microstructural development of Ti matrix during a non-equilibrium SLM process, featured by the prevailing of refined grain size, enhanced fraction of LAGBs, and week texture intensity.

**Effect of Printing Parameters on Microstructural and Interfacial Characteristics**

We conduct intensive experiments on process optimization and find that the microstructural development of reinforcement is very sensitive to the applied printing parameters. Although SLM at a relatively low laser power of 150 W also leads to the in situ formation of TiC reinforcing phase, as confirmed by SAED pattern (Figure 2G), its microstructure experiences a substantial change into a dendritic structure with a dendrite trunk length larger than 6 μm (Figure 2F), completely losing the novel nanoscale lamellar structure formed at a laser power of 350 W (Figure 2B). The C and Ti elemental distribution from the dendritic arm of reinforcement exerts a significant effect on microstructural development of Ti matrix during a non-equilibrium SLM process, featured by the prevailing of refined grain size, enhanced fraction of LAGBs, and week texture intensity.

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**Figure 2. Microstructural Characterizations of Laser-printed Ti-based Samples Using Different Laser Powers**

Laser powers of (A–D) 350 W and (E–H) 150 W are used, while laser scan speed of 1,300 mm s⁻¹ is fixed.

(A and E) Cross-sectional EBSD IPF map of Ti-based samples showing grain morphologies and orientations.

(B) SEM images showing a dense uniform dispersion of the significantly refined reinforcing phase formed at a laser power of 350 W. The inset acquired at a higher magnification highlights the lamellar nanostructure of the reinforcement.

(F) SEM image showing the presence of dendritic reinforcing phase at a laser power of 150 W.

(C and G) Bright-field TEM images and SAED patterns (on white dotted circles) of the lamellar reinforcement formed at 350 W and the dendritic arm formed at 150 W.

(D and H) EDS line scan across reinforcement and matrix (along green lines A₁B₁ and A₂B₂) showing concentration distributions of C and Ti elements.
carbon during non-equilibrium laser solidification determine the morphologies of TiC. SLM at a laser power of 150 W produces a relatively low thermal gradient of $2.01 \times 10^7$ K m$^{-1}$ (Figure S1B) and attendant weak activity of solute atoms, building up a solute boundary layer at the solid-liquid interface owing to a high carbon solute concentration. The constitutional supercooling thus occurs, causing a morphological instability of the solid-liquid interface (Kim, 1978). The amplitude of the perturbation increases with the TiCx crystal growth proceeds. The finally developed TiC phase accordingly demonstrates a dendritic growth mechanism.

Of particular interest is that laser power has an essential influence on growth morphologies of TiC reinforcing phase. A reasonable increase in laser power favors the morphological transformation of TiC from a coarsened dendritic structure to a refined nanoscale lamellar structure (Figure 2B versus 2F); nevertheless, the TiC becomes re-coarsened into a rod-like shape at an excessive laser power of 450 W (Figure S4C), caused by an elevated thermalization of laser energy and attendant enhanced growth kinetics of TiC nuclei. We also disclose that, as a sufficiently high laser power of 350 W is optimized, the lamellar structured TiC reinforcement is generally produced at various laser scan speeds. Nevertheless, we still discern a minor morphological difference. A proper increase in laser scan speed refines the TiC lamellae and homogenizes its distribution in the matrix (Figure S4D versus S4B); however, an excessively high scan speed of 1,700 mm s$^{-1}$ produces the re-coarsened and agglomerated TiC lamellae (Figure S4F). The elevated Marangoni convection and liquid capillary instability in the molten pool formed at a higher laser scan speed intensify the microstructural heterogeneity of as-printed Ti-based nanocomposites.

We demonstrate that the development manner of in situ TiC significantly affects its atomic bonding ability with Ti matrix. The atomic map of the interface between dendritic TiC and Ti matrix, consisting of (0002) planes of Ti and (200) planes of TiC, illustrates an evidently sharp interface (Figures 3A and 3B) and attendant disordered and unfavorable dendrite/matrix atomic bonding (Figure 3C). We measure the spacing between the adjacent lattice planes and the lattice misfit of two phases was calculated to be 9.2% in this instance. By contrast, the interface between lamellar TiC and Ti matrix is considerably coherent (Figures 3D and 3E), indicating a continuous and metallurgically compatible atomic bonding (Figure 3F). By measuring the interplanar spacing of (101) planes of Ti matrix and (200) planes of TiC lamella, we find that the lattice misfit between the nanoscale lamellar TiC and Ti matrix decreases significantly to 1.9%.

