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Towards Smart Electrolytic Plasma Technologies: an Overview of Methodological Approaches to Process Modelling

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Abstract
This paper reviews the present understanding of electrolytic plasma processes (EPPs) and approaches to their modelling. Based on the EPP type, characteristics and classification, it presents a generalised phenomenological model as the most appropriate one from the process diagnostics and control point of view. The model describes the system ‘power supply – electrolyser – electrode surface’ as a system with lumped parameters characterising integral properties of the surface layer and integral parameters of the EPP. The complexity of EPPs does not allow the drawing of a set of differential equations describing the treatment, although a model can be formalised for a particular process as a black box regression. Evaluation of dynamic properties reveals the multiscale nature of electrolytic plasma processes, which can be described by three time constants separated by 2-3 orders of magnitude (minutes, seconds and milliseconds), corresponding to different groups of characteristics in the model. Further developments based on the phenomenological approach and providing deeper insights into EPPs are proposed using frequency response methodology and electromagnetic field modelling. Examples demonstrating the efficiency of the proposed approach are supplied for EPP modelling with static and dynamic neural networks, frequency response evaluations and electromagnetic field calculations.

Keywords: Electrolytic plasma processing; Smart technology; Phenomenological model; Process diagnostics; Process control; Frequency response; Electromagnetic field

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<tbody>
<tr>
<td>AC</td>
<td>alternating current</td>
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<tr>
<td>COMSOL</td>
<td>commercial multiphysics modelling software package</td>
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<td>DC</td>
<td>direct current</td>
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<td>EIS</td>
<td>electrochemical impedance spectroscopy</td>
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<td>EP</td>
<td>electrolytic plasma</td>
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<td>EPCS</td>
<td>electrolytic plasma coating stripping</td>
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<td>EPHT</td>
<td>electrolytic plasma heat treatment</td>
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<td>EPP</td>
<td>electrolytic plasma process/processing</td>
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<td>EPPo</td>
<td>electrolytic plasma polishing</td>
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<td>EPSC</td>
<td>electrolytic plasma surface cleaning</td>
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<td>FR</td>
<td>frequency response</td>
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<td>GRNN</td>
<td>general regression neural network</td>
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<tr>
<td>Hi-PIMS</td>
<td>high-power impulse magnetron sputtering</td>
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<tr>
<td>NARX</td>
<td>nonlinear autoregressive network with exogenous inputs</td>
</tr>
<tr>
<td>NDR</td>
<td>negative differential resistance</td>
</tr>
<tr>
<td>PEC/N</td>
<td>plasma electrolytic nitrocarburising</td>
</tr>
<tr>
<td>PE-CVD</td>
<td>plasma enhanced chemical vapour deposition</td>
</tr>
<tr>
<td>PEO</td>
<td>plasma electrolytic oxidation</td>
</tr>
<tr>
<td>RMS</td>
<td>root mean square (value)</td>
</tr>
<tr>
<td>SCR</td>
<td>surface charge region</td>
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<tr>
<td>VGE</td>
<td>vapour-gaseous envelope</td>
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1. Introduction

Electrolytic plasma processes (EPPs) have recently gained significant attention from coating and metal finishing industries due to their capability to considerably enhance surface properties and environmental compatibility [1, 2]. Being a progression of conventional electrochemical treatments into high voltages, EPPs feature electrolyte boiling and/or discharging phenomena in the vicinity of the working electrode [3, 4]. They are also known to exhibit highly non-linear behaviour manifested in current-voltage and boiling characteristics featuring inflections, extremes and sometimes negative slope regions [1]. These non-linearities may explain why EPPs could in practice surpass their conventional counterparts but represent major challenges for their theoretical description. As will be discussed below, although a range of partial EPP models has been developed, none of them covers all varieties of processing conditions ranging from plasma electrolytic oxidation (PEO) [5] to coating stripping (EPCS) [6] and from heat treatment (EPHT) [7] to surface cleaning (EPSC) and polishing (EPPo) [8].

In the meantime, electrolytic plasma treatments are highly energy intensive and achievement of optimum surface properties with minimum energy and time consumption is extremely important from both economic and environmental viewpoints. The solution of such problems in advanced manufacturing is often achieved by the use of ‘intelligent’ or ‘smart’ technologies incorporating elements of active diagnostics and feedback loops that would allow real time adjustment of process parameters according to a pre-determined model linking them to desirable surface properties (Fig. 1). Moreover, variability of modern manufacturing requires technology to be adaptive, with process parameters being easily adjustable to different batch sizes and part geometries; this is also addressed via incorporation of generic models of the processes into control circuits. It is not surprising to see smart elements, such as active plasma diagnostics and variable process parameters, in recent developments of plasma-based surface engineering technologies, e.g. PE-CVD [9, 10]
and Hi-PIMS [11, 12]; for EPPs however this still remains in a rudimentary state despite some recent progress [13].

This review puts together information regarding electrolytic plasma processes in a structure which helps in understanding their generic features and corresponding electrical characteristics that are important in different practical applications as well as retrieving observable and controllable variables that contribute to the process diagnostics and control. It is also intended to provide a perspective into further possible developments based on the phenomenological approach as well as achieving better insights into electrolytic plasma processing.

2. Classification of electrolytic plasma processes and corresponding models

2.1. Brief characteristics of electrolytic plasma processes

Electrolytic plasma treatments were introduced in the 1950-s [14, 15], although the first reports of underlying physical effects and phenomena date back to the 19th century [16]. These processes are used for surface finishing, including case hardening, nitriding and carbonitriding, [3, 7, 17-20], surface oxidation [21-25], cleaning [26, 27] and polishing [28-30] as well as stripping of defective coatings [6, 31].

EPPs operate in the voltage range 80-1000 V which is higher than that for conventional electrochemical treatments. Aqueous solutions of salts, acids or alkali with low concentration (from 1 to 20%) are used in most cases, and these electrolytes are usually non-toxic and easy to recycle. In the beginning of the process, high current density at the surface of the working electrode (which is usually smaller than the counter electrode) leads to intensive electrolyte evaporation; electrochemical gas evolution and oxide formation also take place.

Depending on oxide film conductivity and semiconductor properties, the formed Schottky type metal-oxide-electrolyte two junction system can be forward or reverse biased [32, 33]. With the forward biased junction, the EPP features a vapour gaseous envelope (VGE) surrounding the working electrode. Most of the voltage drop occurs
across the VGE which is a non-stationary object with a high field, promoting a plasma discharge in the gas media. With the reverse biased junction, the EPP on the anode features formation of a stationary oxide film which attains the majority of the voltage drop. Upon achieving a breakdown field strength in this thin film, microdischarges appear on its surface. These phenomena distinguish EPPs from other electrolytic processes and justify the use of term ‘plasma’.

Visualisation of EP processes has been improved by the availability of high speed digital video recording technology. For the processes with the VGE, typical images are presented in Fig 2 [34, 35], illustrating a progression from gas bubble evolution (a) to VGE formation (b) and microdischarge development (c,d) which results in surface polishing. Typical images of microdischarges in the pores of oxide films are provided in Fig. 3, demonstrating that both average size and population density microdischarge events evolve with treatment time in a complex manner [36-39].

The main characteristics of EPPs are the current-voltage relationship [1, 40, 41] and boiling characteristics [1, 42]. Both are substantially non-linear and exhibit at least one range with a negative slope. This indicates that EPPs are complex and non-linear systems that can be described by neither electrochemical nor plasma models alone; therefore, a synergetic approach should be adopted for modelling of these processes.

2.2. Process classification
EPPs can be categorised by several features as shown in Fig. 4 and discussed below.

2.2.1. Working electrode polarity
By the polarity of the working electrode, cathode [43] and anode [44] treatments can be distinguished, which are different in the electrochemical processes involved and the electron emission mechanism responsible for discharge generation.

