Developing highly nanoporous titanate structures via wet chemical conversion of DC magnetron sputtered titanium thin films

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Titanate structures have been widely investigated as biomedical component surfaces due to their bioactive, osteoinductive and antibacterial properties. However, these surfaces are limited to Ti and its alloys, due to the nature of the chemical conversion employed. The authors present a new method for generating nanoporous titanate structures on alternative biomaterial surfaces, such as other metals/alloys, ceramics and polymers, to produce bioactive and/or antibacterial properties in a simple yet effective way. Wet chemical (NaOH; 5 M; 60 °C; 24 h) conversion of DC magnetron sputtered Ti surfaces on 316L stainless steel were investigated to explore effects of microstructure on sodium titanate conversion. It was found that the more equiaxed thin films (B/300) generated the thickest titanate structures (ca. 1.6 μm), which disagreed with the proposed hypothesis of columnar structures allowing greater NaOH ingress. All film parameters tested ultimately generated titanate structures, as confirmed via EDX, SEM, XPS, XRD, FTIR and Raman analyses. Additionally, the more columnar structures (NB/NH & B/NH) had a greater quantity of Na (ca. 26 at.%) in the top portion of the films, as confirmed via XPS, however, on average the Na content was consistent across the films (ca. 5–9 at.%). Film adhesion for the more columnar structures (ca. 42 MPa), even on polished substrates, were close to that of the FDA requirement for plasma-sprayed HA coatings (ca. 50 MPa). This study demonstrates the potential of these surfaces to be applied onto a wide variety of material types, even polymeric materials, due to the lower processing temperatures utilised, with the vision to generate bioactive and/or antibacterial properties on a plethora of bioinert materials.

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1. Introduction

Medical implants’ reliance on extracellular tissue growth, is fundamentally restricted by the success of the surface modification
used to enhance their inherently bioinert surfaces. Despite titanium (Ti) and its alloys’ (specifically Ti-6Al-4V) advantages regarding corrosion resistance, and their minimal allergenic and immunogenic potential compared to previously used cobalt-chrome (Co-Cr-Mo) alloys or 316L stainless steel (SS), they require sufficient modification to ensure a more natural adhesion (integration mechanically and chemically achieved through bone ingrowth rather than through cementation) at the implant-tissue interface [1].

Presently, surface modification of metallic implants used clinically is performed exclusively by coating (in line with the Food and Drug Administration’s (FDA) regulatory processes) with hydroxyapatite (HA) through high-temperature (>15,000 K plasma temperature [2]; >1500 K droplet temperature [3]) plasma spraying methodologies [4]. By propelling molten particles, generated via injection of powder through a plasma torch, towards the implant surface, a coating forms through rapid quenching and solidification. However, this process causes residual stresses and crystallisation within the produced coating [5], and can often modify the underlying substrate. Over time, the residual stresses encourage propagation of micro-cracks, thereby reducing the long-term stability of the coating in vivo. This can lead to delamination and particle formation which, in turn, can lead to loosening of the implant [6].

An alternative methodology, which aimed to circumvent the issues with plasma spraying, was developed by Kokubo et al. [7–9]. Bioactive titanates (particularly sodium titanate: Na$_2$Ti$_x$O$_{2x-1}$; $x = 1.2, 3$ …), which are generated through simple wet-chemical routes at much lower processing temperatures (60 °C), demonstrated tailorability through cationic exchange, as well as nanoporous morphology for potential bone ingrowth. By facilitating ion-exchange reactions in vitro and in vivo, apatite formation was possible, enabling natural adhesion at the implant-tissue interface, with successful clinical deployment in 10,000 patients in Japan (98% implant survivorship 10 years post-implantation [10]). Titanate surfaces have also been considered for antibacterial applications due to their ion-exchangeability, with silver (Ag) [Lee et al. [11]] and more recently gallium (Ga) (Wadge et al. [12] and Yamaguchi et al. [13]), to facilitate a two-in-one bioactive and antibacterial surface, being notable examples.

Translation of these surfaces on alternative substrate materials (other metals/alloys, polymers, and ceramics), however, has been limited, specifically in the biomedical sector. Various research groups have investigated reactive magnetron sputtering of titanate structures [14], however no research investigating wet-chemical conversion of direct current (DC) magnetron sputtered Ti thin films has yet been published. The overall aim of our research is to develop this technology for a wide range of biomaterials, enabling functionalisation of surfaces tailored to specific applications. The wet chemical method used, as outlined by Kokubo and his team [7–9], relies upon substrate submersion in concentrated (>5 M) sodium hydroxide (NaOH), enabling simplistic and low energy production of sodium titanate structures. This technique is appropriate for the intended application, since it is scalable, enables full conversion of complex geometries, is not energy intensive and does not require complicated costly equipment. The objectives of this study were to investigate the effect of initial surface parameters (film density, structure factor etc.) on the wet-chemical conversion to titanate structures, and explore the possibility of producing titanate structures on alternative biomedical materials.

2. Methodology

2.1. Substrate preparation

316L stainless steel (SS) discs (1 mm thick; 10 mm diameter; >99.9% purity; Smiths Metals Ltd., Nottingham) were polished using abrasive silicon carbide (SiC) paper, with sequentially higher grit from P240 to P4000, followed by colloidal silica polishing (ca. 0.06 μm particle size) on MD-Chem polishing pads. Polished fused silica discs were utilised for surface and cross-sectional SEM observations. The discs were then cleaned ultrasonically in acetone (99.8% purity, Honeywell), followed by Industrial Methylated Spirit (IMS, purity 99.9%, Sigma-Aldrich) for 10 min each, and subsequently air dried, prior to sputtering.

