Mixed oxide growth on combinatorial aluminium–gadolinium alloys — a thermodynamic and first-principles approach

Khurram Shahzad1 · Cezarina Cela Mardare1,2 · Andrei Ionut Mardare1 · Achim Walter Hassel1,2,3

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Abstract

Metal surfaces covered with oxides have attracted considerable scientific attention in various applications. In particular, anodic films fabricated by cost-effective anodizing have been widely used in nano-structured engineering to provide various surface functionalities. However, understanding of alloy film stability, having individual elements with widely varying structures and morphologies, is very limited due to lack of thermodynamic information and effects of electrolyte chemistry. This requires many tedious efforts on a trial and error basis in selecting suitable electrolytes that can produce the protective film at high efficiency on alloys having mixed chemistries. It is, therefore, crucial to develop a combination of high throughput theoretical analysis and automated rapid localized electrochemical probing that provides a fast and simple solution for electrolyte choice and paves the way to the remarkable expansion of industrial applications of oxides. Herein, we demonstrate that combinatorial Al–Gd alloys covering 1.0 to 10.0 at.% Gd can be oxidized into ultra-thin anodic films of controlled thickness through a selection of electrolyte based on thermodynamics (phosphate buffer with a pH of 8.20). We propose that growth of anodic films on alloys at high efficiency is possible if Gibbs free energy minimization criteria would be systematically contemplate.

Keywords Anodizing · Rare earths · Combinatorial · Scanning droplet cell microscopy

Introduction

The “1-to-1” approach involving one system, one calculation, and one experiment at a time is now regarded as the traditional, time consuming, and outdated method in materials research, when there is an ever-increasing need for strategies to develop advanced materials with novel functionalities [1, 2]. The intelligent strategies play a crucial role in technologies ranging from electronics, energy generation and storage, aerospace, medicine, and defence industry [3–5]. To achieve these technologies, attention has been given recently to integrate high-throughput first-principles calculations and combinatorial methods to obtain robust, extrapolative tools for gaining insights into structure–property–composition relationships in order to reduce the experimental efforts, time, and cost in the materials discovery process [6–13].

Ab initio or first-principles computational methods are becoming important in new materials discovery [14]. Scheming fabrication strategies using high-throughput computational approaches provide numerous benefits over traditional experimental-based materials discovery methods. Estimating material’s fundamental properties is often faster with computation tools, when compared to the experiment, and properties of thousands of materials can be calculated in a relatively short period of time. Additionally, it is often trivial in calculations to simulate the effects of chemical substitutions or lattice strain, but achieving these through experiments could take many months of painstaking laboratory work. Thus, computations are effective tools for designing new materials [15–17]. Directly linked to this, combinatorial approaches are defined by the development of a “library” that comprises a material parameter of

1 Institute of Chemical Technology of Inorganic Materials, Johannes Kepler University Linz, Altenberger Str. 69, 4040 Linz, Austria
2 Danube Private University, Steiner Landstrasse 124, 3500 Krems-Stein, Austria
3 Christian Doppler Laboratory for Combinatorial Oxide Chemistry (COMBOX) at the Institute for Chemical Technology of Inorganic Materials, Johannes Kepler University Linz, Altenberger Str. 69, 4040 Linz, Austria
The applications of these intelligent strategies are important, in particular, to rapidly explore the lesser-known and rarely explored rare-earth elements (REEs) and their alloys. The understanding and fabrication of REEs and their alloys through the electrochemical route are very limited due to the lack of both experimental and theoretical literature. As an example, very few efforts have been made to explore the properties of the central atom of lanthanoid series, the gadolinium (Gd), and its oxide (Gd₂O₃). Gadolinium ortho-aluminate (GdAl₂O₄) mixed oxide possesses excellent magnetic and antiferromagnetic properties [20–22]. The pure oxide Gd₂O₃ is a wide bandgap material (5.37–6.37 eV) [23] and meets the low leakage current requirement in CMOS applications [24]. The superior optical properties of Gd₂O₃ thin films make them efficient for use in optical applications [23, 25, 26]. Hong et al. reported that Gd₂O₃ is an excellent dielectric material to passivate GaAs surface, with a dielectric constant of 10 for epitaxial cubic phase and a low leakage current density of 10⁻⁹–10⁻¹⁰ A cm⁻² at zero bias [27], while Zhou et al. have found out a value of 20 for Gd₂O₃ thin-film [28]. This indicates that changing the substrate and growth method affects the anisotropic polycrystalline film growth, which can enhance its dielectric constant. The dielectric constant of Gd₂O₃ can be further used to modify the dielectric behaviour of aluminium-based alloys. The dielectric constant is one of the factors responsible for capacitance enhancement, and portable electronic devices have increased the demand for small-size electrolytic capacitors with high capacitance, low equivalent series resistance, and low leakage current [29, 30]. Materials with high dielectric constants are also crucial in many areas of modern microelectronics for miniature supercapacitors, infrared detectors, radar absorbing coatings, and holographic devices. Thus, there is a growing interest in the synthesis and precise characterization of materials with high dielectric constants and low leakage currents.

