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Anisotropy, band-to-band transitions, phonon modes, and oxidation properties of cobalt-oxide core-shell slanted columnar thin films

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Highly ordered and spatially coherent cobalt slanted columnar thin films (SCTFs) were deposited by glancing angle deposition onto silicon substrates, and subsequently oxidized by annealing at 475 °C. Scanning electron microscopy, Raman scattering, generalized ellipsometry, and density functional theory investigations reveal shape-invariant transformation of the slanted nanocolumns from metallic to transparent metal-oxide core-shell structures with properties characteristic of spinel cobalt oxide. We find passivation of Co-SCTFs yielding Co-Al2O3 core-shell structures produced by conformal deposition of a few nanometers of alumina using atomic layer deposition fully prevents cobalt oxidation in ambient and from annealing up to 475 °C. © 2016 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4941399]

Transition metal oxides constitute an interesting class of materials in solid state physics, with numerous attributes that span dielectric, semiconducting, ferromagnetic, and ferroelectric properties. Much research effort has focused on this vast and still widely uncharted field of oxide compounds. In addition to their unique and useful bulk properties, in recent years, strong interest has developed in shaping these materials into topographies with nanoscale dimensions. One example is core-shell nanostructures whose optical, electrical, and magnetic properties are attractive for novel device architectures in applications such as solar cells and sensors.1–5 Nanostructures and core-shell structures that incorporate the transition metal semiconductor cobalt oxide have been shown to be effective as anodes with largely increased surface area in lithium ion batteries, biosensors, and in electrochemical catalysis.6–11 This is due to the wide tunability of electrical and magnetic properties as a function of temperature and stoichiometry. Oxidation of cobalt may result in different forms depending on growth conditions. Rock salt structure cobaltous oxide (CoO) is known to be present at high temperatures from decomposition of cobaltic oxide (Co2O3) or spinel structure mixed-valence compound CoIIICoIII O4 (Co3O4) while at lower temperatures increased oxygen absorption produces Co3O4.12–14

In this paper, we report on fabrication and characterization of highly regular and coherently arranged few-nanometer-sized columnar core-shell structures consisting of metal cores and metal-oxide shells (Figs. 1(b)–1(d)). Specifically, we investigate Co nanocolumns and how their optical, vibrational, and electronic properties change upon oxidation. We find that cobalt nanocolumns can be almost completely transformed into Co3O4 while fully retaining their shape. Thereby, we have produced highly anisotropic spatially coherent cobalt oxide nanocolumnar thin films, which may be useful for optical and/or electrochemical device architectures. We further find that passivation of cobalt nanocolumns by an ultra-thin conformal overlayer of alumina fully inhibits oxidation of the cobalt nanocolumns at normal ambient as well as during our annealing procedure similar to our previous results with conformal coating of a few layers of graphene.15–17

Highly ordered, spatially coherent slanted columnar thin films (SCTFs) can be fabricated by glancing angle deposition (GLAD), which utilizes particle flux at oblique incidence angles.18 These films exhibit extreme birefringence and dichroism properties that can be tailored by choice of deposition parameters.19 An ultra-thin and conformally overgrown passivation layer via atomic layer deposition (ALD) can be employed to adjust physical and chemical surface properties of SCTFs.20–22 ALD is a chemical vapor deposition process which provides precise, self-limiting, monolayer growth of materials by cycling a combination of precursor, ozone, or plasma.

Mueller matrix generalized ellipsometry (MMGE) determines the anisotropic optical properties of SCTFs by measurement and model analysis of the 4 × 4 Mueller matrix M, where M describes the change in polarization of light after interaction with a sample (Fig. 1(a)).23,24 An appropriate physical model then allows for accurate description of the optical properties of the film. The dielectric function of a uniform flat film may be determined using a homogeneous isotropic layer approach (HILA). For anisotropic films such as SCTFs, the dielectric functions along all three major polarizability axes Nα, Nβ, and Nγ (Fig. 1(e)) may be obtained using a homogeneous biaxial layer approach (HBLA). This method assumes that the material is homogeneously distributed along each distinct axis and does not allow for determination of material constituents.25 For columnar core-shell structures, the anisotropic Bruggeman effective medium approximation (AB-EMA) can be used to model the dielectric function, εeff(ω), of all three major polarizability axes, j = a, b, c, from a number of “bulk” dielectric functions in addition to the constituent fractions, fj, according to
\[ \sum_{n=1}^{m} f_n \frac{\varepsilon_n - \varepsilon_{\text{eff},j}}{\varepsilon_{\text{eff},j} + L_j^D (\varepsilon_n - \varepsilon_{\text{eff},j})} = 0, \]  