2.2. DC magnetron sputtering

To produce the Ti thin films, a TEER UDP-650 type 2 unbalanced magnetron sputtering rig was used, fitted with a solid Ti target (>99.6% purity; 175 × 380 mm, Teer Coatings Ltd.). All substrates were fixed to a rotating sample holder using double-sided polyimide Kapton tape, with samples mounted at a set substrate to target distance (55 ± 1 mm) and were rotated at 5 RPM. Initially, the chamber was pumped down using consecutive rotary and diffusion pumping stages to a base vacuum of <1 × 10⁻⁵ Torr, before introducing Ar gas (purity 99.999%; Pureshield BOC®) at a rate of 20 standard cubic cm/min (sccm). Prior to deposition, all samples were bias cleaned using a pulsed DC bias at a pulse width of 250 kHz, a step time of 500 ns, and a bias voltage of −100 V for 15 min. Various processing parameters were varied to induce structural difference in the produced Ti films, however, target power density was maintained at 2.38 W cm⁻² (5 A, −300 V) to minimise sputtering times. Two parameters that were varied during the sputtering process; the applied bias to the substrates (either 0 or −100 V) and substrate temperature, which was controlled using a mounted heater within the vacuum chamber, it was either off or operated at 150 or 300 °C. Processing parameters and matching sample codes are shown in Table 1.

2.3. Sodium hydroxide chemical treatment

A ca. 5 M NaOH solution was prepared using NaOH pellets (ca. 19.99 g per 100 mL of distilled water; 99.9% purity; Sigma-Aldrich). 10 mL aliquots were then dispensed into polypropylene containers, with individual discs suspended in each. These were then sealed and heated in a low temperature furnace at 60 ± 2 °C for 24 h. Upon removal, samples were washed in deionised water for 60 s, before compressed air drying followed by storage within a desiccator until testing.

| Sample Code | Target Current/A | Substrate Bias/V | Substrate Heating/°C | Coating Time/min | Calculated Sputtering Rate/mm min⁻¹ |
|-------------|------------------|------------------|----------------------|-----------------|-----------------------------------|
| NB/NH       | 5                | 0                | 0                    | 263             | 14.8 ± 0.11                       |
| B/NH        | 5                | −100             | 0                    | 278             | 14.0 ± 0.12                       |
| B/150       | 5                | −100             | 150                  | 278             | 13.3 ± 0.15                       |
| B/300       | 5                | −100             | 300                  | 278             | 13.2 ± 0.08                       |
2.4. Materials characterisation

2.4.1. Scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDX)

Scanning electron micrographs and surface compositional analysis were obtained with a JEOL 7100F Field-Emission Gun Scanning Electron Microscope (FEG-SEM) with an attached Energy Dispersive X-ray (EDX) Spectrometer (Oxford Instruments). A working distance of 10 mm was maintained with acquisitions utilising a beam voltage of 15 kV. Qualitative pore/void area and frequency analysis was conducted using the ‘analyse particle’ plugin in ImageJ, from high contrast SEM images for good accuracy, with a threshold set at ca. 45%.

Cross-sectional SEM was conducted to determine the thickness of all converted and unconverted layer thicknesses, with error measurements taken from 5 separate areas, with 3 replicate measurements taken for each area ($n = 15$). Fused silica discs were utilised for these measurements, as shattering of the discs using a diamond-tipped scribe enabled clear imaging of the cross-sectional morphology. Statistical significance was calculated using a one-way analysis of variance (ANOVA) assessment.

2.4.2. Focussed ion-beam scanning electron microscopy (FIB-SEM)

FIB-SEM was used to mill out sections of the sample for further cross-sectional analysis. An FEI Quanta 200 3D (FEI, Portland, OR) system was used, with Ga+ ion milling performed at 30 kV, and currents ranging from 1 nA to 30 pA. Standard FEI gas injectors were used to deposit platinum (Trimethyl(methylcyclopentadienyl)platinum(IV); 25°C crucible temperature), which was introduced onto the sample via an internal gas injector. Samples were imaged on the JEOL 7100F FEG-SEM utilising the same parameters as Section 2.4.1, with a 40° stage tilt.

2.4.3. X-ray diffraction (XRD) and texture coefficient calculation

Crystalline structures, particularly noting preferred orientation in specific crystallographic planes, were assessed using a Bruker D8 advance XRD spectrometer, utilising a parallel beam geometry and glancing angle beam parameters as follows: Cu Kα source; $\lambda = 1.5406\text{ Å}$; 40 kV; 35 mA. Spectra were measured over a 2θ range of 20–60°, with a step size of 0.015° (2θ), a glancing angle of 1.2°, and a dwell time of 16 s. Relative peak intensities were calculated from the raw spectral data in Bruker’s DIFRAC.EVA software, and the texture coefficient values were obtained using the Harris equation [15]:

$$T_i(h,k,l) = \left(\frac{l_{\text{m}}(h,k,l)}{l_0(h,k,l)}\right)^{1/n}$$

where $l_{\text{m}}$ is the measured diffraction intensity, $l_0$ is the diffraction database intensity value, and $n$ is the number of peaks being calculated, in this case $n = 4$.

2.4.4. X-ray photoelectron spectroscopy (XPS)

For X-ray Photoelectron Spectroscopic (XPS) measurements, a VG ESCALab Mark II X-ray photoelectron spectrometer with a monochromatic aluminium (Al) Kα X-ray source was utilised. Beam incidence angle was maintained at ca. 30°, with both survey and high-resolutions scans conducted. For calibration, high-resolution scanning of the C 1s spectral peak, charge corrected to 284.8 eV, was conducted. Acquisition parameters used were as follows: scan number = 5; dwell times of 0.2 and 0.4 s for survey and high-resolution scans, respectively; and a step size of 1 and 0.2 eV for survey and high-resolution scans, respectively. Survey scans were measured over a range of 0–1200 eV, with all scans analysed using CasaXPS software constraining all deconvoluted peaks of the same elements to the same Full Width at Half Maximum (FWHM), with additional area ratio constraints maintained for spin-orbit splitting (or j-j coupling).

