Modeling polycrystalline electrode-electrolyte interfaces:
The differential capacitance

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Abstract

We present and analyze a model for polycrystalline electrode surfaces based on an improved continuum model that takes finite ion size and solvation into account. The numerical simulation of finite size facet patterns allows to study two limiting cases: While for facet size diameter $d_{\text{facet}} \to 0$ we get the typical capacitance of a spatially homogeneous but possible amorphous or liquid surface, in the limit $L_{\text{Debye}} \ll d_{\text{facet}}$, an ensemble of non-interacting single crystal surfaces is approached. Already for moderate size of the facet diameters, the capacitance is remarkably well approximated by the classical approach of adding the single crystal capacities of the contributing facets weighted by their respective surface fraction. As a consequence, the potential of zero charge is not necessarily attained at a local minimum of capacitance, but might be located at a local capacitance maximum instead. Moreover, the results show that surface roughness can be accurately taken into account by multiplication of the ideally flat polycrystalline surface capacitance with a single factor. In particular, we find that the influence of the actual geometry of the facet pattern in negligible and our theory opens the way to a stochastic description of complex real polycrystal surfaces.

1 Introduction

The theoretical and experimental investigation of polycrystalline electrode surfaces provides valuable information for the understanding of the performance for many types of real electrochemical devices. Fundamental concepts like the structure of the electrochemical double layer, double layer capacitance, Faradaic reaction rate equations and others are derived and verified experimentally for single crystal or liquid metal electrodes. In either case, the physicochemical properties of these electrodes surfaces can be idealized as entirely homogeneous. In contrast, real solid metal electrodes are most often polycrystalline and therefore they expose different crystal facets to the electrolyte, leading to the inhomogeneity of their physicochemical properties.

In this paper we propose a mathematically sound approach which allows to derive characteristics of polycrystalline electrodes from a thermodynamically well founded model. We focus on double layer capacitance and potential of zero charge as equilibrium properties. In particular, we aim at the characterization of an ideal polycrystalline surface which is characterized by a large ratio between facet size characterized by the diameter $d_{\text{facet}} \to 0$ and the Debye length $L_{\text{Debye}}$ of the electrolyte. In this regime, the influence of edge effects related to facet boundaries can be neglected such that compact asymptotic equations defining potential of zero charge and potential dependent double layer capacitance can be derived. Numerical simulations of the spatial charge distribution in the boundary layers confirm that contributions from facet boundary effects indeed are marginal. Ultimately, we replace discrete facet configurations by probability distributions of facets with identical physical properties. This allows a very compact reformulation of the polycrystalline electrode properties in terms of convolution integrals.
Figure 1: Left: Computed boundary layer capacity of a single crystal surface for different concentrations of a non-adsorbing electrolyte. Right: Measured capacitance (Fig. 3.a of [VP86], reprinted the permission of Elsevier)

Surfaces of polycrystalline electrodes in general exhibit a complex unstructured and random pattern of facets with different crystallographic orientation as sketched in Fig. 2 left. It is well known that the electron work function depends on the crystallographic orientation of a surface [HS79, TD95]. As a consequence, the differential capacity and the corresponding potential of zero charge (PZC) of the surface in contact to an electrolyte depend on the crystallographic orientation as well, cf. [Val81, Val82, Val89].

The development of the classical Gouy-Chapman-Stern-Grahame theory of the electrochemical double layer [Gra47] strongly relied on experimental measurements at mercury electrodes. Since these electrodes consist of a liquid metal, their clean surface in contact with an electrolyte typically can be considered as ideally smooth and physical homogeneous. Thus, when trying to transfer the theory from liquid to solid metal electrodes, first an awareness about the relevance of surface roughness and individual properties of surfaces facets of different crystallographic orientation needed to be established, cf. [Fru67]. Valette and Harmelin [VH73] first related careful capacity measurements of silver single crystal surfaces with the corresponding experimental results for polycrystalline silver electrodes. Although not explicitly written in [VH73], a relation of the form (1.1) for the capacity of the polycrystalline surface was assumed. In [BUL80, Vor81], equation (1.1) is derived and analyzed on the basis of equivalent circuits derived from the Gouy-Chapman-Stern-Grahame model.

In this work, we apply a recently derived model framework on the basis of continuum non-equilibrium thermo-electrodynamics to describe a polycrystalline electrode-electrolyte interface cf. [DGM13, LGD16]. This model is capable to predict qualitatively and quantitatively the differential capacity of single crystal electrodes with respect to the applied voltage and with respect to the salt concentration (see Fig. 1). For a single crystal electrode in contact with a non-adsorbing electrolyte, the capacity is exclusively due to the charging of the double layer. As long as the bulk electrolyte concentration is not too large, it shows the typical symmetric camel shape capacitance curve, see Fig. 3 (left), where the potential of zero charge (PZC) is located at the local capacity minimum and does not change with salt concentration. The absolute value of the PZC depends on the crystallographic orientation of the metal surface, and differences between two PZC values are known to correspond well to the respective differences of the electron work function [TD95, LGD16].

For a polycrystalline electrode surface, we derive expressions of the double layer charge, the potential of zero charge, and the differential capacitance and compare these to the (theoretically) more simple and well understood case of single crystal surfaces. The model incorporates the pressure in the electrolyte as a variable and the momentum balance, which ensures thermodynamic consistency. Further it allows to account for volume exclusion effects of all species, and especially of the solvation effect. For a review
of the development of generalized Nernst-Planck models we refer to [KBA07]. Within some reasonable scaling it is a remarkable and non-obvious feature of the resulting model that double layer charge, and subsequently the potential of zero charge, as well as the differential capacity can be determined from solutions of an algebraic equation system, rather than solving space dependent differential equations. This even holds true for a stochastic description of a polycrystalline electrode surface, which we show within this work.

The inhomogeneity of the surface capacitance is commonly believed to cause so called ‘frequency dispersion’ in electrochemical impedance spectroscopy, cf. [Paj94], and the effects of surface inhomogeneity can be expected to have even more serious impact in the presence of Faradayic reactions. We plan to investigate these non-equilibrium aspects in follow-up research.

**General approach.** We model the polycrystalline electrode surface by a regular surface pattern with \( N > 1 \) different types of facets. Each facet corresponds to certain crystallographic orientation. To each facet \( \Sigma_i, 1 \leq i \leq N \), we assign the surface fraction \( s_i = \frac{| \Sigma_i |}{| \Sigma |} \) with \( \sum_{i=1}^{N} s_i = 1 \), where \( | \cdot | \) denotes the surface area, and a value of the electron work function. We show, based on our thermodynamic model of Section 2, that the electric potential \( \varphi^s \) on each facet \( \Sigma_i \) is related to the work function, or in our notation, to the chemical potential \( \mu^s \) of surface electrons of the respective surface orientation.

The polycrystalline electrode \( \Sigma = \bigcup_i \Sigma_i \) is then considered to be in contact to some electrolytic solution \( \Omega^E \), which is in thermodynamic equilibrium described by an improved Poisson-Boltzmann equation with facet-wise constant boundary values for the electric potential. The adjacent space charge layer in the electrolyte can be computed numerically. Figure 2 (right, bottom) shows a 3D numerical computation of the electric potential iso-surfaces in the electrolyte \( \Omega^E \) for a periodical checkerboard electrode surface. Based on such numerical solution we can compute, (i) the overall double layer charge \( Q_{\text{poly}} \) as function of the applied voltage \( E \), (ii) the potential of zero charge \( E_{\text{PZC}} \) of a polycrystalline electrode-electrode interface, and (iii) the corresponding double layer capacity \( C_{\text{poly}} \) (see Figure 2 (right, top)). This approach is valid for arbitrary geometries and facet sizes The only assumption here is that the intersecting lines between two facets do not have independent thermodynamic quantities.

For a non-adsorbing electrolyte\(^1\) we have

\[
\begin{align}
Q_{\text{poly}} &= \frac{1}{| \Sigma |} \int_{\Omega^E} q(x) \, dx = Q_{\text{poly}}^\text{BL}(E) \\
E_{\text{PZC}} &= E \quad \text{s.t.} \quad Q_{\text{poly}}^\text{BL}(E) = 0 \\
C_{\text{poly}} &= \frac{dQ_{\text{poly}}^\text{BL}(E)}{dE} =: C_{\text{poly}}^\text{BL}(E)
\end{align}
\]  

(1.1)

A scale analysis of the boundary value problem yields two characteristic length scales of the model, the typical diameter \( d_{\text{facet}} \) of the facets \( \Sigma_i \) and the Debye length \( L_{\text{Debye}} \) of the electrolyte.