(1)

with the requirements that \( \sum f_n = 1 \) and \( \sum L_j^D = 1 \), where \( L_j^D \) are the depolarization factors along the major polarizability axes, \( N_a, N_b, \) and \( N_c \). For the AB-EMA model, we assumed three different constituents: void, cobalt, and oxide, either the well documented \( \text{Al}_2\text{O}_3 \) deposited by ALD or the as yet to be described cobalt oxide. The optical constants of the cobalt SCTF constituent were determined from the as-deposited sample immediately after deposition, where there was assumed to be no oxide material present. These were then kept constant during the search for the unknown oxide optical constants of the core-shell structures. Furthermore, we previously reported that SCTFs exhibit monoclinic properties, which can be accounted for by defining an angle \( \beta \) between polarizability axes \( N_b \) and \( N_c \) (Fig. 1(e)). During best-match model calculations, model parameters are varied in order to calculate data that matches the experimental data as accurately as possible. Values of the complex dielectric function of an unknown material can be found by a wavelength-by-wavelength regression analysis. Electronic band-to-band transitions cause critical point (CP) features in the dielectric function spectra. The imaginary contribution, \( \varepsilon_2(E) \), can be modeled conveniently using Gaussian lineshapes with best-match parameters amplitude, \( A \), center energy, \( E_n \), and broadening, \( B_r \), and the real contribution, \( \varepsilon_1 \), is obtained from Kramers-Kronig transformation.

\[ \varepsilon_2(E) = A \left( e^{-\left(\frac{E-E_n}{B_r}\right)^2} - e^{-\left(\frac{E-E_n}{B_r}\right)^2} \right), \]  

(2)

\[ \sigma = \frac{B_r}{2\sqrt{\ln(2)}}, \]  

\[ \varepsilon_1(E) = \frac{2}{\pi} P \int_0^\infty \frac{\xi \varepsilon_2(\xi)}{\xi^2 - E^2} d\xi. \]  

(3)

Raman spectroscopy was used to differentiate between the cobalt oxide phases; however, it has been shown that at high excitation laser energy in normal ambient, \( \text{CoO} \) can further oxidize to \( \text{Co}_3\text{O}_4 \). Care must be taken to use a low enough intensity so as to not change the material investigated. Near the Brillouin zone center \( (k = 0) \) of \( \text{Co}_3\text{O}_4 \) (space group \( \text{O}_h \)) normal modes are given from \( \Gamma = A_{1g} + E_g + 3F_{2g} + SF_{1u} + 2A_{2u} + 2E_u + 2F_{2u} \), where the \( A_{1g}, E_g, \) and the triple degenerate \( 3F_{2g} \) are the only Raman active modes.

Density functional theory (DFT) calculations of the Raman spectrum of \( \text{Co}_3\text{O}_4 \) were performed using plane-wave code Quantum Espresso (QE). Local density approximation functions by Perdew and Zunger (PZ) and norm-conserving pseudopotentials from the QE library were used, with the cut-off for the electronic wave-function set at 80 Ry, and a \( 4 \times 4 \times 4 \) Monkhorst-Pack grid shifted by half of the simulation cell for the Brillouin-Zone integrations. The primitive unit cell was first relaxed in order to reduce forces on the ions. The system is considered to be at equilibrium when the forces on the ions were approximately \( 1.0 \times 10^{-5} \) Ry/bohr \((-0.0003 \text{ eV/A})\). The phonon frequencies and Raman activities were...
computed at the Γ point for the relaxed structure using density-functional perturbation-theory for phonons\textsuperscript{34} and the second-order response for Raman activities.\textsuperscript{35} Cobalt SCTF were deposited on Si (100) substrates using GLAD as described previously,\textsuperscript{25} and at room temperature under ultra-high vacuum with an oblique angle of incidence of 85°. Half of the samples were passivated by depositing a conformal layer of Al\textsubscript{2}O\textsubscript{3} using documented ALD techniques\textsuperscript{20,21} by subsequent cycling of trimethylaluminum (TMA) and 18.3 M deionized water (Fiji 200 ALD Reactor, Cambridge NanoTech). The samples were held at 150°C temperature under vacuum, and 55 cycles were deposited with a rate of approximately 0.9 Å/cycle.\textsuperscript{20} All samples were characterized immediately after deposition using SEM and MMGE to determine initial thickness, column diameter, and optical properties. MMGE data were measured in the spectral range of 300 to 1650 nm at four angles of incidence, \( \phi = 45°, 55°, 65°, \) and 75°, and over a full azimuthal rotation of the sample by 6° increments (RC2, J. A. Woollam Co, Inc.). All samples were then placed flat on a sample mount plate under high vacuum and heated gradually to 475°C by increments of 25°C every 30 min, held at 475°C for 2 h, then suddenly exposed to normal ambient and allowed to cool to room temperature. SEM and MMGE investigations were then repeated, and Raman spectroscopy was performed with a 532 nm excitation laser (DXR Raman Microscope, ThermoScientific).

