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Reactive pulsed direct current magnetron sputtering deposition of semiconducting yttrium oxide thin film in ultralow oxygen atmosphere: A spectroscopic and structural investigation of growth dynamics



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ABSTRACT

An experimental investigation was conducted to explore spectroscopic and structural characterization of semiconducting yttrium oxide thin film deposited at 623 K (\pm 5K) utilizing reactive pulsed direct current magnetron sputtering. Based on the results obtained from both x-ray diffraction and transmission electron microscope measurements, yttrium monoxide is very likely formed in the transition region between β -Y₂O₃ and α -Y₂O₃, and accompanied by the crystalline Y₂O₃. Resulting from either the low energy separation between 4d and 5s orbitals and/or different spin states of the corresponding orbitals' sublevels, the stability of monoxide is most presumably self-limited by the size of the crystal in thermodynamic terms. This behavior develops a distortion in the structure of the crystal compared to the metal oxide cubic structure and it also effectuates the arrangement in nanocrystalline/amorphous phase. In addition to this, spectroscopic ellipsometry denotes that the semiconducting yttrium oxide has the dominant, mostly amorphous, formation character over crystalline Y₂O₃. Our purpose, by means of the current findings, is to advance the understanding of formation kinetics/conditions of yttrium with an unusual valency (2+).

1. Introduction

Because of their unusual valence band electronic configuration (4f $^{n-1}$ 5d $^{0, 1}$ 6s 2 ; n = 1-15), rare earth elements (REE) have become, over the last decades, one of the major interesting research subjects in science and crucial components in high-tech industry/products such as catalysis field (e.g. Ce) [1], hybrid engines (e.g. Dy) [2], neutron detection (e.g. Sm, Eu) [3,4], and fusion reactors/nuclear plants (e.g. Y) [5-7]. 3+ is the most common oxidation number exhibiting three phases as cubic (α phase), monoclinic (β phase) and hexagonal (γ phase), whereas the oxidation states of rare-earth metal could vary between 2+ and 4+ [8-13]. The aforementioned crystals exhibit total five multiform alterations in accordance with the ionic radius of REE and the temperature [10]. Arising from the multifaceted nature of trivalent oxide of Y structures (Y₂O₃) such as high melting point (~2700 K), high optical excitation threshold (~5.6 eV), high relative permittivity (~16) and mechanical and chemical stability, previous studies have primarily concentrated on the dielectric/catalytic properties [14-20]. Because the rare-earth elements that have the valence of 2+ exhibit an exclusive behavior, the acquisition of monoxide rare-earth metals has started to attract the attention of many research groups [21-23]. Putting particular emphasis on three analyses, which are the study fields of the valence fluctuating state of pulsed laser evaporated SmO [24], the investigation of the first-principle study of superconductivity in LaO [25] and the first-principle calculations on superconductor LaO [26], may express the importance of the entire topic. Besides that, the first serious discussions and analyses of divalent yttrium (YO) in solid-phase, emerged in 2016 by Kaminaga et al. [27]. To the best of the authors' knowledge, no report on divalent yttrium oxide deposition by reactive pulsed-DC magnetron sputtering has been found. This study was arisen from the lack of research on the oxidation dynamics of yttrium. The uniqueness of this study exists in the fact that first-time solid phase divalent yttrium (YO) was produced by reactive pulsed-DC magnetron sputtering. And for the first-time, refractive index dispersion of YO and Y2O3 mixture is obtained. One of the long-term implications of this study will impact the understanding of how oxide dispersion strengthened

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(ODS) nanoparticles are formed in fusion reactors [28,29].

2. Experiment

In the current research, reactive pulsed-DC magnetron sputtering and e-beam evaporator (for reference Y metal thin film deposition) attached to the multi-functional vacuum cluster tool, which was kept at a high vacuum level (~ 6.00×10^{-5} Pa) during the entire procedure, were operated in ISO 8 class clean room environment. The deposition chambers were baked out for 72 h at 368 K (\pm 5K). In addition to this, for the case of high-temperature deposition, the substrates/substrate holder and close surrounding of the holder were heated up to 673 K (\pm 5K) for 6 h. The substrates were cleaned in the ultrasonic bath with acetone, isopropanol, and DI water, respectively (15 min each) and dried with N2 gas. The samples were loaded and unloaded to/from the chambers using the robotic arm without disturbing the experiment environment. During the magnetron sputtering, the substrates were located 13 cm away from the target. The angle between the target and the substrate was fixed between 45 and 50°. The hysteresis loop of the reactive process, in the range of 0-6 sccm, was characterized in which the discharge was stabilized for 5 min in each step (Fig. 1). The plasma optical emission spectra (200-1100 nm) were measured by a plasus emicon mc spectrometer using the optical fibre probe which overlooks the discharge parallel ≈ 2 cm above the target surface. Residual and process gases (P_{Ar} , P_{O_2} , P_{H_2} , P_{H_2O}) were monitored (Table 4) by means of the process eye professional residual gas analyzer (mks instruments). The deposition of yttrium/yttrium oxide thin films, approximately 400 nm in thickness, were carried out using the metallic (% 99.99 purity) target onto Si (001) and soda-lime glass substrates. The deposition of the films was performed towards the increasing direction of oxygen flow demonstrated into the hysteresis loop (Fig. 1). The partial oxygen pressure was varied between 2.31 \times 10^{-3} and 2.13 \times 10^{-6} Pa (~5.33 \times 10^{-1} Pa, total deposition pressure). During the e-beam evaporation, the substratetarget distance was fixed as 26 cm. The metal thin films' deposition (approximately 300 nm in thickness), at 298 K (\pm 5K) was carried out using the metallic (% 99.99, purity) yttrium pieces onto Si (001) and soda-lime glass substrates. The partial pressure of oxygen (residual gas) was detected $\sim 2.00 \times 10^{-7}$ Pa. The vacuum level/atmosphere is controlled by throttle valve which is located between turbomolecular pump (TMP) and process chamber and by inletting process gasses to the chambers in both cases (magnetron sputtering and e-beam evaporation). The substrates were rotated by 10 rpm in both deposition methods. No post-treatment was applied to the thin films. Growth parameters are

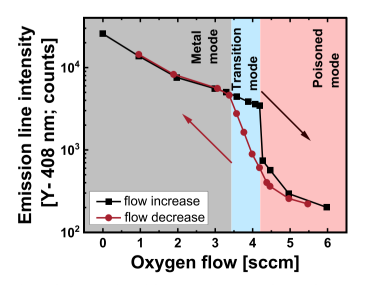


Fig. 1. The hysteresis loop of the reactive deposition in the range between 0 and 6 sccm.

given in detail, in Table 4.