If the facet or facet diameter \( d_{\text{facet}} \) is large compared to the Debye length \( L_{\text{Debye}} \), the double layer charge \( Q_{\text{poly}} \) can be obtained as a sum of the single facet boundary layer charges \( Q_{\text{BL}} \), weighted by their respective surface fraction \( s_i \). Subsequently, the potential of zero charge \( E_{\text{PZC},\infty} \) can be determined from an algebraic equation system, and the differential capacity \( C_{\text{poly}} \) obeys also an

\(^1\)Note that specific adsorption is also discussed within this work.
Figure 2: Sketch of the facet structure of a polycrystalline surface (left) and FEM computation of the electric potential in the electrolyte in contact with a checkerboard polycrystalline surface (right) with corresponding double layer charge $Q_{\text{poly}}$ and capacity $C_{\text{poly}}$ as function of the applied voltage $E$.

Figure 3: Double layer charge and differential capacitance of a polycrystalline electrode determined from (1.2) with $s_1 = s_2 = \frac{1}{2}$.

algebraic representation, i.e.

\[
\begin{align*}
\text{(M1)} \quad \frac{d}{dt} \frac{\Delta E_{\text{poly}}}{E_{\text{Debye}}} &\to \infty \\
Q_{\text{poly}} &= \sum_{i=1}^{N} s_i \cdot \hat{Q}_{\text{poly}}^{BL} \left( E + \frac{1}{\epsilon_0} \mu_e^i - E_{\text{ref}} \right) =: Q_{\text{poly,}\infty}^{BL}(E) \\
L_{\text{poly,}\infty}^{PZC} &= E \text{ s.t. } Q_{\text{poly,}\infty}^{BL}(E) = 0 \\
C_{\text{poly}} &= \sum_{i=1}^{N} s_i \cdot \hat{C}_{\text{poly}}^{BL} \left( E + \frac{1}{\epsilon_0} \mu_e^i - E_{\text{ref}} \right) =: C_{\text{poly,}\infty}^{BL}(E),
\end{align*}
\]

where $\hat{Q}_{\text{poly}}^{BL}$ and $\hat{C}_{\text{poly}}^{BL}$ are nonlinear functions specified in section 2. The construction of the capacity $C_{\text{poly,}\infty}^{BL}$ of a polycrystalline electrode in the case of facets with two equal surface fractions is illustrated in Fig. 3 (right). Remarkably, the potential of zero charge $E_{\text{poly,}\infty}^{PZC}$ of this polycrystalline surface in this case is located at the capacity maximum and not at a local minimum of the curve!

For realistic polycrystalline electrodes, the consecutive labeling of all facets might be impractical. We propose thus a stochastic description of the surface in terms of a probability density $f$, modeling the surface fraction as function of the surface chemical potential of electrons. We derive in the limit
Figure 4: Left: densities of the surface chemical potential $\mu_s$ given by normal distributions with different standard deviation. Right: Capacity curves for the respective densities (solid —) and for the planar single crystal case (dotted line · · ·). For large $\sigma$, the the capacity has a single maximum unlike the camel shape for perturbations of the single crystal surface by small $\sigma$.

In the opposite limiting case, for a vanishing facet diameter $d_{\text{facet}} \to 0$, the potential of zero charge is given as the surface fraction weighted average of the corresponding single crystal quantities, while the charge $Q_{\text{poly}}$ and the capacity $C_{\text{poly}}$ remain non-linear expressions identical to the ones of a typical single crystal surface, i.e.

\begin{equation}
\begin{aligned}
Q_{\text{poly}} &= (f \ast Q^{BL}(E)) =: Q^{BL}_{\text{poly,0}}(E) \\
E^{PZC}_{\text{poly,0}}(E) &= E^{ref} \quad \text{s.t.} \quad Q^{BL}_{\text{poly,0}}(E) \neq 0
\end{aligned}
\end{equation}

\begin{equation}
\begin{aligned}
C_{\text{poly}} &= (f \ast C^{BL}(E)) =: C^{BL}_{\text{poly,0}}(E) \\
E^{PZC}_{\text{poly,0}}(E) &= E^{ref} - \frac{1}{e_0} \sum_{i=1}^{N} s_i M_i \Sigma_i =: E^{PZC}_{\text{poly,0}}(E)
\end{aligned}
\end{equation}

Outline. In the next Section 2, we first introduce the general equilibrium conditions of the applied continuum model and appropriate material models. Then, the differential capacity of a single crystal surface is introduced and characterized based on the model and a model for polycrystalline surfaces is developed. Section 3 is devoted to an assessment of the results obtained by numerical solution of the boundary value problem in 2D and in 3D for periodically patterned surfaces of finite size facets. In particular, the limiting behaviour for extreme facet size parameters and the influence of surface roughness on the length scale of the facets are analyzed. Section 4 contains a discussion of the
polycrystal model in the case of large facet size, including the effects of adsorption and the stochastic description of polycrystalline surfaces. We close with some concluding remarks in Section 5.

2 Continuum model

We use a continuum model that is based on a coupled volume-surface description of charged and possibly reacting mixtures coupled with the electromagnetic field, see [DGM13, LGD16, DGM18]. In this model, as a convention, each quantity defined in the volume has a corresponding quantity confined to the electrode surface $\Sigma$. We use the same letters for bulk and surface quantities and indicate the surface quantities by an underset 's'. For a surface $\Sigma$ we choose the orientation of the normal vector $\nu$ that we denote by $\nu$. Then, we denote the adjacent bulk volumes by $\Omega^E$ and $\Omega^M$, such that $\nu$ is the outer normal of $\Omega^E$. For a generic function $u$ in the volume, we can define the trace and jump at $\Sigma$ by

$$u^{| \Sigma}_K = \lim_{x \in \Omega^K \to \Sigma} u(x), \quad K = E, M \quad \text{ and } \quad [u]_\Sigma = u^{| E}_\Sigma - u^{| M}_\Sigma. \quad (2.1)$$

Bulk positions far away from the surface $\Sigma$ are denoted by $x^K$ and the evaluation of some function $u(x)$ the bulk points is denoted by $u^K = u(x^K), K = E, M$.

We restrict our considerations to the isothermal case, where the temperature $T$ enters the equations only as a constant parameter and on the surface there holds $T_s = T^{| E}_\Sigma = T^{| M}_\Sigma$. Moreover, we only consider the electrostatic case where the electric field is given as $E = -\nabla \varphi$. Excluding surface dipoles, the electric potential $\varphi$ is continuous at the surface, i.e. we can set $\varphi_s = \varphi^{| E}_\Sigma = \varphi^{| M}_\Sigma$.

Description of mixtures. To refer to the constituents of a mixture on the surface $\Sigma$ or the adjacent volume domains $\Omega^K$, we use according index sets $I_\Sigma$, resp. $I_K$ of species. For each constituent, we denote the molecular mass by $m_\alpha$, the partial molar volume by $\upsilon_{\text{ref}}^\alpha$, the charge number by $z_\alpha$, and the number density of particles per m$^3$ by $n_\alpha$, where $\alpha \in I_K$ or $\alpha \in I_\Sigma$, respectively. Then, we introduce the partial mass density $\rho_\alpha = m_\alpha n_\alpha$. The free charge density is ($K = M, E$)

$$q = \sum_{\alpha \in I_K} z_\alpha n_\alpha, \quad q_s = \sum_{\alpha \in I_\Sigma} z_\alpha n^s_\alpha. \quad (2.2)$$

2.1 Model equations

The continuum model relies on balance equations in bulk and surface that are universal, i.e. independent of the considered material. However, the equations contain material dependent constitutive quantities which need appropriate modeling. By application of an entropy principle, cf. [DGM18], we guarantee non-negative entropy production. The key ingredients for the material modeling are the free energy densities $\rho^\psi, \rho^s_\psi$ in the bulk and on the volume, respectively. Within this work, we assume a simple material model with a constant dielectric susceptibility $\chi$ and thus consider free energy densities of the still rather general structure

$$\rho^\psi = \rho^\psi(T, (n_\alpha)_{\alpha \in I_K}) - \frac{1}{2} \chi \varepsilon_0 |\nabla \varphi|^2, \quad \rho^s_\psi = \rho^s_\psi(T_s, (n^s_\alpha)_{\alpha \in I_\Sigma}). \quad (2.3)$$

$^2$The notation indicates that the definition can be made for both domains $\Omega^E$ and $\Omega^M$ in analogous manner. If $u$ is not defined in either $\Omega^E$ or $\Omega^M$, the respective trace value is set to 0.