2.4.5. Raman spectroscopy

For Raman spectroscopy measurements, a HORIBA Jobin Yvon LabRAM HR spectrometer was utilised. Beam and acquisition parameters were as follows: 532 nm laser (25 mW power); 50× objective lens; 300 μm confocal pinhole; 600 lines/mm rotatable diffraction grating with a path length of 800 mm, for simultaneous scanning of multiple Raman shifts. A SYNAPSE CCD detector (1024 pixels, thermoelectrically cooled to −60 °C, was utilised for spectra detection. Calibration of the instrument was conducted using both a standard Si (1 0 0) reference band (520.5 cm$^{-1}$) and the Rayleigh line (0 cm$^{-1}$). Spectral acquisition was conducted for 10 s, with 10 accumulations per sample.

2.4.6. Fourier transform infrared spectroscopy (FTIR)

A Bruker Tensor FTIR spectrometer with an Attenuated Total Reflectance (ATR) attachment containing a diamond crystal/ZnSe lens was used for infrared absorbance measurements. A spectral range of 4000–500 cm$^{-1}$ wavenumber was observed and maintained for all acquisitions.

2.4.7. Surface roughness – Optical profilometry

Surface roughness quantification was conducted using focus variation microscopy performed on an Alicona Infinite Focus instrument. A 50× objective lens was used in all cases, with raw data being levelled using the least squares plane method. 3D area surface texture parameters were extracted using the ISO 25178-2 standard [16]. A vertical resolution of 70 nm was utilised, with a horizontal resolution of 2 μm.

2.4.8. Pull off adhesion strength

The adhesion strength of sputtered Ti films was tested using a portable adhesion testing (P.A.T handy (DFD instruments)) unit in accordance with ASTM-D4541-17 [17]. Stubs (2.8 mm radii) were adhesively bonded to the coated samples using DFD® E1100S epoxy, which was cured for 60 min at 140 °C, and then left to cool to room temperature. All discs were cleaned with compressed air, and air bubbles formed between the stub and disc were carefully removed through pressing of the stubs. Any excess epoxy following curing was removed via a cylindrical cutting tool supplied with the equipment. Analysis of the failure sites was conducted using a Nikon LV100ND upright light microscope.

2.4.9. Scratch adhesion testing

Analysis of both unconverted and titinate converted coating failure was conducted in accordance to BS EN ISO 20502:2016, using a CETR UMI multiple specimen scratch testing system equipped with a Rockwell C indenter. Prior to each scratch, the indenter was cleaned with isopropyl alcohol. Modified testing regimes were utilised due to some failure modes being considerably less than the standard 30 N, making quantification difficult. Therefore, loads were applied over a distance of 3 mm, with pre-loading of 0.5 N and progressive loading up to a value of 15 N over a period of 180 s. The failure sites were again analysed using a Nikon LV100ND upright microscope.

3. Results

3.1. Topographical, structural and compositional analysis

3.1.1. SEM and EDX plus XPS compositional analysis

As observed in the SEM micrographs (Fig. 1A–D) the frequency (ca. 4.1, 3.0, 1.8, and 0.6 μm$^{-2}$) for NB/NH, B/NH, B/150 and B/300
samples, respectively) and area (ca. 3713.7, 2541.3, 1436.8, and 1028.6 nm², for NB/NH, B/NH, B/150 and B/300 samples, respectively) of surface voids appear to diminish from the NB/NH sample to the B/300 sample (see supplementary data Fig. S1). The sputtering time was increased from 263 min for the NB/NH sample (Fig. 1E) to 278 min for the B/NH sample (Fig. 1F) to maintain the same coating thickness (ca. 3.9 µm), suggesting the increased density of the coating given the similar rate of deposition. Furthermore, through the application of substrate heating at 150 and 300 °C, it is clear the thickness of the produced films at the same sputtering time decreased to ca. 3.7 µm for the B/150 and B/300 samples (Fig. 1G–H).

Clear surface alterations are noted for the titanate samples, with nanoporous morphologies consistent with sodium titanate surfaces (Fig. 1I–L). It is also noted that the pore size (defined as the area enclosed by the larger strutted structures) appears to increase with increasing density of the produced Ti coating, however, the large error associated with the B/150 TC sample in the supplementary data may affect its significance (see supplementary data Fig. S2). In addition to the titanium coating, the thickness of the titanate converted films were also assessed. When observing the thickness of the titanate converted portion of the Ti coating, it is evident that this region increases in depth through the application of a bias (from the NB/NH sample (ca. 1.12 µm) to the B/NH sample (ca. 1.20 µm); Fig. 1M & N), as well as the increase in substrate temperature (from the B/150 sample (ca. 1.20 µm) to the B/300 sample (ca. 1.63 µm); Fig. 1 O & P). The increase in depth from all titanate converted samples to the B/300 TC sample is statistically significant (p < 0.0001), however, the increase from the NB/NH_TC, to both the B/NH_TC and B/150_TC samples exhibited no statistical significance.

Elemental analysis via EDX (Table 2) demonstrated similar sodium (Na) inclusion of ca. 7.6, 8.9, 5.6 and 7.6 at. % for the NB/NH_TC, B/NH_TC, B/150_TC, and B/300_TC samples, respectively. Both Ti and O content remained constant, with small inclusions (<1 at. %) of Ca. In comparison, elemental analysis determined via XPS showed a large increase in Na for all samples, with 25.6, 26.8, 16.1, and 17.8 at. % for NB/NH_TC, B/NH_TC, B/150_TC and B/300_TC, respectively. The surface composition examined by XPS showed that Ti and O content remained similar for the NB/NH_TC, B/NH_TC samples, but was higher for the B/150_TC and B/300_TC samples.

