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Fabrication of PTFE/Nomex fabric/phenolic composites using a layer-by-layer self-assembly method for tribology field application

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Abstract: Fabric composites are widely applied as self-lubricating liner for radial spherical plain bearings owing to their excellent mechanical and tribological properties. Nevertheless, the poor interfacial strength between fibers and the resin matrix limits the performance of composites utilized as tribo-materials. To overcome this drawback, a mild layer-by-layer (LbL) self-assembly method was successfully used to construct hybrid fabric composites in the present work. In addition, this investigation addressed the effect of self-assembly cycles on the friction and wear behaviors of hybrid fabric composites under dry sliding condition. The results demonstrate that fabric composites with three or more self-assembly cycles have significantly enhanced surface activities and anti-wear performances. The results obtained in this work can provide guidance in the preparation of self-lubricating liner composites and highlight how the LbL self-assembly techniques could influence the properties of hybrid fabric composites.

Keywords: LbL self-assembly; hybrid fabric; wear; friction

1 Introduction

Because of high load carrying capacities, maneuverability and maintenance-free properties, self-lubricating bearings are widely applied in aerospace, trains, marine and power generation [1–3]. In general, self-lubricating bearings incorporate three components—a metal inner ring, an outer race, and a self-lubricating liner between them. The life-span and stability of self-lubricating bearings during sliding processes are significantly influenced by the self-lubricating materials that constitute the liner. Among different candidate materials, hybrid fabric/polymer composites have been extensively utilized as self-lubricating liners, owing to outstanding chemical resistance, light weights and self-lubrication properties especially under dry sliding conditions [4–6]. The self-lubricating fabrics are produced by a weaving process. PTFE and aramid fibers are usually included in a two-layer warp fabric, as they possess the high specific strength of Nomex fibers and outstanding self-lubrication properties of PTFE fibers [7–9]. Furthermore, phenolic are widely used as adhesive in such composites. The performance of composites utilized as tribo-materials is highly influenced by the efficiency of the friction stress transference between the phenolic matrix and the self-lubricating fabric. However, the fiber-phenolic resin interface adhesion strength is rather weak because of the intrinsically smooth surfaces and inert chemical structure of weaving fibers. Hence, surface modification of hybrid fabrics is vital. Numerous approaches, such as plasma treatment, chemical grafting and whiskerization,
have been utilized to change the surface properties of high-performance fibers for fabric reinforced polymer matrix composites [10−12]. Nevertheless, with the increasing demands for high-performance tribomaterials in industries, single surface modification of hybrid fabrics is no longer sufficient.

Layer-by-layer (LbL) methods have been widely applied for construction uniform films in the fields of interfacial modification, separation, drug delivery, among others [13−15]. Traditionally, LbL assembly was carried out sequentially adsorbing oppositely charged materials onto a substrate. Zhou et al. [16] introduced MgAlFe layered double hydroxide (LDH) and SiO$_2$ onto the surface of aramid by a green LbL self-assembly method, which resulted in improvements in the surface activity and UV resistance of composites. Yu et al. [17] showed that the flame retardancy of the LbL modified Ramie fabric reinforced unsaturated polyester resin composites was improved by increasing the LbL assembly cycles. This was primarily attributed to the early decomposition of the LbL coating on the fabric.

Furthermore, the incorporation of functional fillers was shown to be effective in improving the friction and wear properties of hybrid fabric liner composites [18−19]. Two-dimensional (2D) nano-sheets, consisting of single or few atomic layers, such as graphene oxide (GO), molybdenum disulfide (MoS$_2$), graphitic carbon nitride (g-C$_3$N$_4$), boron nitride, have gained considerable attention owing to their excellent performances when utilized solely or in composites form [20−22]. Among these 2D nano-materials, GO has superior physical structural properties while at the same time, possessing numerous functional groups in the edges that can provide active sites for further surface modification [23].

