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Nanostructured laminar tungsten alloy with improved ductility by surface mechanical attrition treatment

Hong-Yan Guo1,2,3, Min Xia2, Lap-Chung Chan3, Kun Wang1, Xiao-Xin Zhang2, Qing-Zhi Yan2, Man-Chao He1, Jian Lu3,5 & Chang-Chun Ge2

A nanostructured laminar W-La2O3 alloy (WL10) with improved ductility was prepared using a surface mechanical attrition treatment (SMAT). φ1.5 mm ZrO2 WL10 balls subjected to SMAT (called φ1.5 mm ZrO2 ball SMATed WL10) samples possess the best surface profile and excellent integrated mechanical properties (the ductile-brittle transition temperature (DBTT) value decreases by approximately 200 °C, and the bending strength decreases by 100 Mpa). A highly dense group of laminates was detected near the surface of the φ1.5 mm ZrO2 ball SMATed WL10 sample. The SMATed WL10 laminates were composed of a micro-grain layer, an ultrafine-grain layer and a nanosized-grain layer. The nanostructured laminar surface layer of the φ1.5 mm ZrO2 ball SMATed WL10 sample is approximately 1–2 μm. The top surface of the WL10 plates with and without the SMAT process possesses residual compressive stress of approximately −883 MPa and −241 MPa, respectively, in the y direction and −859 MPa and −854 MPa, respectively, in the x direction. The SMAT process could be a complementary method to further improve the toughness of tungsten-based materials.

Tungsten (W) and its alloys are now considered to be the most promising candidates for plasma facing materials (PFMs) in future fusion reactors because of the high melting point, high thermal conductivity, low deuterium/tritium retention and low sputter rates. However, these materials exhibit serious types of embrittlement, such as low-temperature brittleness, high-temperature or recrystallization brittleness and radiation-induced brittleness and hardness1,2. Many methods have been developed to improve the performance of tungsten. Nanostructured (NS) W materials seem to be promising for nuclear applications since the large number of grain boundaries and dislocations in nanostructured W may result in better mechanical properties; in addition, grain boundaries could act as sinks for interstitial atoms, thus diminishing the degradation of the mechanical and thermal properties3,4. Many methods, such as powder metallurgy (PM), severe plastic deformation (SPD) (including equal-channel angular extrusion (ECAE)6,7 and high pressure torsion (HPT)8) and surface machining9, have been employed to refine the grains to ultrafine-grained (UFG) (>100 nm) or nanoscale sizes (<100 nm) to improve the mechanical properties or irradiation resistance performance of W9. However, nanostructures generally possess low ductility. Another method is to introduce extrinsic toughening mechanisms to prevent the propagation of cracks; these methods include reinforcement induced by laminates, fibres, whiskers, and particles, which primarily act behind the crack tip and locally shield it from the driving force10,11. A solution to improve the ductility of tungsten or tungsten alloys is to fabricate laminar nanostructured materials that could deflect and re-nucleate cracks. Residual

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stress could often be induced in laminar structures during the fabrication process. Well-designed residual surface compression may prove to be extremely useful in inhibiting the growth of defects. In this study, we first apply the surface mechanical attrition treatment (SMAT) to bulk W–La2O3 (1 wt%) (WL10) to fabricate a nanostructured laminar tungsten alloy with improved ductility. WL10 is generally considered an important structural material in plasma facing components and has been shown to exhibit better mechanical properties, higher recrystallization temperature, and greater toughness than pure tungsten. SMAT is a manufacturing process that is based on the impact of spherical projectiles onto the sample surface and is proposed as an effective method to achieve grain refinement down to the nanometre scale in the top surface layer; this method has been applied to many metals and alloys. Significant enhancements in the mechanical properties, such as tensile strength, fatigue limit, friction, and wear resistance, have been achieved with this method. SMAT can be used to prepare nanostructure layers without an interface in bulk materials, resulting in a good bonding strength between surface layer and matrix. However, only a surface layer containing the nanocrystalline structure can be obtained using SMAT, and this method can only be used to process a sample with a flat surface of limited size. In this manuscript, the extremely large strain and strain rate induced by the SMAT process were designed to store more dislocations in the intrinsic grains at room temperature, refine the grains into the nanograin size, generate a laminar structure, and induce residual stress; these properties are expected to improve the mechanical properties and irradiation resistance of WL10.