3.1.2. FIB-SEM

Additional cross-sectional information of the samples was determined via FEG-SEM of a FIB milled titanate converted sample. The micrograph (Fig. 2A) demonstrates clear interconnected nanoporosity, with homogenous distribution of sodium (Fig. 2C).

**Table 2**

EDX and XPS data of titanate converted DC magnetron sputtered Ti films. Values given with standard error values (n = 3). All unconverted samples contained 100% Ti, with titanate converted samples also containing trace amounts of Ca (<1 at.%).

| Sample    | EDX     | XPS            |
|-----------|---------|----------------|
|           | Na(at.%) (EDX) | O(at.%) (EDX) | Ti(at.%) (EDX) | Na(at.%) (XPS) | O(at.%) (XPS) | Ti(at.%) (XPS) |
| NB/NH_TC  | 7.6 ± 0.1 | 61.0 ± 0.1 | 30.8 ± 0.1 | 25.6 ± 0.6 | 56.9 ± 0.6 | 17.5 ± 0.3 |
| B/NH_TC   | 8.9 ± 0.1 | 57.3 ± 0.5 | 33.2 ± 0.6 | 26.8 ± 0.7 | 55.4 ± 0.6 | 17.8 ± 0.3 |
| B/150_TC  | 5.6 ± 0.1 | 57.6 ± 0.2 | 36.9 ± 0.2 | 16.1 ± 0.4 | 62.4 ± 0.5 | 21.5 ± 0.3 |
| B/300_TC  | 7.6 ± 0.1 | 60.1 ± 0.2 | 32.2 ± 0.1 | 17.8 ± 0.4 | 62.4 ± 0.5 | 19.9 ± 0.3 |
across the surface of the sample, and homogeneous distribution of titanium within the coating (Fig. 2F). Sodium and oxygen (Fig. 2C & D) penetration appears to decrease through the through thickness of the coating. Vertical growth of the titanate struts are clear on the insert micrographs.

3.1.3. XRD and texture coefficient

XRD spectral data (Fig. 3A) demonstrated for the NB/NH sample, preferred orientation in the (0 0 2) plane of HCP Ti (PDF 00–044-1294) at 38.4° 2θ, with additional smaller peaks at 40.2° and 53.0° 2θ corresponding to the (1 0 1) and (1 0 2) planes, respectively; no other peaks were observed. However, through the application of a bias to the B/NH sample, the relative intensities of the peaks, and number of peaks increased, with a new peak at 35.2° 2θ corresponding to the (1 0 0) plane. The peak corresponding to the (1 0 1) plane increased in intensity. Continuing the trend, through application of substrate heating in both the B/150 and B/300 samples, peaks corresponding to the (1 0 0) and (1 0 1) planes saw an increase in their relative intensities, with the (1 0 1) peak having the highest relative intensity for both samples. Following titanate conversion (Fig. 3B), 3 additional broader peaks were detected on all samples at ca. 24.2, 28.4 and 48.2° 2θ, assigned to titanate species (PDF 00-022-1404 and PDF 00-025-1450), however direct assignment was difficult due to the amorphous nature of the peaks.

Further analysis of the change in preferred orientation was conducted through calculation of texture coefficients \( T_c \) (Fig. 4). Using Harris’ equation, the degree of preferred orientation can be numerically calculated through ratios between measured peak intensities and database intensities for the same material. A \( T_c \) of 1 would indicate an equiaxed sample equating to the material’s database structure, whilst a texture coefficient of \( n \) (\( n \) being the number of peaks calculated; in this case \( n = 4 \)) would indicate complete preferred orientation in that plane. For the NB/NH sample, as seen with the XRD data, the crystalline structure was preferentially orientated in the (0 0 2), with a \( T_{c(002)} = 3.39 \). This value reduced to 2.66 in the B/NH sample, showing a subtle shift from the more columnar structure exhibited in the NB/NH sample. Further reductions of \( T_{c(002)} \) to 2.04 and 1.54 occurred for the B/150 and B/300 samples, respectively, demonstrating further shifts from preferred orientation in the columnar grown NB/NH sample, to more equiaxed structures as \( T_c \to 1 \).

3.1.4. X-ray photoelectron spectroscopy (XPS)

XPS quantification showed significant differences in the high-resolution Ti 2p, O 1s, and Na 1s spectra, as well as subtle alterations in peak intensity/presence, and elemental quantities. In the survey spectra (Fig. 5A), a significant difference from the control to all Ti coated samples was noted, with consistent peak positions and peak intensities of all Ti and O photoelectron emissions. Following NaOH treatment, peaks corresponding to Na were present with individual elemental maps of (C) sodium, (D) oxygen, (E) platinum (introduced from the FIB milling process), and (F) titanium.
detected (most notable being the Na 1s at ca. 1071 eV and Na KLL Auger peak at ca. 498 eV), with significant increase in intensity of the Ti LMM peaks, which overlap with Na 1s spectra.

Concerning the high-resolution spectra, noticeable differences in the O 1s peak (Fig. 5B) deconvolutions and subsequent positions were identified. The 316L SS control peak contained 3 constituent peaks located at 530.1 (39.0%), 531.6 (48.9%), and 533.1 (12.1%) eV, corresponding to O^{2-} (M^xO_y; M = Cr, Fe, etc.), OH^{-} (M^{3+}(OH)_{x}; M = Cr, Fe, etc.), and H_2O chemical states. For the Ti coated samples, the three constituent peaks remained with similar peak positions, however, their relative proportions differed; O^{2-} (79.1 ± 0.7%; Ti^{4+}O), OH^{-}/defective oxides (15.6 ± 0.5%; hydroxides/Ti^{3+}O/Ti^{2+}O), and H_2O (5.3 ± 0.5%). Following titanate conversion, an additional constituent peak appeared at ca. 535.2 eV, which matched the Na KLL Auger peak. Again all other peaks remained in their position at 530.0, 531.7, and 533.1 eV corresponding to O^{2-} (75.2 ± 0.4%; Ti^{4+}O), OH^{-} (11.6 ± 0.4%; hydroxides), and H_2O (5.3 ± 0.6%; experiencing a subtle shift). Variations in the relative intensities were noted as above for the Ti coated samples, with lower proportions of the OH^{-}/defective oxide (15.6 ± 0.5%) photoelectron peak.

