

Bioinspired metal-polymer thin films with varying hydrophobic properties

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Abstract Nanocomposites involve the inclusion of one material into the layers of another material at a nanoscale level. Inspired by nature, nanocomposites material systems offer functionalities over their bulk forms which in some cases have evolved over millions of years. Here, thin film coatings have been fabricated by PVD sputtering, comprising a soft PTFE phase which is combined with a hard metallic NiTi phase. A series of coatings with PTFE ranging from 10 to 75 vol% have been prepared, and their surface energies and microstructures investigated. The surface energy of the nanocomposite films changes with the PTFE content, falling in the range between PTFE and NiTi with water contact angles between 80° and 102° for a thin film with 25 and 75 vol% of PTFE, respectively. Here, both TEM and EDX reveal PTFE forming along NiTi column boundaries. Coatings with PTFE content greater than 50 vol% failed due to a build-up of intrinsic stress. The degree of hybridization between NiTi and PTFE was found to be dependent on the PTFE layer thickness. SEM analysis of this coating reveals PTFE at the surface embedded within the NiTi matrix.

Keywords Nanocomposites, PVD Sputtering, Thin Films, Biomimetics, Sensors, Wear

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Introduction

Bioinspired material designs with multifunctional characteristics aim to offer properties similar to their counterparts found in nature, which in some cases have evolved over millions of years. For example, natural tissue comprises layers of hard and soft tissue material called tropocollagen and hydroxyapatite are organized into staggered arrays to form collagen fibrils.¹ The ratio between the soft and hard material phases varies according to the tissue's functionality, e.g., muscular or epithelial tissue.² The layers form a compact but strong and flexible lamella with load bearing, lubricious, and wear resistant characteristics. The soft material phase has elastomer characteristics and provides a dampening effect, whereas the hard material phase provides strength and load bearing characteristics. Another example involves micturition, a urethral function that relaxes the urethra voluntarily when it senses the bladder is full.³ Here, the combination of the surface energy of the material, which determines the water contact angle (WCA), and the frictional properties of the tissue plays a significant role in defining the wetting and fluid flow characteristics down the urethral tissue.^{3–5} This principle could be applicable to cases where the ability to modify a surface between hydrophilic and hydrophobic can, for example, be used to increase engine component lifetimes. In a modern stop-start engine for example, increased wetting during the *stop*-phase could be used to prevent premature engine wear by always maintaining a thin oil film layer in areas where there is potential of metal-to-metal contact. Since the WCA of a material affects its wetting properties, the ability to modify this characteristic by the way of incorporating a secondary low-surface energy material is investigated. In fact, modifying the surface energy of a metal layer with the addition of a hydrophobic layer has been reported before, where tungsten is coated with PTFE to a form a hydrophobic

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surface.⁶ Incorporating a low-surface energy material such as PTFE with NiTi as the initial choice of materials in this study is driven by their biocompatibility and the need for intelligent in vivo drug delivery applications, where current challenges include the need to administer the correct dose and quantity of drugs when most needed by a patient.⁷ Thin films comprising a soft PTFE phase, which has hydrophobic characteristics, have been combined with a harder NiTi metal phase, a shape memory alloy (SMA) material that offers shape memory functionality in response to heat or stress. It is well documented that SMAs transform from a low temperature, low allotrope martensite B19' monoclinic phase, to a high temperature, high-symmetry allotrope austenite B2 cubic phase. NiTi undergoes such a B2-B19' phase transformation but is limited to a maximum operating temperature of around 100°C.⁸ Thus, the combination of the material biocompatibility and the shape memory functionality could, in theory, be applicable in drug delivery systems as part of a nano-fluid sensor which triggers a dose of drugs that is administered in response to a patient's body temperature. Thin films are prepared by sputter physical vapor deposition (PVD) as this allows for the control of individual film thicknesses at a nano-level, permitting a greater control of film microstructure.⁹ The objective is to modify the surface energy by varying the amount of PTFE and NiTi content in the coating and investigate the resultant wetting and microstructural properties. This study continues our initial research on such coatings where the tribological properties such as the coefficient of friction were measured and the concept of introducing a polymer-shape memory alloy nanocomposite film was initially proposed.¹⁰