Cross-sectional SEM images are presented in Figs. 1(b)–1(d). Approximations for the overall thickness of the SCTF (t), as well as the slanting angle (\( \theta \)) and diameter of the posts (d) are summarized in Table I. Upon annealing the column diameter of the non-passivated SCTF increases approximately twofold due to oxide growth on all exposed surfaces, while the column diameter of the passivated sample is approximately 5 nm larger than that of the as-grown columns due to the alumina overgrowth. Otherwise, no significant structural changes occur upon annealing or passivation.

Fig. 1(a) compares MMGE data versus sample rotation with HBLA best-match model calculated data for as grown, non-passivated annealed, and passivated annealed SCTFs. It can be seen that the data from the passivated annealed SCTF very closely resemble those of the as grown SCTF, signifying that both SCTFs remained nearly structurally and optically identical. While data from the non-passivated annealed SCTF reveal some deviation from the former two, major anisotropic properties of the SCTFs remained similar upon oxidation. This is seen by observing where the zero points in the off-block diagonal elements in the Mueller Matrix data in Fig. 1(a) occur, in which the slanted columns are aligned within the plane of incidence causing pseudo-isotropic points.\textsuperscript{25} For all SCTFs, pseudo-isotropic points occur at the same azimuthal orientations. Best-match AB-EMA model parameters are summarized in Table I. Before annealing, the SCTF is assumed to have no oxide formation yielding a void fraction of about 75%. It is observed that after annealing, the original metal SCTF is almost fully transformed into metal oxide forming a shell around a reducing core with remaining diameter of

<table>
<thead>
<tr>
<th>Parameter</th>
<th>As grown</th>
<th>Co-Co\textsubscript{3}O\textsubscript{4} Annealed</th>
<th>Co-Al\textsubscript{2}O\textsubscript{3} Annealed</th>
</tr>
</thead>
<tbody>
<tr>
<td>t (nm)</td>
<td>99(9)</td>
<td>97(8)</td>
<td>98(4)</td>
</tr>
<tr>
<td>( \theta ) (°)</td>
<td>64(4)</td>
<td>63(4)</td>
<td>64(3)</td>
</tr>
<tr>
<td>d (nm)</td>
<td>17(2)</td>
<td>26(6)</td>
<td>22(4)</td>
</tr>
<tr>
<td>t (nm)</td>
<td>83.4(1)</td>
<td>82.9(1)</td>
<td>97.2(1)</td>
</tr>
<tr>
<td>( \beta ) (°)</td>
<td>61.4(1)</td>
<td>61.7(1)</td>
<td>64.2(1)</td>
</tr>
<tr>
<td>Co %</td>
<td>79.6(1)</td>
<td>83.2(1)</td>
<td>77.5(1)</td>
</tr>
<tr>
<td>Co\textsubscript{3}O\textsubscript{4} %</td>
<td>24.9(1)</td>
<td>2.3(1)</td>
<td>27.6(1)</td>
</tr>
<tr>
<td>Al\textsubscript{2}O\textsubscript{3} %</td>
<td>...</td>
<td>42.6(1)</td>
<td>...</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>16.8(2)</td>
<td>...</td>
</tr>
</tbody>
</table>