For the Ti 2p doublet deconvolutions (Fig. 5C), there was a clear alteration in the number of constituent peaks following titanate conversion of the produced Ti films. Initially, the Ti coated samples contained Ti in 4 different valence states; Ti^{4+} (69.9 ± 1.5%; 2p3/2 = ca. 458.6 eV), Ti^{3+} (9.9 ± 0.5%; 2p3/2 = ca. 457.1 eV), Ti^{2+} (4.5 ± 0.2%; 2p3/2 = ca. 455.3 eV), and Ti^{0} (15.7 ± 1.1%; 2p3/2 = ca. 453.4 eV), representing different titanium oxide and hydroxide states (TiO, Ti_{2}O_{3} and TiO_{2}) as well as metallic titanium. Post-titanate conversion, the number of constituent peaks dropped to only contain Ti in its 4+ valence state (2p3/2 = ca. 458.4 eV).

In the high-resolution spectra of the range for Na 1s (Fig. 5D), a peak was present in both the pre- and post-titanate converted samples, however, this was due to the overlapping of the Ti LMM Auger peaks with the Na 1s spectra. The intensity of the peak increased for the titanate converted samples, demonstrating a peak at ca. 1071.4 eV, quantified as Na 1s (Na^+O; 56.6 ± 6.9%).

### 3.1.5. Raman spectroscopy

As seen in Fig. 6A, there are clear differences in peak positions from the pre-treated samples, to the titanate converted films. For all pre-treated samples, the peaks presented were all attributed
to TiO₂ with anatase phases in the lower Raman shifts (<300 cm⁻¹). Following NaOH treatment, alternate peaks at 285, 440, and 900 cm⁻¹, with broad peaks around 630–690 cm⁻¹ were delineated as a form of sodium hydrogen titanate (NaₓH₂-xTi₃O₇).

3.1.6. Fourier transform infrared spectroscopy (FTIR)

The FTIR spectrum for all converted titanate samples is shown in Fig. 6B. The unconverted Ti films did not present any discernible peak from the background measurements and, therefore, have not been shown. The broad band between 3000 and 3500 cm⁻¹ is attributable to fundamental O—H stretching groups of chemisorbed water. Additionally, there may be peaks consistent with Ti interaction with —OH groups contained within this broad peak. The peak at 1650 cm⁻¹ is also related to O—H bonding. The doublet peak located at 1410 and 1440 cm⁻¹ is characteristic of C—H deformations of adventitious carbon and the peaks present between 850 and 900 cm⁻¹ are attributable to Ti—O and Ti—O—Ti bonding, which when factored with the previously mentioned peaks points toward titanate formation.

3.1.7. Optical profilometry

To determine sample topography and roughness, optical profilometry was utilised. Sample roughness data via optical profilometry was detailed in Fig. 7. It is clear that the polished SS substrate exhibited a reduction in surface roughness (Sa = 129 nm for Control, which reduced to <40 nm) following sputtering of the Ti films. The roughness of the samples then increased significantly from ca. 23.5, 40.0, 27.9, and 35.8 nm to 140.6, 94.4, 90.4, and 52.6 nm, for the NB/NH, B/NH, B/150, B/300, and NB/NH_TC, B/NH_TC, B/300, B/150, BNH, NB/NH, NB/NH, B/300, and B/150 TC, B/300 TC samples, respectively, with the exception of B/300 (whose Sa only subtly increased), following titanate conversion (B/300 TC).
3.2. Mechanical testing

3.2.1. Pull off adhesion strength

Coating adhesion strength was determined using pull off adhesion testing. It was noted that through the application of a substrate bias (from NB/NH to B/NH), as well as the introduction of substrate heating (B/150 and B/300), the coating adhesion strength on the polished substrates decreased, with strengths of 42.2, 30.5, 14.5, and 5.7 MPa for the NB/NH, B/NH, B/150, and B/300 samples, respectively. Additionally, the incidence of full and partial coating failure increases with bias and substrate heating (Table 3 & Fig. 8). After subsequent titanate conversion, the coating adhesion strength significantly decreased compared to the unconverted samples, seeing a reduction from ca. 42 to 14 MPa for the NB/NH and NB/NH_TC samples, respectively. Similar trends are observed for all other titanate samples, with a clear reduction from the unconverted Ti samples to the titanate converted, for the same parameter type (Table 3).

3.2.2. Scratch adhesion testing

Scratch adhesion analysis was conducted as a complimentary technique to assess the relative failure loads and the associated failure modes of the produced films. It is clear there is a reduction in the required applied load to cause coating cracking and hence spallation from the NB/NH sample to the B/300 sample, as well as following titanate conversion for all sample types. Both the NB/NH and B/NH samples exhibited no spallation (Lc3) up to the applied 15 N load. However, the B/150 and B/300 samples exhibited spallation at ca. 2.9 and 1.6 N, respectively. Tensile cracking (Lc1) was observed for all sample types with loads of ca. 2.6, 2.2, 1.6, and 1.3 N for the NB/NH, B/NH, B/150, and B/300 samples, respectively, agreeing with the above trend (Table 4 & Fig. 9). Following titanate conversion, the samples were limited to ductile perforation leading to interfacial delamination. The trend between the samples is the same as that of the unconverted samples, with the NB/NH_TC sample demonstrating the highest load for spallation (Lc3; ca. 4.3 N) and the B/300_TC sample exhibiting the lowest (Lc3; ca. 1.0 N).