Method

The coatings were prepared in a Leybold L560 sputtering machine equipped with two 3-inch-diameter magnetrons-NiTi, powered by a dual-output Advanced Energy Pinnacle® pulse dc supply operating at 200 kHz at a 1.6 µs duty factor, and PTFE powered by an RF power supply operating at 13.56 MHz. Sputtering power densities ranged between 3.5 and 5.3 W/cm^2 . Prior to deposition, the system was turbopumped to a base pressure around 1×10^{-6} mbar, and then Ar gas flowed at a controlled rate to realize the experimental process pressures. The targets were sputter-cleaned (presputtering) for 5 min prior to each sputtering run to remove any impurities. The nanocomposite films were prepared by one of three system operating modes-stationary or rotational, details of which follow, or by the familiar co-sputtering route. In the stationary mode of deposition, the substrate is placed above a magnetron and the target material sputtered for a preset time to produce the coating. The substrate is then moved to the second magnetron and again sputtered for a predetermined time period, producing a thin film coating by the classical layer-by-layer approach.¹¹ Modifying the hybrid nature of the nanocomposite was achieved by varying the volume fraction of PTFE within the thin film, realized by altering the time spent over each magnetron, which in turn altered the layer thickness of either NiTi or PTFE. This approach follows from previous work by this group on multilayer thin film coatings,¹² where the wavelength (λ) is defined as the sum of the two layers, namely $T_{\rm P}$ and $T_{\rm M}$, corresponding to the thickness of the polymer and the metal layers, respectively.

$$\lambda = T_{\rm P} + T_{\rm M} \tag{1}$$

The volume (%) of the polymer content in the coating $(V_{\rm P})$ could then be varied according to:

$$V_{\rm P}(\%) = \frac{T_{\rm P}}{(T_{\rm P} + T_{\rm M})} \times 100 \tag{2}$$

Layered coatings with $V_{\rm P}$ ranging between 10% and 70% have been fabricated with a $\lambda = 200$ nm. In the rotation mode of deposition, the substrate table is rotated continuously during deposition over both magnetrons at a controlled speed, producing an interrupted staggered arrangement of nano-layers with nano-level interplanar spacing.¹³ The layer thicknesses are defined by the time resident over each of the targets: high rotational speeds reduce the time spent above each target and hence lead to thinner layers, whereas low rotational speeds increase the resident time over each target thereby increasing the layer thickness. In the co-sputter route, the substrate is placed between the PTFE and NiTi targets, and deposition occurs by operating both targets simultaneously. The coatings were prepared to realize a total thickness of 2 µm. Film composition is controlled by modifying the target power densities. Glass and Si wafers were used as substrates to prepare the nanocomposite thin films, which were cleaned prior to deposition using IPA. Static water contact angle (WCA) measurements were taken by the sessile droplet method, using a syringe filled with deionized water with a volume of 1.0 μ L at dosing rate of 2.0 μ L/ s. An adjustable cold light source was used to illuminate the sample. The instrument uses SCA20 software to capture images and calculate the WCA. An SEM system, FEI XL30 SFEG equipped with energydispersive X-ray spectroscopy (EDX) operated at 20 kV, was used to investigate the film surface and elemental composition, respectively. TEM images were taken by a JOEL 2000FX TEM equipped with an Oxford Instruments EDX detector, housed at Loughborough University. Samples for TEM were prepared by focused ion beam (FIB), FEI FIB200, which uses gallium ions at a beam current ranging between 1 and 11,500 pA.