$^3$Where not necessary, we omit the labels $E$ or $M$ for the volume domains.
The chemical potentials are defined as
\[ \mu_\alpha = \frac{\partial \rho \hat{\psi}}{\partial n_\alpha}, \quad \mu_s = \frac{\partial \rho s \hat{\psi}}{\partial n_s}. \] (2.4)

By means of Gibbs-Duhem relations, we introduce the material pressure \( p \) and surface tension \( \gamma_s \) as
\[ p = -\rho \hat{\psi} + \sum_{\alpha \in \mathcal{I}_E} n_\alpha \mu_\alpha, \quad \gamma_s = \rho_s \hat{\psi} - \sum_{\alpha \in \mathcal{I}_S} n_s \mu_s. \] (2.5)

In the following, we neglect gravitation and restrict the presentation to planar interfaces with no tangential transport on the surface.

**Equilibrium bulk equations.** The equilibrium of an incompressible mixture in a bulk domain \( \Omega \) is characterized by
\[-(1 + \chi) \varepsilon_0 \Delta \varphi = q, \quad \nabla \cdot \mathbf{p} = -q \nabla \varphi, \quad 0 = \nabla \left( \mu_\alpha + z_\alpha \varepsilon_0 \varphi \right) \quad \text{for } \alpha \in \mathcal{I}_K. \] (2.6a, 2.6b, 2.6c)

The Poisson equation (2.6a) determines the electric potential, (2.6b) is the momentum equation, (2.6c) states that the electrochemical potentials of all species are constant in equilibrium. We remark that these equations are not independent, but (2.6a) and (2.6c) together with (2.5)(left) already imply (2.6b).

**Equilibrium surface equations.** On the surface \( \Sigma \) between the bulk domains \( \Omega^\pm \) the equilibrium is given by
\[-\left[ (1 + \chi) \varepsilon_0 \nabla \varphi \right] = q_s, \quad \left[ (p \mathbf{1} + (1 + \chi) \varepsilon_0 (\nabla \varphi \otimes \nabla \varphi - \frac{1}{2} |\nabla \varphi|^2 \mathbf{1})) \mathbf{\nu} \right] = 0, \quad \mu_s = \mu_\alpha |_{\mathcal{S}_K} \quad \text{for } \alpha \in \mathcal{I}_K, K = E, M. \] (2.7a, 2.7b, 2.7c)

### 2.2 Specific material model

**Bulk electrolyte material model.** The electrolyte in the domain \( \Omega^E \) is a mixture of several species, one of them being the solvent that we refer to by the index \( \alpha = 0 \). The index set of the electrolytic species is denoted by \( \mathcal{I}_E \). The mole fractions \( y_\alpha \) of the constituents are defined as
\[ y_\alpha = \frac{n_\alpha}{n}, \quad \text{with} \quad n = \sum_{\beta \in \mathcal{I}_E} n_\beta. \] (2.8)

In many solvents, ions are solvated, meaning that a number \( \kappa_\alpha \) of solvent molecules are bound into a solvation shell around a center ion. Thus, the partial volume \( v_{\alpha}^{\text{ref}} \) of a solvated ion in the electrolyte is typically much larger than the partial volume of the solvent \( v_0^{\text{ref}} \). We consider the electrolyte as an incompressible liquid mixture of free solvent molecules, undissociated species and solvated ions.\footnote{This is in contrast to the model of a mixture of undissociated species, the bare center ions and solvent molecules, regardless whether they are free or bound into a solvation shell.}
incompressibility of the mixture is characterized by the constraint
\[ 1 = \sum_{\alpha \in I_E} u^E_{\alpha} n_{\alpha}. \tag{2.9} \]

The free energy for an ideal mixture of solvated ions is derived and analyzed in [DGM13, LGD16, DGLM18]. The chemical potentials in the incompressible limit are
\[ \mu_{\alpha} = g^{\text{ref}}_{\alpha} + v^{\text{ref}}_{\alpha} (p - p^E) + k_B T \ln(y_{\alpha}), \quad \alpha \in I_E, \tag{2.10} \]

Let \( g^{\text{ref}}_{\alpha}, \varphi^E \) and \( p^E \) denote the values of the mole fractions, the electric potential and the pressure at some bulk point \( x^E \) in \( \Omega^E \). In equilibrium, the constant electrochemical potentials according to (2.6c) and the material model (2.10) then imply for the bulk mole fractions
\[ \text{equilibrium: } y_{\alpha} = y^E_{\alpha} \cdot \exp \left( - \frac{z_{\alpha} e_0}{k_B T} (\varphi - \varphi^E) - \frac{v^{\text{ref}}_{\alpha}}{k_B T} (p - p^E) \right) \quad \text{in } \Omega^E. \tag{2.11} \]

Note that the pressure \( p \) in the electrolytic boundary layer can be expressed as a function of the potential difference \( \varphi - \varphi^E \) by solving the condition (2.8) with the representations (2.11), i.e. the non-linear implicit relation
\[ \sum_{\alpha \in I_E} y^E_{\alpha} \cdot \exp \left( - \frac{z_{\alpha} e_0}{k_B T} (\varphi - \varphi^E) - \frac{v^{\text{ref}}_{\alpha}}{k_B T} (p - p^E) \right) - 1 = 0. \tag{2.12} \]

Bulk metal material model. We consider a metal in the domain \( \Omega^M \) as a binary mixture of positive metal ions \( M \) and free electrons \( e \) with \( z_{\alpha} = -1 \), i.e. \( I_M = \{ M, e \} \). For the metal bulk free energy density, we adopt the Sommerfeld model [Som28] and assume incompressibility, cf. [LGD16]. We do not specify the model here, because as a consequence of the following surface material model, it turns out that the metal bulk does not influence any of the results in this paper and therefore can be ignored.

Surface material model. The surface \( \Sigma \) between the domains \( \Omega^E \) and \( \Omega^M \) is considered as mixture of the surface metal ions, surface electrons, electrolytic adsorbates and possibly reaction products of the aforementioned species. The index set of surface constituents is denoted by \( I_\Sigma \) with \( I_E \cup I_M \subseteq I_\Sigma \). Note that we consider on the surface also a solvation effect, whereby each adsorbed ion binds \( \kappa_{\alpha} \) solvent molecules which propagates into the partial molar area \( a^{\text{ref}}_{\alpha s} \).

We adopt the surface free energy model proposed in [LGD16]. In particular, the surface chemical potential of the electrons is assumed to be a constant value depending only on the material and the crystallographic surface orientation. We propose to choose this value related to the electron work function \( W_\Sigma \) of the surface \( \Sigma \). Analogously to the metal volume, we have an incompressibility constraint on the surface stating \( a^{\text{ref}}_{\alpha s} n_M = 1 \), where \( a^{\text{ref}}_{\alpha s} \) is the partial area of surface metal ions. On the electrolyte side, we have to account for adsorption from the volume. Since the surface is not necessarily completely covered with adsorbates, we introduce a number density of surface vacancies via
\[ n_s = n_M - \sum_{\alpha \in I_\Sigma \setminus I_M} a^{\text{ref}}_{\alpha s} n_{\alpha}. \tag{2.13} \]
Then, we define the surface fractions of vacancies and adsorbates by

$$n_s = n_s V + \sum_{\alpha \in \mathcal{I}_S \setminus \mathcal{I}_M} n_s \alpha,$$

$$y_V = \frac{n_V}{n_s} \quad \text{and} \quad y_\alpha = \frac{n_\alpha}{n_s} \quad \text{for} \ \alpha \in \mathcal{I}_S \setminus \mathcal{I}_M. \quad (2.14)$$