**Nanoscale Reinforcement Induced Microstructural Development of Ti Matrix**

We further performed X-ray diffraction (XRD) to investigate the constitutional phases and examine the grain refinement and lattice mismatch in SLM-processed Ti-based nanocomposites. The strong diffraction peaks corresponding to TiC and hexagonal close-packed (hcp) Ti are generally identified in the specimens printed at different laser powers. Here we determine that starting from 1.0 wt % CNTs, the content of in situ TiC$_0.3$ in 350-W-printed Ti-based specimen increases dramatically to 16.1 wt %. A close look at the diffraction peak for TiC (111) (350 W, 1,300 mm s$^{-1}$), which locates at 2$\theta$ of 36.451°, reveals that the lattice parameter of TiC is $a = 0.427$ nm. This observation confirms that the in situ formed nanoscale substoichiometric titanium carbide is TiC$_{0.3}$ (Harada and Ohkoshi, 1997). Because of the considerably short laser melting time, the carbon atoms do not have sufficient time to diffuse into the TiC$_0$ lattice to form stoichiometric TiC (Al-Mangour et al., 2017). A large number of interstitial carbon sites therefore remain vacant. This can explain why the incorporation of a low-weight fraction of CNTs to Ti can achieve a highly concentrated precipitation of TiC$_0$ nanoscale reinforcing phase within matrix. The XRD characterization within a small 2$\theta$ range (inset in Figure 4A) illustrates that, as relative to the standard diffraction peaks for hcp-Ti ($\alpha$-phase) located at 2$\theta$ of 35.1°, the exact 2$\theta$ locations of the diffraction peaks for hcp-Ti in laser-printed nanocomposites generally shift to lower 2$\theta$ angles, indicating an increase in interplanar distance based on the Bragg’s law (Gu et al., 2012a). Normally, the occurrence of martensitic transformation is accompanied by microscopic volume expansion, thereby enhancing the distortion of lattice and resultant lattice constant. Our quantitative calculation reveals the lattice parameters $a = 0.29529$ nm and $c = 0.46912$ nm for 350-W-printed specimen, which are apparently larger than the standard lattice parameters $a_0 = 0.2944$ nm and $c_0 = 0.4678$ nm of $\alpha$-Ti (Rostoker, 1952). These hcp-Ti diffraction peaks accordingly correspond to martensitic $\alpha'$-phase, induced by the direct transformation of $\beta$-Ti to $\alpha'$-Ti during laser rapid melting/solidification process (Collins et al., 2016). It is worth noting that a more significant shift of $\alpha'$-Ti peak to a lower 2$\theta$ angle occurs in laser-printed Ti-based nanocomposites using a higher laser power of 350 W, which implies a lager degree of lattice distortion caused by the incorporation of densely dispersed nanoscale lamellar TiC$_0$ in the Ti matrix.
Lattice strain is a measure of the distribution of lattice constants arising from crystal imperfections such as lattice distortion and dislocations (Yogamalar et al., 2009). To quantitatively analyze the lattice strain, we display the modified Williamson-Hall plots of $\sin \theta$ versus $b_{hkl} \cos \theta$ in Figure 4B, where $b_{hkl} \cos \theta$ is the width of XRD diffraction peaks and $\theta$ is the Bragg angle. From the linear fit to the data, we extract the lattice strain and grain size of $\alpha$-Ti from the slope and Y-intercept of the fit line, respectively, where the lattice strain is assumed to be uniform in all crystallographic orientations. We disclose that the application of a higher laser power gives rise to a larger lattice strain of the Ti matrix, as the slope of the red fit line (0.52107 for 350-W printed specimen) is larger than that of the blue fit line (0.18895 for 150-W printed specimen). Meanwhile, a smaller value of Y intercept of red line implies that the grain refinement effect is more significant in the specimen printed at 350 W, which is consistent with the EBSD data (Figure S2A versus S3A). A high-density distribution of in situ formed TiC lamellae ahead of the solidification front in the molten pool provides numerous low-energy-barrier heterogeneous nucleation sites for the Ti matrix, favoring the equiaxed growth of crystals and attendant ultrafine microstructure. We observe the significantly refined, lath-shaped martensitic microstructure of $\alpha$-Ti matrix (Figure 4C) and the high-density dislocations within the martensitic laths (Figures 4D and 4E). Owing to the incorporation of densely distributed nanoscale in situ TiC reinforcement, the $\alpha$-Ti matrix accordingly demonstrates a combined effect of large lattice distortion, significant grain refinement, and high-density dislocations.