Structural investigation of the thin films was carried out by a) X-ray diffraction: RIGAKU MiniFlex 600 benchtop X-ray diffractometer with Cu K_{α} X-ray source. The measurements were carried out between 5 and 60° (20) by 0.0050° steps with 0.0025° resolution (with the Bragg-Brentano theta-2theta diffraction geometry). 11,001 data points were collected for per measurement (scan speed 1deg/min). b) Transmission Electron microscopy/SEM: A high-resolution scanning electron microscope/focused ion beam (SEM-FIB) system Helios 5 UX (Thermo Scientific) was operated at 2 kV using the Through-the-lens detector (TLD). For lamella preparation, samples were sputter-coated with a 30 nm thick gold layer followed by a 300 nm thick FIB deposited Pt layer to protect the surface from ion beam damage. The extracted lamellas were further analyzed by the transmission electron microscope (TEM, Tecnai G20, FEI) operated at 200 kV. Optical characterizations of the films were performed employing spectroscopic ellipsometer (Woollam RC2) in the spectral range from 210 to 1690 nm (or from 5.9 to 0.7 eV), and spectrometer (Agilent, Cary 7000) in the spectral range from 200 to 2500 nm. The thickness of the films was measured by profilemeter (Vecco dektak 150) and spectroscopic ellipsometer (Woollam RC2). The temperature-dependent electrical measurements between 90 and 350K, were carried out by Hall measurement system (Ecopia 5000). Chemical state analyses were carried out employing X-ray photoelectron spectroscopy (Escalab 250XI, ThermoFisher). The base pressure, with the charge neutralizer switched on during spectra acquisition, was better than 10⁻⁵ Pa achieved by rotary and turbomolecular pumps. The calibration and linearity of the binding energy scale was confirmed by measuring the positions of Ag $3d_{5/2}$, Au $4f_{7/2}$ and Cu $2p_{3/2}$ to be at 368.21 eV, 83.93 eV and 932.58 eV, respectively. The FWHM of Au 4f_{7/2} peak was better than 0.58 eV. The samples were loaded into the XPS chamber under the laboratory conditions (~300K, 1.013 \times 10⁵ Pa (1atm)). The samples were sputter-cleaned, Ar⁺ gun with the ion energy 2.0 keV was used for 30 s before the measurements. The size of the cleaned area was 2 \times 2 mm; the incidence angle was 30° with respect to the normal of the surface. The size of the analyzed sample area was 650 \times 100 µm. The excitation source was monochromated Al K_a radiation (h ν = 1486.60 eV), operating water-cooled anode at 150 W. The recorded spectra involve survey, core level, and valence band spectrum. The spectra were acquired sequentially. Complementary work function measurements were performed by UPS with monochromated He-I (see the supplementary material). The binding energy calibrations of adventitious carbon (C_{1s}) of the related samples were performed with respect to the vacuum level, as suggested by G. Greczynski and L. Hultman [30] (see the supplementary material) and all the spectra were shifted, accordingly. In addition, the depth profiles of H atoms were measured by secondary ion mass spectrometry (SIMS) using Cameca IMS 7f microanalyzer. The measurements were performed with the 15 keV Cs⁺ primary beam rastered over the 200 \times 200 μ m² area and only the central part of the crater was used to collect the SIMS signal. The depth conversion of the recorded profiles was performed by measuring the sputtered crater in the films assuming a constant erosion rate. The vibrational spectroscopy analyses were performed by a vacuum Fourier transform infrared spectrometer (Bruker Vertex 80v) with 4 cm⁻¹ spectral resolution.

3. Results and Discussion

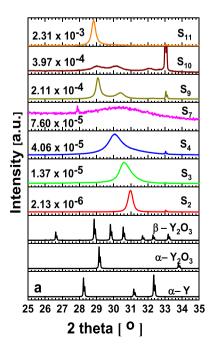
3.1. Structural characterization

This research is primarily separated into two sections. In the scope of the first step, the structural transformation of yttrium/yttrium oxide thin films deposited at 298K (\pm 5K), is investigated basically. The peaks detected for **S**₂, **S**₃, and **S**₄ samples that show metallic type electrical conductivity at approximately 31° (20) and the small peak (which is almost not visible for **S**₃) at approximately 33° (20), point out that metallic yttrium crystal arrangement is most probably accompanied by