**Experimental Methods**

The material used in this investigation was a rolled WL10 plate (120 mm × 100 mm and 4 mm in thickness) of commercial purity (Beijing Tian-long Tungsten & Molybdenum Co., LTD). The setup and procedures of the SMAT process are described in the literature. In this study, SMAT applied to WL10 was performed under vacuum at room temperature. Different balls (shots) with different diameters were used to study the SMAT process on WL10, and details of the SMAT parameters are described in Table 1. Both sides of each WL10 plate were treated. A cross-sectional TEM lamella was prepared using a focused ion beam (FIB) with a Zeiss Ultra 55. Transmission electron microscopy (TEM) observations were carried out on JEOL2010 and FEI Tecnai F20 microscopes operated at 200 kV. Three-point bending (3PB) specimens of 34 mm × 4 mm × 3 mm (length × width × height) were produced to measure the bending strength (see Fig. 1 for the cutting sketch of the specimens). Vickers micro-hardness tests were performed on the polished surface under a load of 200 g for 15 s. For the Charpy impact tests on specimens with dimensions of 27 mm × 4 mm × 3 mm (length × width × height), a notch depth of 0.1 mm and a notch root radius of 0.1 mm were created on the WL10 samples (see Fig. 1 for the cutting sketch of the specimens). All Charpy impact test samples were notched and prepared in the L-S direction (where “L” is longitudinal, which is in the rolling direction, and “S” is short transverse, which is the direction of the thickness of the plate). All samples were cut by electrical discharge machining. The specimens were heated to 673 K, 698 K, 723 K, 773 K, 873 K, and 973 K in an argon atmosphere and then pushed to the support outside the furnace and immediately hit by a striker so that the samples were exposed to the air for a very short period. Because the contact time is short, the effects of air can be ignored. Three samples were tested for each temperature. Residual stresses

| Ball size/mm | WC | Steel | ZrO2 |
|--------------|----|-------|------|
| Amplitude %  | 80 | 80    | 80   |
| Time/min     | 30 | 30    | 30   |
| Vibrating frequency/KHz | 20 | 20    | 20   |

**Table 1.** SMAT processes for the WL10 plate.
were measured in the directions perpendicular to the SMATed surface of WL10 using a μ-X360n portable X-ray residual stress analyser (Pulstec Industrial Co., Ltd.)

Results and Discussion
Optimization of the SMAT process. In the SMAT process, the velocity of the balls is a critical parameter in forming the nanostructures since it is significant in determining the strain-rate and localized strain of the material. Generally, a higher velocity results in a higher strain-rate, consequently enhancing the formation of nanostructures. Many parameters may affect the ball velocity in SMAT, such as the ball density, ball size and the power of the ultrasonic generator, as well as the mechanical properties (mainly the hardness) of the sample.

In this study, WC, steel, and ZrO₂ balls with different densities were used to optimize the SMAT process for WL10. Figure 2 shows the surface profiles of WL10 before SMAT (2a) and after WC ball treatment (2b), ZrO₂ ball treatment (2c), and steel ball treatment (2d). The WC ball SMATed WL10 sample possessed the highest surface roughness; therefore, the steel and ZrO₂ ball-treated WL10 surface seem to be the most appropriate. The mass loss of different ball SMATed WL10 samples is shown in Table 2, revealing that the weight loss of the WC ball SMATed WL10 was up to 72 g (the most serious), which is consistent with the surface profile shown in Fig. 2(b). The surface profile and weight loss results indicated that the amount of energy transmitted by the WC balls is too high since lighter balls can attain a higher ceiling impact velocity. However, the amount of energy transmitted by a light ball is less than that of a ball with a larger mass. Thus, the WC ball can be eliminated for the WL10 in the SMAT process.

