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Exploring the tribological behavior of Ti/Al-DLC/PAO/graphene oxide nanocomposite system

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ABSTRACT

The role of graphene oxide (GO) as an oil-based additive on the tribological behavior of Ti/Al-DLC films and its dependence on GO concentration were systematically investigated. The results revealed that, compared to the additive-free case, the system with a GO concentration of 0.003 wt% achieved remarkable friction reduction and wear resistance, which were decreased by 38.4% and 23.5%, respectively. This is attributed to the synergistic effect of tribo-induced graphitization and the adsorption of GO on the worn surface. However, when the GO concentration was further increased, it induced the clumping of the GO additive, causing increased friction coefficient and severe abrasive wear of the worn surface.

1. Introduction

In modern industry, the moving assemblies used in aerospace, aviation, military, and other engineering fields are inevitably subject to friction and wear, resulting in energy dissipation and material losses [1-4]. Therefore, it is important to minimize friction and reduce wear, thus improving machine efficiency. However, owing to the complex and changeable working environment, single-liquid lubricants cannot satisfy the rigorous requirement of mechanical components for reducing friction and extending the machine service lifetime. Many previous studies [5-8] have confirmed the solid-liquid lubricating system, which combines solid lubricating films/coatings with fluid lubricants to be an effective method of reducing friction. In comparison with the traditional hard anti-wear coating and soft lubricating protective coating, a diamond-like carbon (DLC) film, exhibits superior comprehensive performance as a high-performance solid lubricating film [9-11]. In particular, co-doping with binary metals into DLC film can further endow the film with enhanced mechanical and tribological performances [12–15].

As oil-based lubricant additives for liquid lubricants, graphene and its derivatives have attracted attention because of their ultrathin 2D laminated structure, superior load-bearing capacity, as well as excellent friction reduction and anti-wear capabilities [16-18]. Zhang et al. [19] reported that the lubricating oil with oleic acid-modified graphene (0.02.0.06 wt%) exhibited 17% and 14% reduction in friction coefficient (COF) and wear scar diameter (WSD), respectively. Eswaraiah et al. [20] reported a type of engine lubricant oil with a graphene nanoadditive which showed a decrease of 80% in COF and a decrease of over 30% in wear rate. Similarly, Wu et al. [21] evaluated the tribological behavior of Si₃N₄ ceramic/GCr15 sliding pairs lubricated by 4010 AL base oil containing graphene oxide (GO) nanosheets and found that as the GO concentration was 0.5 wt%, the tribopairs exhibited excellent antifriction and wear-resistant properties owing to the formation of a protective tribofilm, which adhered to the worn surface to prevent the direct contact of tribopairs. Moreover, Zhang et al. [22] illustrated that multilayer GO, which functions as a Mobil DTE oil additive, can reduce

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Fig. 1. (a) Digital photographs of GO-dispersed PAO oils with different GO concentrations after aging for 5 h. Laser particle-size distributions of (b) PAO-GO1, (c) PAO-GO2, (d) PAO-GO3, and (e) PAO-GO4 oils.

the average COF by 67.37%.

Although a considerable number of studies have been focused on the tribological properties of steel/steel or ceramic/steel pairs lubricated with graphene and its derivatives as oil-based additives, the effects of graphene-based additives on metal-doped DLC/steel pairs remain unclear. In previous studies [23–26], we have fabricated the Ti/Al co-doped DLC (Ti/Al-DLC) film using a hybrid ion beam (HIB) deposition system, which showed superior antifriction and wear-resistant behavior under dry conditions. Therefore, in the present work, we further evaluate the tribological behavior of Ti/Al-DLC film composited with poly-alpha olefin (PAO) as the base oil and GO as the additive. We focused on its dependence on the GO concentration and discussed the underlying friction mechanism with respect to the structural analysis of the friction interface.

