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Thermal Stability of CrWN Glass Molding Coatings after Vacuum Annealing

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Abstract: CrWN glass molding coatings were deposited by plasma enhanced magnetron sputtering (PEMS). The microstructure and thermal stability of these coatings were investigated by X-ray diffraction, X-ray photoelectron spectroscopy, scanning electron microscope, transmission electron microscope, atomic force microscope and nanoindentation tests. The as-deposited coating exhibited an aggravated lattice expansion resulting in a constant hardness enhancement. The vacuum annealing induced surface coarsening and the spinodal decomposition of the coating accompanied by the formation of nm-sized c-CrN, c-W₂N, and h-WN domains. The annealed coating with low W content had mainly a face-centered cubic (f.c.c) matrix, strain fields caused by lattice mismatch caused hardness enhancement. Following an increase in W content, the annealed coating showed a mixed face-centered cubic (f.c.c) and hexagonal close-packed (h.c.p) matrix. The large volume fraction of h-WN phases seriously weakened the coating strengthening effect and caused an obvious drop in hardness.

Keywords: CrWN coatings; microstructure evolution; spinodal decomposition; thermal stability; hardness; plasma enhanced magnetron sputtering

1. Introduction

Optical glass lenses have a broad range of applications (e.g., telescopes, cameras, medical equipment, and high-power lasers) because of their good refractive index, excellent chemical stability, light permeability, and high image quality [1–3]. Conventional manufacturing technologies include several techniques (e.g., precision grinding, and ultra-precision lapping and polishing), but these methods are time-consuming and difficult to use for the fabrication of complex shapes [4]. Precision molding, however, has low cost, high efficiency, can be applied to net forming, and is environmentally friendly. These characteristics have attracted great interest from optical manufacturers [5–7]. Unfortunately, the adhesive wear of glass blanks greatly degrades the molding life of the forming die and eventually decreases the quality of the optical glass component surface [8,9].

It has been reported that protective coatings can effectively improve the anti-sticking and anti-wear performance of forming dies [10–18]. Precious metal coatings are widely used in molding manufacturing due to their excellent anti-sticking and anti-oxidation properties [10–12], but they are also expensive for their wide industrial applications. Diamond-like carbon (DLC) films exhibit good self-lubrication and anti-wear performance, however, their poor thermal stability greatly limits any molding application [13,14]. Recently, the application of transition metal nitride coatings to glass molding has received great interest due to the favorable chemical inertness and anti-wear properties of these materials [15–18]. Most current studies focus on the anti-sticking and anti-oxidation properties of transition metal nitride coatings. WCrN coatings have been reported to have good anti-sticking properties at 400 °C, but also exhibited poor anti-sticking performances at 500 °C due to the formation of oxides [19]. CrWN coatings have been reported to experience severe mechanical degradation and coarsening in high temperature nitrogen atmosphere due to the formation of WO₃ phases [20].

The life of glass molding coatings is determined mainly by their anti-sticking and anti-oxidation properties, as well as by their thermal stability. However, relatively little work has been published on the thermal stability of this type of coatings. In this study, CrWN coatings with different W contents are synthesized using PEMS. Detailed characterizations of these coatings were performed to study the evolution of their microstructures, as well as the mechanical properties of the as-deposited coatings after vacuum annealing. The potential effect of the W content on the thermal stability of the annealed coatings will be discussed systematically in the following sections.

2. Materials and Methods

CrWN coatings having different W contents were synthesized by PEMS [21]. Pure W (99.9%) and Cr (99.6%) were used as sputtering targets. A silicon wafer and cemented carbides (WC-8 wt % Co) were used for the substrate. The samples were mechanically polished using a diamond paste and ultrasonically washed using pure ethanol. High purity nitrogen and argon were utilized for the working atmosphere. The samples were placed on a rotated holder, moving at a rotating speed of 2 rpm. The chamber was evacuated to a base pressure of 5×10^{-3} Pa and then heated to 300 °C. Prior to coating deposition, the samples were cleaned by Ar⁺ sputtering to remove any residual pollution and the native oxides. The sputtering parameters were as follows: bias voltage of –120 V, argon flow of 140 sccm, sputtering time 60 min. The CrWN coatings were synthesized in a mixed atmosphere of nitrogen and argon. The chemical composition of the as-deposited coating was varied by adjusting the powers of the Cr and W targets. In this case, the deposition parameters were displayed as follows: bias voltage of –50 V, argon flow 100 sccm, nitrogen flow 100 sccm, and deposition time 100 min. The deposition parameters are listed in Table 1.