 β -Y₂O₃ (Fig. 2a). Although, as the partial oxygen pressure level increases from 2.13×10^{-6} up to 4.06×10^{-5} Pa, the contribution from the oxide phase becomes dominant (while the peak located at approximately 31° (2 θ) shifts towards 30° (2 θ), the intensity of the peak at approximately 33° (20) increases); the thin films obtained, represent the same type electrical conductivity (Fig. 11). In the region between 2.11×10^{-4} and 3.97×10^{-4} Pa (the related samples are \textbf{S}_{9} and $\textbf{S}_{10}), \, \beta\text{-}Y_{2}O_{3}$ is quite likely formed along with α -Y₂O₃. On the other hand, the single peak detected for sample S_{11} could be responsible for simultaneous formation of α and β -Y₂O₃ or only α -Y₂O₃ (the slight shift of diffractogram(s) to lower 20 values can be observed compared to the reference peak positions given). In the region $\sim 6 \times 10^{-5}$ Pa (S₇), the film develops an amorphous structure. The deposition rates of the thin films are given in Fig. 2b. The results obtained from XRD revealed that, even at low pressure levels, yttrium tends to form oxide thanks to its high oxygen affinity. In the second step (which is the core of this article), in order to understand the effect of temperature and/or partial oxygen pressure level on the formation dynamics (phase kinetics), we deposited thin films at high temperature (\sim 673 K) with similar partial oxygen pressure levels used in the amorphization and in fully oxidized regions. And we used metallic yttrium thin film as a reference (detailed experiments conditions are given in Table 4). The main peak we observed at 29.065° (2 θ) in Fig. 3a, most likely corresponds to α phase Yttria (222) with the lattice parameters calculated as a = b = c = 10.6340 Å. Our results are compatible with the literature in which the lattice parameters were calculated a = b = c = 10.6431 Å [31-33]. In addition to that, the peaks at approximately 20.405°, 39.615°, 49.930°, and $\approx 60^{\circ}$ (20) in Fig. 3a, denote the reflection of the x-rays from the corresponding planes in α -Yttria quite likely [33]. In the case of the metallic thin film (Fig. 3a), the major peak at 31.015° (20) indicates the formation of metallic yttrium crystal (possibly α -Y) [34,35]. If we focus on sample **S**₆, the peak that appears at 36.415° (20) in Fig. 3a and b, most likely corresponds to YO (002) [21,27,36], which is consistent with the findings by Losego et al. [37] in which YbO is deposited by molecular beam and Kaminaga et al. [27] where YO is grown by a pulsed laser. In addition to that, the peak that appears at approximately 31° (2 θ) (the shoulder between ~ 30.5 and \sim 31.7° (2 θ)) in the same figure could be responsible for YO ((111), 31.319° (2 θ) [27]) and/or β -Y₂O₃ (PDF Card; 00-044-0399). The results obtained from the current investigation, indicate the formation of nanocrystalline/amorphous YO accompanied by β -Y₂O₃ and/or α -Y₂O₃ (Fig. 3). The interplanar spacing (d) between the corresponding planes was calculated using the data obtained from XRD both in this research (R) and the literature (L) where d = 0.3072 (L) [33], d = 0.3064 nm (R) for α-Y₂O₃ (222), and 0.2485 nm (L) [27], 0.2465 (R) for YO (002). In the interest of clarity, further investigations on morphology and nanostructure were performed for semiconducting thin film by employing HR-SEM (Fig. 4a) and TEM (Fig. 4b). Fig. 4b; r1, r2 and r3 demonstrate the different regions of TEM images of the lamella extracted from S₆. The separation between the regions is \sim 50 nm. r1', r2', and r3' are the focused areas of the corresponding part (the yellow square) in each region. r1' FFT, r2 FFT, r2' FFT, and r3' FFT are the fast Fourier transformations (FFT) of the designated regions. Fig. 4b r2' FFT uncover the formation of amorphous phase that might contain different yttrium oxide phases in unison (some of the possible phases are YO, Y₂O₃). Fig. 4b r2 FFT shows the simultaneous formation of amorphous and crystalline phases. The calculated d-spacing (\sim 0.3 nm) may, considering the interpretation of the spectroscopic ellipsometry and XRD data. correspond to YO. On the other hand, the calculated lattice spacing in Fig. 4b r1' and r3', could indicate the formation of other crystalline phases detected by XRD (β-Y₂O₃ (PDF Card: 00-044-0399) and α-Y₂O₃ (PDF Card: 01-089-5592)). In spite of that, detailed investigations employing high resolution TEM (HR-TEM), selected area electron diffraction (SAED) and synchrotron based x-ray diffraction (SXRD) are required to create more precise relation between the data obtained (will be obtained) from electron microscopy/diffraction and x-ray diffraction.

3.2. Spectroscopic characterization

The main ellipsometric angles Ψ and Δ were measured at the incident angles from (55–85)° with 5° step. Refractive index *n* and extinction coefficient *k* dispersion curves were modelled using the Drude (DO), Gaussian (GO), Tauc-Lorentz oscillator (TLO) and Herzinger-Johs parameterized semiconductor (HJPS) oscillator functions [38]. The optical properties of Si/native SiO₂ substrates were obtained from SE measurements to clean substrates without the film. The surface roughness was modelled by utilizing the Bruggeman effective medium approximation (EMA) [39]. The optical gradient of *n* and *k* was calculated by dividing the film layer into sub-layers with smaller thicknesses



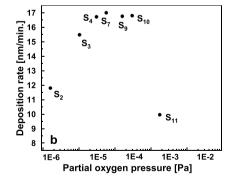


Fig. 2. a- Diffractogram of yttrium (oxide) thin films deposited at room temperature onto Si (001) with different partial oxygen pressure levels (Pa). Reference data was extracted from JCPDS PDF Card; 00-033-1458 (α -Y), 01-089-5592 (α -Y₂O₃), 00-044-0399 (β - Y₂O₃)

b- Deposition rates of the relevant samples with respect to the partial oxygen pressure. The power applied onto the target, and the deposition pressure levels were kept constant. Detailed experiment conditions are given into Table 4.

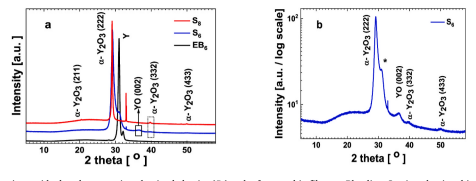


Fig. 3. Diffractogram of yttrium oxide that shows semiconducting behavior (*S*₆) and reference thin films: **a**- Blue line: Semiconducting thin film (*S*₆), Red line α -**Y**₂**O**₃ (*S*₈),: Black line: metallic Y (*EB*₆); **b**- The thin film shows semiconducting behavior in logarithmic scale. The peak at approximately 31 ° (2θ) may result from YO (111) [27] and/or β- **Y**₂**O**₃ (PDF Card; 00-044-0399).