The comprehensive understanding of oxide behaviour in a solution can be enhanced by combining theoretical predictions with localized measurements. The localized analysis is critical as the dielectric properties are generally dependent on the material geometries and crystalline orientations [31]. At microscale, properties may be very different than those at macroscale. It is therefore crucial to probe these properties at a scale where inhomogeneities may be specially resolved. The development of scanning droplet cell microscopy (SDCM), in particular, has allowed for spatially resolved electrochemical characterization of surfaces [32–34]. Coupling SDCM with electrochemical impedance spectroscopy (EIS) may even determine the grain-specific responses and has superior control of the working area [31–36].

In this study, we take advantage of computational and combinatorial strategies to explore the composition, structure, and properties relationship systematically in Al-Gd alloys. For this purpose, the predictions of possible stable phases in a given electrochemical system were obtained from conventional potential-pH (or E-pH) diagrams and DFT simulated compositional phase diagrams and E-pH diagrams. This permits a rapid identification of anodizing electrolytes over a wide pH range for fabrication of Al–Gd alloys in an ultra-fast manner and calculations of material specific parameters.

**Experimental methods**

Laterally graded Al–Gd alloy library was obtained by thermal co-deposition of the Al and Gd from two separated sources on borosilicate float glass substrates (VWR International GmbH, Germany) having dimensions of 2.6 × 7.6 cm². A thermal co-evaporator with a base pressure of 10⁻⁵ Pa was used for co-deposition of Al and Gd. The compositional control along the library was obtained by individually adjusting the deposition rate of each source. Specific details of the hardware used for library deposition were previously reported [37]. A low Gd amount was desired along the library, thus deposition rates of 0.9 and 0.1 nm s⁻¹ were used for Al and Gd, respectively, to obtain a compositional spread of approximately 9 at.% along the length of the substrate. A total film thickness of 350 nm was obtained at the middle of Al–Gd library. It is worth mentioning here that the compositional spread of a library is limited by the particularities of the cosine law governing the thickness uniformity of alloying element taken separately along the substrate. For Gd contents above 10 at.%, a new compositional spread needs to be prepared. However, higher Gd contents would result in formation of various intermetallics, which is not desired here.

Immediately after Al–Gd library deposition, scanning energy-dispersive X-ray (EDX) spectroscopy was used for compositional mapping of the Al–Gd spread. Each alloy was measured at discreet surface locations and their quantitative analysis was performed using IDFix software (remX GmbH, France). The Gd amount varied between 1.0 and 10.0 at.% along the library, while the Al concentration varied complementarily between 99.0 and 90.0 at.%. This defines...
a compositional resolution of the Al–Gd thin film library of 0.12 at.% mm$^{-1}$. For an usual requirement of 1 at.% precision for alloy identification, more than 7 mm are thus available on the surface of the library for characterizing a single Al–Gd alloy. This resulting area is quite convenient for being addressed by other investigation techniques for Al–Gd alloy property mappings. X-ray diffraction (XRD) was used to map the crystallographic particularities of Al–Gd alloys as a function of their composition. The experiments were performed in $\theta$–$2\theta$ (Bragg–Brentano) and $\omega$–$2\theta$ with $\omega = 3^\circ$ (grazing incidence) geometries with CuK$_\alpha$ radiation. Additionally, the surface morphologies of individual Al–Gd alloys (defined by EDX mapping) were characterized by scanning electron microscope (SEM—Zeiss Gemini 1540 XB) before and after anodization.

The electrochemical measurements on thermally co-deposited Al–Gd films were carried out using the SDCM in contact mode [32] at ambient temperature and pressure. The SDCM is a three-electrode system containing two channels in V-geometry, serving as inlet and outlet for the electrolyte. A Pt wire positioned in the electrolyte supply compartment served as counter electrode while a Hg/Hg$_2$Cl$_2$ µ-reference electrode was placed in the drain compartment for avoiding chloride contaminations [38]. The Al–Gd thin film library deposited on glass substrates served as working electrode. The addressed area (0.049 cm$^2$) was determined by SEM examination of anodically grown films. A steady flow of electrolyte is provided by a peristaltic pump that works as both electrolyte supply and drain at a constant rate.