The chemical potentials of the surface species are

$$\mu_s = \psi_s^{\text{ref}} + k_B T \ln \left( y_\alpha \right) - \frac{a_\alpha \gamma s}{k_B T},$$

$$\mu_M = \psi_M^{\text{ref}} + k_B T \ln \left( y_V \right) - \alpha \rho M \gamma s,$$

$$\mu_e = \mu_e^{\text{ref}} \quad (= \text{const.}). \quad (2.15)$$

With respect to the same bulk point in the electrolyte as above, i.e. the values $y_E^{\alpha}$, $\varphi E$ and $p E$, the surface mole fraction of the vacancies, the electrolytic adsorbates, and the surface reaction products, have the representations in terms of the potential difference $U = \varphi |_{\Sigma} - \varphi \infty$,

$$y_V = \exp \left( - \frac{a_V^{\text{ref}}}{k_B T} \gamma s \right), \quad (2.16a)$$

$$y_\alpha = y_E^{\alpha} \exp \left( - \frac{\Delta g_\alpha}{k_B T} - \frac{e_0 z_\alpha U}{k_B T} - \frac{a_\alpha \gamma s}{k_B T} \right), \quad \text{for} \ \alpha \in \mathcal{I}_E, \quad (2.16b)$$

$$y_\beta = \prod_{\alpha \in \mathcal{I}_E} (y_E^{\alpha})^\nu_{\alpha \beta} \exp \left( - \frac{\Delta g_\beta}{k_B T} - \frac{e_0 z_\beta \sum_{\alpha \in \mathcal{I}_E} \nu_{\alpha \beta} z_\alpha U}{k_B T} - \frac{a_\beta \gamma s}{k_B T} \right), \quad \text{for} \ \beta \in \mathcal{I}_S \setminus \mathcal{I}_E \setminus \mathcal{I}_M, \quad (2.16c)$$

where the amount of adsorbates and reaction products on the surface is controlled by the corresponding Gibbs energies defined by

$$\Delta g_\alpha = \psi_\alpha^{\text{ref}} - (\psi_\alpha^{E} + \psi_\alpha^{\text{ref}} \rho E) \quad \text{for} \ \alpha \in \mathcal{I}_E, \quad (2.17a)$$

$$\Delta g_\beta = \psi_\beta^{\text{ref}} - \sum_{\alpha \in \mathcal{I}_M} \nu_{\alpha \beta} (\mu_\alpha^{M} + e_0 z_\alpha U^{M}) - \sum_{\alpha \in \mathcal{I}_M} \nu_{\alpha \beta} (\psi_\alpha^{E} + \psi_\alpha^{\text{ref}} \rho E) \quad \text{for} \ \alpha \in \mathcal{I}_E. \quad (2.17b)$$

The above representations and the definition (2.14) of the surface mole fractions yield an algebraic equation which determines the surface tension $\gamma_s$ as a function of $U$.

### 2.3 Planar single crystal surface

It is well known that at the surface $\Sigma$ between a metal and an electrolyte a charged double layer exists, which exponentially decays into the electrolyte bulk and has a width in the order of the Debye length

$$L_{\text{Debye}} = \sqrt{\frac{(1 + \chi) \varepsilon_0 k_B T}{e_0^2 n^{\text{ref}}}}. \quad (2.18)$$

For an electrolyte with a salt concentration of $0.1 \text{mol/L}$, the Debye length is in the order of one nm. If the dimension of the considered experimental setup is considerably larger than $L_{\text{Debye}}$, say in the range of $1 \mu\text{m}$, it often is reasonable to consider the surface $\Sigma$ as an infinite planar surface and the electrolyte domain $\Omega^E$ as a half space. In this simple geometric setting, all quantities can only depend
on the spatial coordinate normal to \( \Sigma \), which we denote in the following by \( x \). For a flat surface \( \Sigma \), we can write the electrolyte up to the bulk point \( x^E \) as \( \Omega^E = \Sigma \times [0, x^E] \). Then, we define the boundary layer- and the surface charge density by

\[
Q_{BL} = \int_0^{x^E} q \, dz \quad \quad Q_s = -e_0 \sum_{\alpha \in I_{\Sigma} \setminus I_M} z_\alpha n_\alpha . \tag{2.19}
\]

In equilibrium, both quantities are completely determined by the potential difference \( U = \varphi_s - \varphi^E \).

It is a remarkable feature of the applied material model in Sect. 2.2 that it is possible to determine the boundary layer charge density \( Q_{BL} \) as function of \( U \) without needing to spatially resolve the bulk equations (2.6)–(2.7), i.e.

\[
Q_{BL} = \text{sgn}(U) \sqrt{2\varepsilon_0(1 + \chi)}(\hat{\rho}(U) - p^E) =: \hat{Q}_{BL}(U) . \tag{2.20}
\]

where \( \hat{\rho}(U) \) is the material pressure determined from (2.12). Moreover, all surface mole fractions are determined according to (2.16) and we get for the surface charge density

\[
\hat{Q}(U) = -\sum_{(\alpha \in I_{\Sigma} \setminus I_M)} z_\alpha \epsilon_0 y_\alpha + \sum_{(\alpha \in I_{\Sigma} \setminus I_M)} \nu_{\alpha\beta} z_\alpha \epsilon_0 y_\beta \tag{2.21}
\]

Thus, we can introduce the differential capacity as

\[
\hat{C}(U) = \frac{d}{dU} \left( Q_{BL}(U) + Q_s(U) \right) = C_{BL}(U) + C_s(U) \tag{2.22}
\]

Under the standard assumption that the potential difference between an ideally non-polarizable reference electrode and the bulk electrolyte remains constant under potential variations, it is possible to show \cite{LGDT16} that

\[
U = \varphi_s - \varphi^E = E + \frac{1}{\epsilon_0} \mu_e - E^{\text{ref}} \quad \text{with} \quad E^{\text{ref}} = \text{const} . \tag{2.23}
\]

where \( E \) is the applied voltage in a three electrode setup, \( \mu_e \) the surface chemical potential of electrons, and \( E^{\text{ref}} \) a constant which depends on the actual reference electrode. For a specific surface \( \Sigma \), the differential capacity in the absolute scale \( E \) is given by an appropriate shift of (2.22), i.e.

\[
C(E) = \hat{C} \left( E + \frac{1}{\epsilon_0} \mu_e - E^{\text{ref}} \right) . \tag{2.24}
\]

2.4 Modeling of planar polycrystal surfaces

We consider now a planar surface \( \Sigma \) which is composed of \( N \) facets \( \Sigma_i \), i.e. \( \Sigma = \bigcup_{i=1}^N \Sigma_i \) as sketched in Fig. 2. We denote by \( |\Sigma_i| \) the area of the facet \( \Sigma_i \) whereby the total area of \( \Sigma \) is \( |\Sigma| = \sum_{i=1}^N |\Sigma_i| \) and surface fraction \( s_i := \frac{|\Sigma_i|}{|\Sigma|} \). We make the assumption

(A1) The facet boundaries \( \partial \Sigma_i \cap \partial \Sigma_j \) for \( i \neq j \) constitute no independent thermodynamic entities.

\(^5\)The typeface \( \hat{Q} \) indicates that this is a function \( U = \varphi_s - \varphi^E \) and not the applied voltage \( E \).
Thus, each facet $\Sigma_i$ can be treated as a finite size planar single crystal surface, and we can apply the material model of Sect. 2.2 on each facet $\Sigma_i$ separately. Since the surface density of metal ions depends on the crystallographic orientation of the surface, the material parameter $a_M$ is different for each individual facet $\Sigma_i$. As a consequence, in the absence of adsorption to the surface, the surface tension $\gamma$ differs between the facets $\Sigma_i$. In addition, we have to assign to each facet $\Sigma_i$ a constant value of $\mu_s$, which has to be related to the electron work function on a surface of the respective crystallographic orientation. Because all facets are connected by electric conductors, the electrochemical potential of the electrons is constant over all grains $\Omega_i$ adjacent to the respective facets $\Sigma_i$, i.e.

$$\left.\left(\mu_e - e_0\varphi\right)\right|_{\Sigma_i} = \left.\left(\mu_e - e_0\varphi\right)\right|_{\Omega_i} = \left.\left(\mu_e - e_0\varphi\right)\right|_{\Sigma_j}. \tag{2.25}$$

Between each two surfaces, we thus have the pairwise potential difference

$$\varphi|_{\Sigma_j} - \varphi|_{\Sigma_i} = \frac{1}{\varepsilon_0} \left(\mu_e|_{\Sigma_j} - \mu_e|_{\Sigma_i}\right). \tag{2.26}$$