and applying the EMA considering the films as a mixture of voids to vary the *n* and *k* values from the bottom to the top of the film [40]. Optical band gap E_{q} , for allowed direct transition, is obtained as a fitting parameter from TLO for S_6 sample and from HJPS for S_8 . The mean square error (MSE) between the modelled and experimental SE data was obtained in the range of 2-10. SE experimental data model-based regression analyses were performed by the Woollam software CompleteEASE®. The measured spectroscopic ellipsometry data for all samples are presented in Fig. 5. The absence of oscillation in (Ψ, Δ) for pure Y (\mathbf{EB}_6) and for the sample with crystalline mixture (\mathbf{S}_6) shows that these materials have high extinction (and absorption) coefficient k and that the thickness of these films is above 300-400 nm. Yttrium has lower values of Ψ in respect to the sample with the crystalline mixture that signifies pure Y, which has lower refractive index n, and higher k in respect to the samples with the crystalline mixture. The numerous high oscillations in (Ψ, Δ) spectra for **S**₈ sample, shows that this film is transparent for almost the entire spectral range with approximately 400 nm thickness. The decrease in oscillations above 3 eV, signifies that this material is starting to absorb the light, slightly. Fig. 6 illustrates the difference in the modelled spectra for Y2O3 samples, if the absorption is considered (or not considered) at E below optical band gap E_g . The refractive index n and extinction coefficient k as a function of photon energy E for Y, Y₂O₃, and a crystalline mixture of YO and Y₂O₃ materials, are shown in Fig. 7. Typical metallic characteristics in (n, k) curves can be seen for Y: *n* and *k* increase with the decrees of *E* from 3 eV to 0.7 eV, and it has relatively high k values (>1) at the spectral range above 3 eV (black curves of **EB**₆ in Fig. 7). The material that contains a mixture of semiconducting YO and dielectric phases of and Y₂O₃ (blue curves of S₆ in Fig. 7) has k values lower at the entire spectral range compared to k for pure Y. Typical dielectric (n, k) curves can be seen for Y₂O₃ sample: (n, k) goes up with the increase of E (red curves of S₈ in Fig. 7). The values of the films' thickness (d), surface roughness (Sr), optical band gap (E_g), *n* and *k* at 550 nm for Y, Y₂O₃, and for a crystalline mixture of YO and Y₂O₃, are summarized in Table 1. Optical properties of Y, YO and Y₂O₃ found in the literature, are given in Table 2. There are no reports in the literature about n values for YO or other crystalline mixtures that contain YO. The optical E_g value obtained from the TLO for crystalline mixture of YO/Y₂O₃ is (0.30 \pm 0.21) eV and, it is in good agreement with Kaminaga et al. work [27]. The findings of SE are supported by the absorbance spectra (Fig. 10b). The n values for Y are higher, with respect to the values reported in the literature that could be related with deposition conditions which affect the film thickness, structure and quality [41]. E_g obtained from HJPS for Y_2O_3 thin films, comply with the data in the literature [42–44]. The presence of the absorption for Y_2O_3 structure at the visible spectral range (between 2 and 4 eV in Fig. 7b) is related to the variation of the (n, k) within the depth of the film due to the inhomogeneities, e.g., porosity, defects, compositional variation in the film (Fig. 8). Absorption above zero at visible range was also observed in the work of Mudavakkat et al. [44] and Kaminaga et al. [27]

XRD analyses of S_6 evidenced the presence of crystalline mixture (Fig. 3). Thereby, the (n, k) dispersion curves for S_6 should represent the effective (n, k) values as a mixture of both Y_2O_3 and YO phases, theoretically. Based on that, the interpretation of SE data as EMA mixture of Y2O3 and YO (Model 1) and Y2O3, Y and YO (Model 2) was performed to obtain (1) the volume fraction of these mixtures and (2) to obtain physical (n, k) dispersion curves for YO. Two EMA models are illustrated in Fig. 9. The introduction of pure Y in one of the EMA models was carried out to check the reliability of the simulation. Both models gave the same fit and the same MSE to the experimental data (Table 3). The volume fraction of YO obtained by SE is high with respect to the values one could expect from XRD data: the peak of YO phase has low intensity with respect to Y2O3 phase. High volume fraction of YO obtained by SE for the sample S_6 could be explained by the dominant presence of amorphous YO phase. However, the extinction coefficient obtained for YO in both models gave comparable values to pure Y. Such observation could suggest that the film should contain a small amount of metallic Y (like in Model 2, for example), which is a contradictory information to XRD data. Therefore, the effective (n, k) curve previously obtained for S_6 sample (Fig. 7), is a close representation of the physical (n, k) values for YO. The contribution of Y_2O_3 to the SE spectra of S_6 is not evident: there are practically no oscillations of (ψ, Δ) compared to Y₂O₃ sample S₈. Sample S_6 has a rather higher thickness (~500 nm) and higher k values \sim 0.5-1 at the entire spectral range that corresponds to high absorption values $\sim 10^5$ cm⁻¹. Thus, SE is not sensitive to the whole thickness of S₆ sample, since the light is not passing through the entire depth of the film. Both high k and high thickness are affecting (Ψ, Δ) spectra which has quite low oscillations. It could be speculated that the presence of Y_2O_3 , seen in XRD spectra, is pronounced more for the first 100-200 nm of the film, which is "invisible" for SE due to high k of S_6 sample near the surface (last 100-200 nm). On the other hand, the contribution from Y_2O_3 cannot be seen in the k curves obtained (Fig. 7b) and also in the corresponding absorption α curves (Fig. 10), too. This could also be related to the fact that the E_g values Y_2O_3 are approximately 6 eV and thus *k* and α are not still increasing rapidly. In the work of Kaminaga et al. α has already reached ~13 \times 10⁵ cm⁻¹ at approximately 6 eV, while in our case α reaches $\sim 10^5$ cm⁻¹ at same proximity, which is one order less. Thus, α curves of the sample with crystalline mixture by Kaminaga et al. are also affected by Y2O3 contribution, probably due to lower E_g values of Y_2O_3 , and high volume fraction of Y_2O_3 in the sample. It should be noticed that; α values for S_6 are higher with respect to Kaminaga et al. (Fig. 10) due to a major contribution from the YO phase that supports the conclusions already drawn from EMA simulations, considering XRD data and SE analyses carried out aforehand. It could be concluded that; effective (n, k) curves obtained for S₆, are significantly dominated by the amorphous semiconducting YO, and are close representations of the physical (n, k) values for YO. Nevertheless, additional