Results and discussion

PTFE thin films

The energetics of the sputtering process can affect the polymerization of the complex functional groups in PTFE.^{14,15} An important factor to establish first is whether sputtering retains the polymerization of the deposited material.^{16,17} To ascertain the polymerized content, Fourier transform infrared spectroscopy (FTIR) was conducted on a number of PTFE films deposited by RF sputtering under various processing conditions. The main features of an FTIR spectrum from PTFE are the strong absorption bands between 1000 and 1400 cm⁻¹, characteristic of C–F_X (X = 1, 2) stretching vibrations. An analysis of the resultant FTIR spectrum obtained in this study is shown in Fig. 1. The area under the CF_X (X = 1, 2) peaks is associated with highly polymerized PTFE.¹⁸ The area under the peak at 900 cm^{-1} is associated with CF₃ and is related to the polymer chain ends, where more monomeric, fragmented, or highly branched PTFE would have more CF_3 .¹⁹ Thus, the greater the area under this peak, the lower the polymerized content. An ideal PTFE film prepared by a sputtering route would therefore have a large area under the CF_X peak compared to the area under the CF_3 peak. Additionally, the presence of CF_3 and CF₂ radicals in sputtered PTFE thin films contribute to reducing the surface energies, thereby resulting in a hydrophobic surface.²⁰ The optimum polymerization for RF-deposited films in this study was at a power of 150 W and a pressure of 15 mbar (presented in Fig. 1), since these processing conditions minimized the degree of crosslinking and unreacted monomers which would produce less continuous films, supporting previous studies by Stelmashuk.²¹ These process conditions were adopted for subsequent fabrication of the intercalated PTFE-NiTi thin films. The findings support previous studies on sputtered PTFE films where it was determined that the degree of crosslinking is not affected as a result of sputter ion bombardment.^{14,19,22,23} Additionally, it has been noted that films with reduced polymerization had rougher surfaces owing to changes in their functional groups which produced excessive crosslinking and a high proportion of unreacted monomers.^{18,19,24,25}

NiTi thin films

Pulse-DC sputter-deposited intermetallic NiTi thin films are amorphous and require subsequent post-heat treatment and annealing to realize reversible martensitic to austenitic (B2–B19') phase transformations.²⁶ NiTi thin film coatings with the Ti content varying between 40 and 54 at% have been prepared. The temperatures at which the martensite and austenite phases begin, peak, and end have been measured by differential scanning calorimetry (DSC). The data plotted in Fig. 2 show only the austenitic peak phase temperature, which increases proportionally with increasing Ti content in near equiatomic NiTi, supporting other studies,^{8,27} and demonstrating an upper transition temperature that can be varied in the range -20 to 140° C.

The NiTi composition making up the nanocomposite films was selected as 47 at% Ti, since this composition realized a peak austenitic phase transformation close to body temperature. As the nanocomposite films cannot be heat treated due to the low melting temperature of PTFE, the phase transformations in NiTi are instead triggered by NiTi's pseudoelastic characteristics.²⁶

Nanocomposite thin films

The thin films prepared in this study are presented in Table 1 along with the measured WCA and their mode of fabrication. A thin film of RF-sputtered PTFE is hydrophobic having the highest measured WCA of 108° . On the other hand, a thin film of pulse-DC sputtered NiTi exhibits hydrophilic characteristics with a measured WCA of 55° .



Fig. 1: FTIR spectra of a PTFE film fabricated on a glass substrate. The strong absorption bands between 1000 and 1400 cm⁻¹ relate to C–F_X (X = 1, 2) stretching vibrations: RF sputter-deposited film at 150 W and 15 mbar chamber pressure



Fig. 2: A plot of the change in austenitic phase transformation temperature with increasing Ti in NiTi. This linear increase in phase transformation temperature with Ti content has been reported by other studies²⁷

Table 1: PTFE-NiTi nanocomposite thin films investigated in this study

Sample ID	V _p (%)	WCA (°)	Deposition mode
PTFE	100	108	_
NiTi	_	55	_
A	10	88	Layered
В	25	79	Layered
С	50	68	Layered
D	75	102	Layered
E	35	$96 \rightarrow 52$	Co-sputtered
F	54	$102 \rightarrow 43$	Co-sputtered
G	25	99	Hybrid ^a
Н	10	87	Rotated

 V_{P} represents the vol% of PTFE in the coating. Films with a PTFE content greater than 30% subsequently failed due to an increase in stress build-up. The WCA of samples labeled here as E and F dropped during the measurements

^a Film deposition by first layering and then co-sputtering

Figure 3 shows photographs of a droplet of water on the surface of the films and is used to determine the WCA. The deposition process was intentionally terminated on the NiTi layer in all films produced by the layered approach. The thin films have a measured WCA ranging between bulk PTFE and bulk NiTi, suggesting a varying degree of hybridization between the films and the deposition approached adopted. An increase in the PTFE content $(V_P\%)$ in the coatings prepared via a layered route increases the measured WCA, approaching that of bulk PTFE in a thin film with 75 vol% PTFE. The measured WCAs for both samples C and D, having NiTi as their top layers, differ with C having a lower WCA than D, suggesting that there is a greater degree of hybridization between PTFE and NiTi in sample D.