**Patterned planar surface.** As in the single crystal case before, we can expect boundary layers that decay exponentially from the surface $\Sigma$ into the electrolyte bulk. The length scale of the boundary layer is given by the Debye length according to (2.18) and the electrolyte bulk can be characterized by certain constant far field values $y^E_\alpha$, $\varphi^E$ and $\rho^E$ in a distance from the surface that is large compared to $L^\text{Debye}$. We assume that the electrolyte in $\Omega^E$ is not influenced by any other boundary layer than the one at $\Sigma$. Given the applied potential $E - E^{\text{ref}}$ and a pattern of facets $\Sigma_i$ with according constant values of $\mu_e$, we choose some value $\tilde{\mu}_e$ and set $\varphi^E = -(E - E^{\text{ref}}) - \frac{1}{\varepsilon_0} \tilde{\mu}_e$. Then, the state in the electrolyte in $\Omega^E$ is given by the boundary value problem

$$-(1 + \chi)\varepsilon_0 \Delta \varphi = q(\varphi, \rho) \quad \text{in } \Omega^E, \tag{2.27a}$$

$$1 = \sum_{\alpha \in I_E} y^E_\alpha(\varphi, \rho) \quad \text{in } \Omega^E, \tag{2.27b}$$

$$\varphi|_{\Sigma_i} = \frac{1}{\varepsilon_0} \left(\mu_e|_{\Sigma_i} - \tilde{\mu}_e\right) \quad \text{on } \Sigma, \tag{2.27c}$$

$$\nabla \varphi \rightarrow 0 \quad \text{far away from } \Sigma, \tag{2.27d}$$

where $y^E_\alpha(\varphi, \rho)$ is given by (2.11) for $\alpha \in I_E$ and $q$ determined from $y^E_\alpha$ according to (2.2) and (2.9). The surface charge density can be determined by (2.21) on each facet $\Sigma_i$ separately and we get

$$Q^E|_{\Sigma_i} = \hat{Q}^E s \left(E - E^{\text{ref}} - \frac{1}{\varepsilon_0} \tilde{\mu}_e|_{\Sigma_i}\right). \tag{2.28}$$

Thus for the surface $\Sigma$, we conclude

$$Q^\text{poly}(E) = \sum_{i=1}^N s_i \hat{Q}^E \left(E - E^{\text{ref}} - \frac{1}{\varepsilon_0} \tilde{\mu}_e|_{\Sigma_i}\right). \tag{2.29}$$

---

6One has to be careful with an interpretation of this fact. Surface tension of planar solid surfaces is difficult to measure and one has to distinguish between the 'thermodynamic' surface tension and the 'interfacial tension', cf. [DGLM18].

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Upon solution of the boundary value problem, the boundary layer charge density is computed from

\[ Q_{\text{BL}}^{\text{poly}}(E) = -\frac{1}{|\Sigma|} \int_{\Omega^e} q(x) \, dx , \]  

Finally, the total surface charge is

\[ Q_{\text{poly}}(E) = Q_{\text{BL}}^{\text{poly}}(E) + Q_s^{\text{poly}}(E) \]

and the differential capacity of the patterned surface is computed by taking the derivative with respect to the applied potential \( E \).

**Ensemble of planar single crystal surfaces.** The total boundary layer charge density can be decomposed into the contributions adjacent to the different facets \( \Sigma_i \), viz.

\[ Q_{\Sigma_i}^{\text{BL}} = \sum_{i=1}^{N} s_i Q_{\Sigma_i}^{\text{BL}} \quad \text{with} \quad Q_{\Sigma_i}^{\text{BL}} = -\frac{1}{|\Sigma_i|} \int_{\Omega^e(\Sigma_i \times R)} q(z) \, dz . \]  

Since boundary effects propagate into the interior with a length scale according to the Debye length \( (2.18) \), boundary effects due to the finite size of \( \Sigma_i \) can be neglected once the characteristic facet diameter is sufficiently large, i.e. \( L_{\text{Debye}} \ll d_{\text{facet}} \). Then, the charge density in front of \( \Sigma_i \) is well approximated by the boundary layer charge \( (2.20) \) at a planar single crystal with the same material parameters as \( \Sigma_i \), i.e.

\[ \hat{Q}_{\Sigma_i}^{\text{BL}}(E - E_{\text{ref}} + \frac{1}{e_0} \mu_e) \to 0 \quad \text{for} \quad \frac{L_{\text{Debye}}}{d_{\text{facet}}} \to 0 . \]  

Thus, in the limit of large facets, the boundary layer charge density \( Q_{\text{poly}}^{\text{BL}} \) of the polycrystal is given by the according density of an ensemble of non-interacting (infinitely large) planar single crystal surfaces, all in contact with the same electrolyte, viz.

\[ Q_{\text{poly}}^{\text{BL}}(E) = \sum_{i=1}^{N} s_i \hat{Q}_{\Sigma_i}^{\text{BL}}(E - E_{\text{ref}} + \frac{1}{e_0} \mu_e) , \]  

The boundary layer capacity related to \( (2.33) \) is given by \( (1.1) \). As \( (2.28) \) can be applied independent of the size of \( d_{\text{facet}} \), we conclude

\[ C_{\text{poly}}(E) = \sum_{i=1}^{N} s_i \hat{C}(E - E_{\text{ref}} + \frac{1}{e_0} \mu_e) . \]  

### 3 Surfaces of finite size pattern and approached limit relations

In order to keep the focus of this Section sharp, we assume here that the electrolyte is non-adsorbing. If in addition also adsorption has to be take in to account, surface charge and surface capacity can be computed in purely algebraic manner by a post-processing step applying \( (2.29) \) and its derivative with respect to the applied potential, respectively.

#### 3.1 Pattern of two equally sized facets

We consider first the most simple configuration of a symmetric bi-crystalline surface, where \( N = 2 \), \( s_1 = s_2 = 1/2 \). We choose an average potential \( \bar{\mu}_e \) such that \( \mu_e|_{\Sigma_1} - \bar{\mu}_e = - (\mu_e|_{\Sigma_2} - \bar{\mu}_e) = -0.1 \text{eV} \).
Table 1: Physical constants and parameters used in the numerical simulations

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dielectric constant ($\varepsilon_0$)</td>
<td>$8.85418781762 \times 10^{-12} \text{C}^2/(\text{V} \cdot \text{m})$</td>
</tr>
<tr>
<td>Boltzmann constant ($k_B$)</td>
<td>$1.3806488 \times 10^{-23} \text{J}/\text{K}$</td>
</tr>
<tr>
<td>Elementary charge ($\varepsilon_0$)</td>
<td>$1.602176565 \times 10^{-19} \text{C}$</td>
</tr>
<tr>
<td>Avogadro number ($N_A$)</td>
<td>$6.02214129 \times 10^{23} \text{mol}^{-1}$</td>
</tr>
<tr>
<td>Temperature ($T$)</td>
<td>$298.15 \text{K}$</td>
</tr>
<tr>
<td>Reference pressure ($p_{\text{ref}}$)</td>
<td>$1 \times 10^5 \text{Pa}$</td>
</tr>
<tr>
<td>Dielectric susceptibility ($\chi$)</td>
<td>$15$</td>
</tr>
<tr>
<td>Solvation number ($\kappa_\alpha$)</td>
<td>$10$</td>
</tr>
<tr>
<td>Reference number density ($n_{\text{ref}}$)</td>
<td>$15.5 \text{mol}/\text{L}$</td>
</tr>
<tr>
<td>Specific volumes ($\nu_{\alpha}$)</td>
<td>$(1 + \kappa_\alpha) \nu_{\text{ref}}$</td>
</tr>
</tbody>
</table>

Figure 5: Left: profile of the electric potential for different diameters of the surface stripes. Right: capacity curves for different electrolyte concentrations, where solid lines (—) refer to the algebraic solution according to (1.1), FEM solution for facet size $d_{\text{facet}} = 3.07 \text{ nm}$ is indicated by markers (+), and for $d_{\text{facet}} = 12.28 \text{ nm}$ by dashed lines (- - ).