**Tensile Properties and Strengthening Mechanisms**

We perform tensile tests and compare the results with specimens produced from CP-Ti by conventional processing methods (Figure 5A). The laser-printed in situ Ti-based nanocomposites with a small amount (0.5–1.0 wt %) CNTs addition have high tensile strengths, which are almost two times stronger than those of conventional coarsen-grained CP-Ti parts. Of particular significance is that the laser-printed in situ Ti-based nanocomposites maintain sufficiently high ductility, demonstrating a significantly elevated balance.
between strength and ductility. Therefore, tailoring the in situ formation of nanoscale TiC reinforcement throughout the Ti matrix dramatically improves the overall mechanical properties without sacrificing the ductility of the nanocomposites. We provide in Figure 5B five representative tensile engineering stress-strain curves of laser-processed Ti-based nanocomposites and CP-Ti extracted from Figure 5A. With an addition of 0.5 wt % CNTs, a simultaneous improvement in tensile strength of 799 MPa and fracture elongation of 21% is achieved for laser-printed specimen at 350 W, compared with the laser-printed CP-Ti prepared at the same processing parameter (tensile strength of 735 MPa and elongation 19.6%). The Young’s moduli deduced from tensile stress-strain curves of laser-printed Ti-based nanocomposites with 0.5 wt % CNTs vary from 96 to 107 GPa. With the additive CNTs increased to 1.0 wt %, laser-printed nanocomposites yield a better strength of 912 MPa by a slight sacrificing of elongation to 16%. Since more nanoscale TiCx reinforcement form within laser-printed Ti-based nanocomposites, the Young’s modulus increases to a higher value within 104–118 GPa. As relative to coarse dendritic TiC with an intrinsic brittle nature formed at a low laser power (Figure 3A), the homogeneously dispersed, ultrafine nanoscale TiC reinforcement obtained via a high-laser-power printing can effectively transfer load owing to their favorable interfacial atomic bonding with α-Ti matrix (Figure 3D–3F). Here we apply a key performance indicator “product of strength and elongation” (PSE) to indicate the balance between strength and ductility. The PSE of the optimally printed Ti-based nanocomposites reaches 16.8 GPa-%, showing a maximum 350% improvement in PSE as relative to conventionally processed unreinforced CP-Ti.

To understand the deformation behaviors and strengthening mechanisms of laser-printed in situ Ti-based nanocomposites, we perform microstructural characterizations on tensile samples at different strain levels. Dislocation slipping plays a dominant role in the deformation at a relatively low tensile strain of 3%. The nanoscale lamellar TiC reinforcing phase tends to hinder the movement of dislocations and activates the Orowan effect,
resulting in the trapping and pileup of dislocations around TiC lamellae with a further deformation (Figure 6A). It is known that the strengthening mechanism through grain refinement predominates for SLM-processed CP-Ti without any reinforcement and the grain refinement normally leads to reduction of twining propensity (Attar et al., 2014). Differently, a small amount of deformation twins, which are generally present near the zones with a considerably high dislocation density, are also observed in the present Ti-based nanocomposites in this strain level (Figures 6B and 6C). As the strain level increases to 8%, twinning becomes a predominant mechanism responsible for the further deformation. More importantly, these twins show a nano-sized structural feature (nano-twins). The nano-twins restrict the transmission of dislocations from one side of the twin boundary (TB) to the other (Zhu et al., 2007). Therefore, compared with the low strain level, the pileup of dislocations becomes more significant near the TiC lamellae, further producing the stress concentration zones under the action of a tensile load (Figure 6E). Numerous sets of nano-twins nucleate and develop around the lamellar TiC (Figures 6D and 6F) when the stress concentration reaches the twin critical value (Zhu et al., 2007). Once the deformation dominated by dislocation slipping is hard to proceed, the subsequently present nano-twins lead to the formation of twinning-induced lattice reorientation, where TBs can act as slip planes on which dislocations move and pile up. The dislocation slip along TBs releases effectively the stress concentration at the dislocation-TB intersection region, thereby accumulating a large amount of plastic strain for a high density of TBs (Lu et al., 2009). This slip-within-twin mechanism favors an additional plastic deformation, significantly improving the deformability and achievable ductility of Ti-based nanocomposites.