Therefore, the aim of this study is to improve the interfacial activity of hybrid fabric and simultaneously anchor GOs on fibers surfaces using a mild and green LbL method. Specifically, GO/PAMPA and PDDA were alternately self-assembled on the surfaces of hybrid fabrics. X-ray powder diffraction (XRD), Raman spectroscopy, X-ray photoelectron spectroscopy (XPS) and scanning electron microscopy (SEM) results are used to prove the formation of even GO coatings on PTFE and Nomex fiber surfaces using LbL self-assembly techniques and significant enhancement in the wettability of the lubricating and bonding surface. Moreover, the wear mechanisms are examined based on the characteristics of the worn surfaces of hybrid fabric composites and steel counterpart pin.

2 Experimental

2.1 Materials

PTFE and Nomex fibers were purchased from DuPont Plant. The volume ratio of PTFE to Nomex in fabric is 1:3. Poly (2-acrylamido-2-methyl-1-propane sulfonyl acid) solution (PAMPA, $M_w = 2000000$, 15 wt% in H$_2$O) was purchased from Sigma-Aldrich, while poly (diallyl ammonium Chloride) solution (PDDA, $M_w = 1000000$, 35 wt% in H$_2$O) was purchased from Meryer Chemical Technology Co., Ltd. Graphene oxide (GO) was provided by Nanjing XFNANO Materials Tech Co., Ltd, China. The adhesive resin was provided by Shanghai Xing-Guang Chemical Plant (China).

2.2 Characterizations

XRD analysis was performed on an XRD system (Philips Corp., The Netherlands), operating with Cu-K$_\alpha$ radiation at a scanning rate of 0.5° per second over a 2θ range of 10° to 80°. Raman spectroscopy (in a LabRAM HR800 system with 532 nm laser excitation), was employed to characterize the structural features of the LbL modified hybrid fabric. The surface morphology and microstructure of graphene oxide and hybrid fabric were observed using field emission scanning electron microscopy (FE-SEM, JSM-6701F, JEOL, Japan). The morphologies of the worn surfaces of the composites and their counterpart steel pins were analyzed on a JSM-5600LV scanning electron microscope (SEM). X-ray photoelectron spectroscopy (XPS) measurements were conducted on a VGESCALAB210 spectrometer. A drop-shaped analyzer (KrüssDSA100, Krüss Company, Ltd., Germany) was used to measure the contact angle (CA) of water droplets (5 μL) at ambient temperature. The tribological properties were studied using Xuanwu-III friction and wear tester.

2.3 Preparation of LbL modified PTFE/Nomex fabric and composites

To enhance the charges on the fiber surface, the
PTFE/Nomex fabric was treated with air-plasma (30 W, 5 min) and dipped into a dispersion containing 0.1 wt% negatively charged PAMPA and 0.1 wt% GO solution at PH 10, for 10 min. Subsequently, the hybrid fabric was rinsed with distilled water and then dried in an oven at 80 °C. The obtained fabric was then dipped into the PDDA solution at pH 10 for 10 min, washed with distilled water and dried in an oven at 80 °C. This process is regarded as one self-assembly cycle, and the resultant fabric is coded as a LbL-fabric. The process was repeated \(n\) times to obtain different fabrics with different number of self-assembly cycles, designated as LbL-n-fabric \((n = 3, 6, 9, 12)\). The fabric/phenolic composites were fabricated based on the method used in our previous work [18].

3 Results and discussion

3.1 LbL film modified hybrid fabric (LbL-fabric) characterization

The structures of the virgin fabric, LbL-6-fabric and GO were examined by XRD. Figure 1(a) shows the XRD pattern of the virgin fabric. The occurrence of a highly eminent peak located around the 18.2° region, is consistent with the high crystallinity of PTFE fibers [23]. The XRD pattern of the Nomex fibers displays two relatively lower intensity peaks at around 23.52° and 27.33° which is consistent with its reported values [24]. Compared to the virgin fabric, the intensity of the diffraction peaks for the LbL-6-fabric obviously reduced. This is attributable mainly to the GO on the fiber surfaces (see Fig. 1(b)). Figure 1(c) illustrates the XRD spectrum of the GO. A typical single strong peak [25] appeared at approximately 11.35°. Because the XRD signal of the PTFE fibers is very intense and only a small amount of GO is anchored on the fiber surface, the surface changes of LbL-6-fabric is difficult to examine using this technique.