2. Experimental details

2.1. Liquid lubricant

The liquid lubricant consisted of PAO oil as the base oil and GO as the lubricant additive; high-purity GO was purchased from Nanjing Jicang Nano Technology Co., Ltd., and its thickness was approximately 1–2 nm. The PAO oil, provided by Shell China Ltd (Shanghai), was mainly composed of hydrocarbons. To prepare the blended lubricants, GO powder was first added to the PAO-based oil and then subjected to ultrasonic dispersion for 180 min. The GO concentration ranged from 0 wt %, 0.003 wt%, 0.006 wt%, 0.012 wt%, to 0.018 wt%, and the corresponding lubricants were abbreviated as Pure oil, PAO-GO1, PAO-GO2, PAO-GO3, and PAO-GO4, respectively.

2.2. Film deposition

Ti/Al-DLC films were deposited on a silicon P (100) wafer using the HIB technique, which was composed of a linear anode-layer ion source (ALIS) supplied with C_2H_2 gas for DLC deposition and a direct current magnetron sputtering (DCMS) device provided with a $Ti_{50}Al_{50}$ (99.99%) target. Prior to deposition, the substrates were ultrasonically cleaned in ethyl alcohol and then etched in Ar^+ bombardment for 20 min to remove the adhered contamination. During deposition, a Ti/Al interlayer was first deposited on the substrate under conditions of 55 sccm Ar, a sputtering current of 4 A, bias voltage of -100 V, and deposition time of 16 min [25]. After that, 70 sccm Ar and 10 sccm C_2H_2 were supplied to



Fig. 2. (a) TEM micrographs, (b) HR-TEM lattice plane and corresponding SAED diagram of GO, (c) AFM image, and (d) corresponding cross-sectional profile of 2D AFM image of as-prepared GO additive.

DCMS and ALIS simultaneously to deposit the Ti/Al-DLC film; the ALIS current, sputtering current, negative-bias voltage, and deposition time were 0.2 A, 2.5 A, -50 V, and 60 min, respectively. Details of the deposition parameters can be obtained in our previous studies [23,27].

2.3. Microstructure characterization

The chemical structures of GO, PAO, and PAO-GO were characterized using Fourier transform infrared spectra (FTIR, NICOLET 6700) with the wave number ranging from 400 to 4000 cm⁻¹. The morphology and thickness of GO nanoplates were obtained by the high-resolution transmission electron microscopy (FEI Tecnai F20) and scanning probe microscope (Veeco Dimension 3100 V, SPM). The particle-size distribution of each oil sample was tested by the dynamic light scattering particle size analyzer (Zetasizer Nano ZS, ZETA). Furthermore, the profile of wear tracks was recorded using a surface profilometer (Alphastep IQ, US) through a linear scanning mode. The field emission scanning electron microscopy (SEM, FEI Quanta FEG 250) combined with the X-ray energy dispersive spectroscopy (EDS) was utilized to analyze the surface morphologies of wear scar on the contact ball and the wear track on the film, respectively. The carbon bond structure of wear track was also investigated by Raman spectroscopy (Renishaw, inVia-reflex) with the laser wavelength of 532 nm.

2.4. Tribological testing

The tribological properties of the Ti/Al-DLC/PAO/GO composite system were evaluated using a pin-on-disk tribometer (Anton Paar, TRB3) at room temperature (20 \pm 5 °C). A grinding GCr15 steel ball (HRC60) with a diameter of 6 mm was selected as the mating material. Each test was undertaken at a rotating speed of 320 rpm, an applied load of 5 N, a radius of 1.5 mm, and sliding distance of 4500 laps. In addition, each sample test was repeated thrice to calculate the average COF (μ) and wear rate (W). After the tests, the surfaces of the counterpart and film were ultrasonically cleaned with acetone. The wear rate of each sample was calculated using the Archard equation:

$$W = V / (L \times N) \tag{1}$$

where *L* and *N* were the total sliding distance and normal load separately; *V* corresponded to the wear-induced volume loss, which was calculated by the integral of cross-sectional profile of wear track. While the wear rate of mating ball was calculated as follows [28]:

$$V = \left(\frac{\pi h}{6}\right) \left(\frac{3D^2}{4} + h^2\right),\tag{2}$$

$$h = R - \sqrt{R^2 - \frac{D^2}{4}},$$
 (3)

where h was the height of wear material of the grinding ball; D and R



Fig. 3. (a) Raman spectrum of as-prepared GO; (b) FTIR spectra of GO, PAO, and PAO-GO, respectively.

were the diameter of wear scar and the radius of mating ball, respectively.