All measurements were carried out using self-developed Labview software that controls the scanning of the SDCM tip along the library (using an XY translation stage) and monitors the applied force during contact. In this way, a single droplet of electrolyte is moved along the surface. At each unique XY position which also corresponds to unique alloy composition, open-circuit potential (OCP) was measured for 180 s in 0.2 M phosphate buffer having a pH value of 8.20 (Fig. 1). The buffer solution was obtained by mixing anhydrous potassium monobasic and sodium phosphate dibasic heptahydrate. Following OCP recording, Al–Gd alloys were locally anodized potentiodynamically to 10 V. Film resistance and dielectric properties of grown films were determined by simultaneously coupling EIS with cyclic voltammograms (CV) using a Compact Stat electrochemical interface system (IVIUM Technology, The Netherlands) with an AC amplitude of 50 mV in a frequency range of $10^2$ to $10^{-1}$ Hz. All impedance spectra in the present study were recorded at 0 V vs. SHE. The present study was conducted using the constant electrolyte flow from inlet to outlet compartments within the SDCM body. This ensured fresh electrolyte at the contact with the working electrode during combined CV-EIS measurements. The protocols followed in the electrochemical study is as follows: (1) positioning of cell at a given location on the substrate and OCP recording, (2) anodic sweep (100 mV s$^{-1}$) from OCP value measured before, (3) EIS measurement of grown film, (4) looping of (1—3) to 10 V at 1 V incremental step. A more detailed description of the microelectrochemical flow measurement can be found elsewhere [6, 32, 39]. The thermodynamic data and electrochemical reactions used in the present study were taken from different sources [40–46]. Information regarding formation energies of species was taken from an open source web application [47]. The E-pH diagrams from first-principles DFT calculations were constructed from the high-throughput infrastructure developed by Jain et al. [46, 47], where experimentally determined free energies of aqueous ions and DFT energies for solid phases were used to construct multi-element compositional phase diagrams and E-pH diagrams. First-principles Car-Parinello molecular dynamics was used for simulations of aqueous states [48]. All E-pH diagrams were calculated at 298 K and atmospheric pressure for dissolved concentrations of species per litre of water.

Results and discussion

Film structure and morphology

The crystallographic properties of individual alloys were examined in both $\theta$–$2\theta$ and $\omega$–$2\theta$ modes along the Al–Gd library. XRD analysis was performed at various Al:Gd ratios and selected XRD patterns are presented in Fig. 2 as a function of Gd concentration. Aluminium possesses fcc symmetry, whereas Gd has a hexagonal structure. Additional phases other than Al were not found and the addition of Gd did not induce tri-aluminide intermetallic phase. Significant reduction in Al(111) peak intensity is observed when mapping from higher to lower Gd concentration. This could be due to variation in Al and Gd ratios as well as the gradual loss of crystallinity which indicates the influence of Gd concentration.
The binary Al–RE phase diagram and detailed analysis of Al–RE systems by Gschneidner et al. [49] indicate that GdAl₃ tri-aluminide intermetallic phase is formed between 0 and 25 at.% Gd addition to the Al matrix. The formation of the REAl₃ phase is linked with the transformation of the structure from predominately hexagonal to cubic for the light lanthanoids having large radii (La, Ce, Nd, Pr, Sm, and Gd). Contrary to binary Al–Gd phase diagram and Gschneidner et al. prediction, the absence of the GdAl₃ phase may be attributed to the low sensitivity of the XRD analysis at nm-depth scale. Additionally, Al rich-phase predominates in the alloy matrix, and it is likely that the presence of GdAl₃ is eclipsed by the Al-rich matrix.

The surface morphologies along the Al–Gd compositional spreads before and after electrochemical studies were examined by SEM. The SEM images prior to anodizing were taken in the vicinity of anodizing spot for each composition. Figure 3 shows the map of the observed morphologies from 1.5 to 10.0 at.% Gd, and the composition of each investigated region is given at the upper left corner. The microstructural analysis discloses a gradual change in surface morphology as the Gd content in the Al matrix increases (Fig. 3a) and can be divided into three distinct parts. The typical Al grains can be seen in the first line of SEM images, ranging from 1.5 to 2.5 at.% Gd. The suppression of large grains and gradual formation of mixed morphology are evident from 3.5 to 5.5 at.% Gd (second line), where the imprint of surface grains and grain boundaries still can be seen. A noticeable change in the surface morphology and distinct features with thick upright needles can be observed from 7.0 to 10 at.% Gd. Moreover, the change in surface features with Gd addition agrees well with XRD analysis, where the reduction in peak intensity and loss of crystallinity were attributed to variation of the Al:Gd ratio.

Surface morphologies after anodizing reveal a significant change and show a network of bright lines on the original Al grains, as can be readily seen from the SEM image in Fig. 3b. Because of their strong adsorption on the metal surface, the possibility of phosphate precipitates cannot be overlooked. No signs of localized corrosion or pitting was found in SEM observation.

**Electrochemical studies**

Composition and structural complexity in binary Al–RE alloys render the electrochemical measurement unstable [50]. One alloy constituent may dissolve preferentially in a certain pH range leaving the other constituent unaffected. Structure and morphological complexity due to lattice mismatch and a large difference in atomic and ionic radii further complicate the understanding of dissolution susceptibility in a specific pH range. Consequently, predicting thermodynamically auspicious conditions for anodizing multi-component systems is a meticulous task due to a large number of competing reaction pathways and end products. Additionally, the classical E-pH diagrams do not address the alloy–H₂O systems, thereby producing a gap in the theoretical understanding of Al–RE alloys [40, 41, 51, 52]. Hence, a theoretical framework is also developed in the present study, which can readily foresee the thermodynamic feasibility of reactions, stable phases in a complex Al–Gd–P–O–H system. For instance, conventional E-pH diagrams do not reveal the presence of AlPO₄ and GdPO₄ phases in Al–H₂O and Gd–H₂O aqueous systems. Consequently, calculations of material specific parameters and analysis of film properties would remain incomplete without considering missing phases in conventional E-pH diagrams.