Conclusions
The SLM 3DP process, owing to its unique material additive manufacturing philosophy and highly non-equilibrium physical and chemical metallurgical nature, provides a one-step approach to create complex structured, new nanocomposite components with unique reinforcement architectures and elevated mechanical properties. With the aid of SLM 3DP, an addition of a small amount of CNTs (1.0 wt %) to CP-Ti enables a dramatic in situ formation of Ti-based nanocomposites having a dense uniform dispersion of 16.1 wt % nanoscale lamellar TiC reinforcement. An enhanced tensile strength of up to 912 MPa and a high fracture elongation of 16% are simultaneously achieved for laser-printed in situ Ti-based nanocomposites, showing a maximum 350% improvement in PSE as relative to conventional CP-Ti. Strengthening of Ti-based nanocomposites is based on coherent interface and minimal lattice misfit between nanoscale TiC lamellae and Ti matrix through a synergistic function of multiple strengthening mechanisms including dislocation slipping and deformation twinning. A great balance between strength and ductility is achieved via laser-tailored formation of nanoscale reinforcement and interfacial regulation. Our approach hopefully broadens the structural applications of CNTs, whose worldwide production capacity has exceeded several
thousand tons per year, in load-bearing Ti-based engineering components via laser-tailored reorganization with Ti materials that are served as cornerstone of modern aircraft, aerospace, and energy industries.

Limitations of the Study

For reaction-based 3DP method for formation of TiC-reinforced Ti-based nanocomposites, the influence of the initial carbon source on laser-tailored formation of TiC reinforcement needs to be further considered. Besides the CNTs being studied in this work, the nanoscale carbon sources can be broadened to graphite nanoparticles and even graphene. The effects of the type, structure, and size of the carbon matter on the laser printability and resultant microstructural and mechanical properties of laser-printed Ti-based nanocomposites deserve to be further studied. Second, besides the matrix metal titanium studied in this work, other metals having carbide-forming ability can also be mixed with carbon sources for laser printing to yield the carbide-reinforced metal matrix nanocomposites. Third, as a unique heat source, laser plays an important role in tailoring in situ reaction and forming metal matrix nanocomposites. In addition to the fiber laser being applied in this study, more new lasers can be applied for more potential breakthroughs in microstructure control and performance enhancement.

Resource Availability

Lead Contact

Further information and requests for materials should be directed to and will be fulfilled by the lead contact, Dongdong Gu (dongdonggu@nuaa.edu.cn).
Materials Availability
All new unique reagents generated in this study are available from the lead contact with a completed materials transfer agreement.

Data and Code Availability
The published article includes all datasets generated or analyzed during this study.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101498.

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AUTHOR CONTRIBUTIONS
D.G. conceived the idea and designed the study. D.G., H.C., D.D., and L.X. prepared nanocomposite powder. D.G., H.C., D.D., H.Z., T.Z., and C.H. conducted laser additive manufacturing experiments. H.C., D.D., and C.M. developed theoretical models and methods for simulation. H.C., D.G., C.M., and K.L. characterized microstructures and properties. D.G., H.C., A.G., and R.P. analyzed the data. D.G. wrote the paper. D.G. and R.P. supervised the whole work.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Carbon Nanotubes Enabled Laser 3D Printing of High-Performance Titanium with Highly Concentrated Reinforcement

