Thermostimulated luminescence of plasma electrolytic oxidation coatings on 6082 aluminium surface

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ABSTRACT
For decades, plasma electrolytic oxidation (PEO) coatings are actively studied and applied to protect the surface of various valve metals from chemical or mechanical damage. However, over the last couple of years intense research is being done to explore additional possibilities of the PEO coatings apart from their classical application. One of the possible additional uses is thermostimulated luminescence dosimetry that is already widely applied for environmental and health monitoring. This research proposes a method to produce a novel functional coating on aluminium surface exhibiting intense thermostimulated luminescence signal that could be used for dosimetry of ionizing radiation. The result was achieved using plasma electrolytic oxidation with modified electrolyte to introduce carbon ions into the oxide thus inducing defects in the crystalline structure of the coating. Al6082 aluminium alloy was used as a substrate, KOH and ethanol mixture as an electrolyte. A bipolar pulsed regime was used for 15 min. The obtained coating combines the desired luminescence properties with a good mechanical stability due to the relatively hard cubic phase of the coating. Scalability of the technology and low production cost makes the coatings prospective for various practical applications.

1. Introduction

Plasma electrolytic oxidation (PEO) is an electrochemical process used to obtain thick (Snizhko et al., 2004) oxide coatings on various valve metals. Although this technology was developed more than 40 years ago (Yerokhin et al., 1999) (Yerokhin et al., 1998) it is still under development and research because of its scientific and commercial value. As the main use of the coatings is to protect metal from the environment, an extensive research is done to improve their mechanical (White et al., 2016) and chemical properties (Liang et al., 2009) or even to produce thermal barriers (Curran et al., 2007).

One of the trends of PEO studies is quickly gaining traction: the addition of the luminescence properties to PEO coatings. Various studies were performed to obtain luminescent coatings on aluminium (Smits et al., 2015), zirconia (Stojadinović et al., 2016) or zinc (Grigorjeva et al., 2015) with the most significant difference between them being the method of coating modification to introduce rare earth (RE) ions or other atoms into the structure of coatings. A large number of variable parameters (Kalkanci and Kurnaz, 2008) (electrolyte, electrodes, various electrical parameters, material used, electrolyte temperature etc.) allow for many ways to implement the doping of the coating. The most straightforward approach uses an already doped material (Smits et al., 2015); however, due to the cost of RE-doped metal and the lack of optimisation this approach is useable only for scientific interest. Another approach is to use the recently developed pore filling method (Zolotarjovs et al., 2016), but despite the great versatility of this method it is not practical for commercial use due to the multi-step nature of the process. Finally, using modified electrolyte (adding some concentration of dopants to the electrolyte mixture) proved to be the most commercially viable and effective approach (Stojadinović et al., 2016); however, there are no results with real commercial value yet.

The main aim of this research is to evaluate the possibility of using PEO coatings for dosimetric applications based on thermostimulated luminescence. Fig. 1 demonstrates the proposed concept of the application. To solve the problem of making a good dosimeter using the PEO method, one can look for inspiration at the already existing dosimeters. One of the most popular commercially available dosimeters is “TLD-500” - alumina doped with carbon. Since the first research of this material as a TSL dosimeter done in 1990 by Akselrod et al. (Akselrod MKortov and Kravetsky, 1990), it was evident that such a material can perform well as a personal TSL dosimeter because it has an intense,
narrow TSL peak (Yang et al., 2008), the intensity of which is proportional to the absorbed dose in a wide range (10^{-6}Gy-1Gy gamma-dose) (Akselrod MKortov and Kravetsky, 1990). An additional advantage, this material can be produced by many different methods (Yong et al., 2013). On top of that, recent studies demonstrate the possibilities and advantages of optically stimulated luminescence dosimetry on TLD-500 (Tanir et al., 2012) (Meriç et al., 2015) (Akselrod et al., 2006).

The PEO process was developed mostly to improve the mechanical properties of the coating therefore there is an abundance of studies where various crystalline phases of alumina were obtained, including the hardest alumina phase (α, corundum) (Khan et al., 2010) (Dehnavi et al., 2014). However, the TSL signal from an unmodified PEO coating in alpha - or any other phase is negligible and cannot be directly used for dosimetry. Therefore, the main problem to overcome is the doping of alumina coating with carbon. As discussed previously, the most commercially viable method is by using modified electrolyte, thus a highly soluble substance should be used as a source of carbon.