On the surface, we thus have the boundary values $\varphi_{|_{\Sigma_i}} = \Phi_i$ for $i = 1, 2$ with $\Phi_1 = -\Phi_2 = -0.1 \text{V}$. The simplest realization of this configuration consists of a pattern of parallel stripes with alternating prescribed potential. Here, we let $d_{\text{facet}}$ denote the width of the stripes. With an appropriate choice of the coordinate system, the boundary values can be described by a 1D-function such that

$$
\varphi(x) = \begin{cases} 
\Phi_1 & \text{for } 2|x| < d_{\text{facet}}, \\
\Phi_2 & \text{for } 2|x - d_{\text{facet}}| < d_{\text{facet}},
\end{cases}
$$

and periodic continuation. Accordingly, the potential and the charge density in the electrolyte domain $z > 0$ can be determined from a 2D-FEM computation by integration of the free charge $q$ in space according to (2.30). The choice of the boundary values implies that in the far field the potential approaches $\varphi \to \varphi^E = -\left(E - E_{\text{ref}}^f\right) - \frac{1}{\varepsilon_0} \bar{\mu}_s$. Plots of the electric potential for $E = E_{\text{ref}}^f - \frac{1}{\varepsilon_0} \bar{\mu}_s$ and an electrolyte with a bulk concentration of $0.1 \text{ mol/L}$ and the remaining parameters according to Tab. 1 are given in Fig. 5 (left). In the upper plot, where the facet size was chosen as $d_{\text{facet}} = 10L_{\text{Debye}} \approx 12.28 \text{ nm}$, one can observe that the piece-wise constant boundary data to a large extent propagates into the electrolyte and decays with increasing distance to the boundary. To the contrary, in the lower plot, where $d_{\text{facet}} = 5L_{\text{Debye}} \approx 3.07 \text{ nm}$, the profile of the potential is dominated by facet boundary effects such that the regions of parallel iso-lines almost disappear.

Next, the applied potential is varied in positive and in negative direction and the boundary value problem
From this solution, the free charge density $q$ and $Q_{\text{BC}}(E)$ are computed. Fig. 5 (right) shows the resulting capacity for different electrolyte concentrations in dependence of the facet size $d_{\text{facet}}$. We observe for increasing facet size $d_{\text{facet}}$ a convergence of the computed capacity curves to the algebraic solution (1.1). As expected, the convergence is slower for more dilute electrolytes where the Debye length is comparably larger. We conclude that the relevant quantity for the convergence is the facet size relative to the Debye length $L_{\text{Debye}}$. Thus, for the algebraic result (1.1) to be a good approximation, a polycrystalline surface has to be more coarse faceted when used in connection with a dilute solution, than in the case of a more concentrated solution. Another observation is, that in the potential range outside of the lowest and the largest position of the maxima, the algebraic expression is always a good characterization of the polycrystalline surface, independent of the facet size.

In addition, 3D-FEM simulations with a checkerboard pattern of the surface were performed where the boundary values are given by periodic continuation of

$$
\varphi_s(x, y) = \begin{cases} 
\Phi_1 & \text{for } \max(|x|, |y|) < d_{\text{facet}}, \quad \text{sgn}(xy) > 0, \\
\Phi_2 & \text{for } \max(|x|, |y|) < d_{\text{facet}}, \quad \text{sgn}(xy) < 0.
\end{cases}
$$

The observations from the 3D computations are analogous as for the 2D case. Since for given $d_{\text{facet}}$, the length of the contact lines in the checkerboard pattern is double compared to the striped pattern represented by the 2D case, convergence for increasing facet size is a bit slower here. Together, the 2D and the 3D results indicate that the shape of the facets is not relevant as long as the facet diameter $d_{\text{facet}}$ is sufficiently large compared to the Debye length $L_{\text{Debye}}$. We conclude that already the covered surface fraction of the facets and the corresponding single crystal capacities fully determine the capacity of a polycrystalline surface of sufficiently large facet size.

As FEM simulations in 3D are computationally expensive, we used anisotropic grids with a mesh grading with respect to the a-priori known structure of the potential, see Fig. 6. To check that the grid is sufficiently fine, we compared the resulting capacity curves from numerical solutions of the boundary value problem (2.27) on different refinement levels of the initial mesh. Refined mesheson the refinement levels 3, 4 and 5 consisted of 2 601, 18 513 and 139 425 nodes, respectively. Fig. 6 shows the convergence of the computed capacity curves when refining the mesh.
Figure 7: Left: capacity curves for 0.1 mol/L electrolyte and different facet size parameters (dashed lines - - ). Limits for large facet diameter (blue, solid —) and vanishing facet diameter (red, solid —). Right: lines show the position of the PZC of the patterned surface in dependence of the facet size $d_{\text{facet}}$ for different bulk concentrations. The circles on the to indicate the values of $E^{\text{PZCpoly}}$ according to (1.1) that are approached for $d_{\text{facet}} \to \infty$ and the dashed line refers to $E^{\text{PZCpoly}}$,0 according to (1.4).

3.2 Characterization of PZC and limiting behaviour

In the previous section, we did not discuss the PZC for the patterned polycrystalline surface. The reason is that due to the symmetry, i.e. $s_1 = s_2$, it always coincides with the chosen reference potential. In order to introduce some non-symmetry, we slightly modify the configuration to $N = 2$, $s_1 = 1/3$, $s_2 = 2/3$, and again set $\Phi_1 = -\Phi_2 = -0.1$V. For a 2D-FEM simulation of this configuration, we choose

$$
\varphi_s(x) = \begin{cases} 
\Phi_1 & \text{for } 2|x| < d_{\text{facet}}, \\
\Phi_2 & \text{for } |x - \frac{3}{2} d_{\text{facet}}| < d_{\text{facet}},
\end{cases}
$$

(3.3)

with periodic continuation of the boundary data. As in the previous example, we again observe that the capacity of the patterned surface approaches (1.1) for large facet sizes $d_{\text{facet}}$, see Fig. 7 (left). From the computed boundary charge as a function of the applied potential according to (2.30), we can now determine the potential of zero charge of the patterned surface with finite $d_{\text{facet}}$. We observe that for a fixed facet size $d_{\text{facet}}$, the PZC is not constant but increases with the bulk electrolyte solution, cf. Fig. 7 (right). Moreover, the potential range, in which the PZC varies with respect to the bulk concentration, gets wider for increasing $d_{\text{facet}}$. As a limit for $d_{\text{facet}} \to \infty$, the PZC reaches the values we get from the surface fraction weighted averaging of single crystals according to (1.1). Since the capacity of a planar single crystal is approximately quadratic at its PZC, the boundary layer charge behaves almost cubic. Thus, one should not expect, that the linear combination of single crystal capacities would attain its PZC at the respective linear combination of the individual PZCs.

In addition, another limit is of interest, however more from a theoretical point of view. Let us consider $d_{\text{facet}} \to 0$, although a pattern at the atomic length scale or even blow that scale can not be realized in practice. We observe, that for $d_{\text{facet}} \to 0$, the capacity curve changes from the three-maxima shape according to (1.1) into the typical two-maxima or camel shape that is well known from the planar single crystal. In fact, for $d_{\text{facet}} \to 0$, the capacity converges to the capacity of a planar single crystal surface where the PZC is determined as the sum of the PZCs on the contributing facets weighted by their respective surface fraction $s_j$ according to (1.4). Moreover, this limit of the PZC is independent of the bulk electrolyte concentration.
Figure 8: Electric potential for an electrolyte of 0.1 mol/L salt concentration in contact with a symmetric bi-crystal surface at the applied potential $E = E_{\text{ref}} - \frac{1}{e_0} \bar{\mu} s_e$. Top: flat surface, i.e. $h_{\text{max}} = 0$, bottom: differently oriented facets with $h_{\text{max}} = 1.54 \text{nm}$.

### 3.3 Surface roughness

Because of surface roughness, the actual surface area of real surfaces is larger compared to their ideally plain counterparts which are characterized by their so called visible surface area. As a measure of this effect, the surface roughness factor $W$ is introduced as the quotient of the real surface area over the visible. Surface roughness appears on a large range of different length scales, from `physical inhomogeneity' on the atomic length scale to 'geometric roughness' on a macroscopic scale of several $\mu$m. A lot of attention was devoted to the analysis of `geometric' surface roughness which in experiments may be caused e.g. by electrode pre-treatment like polishing. In the context of electrochemical impedance analysis, roughness is then often analyzed in the framework of fractal geometry, where on the larger scales, an interaction of the boundary layer with bulk transport properties of electrolyte has to be expected.

We want to concentrate on the effect of surface roughness on the boundary layer capacitance and therefore consider only deformations of an ideally plain surface that is on a length scale of the facet size $d_{\text{facet}}$. We return to the setting of the symmetric bi-crystal of Sect. 3.1 and let the surface be given by

$$
\frac{d_{\text{facet}}}{h_{\text{max}}} h(x) = \begin{cases} 
    x & \text{for } 2|x| < d_{\text{facet}}, \\
    (d_{\text{facet}} - x) & \text{for } 2|x - d_{\text{facet}}| < d_{\text{facet}},
\end{cases}
$$

where the parameter $h_{\text{max}}$ is the maximal elevation of the surface over the supporting pane, see Fig. 8.