The electrolysis of aqueous solutions in EPP exhibits several electrode processes. Common anode processes include oxygen evolution (1) and metal
oxidation (2). Depending on electrolyte chemical activity, the latter could result in either surface dissolution or formation of an anodic oxide film (3). Hydrogen evolution (4) and/or cation reduction (5) occur on the cathode surface:

\[
\begin{align*}
2H_2O - 4e^- & \rightarrow O_2↑ + 4H^+ \\
Me - ne^- & \rightarrow Me^{n+} \\
xMe^{n+} + \frac{3}{2}xnH_2O & \rightarrow Me_xO_n + xnH_3O^+ \\
2H^+ + 2e^- & \rightarrow H_2↑, \\
Cat^{n+} + n\bar{e} & \rightarrow Cat^0\downarrow
\end{align*}
\]

For cathode EPP, the working electrode area must be less (by at least an order of magnitude) than that of the anode, to provide high local current densities promoting electrolyte boiling. This, together with hydrogen evolution, contributes towards formation of VGE, whereas deposition of reduced cations changes electrode surface composition. The metal electrode in the cathode treatments can be heated up to 1000 °C, which enables plasma electrolytic carburising and nitrocarburising (PEC/N) and other diffusion-based treatments [45, 46]. For the anode EPPs, the electrode area ratio is less critical. Instead surface passivation of the working electrode is important which can lead to the growth of oxide films according to reaction (3), giving rise to a wide variety of PEO processes [47-49], or if (3) is overcome, to the surface cleaning and polishing to a mirror finish [50] as well as case hardening [51, 52], nitriding [3] and coating stripping [6].

Pulsed EP treatments, wherein the voltage is supplied as rectified sine or square waves, have been gaining significant attention since early 1980’s. AC and pulsed bipolar EPPs, in which anodic and cathodic polarisation pulses are periodically interleaved, thus combining the benefits of the both treatment modes [53, 54], often appear to be more effective than DC ones. The bipolar modes have more degrees of freedom and are therefore harder to optimise, but can provide better performance of the modified layer [18, 55, 56].
2.2.2. Metal substrate type and the main voltage drop in the system

For PEO treatments, the type of metal substrate is of high importance as various oxides can be formed on the anode surface according to (3) [32]. Valve metals (Al, Ti, Mg, Ta, Zr) form n-type semiconductor oxide films due to the presence of anionic vacancies at the metal-oxide interface and a space charge region (SCR) within the film [57]. The SCR width depends on the concentration of charge carriers, oxide relative permittivity, interfacial potential drop, magnitude and polarity of applied voltage. At the anodic polarisation, the surface layer exhibits low conductivity (the valve effect) and the major voltage drop occurs across it. With the SCR thickness usually being less than a micron, the electric field there at voltages typically applied in EP treatments can reach breakdown values (~ 10^7 V/cm) [33]. PEO treatments can therefore be easily implemented to grow protective coatings on the valve metals [2, 58]. For other metals, such as Fe, Cr, Ni, W, Cu etc., that either form p-type oxide films (3) or do not form stable oxides at all (2), high anodic potentials result in VGE formation, promoting non-oxidising treatments, e.g. polishing [59], although recent developments of fluorine containing electrolytes also help in carrying out polishing of valve metals [60].

2.2.3. Processing conditions and required surface properties

EPPs can also be categorised into several types of processes (as indicated in Sections 1 and 2.1) intended to achieve different surface characteristics and properties, requiring different processing conditions and parameters to be controlled.

EPHT [3, 46] requires film boiling in the VGE. This results in thermal shielding of the working electrode; thus, the power dissipating in the VGE due to Joule heating contributes to a rapid increase of the surface temperature up to ~1000°C in several seconds [61]. If the treatment is terminated at this stage, case hardening can be implemented after the voltage is cut off and the working electrode is left in the electrolyser. Fig. 5a shows typical cross section of a surface layer after EPHT; a decrease of the grain size is evident in this image thus confirming the case hardening
For prolonged EPHTs, the bulk of the working electrode can be heated up in order to achieve a desired microstructure and/or phase transformation.

PEO is used to form oxide ceramic surface layers with different thicknesses, phase compositions, protective and functional properties [1, 2]. Typical coating morphologies are shown in Fig. 5b and 5c for Al and Ti alloy substrates [62, 63]. Localised microdischarges contribute to the formation of high temperature oxide phases, such as rutile and corundum, within the coating, providing hardness and chemical stability [64, 65]. Porosity could provide the coating with useful functionality, e.g. by increasing specific surface area; moreover it may contribute to internal stress state and improve coating adhesion by reducing its global stiffness, but could also affect such important properties as wear and corrosion resistance [66].

EPPO requires bubble boiling in the VGE [67], which helps to decrease the surface roughness down to \( Ra \approx 0.05 \) \( \mu \)m. Although the initial surface state defines the minimal roughness achievable by the treatment, a 4 to 5 fold improvement in the surface finish can be realised [68]. A typical surface morphology after EPPO is shown in Fig 5d; traces of original surface topology can still be discerned, smoothed significantly by the treatment [13].

EPSC removes rust, oxide scale or organic contaminants [8, 26, 27]. It operates at cathodic polarisation of the working electrode. Spark discharges in the VGE provide cavitation effects at the electrode surface as well as heating, melting and resolidification of a thin surface layer [8]. As a result, the contaminants are removed by pressure waves or are decomposed in the plasma discharge by thermolytic and/or radiolytic mechanisms. As shown in Fig. 5e, a typical surface morphology exhibits micro-craters and spheroidal nodules [8], resulting in a typical 3 to 4 fold increase in surface roughness. This however can be mitigated by application of pulsed current providing control over the impact of plasma discharge on the surface [27]. Formed amorphous or nanostructured surface layers could increase corrosion and erosion resistance [26] but reduce the fatigue strength of the material [27].

EPCS operates at the anodic polarisation of the working electrode under the transient type of boiling in the VGE [6, 13]. These conditions are similar to the EPPO
except for anodic dissolution which is intensified by the boiling conditions, hence surface composition, roughness and dimensional tolerance are important. Typically, an aluminide coating can be stripped at a rate of 2 to 3 \( \mu \text{m}\cdot\text{min}^{-1} \) [69]. The stripping proceeds gradually (Fig 5f), with no pitting corrosion developing on the surface. After the stripping, a new coating can be applied on the component, thus prolonging its operating life.

### 2.2.4. Process arrangement

Electric energy consumed during EPP lies within the range 300 to 500 W/cm\(^2\) (4 to 5 kW·h·m\(^{-2}\)·\(\mu\text{m}^{-1}\)) [25]. Therefore, a technological constraint of \( \leq 0.5 \text{ m}^2 \) is usually imposed on the treated surface area. Smaller surfaces can be treated as a whole, using bath immersion configuration (Fig. 6a), whereas spraying processes carried out with a specially designed counter-electrode tool (Fig. 6b) are usually applied to sequentially treat larger surface areas. Most EPPs are implemented in the bath; in spraying processes, the electrolyte is pumped through holes focused by a dielectric nozzle contacting the workpiece. The VGE appears where the electrolyte touches the treated surface as shown in Fig. 5 for EPHT. A range of counter-electrode tool designs has also been developed for PEO of large scale marine components [70].

### 2.2.5. Power supply type

All EPPs require high voltage and high current power supplies for their realisation. For the processes requiring stable VGE (i.e. diffusion, heating and case hardening, stripping and polishing), voltage-controlled sources are normally used, i.e. the power supplies with low output impedance achieved by a large capacitor connected in parallel with the electrolyser or/and by application of voltage stabilising feedback loops. These requirements were well established in early works [71, 72]. EPPs involving oxide film growth can also be sustained by a voltage-controlled source. If the voltage is stabilised at a certain level, the current changes in time depending on the electrolyte temperature, surface properties and other process parameters. The typical shape of the current curve is shown in Fig. 7a. For the
processes with VGE, the current decay can be attributed to the electrolyte heating if no cooling is provided; for oxidation treatments, the current decreases due to the layer growth causing an increase in its resistance, if no structural defects are formed within it. For the latter processes, current-controlled sources can be preferred provided that the relation of oxide growth to Faraday’s law within the operation window is confirmed. Such supplies contain power inductors in series with the electrolyzers and feedback circuits for current stabilisation [73]. The process evolution is usually characterised by the voltage transient behaviour (Fig. 7b) which can be linked to the coating growth.

For pulse and bipolar EPPs, such categorisation also stands, since the pulse unit itself is fed by a voltage or current source. These processes can be described by the behaviour of average or RMS values of corresponding electrical parameters as shown in Fig. 7, although a more sophisticated analysis of dynamic voltage-current curves and/or voltage/current transients may also be performed [41, 74-76].