To keep the research as close to practical applications as possible, only commercially viable materials and methods were used in the present study: doping through electrolyte, refraining from the use of rare earth ions, and selecting one of the most popular aluminium alloys (Al6082) as the substrate.

2. Experimental details

2.1. Samples

The substrate material (Al6082) consists of 95–98% aluminium with the addition of Cr, Cu, Fe, Mg, Mn, Si, Ti, Zn and other metals with concentrations varying from 0.1% to 1.3%. In this study, a 3 mm thick aluminium sheet was cut into samples of size 25 × 25 × 3 mm³, each having the total surface area of 15.5 cm². All samples were rinsed in acetone and ethanol and dried before the PEO process.

2.2. PEO setup

Custom 5 kW bipolar pulse electric generator ELGOO PEO V3 was used. The device is externally controlled via PC which allows great repeatability and high customization of parameters including pulse timing, voltage, current and their change in time. In this study, samples were prepared using roughly estimated initial parameters and then optimized for best TSL intensity. Only the best sample is presented in this paper.

The electrolyte consisted of 3 components: deionized water, 2 g/L KOH and 60 ml/L ethanol. The additive we chose for this study is ethanol due to its high solubility; however, other carbon – containing chemicals could be used as well. Electrolyte was contained in a double walled water-cooled reaction chamber thus a stable 20 ± 5 °C temperature was maintained.

Square voltage-limited non-symmetrical 5 ms pulses were used with the voltages of +700 V and −233 V. 1 ms pauses were set in-between pulses. The AC regime with the 1/3 ratio of positive and negative pulses was chosen to reduce porosity (Xin et al., 2006). Current density remained constant with minimal decline over the course of 15 min and was equal to 0.6 kA m⁻². Pt plate was used as a counter electrode. Processing time was set to 15 min. The limit on the processing time was chosen based on the optimisation procedure with the luminescence intensity as the main parameter. The observed decrease of the TSL intensity after 15 min is mainly due to the degradation of the electrolyte – evaporation of the ethanol as well as local increase in temperature causing the irreversible effects with slight colour change to yellow.

2.3. Measurements

TSL luminescence measurements were performed on a custom TSL setup consisting of: heating element (heating speed – 12 K/s, range: 293 K-700 K), vacuum chamber (Ilmvac turbomolecular pump providing vacuum higher than 10⁻¹¹ Torr), X-ray tube with W anode operating at 30 kV, 10 mA and detection system with 2 separate modules - first one Andor DU-401 A-BV IDus CCD camera coupled with Andor Shamrock B-303i spectrograph and the second one - Hamamatsu PMT attached directly to the chamber. Dual registration system allows registering the whole spectral distribution of TSL with CCD as well as registering low intensity integral signals with PMT. Spectra measured with CCD were not corrected to accommodate differences in detection at different wavelengths as only the comparison study was performed. Slight differences between XRL and TSL spectral distribution in the PEO sample are partly caused by the different monochromator slit widths – 50 μm in case of XRL (spectral resolution 1 nm) and 2000 μm during TSL measurement (spectral resolution of 15 nm).

The X-ray diffraction (XRD) spectrometer PANalytical XPert Pro was used to determine crystalline structures. Scanning electron microscopy (SEM) was performed on Phenom PRO desktop SEM.

Consumer-grade CMOS camera (Canon EOS 1100D) with 35s exposure time was used for evaluation of practical application of PEO coating (see Fig. 6). A uniform heater BIOSAN MSH 300 with a surface temperature of 300°C was used to heat the samples during the measurement.