Table 1. Experimental details of as-deposited coatings.

Samples	Power of Cr Target (W)	Power of W target (W)	Time (min)
C1	5000	4000	65
C2	2700	4000	100
C3	1600	5100	100

The compositions of the coatings were determined by an energy dispersive spectrometer (EDS) combined with a scanning electron microscope (SEM, Zeiss ΣIGMA HD, Oberkochen, Germany). Moreover, the crystalline structures of the coatings were investigated by X-ray diffraction (XRD, X'Pert Powder, Malvern Panalytical, Malvern, UK) using a Cu K α radiation source with parallel beam. The incident angle was of 1°, while the diffraction angle was scanned from 20° to 90°. Additionally, X-ray photoelectron spectroscopy (XPS, ThermoFisher, K-Alpha+, Waltham, MA, USA) was employed to detect the chemical states of the coatings. The data were collected after 30 s from etching to remove any contaminants adsorbed on the coating surface. A SEM (Zeiss ΣIGMA HD) was employed to characterize the surface morphology of the coatings, while their microstructure was

investigated with a transmission electron microscope (TEM, FEI Titan Cubed Themis G2 300, ThermoFisher, Waltham, MA, USA). Moreover, atomic force microscopy (AFM, Oxford MFP-3DInfinity, Abingdon, UK) was employed to evaluate the surface roughness of the coatings: the scanning area of each image was of $10 \times 10 \mu\text{m}$. The mechanical properties of the coatings were evaluated by using a nm-indentation system (Hyston TI950, Bruker, Billerica, MA, USA): five indentations were performed on each sample; additionally, the hardness and elastic modulus selected at a depth of corresponding to 1/10 of the coating thickness, as to minimize the negative effect of the substrate. Finally, vacuum annealing was conducted in a heat treatment furnace at a pressure of 0.1 Pa (RTP-500, Beijing Ruiyisi Technology Co.LTD, Beijing, China), the annealing temperature and time were of $650 \text{ }^\circ\text{C}$ and 300 min, respectively.

3. Results

Table 2 summarizes the chemical compositions of the Cr-W-N coatings synthesized using different sputtering powers. The Cr content varied from 13.9 ± 0.7 at % to 35.9 ± 2.3 at %, the W content varied from 21.8 ± 5.0 at % to 42.1 ± 6.9 at %, and the N content varied from 40.2 ± 4.8 at % to 43.1 ± 4.4 at %. The oxygen content fluctuated between 0.9 ± 0.4 and 2.1 ± 0.6 at %.

Table 2. Chemical compositions of as-deposited coatings.

Sample	Cr at %	W at %	N at %	O at %
C1	35.9 ± 2.3	21.8 ± 5.0	40.2 ± 4.8	2.1 ± 0.6
C2	26.4 ± 1.4	31.9 ± 6.1	40.3 ± 4.5	1.4 ± 0.5
C3	13.9 ± 0.7	42.1 ± 6.9	43.1 ± 4.4	0.9 ± 0.4

Figure 1 shows the XRD patterns of the as-deposited coatings. The main diffraction peaks were observed at diffraction angles of 37.21° , 43.27° , 62.85° , 75.64° and 78.73° , corresponding to the (111), (200), (220), (311) and (222) planes of both the c-CrN and c-W₂N phases [22]. The value of these diffraction peaks slightly shifted to lower angles with the increase of W content. This change likely originated from lattice expansion due to the solid solution of W atoms [20,23,24]. In addition, the coatings exhibited a (111) preferential orientation, although the intensity of the strong (111) peak was considerably lower as the W content is increased from 21.8 ± 5.0 at.% to 42.1 ± 6.9 at.%, The results are in accordance with those of previous studies, indicating that the doping of a small amount of W to CrN can induce an evolution of the texture from a preferential orientation of (111) to one of (200) [20]. Moreover, the weak (222) peak is very wide and close to the diffraction angle of (311) peak, leading to the asymmetry of the (311) peak.