Figure 3 shows the 3PB bending strength of the commercial WL10 (920 MPa), ZrO₂ (760 MPa), WC (740 MPa), and steel ball (700 MPa) SMATed WL10 samples. A decrease in the bending strength was observed in all SMATed samples. In addition, the microhardness was tested in this work since it could reflect the relative density and the strain-hardening effects.

The microhardness distributions in the samples after SMAT was applied to various ball compositions are depicted in Fig. 4. As seen, the microhardness of the WL10 tested from the top surface to 60 μm is lower than the mean value of the matrix, revealing that the surface may possess defects induced by machining. Additionally, all the SMATed samples possess the lowest microhardness at the positions of 120–140 μm beneath the top surface, indicating the existing of microcracks. With microhardness positions above 140 μm, the ZrO₂ ball SMATed

| Parameter | WC | Steel | ZrO₂ |
|-----------|----|-------|------|
| Ball size/mm | ϕ2 | ϕ3 | ϕ1.5 |
| Δm/g | −72 | −3 | −3 |

Table 2. Weight loss of different ball SMATed WL10 samples. “−” reference the SMATed sample mass loss.
WL10 possesses a mean microhardness higher than that of the untreated WL10. In contrast, the WC and steel ball SMATed WL10 samples have a mean microhardness lower than that of the untreated WL10. These results indicate that the ZrO2 ball resulted in the best strain-hardening effect on the WL10. Since the ZrO2 ball has a smaller density than that of the WC steel ball, it can attain a higher velocity, and its sufficient weight enables a higher impinging energy to be carried to the sample surface. Therefore, the ZrO2 ball could be more efficient than the WC steel ball. Based on the surface profiles and the 3PB bending strength and microhardness results, we selected the ZrO2 ball to further study the mechanical properties and microstructures of SMATed WL10.

Mechanical properties and microstructure of the ϕ1.5 mm ZrO2 ball SMATed WL10. The ductility of the WL10 and the ϕ1.5 mm ZrO2 ball SMATed WL10 was characterized using Charpy impact tests. Since the commercial rolled tungsten sample was anisotropic and since the ductile-brittle transition temperature (DBTT) strongly depends on the orientation of the specimen, the Charpy curve may not be suitable to estimate the DBTT value. In this case, the sample profile with fracture features was more acceptable to reveal the DBTT shift. The corresponding absorbed energy as a function of the test temperature of the WL10 and ϕ1.5 mm ZrO2 ball SMATed WL10 is listed in Table 3. Figure 5 shows the sample profile and reveals the appearance of the fractures. As seen, the estimated DBTT of the WL10 was approximately 973 K compared with the DBTT value (between 698 K and 723 K) of the ϕ1.5 mm ZrO2 ball SMATed WL10. Of particular interest, the fracture sample of the SMATed WL10 possesses a thin tail layer on the un-notched surface at 698 K, 723 K, and 773 K, which was different from the delamination feature from un-SMATed WL10 at 873 K and from the features listed in other reports. The SMATed sample at 773 K was almost fractured, except for the SMAT-induced thin layer connecting the two fractured pieces.

Figure 3. 3PB bending strength of different ball SMATed WL10 samples.

Figure 4. Microhardness of different samples taken from the cross-sectional plane in the depth direction. “D” references the distance beneath the top surface.

Figure 6(a) shows the cross-sectional SEM image of the untreated commercial WL10. Several microcracks were observed near the surface, which may be attributed to the rolling effect during the deformation process. For the ϕ1.5 mm ZrO2 ball SMATed WL10, as indicated in Fig. 6(b), a highly dense group of laminates was detected near the surface. The laminar structure is further demonstrated in Fig. 6(c) where the thickness of the individual laminae was approximately 1–2 μm and the width was approximately 50 μm. In addition, many microcracks...
Table 3. The absorbed energy as a function of the test temperature of the WL10 and the ϕ1.5 mm ZrO₂ ball SMATed WL10.