The element analysis of coating was performed using transmission electron microscope (TEM, Tecnai G2 F20, FEI) operated at 200 kV. The powder-like TEM samples were prepared by mechanical scraping from coating using diamond cutter and placed on Lacy carbon film grid AGS166 (Agar Scientific).
3. Results and discussion

3.1. Coating characterisation

As it was mentioned previously, undoped alumina PEO coating does not exhibit TSL signal, therefore the first task is to incorporate carbon in the structure. Several approaches were evaluated as the pilot experiments for this study; however, the best result was obtained by the modification of the electrolyte with highly soluble organic substance – ethanol. To verify the implementation of the carbon ions in the coating the TEM EDX measurements were performed (Fig. 2). By analysing and comparing the intensity of $K_{\alpha}$ line of carbon at 0.277 KeV it is evident that the carbon atoms are indeed incorporated in the structure. The wide variations of carbon concentration in doped coatings (Fig. 2, inset) are ascribed with non-uniform distribution of carbon ions in the coating. Even though electron beam for EDX measurements was focused in small areas without any Lacey film underneath the particle, undoped samples still show a small carbon signal, which originates from electron multiple scattering, similarly to the Cu signal from Cu grid.

The knowledge about electronic processes in the sample related to visible luminescence output is crucial for analysis of any possible dosimeter, therefore multiple TSL measurements were performed for both the reference sample (TLD-500) and the obtained C-doped PEO coating (Fig. 3). The PEO coating exhibits a broad TSL peak extending from 300 K up to 630 K (full width at half maximum, FWHM = 150 K). Its width indicates a very broad trap distribution in the lattice, covering a wide energy range. To compare the coating with the existing technology, TSL of TLD-500 was measured and the obtained TSL glow curve and spectral distribution coincide well with the literature (Kalita and Chithambo, 2017a) (Palan et al., 2016) (a Weinstein et al., 2002) – with a characteristic luminescence band at 400 nm and a TSL peak at 440 K associated with F and F$^+$ centres. When comparing both samples (PEO coating and TLD-500) one can see that although the general positions of the TSL maxima are similar, their FWHMs are drastically different. The width of the TSL peak is directly correlated with the energy distribution of a trap – the smaller the FWHM, the more defined activation energy is. Since the carbon-doped PEO coating exhibits extremely broad TSL peak, one can conclude that the possibilities of a localized electron to recombine are broad which is due to irregularity of the atomic structure present in the coating. The difference between highly ordered crystalline structure of TLD-500 and mostly amorphous structure of the coating is evident.

The difference of FWHM of the PEO sample and TLD-500 should be studied in finer details to better understand the nature of the traps.
which was done by measuring the spectral distribution of TSL signal from the samples. Fig. 4 shows the spectral distribution for the TSL maximum at 440 K for both the PEO coating and the TLD-500 samples as well as the X-ray induced luminescence (XRL) spectrum measured from PEO sample under irradiation. (Fig. 4). Although trap centres in the PEO coating and TLD-500 crystal are similar, the luminescence main peak position varies drastically. The luminescence maximum at 420 nm of TLD-500 was studied in depth by Akseidzov et al. and is described as the F\(^{−}\) centre charge compensated by C\(^{2+}\) ions in the α-Al\(_2\)O\(_3\) matrix (Akseidzov et al., 1999). However, this is not the case for the PEO coating – luminescence is shifted to the green region to 530 nm. Similar case was observed by Rodriguez et al. for Al\(_2\)O\(_3\) doped with Mg and C (Rodriguez et al., 2011). Here, F\(_2\)\(^{2−}\) (2 Mg) centre is a result of charge compensation for the substitutional Mg\(^{2+}\) ion in Al\(^{3+}\) site. Mg ion is present in the substrate (aluminium alloy) as well as its presence in the coating is confirmed by the TEM EDX measurements (Fig. 2).

The PEO coating exhibits also the complex red band - a combination of various metal oxide luminescence bands. As the starting alloy (Al\(_6\)O\(_8\)) contains many additives to improve its mechanical properties, they are expected to incorporate in the obtained coating too. TEM EDX results confirm the presence of other (apart from Al) metal ions, like Mg that are implemented in the coating from the substrate metal.

As an example, Mg (Kalita and Chithambo, 2017b), Mn (Martínez-Martínez et al., 2005) or Cr (Salek et al., 2016) oxides exhibit luminescence in the red region of the spectrum; however, due to the amount of co-dopants in the coating it is hard to distinguish any particular lines apart from the most intense Cr\(^{3+}\) R-line doublet at 693–694 nm.