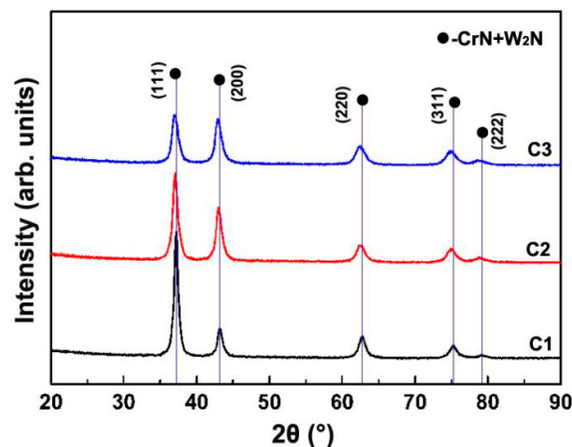


Figure 1. XRD patterns of as-deposited coatings.

Figure 2 shows the XPS spectra of Cr2p, W4f, N1s and O1s in sample C2. As shown in Figure 2a, the binding energies of the Cr2p_{3/2} and Cr2p_{1/2} peaks centered at 574.7 eV and 583.4 eV, which correspond to the Cr-N state [25,26]. Meanwhile, the binding energies of Cr2p_{3/2} and Cr2p_{1/2} peaks at 576.6 and 586.3 eV correspond to the Cr-O states [27]. The binding energies of W4f_{7/2} and W4f_{5/2} peaks located at 31.8 eV and 33.9 eV (Figure. 2b), which are associated with W-N binding state [28,29]. The binding energies of W4f_{7/2} and W4f_{5/2} peaks at 35.6 eV and 37.8 eV are assigned to the W-O state [30]. Fig 2(c) displays that the binding energies of N1s peaked at 396.9 eV, 397.7 eV and 399.8 eV, which can be assign to Cr-N and W-N binding states, respectively [25,28]. The O1s spectrum in Fig. 2(d) shows that the binding energies of the O1s peaked at 530.1, 530.7 and 531.8 eV, which can be assign to Cr-O, W-O and H-O states, respectively [31–33].

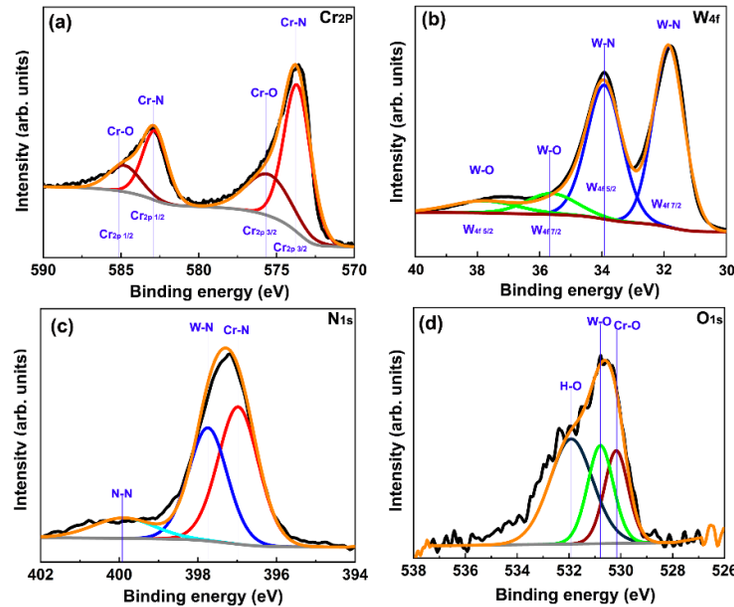


Figure 2. XPS spectra of sample C2: (a) Cr2p, (b) W4f, (c) N1s, (d) O1s.