2.3. Model classification

The EPP models can be arranged into groups as shown in Fig. 8.

VGE models using differential equations have been proposed for polishing and heating processes which do not feature formation of oxide layers during the treatment. These quite complex theoretical models involve variable thermal-physical parameters related to the electrolyte flow [67] and VGE shape [77].

A wide range of phenomenological models have been developed for EPHT and PEO processes. Accounting for electrolyte boiling characteristics [42] helps to explain the complex dynamics of the electrode temperature which could rise from 100 to 1000 °C in a few seconds. Further development provided a group of heating modes [7] which can be used to better understand how to control the electrode temperature during heat treatment. Relationships among different EPPs discussed in [1] form a phenomenological basis for EPHT, PEO and other processes. A phenomenological model of coating stripping [78] can also be used to understand
various EPPs. Microdischarge evolution in the pores of oxide layers and oxide crystallisation phenomena during PEO processes have been discussed in [79-81].

One group of theoretical models focuses on metal-dielectric-electrolyte interfaces [33] and analyses of impedance spectra [82, 83]. This group is tightly connected with equivalent circuit models which can be obtained from the impedance spectra and current-voltage characteristics. The circuits formed by resistive elements [84] are mainly used for DC EPPs, whereas those made by resistive and capacitive elements [85, 86] can be used for pulsed PEO and the nonlinear circuits [80] which apply to AC PEO conditions.

There are also empirical models which include the approximation of voltage transients are used for galvanostatic DC PEO processes [87]. They also include approximations of dynamic and pulsed current-voltage characteristics, used for AC and pulsed PEO processes respectively [41, 81]. Empirical regression models help to estimate dependencies between the process parameters (voltage, current density, electrolyte temperature) and the surface properties, e.g. coating thickness and roughness. A range of such models can be found in the literature for different EPPs [68, 74].

Thus the most developed models are those for EPHT and PEO processes, for which phenomenological and empirical approaches dominate. However empirical models provide an insufficient level of generality and the phenomenological approach appears to be more generally applicable. The main problem with the latter approach lies in a lack of means to formalise the behaviours observed in experimental studies. For example, the descriptive phenomenology of microdischarges during PEO processes has not changed since 1990s [79] to 2000s [36] and to recent publications [88], but no general models quantitatively describing the process have been proposed yet. Therefore, the phenomenological approach pursued here includes two steps: a study of the generalised structure of the model and its numerical solution for selected EPPs using modern regression tools.
3. Phenomenological modelling of electrolytic plasma processes

3.1. Generalised structure

Based on the above analysis, a generalised phenomenological model structure has been proposed (Figs. 9 and 10). The structure is a further development of the phenomenological model presented earlier [78]. The model considers integral EPP characteristics at a macro level for volumes substantially exceeding the Debye radius. The generalised phenomenological model is based on the systematic analysis of EPP mechanisms, and their further decomposition into subsystems.

The following EPP features were considered in the model: oxidation, heating, polishing, and coating stripping, all in aqueous electrolytes. The model describes the system ‘power supply – electrolyser – treated surface’ as a system with lumped parameters which characterise integral properties of the surface layer and integral parameters of the process.

3.1.1. Major subsystems

The upper level of the model contains the major subsystems joined into two contours: (i) electric current flow: working electrode, its surface layer, vapour gaseous envelope, electrolyte, counter electrode and power supply; (ii) heat exchange: electrolyte, heating and cooling sources. The structure of the first contour reflects the fact that the voltage provided in EPP generally comprises thermodynamic kinetic and ohmic components. The former two are essential to achieve potentials at which the discharge can be sustained in principle and ensure that the desired process proceeds with required rate; these are addressed by the surface layer and VGE subsystems. The ohmic component is associated with electrical characteristics other subsystems as discussed in Section 3.1.3. The structure of the second contour reflects the need to maintain the required temperature in the system during processing. The first level describes processes in the subsystems (Fig. 9), the second and the third levels contain electrical and other measurable and controllable characteristics of the subsystems (Fig. 10).
3.1.2. Processes in the subsystems

On the first level of the model, electrodes 1 and 5 feature anodic and cathodic processes typical of any electrochemical treatment. If the working electrode is positively polarised, it features anode processes 1.1.1, whereas the counter electrode features cathodic processes 5.1.1, and vice versa (1.1.2 and 5.1.2 respectively). Additionally, the processes of thermionic emission 1.1.3 specific to cathode EPPs may occur on the negatively polarised working electrode. Moreover, during cathode EPHT the working electrode also features metallurgical processes 1.1.4.

The processes in the electrolyte 4 include processes 4.1.1 that are similar for any kind of electrochemical treatment at any polarisation, and processes 4.1.2-4.1.4 characteristic of EPPs only. The latter reflect phenomena occurring at the partial electrolytic cathode and anode formed by the VGE-electrolyte interface during anodic and cathodic polarisation of the working electrode, respectively. Some processes in this category are common (4.1.2), other depend upon electrode polarity. For example, formation of hydrated electrons and their emission into the VGE with further avalanche evolution of an electric discharge in it occur only at the electrolytic cathode 4.1.3. Otherwise, the electron emission proceeds from the surface of the metal electrode and the current in the VGE flows in the opposite direction. This leads to much more intensive radiolysis at the electrolytic anode 4.1.4.

The processes in the VGE 3 are characteristic to EPPs only. Boundary processes 3.1.1 contribute to the VGE formation and evolution during plasma electrolysis. Importantly, in some cases, e.g. during PEO, these could not be intensive enough to form a continuous VGE. A vapour gaseous envelope can be characterised by three types of boiling, with changes in the boiling type occurring intermittently according to the boiling curve [42]. The processes within VGE 3.1.3 and plasma processes 3.1.4 describe the influence of the vapour gaseous envelope on the treated surface. The VGE can be perceived as analogous to the electrode gap in electrochemical machining; therefore, the process control for EPPs with VGE is predominantly a control of the vapour gaseous envelope properties.
The processes in the surface layer 2 are the most complex in the system, and they are individual for any EPP type. These include surface layer formation 2.1.1 by different mechanisms similar to electrochemical treatments, as well as plasma-assisted and metallurgical processes 2.1.2 and 2.1.3 are specific to EPPs. The latter transformational processes define the final properties of the surface layer after the treatment.

Note that control of processes 2.1 in the surface layer can be achieved only indirectly via the processes 3.1, 4.1, 1.1 and 5.1 (in descending order of influence) by controlling the electric power supply 6 and the heat power supply 7, provided that the properties of the subsystems 1–5 are identified.

3.1.3. Electrical characteristics of the subsystems

The second layer in the structure of the phenomenological model comprises the electrical characteristics of the subsystems (Fig. 10). This arrangement reflects the fact that the electric power is the major driving force for surface modifications during EPPs. For all subsystems, the electric phenomena, equivalent circuit and electromagnetic field properties should be considered. Note that within the phenomenological approach to the analysis of integral characteristics, the electromagnetic field properties are considered as averaged over the corresponding subsystem.

Electrical characteristics of both electrodes 1 and 5 are defined by corresponding resistances which are often considered insignificant due to electronic conductivity prevailing in metals. The interfacial electric double layer appearing at the counter electrode is represented as a voltage source in the equivalent circuit. The other dissimilarity appears in the average current densities, because the working electrode usually has at least an order of magnitude smaller surface area. At the same time, electrolyte 4 has ionic conductivity which is typically an order of magnitude lower and shows almost resistive impedance behaviour in the frequency band from several Hz to tens of kHz. Defining the current density distribution in the electrolyte would
help in assessing the net electrolyte resistance for a certain configuration of the electrolyser.

Electrical characteristics of the VGE 3 are dominated respectively by ionic and electronic conductivity in the absence and presence of a plasma discharge. These conductivity mechanisms define the value of a corresponding resistive element in the equivalent circuit. One of the main features of the VGE is negative differential resistance (NDR) which appears in certain regions of current-voltage characteristics [1] owing to the following two reasons: (i) the VGE thickness increases with increasing voltage due to higher Joule heat, and the current decreases; (ii) current-voltage characteristics of plasmas developed in electric discharges that occur in the VGE may exhibit NDR. If a continuous VGE is present, almost all supplied voltage drops across it and this induces a strong electric field in the VGE, initiating a distributed electric discharge visible by an unaided eye.