| Test temperature & Samples | WL10 (Absorbed Energy, J) | ϕ1.5 mm ZrO₂ ball SMATed WL10 (Absorbed Energy, J) |
|----------------------------|---------------------------|--------------------------------------------------|
| 400°C                      | 0.375                     | 0.875                                            |
| 425°C                      | —                         | 1                                                |
| 450°C                      | 0.375                     | 1.75                                             |
| 500°C                      | 0.375                     | 3.25                                             |
| 600°C                      | 0.375                     | 5.25                                             |
| 700°C                      | 5.25                      | 8.75                                             |

Figure 5. The Charpy impact test results of the WL10 and the ϕ1.5 mm ZrO₂ ball SMATed WL10.

Figure 6. Cross sectional profile of WL10. (a) Untreated commercial WL10, (b) ϕ1.5 mm ZrO₂ ball SMATed laminates observed near the surface, (c) enlarged SEM image of laminates in (b), and (d) cross sectional nanostructured surface layer taken from the FIB sample.

were observed accompanying the laminates. Figure 6(d) shows the cross-sectional nanostructured surface layer taken from the FIB sample. As shown in the figure, after the SMAT process on the WL10 plate, a nanosized-grain layer with a mean grain size of approximately 50 nm and a neighbouring UFG layer with a mean grain size of less
than 100 nm form on the surface with a total thickness of approximately 1–2 μm in which the grain size gradually increases as a function of the depth beneath the surface. These results revealed that the SMATed WL10 laminates were composed of a micro-grain layer, a UFG layer and a nanosized-grain layer.

TEM images of the top surface layer of the φ1.5 mm ZrO₂ ball SMATed WL10. The nanostructured surface layer was further revealed through TEM characterization. Figure 7(a) shows the cross-sectional TEM micrograph of the φ1.5 mm ZrO₂ ball SMATed WL10 sample, taken from the SMATed top surface, as indicated in Fig. 6(d). As seen in the figure, the gradient structure results from a gradual decrease in the applied strain with an increasing depth of the deformed layer, from the top surface to the substrate. Elongated W grains can be observed closest to the surface. Individual grains were prepared with sizes less than 50 nm and are separated by high-angle grain boundaries. Most of the grains are heavily strained and contain a high density of dislocations. The selected area electron diffraction (SAED) patterns of the SMATed layer in Fig. 7(b) are shown in circles, indicating the large fraction of high-angle grain boundaries. Additionally, some diffraction spots are split, which

Figure 7. (a) Cross-sectional TEM micrograph of the φ1.5 mm ZrO₂ ball SMATed WL10 and (b) the SAED pattern taken from (a).

Figure 8. TEM images of dislocations and dislocation loops including (a) bright field and (b) dark field.
could be caused by the existence of low-angle grain boundaries. The azimuthal spreading of the diffraction spots is approximately 3–5°, indicating the existence of high internal strain. The high density of dislocations was further revealed in Fig. 8, as shown in the bright field and dark field TEM images of the W sample surface. The dislocation density of the W grains was approximately $3.5 \times 10^{12} \text{ cm}^{-2}$. According to the microhardness results, the bending strength, cross-sectional profiles and micrographs, the decrease in the bending strength of the ϕ1.5 mm ZrO$_2$ ball SMATed WL10 may contribute to the existence of microcracks in the laminar structure. However, in the case of the toughness of the ϕ1.5 mm ZrO$_2$ ball SMATed WL10 sample, the given value of the DBTT in Fig. 5 should be lower (that is, the SMATed sample was tougher than the given data) since a notch depth of 1 mm and a notch root radius of 0.1 mm were created on the SMATed WL10 (the striker hit the side, which was cut first). Therefore, only one side contributed to the decrease of the DBTT in Fig. 5. We prepared un-notched Charpy test samples; however, the ϕ1.5 mm ZrO$_2$ ball SMATed WL10 was so tough that even the striker (25J) could not destroy it at room temperature. Therefore, the SMAT process was expected to be effective in decreasing the DBTT of the WL10.