Figure 3 shows the surface images of the as-deposited coatings. Sample C1 exhibited a granular morphology, as shown in Figure 3a, plenty of micropores were distributed around grain boundaries, indicating a loose and coarse growth structure. The surface features changed considerably with an increase of the W content. As shown in Figure 3b, sample C2 exhibited a dense surface morphology, plenty of nm-sized grains were closely packed in large cluster particles. Sample C3 also showed a dense surface (Figure 3c). There were plenty of cluster particles consisting of numerous fine grains, but the sizes of such cluster particles decreased considerably in comparison to sample C2.

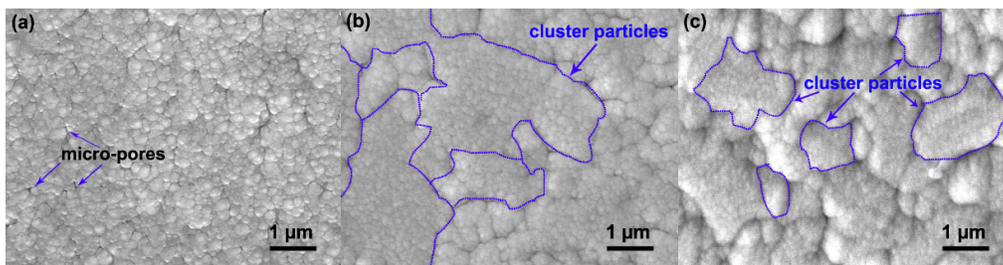


Figure 3. Surface SEM images of as-deposited coatings: (a) C1, (b) C2, (c) C3.

Figure 4 shows the cross-section TEM images of sample C2. The dark field TEM image in Figure 4a displayed a distinct columnar grain morphology. Moreover, a polycrystalline cubic structure with (111), (200), (220), and (311) reflections was identified according to the selected area diffraction pattern

(SADP). The HAADF STEM image in Figure 4b showed an obvious two-layered structure with alternate bright (W_2N) and dark (CrN) contrast according to the element mapping image. These multilayer structure mainly originated from the rotation of sample holder during the coating deposition. Meanwhile, these nano-multilayers exhibited a typical coherent epitaxial growth mode (Figure 4c). The excellent coherence relations were clearly kept between W_2N and CrN phases, as shown in Figure 4d, the d-spacing values were 0.145 nm for the CrN phase and 0.146 nm for the W_2N phase. These similar d-spacing values were assigned to the (220) planes. Additionally, plenty of W atoms were dissolved in the CrN matrix, and some Cr atoms were also identified in the W_2N matrix. Both of which indicated that a significant solid solution effect occurred during the coating deposition.

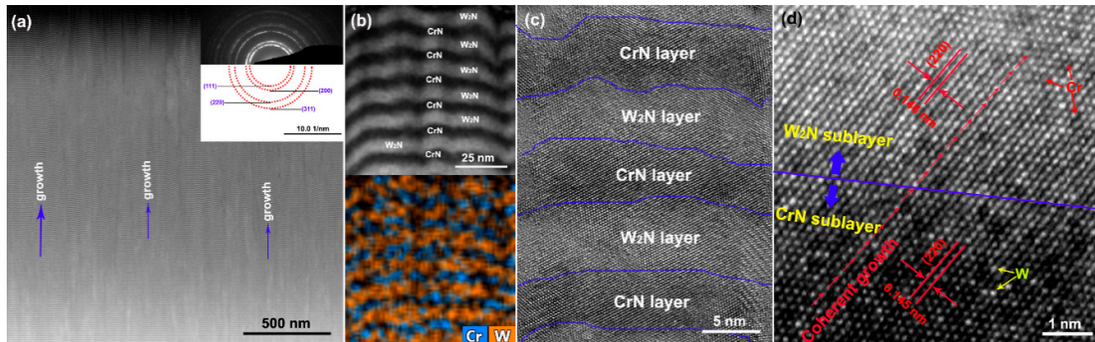


Figure 4. Cross-section TEM images of sample C2: (a) TEM image with corresponding SADP, (b) HAADF STEM image and element mapping, (c) and (d) HRTEM images.