Electrical characteristics of the surface layer 2 depend on its conductivity which can be of electron (for metals and some oxides, nitrides and carbides), electron-hole (for oxides having semiconductor properties) or ion (for microdischarges in pores) type. Relevant resistances in equivalent circuits can correspond to electronic and ionic conduction, whereas NDR could reflect the characteristics of electric discharges in the pores, and capacitances can be attributed to surface layers with low specific conductivity. If a continuous oxide layer is present on the surface, almost all supplied voltage will drop across it, initiating a strong electric field which would trigger localised microdischarges in the layer pores and other defects.

Unlike the solid surface layer with low specific conductivity, the VGE is a ‘quasi-stationary’ object existing only during the treatment. Therefore, if the oxide film is present on the electrode surface before the treatment, a continuous VGE is not usually formed. This differentiation can be illustrated by distinct differences between the two anodic DC treatments, EPPo, wherein a bubble boiling within the VGE and a glow discharge at the electrolytic cathode are observed, and PEO featuring microdischarges localised in the coating pores. Advanced EPPs, e.g. bipolar pulsed PEO or EPCS of a TiN coating, combine voltage drops within the VGE and the
dielectric surface layer at different stages of the treatment. The models of these processes are the most complex. Moreover, in such complex systems, it is not straightforward to distinguish the VGE from the surface layer and unambiguously identify the type of discharge and the subsystem where it primarily evolves.

3.1.4. Measurable and controllable characteristics of the subsystems

Each subsystem can be characterised by parameters measurable during the treatment (observable) and after it (unobservable). Both of them are controllable since they can be influenced by variations in voltage, current and electrolyte temperature imposed by an operator.

The electrodes 1 and 5 have different requirements. If the counter electrode has to maintain its chemical and phase composition 5.3.1, the requirements to the working electrode – i.e. the component being treated – are more formalised in terms of physical, chemical and mechanical properties 1.3.1 that are measurable only after the treatment. During the process, however, it is possible to measure only a limited number of characteristics, such as electrode temperature 1.3.2. For the electrolyte 4, the majority of its electrochemical, optical and other properties 4.3.2 can be measured during the treatment, while the chemical composition 4.3.1 could be accurately assessed only after the treatment. Since VGE 3 does not exist apart from during the EPP, only in-situ measurements 3.3.2 are possible. Significant information could be gained from visualisation techniques which supply spatial characteristics [37, 89], type of boiling [90], optical emission intensity [4, 91, 92] and spectral content [38, 93, 94] associated with the VGE. Acoustic techniques could be also informative [95, 96].

Measurable properties of the surface layer 2 are the required EPP target characteristics. Usually they are assessed by advanced physical, chemical and mechanical techniques; therefore, their conventional measurement in-situ is impossible 2.3.1. These target characteristics could also overlap with the properties of the working electrode 1.3.1; however, the latter commonly act as constraints (e.g. the constraint on the substrate phase composition change during the deposition or
stripping of a coating). Therefore, to retrieve observable and controllable variables contributing to the EPP diagnostics and control, it is necessary to apply indirect techniques wherein information regarding properties 2.3.1 is extracted using evolution of estimates 3.3.2, 5.3.2, 1.3.2, 4.3.2 obtained e.g. from measured electrical characteristics 1.2.2, 2.2.2, 3.2.2, 4.2.2, 5.2.2 via justified equivalent circuits [75, 97].

3.2. Formalisation possibilities

The complexity of the electrolytic plasma system does not currently allow formalisation in the form of differential equation sets describing the treatment of real components in industrial electrolytes. These sets cannot be formalised comprehensively and therefore still remain unresolved. As a result, a phenomenological approach to EPP description and black box regression models using experimentally obtained data appear to be justified. These models can be realised using different means of regression analysis – from linear regression equations to neural networks [98, 99]. All these means perform a non-linear transformation of the space of inputs into the space of outputs.

A typical model structure is shown in Fig. 11a. Column vector of inputs $\mathbf{X}$ corresponds to the parameters of electrical and heating power supplies 6 and 7 in Figs. 9 and 10; these parameters must be controllable in order to have reproducible outputs. The inputs could include electrical regimes (e.g. DC voltage value, or pulse amplitude, frequency, duty cycle and other parameters describing the voltage waveform) and electrolyte temperature (either initial value or that stabilised by a thermostat). Procedures of data scaling to the range $[-1; +1]$ and descaling help to improve the model quality and contribute to the significance analysis of the input variables (Fig. 11b).

The time variable $t$ can also be an input of a regression model; this model is of a static type, and the physical meaning of this variable is the treatment duration. For a dynamic model the time variable should not appear in the input. Instead, within the phenomenological approach pursued here, dynamic models are obtained for finite
differences, and the time variable is included indirectly as a discrete time step as in [100].

The column vector of outputs $\mathbf{S}$ comprises the surface properties after the treatment and corresponds to blocks 2.3.1 and 1.3.1 in Fig. 10. These output variables evolve in long time scale so that corresponding time constants range into several minutes. The column vector of outputs $\mathbf{C} = [\mathbf{C}_1; \mathbf{C}_2]$ includes variables measurable during the electrolytic plasma process, and it could contain characteristics from blocks 1.3.2, 3.3.2, 4.3.2 and 5.3.2 in Fig. 10. Instantaneous values of these characteristics form vector $\mathbf{C}_1$; they evolve in short time scale so that corresponding time constants range into milliseconds. However, their integral (averaged over time) values which form vector $\mathbf{C}_2$ evolve more slowly. These functions are the most informative for the process diagnostics, although the optimal averaging parameters must be considered individually for a particular EP process, and this task is very complex and at the state of the art [101].

For characteristics 5.3.2.1 and 5.3.2.2 (Fig. 10) – current and voltage are connected by Ohm’s law, and they cannot both together be inputs or outputs. If the voltage is an input, the current is an output, and vice versa. Here we consider the voltage as an input and the current as an output, which is typical for EPPs with the VGE. For PEO type processes, the opposite consideration can sometimes be beneficial but this does not alter the concept of the phenomenological model.

A particular composition of vectors $\mathbf{X}, \mathbf{S}$ and $\mathbf{C}$ is defined by measurement tools available to a researcher; however, the most crucial ones for investigation and modelling of the electrolytic plasma processes should be considered as follows:

- vector $\mathbf{X}$: average, effective (RMS) and instantaneous (waveform) values of voltage 5.3.2.2; average (over volume) electrolyte temperature 4.3.2.1;
- vector $\mathbf{S}$: phase composition 2.3.1.1 and 1.3.1.4; coating thickness 2.3.1.7; roughness 2.3.1.4; specific weight change 1.3.1.5; corrosion properties 2.3.1.3; fatigue strength 1.3.1.2;
- vector $\mathbf{C}$: average, effective (RMS) and instantaneous (waveform) values of current 5.3.2.1; working electrode temperature 1.3.2.1; type of VGE boiling 3.3.2.2.
Expansion of vector $S$ provides a deeper understanding of surface layer transformation as a result of EPP. Expansion of vector $C$ contributes to the EPP diagnostics and control possibilities. Any expansion of these vectors improves the understanding of the process mechanism.

3.3. Dynamic properties of EPPs as multiscale systems

Now consider changing rates of EPP characteristics and evaluate corresponding time constants as the first order measures of the system inertia. Evaluation of dynamic properties for various EPPs is shown in Table 1.

The most common data concerning EPP dynamics describe the surface properties comprising vector $S$. Typical examples are collated in Fig. 12 as time dependencies for EPPo [68], PEO [39] and EPCS [69] processes at optimal values of voltage ($U$) and electrolyte temperature ($T$). As seen from Fig. 12, PEO and EPCS processes, which are intended to produce a change of the modified layer thickness, run at approximately constant rate from 1 to 3 $\mu$m/min. The EPPo process which primary objective is to reduce surface roughness $Ra$ has much higher rate in the beginning of the treatment, and this can be described by an exponential fall [68]. A typical EPP duration reaches tens of minutes, with the longest PEO treatments lasting up to several hours [25, 73, 102]. Therefore, for vector $S$, the time constant value ranges into hundreds of seconds (Table 1); this corresponds to the process evolution in the long time scale and can be adequately described with a sampling period in the range of minutes.