3. Results and discussion

3.1. Stability of GO-dispersed oil

GO-containing oil was prepared through a physical method without any extra dispersing agent. In order to confirm the stability of GO in PAO oil, Fig. 1(a) depicts the digital photographs of all GO-dispersed PAO oils with different GO concentrations after aging for 5 h. For comparison, pure PAO oil is also provided. Notably, there is no obvious sedimentation and separation of each observed oil sample, illustrating that the GO exhibits good dispersity and stability in the PAO oil. Fig. 1(b)–1(e) further display the laser particle-size distribution of each sample. It can be seen that as the GO concentration increases from 0.003 wt% to 0.018 wt%, the average particle size of GO increases from 21.26 to 45.87 nm. On the one hand, this proves that the GO can still remain at nanoscale sizes and good dispersion in PAO oil. On the other hand, the wide-laser particle-size distribution of GO indicates that GO mainly presents a particle-aggregation state rather than a monodisperse state in PAObased lubricant oil.

3.2. Structure and morphology of GO additive

Fig. 2 demonstrates the structure and morphology of as-prepared GO. In Fig. 2(a), numerous wrinkles and curls exist in the GO nanoplates, which are mainly related to the degree of oxidation of GO and the surface energy [29], suggesting the existence of structural defects. The HRTEM lattice micrograph and corresponding selected area electron

diffraction (SAED) diagram of GO are depicted in Fig. 2(b). Notably, the thickness of the GO nanoplates is approximately 2 nm, indicating the multilayer structure of GO [30]. In addition, the sharp crystalline diffraction rings are identified as (222) and (600) reflections of GO. Fig. 2(c) and (d) show the AFM morphology and cross-sectional profile, in which the white horizontal line is illustrated in 2D AFM morphology. It can be seen that the average thickness of GO comes out to be 1.89 nm, which is consistent with the TEM result.

Fig. 3(a) shows the Raman spectra of GO additive with a wavelength range of 1000–2000 $\rm cm^{-1}$. It can be fitted into two prominent peaks: one is the D band at 1359 cm^{-1} corresponding to disordered sp³-C caused by the structural defects of GO; the other is the G band located at 1591 cm⁻¹, which is associated with the in-plane vibration [31]. In addition, the intensity ratio of D peak to G peak (I_D/I_G) is closely related to the disorder degree of graphite materials [22,32]. In this case, the I_D/I_G ratio in GO is approximately 1.18, implying the structural defects of GO caused by the oxidation of functional groups. This is consistent with the TEM analysis. To further explore the structure of the functional group in GO and its interaction with PAO oil molecules, the FT-IR characterization is conducted, as shown in Fig. 3(b). For the GO, the absorption bands located at 1105 cm⁻¹ and 1733 cm⁻¹ are assigned to C-OH and C=O stretching vibrations of hydroxyl and carboxyl groups, respectively; the band at 1625 cm⁻¹ is attributed to the unoxidized sp² C=C stretching vibration, whereas the bands at 3415 cm⁻¹ and 1397 cm⁻¹ originate from the stretching and bending vibrations of -OH, respectively [33]. For the pure PAO oil, the strong absorption bands in the region of 2855 cm⁻¹ to 2970 cm⁻¹ correspond to the stretching vibration of $-CH_2$ in alkyl chains; the bands at 1360 cm⁻¹ and 1466 cm⁻¹ are related to the vibrations of C-H group in PAO oil molecules [34-36]. After dispersing GO in PAO oil, each sample also shows the same peaks



Fig. 4. (a) Friction curves and (b) average COF values of Ti/Al-DLC films lubricated by pure and GO-dispersed PAO oils, respectively.