Figure 5 shows the AFM images of the as-deposited coatings. Here sample C1 showed a relatively rough surface with a high Ra value of 10.841 nm (Figure 5a). The roughness value decreased considerably with the increase in W content. Figure 5b shows a smooth surface for sample C2, having a low Ra value of 2.417 nm. Figure 5c indicates a similar Ra value for sample C3 (2.698 nm). Based on the XRD, SEM, TEM, and AFM results, we can infer that the CrWN coating has an improved surface quality, although it undergoes aggravated lattice expansion under increasing of W content.

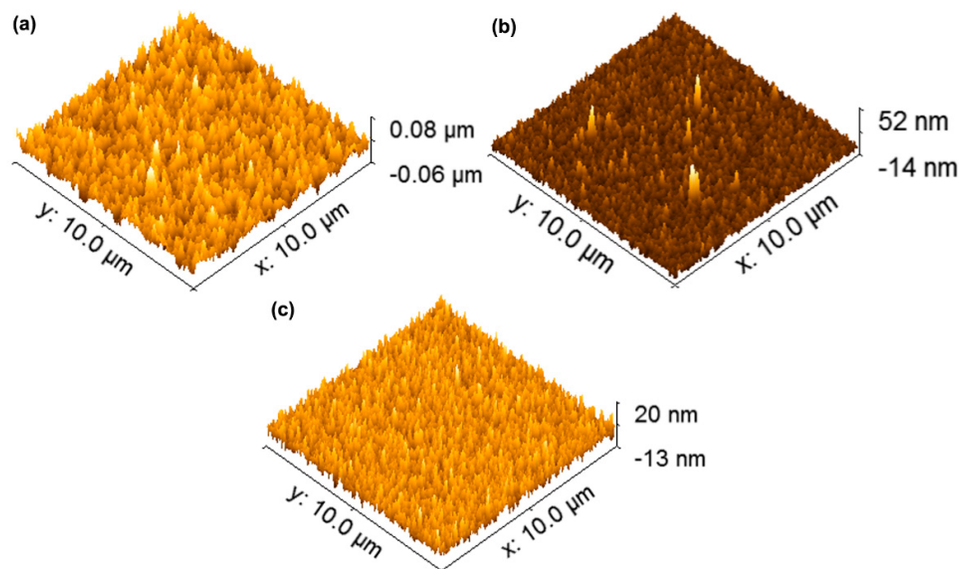


Figure 5. AFM images of as-deposited coatings: (a) C1, (b) C2, (c) C3.

Figure 6 shows the XRD patterns of the vacuum annealed coatings. The annealed samples C1 and C2 exhibited similar phase structures to those of the as-deposited coatings. Strong diffraction

peaks of mixed c-CrN and c-W₂N phases can be seen in the XRD patterns. Simultaneously, weak peaks of h-WN phases can be seen in these annealed coatings, implying a slight phase decomposition. However, the phase decomposition is aggravated significantly with the increase of W content. Apart from the original diffraction peaks of the mixed c-CrN and c-W₂N phases, the diffraction intensity of the h-WN peak was found to increase significantly in the annealed sample C3, indicating the formation of a large volume fraction of h-WN phases.

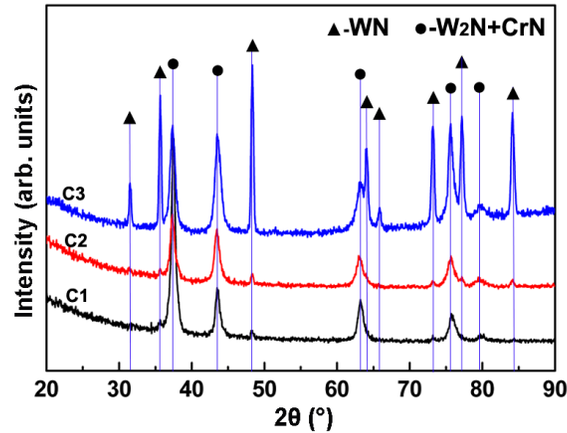


Figure 6. XRD patterns of vacuum annealed coatings.