Another type of the data represented in the voltage and current waveforms included into vector $C_1$ are instantaneous values of the measured variables. For a PEO treatment at 50 Hz, typical waveforms are shown in Fig. 13a [103]. For potentiostatic DC EPPo, a typical waveform of the current is depicted in Fig. 13b [104]. Spectral analysis of the EPPo waveforms provides an effective bandwidth characterising EPP: from 0 to 10 kHz (Fig. 14); this is consistent with the data reported elsewhere [4, 71, 105, 106]. Obtained from the power spectral density $P$, the time constant estimates for the short time scale in the EPP range from hundreds of
microseconds to hundreds of milliseconds (Table 1). Data acquisition with appropriate sampling rates of tens and hundreds of kHz generates a significant volume of often noisy data which needs statistical signal processing.

The third type of data comprises RMS voltage and current chronograms, i.e. averaged values of the instantaneous waveforms or instrument readings included into vector $C_2$. Fig. 15 (a, b) shows typical voltage and current chronograms for PEO processes with stabilised RMS current and voltage respectively [107, 108], whereas Fig 15(c) presents a typical current chronogram for an EPP with a VGE [6]. The curve shapes which are generally similar to those in Fig. 7 help in the evaluation of time constants for the characteristics in vector $C_2$ (Table 1). Corresponding values range into tens of seconds, which provides a compromise time scale to enable formal EPP description and modelling.

For better understanding, diagnostics and control of EPPs, new controllable and observable parameters with time constants close to those for the variables in vector $S$ should be found by appropriate integration techniques in electric, acoustic or optical characteristics and put into vector $C_2$. This is a new, important and challenging problem, solutions for which just started to appear [95, 97, 109, 110].

Thus, electrolytic plasma processes are multiscale systems described by three time constants which are separated by 2-3 orders of magnitude and correspond to different groups of characteristics in the phenomenological model. To obtain a finite difference dynamic model for a particular EPP, the discrete time step size $\Delta t$ should correspond to the time constants for vector $C_2$, ranging from seconds to tens of seconds. To put together the multiscale data from vectors $S$, $C_1$ and $C_2$, the data from vector $C_1$ must be averaged, and the data from vector $S$ must be interpolated. Correctness of the data representation for vector $S$ is ensured by the slow rate of surface layer evolution during the treatment. Correctness of the data representation for vector $C_1$ is ensured by the ergodicity interval for the instantaneous values being similar to the step size of the discrete compromise time scale [86].

3.4. Regression modelling and design of experiments
The black box concept used for the phenomenological modelling (Fig. 11) is tightly connected with experimental design methodology [111]. Various researchers use different designs, from full factorial to complex composite, including Taguchi analysis [112, 113]. Design of EPP experiments could be performed to address the following aims:

1) process mechanism investigation;
2) establishment of optimal treatment conditions;
3) development of nonlinear models of the process;
4) development of process diagnostics methods.

To develop verified EPP models including diagnostic information, detailed research into a wide range of factors is required. Due to process non-linearity, the application of full factorial designs with more than two factor levels is inevitable. Such designs, including at least voltage, electrolyte temperature and treatment duration as the factors, are quite laborious. Investigation of modern EPPs that employ pulsed bipolar current also requires the factors of the pulse shape and frequency, to be taken into account and this increases the design matrix dimension yet further. Comparison of studies based on different experimental designs, e.g. central composite [78], fractional factorial [86] and full factorial [39, 69], shows the following. The surface properties evolution analysis is seriously impeded if the experimental points are absent or shifted from the factor levels of the full factorial design. Mathematical models, even those based on neural networks, do not allow the restoration of sufficient information regarding the surface state. Overcoming these obstacles require additional experiments, so that their total number could eventually reach that in the full factorial design. Therefore, in the state space cross-section corresponding to the technological factors, a full-factorial design is highly recommended.

The situation with the factor of process duration is slightly different. When investigating significantly nonlinear EPPs, e.g. TiN coating stripping [6, 78], the treatment history of each sample is of high importance, because neither surface nor VGE state could be replicated when the process is resumed after termination.
individual experiments are required. However, for near stationary EPPs, the surface layer is persistent after the treatment termination, and the results of experiments with and without termination would be statistically similar. For processes, such as PEO, EPPo and EPCS, it is possible to obtain several experimental points by using one sample which is sequentially treated and measured after each run step. This approach significantly reduces the required number of samples but relies on a robust measurement technique.

According to the experimental design, measurements of the surface properties are required after each planned treatment duration. This requires process termination and manipulation with the samples, which can take significant time. Employing indirect methods of in-situ surface state evaluation helps in reducing the design dimension over the process duration factor.

The problem of an increased number of experiments in order to investigate the effects of frequency and other pulse parameters could be partly resolved using the frequency response (FR) approach which utilises the ergodicity time for a slow non-stationary process for performing measurements over frequency [101]. In other words, the frequency scans run in the ‘fast’ time for the system which properties gradually evolve in the ‘slow’ time as shown in Section 3.3. Therefore, the experimental design dimension can be collapsed over frequency so that the scans occur in every experiment, thus, contributing to saving samples and runs.

Finally, in EPP studies, it is advisable to apply full factorial designs, wherein, for every combination of factor levels corresponding to technological parameters, individual experiments are performed using different samples for strongly non-stationary EPPs; for near-to-stationary processes, this can be done using a single sample. Dependencies on the pulse parameters, e.g. frequency, can be investigated within every experimental run due to the time scale difference between these parameters and surface properties.

3.5. Frequency response approach to the process diagnostics
One of the main problems that slow down industrial EPP applications is a lack of diagnostic tools that provide possibilities for process monitoring and control, i.e. tools that could join vectors $S$ and $C$ together. The reason for the diagnostic tools not being commonly accepted and widely used is that the surface properties of the treated components are unavailable for measurements during the treatment, and, therefore, they are considered unobservable (but controllable) state variables of the system. As seen from the phenomenological model structure (Fig.10, block 3.3.2), several approaches to the process diagnostics are available, e.g. based on the analysis of electrical [104, 114, 115], optical [38, 91, 93, 94, 116] and acoustic [95] characteristics, video imaging [36, 37], and their various combinations [4, 96, 117]. Only a few however are developed enough to be employed in a real-time process control system, e.g. voltage-current characteristics [73, 110], power spectral density of the current [100], and characteristic line intensities of discharge optical emission spectra [109].

All of the above diagnostic approaches belong to the class of passive system identification. Recently, an active identification approach was developed on the basis of the frequency response measurements, both for large and small signal modes [39, 86]. It was shown that this method allows direct identification of the surface layer properties, e.g. barrier layer specific capacitance [83], and also indirect identification of non-electric properties, such as coating thickness and roughness [86]. The methodology of data acquisition and signal processing within this approach is presented in [101]; whereas the methodology of phenomenological modelling based on diagnostics information is discussed below.

The framework of EPP diagnostics and modelling using the frequency response approach (Fig. 16) runs through blocks 1 to 6 around the core of plasma fundamentals 7 [118]. Within the framework, the EP process 1 comprises the subsystems shown in the phenomenological model structure (Fig. 9). These subsystems can be assessed through the frequency sweep technique 2 which can be operated in the small signal or large signal modes. This imposes certain limitations on the waveform types, sweep modes and data acquisition parameters. Supported and extreme situations (block 3)
formalise the bounds within which the FR approach can be applied to verify evidence of processing conditions and surface properties in electrical characteristics. The frequency response analysis and simulation includes application of spectral methods, including electrochemical impedance spectroscopy (EIS) approaches to the equivalent circuit structural and parametric identification resulting in forming vector \( C \) in the model. Besides conventional resistive and capacitive components from EIS, negative differential resistance and distributed elements may appear in the equivalent circuits to fit complex shaped spectra. This step is tightly connected with the surface analysis and characterisation. Information obtained through the coating thickness, morphology, functional properties evaluation, as well as phase and chemical composition assessment forms vector \( S \) in the model. Uncovering regularities connecting blocks 4 and 5 using correlation and other types of statistical analyses formalises the ways to connect the process characteristics and the surface state. Therefore, block 6 – process diagnostics and control using in-situ and ex-situ methods can be realised, leading the development of the electrolytic plasma processes to a new horizon.