In order to find a compromise pH range for Al–Gd alloys in phosphate buffer, the conventional E-pH diagrams Al–H₂O, Gd–H₂O, P–H₂O systems were simulated from an experimental database [40–44] and superimposed on each other. Figure 4a shows the superimposition of Al–H₂O and Gd–H₂O systems, while Fig. 4b reveals the superimposition of Gd–H₂O and P–H₂O systems. A different stability domain for Gd(OH)₃ and Gd₂O₃ arises because of the difference in free energies for hydroxide and oxide formations (see Table 1). In other words, considering the metal hydroxide alone is not sufficient to predict the pH range even for a single metal. Figure 4a discloses that the domain of Al₂O₃ stability terminates at around pH 9.0 and shifts towards lower pH range as the dissolved Al concentration...
in the solution reduces. Additionally, passive domains in the Gd–H₂O system begin at around pH 9.0 and moves towards higher pH values when the concentration of dissolved Gd is decreased in the solution. Simultaneous oxidation of Al and Gd alloys at high efficiency or at a nearly equal amount of dissolved concentrations of both metals is not thermodynamically feasible in the aqueous electrolyte at any pH value. Consequently, oxide-forming species such as phosphate and borate are needed to shift the passive domains of both metals. However, it was further found out that superimposing conventional pH diagrams of metal–H₂O systems with oxide-forming species does not modify the E–pH diagrams. One reason for this is that superimposed diagrams do not incorporate the reactions between metals and oxide-forming species. Thus, it is not possible to progress in the understanding of an alloy anodizing in a complex aqueous environment.
medium by just superimposing a series of conventional Pourbaix diagrams, and this may often lead to false conclusions in predicting aqueous corrosion. Moreover, extracting thermodynamic information from published literature is a very time-consuming effort. Many alloys exhibit intriguing physical and chemical properties resulting from the complex interactions between the constituent’s elements. Several known or unknown non-equilibrium compositions and structures can form in a material during its lifetime and at present, it is challenging to accurately simulate all of the relevant phases in a given system. In order to overcome this limitation, there is a need for an advanced approach to predict the realistic complex systems in a high-throughput manner. Therefore, to maximize the prediction of possible phases in the Al–Gd–P–H–O complex system, the grand canonical linear programming (GCLP) method [7, 9] and

| Chemical and electrochemical reactions | ΔG (kJ mol⁻¹) | Species | ΔG f (kJ mol⁻¹) |
|---------------------------------------|--------------|---------|----------------|
| H₂O                                  | 474.27       | H₂O     | −237.370       |
|                                       |              | H⁺      | 0              |
|                                       |              | e⁻      | 0              |
| Gd–H₂O system                         | 659.71       | Gd³⁺    | −659.710       |
| Gd³⁺ + 3H₂O → Gd(s)                   | 89.01        | Gd(OH)₃(s) | −1282.81     |
| Gd³⁺ + 2H₂O→ 4H⁺ + Gd(OH)₄⁻           | 196.28       | Gd(OH)₄⁻ | −938.170      |
| 2Gd³⁺ + 3H₂O→ Gd₂O₃ + 6H⁺             | 306.97       | Gd₂O₃   | −1724.56       |
| Gd–H₂O–PO₄³⁻ system                   | −118.11      | Gd(PO₄)₂³⁻ | −2827.22     |
| 2H⁺ + Gd³⁺ + PO₄³⁻ → GdH₂PO₄²⁺        | −127.63      | GdH₂PO₄²⁺ | −1812.04      |
| Gd³⁺ + PO₄³⁻ → GdPO₄(s)               | −146.18      | GdPO₄(s) | −1830.59       |
| PO₄³⁻ –H₂O system                     | −75.26       | PO₄³⁻   | −1024.74       |
| H⁺ + PO₄³⁻ → HPO₄²⁻                   | −70.44       | HPO₄²⁻  | −509.020       |
| Al–H₂O system                         | 481.0        | Al³⁺    | 481.0          |
| Al³⁺ + 3H₂O→ Al(s)                    | 59.23        | Al(OH)₃(s) | −1137.6     |
| Al³⁺ + 4H₂O→ 4H⁺ + Al(OH)₄⁻           | 129.52       | Al(OH)₄⁻ | −310.92       |
| 2Al³⁺ + 3H₂O→ Al₂O₃ + 6H⁺             | 91.81        | Al₂O₃   | −1582.3        |
| Al–H₂O–PO₄³⁻ system                   | −119.76      | AlPO₄(s) | −1625.50       |