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Figure S1. Finite volume method (FVM) simulation of thermal evolution behavior of molten pool during SLM of 1.0 wt.% CNTs/Ti nanocomposite powder using different laser powers. (A) Profiles of temperature evolution and attendant cooling rate of molten pool during solidification. (B) Profiles of temperature versus distance from the center of molten pool and the corresponding temperature gradient at solid/liquid interface within molten pool. Parameters for simulation of SLM of Ti-based nanocomposite powder in the FVM model included: laser spot size of 70 μm, laser powers of 150 W and 350 W, laser scan speed of 1,300 mm s⁻¹, hatch spacing of 40 μm, powder layer thickness of 40 μm and ambient temperature of 300 kelvin (K). Related to Figure 2.
Figure S2. Crystallographic analysis of grains and grain boundaries (GBs) in SLM-processed Ti-based sample using a laser power of 350 W and a scan speed of 1,300 mm s\(^{-1}\). (A) Cross-sectional EBSD GB map on the ultrafine equiaxed grains with high-angle grain boundaries (HAGBs) (blue), low-angle grain boundaries (LAGBs) (green) and subgrain boundaries (grey) superimposed. The building direction of the sample in this image is vertical. (B) Distribution of grain sizes of as-built Ti-based sample revealed by EBSD measurement. (C) Pole figures (above) and inverse pole figures (below) for as-printed Ti-based sample. Related to Figure 2.
Figure S3. Crystallographic analysis of grains and grain boundaries (GBs) in SLM-processed Ti-based sample using a relatively low laser power of 150 W and a scan speed of 1,300 mm s\(^{-1}\). (A) Cross-sectional EBSD GB map on the typical columnar grains with HAGBs (blue), LAGBs (green) and subgrain boundaries (grey) superimposed. The building direction of the sample in this image is vertical. (B) Distribution of grain sizes of as-built Ti-based sample revealed by EBSD measurement. (C) Pole figures (above) and inverse pole figures (below) for the printed Ti-based sample. Related to Figure 2.
Figure S4. Effects of laser power and scan speed on microstructural development of in-situ TiC reinforcement in SLM-processed Ti-based samples. (A) The refined dendritic structure at 250 W; (B) the considerably refined nanoscale lamellar structure at 350 W; (C) the re-coarsened rod-like structure at 450 W. An optimal laser scan speed was kept constant at 1,300 mm s$^{-1}$ for this batch of specimens in order to highlight the exclusive influence of laser power. (D) The agglomerated and coarsened TiC lamellae at 500 mm s$^{-1}$; (E) the gradually refined and dispersed TiC lamellae at 900 mm s$^{-1}$; (F) the slightly agglomerated and coarsened TiC lamellae at 1,700 mm s$^{-1}$. A sufficiently high laser power of 350 W was optimized for this batch of experiments in order to realize a general formation of lamellar structured TiC reinforcing phase. Related to Figure 2 and Figure 3.
Figure S5. Effects of CNTs contents and laser printing parameters on yield strength ($\sigma_y$), ultimate tensile strength (UTS, $\sigma_b$) and fracture elongation ($\delta$) of the tensile samples reported in Figure 5A. (A) Ti-based samples with 0.5 wt.% CNTs addition. Laser printing parameters are: A1 sample, laser power $P=350$ W, scan speed $v=1,300$ mm s$^{-1}$, hatch spacing $h=50$ μm; A2 sample, $P=350$ W, $v=1,500$ mm s$^{-1}$, $h=50$ μm; A3 sample, $P=375$ W, $v=1,700$ mm s$^{-1}$, $h=50$ μm; A4 sample, $P=400$ W, $v=1,800$ mm s$^{-1}$, $h=50$ μm. (B) Ti-based samples with 1.0 wt.% CNTs addition. Laser printing parameters are: B1 sample, $P=350$ W, $v=1,300$ mm s$^{-1}$, $h=40$ μm; B2 sample, $P=375$ W, $v=1,400$ mm s$^{-1}$, $h=40$ μm; B3 sample, $P=375$ W, $v=1,600$ mm s$^{-1}$, $h=40$ μm; B4 sample, $P=425$ W, $v=1,800$ mm s$^{-1}$, $h=40$ μm. Related to Figure 5.
TRANSPARENT METHODS

Materials

Commercially pure (CP) titanium (Ti) powder. CP-Ti powder (American Society for Testing and Materials, ASTM, Grade 1) was supplied by TLS Technik GmbH, Germany. The powder consisted of Ti (balance), N (0.005%), C (0.01%), H (0.002%), Fe (0.16%) and O (0.09%), in weight percent (wt.%). The CP-Ti powder had a spherical shape and a particle size distribution of 15–53 μm, with a mean particle size of 39.6 μm.

Carbon nanotubes (CNTs) powder. Multi-walled CNTs powders with a purity larger than 95% were supplied by abcr GmbH, Germany and Chengdu Organic Chemicals Co. Ltd., Chinese Academy of Sciences, China. The CNTs powders had an outside diameter of 8–15 nm, a length of 10–50 μm and a specific surface area (SSA) larger than 180 m²/g.