Finally, this framework offers a path for further research into EPP fundamentals and provides a powerful tool for process diagnostics and control, thus contributing towards smart electrolytic plasma technologies.

3.6. Electromagnetic field modelling

EPP implementation is partially hampered by a lack of mathematical models allowing the estimation of the treatment uniformity. The VGE formed around the working electrode is usually considered relatively thin (\( \leq 10^{-3} \) m), providing the highest electrical resistance in the circuit. Therefore, the voltage drop in the electrolyte is often neglected, and the current density at the electrolyte – VGE interface is averaged and considered constant [6, 121]. While the former is fair for the majority of EPPs with a VGE, the latter holds only when using the simplest electrode shapes and layouts, e.g. coaxial cylindrical cells. A very limited number of EPP related publications discuss primary and/or secondary current density distributions in
the electrolyte and the VGE respectively [73, 122]. This is a growing area for research [123].

Electromagnetic field modelling is well-established for the analysis of electrochemical processes. It is widely used for simulation and structural optimisation of electrolyzers with liquid metals, especially for Al reduction [124], cathodic protection systems [125], fuel cells and batteries [126], anodic coatings [127] and other applications. The nonlinearity of EPPs discussed earlier could be a reason why it has not so far been explored for this group of processes. Solution of field problems for EPPs would contribute to the phenomenological modelling, e.g. by determining the net electrolyte resistance as a boundary condition for the frequency response analysis or specifying the boundaries within the phenomenological approach that treats the process as a system with lumped characteristics.

3.7. Perspective on phenomenological modelling and research of electrolytic plasma processes

The advancement of electrolytic plasma technologies relies strongly on availability of adaptive process control systems (Fig. 1 (b,c)) tailored to particular processes. Development of essential elements of such systems constitutes further research needs in this field. These include development of:

- dynamic reference models for individual processes;
- new diagnostic tools linking observable process parameters and features with key surface characteristics and properties;
- process identification models based on the diagnostic tools and methods;
- robust feedback algorithms and appropriate hardware for their realisation.

The modelling aspect holds the key to further progression in that direction and the proposed phenomenological approach could significantly contribute to this. Developments in EPP modelling and associated research may proceed along the following possible paths. First of all, most researchers model surface properties using regression analysis which results in static models. The proposed approach would also help in the development of dynamic models essential for process control.
The other important research direction is development of electrical, optical and acoustical methods for process diagnostics and incorporation of corresponding characteristics of into EPP models. This includes frequency response interpretations [128, 129], statistical spectral analysis of electrochemical noise in the voltage and current [130], microdischarge image analysis [131], spectral analyses of optical [93, 116] and acoustic emission [96]. These analyses should lead to identification of informative characteristics and establishment of their correlations with surface properties.

Electromagnetic field problems are starting to attract attention. This includes primary current density distribution in the electrolyser and secondary distribution in the vapour gaseous envelope and the modified layer. These problems are challenging due to the multiscale character of the object and non-linearity of the boundary conditions in the vicinity of the treated surface. This should facilitate electrolyser design, including optimisation of electrode layouts and electrolyte management systems for treatment of components with complex geometries.

Yet another important set of modelling problems whose solution could contribute to the EPP development includes electric processes in power supplies. Dynamic models of electrical process characteristics, i.e. voltage-current diagram and frequency response, should enable design of specialised power supplies which could be stable under such non-linear loads as VGEs and plasma discharges featuring NDR. This could contribute to the design and manufacturing specialised EPP equipment and scaling up various processes.

The next important route for development is in combining with other electrochemical processes in terms of the techniques used for the process diagnostics and modelling. Quantitatively it is only voltage that separates conventional electrochemical processes from EPPs. Therefore, the approaches successfully applied in electrochemistry could in principle be adopted to study EPPs should the equipment capabilities permit. An example of such bridging is in-situ impedance spectroscopy proposed by the authors in [86] and later used elsewhere [129].
Ultimately, using this approach to bridge between plasma physics, materials science, electrochemistry, electrical engineering, computational mathematics and computer science would contribute to the development of smart electrolytic plasma technologies capable of addressing key challenges of modern manufacturing.

4. Examples of EPP phenomenological modelling

4.1. Regression neural network modelling

Let us consider an example of phenomenological modelling of EPCS of diffusion aluminide coating from nickel superalloy. Experimental details and resulting surface properties are described elsewhere [31, 69].

A full-factorial experimental design (Table 2) with a total of 36 physical experiments was implemented. Nine runs were made at all combinations of the voltage and electrolyte temperature with the levels selected (Fig. 17a). According to the model structures shown in Fig. 11, a static regression model was developed in Matlab using a general regression neural network (GRNN) [99]. The network structure and its Matlab representation are shown in Fig. 18a,c. It has 54 neurons in the hidden layer and 2 neurons in the output layer. This network was trained to transform the input vector of variables $X=\begin{bmatrix} U; T \end{bmatrix}$ and $t$ into the output vector $S=\begin{bmatrix} h; Ra \end{bmatrix}$, with coefficient of determination $R^2=0.99$, using 54 learning examples derived from the physical experiments. The modelling results shown in Fig. 19 indicate that the solid lines of the model curves stay within the confidence intervals designated by the dashed lines; therefore, the model is adequate. The model is clearly non-linear and the application of conventional regression equations could not succeed in creating an adequate model; however, this is not the case for the neural networks.

For the dynamic model, a nonlinear autoregressive network with exogenous inputs (NARX) was used. This is a recurrent dynamic network, with feedback connections enclosing several layers of the network [132]. Its structure is shown in Fig. 18b, where $Z^{-1}$ block in the feedback loop represents one step time delay. This
network has vector $\mathbf{X}$ in the input, and vector $\mathbf{S}$ in the output. Time $t$ is included indirectly as the time step $\Delta t = 1$ min. The Matlab representation of the model is depicted in Fig. 18d, which shows that current values of input vector $\mathbf{X}$ are passed with zero delays into the hidden layer. The output values of vector $\mathbf{S}$ are also fed to the input of the hidden layer with one-step delay. This means that the network calculates its output from current values of voltage and electrolyte temperature, and from the surface properties estimated in the previous time instant. This makes the model dynamic. After optimisation, the network has 20 neurons in the hidden layer and 2 neurons in the output layer.

To train the dynamic neural network model, a numerical experiment with the design shown in Fig. 17b was performed. The model resolution was increased using the numerical experiment containing 289 runs with $\Delta t = 1$ min, which is an 8-fold finer resolution than the physical experiment. This produced a training set with 6069 learning examples. This NARX network was trained using the Levenberg-Marquardt algorithm with performance validation in order to avoid overtraining [99]. The resulting dynamic neural network model has the coefficient of determination $R^2=0.99$, providing the same modelling results as the static model for constant input values as shown in Fig. 19 and in Fig. 20b,c, line 1.

The difference between the static and dynamic model appears for the inputs that change in time. The reaction of the models to a voltage step from 300 to 400 V at $t=5$ min is shown in Fig. 20. The static model shows a step of sudden coating growth but this would be physically impossible during the coating stripping process. The dynamic model does not show any step; only the rate of the coating stripping changes according to the process mechanism [69]. Therefore, the dynamic EPP model derived by the phenomenological approach is capable of estimating the surface properties during the treatment under arbitrary variations of inputs (within the limits imposed during the model construction), thus contributing to understanding of EPP mechanisms, process diagnostics and control and ultimately promoting development of smart electrolytic plasma technologies.
4.2. Frequency response modelling

The frequency response process diagnostics and modelling provide useful tools for the revealing hidden regularities underlying EP phenomena and in-situ estimation of surface properties. Examples of the FR analysis for different substrate materials (Al and Ti), and for different surface macro morphology (dense and porous Al samples) are provided in Figs. 21(a-c).