and attributed to defect states in the band gap. The decrease of *n* toward

the surface of the films is related to the increase of the film porosity. The

a

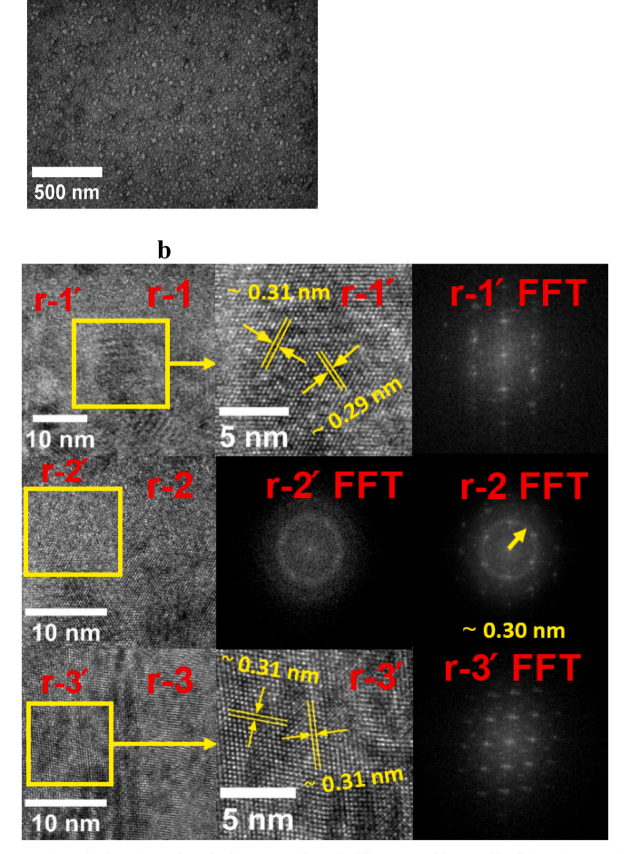


Fig. 4. a- HR-SEM image of S_6 : b- TEM images of S_6 ; **Left column:** r1, r2, and r3 are the different regions of the extracted lamella (separation \sim 50 nm). **Middle column:** r1', and r3' are the focused areas of the indicated regions in the left column and r-2' FFT is the fast Fourier transformation (FFT) of region r-2'. **Right column:** r-1' FFT, r-2 FFT and r-3' FFT are the fast Fourier transformations of the corresponding regions given in the left column.

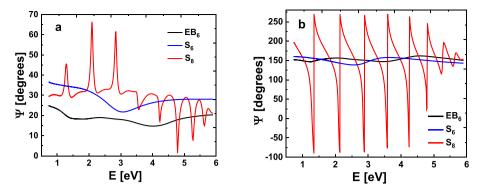


Fig. 5. Main ellipsometric angles Ψ (a) and Δ (b) as a function of photon energy *E* for three different samples.

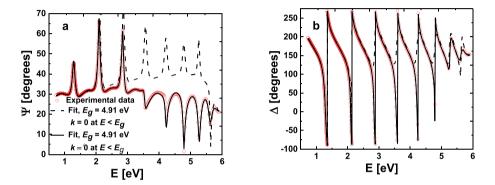


Fig. 6. Experimental (symbols) and modelled (lines) main ellipsometric angles Ψ (a) and Δ (b) as a function of photon energy *E* for three different samples. Two models are given to show the difference in spectra if absorption at *E* bellow E_g is taken (or not taken) into account.

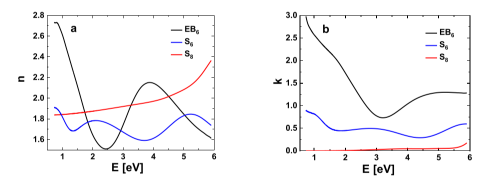


Fig. 7. Refractive index *n* and extinction coefficient *k* as a function of photon energy *E* for three different samples: Y (*EB*₆), sample with a crystalline mixture of YO and Y₂O₃ (*S*₆), and Y₂O₃ (*S*₈).

Table 1

Obtained values of the different films on Si: d - thickness of the films, S_r - surface roughness, E_g - optical band gap, n and k at 550 nm, oscillators used to model complex dielectric function of the films.

Film	d, nm	S _r , nm	E_g , eV	n	k	Oscillators
Y YO/Y2O3 Y2O3	$\begin{array}{l} 200^{a)} \\ 500^{a)} \\ 417.4 \pm 0.5 \end{array}$	$^{-}$ - 12.8 \pm 0.2	$^-$ 0.30 \pm 0.21 5.86 \pm 0.64	$\begin{array}{c} 1.533 \pm 0.001 \\ 1.78 \pm 0.05 \\ 1.89 \pm 0.02 \end{array}$	$\begin{array}{c} 1.437 \pm 0.001 \\ 1.47 \pm 0.02 \\ 0.0101 \pm 0.0004 \end{array}$	1 DO, 3 GO 1 DO, 1 TLO, 3 GO 1 HJPS, 2 GO

^{a)} The film is not trispirane for the given SE spectral range: the thickness is obtained with profilometer.

studies on thinner samples (~200 nm) fabricated under the same conditions as S_6 , will help to have a deeper understanding about the YO and Y_2O_3 phase formation dynamics, volume fraction and physical (*n*, *k*) dispersion curve characteristics for both crystalline and amorphous YO.

The main purpose of employing XPS in this research is the chemical state identification of yttrium (Y3d) in EB_6 , S_6 , and S_8 samples. Fig. 12a represents Y3d spectrum that belongs to EB_6 ; the peak detected at

approximately 155.90 eV, points out the formation of Y^0 (Y3d_{5/2}). The result obtained is consistent with the findings of the previous study by Mongstad, T. et al. [45]. Additionally, Fig. 12a' is the O1s spectrum of the coinciding Y3d spectrum. As it can be clearly understood, no peak has been detected. Fig. 12b shows Y3d spectrum of the sample named as **S**₆. The peak observed at 156.49 eV signifies the formation of Y²⁺ (Y3d_{5/2}). The results obtained are compatible with the literature findings 156.40

Table 2

Refractive index at 550 nm and optical band gap for Y and Y_2O_3 thin films reported in literature.