Figure 7 shows the XRD patterns and cross-section TEM images of as-deposited and annealed sample C2. Compared with the as-deposited coating, the annealed coating remained stable structure of CrN and W₂N phases, but their diffraction peaks obviously shifted to higher angles, indicating a mitigating lattice expansion. Meanwhile, the as-deposited coating showed clearly two-layered structure with alternate bright (W₂N) and dark (CrN) contrast (Figure 7a). By contrast, the annealed coating exhibited obviously three-layered structure with bright (W₂N), gray (mixed CrN-W₂N), and dark (CrN) contrast, as shown in Figure 7b, a distinct gray diffusion layer forms between CrN and W₂N sublayers. The partial enlarged image in Figure 7b reveals that the excellent coherence relations were kept in the diffusion layer. Apparently, the vacuum annealing effectively drove the decomposition of the supersaturated solid solution resulting in the formation of a diffused solid solution matrix.

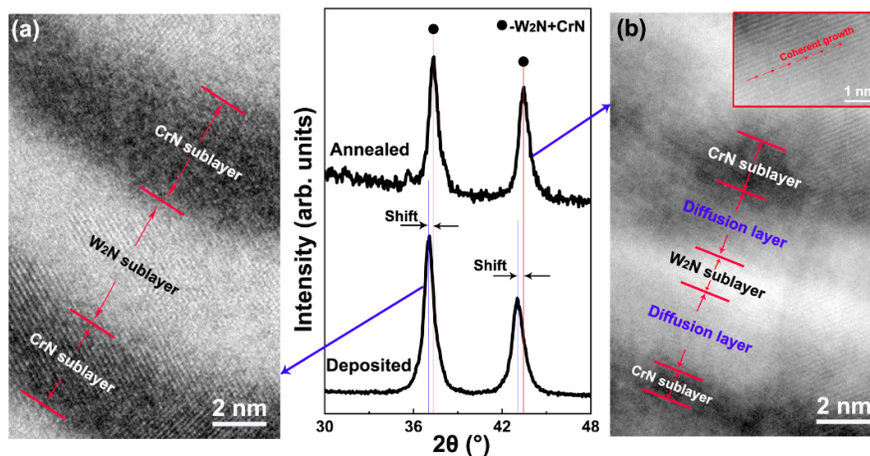


Figure 7. XRD patterns and cross-section TEM images of as-deposited (a) and annealed (b) sample C2.

Figure 8 shows the surface images and EDS spectra of the vacuum annealed coatings. As shown in Figure 8a, the annealed sample C1 exhibited a granular surface morphology with plenty of newly

grown nm particles. The O content in this coating was of 9.6 ± 1.7 at %. Meanwhile, the surface features of the annealed samples C2 and C3 were slightly different from those of the as-deposited coating (Figure 8b,c). These nm-sized grains in fact grew, leading to a visible coarsening of the cluster particles. The O contents were of 7.4 ± 1.3 at % and 12.1 ± 1.7 at % for annealed samples C2 and C3, respectively. The SEM characterizations clearly confirmed that these coatings suffered slight oxidation damages during the vacuum annealing.