This results in a surface roughness factor of

$$
W = \sqrt{\left(\frac{d_{\text{facet}}}{h_{\text{max}}}\right)^2 + \left(h_{\text{max}}\right)^2},
$$

We performed 2d FEM simulations of the boundary layer in dependence of the applied potential and the electrolyte concentration varying the surface roughness parameter relative to the Debye length as $h_{\text{max}} / L_{\text{Debye}} = \frac{5}{8}, \frac{5}{4}, \frac{5}{2}$. The respective surface roughness factors according to (3.5) are $W = 1.002, 1.0078, 1.031$. We observe that with increasing $h_{\text{max}}$, the capacity curves deviate stronger from the calculated capacity of the perfectly smooth surface, see Fig. 9 left. Moreover, Fig. 9 right indicates that surface roughness can be very accurately taken into account by multiplying the surface roughness factor $W$ to the capacity of the perfectly plain surface, i.e.

$$
C(E) = W(h_{\text{max}}) C_{\text{poly}}(E).
$$
4 Discussion of the idealized polycrystal

Within this discussion, we always apply the limit of large facet size $L_{\text{Debye}} \ll d_{\text{facet}}$, keeping in mind, that this limit is already well approached for $d_{\text{facet}} = 10L_{\text{Debye}}$.

4.1 Difference of PZC

In the previous section, we have kept potential difference between the two considered facets constant (viz. 0.2 V). However, this difference is a material dependent characteristic quantity that varies between different materials or different crystallographic orientations. While differences of the PZC, respective $\mu_{c_e}$, between differently oriented surfaces are considered minor for some metals, e.g. 'low melting metals' like Cd, Sn, Pb, Zn and Bi, cf. [Vor81, TD95], considerable differences between the low index facets are reported for e.g. Ag, Au, Cu, Pt and W, cf. [DKW15]. Thus, we examine the situation of an bi-crystal with equal surface fractions $s_1 = s_2 = 1/2$ and vary the difference of the surface chemical potential of the electrons $\Delta \mu_{c_e} = \mu_{c_e}|_{S_1} - \mu_{c_e}|_{S_2}$ between 0eV and 0.8eV. This corresponds to varying the boundary values such that $0 \leq \Phi_1 - \Phi_2 \leq 0.8V$. We observe, that for $\Phi_1 - \Phi_2 < 0.115V$ the capacity retains the typical camel shape. For $0.115V < \Phi_1 - \Phi_2 < 0.375V$ there is a three maximum shape, whereas for larger potential differences between the facets, we have a four-maximum configuration that allows to identify the local extrema of the individual facets, see Fig. [10]. Once more, we stress that experiments on should be aware of the fact that for a polycrystalline surface the PZC in general is not found at a local capacity minimum! For sufficiently large potential difference $\Phi_1 - \Phi_2$, the PZC will in general be between the inner two local maxima of the capacity but only for specific configuration of the surface fractions $s_i$, the PZC is located at a local capacity minimum. Even more, for moderate $\Phi_1 - \Phi_2$, the PZC is located near a local capacity maximum!

4.2 Adsorption to the surface

To study the effect of adsorption, we return to the simple configuration of a symmetric bi-crystal in Sect. [3.1] i.e. we choose $N = 2, s_1 = s_2 = 1/2$. We assume that the adsorption energies $\Delta g_\alpha$ do...
Figure 10: Capacity curves for varying difference $\Delta \mu_e = \mu_e|_{S_1} - \mu_e|_{S_2}$ for a bi-crystal with equal surface fractions $s_1 = s_2$.

Figure 11: Left: Capacity curves for an adsorbing electrolyte and different bulk electrolyte concentrations where the higher peak at potentials more positive than the PZC is due to adsorption of anions. Right: Construction of the capacity of a polycrystal surface as the sum of the single crystal capacities of the contributing facets weighted by their surface fraction.

not differ between the different facets. In particular, by the choice of

$$\Delta g_A = -0.25\text{eV}, \quad \Delta g_C = +1\text{eV}, \quad \Delta g_S = -0.0735\text{eV}.$$ (4.1)

we assume, that cations effectively do not adsorb while anions and the solvent are allowed to adsorb to the surface. For simplicity and clarity of presentation, we do not consider surface reactions like solvation shell stripping or electron transfer reactions. An extension is straightforward by applying (2.16c) for the reaction products and (2.21) for the surface charge. According to (2.29), the surface capacity of a patterned surface only depends on the material parameters of the contributing facets and their surface fractions $s_i$, but not on the facet geometry or the facet size parameter $d_{\text{facet}}$. Thus, we can combine (1.1) and (2.29) to conclude that for large facet size, the capacity of a perfectly plain polycrystal surface is

$$C_{\text{poly}} = \sum_{i=1}^{N} s_i \hat{C} \left( E - E^{\text{ref}} + \frac{1}{\varepsilon_0} \mu_e |_{\Sigma_i} \right).$$ (4.2)

As already known for single crystal surfaces, if there is adsorption from the electrolyte, the PZC depends on the electrolyte and its bulk concentration, cf. e.g. [LGD16]. The same also holds true for the polycrystal and we conclude from (1.1) and (2.29) that for $L_{\text{Debye}} < d_{\text{facet}} \rightarrow \infty$, the determination of

In the case $d_{\text{facet}} \rightarrow 0$ the analogous result holds true, due to (1.4) and (2.29).
\[ 0 = \sum_{i=1}^{N} s_i \hat{Q} \left( E_{\text{poly}}^{PZC} - E_{\text{ref}}^{PZC} + \frac{1}{\varepsilon_0 \mu_c} |\Sigma_i| \right). \] (4.3)

4.3 Multiple different facets

The above results can easily be extended to more complex polycrystalline surfaces with many facets of different type. We consider the case of \( N = 4 \) different types of facets \( \Sigma_1, \Sigma_2, \Sigma_3, \Sigma_4 \) with the respective surface fractions \( s_1 = 4/9, s_2 = s_3 = 2/9, s_4 = 1/9 \). Also, we want to have junction points where more than two different facets meet. One possible simple realization on a 2D surface is the periodic symmetric continuation of the pattern sketched in Fig. 12. The boundary data is given as \( \varphi = \Phi_1 \) on \( \Sigma_i \), for \( i = 1, 2, 3, 4 \). We choose \( \Phi_1 = -\Phi_2 = -0.1 \) V. First, we let \( \Phi_4 = \Phi_1 \) and \( \Phi_3 = \Phi_2 \) and get slightly non-symmetric capacity curves\(^8\), see Fig. 12. Next, we change \( \Sigma_3 \) such that \( \Phi_3 = -.061 \) V while still \( \Phi_4 = \Phi_1 \). Finally, set \( \Phi_4 = .085 \) V and keep the remaining boundary values as before, see Fig. 12 bottom line.

As in Section 3.1 we checked convergence of the capacity calculated from 3D-FEM simulation for finite facet size \( d^{\text{facet}} \) to the limit equation (1.1) for \( d^{\text{facet}} \to \infty \) and grid convergence of the numerical solutions.

\(^8\)This situation is equivalent to the slightly non-symmetric case \( N = 2 \) and \( s_1 = 5/9, s_2 = 4/9 \).
4.4 Stochastic distribution of surface patterns

A precise measurement of all surface facets and their corresponding work functions is extremely expensive, if possible, and even measurements on very similar facets may contain some scatter. Hence, the input values for the mathematical model of the foregoing section can only be determined to a certain precision, which has to be taken into account when realistic polycrystalline surfaces are to be described. We sketch therefore a stochastic description of the polycrystalline surface based on the derived deterministic model and provide some numerical examples of the double layer capacity.