The assumption of the irregular structure in the coating is based on the relatively short PEO time (15 min are usually not enough for the formation of corundum) as well as on the FWHM of TSL. To address this assumption, the XRD spectra of coatings obtained with and without ethanol in electrolyte (all other parameters were kept unchanged) were compared (Fig. 5). One can see that apart from the main aluminium peaks at 38.44, 44.69 and 65.07 deg. (from the substrate), several others are present in the diffraction pattern of the coatings. The small peaks visible in 39–44° region (as well as peak at 48.7°) originate from the different phases present in the alloy itself (Al\(_6\)O\(_8\)), e.g. Mg2Si phase and various intermetallic phases. After comparing data to the XRD database (Graulis et al., 2009), it was concluded that peaks at 45.87 and 66.86 deg. are due to the presence of eta- or gamma-phase in the alumina coating and correspond well with the JCPDS cards #29-0063 and #50-0741 (marked with arrows on Fig. 5). Some additional crystalline structure is present in the sample without ethanol and the addition of ethanol in electrolyte prevents the crystalline phase from occurring due to the increased defect concentration from incorporation of C atoms.

Although many studies reported the possibility to obtain alpha-phase aluminium oxide with PEO using longer processing time (Yerokhin et al., 2003) (Tian et al., 2002), the optimisation of the parameters in this study revealed the loss of TSL intensity with increased PEO time, therefore this approach does not allow synthesis of trigonal alumina dosimetry coating. Additional studies on optimizing the electrolyte composition, temperature and time are needed to obtain α-phase alumina by PEO.

The thickness and morphology of the “ethanol-processed” coating were studied using SEM. (Fig. 6). It is evident that it looks like classical (undoped) PEO coating with a thickness of about 20 μm. Although some porosity is present, it is characteristic for PEO and can be used to an advantage (Curran and Clyne, 2006). The presence of ethanol in electrolyte does not seem to affect the coating morphology.

### 3.2. Practical application evaluation – delayed radiation mapping

By analysing the output intensity of the best PEO sample a conclusion can be drawn that coatings of this type exhibit easily measurable intense TSL signal in the visible light region. The possibility to use coatings as radiation detecting screens is demonstrated (Fig. 7). The concept is to apply the PEO coating to a large flat surface (e.g. aluminium sheet) and use it to get the distribution of ionizing radiation in experimental/medical/industrial setups. Due to the limitations of the power supply used in this study, the maximum surface area of the samples is limited to 15.5 cm\(^2\). To overcome this limitation, 4 identical samples were prepared and arranged in a grid (2 by 2 samples) giving the total active area of 25 cm\(^2\) (one side of the whole 2 × 2 arrangement). The grid was then irradiated with X-rays for 30 min (W anode, 30 kV 10 mA) from 10 cm distance. After the irradiation, the grid was placed on a pre-heated uniformly hot surface (T = 300°C) and the luminescence from the grid was measured using a CMOS sensor in a commercially available camera with a long exposure (35s). Fig. 7 shows the obtained colour-corrected image.

In addition, by analysing a colour version of the image one can see that in the irradiated part of the grid tiles have a non-uniform colour distribution of the luminescence with the edges of the tiles emitting orange-red light and middle parts emitting green luminescence. The luminescence at the edges corresponds to the red band previously connected with ions of various metal additives in alumina coating, while the middle part of the tiles is dominated by the intrinsic luminescence of alumina. The difference between the luminescence output at the edges and in the middle is explained by the increased current density in the regions close to the sharp edges of the tiles.
Fig. 7. TSL integral output image of X-ray irradiated C-doped PEO coating, taken by camera with CMOS sensor.

4. Conclusions

A new functional material was developed combining advantages of PEO alumina coatings (high mechanical and chemical stability, strong adhesion, ease of production) with an intense TSL signal, partly similar to the TSL of alumina-based dosimeters. Commercially viable approach to the TSL of alumina-based dosimeters (high mechanical and chemical stability, strong adhesion, ease of production) with an intense TSL signal, partly similar to the TSL of alumina-based dosimeters.

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References