Raman spectroscopy was adopted to investigate the structure of the lubrication and bonding surface of the hybrid fabric. Figure 2 shows the Raman spectra of the virgin and LbL-6-fabric. In the Raman spectra of the virgin fabric (Fig. 2(a)), the weak peaks at 278 cm\(^{-1}\) and 1,304 cm\(^{-1}\) correspond to the different modes of the CF\(_2\) groups [23], while the strongest peaks centered at 1,000 cm\(^{-1}\), 1,250 cm\(^{-1}\), 1,339 cm\(^{-1}\), 1,542 cm\(^{-1}\) and 1,602 cm\(^{-1}\) correspond to the C-C bands in the aromatic

![Fig. 1 XRD patterns: (a) virgin fabric; (b) LbL-6-fabric; (c) graphene oxide.](http://friction.tsinghuajournals.com)
rings of Nomex fibers [24]. The vibration at 1,649 cm\(^{-1}\) position is attributable to the band of C=O group in the main fiber structure of Nomex. However, nearly all distinctive peaks of the PTFE and Nomex fibers disappear when the hybrid fabric is modified by the LbL technique (GO/ PAMPA/PDDA) (see Fig. 2(b)). Two new broad peaks appear at 1,362 cm\(^{-1}\) and 1,590 cm\(^{-1}\) and correspond to the D and G bonds of GO, respectively. This result suggests that the GO have been successfully deposited on fibers surfaces.

To confirm the above statement, XPS spectra of the virgin and LbL-6-fabrics were obtained. The virgin fabric spectra of Fig. 3(a) show four bands, which are correspond to C1s, N1s, O1s and F1s at 283 eV, 400 eV, 529 eV and 689 eV, respectively. The F1s band of the LbL-6-fabric disappear, a situation attributable mainly to the lower fraction of PTFE in the hybrid fabric (PTFE to Nomex volume ratio is: 1:3). Two high intensity peaks at 283 eV and 529 eV (Fig. 3(b)) are attributable to C1s and O1s, respectively. However, for LbL-6-fabric, small band assigned to S2p at 167 eV is observed, suggesting that PAMPA exists on the surfaces of the LbL-6-fabric.

The morphologies of the Nomex and PTFE fiber surfaces were observed by SEM. As shown in Figs. 4(a) and 4(d), the surfaces of the as-received Nomex and PTFE fibers exhibit relatively clean and smooth morphologies. While the LbL-6-Nomex fiber (Figs. 4(b) and 4(c)) presents a highly homogeneous distribution of GO over the Nomex surfaces, the GO distribution is less-homogeneous, for the LbL-6-PTFE fiber, owing to the extremely smooth and low surface energy of the PTFE surface. These results are in agreement with the Raman spectroscopy and XPS results. Figures 4(g)–4(i) show the contact angles of the virgin and LbL-6-fabric. It can be seen that the contact angle of the lubricating and bonding surfaces of the virgin fabric is 117.4 °and 104.8°, respectively (see Figs. 4(g) and 4(h)). On the contrary, the lubricating and bonding surfaces of the LbL-6-fabric exhibited superhydrophilicity, which suggests that the LbL-6-fabric has excellent wettability. Previous work [26], has shown
that outstanding wettability is beneficial to enhancing the interfacial strength between fibers and resin matrices [26].

3.2 The influence of LbL modification on tribological properties of hybrid fabric composites

Figure 5 shows the friction and wear properties of virgin fabric and LbL-n-fabric composites, including friction coefficients and wear rates. As can be seen from Fig. 5(a), the friction coefficient of fabric composites increases with the number of self-assembly cycles, until a limiting point after which it decrease. Like in previous work [27], the addition of GO does not reduce the friction coefficient of the hybrid fabric composite, while obviously improving the wear resistance. The LbL self-assembly increased the friction coefficient from 0.0778 to 0.137, whereas the wear rate was significantly reduced from 1.231 to 0.551.