The coatings were obtained on disk samples with nominal area \( A = 15 \text{ cm}^2 \) in a conventional alkaline electrolyte with specific conductivity \( 8 \text{ mS} \cdot \text{cm}^{-1} \) in pulsed bipolar mode with the frequency sweep from 20 Hz to 20 kHz at the following voltages: \( U_{\text{DC}} = 225 \text{ V}; U_{\text{AC}} = 305 \text{ V} \). The resultant coating thickness on dense samples was \( h = 20 \mu \text{m} \). Other experimental details can be found elsewhere [86, 133]. A typical surface morphology is shown in the insets in Fig. 21.

Nyquist and Bode plots in Fig. 21 demonstrate that the impedance of the PEO process in all the cases exhibits resistive-capacitive behaviour (the phase angle is negative). The Nyquist plots show elliptical arcs which differ from conventional semicircle plots for RC circuits [134]. Also, the plots have negative slope at lower frequencies, which can be explained by microdischarge plasma impedance featuring NDR.

A comparison of impedance spectra for PEO-Al and -Ti reveals a noticeable difference below 2 kHz, with higher absolute values of the phase angle for PEO-Ti making the semicircle narrower. This can be explained by fitting the spectra to a simple model circuit shown in the inset to Fig. 21. The fitting results collated in Table 3 indicate that whilst \( R_p \) values are similar, the capacitance \( C \) for the two materials interfaces is different, being significantly higher for Ti than Al. According to the estimate of capacitance as \( C = \varepsilon_r \varepsilon_0 A/h \), it can be influenced by the relative permittivity \( \varepsilon_r \) of corresponding oxides as indicated in Table 3. This opens up a possibility for in-situ monitoring of phase transitions during formation of PEO coatings, if the new phase has significantly different dielectric properties. Such an approach can also provide guidance on how to adjust current pulse parameters in order to control the reactive component of the current with changing impedance of
the system. Moreover this could be useful to achieve a better understanding of frequency effects that are commonly observed and studied in AC and pulsed current EPPs [56, 76, 135-141], yielding partial and often controversial inferences.

A comparison of the impedance spectra obtained for dense and foam aluminium substrates shows that both $R_p$ and $C$ are higher for the foam. Taking into account that the actual surface for the foam samples is much higher than the nominal area, this ratio can be assessed as a ratio of the capacitances: $A_{\text{dense}}/A_{\text{foam}} = C_{\text{dense}}/C_{\text{foam}} \approx 1/5$. However, the ratio of $R_p$ for the dense and foam substrates is only $\approx 1/2$, indicating that certain diffusion limitations may occur for the foam inner regions, contributing to the process non-linear scaling behaviour.

Thus the application of FR methodology for EPP diagnostics can reveal the characteristics of surface state in-situ, provided that the correlations between impedance spectra and surface properties are sufficiently well understood. This can be facilitated by coupling electrical FR with other observable characteristics, such as optical and acoustic emission.

4.3. Electric field modelling

Knowledge of electric field distribution in the EPP electrolyser is important for assessment of the treatment uniformity. Despite the fact that almost all voltage drops across either the VGE or the oxide layer, the voltage drop over the electrolyte should not be neglected for EPP of complex shape components. Uneven current density in the electrolyser induced by tank size and shape as well as size, shape and position of the electrodes leads to non-uniform current density distribution at across the component surface, resulting in the differential thickness of the modified layer.

An example of the PEO treatment of a component with deep holes is considered in Fig. 22. It could be envisaged that shielding by the component walls would reduce the current density on the inner surfaces. Solving boundary problem of the current density distribution would help in evaluating the deviation of the coating thickness.
The vector of current density $\mathbf{j}$ in the electrolyte is described by a Kirchhoff’s law [142]:

$$\text{div}\mathbf{j} = 0$$

(6)

Taking into account Ohm’s law

$$\mathbf{j} = \sigma \mathbf{E},$$

(7)

where $\mathbf{E}$ is electric field and $\sigma=4.2$ mS cm$^{-1}$ is the electrolyte conductivity, and the relation with electric potential $\phi$

$$\mathbf{E} = -\text{grad}\phi,$$

(8)

the boundary problem can be solved numerically using Laplace equation

$$\text{div} \text{grad} \phi = 0,$$

(9)

employing field modelling software utilising the finite elements method, e.g. COMSOL.

For the setup shown in Fig. 22b, the boundary conditions are as follows

$$\phi=0 \text{ for the counter-electrode,}$$

(10)

$$\phi=500 \text{ V for the sample,}$$

(11)

$$E_n=0 \text{ for the conductive media boundaries,}$$

(12)

where index $n$ denotes normal projection of the electric field vector.

Analysis of the electric field and current density distribution in the electrolyte shown in Fig. 22a, by colour filling and contour lines respectively, reveals particularly high current densities at the edges. This edge effect blocks penetration of electric field in the holes which results in a sharp drop of local current densities at about 3 mm from the mouth of the hole.

As $j_n$ does not change on the sample/electrolyte interface, the local charge density decreases with the hole depth, so as the coating thickness (Fig 22b). The reduction is particularly noticeable during the first 3 mm of the hole depth, which is consistent with the field modelling results in Fig. 22a, whereas on the outer surface, the thickness distribution is relatively even. Thus, the distribution of current density in the electrolyte can explain variations of surface characteristics within the
workpiece and provide estimates of boundary conditions for FR analysis; it should therefore be taken into account during phenomenological modelling.

5. Conclusions

From the review of electrolytic plasma processes and approaches to their modelling the following conclusions can be drawn:

1. Two major EPP types featuring VGE and oxide film formation can be distinguished, with more detailed classification made concerning electrical and technological features of the processes.

2. Models using partial differential equations, existing theories of processes in the surface oxide layer as well as empirical, phenomenological and equivalent circuit approaches have been developed for EPPs. The phenomenological models appear to be the most appropriate for the development of prospective smart electrolytic plasma technologies.

3. A generalised phenomenological model structure has been proposed. The model describes the system ‘power supply – electrolyser – treated surface’ as a system with lumped parameters. This model forms a basis for a deeper EP process understanding and provides a methodology for their modelling, diagnostics and control.

4. As the generic complexity of EPPs does not currently allow the set of differential equations describing the treatments to be solved, the phenomenological model of a particular EPP could be formalised as a black box regression. Within this approach, input and output variables have been analysed and categorised as controllable supply inputs, unobservable surface properties and observable electric, optical and acoustic characteristics. A general algorithm assisting the design of static and dynamic process models for a given EPP has been proposed to formalise the phenomenological model.

5. Analysis of dynamic properties demonstrated that EPPs are multiscale systems which can be described by three time constants separated by 2-3 orders of
magnitude (minutes, seconds and milliseconds) for the unobservable surface properties, integral observable characteristics and instantaneous observable characteristics. It was justified that the discrete model rates should be converted to the range of tens of seconds, for the slower variables by interpolation, and for the faster variables by integration.

6. Since the phenomenological models rely heavily on experimental data, regression modelling and experimental design recommendations have been formulated to optimise the number of experiments required for a particular model.

7. The frequency response approach to diagnostics of unobservable surface properties has been proposed. The approach includes spectral analysis of electrical characteristics, resulting, for example, in impedance spectra of the EP process; the evolution of the spectra is highly correlated with the surface properties; therefore, a framework for the frequency response assessment within the phenomenological approach has been proposed.

8. Electromagnetic field modelling complements the phenomenological approach by providing boundary conditions for the FR estimates in the form of electrolyte resistance, representing the electrolyser impedance at infinite frequency. Moreover, it helps in assessing the treatment non-uniformity for complex shape components.

9. Further research needs concerning phenomenological modelling of EPPs have been discussed. Bridging between plasma physics, materials science, electrochemistry, electrical engineering, computational mathematics and computer science using the phenomenological approach could facilitate development of smart electrolytic plasma technologies.