To investigate the origin of the toughness improvements by the SMAT process, the microstructure of the ZrO$_2$ ball SMATed and Charpy tested WL10 was observed. Figure 9 shows the fracture surfaces of due to the Charpy impact on ϕ1.5 mm ZrO$_2$ ball SMATed WL10. When tested at 673 K, the sample exhibited an almost fully brittle cleavage fracture (see Fig. 9(a)). However, a very thin tail layer was observed on the un-notched and fractured surface, as shown in Fig. 9(b). When the temperature was increased to 698 K, the brittle fracture also dominated (Fig. 9(c)), but the length of the thin tail layer increased. Figure 9(e) shows the ductile fracture at 723 K, indicating that the DBTT is approximately 723 K. The thin tail layer was observed in both Figs 5 and 9(f). Of particular interest, apparent laminar structures were also observed in the SMATed thin tail layer of the fractured surfaces, as

![Figure 9. Fracture surfaces of the Charpy impact for the ϕ1.5 mm ZrO$_2$ ball SMATed WL10; (a,b) 673 K, (c,d) 698 K, and (e,f) 723 K.](image-url)
indicated in Fig. 9(b,d and f). In contrast, no laminar structure was detected in the un-treated commercial WL10, as indicated in the fracture surface at 873 K in Fig. 10(a and b).

Meanwhile, during the grain refinement in the SMAT process, high compressive residual stresses could be simultaneously induced in the surface layer. In this case, a well-designed residual surface compression could be helpful in inhibiting the growth of defects\(^\text{13, 14}\). In this manuscript, the residual stress was measured for the specimens with and without SMAT, as shown in Fig. 11. The top surface of the WL10 plates with and without SMAT possesses a residual compressive stress of approximately \(-883\) MPa and \(-241\) MPa in the \(y\) direction, and \(-859\) MPa and \(-854\) MPa in the \(x\) direction, respectively. The residual compressive stress in the \(x\) direction with and without SMAT exhibited similar values, and the residual compressive stress in the un-treated surface may be attributed to the deforming and rolling effect of the WL10 plate. However, in the \(y\) direction, the residual compressive stress was significantly improved using the SMAT process.

The results revealed that the SMAT process could be used to prepare nanostructured laminar tungsten alloys with improved ductility, and a high compressive residual stress could be simultaneously induced on the surface layer. However, the origin of the good ductility was manifold due to the multi-scale-based deformation mechanisms and fracture mechanisms integrated into the laminates. The following two major factors contributed to the ductility improvements of the SMATed WL10 sample.
could be a complementary method to further improve the toughness of tungsten-based materials. The SMAT process could be the three major factors that contribute to the ductility improvements of the SMATed WL10. The SMAT process enables a higher impinging energy to be carried to the sample surface. The ϕ1.5 mm ZrO2 ball SMATed WL10 sample possesses a smaller density than that of a WC steel ball; therefore, it can attain a higher velocity, and its sufficient weight has a smaller density than that of a WC steel ball; therefore, it can attain a higher velocity, and its sufficient weight could impede advancement of the crack along the direction perpendicular to the interface plane when the first crack was initiated. Therefore, the toughening mechanisms induced by the compressive residual stress and the multiple parallel crack shielding (laminates) were considered to contribute to the ductility improvements of the SMATed WL10.