CNTs/Ti nanocomposite powder. Two batches of CNTs/Ti nanocomposite powder consisting of 0.5 wt.% and 1.0 wt.% CNTs were prepared respectively in a Pulverisette 4 vario-planetary ball mill (Fritsch GmbH, Germany), using a ball-to-powder weight ratio of 2:1 and a rotation speed of main disk of 200 rpm. The operative milling time was set at 4 h and a ball-milling duration of 20 min was followed by an interval time of 10 min, in order to avoid the temperature rise within the grinding bowls. To avoid oxidation of the powder, the milling treatment was under the protection of an argon atmosphere within the bowls.

Selective Laser Melting (SLM)

Laser additive manufacturing of as-prepared CNTs/Ti nanocomposite powder was performed on two SLM machines developed by Nanjing University of Aeronautics and Astronautics (NUAA) and Fraunhofer Institute for Laser Technology (ILT), respectively. These two machines had the basically same configurations, consisting mainly of a single mode, continuous wave modulated YLR-500-SM ytterbium fiber laser with a maximum
power of 500 W and a spot size of 70 μm (IPG Laser GmbH, Germany), a hurrySCAN 30 scanner (SCANLAB GmbH, Germany) with a scan speed up to 7.0 m/s, an automatic flexible powder spreading mechanism with carbon fibers braided brush, an inert argon gas protection system, and software tools to manage and control 3D printing process. As specimens were to be built, a titanium substrate was fixed on the building platform and fitted precisely with the powder spreading brush. The building chamber was then sealed and the argon gas was fed inside, decreasing the O₂ content below 10 ppm. Afterward, the CNTs/Ti nanocomposite powder was deposited on the substrate by the layering mechanism, with the powder layer thickness of 40 μm. The laser beam was controlled by the software data to scan the powder bed surface selectively, producing a two-dimensional consolidated profile. An ‘island scanning’ strategy was applied by dividing the cross-section of a specimen into islands with a 5 mm × 5 mm size to reduce the vector length. The scan angle (SA) was rotated with 37º in each island between neighboring layers (Figure 1F). Samples of 10 mm × 10 mm × 8 mm blocks were built using a wide range of SLM parameters, i.e., laser powers of 100–400 W and scan speeds of 400–2,000 mm s⁻¹, for SLM parameters optimization by examining densification and microstructures. Specimens of 70 mm (length) × 12 mm (width) × 6 mm (height) tensile blocks were further prepared using the optimized parameters that enabled to achieve a SLM densification larger than 99.5%. The 70 mm × 12 mm rectangular plates of specimens were built parallel to the underlying substrate. The completed specimens were removed from the machine and cleaned of residual powder by compressed dry air.

**X-Ray Diffraction (XRD)**

Phase identification was conducted by XRD using a D8 Advance X-ray diffractometer (Bruker AXS GmbH, Germany) with Cu Kα radiation at 40 kV and 40 mA, using a continuous scan mode at 1°/min. A slower scan rate of 0.01°/min was further applied over 2θ=34°–36° to obtain a more accurate determination of diffraction peaks. XRD was
utilized to evaluate peak broadening with grain size and lattice strain due to dislocations, using the modified form of Williamson-Hall plot as follows (Yogamalar et al., 2009):

$$\beta_{hkl} \cos \theta = \left( \frac{k\lambda}{D_v} \right) + (4\varepsilon \sin \theta)$$  \hspace{1cm} (1)

where $D_v$ is the volume weighted crystallite size, $k$ is the shape factor (0.9), $\lambda$ is the wavelength of Cu Kα radiation, $\beta_{hkl}$ is the instrumental corrected integral breadth of the reflection (in radians) located at $2\theta$, and $\theta$ is the angle of reflection (in degrees). Equation (1) represents a uniform deformation model (UDM), where the strain was assumed to be uniform in all crystallographic directions (Yogamalar et al., 2009). From the linear fit to the data, as shown in Figure 4b, the crystallite size was extracted from the Y-intercept and the lattice strain was deduced from the slope of the fit. The weight fraction of in-situ formed TiC, $W_{\text{TIC}}$, within the TiC/Ti dual-phase system was calculated by XRD analysis as follows (Chung, 1974):

$$W_{\text{TIC}} = \frac{I_{\text{TIC}}/K_{\text{TIC}}}{I_{\text{TIC}}/K_{\text{TIC}} + I_{\text{Ti}}/K_{\text{Ti}}}$$  \hspace{1cm} (2)

where $I_{\text{TIC}}$ and $I_{\text{Ti}}$ are the diffraction intensities of TiC and Ti phases, and $K_{\text{TIC}}$ and $K_{\text{Ti}}$ are constants related to the materials and can be found in JCPDS Card No. 71-0298 and JCPDS Card No. 89-2959.