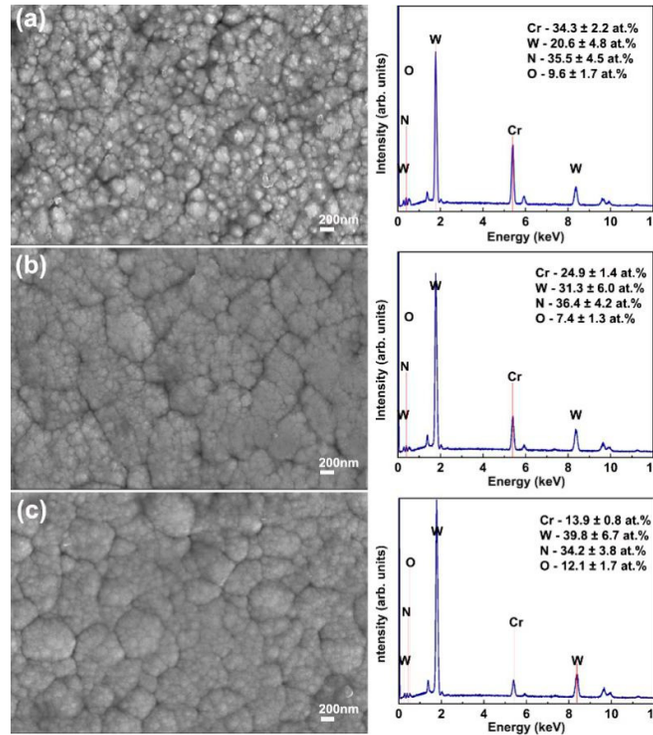


Figure 8. Surface images and EDS spectra of vacuum annealed coatings: (a) C1, (b) C2, (c) C3.

Figure 9 shows the roughness values and AFM images of the as-deposited and annealed coatings. The Ra values of samples C1 and C2 increased from 10.841 to 11.332 nm and from 2.417 to 3.204 nm, respectively; by contrast, the Ra value of sample C3 increased abruptly from 2.698 to 4.945 nm, indicating a severe surface coarsening.

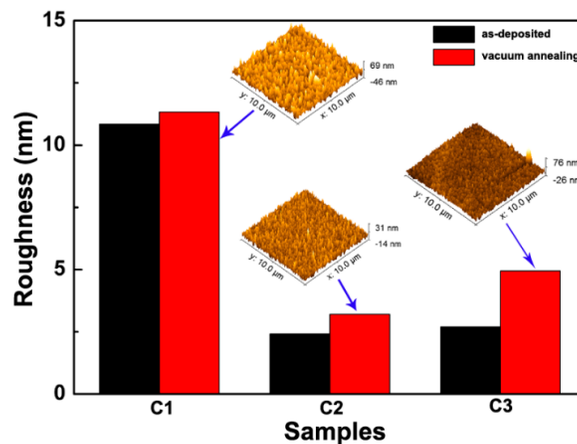


Figure 9. Roughness values and AFM images of as-deposited and annealed coatings: (a) C1, (b) C2, (c) C3.

Based on the SEM and AFM characterizations, we found that the annealed coatings underwent various degrees of surface coarsening, which was likely triggered by surface oxidation. The EDS results further showed the occurrence of slight oxidation erosion during vacuum annealing, which consequently resulted in the formation of Cr–W oxides [20]. Although these oxides are too small to be detected by XRD in Figure 6, but they led to severe surface coarsening because of their loose structure [20,34]. The significant differences in Ra value between the annealed coatings having different W content can be likely attributed to the varying composition of the oxide layers. According to the EDS results (Figure 8), the volume fraction of Cr oxides decrease considerably and was replaced by a rising volume fraction of W oxides under increasing W content. Previously, it has been reported that dense Cr oxides can act as protective layers and inhibit O diffusion into the coating [35], whereas the W oxides usually exhibit a more porous structure [34]. Therefore, an increase in W oxides can eventually lead to a constant increase of the Ra values.

Figure 10 shows the hardness of the as-deposited and annealed coatings. The as-deposited coatings exhibited a constant hardness enhancement under increasing W content. The hardness was 12.9 GPa for sample C1, and slightly increased to 13.7 GPa for sample C2. By contrast, sample C3 showed a higher hardness of 15.5 GPa. The vacuum annealing induced a remarkable age-hardening in sample C1 and C2. The annealed sample C1 showed a slight increase in hardness (from 12.9 to 14.8 GPa), whereas that of the annealed sample C2 increased rapidly (from 13.7 to 21.6 GPa). Meanwhile, the annealed sample C3 underwent a serious hardness degradation (from 15.5 to 10.1 GPa).