![Fig. 4](image-url) a Superimposed E–pH diagram of Al–H₂O and Gd–H₂O systems. The diagrams are constructed at different molar concentrations of dissolved Al and Gd with exponents of 0, −3, −6, and −8 in water at 298 K; b superimposed E–pH diagram of Gd–H₂O and P–H₂O systems for Gd₂O₃/Gd(OH)₃ formation at molar concentration per litre of water with exponent of −6
open-source web application [45–47] were also utilized for analysing the complex ground state thermodynamics. GCLP strategy works by mapping a free energy minimization problem as a linear algebra problem and this allows the computation of very large number of phases in a very high-dimension phase space. The exemplary DFT simulated compositional phase diagrams for the identification of stable or metastable phases in Al–Gd–P–O–H₂ system are provided in the supplementary Figures S1, S2 and S3. An example of implementation of above-mentioned approach to construct E-pH diagrams can be seen in Fig. 5, where stable phases found in compositional phase diagrams of Al–Gd–P–O–H₂ system can be considered for E-pH diagrams of Al–P–H₂O (Fig. 5a), Gd–P–H₂O (Fig. 5b), Al–Gd–P–H₂O (Fig. 5c) aqueous systems. Figure 5d reveals the summary of enthalpy change estimated from DFT calculations for Gd₂O₃–H₂O, Al₂O₃–H₂O, Al₂O₃–Gd₂O₃, Gd₂O₃–P₂O₅, and Al₂O₃–P₂O₅ binary systems. It is obvious that phosphate incorporation remarkably modifies the E-pH diagrams of both constituents of the alloy and provides the protection of Al³⁺ and Gd³⁺ species around pH 6.0–9.0 (Fig. 5c) above 0 V. Consequently, phosphate-containing electrolyte was chosen between pH 6.0 and 9.0. The oxide-forming species such as phosphate cannot be selected by just considering the free energies of species (AlPO₄ and GdPO₄). There must be some thermodynamic pH range at which a specific electrolyte can be used. It is worth-mentioning here that the main objective of superimposed E-pH diagram computations was to identify the electrolyte and pH ranges. A similar framework can be designed for other electrolyte selection such as phosphate-citrate buffer, ammonium pentaborate tetrahydrate.

Following theoretical consideration and electrolyte selection, schematic and expected mixed oxide growth for the present study is shown in Fig. 6, which discloses the use of SDCM along Al–Gd library (a), oxidation of Al and Gd atoms upon application of formation voltage, V and interaction of cations with the phosphate buffer (b, c). The relative sizes of atomic and ionic radii of Al and Gd are noticeable before and after the application of V as the migration rate of cations through the film depends on their sizes and bond energies. The oxidation of alloying elements in Al-binary alloys depends on Gibbs free energy per equivalent for the

Fig. 5  Simulated E-pH diagrams using first-principles DFT methods: a P–Al–H₂O system at ₐₐ₃⁺ = 10⁻⁶ M, ₐ₈₉₅⁻ = 10⁻¹ M; b P–Gd–H₂O system at ₐ₉₈₇⁺ = 10⁻⁶ M, ₐ₈₉₅⁻ = 10⁻¹ M; c P–Al–Gd–H₂O system at ₐ₈₉₅⁺ = ₐ₉₈₇⁺ = 10⁻⁶ M, ₐ₈₉₅⁻ = 10⁻¹ M. d Enthalpy change of oxide formation for Gd₂O₃–H₂O, Gd₂O₃–P₂O₅, Gd₂O₃–Al₂O₃, H₂O–Al₂O₃, and P₂O₅–Al₂O₃ binary oxide phase diagrams using first-principles DFT methods
formation of the alloying element oxide, relative to that of formation of alumina. A more negative free energy value of the alloying element oxide relative to alumina specifies that the alloying element has a higher mobility than Al [53–57]. As Gd$_2$O$_3$ and GdPO$_4$ have higher negative free energy compared with Al$_2$O$_3$ and AlPO$_4$, it is likely that Gd$^{3+}$ will migrate faster than Al$^{3+}$ during anodizing Al–Gd alloys, and both Al and Gd ions would be incorporated into the oxide film and would form units of oxides. Phosphate adsorption on the surface of mixed oxide is suggested in part (d) of the figure, where the expected amorphous character of the anodic film is neglected in order to better suggest mixing. The mixing from the molecular point of view is represented in part (e). Very large sizes of AlPO$_4$ and GdPO$_4$ molecules and closed structure of mixed oxide (GdAlO$_3$) suggest that both AlPO$_4$ and GdPO$_4$ despite their thermodynamic stability may have limited inward migration rate. Because of the constant flow of fresh electrolyte, it is assumed that phosphate-containing species will flush out leaving only a minute quantity of PO$_4^{3-}$ on the film surface as adsorbed species.