Nanostructured laminar WL10 was prepared using a surface mechanical attrition treatment (SMAT). A ZrO2 ball with a diameter of 1.5 mm was used to prepare the SMATed WL10 sample. The SMATed WL10 laminates were composed of a micro-grain layer, an ultrafine-grain layer, and a nanosized-grain layer. The nanostructured laminar surface layer of the SMATed WL10 sample is approximately 1–2 μm. The bending strength of the SMATed WL10 was attributed to the microcracks in the laminates. The top surface of the WL10 plates with and without the SMAT treatment were conditionally induced by the SMAT process (as indicated by the SAED pattern in Fig. 7(b)), which demonstrates the presence of highly dense and well-aligned dislocations. In addition, a large fraction of high-angle grain boundaries was detected near the surface of the SMATed WL10 sample. A highly dense group of lamellae was initiated by the SMAT process (as indicated by the SAED pattern in Fig. 7(b)), which demonstrates the presence of highly dense and well-aligned dislocations. In addition, a large fraction of high-angle grain boundaries was detected near the surface of the SMATed WL10 sample. A highly dense group of lamellae was observed at the fracture surface, as shown in Fig. 6. Thus, most of the cracks were arrested in the nanosized-grain thin tail layer. The existence of the compressive residual stress in the SMAT layer could impede advancement of the crack along the direction perpendicular to the interface plane when the first crack was initiated. Therefore, the toughening mechanisms induced by the compressive residual stress and the multiple parallel crack shielding effects greatly improve the ductility of SMATed WL10.

Table 4 gives a comparison between the present and previously reported data related to the DBTT of tungsten-based materials. Generally, the DBTT of tungsten-based materials decreases by alloying, particle strengthening, and working processes such as rolling, forging, and injection moulding. All of these methods have been demonstrated to effectively improve the ductility of tungsten-based materials. In the case of SMAT processing in this report, the DBTT of the SMATed WL10 sample is approximately 723 K (intuitively seen in Fig. 5 and Table 4), which is much lower than that of the untreated bulk WL10 in this report at 973 K. In this sense, the SMAT process could be a complementary method to further decrease the DBTT value of tungsten-based materials.

Conclusions

Nanostructured laminar WL10 was prepared using a surface mechanical attrition treatment (SMAT). A ZrO2 ball has a smaller density than that of a WC steel ball; therefore, it can attain a higher velocity, and its sufficient weight enables a higher impinging energy to be carried to the sample surface. The ϕ1.5 mm ZrO2 ball SMATed WL10 sample possesses the best surface profile. A highly dense group of lamellae was detected near the surface of the ϕ1.5 mm ZrO2 ball SMATed WL10 sample. The SMATed WL10 laminates were composed of a micro-grain layer, an ultrafine-grain layer, and a nanosized-grain layer. The nanostructured laminar surface layer of the ϕ1.5 mm ZrO2 ball SMATed WL10 sample is approximately 1–2 μm. The bending strength of the SMATed WL10 was attributed to the microcracks in the laminates. The top surface of the WL10 plates with and without the SMAT process possesses residual compressive stresses of approximately −883 MPa and −241 MPa, respectively, in the y direction, and −859 MPa and −854 MPa, respectively, in the x direction. The existence of the micro-grain layer in the laminate, the compressive residual stress and multiple parallel crack shielding (laminates) were considered to be the three major factors that contribute to the ductility improvements of the SMATed WL10. The SMAT process could be a complementary method to further improve the toughness of tungsten-based materials.

Table 4. Comparison between the present and previously reported data related to the DBTT of tungsten-based materials.

| Material/Size | Working process | DBTT(K) | Dimension(mm) | Method  | Ref. |
|---------------|-----------------|---------|---------------|---------|------|
| WL10          | Swaging + Rolling; SMAT | ≤723    | 27 × 4 × 3    | Charpy  | our  |
| WL10          | Swaging + Rolling | 973     | 27 × 4 × 3    | Charpy  | our  |
| WL10          | Swaging + Rolling | 973     | 10 × 10 × 55  | Charpy  | 28   |
| W – 0.5 ZrC (8.5 mm thick plate) | Rolling | 373     | 2 × 2 × 20    | 3PB    | 29   |
| W – 2Y2O3 (2 mm thick, ϕ95 mm) | Hot Forging | 473     | 2 × 2 × 25    | 3PB    | 30   |
| Pure W (10 mm thick plate) | Rolling | 473     | 2 × 4 × 20    | 3PB    | 31   |
| Pure W (4 mm thick) | HIP | 473     | 2 × 3.3 × 20  | 3PB    | 31   |
| W – 0.2 TiC (1 mm thick) | Forging + Rolling | 440     | 1 × 1 × 20    | 3PB    | 32   |
| W – 0.25 Ti – 0.05 C (1 mm thick plate) | Rolling | 260     | 1 × 1 × 20    | 3PB    | 32   |
| W – 1% Y2O3 | Injection molding | 1273    | 3 × 4 × 27    | Charpy  | 33   |
| Pure W | Injection molding | 1173    | 3 × 4 × 27    | Charpy  | 33   |
| W – 0.5 TiC | HIP + Forging | 484     | 1 × 1 × 20    | 3PB    | 34   |
| Pure W (0.1 mm thick foil) | Rolling + Joining | 373     | 4 × 15 × 33   | Charpy  | 35   |