Description		<i>n</i> @550 nm	Eg, eV	Ref.	
Metallic yttrium Y					
Y thin film		~0.1–0.2 1.3 (CXRO tables)	-	[68]	
Yttrium monoxide	YO				
Crystalline mixture of Y_3O_2 and YO fabricated by PLD on CaF ₂ (001) substrate		_	0.1	[27]	
Yttrium oxide Y ₂ O ₃					
Nanocrystalline thin by PVD on quartz		1.79–1.90 (for substrate temperature during film deposition of 323–673 K)	-	[69]	
Thin films grown on to unheated [100] Si wafers with 1.2–2 µm SiO ₂ buffer layer	e-beam deposition	1.718 (process pressure 2.93 \times 10 ⁻² Pa with O ₂ flow rate 70 sccm) 1.741 (8.00 \times 10 ⁻³ Pa, 10 sccm) 1.917 (10.00 \times 10 ⁻³ Pa, no O ₂ flow)		[42]	
	IBAD ¹⁾	1.911 (2.00 \times 10 ⁻² Pa, 25 sccm)			
	HiTUS ²⁾	1.660 (1.09 \times 10 ⁰ Pa, 5 sccm) 1.868 (2.66 \times 10 ⁻¹ Pa, 9 sccm)			
Bulk cubic Y ₂ O ₃		1.934			
Thin films on Si subs	strate	\sim 1.645 (973 K; ion beam, 4.00 × 10 ⁻³ Pa) \sim 1.765 (973 K; IBAD, 150 eV O ₂) \sim 1.895 (room temperature; ion beam, none)		[43]	
Thin films grown by CVD on quartz and Si substrates		1.65–1.73	5.62–5.80	[70]	
Thin films on [100] Si wafers and quartz substrates by radio- frequency magnetron sputtering		~1.70–1.94 (substrate temperature from room temperature to 773 K)	5.91-6.15	[44]	

¹⁾ IBAD – ion beam assisted deposition.

²⁾ HiTUS - reactive sputtering with oxygen in a standard high target utilization sputtering system.

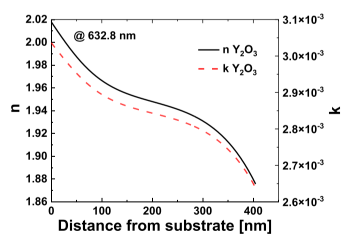


Fig. 8. Nonlinear depth profile for Y_2O_3 sample (S_8) at 632.8 nm (He–Ne laser) wavelength. The film layer was divided into 15 sub-layers (optimal value found for the lowest MSE).

eV (YO) [27] and 156.50 eV (YH_{2.1}) [46]. Further, at the same spectrum, the peak observed at 156.81 eV refers to the formation of $Y^{3+}(Y3d_{5/2})$ [47,48]. Fig. 12c shows Y3d spectrum of sample **S**₈, the peak detected at around 156.85 eV refers to $Y^{3+}(Y3d_{5/2})$ formation which is compatible with the results obtained by Nefedov, V., et al. [48]. In addition, the peak at approximately 155.26 eV most likely indicates $Y^0(Y3d_{5/2})$ at the

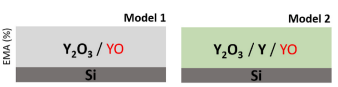


Fig. 9. Two different Bruggeman EMA models applied to fit SE data of S_6 sample to find physical (n, k) dispersion curves of YO and volume fraction of the mixtures. Model 1: film is a mixture of Y_2O_3 and YO. Model 2: film is the mixture of Y_2O_3 , Y and YO. In the models the obtained (n, k) curves for Y and Y_2O_3 where used. Resistivity values and optical band gap for YO was fixed to 0.045 Ω cm and 0.1 eV, respectively, to lower the number of fitting parameters.

same spectrum [49,50] that might result from the interaction between Ar⁺ and the sample. A detailed previous study carried out (for various elements) by G. Greczynski and L. Hultman [51] has reported the observation of slight shift of the spectrum towards lower binding energy values, which we also detected in the current research for S_8 (Y⁰ (Y3d_{5/2})) compared to the literature. Considering the O1s spectrum of samples S₆ (Fig. 12b') and S₈ (Fig. 12c'); Albeit, the peak detected (both Fig. 12b'. and Fig. 12c'.) at approximately 529.20 eV, corresponds to metal-oxygen (In this study Y-O)) [48], the philosophy behind the exchange of the intensities and the positions (528.40 eV-Fig. 12b'.) and (527.50 eV- Fig. 12c') is not clear (relying on the NIST data base). Moreover, additional SIMS measurements were performed to understand H content in the samples S₆ and S₈ (Fig. 6., supplementary material). The concentration of oxygen was measured $\sim 10^2$ higher than hydrogen concentration in both samples. We conclude, the contribution/contamination derived from H must be small enough. The results obtained from both XRD and SE, support the conclusions that come up from XPS measurements.