Fig. 5. Wear rates of Ti/Al-DLC films lubricated by pure and GO-dispersed PAO oils, respectively.

as those in pure PAO oil; however, no absorption peaks of GO can be observed. This phenomenon occurs mainly because when the graphene oxide content is low, the stretching vibration of $-CH_2$ in alkyl chains of the PAO oil has a strong absorption peak (Fig. 3(b)), which shows a shielding effect on the characteristic peaks of graphene oxide.

3.3. Tribological performance of Ti/Al-DLC/PAO/GO system

Fig. 4(a) displays the characteristic friction curves of the system lubricated by pure or GO-dispersed PAO oils. From the figure, compared to pure PAO, adding GO additive into oil could further reduce the COF value, suggesting the positive role of GO additive in the frictionreducing capacity of PAO oil. Furthermore, this change is strongly sensitive to the GO concentration. The average COF during the steadystate friction stage was calculated for each case, as illustrated in Fig. 4 (b). Compared to the pure PAO oil (0.125), when the concentration of GO additive in PAO oil is 0.003 wt%, the minimal COF value of 0.077 is obtained, which is reduced by 38.4%. However, with further increase in the GO concentration to 0.018 wt%, the COF values increases gradually, but it remains lower than that of the pure PAO oil, confirming the synergistic effect of GO additive and PAO oil. Meanwhile, the system with PAO-GO4 undergoes a long running-in process following an acute fluctuation owing to serious interaction between the GO sheets during the sliding process, as will be discussed later.

The wear rates of Ti/Al-DLC film lubricated by pure or GO-dispersed oils are given in Fig. 5. As can be seen that following the increase of GO concentration, the wear rate of composite system first shows a decreasing and then increasing trend, which coincides with the change of COF in Fig. 4(b). When the concentration of GO additive reached 0.003 wt% (PAO-GO1), the system exhibits a superior antiwear behavior and the wear rate is only approximately 2.05×10^{-16} m³/Nm, which is approximately 23.5% lower than that lubricated by pure PAO oil. As the concentration of GO additive further increase to 0.006 wt% and 0.012 wt%, the wear rate conversely increases by 20.5% and 42.9%, respectively. However, when the concentration of GO additive was 0.018 wt% (PAO-GO4), the wear rate could not be calculated owing to the partial exfoliation of the Ti/Al-DLC film.

Moreover, the wear scars on each mating ball were analyzed, including the SEM image of the wear scar for each case and the corresponding WSD and wear rate, as illustrated in Figs. 6 and 7, respectively. Under the pure PAO oil, the wear scar on the steel ball is almost round in shape, and there exist some wide and deep scratches (Fig. 6(a)), indicating that abrasive wear occurs during the rubbing process. Compared with Fig. 6(a), the wear scar lubricated by PAO-GO1 shows a smaller wear scar (Fig. 6(b)) and a smoother worn surface (Fig. 7(a)). The wear rate of the mating ball also reaches the lowest value of 1.16×10^{-5} mm⁻³ (Fig. 7(b)), suggesting that excellent wear resistance was achieved when the GO concentration was 0.003 wt%. However, the WSD and wear rate of counterparts display continuously increasing



Fig. 6. SEM morphologies of wear scars on the mating balls lubricated by (a) pure PAO oil, (b) PAO-GO1, (c) PAO-GO2, (d) PAO-GO3, and (e) PAO-GO4, respectively.



Fig. 7. (a) WSD and (b) wear rate of the mating balls lubricated by pure PAO oil and GO-dispersed oil, respectively.



Fig. 8. Wear tracks and their enlarged views of Ti/Al-DLC films lubricated by (a)(f) pure oil (b)(g) PAO-GO1 (c)(h) PAO-GO2 (d)(i) PAO-GO3, and (e)(j) PAO-GO4, respectively, after sliding processes.



Fig. 9. AFM semicontact images of the wear tracks of Ti/Al-DLC films lubricated by (a) Pure oil, (b) PAO-GO1, (c) PAO-GO2, (d)PAO-GO3, and (e) PAO-GO4, respectively, after sliding processes.