**Potentiodynamic oxide formation**

Anodizing and EIS studies were performed on alloys along the library. Series of cyclic voltammograms with a step-wise increase of $V$ between 1 and 10 V were recorded at 11 different compositions by positioning the tip of SDCM along the Al–Gd compositional spread. The graphical representation of selected CVs is presented in Fig. 7 which corresponds to 110 potentiodynamic scans and EIS spectra. For each composition, a visible current density plateau can be readily observed which defines the charge consumed during the incremental growth process. This is due to the direct relationship between the charge transferred to the working electrode and the oxide volume grown following Faraday’s law. This leads to the average oxide formation factor, $k_{ave}$ which can be defined as

$$k_{ave} = \frac{\Delta d}{\Delta V} = \frac{qK_{ox}}{V}$$

(1)

where $K_{ox}$ is a material constant, $d$ is the oxide thickness, $V$ is anodizing potential, $q$ is the charge consumed, and $\eta$ is the efficiency of the film formation current. $K_{ox}$ can be defined in terms of oxide specific constants (mixed oxide molar mass, $M_{ox}$, oxide density, $\rho_{ox}$, and oxidation number, $z$, and Faraday’s constant $F$) by the following equation:

$$K_{ox} = \frac{M_{ox}}{zF\rho_{ox}}$$

(2)

The molecular masses and densities of mixed anodic films locally grown on the Al–Gd alloys library were estimated using the mixed-matter theory by means of linear distributions between the Al$_2$O$_3$ and Gd$_2$O$_3$. Owing to negligible dissolution of Al and RE in phosphate-containing electrolytes [58–65], a typical passive film current density plateau
and negligible leakage current density response on reversal of potential sweep scans, the efficiency of film formation can be assumed here as 100%. Because of the large sizes of GdPO₄ and AlPO₄, it is expected that phosphate incorporation within the film would be extremely small. The extremely low dissolution of RE in phosphate-containing electrolyte was verified by coupling CV-EIS measurements and ICP-OES analysis in Al–Ho system (~10⁻²⁰ ppb) [64]. The current overshoots present at low potentials may be attributed to the delayed oxide growth due to the space charge layer formation or the sudden reaction of the metal surface to the electrolyte. Therefore, the CV and EIS data are excluded at 1 V in *k*ₐᵥₑ calculation.

Electrochemical impedance spectroscopy

Bode and Nyquist plots for two extreme compositions are shown in Fig. 8. The impedance changes linearly as a function of frequency for all compositions, and gradually rises with *V* implying a linear growth of the anodic film. The direction of arrows shows the increase in anodizing potential. As the film grows, an expected shift of phase angle towards –90° typical of dielectric oxide film is obtained. The value of phase shift, *α* obtained from impedance fitting was above 0.92 in all compositions.

Figures 8a, c show that regardless of film composition, Bode plots on a log scale show similar absolute impedance. The corresponding Nyquist plots on a linear scale in Fig. 8b, d disclose the magnitude reduction in the imaginary component (–Z*ᵢₘ₈*) of impedance without affecting the real component (Z*ᵟ*). This indicates that Gd addition affects the capacitance or dielectric properties and has no appreciable effect on the resistance of passive films. As the capacitance varies inversely with the impedance, it is envisaged that films comprising Al and Gd mixed oxides can be obtained by incorporating Gd with enhanced capacitance into Al matrix while retaining the film resistance, *R*ᵟ of pure alumina. Moreover, the phase shift behaviour reveals that as oxide grows thicker, incremental *V* not only induces a capacitive response, as evidenced by the magnitude of phase shift, but also widens the constant phase element (CPE).

The influence of Gd on dielectric constant can be obtained by a careful determination of capacitance through EIS fitting. Different approaches are used in the literature to calculate the true value of capacitance from CPE without considering the origin of the CPE [66–68]. It is difficult to implement a suitable CPE-C conversion model, thereby, the determination of capacitance from CPE data is often inadequate, leading to the erroneous prediction of physical properties [69–71]. For this reason, special consideration has been taken to understand the CPE behaviour for passive film growth.

The impedance of *R*ₐₑ(◊*R*ᵟ*) circuit exhibiting CPE behaviour having parameters *Q* and *α* can be expressed as

\[
Z = R_{el} + \frac{R_{f}}{1 + Q(j\alpha)^2 R_{f}}
\]

where *R*ₐₑ and *R*ᵟ are the electrolyte and film resistances, respectively. Hirschorn et al. demonstrated the significance of employing the correct formula for a given distribution of time constants as well as impact on film properties [69].
Hsu and Mansfeld [67], on the other hand, considered the frequency, \( \omega_{\text{max}} \) on the top of the depressed semi-circle where the real part \((Z_r)\) lies midway between low and high-frequency x-axis intercepts on the Nyquist plot. As a result, multiple approaches are compared in order to accurately predict the capacitance, \( C \) and \( k_{\text{ave}} \) from CPE having parameters \( Q \) and \( \alpha \). The results for the \( k_{\text{ave}} \) values were also compared with the \( k_{\text{ave}} \) values obtained from Faraday’s law (Table 2). The \( R_f, \alpha, \) and \( Q \) values used in Table 2 were obtained by analysing the EIS data using ZView software.