Figure 6 shows the SEM morphologies of the worn surfaces of hybrid fabric composites sliding against the counterpart pin at 70 MPa and 0.26 m/s. As seen from Figs. 6(a) and 6(e), numerous broken fibers protrude from the worn surfaces of the virgin fabric composite, in addition to the abundant wear debris, distributed around them. These situations indicate that the interfacial bonding strength between the fibers and the phenolic resin matrix is weak. For the LbL-3 hybrid fabric composite, the worn surfaces are relatively smooth with few fiber pull-outs and wear debris on the surface (Fig. 6(b)). Fiber thinning indicates that the interfacial bonding strength between the fibers and the phenolic resin matrix increase after polyelectrolytes and graphene oxide (on a layer by layer basis), were assembled on the hybrid fabric surfaces (Fig. 6(f)). However, the worn surfaces of the LbL-6 fabric composite show evidently different surface morphologies. In comparison with the virgin fabric composite, the LbL-6-fabric is well entrapped in the phenolic resin matrix owing to the deposition of graphene oxide on the fiber surface by the LbL method. As seen from Figs. 6(c) and 6(g), broken fibers and wear debris on the worn surfaces of the LbL-6-fabric composite are clearly reduced owing the improvement in interfacial bonding strength. Figures 6(d) and 6(h) shows the worn surfaces of the LbL-12-fabric composite. In this case, however, many phenolic resins are observed to peel off and numerous fibers are exposed and broken, which demonstrates that the tribological behaviors of hybrid fabric composites do not improve beyond a certain limit, as the number of
LBL self-assembly cycles is increased.

It has been proven by numerous reports that the transfer film formed on counterpart pin surfaces plays a vital role in the tribological properties of polymer matrix composites. In general, high-quality transfer films can prevent or reduce the direct contact between polymer matrix composites and steel counterpart pins. This results in excellent tribological properties. Hence, the morphologies of the counterpart pin surfaces after sliding against the virgin and LbL-6 hybrid fabric composites are shown in Fig. 7. It can be seen from Fig. 7(a) that the transfer film of the virgin composite seems to be lumpy, discontinuous and easily shelled off from the counterpart pin surfaces. Besides, numerous wear debris and furrows are observed on the surface of the counterpart pin, which indicates that the virgin hybrid fabric composite underwent severe wear. However, the incorporation of GO and self-assembly polyelectrolytes by the LbL method into the hybrid fabric composite leads to the formation of a high-quality transfer film. The transfer films formed on the counterpart pin surface were highly homogeneous and continuous (see Fig. 7(b)). Furthermore, Raman spectra analyses proved that the transfer film of the LbL-6 hybrid fabric composite contains GO. It is worth noting that the two prominent peaks at 1,365 cm$^{-1}$ and 1,611 cm$^{-1}$ correspond to the D and G bonds of GO, respectively (see Fig. 8). Hence, it can be concluded that adding GO to the hybrid fabric composite contributed to forming a homogeneous and continuous transfer film on the counterpart pin surface.

4 Conclusions

In summary, the hybrid fabric was modified using a green and mild LbL method, which only influences the chemical composition and morphology of the fiber surfaces. Besides, the effects of the LbL self-assembly cycles on the tribological performances of hybrid fabric/phenolic composites were systematically investigated. The characterization results demonstrate that GO

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Fig. 6  SEM images of the worn surfaces of hybrid fabric/phenolic composites: (a) virgin composite, (b) LbL-3-fabric composite, (c) LbL-6-fabric composite, (d) LbL-12-fabric composite, (e–f) the magnified images of (a–d), respectively.

Fig. 7  SEM images of the worn surfaces of counterpart pin sliding against hybrid fabric composite: (a) virgin, and (b) LbL-6-fabric composite.
Fig. 8 Raman spectra of the steel counterpart pin surface.

successfully anchors on hybrid fabric surfaces and significantly improves the surface wettability. The wear rate significantly reduces when the hybrid fabric is modified using the LbL self-assembly technique. Sliding wear tests indicate that LbL-6-hybrid fabric/phenolic composites exhibit the lowest wear rate. Moreover, a smooth and uniform transfer film is formed on the counterpart pin after sliding against the LbL-6-hybrid fabric composite.

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