However, both samples subsequently failed due to intrinsic stresses, and no further analysis was possible. In fact, it was observed that films with PTFE greater than 30 vol% were all prone to mechanical instability as these films disintegrated soon after the WCA measurements were taken. Further analysis was therefore not possible; however, an analysis of the material flakes suggests that the failure was cohesive, probably due to the intrinsic stressed state of the polymer phase interlayers producing delamination cracks along the polymer and metal interface, 2^{28} which constitutes a cohesive failure. As a route to overcome such in-built stresses, sample G was fabricated by employing a hybrid approach—layered followed by co-sputtering. The thickness of the individual layers was intentionally reduced toward the surface as a route to reduce stress, and a series of 10 PTFE-NiTi layers with a volume fraction of 25% PTFE deposited. PTFE-NiTi was finally co-sputtered to complete film deposition. A TEM image of this film is shown in Fig. 4 and will be discussed later in this paper. Thin films that were deposited by co-sputtering proved unstable as their surface energy changed dramatically over the course of the WCA measurements, samples E and F. The measured WCA dropped by 50% from around 100° in the space of 10–15 s, suggesting that microstructural porosity and instability of the hybrid nanocomposite framework containing 30 vol% PTFE or greater were





Fig. 4: TEM image of sample H fabricated by a co-sputtering and a layered approach. (a) Bright-field image showing PTFE as the lighter bands, (b) dark-field image showing PTFE as the darker bands. (c) A close-up image of the sample showing greater hybridization as the PTFE layer thickness is reduced

Fig. 3: Image of a water droplet on the surface of the nanocomposite films, which was used to measure the water contact angle (WCA). Apart from sample G, the samples were prepared using a layered approach with the process terminating on the NiTi layer. The WCAs are between bulk PTFE and NiTi

responsible. The quantity of water absorbed was measured by dropping a known volume of water on an area of the film. For a 2- μ m-thick film absorbing water over a 0.25 cm² area, the amount of water absorbed was estimated to be approximately 0.017 μ L. The absorbed volume of water represents the vacant volume within the thin film, which equated to around one-third of the total volume of the thin film. This is not attributed to simple evaporation due to the timescales (10–15 s) the measurements were taken over. The measured WCAs for the co-sputtered films suggest hydrophilic characteristics and support a previous study on nano-porous thin films, which concluded that water absorption was greater in films that were hydrophilic and porous.²⁹

Figure 4 shows both the bright and dark-field TEM images of G, which on first viewing appear as distinct layers that decrease in thickness toward the surface. The co-sputtered top layer of the coating fell off, but the layered-deposited coating remained intact. PTFE appears as light-colored bands in (a) and appears as dark bands in (b). A close-up of the bright-field image in (c) shows, in addition to continuous PTFE layers running parallel to the substrate, deposition of further

PTFE along NiTi columnar boundaries. Also, as the layer thickness of PTFE is reduced, there is an increased visible evidence of PTFE forming along the NiTi column boundaries, i.e., there is an increasing degree of hybridization as the PTFE layers and NiTi layers become thinner toward the surface. The increasing level of hybridization is thought to be due to the repulsive interplay of the electronegative fluorine in PTFE and electrons in NiTi, in addition to intra-layer attraction between NiTi metallic cations and PTFE. To further test this hypothesis, a coating with a PTFE volume fraction of 10% (sample H) was prepared.