\textbf{FIG. 2.} Optical constants along major polarizability axes \( N_\alpha, N_\beta, \) and \( N_\gamma \) of non-passivated (a) and (b), non-passivated annealed (c) and (d), and Al\textsubscript{2}O\textsubscript{3} passivated annealed (e) and (f) cobalt SCTFs. Critical point transition energies, identified from the subsequent AB-EMA analysis, are indicated by vertical lines.
approximately 2 nm still above our best-match model parameter uncertainty range (Table I). For Co₃O₄, the density is less than that of pure cobalt; thus, its formation results in an increase in column diameter and a reduction of the void fraction to approximately 55%, from which we suggest a bidirectional oxide growth. Fig. 2 depicts the anisotropic optical constants (refractive index and extinction coefficient). The optical constants along Nₓ, Nᵧ, and Nₗ change between the as-grown and the non-passivated annealed SCTF, while those of the passivated annealed SCTF remain nearly identical to those of the as-grown SCTF. The oxidized cobalt SCTF exhibits strong changes in birefringence and dichroism properties, transforming from highly absorbing (metal-like) to highly transparent (dielectric-like). For polarization along axis Nₓ, the optical constants are typically found nearly identical to the bulk optical constants of the column material, here bulk cobalt. For annealed non-passivated SCTF, we thus indicate the transformation to Co₃O₄ because the optical constants for axis Nₓ are nearly identical to bulk Co₃O₄. The AB-EMA dielectric constants for axis Nₓ are compared to those given by Athey et al., which were obtained using the HILA model for data from a Co₃O₄ thin film deposited using 12 s spray pyrolysis onto soda-lime float glass, included in Fig. 3. These results show close agreement on magnitudes as well as absorption band locations yielding further confirmation that the cobalt oxide is in the form of Co₃O₄. Our CP parameter analysis results yield optical constants for axis Nₓ which are compared to those reported previously in Table II of Raman spectra of the non-passivated annealed SCTF with DFT calculated Co₃O₄ Raman intensities are shown in Fig. 4. We identify peaks that correspond to modes $F_2$, $E_g$, $F_2$, and $A_{1g}$ at 192 cm⁻¹, 488 cm⁻¹, 521 cm⁻¹, 625 cm⁻¹, and 694 cm⁻¹, respectively. We note that the peak at 521 cm⁻¹ is caused by the silicon substrate and the Co₃O₄ peak at 488 cm⁻¹ is subsumed as a shoulder. We also note that the Co₃O₄ peak at 521 cm⁻¹ is not visible as it appears at the same frequency as the strong silicon peak. Surface-enhanced scattering effects cause the secondary Si modes, 2TA at 304 cm⁻¹ and 2TO at 969 cm⁻¹, which pronounce upon oxidation due to the increased SCTF transparency. Fig. 4 also shows examples of mode displacement vectors for $A_{1g}$, $E_g$, and triple-degenerate $3F_2$ modes. We find very good agreement between the experimental and DFT predicted Raman intensities as well as with literature for Co₃O₄. We therefore conclude that the annealing process of non-passivated Co-SCTF leads to nearly full oxidation forming Co-Co₃O₄ core-shell structures. The passivated annealed SCTF does not reveal any mode related to cobalt oxide. We conclude that alumina passivation prevents the Co-nanocolumns from oxidation regardless of thermal annealing up to 475 °C, while a small diameter increase is caused by the alumina passivation layer.

In summary, we investigated the changes in optical anisotropy, the emergence of band-to-band transitions, and phonon modes upon the thermal oxidation of cobalt slanted columnar thin films. Scanning electron microscopy, Raman scattering, generalized ellipsometry, and density functional theory investigations reveal a shape-invariant transformation of cobalt columns from metallic to transparent metal-oxide core-shell structures with properties characteristic of spinel cobalt oxide. The anisotropic optical constants derived for the as-grown and non-passivated annealed cobalt columns

![Graph](image.png)

**Fig. 3.** AB-EMA “bulk” oxide constituent model dielectric function of non-passivated annealed Co-SCTF along with Gaussian curves from CP analysis. A HILA model dielectric function of CoO₃ obtained from data taken on a soda-lime float glass after a 12 s pyrolytic spray deposition by Athey et al. (Reproduced with permission from Athey et al., J. Vac. Sci. Technol. A 14, 685 (1996), American Vacuum Society). Critical point transition parameters of Co₃O₄ obtained in this work from MMGE data analysis of oxidized cobalt SCTF in comparison with selected data reported in literature.
reveal transformation from metal-like to dielectric-like suggesting highly transparent properties of the cobalt oxide nanostructures. From structural investigations, we propose that oxide grows bidirectionally, leaving only a small fraction of a cobalt core as the remaining void fraction diminishes proportionally upon oxide formation. In contrast, we observe insignificant structural and optical changes of the oxide overcoat. We conclude that alumina passivation provides an efficient oxygen barrier onto cobalt nanostructures, which may find use in design of device architectures including cobalt-cobalt oxide core-shell nanostructures for applications in normal ambient.

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41See http://www.xcrysden.org for XCrysDen used for cell and mode rendering in Fig. 4; See also A. Kokalj, *Comput. Mater. Sci.* 28, 155 (2003).