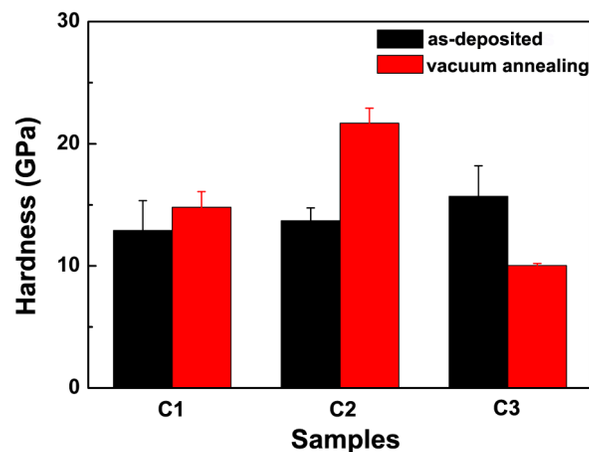


Figure 10. Hardness of as-deposited and annealed coatings.

Based on the XRD and TEM characterizations, it can be inferred that the sputtered atoms or ions triggered an obvious injection effect in the as-deposited coatings, as shown in Figure 4d, a significant solid solution effect appeared in both W_2N and CrN sublayers. According to the previous results [20,23,24], these solid solution atoms induced serious lattice expansion resulting in a visible peak shift as determined from the XRD test (Figure 1). Meanwhile, this lattice expansion was significantly aggravated following an increase in the W content, which provided an obvious strengthening effect and eventually led to a constant hardness enhancement. Moreover, the addition of W to CrN led to an obvious reduction of the particle (i.e., grains or clusters) sizes (Figure 3) and refined their structure, causing a hardness improvement. The annealed coatings with different W contents exhibited significant differences in phase composition, which consequently had a strong effect on their mechanical properties. As determined from the nanoindentation test, the annealing sample C1 and C2 showed a prominent age-hardening, which mainly originated from their microstructure evolution. As identified in Figure 4d and Figure 7a, these doping atoms induced the formation of supersaturated solid solution in the as-deposited coating, but the supersaturated matrix exhibited a non-uniform distribution resulting in limit strengthening effect. By contrast, vacuum annealing induced spinodal decomposition of supersaturated solid solution to form nm-sized c-CrN, c- W_2N ,

and h-WN domains (see Figure 6). Although a small amount of h-WN phases was formed, but the annealed coating showed similar face-centred cubic (f.c.c) structure to that of as-deposited coating. According to the previous age-hardening theories [36–39], strain fields, originating from the lattice mismatch, acted as obstacles for the dislocation movement and caused hardness enhancement. Additionally, thermal activation greatly driven the homogenization of the supersaturated matrix, as evidenced in Figure 7b, a distinct diffusion layer formed between CrN and W₂N sublayers. In comparison to the limited strengthening effect in as-deposited coating, the coherent solid solution diffusion matrix provided strong strengthening effect due to the larger mismatch, and consequently resulted in a profound age-hardening in sample C1 and C2. Nevertheless, the annealed sample C3 underwent serious spinodal decomposition accompanied by the formation of a mixed f.c.c and hexagonal close-packed (h.c.p) matrix structure. The large volume fraction of h-WN phases weakened considerably strengthening effect of the coherent interface and eventually led to a drop of the hardness value.

4. Conclusions

This study investigated the microstructure and thermal stability of CrWN glass molding coatings having different W contents after vacuum annealing. The main conclusions are as follows:

- The as-deposited coatings showed columnar structures consisting of multilayer c-CrN and c-W₂N phases. The preferential orientation of these structures varied from (111) to (200), and the surfaces became smoother under increasing W content.
- The as-deposited coatings suffered from aggravated lattice expansion. The grain size and cluster particle sizes showed an obvious decrease under increasing W content. These changes led to a constant hardness enhancement.
- The annealed coatings underwent minor oxidation and varying degrees of surface coarsening. The roughness value gradually increased due to an increasing volume fraction of W oxides.
- The annealed coatings experienced spinodal decomposition while forming nm-sized c-CrN, c-W₂N, and h-WN domains. The volume fraction of the h-WN phase increased significantly with the increase of W content.
- The annealed coating under low W content exhibited a significant age-hardening due to the strain fields originating from the lattice mismatch. The annealed coating under high W content suffered from a severe hardness degradation because the large volume fraction of h-WN seriously weakened the interface strengthening effect.

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