4. Discussion

4.1. Topographical, structural and compositional analysis

The ability to produce titanate surfaces on alternative materials through chemical conversion of Ti sputtered films, allows improved surface properties of bioinert materials, whilst also broadening the applicability of these surfaces. If appropriate bonding and optimisation of the produced films is met, these surfaces would potentially be utilised in a wide variety of applications, from hip stems, to fracture fixation devices based on metallic and non-metallic implant materials. Specifically, the broadening of this process to incorporate polymeric materials makes this technique advantageous for future biomedical applications.

By varying processing parameters, with the ultimate aim to produce denser, equiaxed films, the effects of oxygen penetration, coating microstructure and relative ingress of the NaOH solution on the formation mechanism of titanate can be explored. Structural dependency has yet to be elucidated and is critical to the results in this study. The application of a substrate bias from the NB/NH sample (5 A target current; 0 V substrate bias), to the B/NH sample (5 A target current, –100 V substrate bias), enables bombardment of the substrate by Ar⁺, with sufficient energy to transfer into the produced coating. This then allows reordering of the atoms into more equiaxed structures; akin to those of bulk metallic materials. However, such a process can also cause re-sputtering of the produced coating and, therefore, optimisation of the substrate bias must be employed. In this study, optimised biasing has previously been...
determined by TEER for the rig set-up used and investigated by Priyadashini et al, with –100 V showing significant grain refinement [18]. Furthermore, application of additional surface heating in the B/150 and B/300 samples (150 °C and 300 °C, respectively) aids in energy transfer, atomic re-ordering, and, hence, its crystalline structure [19,20]. Ultimately, this understanding was confirmed in the average void area/frequency (see supplementary data Fig. S1), XRD (Fig. 3) and subsequent texture coefficient analysis (Fig. 4), with the NB/NH sample exhibiting preferred orientation in the (0 0 2) plane of HCP Ti (PDF 00-044-1294), with a Tc

| Sample  | $S_a$ (nm) | $S_z$ (μm) |
|---------|------------|------------|
| Control | 129.7 (± 16.7) | 3.5 (± 1.5) |
| NB/NH   | 23.5 (± 4.3) | 1.3 (± 0.5) |
| B/NH    | 40.0 (± 9.6) | 3.0 (± 0.6) |
| B/150   | 27.9 (± 4.7) | 1.5 (± 0.9) |
| B/300   | 35.8 (± 8.4) | 3.3 (± 0.3) |
| NB/NH_TC| 140.6 (± 49.9) | 4.2 (± 0.7) |
| B/NH_TC | 94.4 (± 14.7) | 5.5 (± 1.4) |
| B/150_TC| 90.4 (± 37.3) | 6.5 (± 1.6) |
| B/300_TC| 52.6 (± 16.8) | 4.8 (± 0.4) |

Fig. 7. 3D visualisation and table of optical profilometry roughness values of surface topography for control, unconverted and titanate converted samples. $S_a$ and $S_z$ values represent arithmetic mean height and maximum height of surface features, respectively; n = 3.
of 3.39, with the highest measured average void area \((3.71 \pm 0.5) \times 10^{-3} \text{ m}^2\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}\), and void frequency \((4.1 \pm 0.2) \times 10^{-2} \text{ m}^{-1}## Table 3

Pull off adhesion data demonstrating average failure strength of both unconverted and converted Ti films \((n = 8)\). Additionally, the frequency of each failure mechanism has been detailed.

| Sample      | Failure Strength/MPa | Full Interfacial Failure | Partial Interfacial Failure | Adhesive Failure in the Epoxy |
|-------------|----------------------|--------------------------|-----------------------------|------------------------------|
| NB/NH       | 42.2 ± 3.5           | 0                        | 5                           | 3                            |
| B/NH        | 30.5 ± 3.7           | 2                        | 3                           | 3                            |
| B/150       | 14.5 ± 4.8           | 8                        | 0                           | 0                            |
| B/300       | 5.7 ± 0.5            | 1                        | 6                           | 1                            |
| NB/NH_TC    | 13.7 ± 2.0           | 4                        | 4                           | 0                            |
| B/NH_TC     | 10.8 ± 3.5           | 2                        | 3                           | 3                            |
| B/150_TC    | 8.1 ± 2.2            | 5                        | 3                           | 0                            |
| B/300_TC    | 3.0 ± 0.9            | 7                        | 0                           | 1                            |

Fig. 8. Optical microscope images and EDX elemental maps of sample surfaces and pull off stubs, respectively. Typical failure modes of full interfacial, partial interfacial, and full adhesive failure shown. C elemental maps are attributed to the epoxy adhesive, whilst Ti references the failed coating.

Table 4

Failure loads calculated from failure initiation of scratch testing track profiles. Error calculated from 5 samples, in accordance with ISO 20502:2016.