First, the micro-grain layer in the laminate plays a primary role in improving the ductility since its ability to deform plastically was conditioned by the dislocations. In addition, a large fraction of high-angle grain boundaries at the top surface was induced by the SMAT process (as indicated by the SAED pattern in Fig. 7(b)), which could lead to a high strain-hardening rate in the UFG layer, consequently, leading to a high total elongation.12, 26.

Second, the residual stress strongly affects the mechanical properties and the selection of the crack path in the brittle laminates of ceramic-like materials. A well-designed residual stress profile on the surface of the material and a greater initial compression to a depth slightly below the surface can impede the surface crack propagation. This process toughens the materials and leads to multiple cracking, which serves as a forewarning of the final failure. As indicated in Fig. 11, a high compressive residual stress was induced in the nanosized-grain layer, and multiple cracks were observed at the fracture surface, as shown in Fig. 6. Thus, most of the cracks were arrested in the nanosized-grain thin tail layer. The existence of the compressive residual stress in the SMAT layer could impede advancement of the crack along the direction perpendicular to the interface plane when the first crack was initiated. Therefore, the toughening mechanisms induced by the compressive residual stress and the multiple parallel crack shielding effects greatly improve the ductility of SMATed WL10.

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Conclusions

Nanostructured laminar WL10 was prepared using a surface mechanical attrition treatment (SMAT). A ZrO2 ball has a smaller density than that of a WC steel ball; therefore, it can attain a higher velocity, and its sufficient weight enables a higher impinging energy to be carried to the sample surface. The ϕ1.5 mm ZrO2 ball SMATed WL10 sample possesses the best surface profile. A highly dense group of lamellae was detected near the surface of the ϕ1.5 mm ZrO2 ball SMATed WL10 sample. The SMATed WL10 laminates were composed of a micro-grain layer, an ultrafine-grain layer, and a nanosized-grain layer. The nanostructured laminar surface layer of the ϕ1.5 mm ZrO2 ball SMATed WL10 sample is approximately 1–2 μm. The bending strength of the SMATed WL10 was attributed to the microcracks in the laminates. The top surface of the WL10 plates with and without the SMAT process possesses residual compressive stresses of approximately −883 MPa and −241 MPa, respectively, in the y direction, and −859 MPa and −854 MPa, respectively, in the x direction. The existence of the micro-grain layer in the laminate, the compressive residual stress and multiple parallel crack shielding (laminates) were considered to be the three major factors that contribute to the ductility improvements of the SMATed WL10. The SMAT process could be a complementary method to further improve the toughness of tungsten-based materials.

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Prof. Jian Lu, Prof. Chang-Chun Ge, Prof. Qing-Zhi Yan, Prof. Man-Chao He designed the project. Dr. Hong-Yan Guo, Min XIA, Lap-Chung Chan, Xiao-Xin Zhang carried out the experiments. Dr. Kun Wang carried out dislocation density calculation in TEM process. Hong-Yan Guo, Min Xia analysed data and wrote the paper under the direction of Prof. Jian Lu, Prof. Chang-Chun Ge and Prof. Jian Lu, Prof. Qing-Zhi Yan, Prof. Man-Chao He revised the paper. All the co-authors contributed to discussions.
Additional Information

Competing Interests: The authors declare that they have no competing interests.

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