Fig. 10. (a-b) SEM micrographs and (c) EDS spectra of wear track for Ti/Al-DLC film lubricated by 0.018 wt% of GO composite oil.

trends as the GO concentration in PAO oil further increases to 0.018 wt% (Fig. 7) following the presence of many deep ploughs in the wear scars (Fig. 6). These results agree well with the wear rate of the film, as illustrated in Fig. 5.

3.4. Tribological mechanism

Because no obvious transfer layer is observed in the mating balls (Fig. 6), we focus on the structural transformation of the wear track on the film to clarify the underlying friction and wear mechanisms induced



Fig. 11. (a) Raman spectra and (b) fitting results of the wear tracks of Ti/Al-DLC films lubricated by pure and GO-dispersed PAO oils, respectively.



Fig. 12. Raman mappings of selected worn surfaces of films lubricated by (a) pure PAO oil, (b) PAO-GO1, and (c) PAO-GO4, respectively, in which color represents the different I_D/I_G values.

by different GO concentrations. Fig. 8 presents the wear profiles and their corresponding enlarged morphologies of Ti/Al-DLC films lubricated by pure and GO-dispersed PAO oils, respectively, after the sliding processes. For the pure oil case (Fig. 8(a) and (f)), the width of the wear track is 122.4 μ m and some shallow furrows exist inside the wear track, accompanied by the plastic deformation of the worn surface [37], and the surface roughness of the wear track is approximately 2.08 nm (Fig. 9 (a)). As the GO additive is dispersed in PAO oil, the width and roughness of the wear track as a function of GO concentration first decrease and then increase. When lubricated by PAO-GO1, the width of the wear track reaches a minimum value of 119.6 μ m (Fig. 8(b)); the worn surface is shallow and has the lowest surface roughness value (Ra = 0.92 nm Fig. 9(b)) when compared to other lubricating conditions (Fig. 9). Besides, it can be noticed that some black particles adhere to the worn surface (Fig. 8(g)), which is related to the adsorption of GO additives and also indicates the presence of adhesive wear during the sliding process.

However, with further increase in the GO concentration to 0.006 wt % (PAO-GO2) and 0.012 wt% (PAO-GO3), both the width of wear track and dark particles inside the wear trace increase (Fig. 8 (c), (d), (h), and (i)), but some scratches and traces of deformation are visible on the

surface, suggesting extra contribution from abrasive wear. In particular, there are obvious white bumps observed in Fig. 9(c) and (d), which originate from the aggregation of GO additives in the wear track [37]. This is consistent with the SEM analysis (Fig. 8). Furthermore, when the GO concentration reaches 0.018 wt% (Fig. 8(e) and (j)), partial delamination and exfoliation occur inside the wear track. It arises from the excess GO additive, causing aggregation and bonding between the GO sheets during the sliding process. This increases the interfacial roughness (2.35 nm, Fig. 9(e)), cross-links the mated surfaces, blocks the hydrodynamic lubrication of PAO oil, and hinders the sliding of tribopairs [38]. However, compared to the wear rate, the COF remains low without a significant increase (Fig. 4(b)). Therefore, we further explored the morphology and composition of the worn surface, as illustrated in Fig. 10. As shown in the enlarged view of the worn surface (Fig. 10(b)), the dark regions are noticeable on the rubbing surface. Further EDS analysis (Fig. 10(c)) reveals that compared to the as-deposited surface (spectrum 1), these dark areas (spectrums 2 and 3) exhibit an increased amount of carbon on the worn surface, which are attributed to the GO additives adsorbed to the surface [39,40]. Moreover, the adsorption of oil molecules in the worn region of the DLC surface plays a crucial role in



Fig. 13. Schematic of the GO-induced lubricating mechanism.

lowering the COF value and reducing the risk of DLC self-consumption in practical applications.