| Sample      | Tensile Cracking \((L_c1)/N\) | Buckling/Chevron Cracking \((L_c2)/N\) | Spallation \((L_c3)/N\) |
|-------------|------------------------------|----------------------------------------|--------------------------|
| NB/NH       | 2.6 ± 0.1                    | 9.9 ± 0.5                              | –                        |
| B/NH        | 2.2 ± 0.2                    | 6.5 ± 0.7                              | –                        |
| B/150       | 1.6 ± 2.9                    | –                                       | 2.9 ± 0.4                |
| B/300       | 1.3 ± 0.3                    | –                                       | 1.6 ± 0.5                |
| NB/NH_TC    | –                            | –                                       | 4.3 ± 0.4                |
| B/NH_TC     | –                            | –                                       | 2.3 ± 0.4                |
| B/150_TC    | –                            | –                                       | 1.4 ± 0.7                |
| B/300_TC    | –                            | –                                       | 1.0 ± 0.3                |

\[ r(\mu m) = \frac{0.0276 \cdot A \cdot Z_0^{0.67}}{\rho \cdot E_0^{0.89}} \]

where \(A\) is the atomic mass, \(Z\) is the atomic number, \(E_0\) is the electron gun voltage, and \(\rho\) is the material density, \(r = ca. 1.72 \mu m\) for Ti at 15 kV. From the XPS quantity measurements (also Table 2), the more columnar structures retained more Na (25.6 and 26.8 at.% for NB/NH_TC and B/NH_TC, respectively) in the top portion of the coating (X-ray penetration for XPS on the order of a few nm; slightly higher for nanoporous titanate [24]) compared to the external heated samples (16.1 and 17.8 at.% for B/150_TC and B/300_TC, respectively); the diffusion gradient was also evident in the FIB milled sample (Fig. 2). As discussed by Zaraté et al., due to the relative ratios of contained elements quantified from the XPS analysis, the titanate composition can be correlated with obtained values from EDX.
Na$_2$Ti$_2$O$_5$·H$_2$O = 17 at.% Ti, 50 at.% O, 16 at.% Na) was not possible, however, as reported by Zaraté et al., the produced titanate structure may be a mixture of closely related phases (Na$_2$Ti$_3$O$_7$; Na$_2$Ti$_3$O$_7$·nH$_2$O; Na$_2$Ti$_2$O$_5$·nH$_2$O), as confirmed via FTIR (Devi et al., Fig. 6B), XPS (Zaraté et al., Takadama et al. and others described in Takadama’s paper [27,28]; Fig. 5), XRD (Devi et al. [26]; Fig. 3), and Raman (Zhang et al. & Kasuga et al. [29,30]; Fig. 6A). XRD (Fig. 3) of the titanate converted samples exhibited 3 broad peaks at ca. 24.2, 28.4 and 48.2°, which agreed with Devi et al., for Na$_2$Ti$_3$O$_7$, however, there is a lack of clear peaks at ca. 33 and 36°, which may be due to the size of the nanocrystals/amorphous phases present, or the masking of the later peak in the (1 0 0) Ti diffraction peak [26]. A key feature is also noted with respect to the growth of the coating, since the portion of the Ti coating left on all samples following conversion is less than the original coating height (ca. 3.5–4 μm) produced, suggesting a two directional growth of titanate with the outwards growth being dominant; the conversion of titanium to titanate via diffusion results in expansion of the gel layer outwards, much like the expansion of a foam. converse to the original hypothesis that a columnar structure would produce a thicker titanate structure via increased diffusion of the NaOH solution, an alternative hypothesis has been proposed by the authors. As the Ti coating transitions from a columnar to a more equiaxed structure, the fraction of voids present between the crystals reduces (Figure S1). As stated above, the growth of these titanate structures, produced through chemical routes, occurs perpendicular to the surface. However, in the columnar films, perpendicular growth to the crystal surface can cause growth at more oblique angles due to the presence of the voids, resulting in a thinner titanate film. The B/300 sample, being the most equiaxed and therefore containing the lowest void fraction of the films tested, produced a thicker titanate structure. The proposed phenomenon above correlates well to the observed trends of decreasing void fraction (Fig. S1) and increased titanate thickness (Fig. 1) as the Ti coating shifts from columnar (NB/NH) to more equiaxed films (B/300).

The presence of TiO$_2$ peaks in the Raman (Fig. 6A) spectra were to be expected since the Raman spectrometer was utilised in reflectance, hence, the thin passivated TiO$_2$ layer (from ca. 20 Å to a few nm [31–33]) would have the highest intensity. Hanaor et al. described the formation of varying phases of TiO$_2$, particularly noting anatase occurs in pressures less than 20 kbar and below 600 °C [34]. Therefore, the region the sputtering rig is occupying is well within this range, hence, the formation of anatase phases in the Raman spectra is logical. The broad peaks around 415 and 605 cm$^{-1}$ are also in agreement with studies presented by Ocana et al. [35], Hsu et al. [36], and Exarhos [37]. It is worth noting the location of these peaks are similar to that of A$_{1g}$ and E$_g$ rutile modes, whereby Exarhos delineated that both have the same localised structural groups, however, with the additional absence of long-range order [37]. This is further noted in the XPS analysis (Fig. 5), since surface penetration is lower, whereby the produced surface layer contains multiple Ti valence states, corresponding to different titanium oxide forms (TiO$_2$, Ti$_2$O$_3$, and TiO). TiO$_2$ is the most abundant compared to Ti$_2$O$_3$ and TiO states, with the TiO being the least abundant due to thermodynamic favourability, as described by Hanawa et al. [38]. Once treated in the NaOH solution, conversion into sodium titanate was observed, with Raman peaks being noted at ca. 285, 440, 630–690, and 900 cm$^{-1}$ [39–41]. The peaks at 285 and 900 cm$^{-1}$ are in good agreement with intrinsic hydrogen titanate.
(H₂Ti₃O₇) bands found in a study conducted by Rodriguez et al. [42]. This may alternatively correspond to Na₂Ti₃O₇, as detailed by Bamberger et al., which revealed good agreement with the XPS and XRD analyses detailed above [43]. Furthermore, both Ma et al. and Kim et al. quantified the formation of a peak at 440 cm⁻¹, which is indicative of Ti–O bending vibrations, whereby the Ti and O atoms are 6- and 3-coordinated, respectively [44,45]. The broad peak in all TC samples ranging from 630 to 690 cm⁻¹, is characteristic of [TiO₆] octahedral Ti–O–Ti vibrations found in titanate structures, as characterised by Kasuga et al. and Zhang et al. [29,30]. These results are further confirmed from the complimentary FTIR measurements (see Fig. 6B), where Ti–O and Ti–O–Ti bonds were detected at ca. 850 and 900 cm⁻¹, respectively.