10. Examples of EPP phenomenological modelling have been provided for modelling surface properties and their in-situ diagnostics, and for the electric field distribution in the electrolyte and at the sample surface. The examples show the efficiency of the proposed phenomenological approach.
6. Acknowledgements

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### Table 1. Evaluation of dynamic properties for various EPPs

<table>
<thead>
<tr>
<th>Process</th>
<th>Variable</th>
<th>Variable change rate</th>
<th>Time constant (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vector S</strong>&lt;br&gt;Long time scale&lt;br&gt;EPHT [3]</td>
<td>Hardened layer thickness $h$</td>
<td>5…50 µm min$^{-1}$</td>
<td>100…500</td>
</tr>
<tr>
<td>PEO [1]</td>
<td>Coating thickness $h$</td>
<td>1…4 µm min$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>EPPo [68]</td>
<td>Surface roughness $Ra$</td>
<td>0.05…0.2 µm min$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>EPCS [31]</td>
<td>Coating thickness $h$</td>
<td>1…4 µm min$^{-1}$</td>
<td></td>
</tr>
<tr>
<td><strong>Vector C_{1}</strong>&lt;br&gt;Short time scale&lt;br&gt;EPHT [3]</td>
<td>Instant current $i$</td>
<td>0.1…2 A ms$^{-1}$</td>
<td>10$^{-1}$ … 10$^{-4}$</td>
</tr>
<tr>
<td>PEO [4]</td>
<td></td>
<td>0.1…5 A ms$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>EPPo [104]</td>
<td></td>
<td>0.2…1 A ms$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>EPCS [105]</td>
<td></td>
<td>0.1…2 A ms$^{-1}$</td>
<td></td>
</tr>
<tr>
<td><strong>Vector C_{2}</strong>&lt;br&gt;Compromise time scale&lt;br&gt;EPHT [7]</td>
<td>Working electrode temperature $T_c$</td>
<td>30…50 °C s$^{-1}$</td>
<td>10…100</td>
</tr>
<tr>
<td>PEO [87]</td>
<td>Effective (RMS) voltage $U$</td>
<td>2…10 V s$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>PECS [6]</td>
<td>Effective (RMS) current $I$</td>
<td>20…200 mA/s</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Full factorial experimental design

<table>
<thead>
<tr>
<th>Factor</th>
<th>Min. value</th>
<th>Max. value</th>
<th>Step</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voltage U (V)</td>
<td>300</td>
<td>400</td>
<td>50</td>
</tr>
<tr>
<td>Electrolyte temperature T (°C)</td>
<td>50</td>
<td>90</td>
<td>20</td>
</tr>
<tr>
<td>Process duration t (min)</td>
<td>0</td>
<td>20</td>
<td>5</td>
</tr>
</tbody>
</table>

Table 3. FR modelling results for different substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Re (Ω)</th>
<th>Rp (Ω·cm²)</th>
<th>C (F·cm⁻²)</th>
<th>εᵣ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al dense</td>
<td>10</td>
<td>75.2</td>
<td>2.7·10⁻⁷</td>
<td>8…11</td>
</tr>
<tr>
<td>Al foam</td>
<td></td>
<td>132.6</td>
<td>1.3·10⁻⁶</td>
<td></td>
</tr>
<tr>
<td>Ti dense</td>
<td></td>
<td>74.2</td>
<td>1.2·10⁻⁶</td>
<td>80…173</td>
</tr>
</tbody>
</table>
Fig. 1. Feedback concepts using models: (a) Classical control system with observable output $Y$; (b) Control system with reference model. The output $S$ is unobservable and the reference model must be dynamic; (c) Control system with diagnostic loop. Output $S$ is unobservable and output $C$ is observable.
Fig. 2. Stages of vapour gaseous envelope evolution: a – oxygen evolution; b – bubble boiling in the VGE; c – microdischarges in the VGE; d – microdischarge treatment of the surface micro profile; 1- working electrode; 2 – electrolyte; 3 – oxygen (a) and vapour (b) bubbles; 4 – microdischarges [34, 35]
Fig. 3. Side view pictures of aluminium alloy samples at different time of the PEO process: a – few seconds; b – 15 min.; c – 30 min.; d – 45 min. [38]
Fig. 4. Classification of electrolytic plasma processes
Fig. 5. Cross-section of a steel sample after case hardening (a) [40]; micrographs of the top view of PEO coating on Al (40 μm thick) (b) [62] and Ti (13 μm thick) (c) [63] alloys; micrograph of the surface morphology of a stainless steel after polishing (Ra = 0.15 μm) (d) [13]; micrograph showing typical morphology of a EP cleaned steel (e) [8]; cross-section of the aluminide coating being stripped from a nickel superalloy (f) [31]
Fig. 6. Typical arrangements of the equipment used for EP processes in electrolyser (a) [1] and with electrode tool (b) [7]: 1 – window; 2 – mixer; 3 – connecting wires; 4 – exhaust/ventilation system; 5 – grounded case; 6 – power supply unit; 7 – workpiece; 8 – cooling system; 9 – electrolyser-counter electrode; 10 – insulating plate
Fig. 7. Typical current curve for running EP processes from a voltage source (a) and typical voltage curve for running PEO processes from a current source (b):
1-2 – anodisation; 2-3 – sparking; 3-4 – plasma electrolytic oxidation
Fig. 8. Classification of models of EP processes

Models of electrolytic plasma processes

VGE models using partial differential equations
- With respect of electrolyte flow
  - Using Maxwell’s and heat transfer equations
  - Using boiling characteristic

Phenomenological models
- Model-classification of heating mode
- Relationship among processes during plasma electrolysis
- Evolution of microdischarges in oxide pores
- Evolution of microdischarges and crystallisation of oxides

Theoretical models of processes in oxide layer
- Metal-dielectric-electrolyte system model
- Impedance spectroscopy analysis
- Resistive model circuit
- Resistive and capacitive model circuit
- Nonlinear circuit

Equivalent circuit models
- Approximation of voltage transients
- Dynamic current-voltage characteristics
- Pulsed current-voltage characteristics
- Coating thickness, roughness vs. process parameters

Empirical models of operational characteristics

Electrolytic plasma polishing
- Electrolytic plasma heating
- Plasma electrolytic oxidation
Fig. 9. Generalised phenomenological model structure
Fig. 10. Generalised phenomenological model structure (continued)
Fig. 11. Black box EPP model structure (a) and internal structure of the EPP model (b)
Fig. 12. Dynamics of the major surface properties during different EPPs:
EPPo – U=300 V, T=80 °C; PEO – U=550 V, T=20 °C; EPCS – U=350 V, T=70 °C
Fig. 13. Waveforms of voltage and current density during PEO process at 50 Hz (a) [103] and waveform of the AC component of the current during EPPo at 10\textsuperscript{th} second of the treatment (b) [104]
Fig. 14. Statistical spectra of the AC component of the current during EPP of cylindrical (a-c) and square (d-f) samples at different treatment times [104]: a, d – 25 s; b, e – 105 s; c, f – 345 s
Fig. 15. Typical chronograms of RMS current density and voltage for AC PEO with RMS current density stabilisation (a) [107]; chronograms of RMS current density for PEO process in different electrolytes with RMS voltage stabilisation (b) [108] and chronograms of current for EPCS process run with voltage stabilisation with different initial temperatures (c) [6]
Fig. 16. Framework for EPP diagnostics and modelling using frequency response approach
Fig. 17. U-T plane of the experimental design: a – for physical experiment; b – for numerical experiment
Fig. 18. Structures of neural network models (a, b) and their Matlab representations (c, d): 
a, c – static; b, d – dynamic
Fig. 19. Modelling results for the static neural network model:
a, c, e – coating thickness; b, d, f – surface roughness;
a, b – U = 300 V; c, d – U=350 V; e, f – U = 400 V;
1 – T = 50 °C; 2 – T = 70 °C; 3 – T = 90 °C
Fig. 20. Results of modelling a response to a voltage step from 300 to 400 V (a) at T=50 °C with the static (b) and dynamic (c) models of PECS process: 1 – no step; 2 – with the step
Fig. 21. Nyquist (a), (c) and Bode (b), (d) plots of in-situ impedance spectra of PEO-Al and -Ti (a), (b), and for dense and foam Al substrates (c), (d). The insets show typical coating morphology for PEO coating on Ti, equivalent circuit used for the model fitting and typical macro morphology of the coated samples.
Fig. 22. Electric field distribution in the electrolyser (a) for the PEO treatment of Al component having inner holes (b) and coating thickness distribution across the inner and outer surfaces (c)
References


[34] Electrolyte plasma polishing process description, Steklovac, Belarus, 2004.