**Microstructural Characterization**

Laser-printed parts were removed from the building plates using wire electro discharge machining (EDM). Samples for metallographic examinations was ground with 320, 600, 1,000, 1,200 and 2,000 grit sand paper. Final polishing of the samples was performed using 0.5 μm diamond polishing agent. The polished samples were etched with a solution consisting of HF (5 ml), HNO₃ (15 ml) and H₂O (150 ml) for 20 s to reveal microstructures. Microstructures were characterized using a S-4800 field emission scanning electron microscope (FE-SEM; Hitachi, Ltd., Japan). Electron backscatter diffraction (EBSD) was performed in a Nova™ NanoSEM 430 FE-SEM (FEI Company,
USA). A typical scan area of 2 mm × 2 mm on the cross-sections of laser-printed specimens was used for EBSD analysis with a scanning step size of 0.5 μm. EBSD data were analyzed using the Orientation Imaging Microscopy (OIM) analysis software provided by EDAX. Interfaces between grains were divided into three categories: sub-grain boundaries when the misorientation angle was 2°–5°, low-angle grain boundaries (LAGBs) when the misorientation angle was 5°–15°, and high-angle grain boundaries (HAGBs) with the misorientation angle of 15°–180°. To calculate the grain size, each grain was approximated as a sphere to reveal the grain diameter. Transmission electron microscope (TEM) specimens were prepared by standard electropolishing techniques. Bright-field (BF), dark-field (DF) and high-resolution (HR) TEM and corresponding selected area electron diffraction (SAED) analyses were conducted in a FEI Tecnai G2 F20 TEM operated at 200 kV with a point-to-point resolution less than 0.2 nm. The lattice misfit $\delta$ between Ti matrix and TiC reinforcing phase was calculated by:

$$\delta = (d_{\text{Ti}} - d_{\text{TiC}})/d_{\text{Ti}}$$

where $d_{\text{Ti}}$ and $d_{\text{TiC}}$ are interplanar spacings of Ti and TiC phases at the interface. The chemical concentration maps were acquired by energy-dispersive spectroscopy (EDS) in TEM, using a Bruker QUANTAX Silicon Drift Detector (SDD) operated at an accelerating voltage of 200 kV.

**Mechanical Testing**

Uniaxial tensile tests were performed at room temperature in a CMT5205 tensile machine (MTS Industrial Systems, China) equipped with a 100 kN load cell. The dog-bone-shaped tensile samples (gauge length: 25.0±0.1 mm, width: 6.0±0.1 mm, thickness: 5.0 mm) were prepared according to ASTM E8 standard. The extension velocity of the cross head was 0.2 mm/min and the specimens were loaded continuously until fracture. The loading direction of the tensile samples is perpendicular to the SLM building direction. Six tensile specimens were tested for each selected laser printing parameter to yield standard error
bars. In order to disclose the deformation behaviors of laser-printed parts subjected to uniaxial loading, the deformation microstructures at different levels of ~3% and ~8% tensile strains were characterized by a FEI Tecnai G2 F20 TEM operated at 200 kV. A series of BF-TEM, HR-TEM and SAED images were recorded to reveal the deformation behavior and underlying strengthening mechanisms of specimens from multiple scales.

**Numerical Simulation**

The thermodynamic behavior during SLM of CNTs/Ti nanocomposite powder was simulated to reveal the formation conditions for in-situ Ti-based composites during laser rapid solidification. A continuous three-dimensional geometry of Ti-based nanocomposites was established and adaptively meshed by the definition of wall conditions using a Gambit software. The defined geometry was imported into a FLUENT software using a finite volume method (FVM) model. The laser energy with a Gaussian distribution was defined as a heat flux applied onto the free surface of powder bed considering the laser penetration depth. Three categories of coupled heat transfer mechanisms including the heat irradiation to powder layer from laser beam, the heat conduction among powder particles, and the heat convection between molten pool and ambient atmosphere were considered in the model. The energy, mass and momentum transformation were loaded on the top surface and the multiphase interface of the physical model. More details about the FVM model, calculation process and data extraction can be found in our previous work (Dai and Gu, 2015; Gu and Yuan, 2015; Yu et al., 2016).
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