Cumulatively, the compositional and morphological analysis confirmed the titanate conversion of all Ti films, irrespective of the structure factor present, with the more equiaxed structure (B/300_TC) producing the thickest titanate (ca. 1.6 μm), demonstrating the distinct effect of the structure factor on the depth of titanate penetration.

4.2. Mechanical testing

A key consideration for any biomaterial material, which has been coated for improved surface properties, is the adherence of the produced coating, since application-dependant environments may cause fretting, micro-motion, and delamination. As determined via pull-off adhesion testing (Fig. 8 & Table 3), as well as coating scratch testing (Fig. 9 & Table 4), the more columnar NB/NH samples exhibited the greatest adhesion strength on the polished SS 316L samples (ca. 42.2 MPa, comparable to the FDA’s minimum requirement of 50.8 MPa [46]), and despite cracking, did not spall during the scratch testing analysis (Fig. 9 & Table 4), proving good adhesion even on polished substrates. However, following the application of a substrate bias and substrate heating, the failure strength and spallation forces are greatly reduced. As discussed by Shen et al. (in relation to W coatings), ion-bombardment plays a significant role in the stress and microstructure of the produced coating [47]. The production of residual stresses within the coating, occurring via an ‘atomic shot peening’ effect, are higher in the denser, less columnar films, with more columnar films being unable to support large internal stresses [47]. Following titanate conversion, the respective films experienced a reduction in the adhesion strength and failure loads, which is possibly due to ingress of NaOH solution under the films, converting the bottom portion of the coating, which is exacerbated by the smooth nature of the substrate to which the coating is applied.

Despite the mechanical properties presented, the coating adhesion should increase if applied to roughened surfaces, more akin to those found on orthopaedic implants, due to the mechanical interlock present [48]. Furthermore, excessively thick coatings would not be used in practical purposes since, as determined by Vega-Morón et al. [49], the thicker the films used, the weaker the adhesion strength of the produce sputtered coating. Hence, it is clear further studies into the optimisation of these surfaces in terms of structure and thickness are needed, as well as an in-depth view on cellular proliferation/adherence and integration into surrounding tissue in in vitro and in vivo tests, respectively.

5. Conclusions

Presently, literature findings for biomedical titanates have been restricted to Ti-based substrate materials [7,8,50,51]. Despite extensive innovation and success within this research avenue, the ability to translate titanate structures to alternative materials for biomedical applications has yet to be realised within the literature. There is a clear need to produce either bioactive or antimicrobial properties on a wide range of material types for a plethora of biomedical applications. The above results have demonstrated, in detail for the first time, that translation of titanate structures onto alternative surfaces through a combinatorial DC magnetron sputtering and chemical conversion approach is possible, therefore, broadening the applicability of these surfaces for biomedical applications.

The original hypothesis of the work conducted was that variation of the structural properties of DC magnetron sputtered films would result in a higher titanate conversion percentage for columnar grown films compared to more equiaxed counterparts. However, the findings demonstrated a converse trend, whereby more equiaxed films produced the thickest titanate structures, due to the reduction in void fraction and area negating oblique angle growth of the titanate structure, which was characterised in the columnar grown films. The combination of XRD and texture coefficient analysis demonstrated the successful production of graded films from columnar, with preferred orientation in the (0 0 2) plane (NB/NH), to those more akin to equiaxed titanium (B/300). Furthermore, the titanate thicknesses produced ranged from 1.12 to 1.63 μm; thicker than currently found in the literature [51].

Optical profilometry further confirmed the increased surface roughness (Sa) values from the non-titanate converted samples (ca. 23–40 nm) to the titanate converted samples (ca. 52–140 nm). Furthermore, there was a clear trend noted in the film adherence through both the pull-off tests, as well as the scratch adhesion tests. The NB/NH sample types exhibited the greatest film strength (ca. 42 MPa), with decreasing values noted for the higher T value films (B/300 had reduced to ca. 6 MPa). A similar trend was also noted for the titanate converted samples, with the titanate conversion itself resulting in a decrease in film adherence compared to the native sputtered film (e.g. the NB/NH and NB/NH_TC samples exhibited pull-off failure strengths of ca. 42 and 14 MPa, respectively). With careful substrate surface modification through sandblasting or interlayer generation, the adhesion could be improved for these coatings making them viable for biomedical applications; currently the FDA adhesion strength standard for HA sputtered films is 50.8 MPa [46].

Currently, cellular studies are being conducted to further understand the biological impact of such coatings and to assess their efficacy for orthopaedic applications. Additional investigations into the adhesion strength of such films through the application of various interlayers and/or increased surface roughness are also being conducted, with a view to apply the titanate structures onto biodegradable materials. Ultimately, the proof of successful conversion of sputtered Ti films into titanate should open up a wide variety of material types and applications for these structures.

CRediT authorship contribution statement

Matthew D. Wadge: Conceptualization, Methodology, Validation, Investigation, Formal analysis, Resources, Data curation, Writing - review & editing. Burhan Turgut: Validation, Investigation, Formal analysis, Data curation, Writing - review & editing. Bryan W. Stuart: Conceptualization, Methodology, Supervision, Writing - review & editing. Reda M. Felfel: Conceptualization, Supervision, Research, Project administration, Writing - review & editing. Ifty Ahmed: Conceptualization, Supervision, Project administration, Writing - review & editing. James W. Murray: Investigation, Formal analysis, Data curation, Writing. Writing - review & editing. David M. Grant: Conceptualization, Methodology, Supervision, Funding acquisition, Resources, Project administration, Writing - review & editing.
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

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