Vibrational spectrum investigation is carried out in Far-Ir (50-240 cm⁻¹) and Mid-Ir (240-700 cm⁻¹) region (Fig. 13), separately. In Mid-Ir region, we observed 6 peaks at \sim 240, 300, 335, 370, 460, and 555 cm⁻¹ based on the normal vibration mode of cubic sesquioxide yttrium lattice. The results we obtained, are consistent with the conclusions of previous studies [52-55]. We observe the peaks in both cases (sample S₈ (used as reference) and sample S_6). The intensity of the detected peaks demonstrates attenuation for sample S_6 compared to sample S_8 . We interpret that the decrease in the intensity of the peaks is most likely due to the formation of other crystalline phases (in the case study both semiconducting yttrium oxide with 2+ oxidation state and/or β -Y₂O₃), and of amorphization. For the reason that the electromagnetic wave couples with the vibrations of phonons in semiconductors in Far-IR [56], we focus on S_6 thin film in this region. But, it might also lead a challenge. To cope with this problem, we address two well-known phenomena as the strain (stress) accumulated in the thin film during the growth [57,58] and/or the formation of nanocrystalline/amorphous phase we detected in our thin film. We start with strain (stress) accumulation on the thin film. Born effective charge is influenced by strain (stress). This differentiation in the charge results in the electrical permittivity of the material of interest. Consequently, the coupling frequency of the electromagnetic radiation with both phonons and other elements, diversifies. This relation is noticeable for BaO, MgO, SrO, CaO [59]. Considering this phenomenon, the peaks in the region 90-170 cm^{-1} (70–200 µm) may presumably signify the formation of YO. The observation of EuO phonons in a similar region was obtained by Goian, V., et al. [60]. The coupling of electromagnetic wave with the vibrational frequency of the absorbers is common for most of the cubic rare-earth oxides between 90 and 170 cm⁻¹ [61]. Now we can discuss the nanocrystalline/amorphous formation. Normal vibrational modes are known as phonons for crystalline structures. On the other hand, if we deal with the amorphous materials (or nanocrystals), understanding the dynamics of phonons will be formidable [62-65]. One of the most noticeable differentiation in the spectrum is the broadening of peaks [62] (Fig. 13b and a). This might generate the observation obtained by Goian, V., et al.

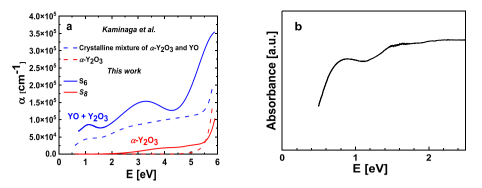


Fig. 10. a) Absorption curves for S_6 and S_8 samples calculated from extinction coefficient values (Fig. 7b) and compared with the results of Kaminaga et al. [27], b) Absorbance spectra of sample S_6 .

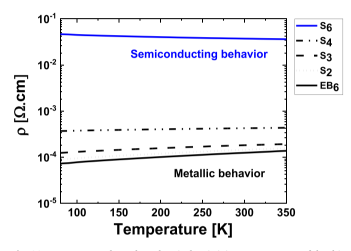


Fig. 11. Temperature-dependent electrical resistivity measurements of the thin films show semiconducting (S_6) and metallic behavior (including reference thin film (EB_6) with respect to the partial oxygen pressure level. Detailed growth parameters are given in Table 4.

Table 3

Summary of the fitting result applying two different Bruggeman EMA models to **S6** spectroscopic ellipsometry data. EMA% is the volume fraction for the corresponding material.

Model	MSE	EMA %		Nr of	(n, k) curves of	
		Y ₂ O ₃	Y	YO	correlating fitting parameters	ΥΟ
1	8.5	58.9		41.1	4	n lover respect to
2		$^{\pm}$ 0.4 48.9 $^{\pm}$ 0.5	$\begin{array}{c} 11.9 \\ \pm \ 0.3 \end{array}$	$^{\pm}$ 0.4 39.5 \pm 0.4	2	all samples; <i>k</i> comparable to Y

[60], more clear and those findings could indirectly point out the monoxide formation in our thin film. In IR spectrum, the coupling of electromagnetic wave with the traverse optical phonons (TO) and longitudinal optical phonons (LO) appears in pairs for the crystalline materials. Whereas in the amorphous materials, traverse optical phonons are more noticeable than the longitudinal optical phonons [63,66,67]. These findings may explain the reason behind the observation of low intensity peaks in the region 90-170 cm⁻¹ (Figs. 13a), and 70–200 μ m (Fig. 13b, the real space representation of the spectrum shown in Fig. 13a, in which, the relevant region becomes more noticeable). Our interpretations acquired from IR spectrum analyses show compliance with the results obtained from both structural and spectroscopic characterizations performed aforehand. Notwithstanding, further research must be carried out in order to construct an understanding of the

crystalline size dependency of the spectrum, and of the effect of accumulated stress (strain) in the structure on the spectrum.

3.3. Electrical characterization

In order to understand the transition dynamics of electrical conductivity from metallic to insulating demeanor, temperature-dependent electrical conductivity measurements were performed. The partial pressure level of oxygen was varied between 2.00 \times 10^{-7} and 2.31 \times 10^{-3} Pa (Fig. 11, and Table 4). The thin films obtained with 4.06×10^{-5} Pa and lower partial oxygen pressure values represent the metallic conductivity. The conductivity of the films increases with the decrease of the oxide phase/oxygen concentration in the host crystal structure (possibly α -Y). This relation becomes more noticeable if Figs. 2, 3 and 11 are considered/compared simultaneously. On the other hand, the pressure levels higher than ${\sim}8.00 \times 10^{-5}$ Pa are suitable for fully oxidize vttrium. The most striking result to emerge from the data is that; \sim 7.55 $\times \ 10^{-5}$ Pa is found as the convenient partial oxygen pressure level for the formation of semiconducting yttrium oxide at high temperature (623 K). The current findings also indicate that the range of the partial pressure level of oxygen for the formation of trivalent yttrium (Y₂O₃) and divalent yttrium (YO), is very sensitive. The aforementioned finding is consistent with the literature, which indicates that, YO shows semiconducting behavior as EuO and YbO [8]. Furthermore, Kaminaga et al. observed the semiconducting behavior for yttrium monoxide (YO) [27].

4. Conclusion

This paper primarily investigates the relation between the oxidation dynamics and structural formation (crystalline or amorphous) of yttrium oxide thin film. The results obtained, confirm the previous findings and contribute to the additional evidence which suggests that; under specific conditions, yttrium can take 2+ oxidation state in solid phase. One of the most significant findings gained from this study is the deposition of yttrium monoxide (YO) acquired from a metal target, which uses the reactive pulsed-DC magnetron sputtering for the first time. The second major finding is that the semiconducting yttrium oxide is presumably formed in the transition region between α phase and β phase yttria at high temperature, and accompanied by the crystalline Y₂O₃. The third finding is that, n dispersion curves are evaluated for the mixture of YO and Y_2O_3 for the first time. Moreover, it was demonstrated that the major contribution in evaluated (n, k) curves of YO and Y₂O₃ mixture, is provided by the presence of amorphous YO. Fourthly, in the case of the thin films that show metallic conductivity, the oxide phase (quite likely β -Y₂O₃) is formed simultaneously with metallic Y crystal, at room temperature. The current results are added to a growing body of literature on the solid-state divalent yttrium arrangement. This research has thrown up many questions in need of further investigation. Further research needs to examine the links between the oxide phase (α and/or