Fig. 11(a) shows the Raman spectra of the wear tracks of Ti/Al-DLC films lubricated by pure and GO-dispersed PAO oils. The quantitative fitting results are shown in Fig. 11(b). As the GO concentration increases from 0 to 0.018 wt%, the contribution of GO to the spectrum gradually increases, as evidenced by the increased D band intensity (Fig. 11(a)). However, the intensity ratio of the D band to G band as a function of GO concentration shows slight change. In addition, the blue shift of the G band is correlative with the average size of sp² domains. This is because the GO easily tears under Hertz contact stress, resulting in disordered GO with a significantly small size [21,41]. These results confirm that the carbon content is increased, as measured with the EDS analysis (Fig. 10 (c)).

Raman mappings inside the worn surfaces with sizes of $20 \ \mu m \times 20 \ \mu m$ were further conducted, as shown in Fig. 12, to disclose the effect of the GO additive on the friction of the Ti/Al-DLC film. The color bar corresponds to different I_D/I_G values at the selected worn area. As can be observed, compared to lubrication by pure PAO oil only (Fig. 12(a)), when the GO concentration in PAO oil is 0.003 wt%, a high I_D/I_G value is obtained, suggesting the enhanced graphitization of the rubbing surface (Fig. 12(b)), which contributes to the antifriction behavior of the film. However, when GO concentration increases to 0.018 wt% (Fig. 12(c)), the excessively high concentration of GO can in turn suppress the triboinduced graphitization owing to the agglomeration of GO additive (Fig. 8) and the increased surface roughness (Fig. 9).

According to the aforementioned analysis, the GO-induced lubricating mechanism is illustrated in Fig. 13. When lubricated by pure PAO oil, direct contact between the GCr15 steel ball and Ti/Al-DLC film generates a high Hertz contact stress, causing the rapid collapse of the oil film. However, the addition of GO additives in PAO oil endowed the film with a sudden decrease in both the friction and wear rate owing to the interfacial graphitization (Fig. 12) and the adsorption of GO nanoplates on contacting surface (Fig. 8). Therefore, the roughness of the rubbing surface was reduced (Fig. 9(e)), smoothing the sliding interface and preventing the cold welding of the mated surfaces, which is also favorable for the hydrodynamic lubrication of PAO oil. However, at high GO concentrations, the homogeneous distribution of GO-dispersed PAO oil was disrupted, and the excess GO additive promoted serious agglomeration (Fig. 9). This caused the increased roughness (Fig. 7) and cross-linking of mating surfaces, aggravating the sliding resistance of both the mating surfaces and PAO oil.

4. Conclusion

In this work, Ti/Al-DLC films were fabricated on silicon substrates using an HIB system, and the effect of the concentration of GO as an additive on the tribological behavior of Ti/Al-DLC films under oil lubrication conditions was systematically explored. The results revealed that GO, as an additive of lubricating oil, played a significant role in enhancing the tribological performance of Ti/Al-DLC films. Compared to the case with pure PAO oil, when GO with concentration of 0.003 wt% was formulated into oil, the system achieved excellent friction reduction and wear resistance performances, decreased by 38.4% and 23.5%, respectively. However, when the GO concentration was further increased, it resulted in an increase in both the COF and wear rate. Microstructure analysis of the wear track indicated that at a low concentration (0.003 wt%), the GO could form a discontinuous and thin tribofilm to smooth the friction interface, and thus, prevent the direct contact between the friction pairs. Combined with the triboinduced graphitization, the shearing strength of the sliding interface was reduced, accounting for the improved antifriction and wear-resistance behaviors. However, at high GO concentrations such as 0.018 wt%, the layered structure of GO was not retained owing to the serious agglomeration, which aggravated the interfacial roughness and cross-linking of mated surfaces. Additionally, the graphitization of friction interface was weakened, leading to the increased shearing barrier following the severe abrasive wear and high friction coefficient.

Declaration of competing interest

We declare that we have no financial and personal relationships with other people or organizations that can inappropriately influence our work, there is no professional or other personal interest of any nature or kind in any product, service and/or company that could be construed as influencing the position presented in, or the review of, the manuscript entitled.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ceramint.2020.12.228.

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