It is also worth mentioning here that Hsu-Mansfeld and Hirschhorn et al. presumed the semi-circle depression model while deriving the mathematical expression for CPE without considering the role of film resistance, \( R_f \) in Eq. (3) [67, 69]. Obviously, the term “\( Q(j\omega)^\alpha \)” in Eq. (3) cannot completely determine the resultant impedance of \( R_{\text{el}} (QR_f) \) circuit, and CPE behaviour in itself cannot be fully understood by neglecting \( R_f \). Therefore, to understand the uniform film growth as a function of \( V \), the impedance spectra are also simulated in parallel by assuming different scenarios as illustrated in Fig. 9.

Case 1 in Fig. 9a illustrates the situation where only \( \alpha \) changes uniformly from 1.0 to 0.5 as commonly observed in the semi-circle depression model. Changing \( \alpha \) without affecting \( Q \) and \( R_f \) does not replicate the typical impedance spectra as generated in uniform film growth (see Fig. 8) and even suppresses the CPE behaviour when \( \alpha \) approaches 1.0 as demonstrated in the phase shift vs. frequency plot. Case 2 describes the resistive film scenario in which the film resistance, \( R_f \) and the ratio \( R_f/R_{\text{el}} \) vary considerably, and CPE parameters \( Q \) and \( \alpha \) contribution are largely suppressed due to the dominating effect of \( R_f \) (Fig. 9b). Under such conditions, the CPE behaviour starts extending towards the lower frequency part of impedance spectra. Finally, case 3 represents the most realistic situation of thin-film growth by anodizing where all \( \alpha, R_f \), and \( Q \) vary in a cooperative manner during film growth (Fig. 9c). The widening of the CPE towards low or high-frequency end of Bode plot will largely depend on the dominant effect from \( Q \) and \( R_f \). The CPE broadening for the highly capacitive film, must occur towards the higher frequency side, while for a resistive film having negligible contribution of \( Q \), it should be extended towards low frequency. Lastly, uniform film growth can be easily understood from the shape of the Bode plot where \( R_f, Q \), and \( \alpha \) are changing as a function of \( V \) in the present study.

The summary of EIS analysis is provided in Fig. 10 and Table 2 in the form of inverse capacitance, \( C^{-1} \), \( k_{\text{ave}} \), and relative permittivity, \( \varepsilon_r \), as a function of \( V \). The \( C^{-1} \) increases linearly with \( V \) (Fig. 10a), as expected from the following equation:

\[
C^{-1} = \frac{d}{\varepsilon_r \varepsilon_0 A} = \frac{k_{\text{ave}} V}{\varepsilon_r \varepsilon_0 A}
\]

(4)

in which, \( \varepsilon_0 \) is the permittivity of vacuum and \( A \) is the surface area.

### Comparison of oxide formation factors

Table 2 indicates that EIS analysis yields \( k_{\text{ave}} \) values ranging from 1.34 to 1.62, while the \( k_{\text{ave}} \) values vary from 1.52 to 1.76 when calculated from Faraday’s law. Comparison of \( k \)-factor obtained from different equations reveals that Hsu-Mansfeld equation \( (C_{\text{ox,M}}) \), Faraday’s law, and normal

| at.% Gd | Oxide formation factor, \( k_{\text{ave}} \)/\( \text{nm} V^{-1} \) |
|--------|----------------------------------|
|        | Hsu-Mansfeld \( C_{\text{ox,M}} = Q(\omega_{\text{max}})^{\alpha} \) | Bryan Hirschhorn \( C_{\text{ox,H}} = Q^2 (R_f)^{1-(\alpha)/\alpha} \) | Bryan Hirschhorn \( C_{\text{ox,H}} = Q^2 (R_{\text{el}})^{1-(\alpha)/\alpha} \) | Faraday’s Law \( d = \frac{M\ell}{A\varepsilon_0 F_p} \) |
| 1.5    | 1.49 | 1.46 | 2.18 | 1.52 |
| 2.0    | 1.50 | 1.49 | 2.37 | 1.66 |
| 2.5    | 1.45 | 1.42 | 2.44 | 1.64 |
| 3.0    | 1.46 | 1.42 | 2.57 | 1.63 |
| 3.5    | 1.42 | 1.39 | 2.83 | 1.63 |
| 4.5    | 1.49 | 1.42 | 3.40 | 1.70 |
| 5.0    | 1.58 | 1.53 | 3.92 | 1.73 |
| 5.5    | 1.61 | 1.58 | 4.46 | 1.76 |
| 6.0    | 1.58 | 1.60 | 4.90 | 1.70 |
| 7.0    | 1.62 | 1.58 | 5.72 | 1.71 |
| 8.5    | 1.58 | 1.42 | 6.48 | 1.72 |
| 9.0    | 1.53 | 1.48 | 7.63 | 1.73 |
| 9.5    | 1.50 | 1.50 | 8.14 | 1.74 |
| 10.0   | 1.34 | 1.31 | 7.64 | 1.75 |
distribution approach ($C_{ox,n}$) yield similar values of true capacitance. Additionally, the surface distribution ($C_{ox,s}$) akin to Brug's formula [68] gives an unrealistically high estimation of $k_{ave}$ values. As the contribution of $R_f$ is negligible in the surface distribution approach, it is probable that the approach will give an erroneous prediction of $k_{ave}$ value when applied to the passive oxide film. Slightly higher $k_{ave}$ values are obtained from the Faraday’s law as the $k_{ave}$ from EIS measurement involves the calculation of film thickness directly resulting from the film, while Faraday’s law requires the anodizing charge (including side reactions) to be used directly for the determination of $d$ values. The additional charge consumed in side reactions such as oxygen gas evolution and metal dissolution affects the $k_{ave}$. Moreover, the mixed matter approach was assumed for the densities and molecular masses of the locally grown oxide film. As the few nm thin films are difficult to characterize, the errors in $k_{ave}$ values from EIS and Faraday’s methods are inevitable. Precise assumptions of material constants, accurate $C_{ox}$ estimation, and negligible side reactions would ensure comparable $k_{ave}$ values from EIS and Faraday’s law.