The bright and dark-field TEM images taken of sample H are shown in Figs. 5a and 5b, respectively. The thicknesses of the PTFE layers are around 30 nm, appearing as white bands running from left to right in 5a and as dark bands in 5b. In regions between the PTFE, within the NiTi layer (bands colored gray), there are visible signs that PTFE appears vertically running along NiTi column boundaries, thus supporting our notion of hybridization. However, it could be argued that such vertical bands could be empty and their gray coloration is in fact an atomic number contrast from a mixture of NiTi and PTFE. A study to investigate this further was undertaken using energydispersive X-ray analysis (EDX) SEM, and the results of which are shown in Figs. 5c and 5d, showing the O and F k- α maps, respectively.



Fig. 5: An intercalated PTFE–NiTi thin film formed when the PTFE layers are reduced to 30 nm, Sample H. (a) The brightfield image shows PTFE (light areas) forming along the column boundaries of the darker, NiTi layers. (b) A dark-field image of the same region of the structure. (c), (d) Energy-dispersive X-ray images of the $k\alpha^1$ X-rays from the O and F, respectively, further supporting the notion of hybridization and where the fluorine in PTFE is observed running vertically between the NiTi column boundaries

By filtering only X-rays from $O_{k\alpha}^1$ and $F_{k\alpha}^1$, Figs. 5c and 5d, respectively, show that the fluorine from the PTFE appears as horizontal bright bands running parallel to the substrate, together with areas within the NiTi layer where fluorine is observed running vertically along NiTi column boundaries. The fluorine map implies the presence of PTFE and confirms the hybrid nature of the nanocomposite. We surmise that this occurs due to the intermolecular forces between the PTFE chains and crosslinked fragments which are arranged in both vertical and horizontal directions.³⁰ Thus, both the TEM images and EDX analysis support our belief that the two materials produce a hybrid nanocomposite material. Evidence of this hybrid material with the PTFE polymer appearing on the

surface of the coating is now presented. A study of the surfaces of the thin film samples A and H was undertaken using SEM, both produced with 10% volume fraction PTFE.

The surface of H comprises of PTFE hybridized within the NiTi matrix and thus accounts for the mixmode measured WCA (Fig. 6a). The surface of this film is compared and contrasted with A, which also has the same measured WCA, but fabricated with thicker PTFE layers (Fig. 6b). The surface of A comprises dense columnar grains, characteristic of a PVD-sputtered film, with EDX analysis of these regions confirming NiTi-rich regions interspersed with PTFE-rich nodules on the film surface. From these images, we can conclude that H is more hybrid in nature than A, which



Fig. 6: Surface morphologies of two samples with the same 10% volume of PTFE, but prepared by (a) Sample H; rotation, (b) Sample A; layered

we understand to be due to the more stable and thinner PTFE and NiTi layers.

Conclusions

This work has demonstrated the successful fabrication of metal-polymer nanocomposite thin films by PVD sputtering. Nanocomposite thin films have been realized by either multilayering or continuously rotating above NiTi and PTFE targets during the deposition process. Co-sputtered films failed due to intrinsic stresses, and further analysis of this co-sputtered microstructure was not possible. Furthermore, films with PTFE greater than 30 vol% were also shown to be unstable. Results from water contact angle measurements reveal that the surface of the nanocomposite films has a WCA that falls between that of a thin film of PTFE and NiTi. Further microstructural analysis using TEM reveals that the hybrid nature increases in films with thin (30 nm) PTFE layers, where EDX Xrays from $O_{k\alpha}^1$ and $F_{k\alpha}^1$ shows PTFE forming along NiTi column boundaries. Surface studies of two films with the same vol% of PTFE but with different PTFE layer thicknesses have different morphologies. The surface of the coating fabricated with 30 nm PTFE layers has visible PTFE at the surface and growing between the voids in the NiTi metal matrix, whereas a coating with thicker PTFE layers has a surface more like that of a PVD-deposited metallic coating (NiTi was the topmost laver).

It is proposed that the characteristics of such films could, in theory, suit a future drug delivery system which responds to a patient's body temperature to administer drugs accordingly. Another potential application is where a hydrophobic film needs to be maintained, for example, in bearings designed for cyclic stop–start applications. Changes in the austenite to martensite transformations in the NiTi alloy could be employed to modify the surface energies in response to the external environment.

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