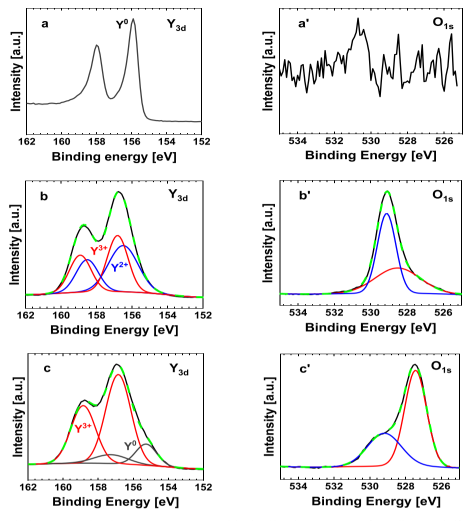


Fig. 12. Y_{3d} (left column) and relevant O_{1s} (right column) XPS measurements of the thin films that represent: a, a'- metallic (*EB*₆); b, b'- semiconducting (*S*₆), and c, c'- insulating (*S*₈) behavior. The dashed lines (green in color) in figure b, b', c, and c' define the fitting results of the spectrum.

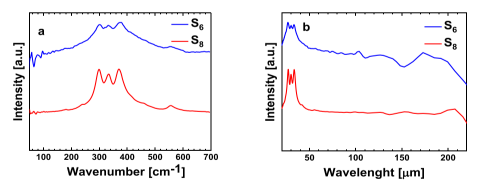


Fig. 13. IR Spectrum of insulating (S_8) and semiconducting (S_6) thin films, a-Scaled in wavenumber, b -Scaled in wavelength.

 β -Y₂O₃) and Y 2+ formation more closely using techniques such as XAS and theoretical modelling, and to establish whether/if obtaining thermodynamically stable single phase semiconducting yttrium oxide (YO) is possible.

CRediT authorship contribution statement

H. Arslan: Writing – review & editing, Writing – original draft, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. I. Aulika: Writing – review & editing, Writing – original draft, Investigation, Formal analysis. **A. Sarakovskis:** Formal analysis, Data curation. **L. Bikse:** Formal analysis. **M. Zubkins:** Resources, Data curation. **A. Azarov:** Formal analysis. **J. Gabrusenoks:** Formal analysis, Data curation. **J. Purans:** Supervision, Resources, Conceptualization.

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.

43

 $\sim 10^{-5}$

 $\sim 10^{-4}$

 $\sim 10^{-5}$

 $\sim 10^{-4}$

Table 4

EB₆

 S_6

S₈

Deposition/Evaporation parameters of the thin films; a-magnetron sputtered thin films: S_2 , S_3 , and S_4 are the thin films that show metallic type electrical conductivity, S_7 , S_8 , S_9 , S_{10} , and S_{11} are the thin films represent insulating character (S_8 is used as main reference sample), S_6 is the thin film shows semiconducting type electrical conductivity. b- EB_6 , e-beam evaporated thin film used as reference metal thin film. c- Partial pressure levels (approximate) of H_2 , H_2O and O_2 of samples EB_6 , S_6 and S_8 .

Sample	Deposition Pressure (Pa)	P_{Ar} Deposition (Pa)	P_{O_2} Deposition (Pa)	Growth Temperature (K)	Power (W/cm ²)
S ₂	${\sim}5.33 imes10^{-1}$	$6.40 imes10^{-2}$	$2.13 imes 10^{-6}$	298 ± 5	7.4
S ₃	${\sim}5.33 imes10^{-1}$	$4.64 imes10^{-2}$	$1.37 imes 10^{-5}$	298 ± 5	7.4
S ₄	${\thicksim}5.33\times10^{-1}$	$4.37 imes10^{-2}$	$4.06 imes 10^{-5}$	298 ± 5	7.4
S ₆	${\thicksim}5.33\times10^{-1}$	$4.92 imes10^{-2}$	7.55×10^{-5}	623 ± 5	7.4
S ₇	${\thicksim}5.33\times10^{-1}$	$4.81 imes 10^{-2}$	7.60×10^{-5}	298 ± 5	7.4
S ₈	${\thicksim}5.33\times10^{-1}$	$4.91 imes 10^{-2}$	$1.26 imes 10^{-4}$	623 ± 5	7.4
S ₉	${\sim}5.33 imes10^{-1}$	$4.09 imes10^{-2}$	$2.11 imes 10^{-4}$	298 ± 5	7.4
S ₁₀	${\sim}5.33 imes10^{-1}$	$3.92 imes10^{-2}$	$3.97 imes10^{-4}$	298 ± 5	7.4
S ₁₁	\sim 5.33 $ imes$ 10 $^{-1}$	$3.55 imes10^{-2}$	2.31×10^{-3}	298 ± 5	7.4
b					
Sample	Crucible	P_{0} , Evaporation (Pa)	Growth Te	mperature (K)	Current on e-Gun (mA)

c						
Sample	Rest/Base gases (Pa)			Process gases (Pa)		
	P_{H_2}	P_{H_2O}	P_{O_2}	P_{H_2}	P_{H_2O}	P_{O_2}
EB ₆	$\sim 10^{-7}$	$\sim 10^{-6}$	$\sim 10^{-7}$	$\sim 10^{-7}$	$\sim 10^{-6}$	$\sim 10^{-7}$

 $\sim 10^{-5}$

 $\sim 10^{-5}$

298 + 5

 $\sim 10^{-6}$

 $\sim 10^{-6}$

 $\sim 2.00 \times 10^{-7}$

 $\sim 10^{-5}$

 $\sim 10^{-5}$

Data	availa	hility
Data	avana	DIIITV

No data was used for the research described in the article.

w

 $\sim 10^{-6}$

 $\sim 10^{-6}$

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.vacuum.2023.111942.

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