Fig. 8 Bode and Nyquist plots recorded during simultaneous CV and EIS measurements along the Al–Gd alloy library during film growth for a, b Al–2.0 at. % Gd alloy, c, d Al–9.5 at. % Gd alloy. The data were recorded after each 1 V incremental in CV to a maximum of 10 V.

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Dielectric constant

The $\varepsilon_r$ (Fig. 10b) was determined from the slope of $V$ plotted against $C^{-1}$. Figure 10b shows that $\varepsilon_r$ values are ranging from 13.5 to 23.0 for Al–Gd alloys indicating that addition of Gd to Al improves the dielectric constant of pure alumina.

Because of material’s constant assumption in mixed matter theory, a 10% error bar is also included in Fig. 10b. $\varepsilon_r$ of anodic films containing mixed oxides lies between the values reported for pure $\text{Al}_2\text{O}_3$ ($\varepsilon_r$ for $\text{Al}_2\text{O}_3 = 9–13$) and pure $\text{Gd}_2\text{O}_3$ ($\varepsilon_r$ for $\text{Gd}_2\text{O}_3 = 10–24$ on single and polycrystalline thin-films) [24, 27]. $\varepsilon_r$ values of both $\text{Al}_2\text{O}_3$ and $\text{Gd}_2\text{O}_3$
in phosphate buffer (pH 8.20) using SDCM approach were found out to be 12.7 and 20.5 respectively. This result is a prominent example of the advantage of combinatorial investigations over common batch specimen production, where the property of interest ($r$ in the present case) can be readily identified for wide range of compositions with high accuracy, since the history of preparation, surface treatment, air contact, and other steps are identical. The variation of the screened property can be solely attributed to the difference in composition.

The efficiency of film formation, composition, and nature of amorphous and crystalline phases within the mixed structure can be estimated by a combination of FIB-TEM, XPS, RBS analyses which in the present case is not possible owing to the ultra-thin anodic films. At such a low value (~1–15 nm), obtaining a smooth RBS spectrum and its further simulation is a very challenging task. No visible sign of gas evolution, extremely low dissolution of rare-earth elements in phosphate buffer solution, and detection of negligible Ho (14 ppb) using the SDCM-ICPSOES approach indicates the negligible consumption of ionic current in various side reactions [59, 64].

Conclusions

In summary, a combination of a high-throughput simulation and high-throughput scanning was investigated in a chosen electrolyte on an Al–Gd compositional spread covering a range from 1.0 to 10. at. % Gd. The results of the present study must be interpreted in terms of used electrolyte and special SDCM geometry. The main conclusions are as follows:

1. Novel E-pH and phase diagrams were simulated by coupling conventional and first-principles approaches, and it directly allowed us to predict the domains of thermodynamic predominance for rapid identification of electrolytes on Al–Gd alloys in a given aqueous system.

2. XRD patterns did not show additional phase formation in mixed alloy chemistries except the reduction of Al(111) peak intensity, revealing the loss of crystallinity in the chosen composition range.

3. From the SEM images of the as-deposited film, three distinct regions were identified: large grains typical of fcc phase in Al-rich region, disappearance of these grains in the middle of the compositional spread, and evolution of upright needle like morphology in alloys of high Gd concentrations.

4. Composition-dependent changes in the film formation factor and relative permittivity, and inverse capacitance were estimated indicating the influence of Gd addition while maintaining the passivation behaviour of alumina. These changes were linear and could be directly correlated to the nanoscopic change of the morphology as a function of alloy composition.

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