Consider the set \( S^{\mu} \) of all possible values of the surface electron chemical potential \( \mu^s_e \) on the surface \( S \), e.g. \( S^{\mu} = \{ \mu^s_1, \ldots, \mu^s_J \} \). We denote with \( \Sigma^\mu \) all facets which have the value \( \mu^s_e \) for \( \mu_s^e \), i.e.

\[
\Sigma^\mu = \bigcup_{i \in I^\mu} \Sigma_i \quad \text{with} \quad I^\mu := \{ i \in \{ 1, \ldots, N \} | \mu_s^i = \mu \}
\]

(4.4)

to switch the facet labeling from a consecutive numbering of all facets to values of \( \mu^s_e \). We may thus write

\[
\sum_{i=1}^N s_i = \sum_{\mu \in S^{\mu}} s_\mu \int_{-\infty}^{\infty} \delta(\omega - \mu) \, d\omega \quad \text{with} \quad s_\mu = \sum_{i \in I^\mu} s_i,
\]

(4.5)

where \( \delta \) is the Dirac-distribution and consider thus

\[
f(\omega) := \sum_{\mu \in S^{\mu}} s_\mu \delta(\omega - \mu)
\]

(4.6)

as probability density since

\[
\sum_{i=1}^N s_i = \int_{-\infty}^{\infty} f(\omega) \, d\omega = 1.
\]

(4.7)

For realistic polycrystalline electrodes, the Dirac-distribution might smear to Normal-distributions, whereby \( f \) becomes a continuous function.

The charge \( Q_{BL}^{\text{poly}} \) stored in the polycrystalline boundary layer thus rewrites as

\[
Q_{BL}^{\text{poly}} = \sum_{i=1}^N s_i \hat{Q}_{BL}^s \left( E - E_{\text{ref}} + \frac{1}{\varepsilon_0} \mu^s_i \right)
\]

(4.8)

\[
= \sum_{\mu \in S^{\mu}} s_\mu \int_{-\infty}^{\infty} \delta(\omega - \mu) \hat{Q}_{BL}^s \left( E - E_{\text{ref}} + \frac{1}{\varepsilon_0} \omega \right) \, d\omega
\]

(4.9)

\[
= \int_{-\infty}^{\infty} f(\omega) \hat{Q}_{BL}^s \left( E - E_{\text{ref}} + \frac{1}{\varepsilon_0} \omega \right) \, d\omega
\]

(4.10)

\[
= (f * \hat{Q}_{BL}^s)(E - E_{\text{ref}}).
\]

(4.11)

With the convolution operator * we obtain thus a very compact typeface for \( Q_{BL}^{\text{poly}} \) in terms of the distribution function \( f(\omega) \), which describes the facet density parametrized in terms of \( \mu^s_e \). The potential
of zero charge $E_{\text{PZC}}^\text{poly}$ is thus (implicitly) defined via $(f \ast \hat{Q}^\text{BL})(E - E_{\text{ref}}) = 0$ and the double layer capacity

$$C_{\text{poly}}^\text{BL} = \sum_{i=1}^{N} s_i \hat{C}^\text{BL} \left( E - E_{\text{ref}} + \frac{1}{\varepsilon_0} \mu_s^i \right)$$

(4.12)

$$= \int_{-\infty}^{\infty} f(\omega) \hat{C}^\text{BL} \left( E - E_{\text{ref}} + \frac{1}{\varepsilon_0} \omega \right) d\omega$$

(4.13)

$$= (f \ast \hat{C}^\text{BL})(E - E_{\text{ref}}).$$

(4.14)

Note that $\hat{Q}^\text{BL}$ and $\hat{C}^\text{BL}$ remain the non-linear functions described in section 2.

In order to show the consequences of the smearing of the convolution operator $\ast$ on the non-linear functions of $C^\text{BL}$ we consider the following example. Consider a normal distribution of facets with respect to its mean value $(\mu_s - \overline{\mu}_s) = 0$ where $\sigma$ is the standard deviation, see Fig. 4.

For small standard deviation like $\sigma = 0.025$, the resulting capacity according to (1.3) is close to the single crystal capacity, see Fig. 4. However, for $\sigma = 0.2$, the local capacity minimum has disappeared and there is only a single capacity maximum. Due to the symmetry of the chosen density $f(\omega)$, the PZC is located at the capacity maximum. We want to stress, that this particular configuration of facets with a broad scatter of the individual facet properties is very different from the limit related to (1.4) where on each finite size part of the surface a large number of accordingly small facets are considered.

Next, we want to investigate the effect of a polycrystal surface mainly covered by facets of low index surfaces like (100), (110) and (111). For the low index facets of Ag we chose values of $\mu_s$ related to the (negative) work function as given in Table 2. To describe a non-ideal Ag polycrystal, we consider a probability density that consists of normal distributions around the potentials of the low index facets and a standard deviation small enough, such that the peaks do not overlap, see Fig. 13 where $\sigma = 0.025$ is chosen. The resulting capacity curve remains close to the capacity of an ideal polycrystal with equal surface fractions $s_i = \frac{1}{3}$ for $i = 1, 2, 3$. Moreover, we perturb the configuration such that an equi-distribution in a 1V potential range is superimposed. As a result, we obtain a much more smooth capacity curve. The pure equi-distribution for $\omega(u)$ results in a capacity similar to the normal distribution with large standard deviation.

Table 2: Recommended values for the electron work function on Ag surfaces according to [DKW15].

<table>
<thead>
<tr>
<th>surface</th>
<th>work function</th>
</tr>
</thead>
<tbody>
<tr>
<td>(110)</td>
<td>4.10 eV</td>
</tr>
<tr>
<td>(100)</td>
<td>4.36 eV</td>
</tr>
<tr>
<td>(111)</td>
<td>4.53 eV</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$\mu_s - \overline{\mu}_s$</th>
</tr>
</thead>
<tbody>
<tr>
<td>+0.2 eV</td>
</tr>
<tr>
<td>-0.06 eV</td>
</tr>
<tr>
<td>-0.23 eV</td>
</tr>
</tbody>
</table>

5 Conclusions

By the approach of numerically solving an improved Poisson-Boltzmann problem for patterned surfaces we provide justification for classical simple equations describing the capacity of polycrystalline surfaces. The classical formulas [BDL80, Vor81] have been developed from simple models of the electrochemical double layer at a time before the measurement of clean single crystal surfaces was brought to perfection by [Val81, Val82, Val89]. The improved double layer model [DGM13, DGM18] was successfully validated.
Figure 13: Left: different probability densities describing the surface chemical potential of electrons on a Ag-polycrystal where the surface is covered by facets close to the three low index facets (100), (110) and (111) (blue), the low index facets and additionally an equi-distribution other facets (red), an a pure equi-distribution of facets (yellow). Right: Capacity curves for the respective densities (solid —) and capacity of the ideal polycrystal of only the three low index facets with equal surface fraction (dotted line · · ·).

against these most reliable experimental data [LGD16] and now allows to turn the attention from the single crystal to the less specific conditions of polycrystalline surfaces.

While for a not too small facet size parameter $d_{\text{facet}}$, the result for the capacity of a polycrystalline surface is the expected surface fraction weighted average, some caution is required when considering PZC. Due to the non-linearity in the relation between applied potential and boundary layer charge, the PZC is not given by a weighted average and moreover, it also varies with the electrolyte concentration. The result for $d_{\text{facet}} \to 0$ can be interpreted in the way that for an amorphous solid material or a liquid material the surface is inhomogeneous on an infinitesimal scale such that the surface properties for each point are independent. Both limits, for large and for small facet size, allow a to get rather clear picture of the electrochemical double layer at patterned surfaces like polycrystal surfaces. Moreover, the results justify the simple correction for rough surfaces by a multiplication with a single geometry dependent factor.

Another interesting aspect is the possibility to develop a stochastic description of complex polycrystalline surfaces. The results indicate that assembly of an electrode from a huge number of particles with different properties can be represented by a distribution function with a large standard deviation. As a consequence, the capacity of such a surface does not show the typical single crystal camel shape or multiple clearly separated maxima but a rather constant capacity over a broader potential range. In light of this result, the function of supercapacitors can be explained by the inhomogeneity of the fine faceted granular electrode surface.

**Outlook.** In the modeling of the patterned surfaces with finite facet size parameter $d_{\text{facet}}$, we neglected distinguished contributions from the facet boundary lines. While the according effects on the capacity might be neglected for large enough facets, accurate description of the finite size facet structure might require an extension of the underlying thermodynamic continuum model to line thermodynamics.

Surface reactions like solvation shell striping of adsorbed ions can be included into the presented models for polycrystalline surfaces in a straightforward manner. The same also holds true for electron transfer reactions, as long as the reaction products do not leave the surface. In the presence of Faradayic currents additional length scales for the non-equilibrium transport of ions have to be taken into account. In non-equilibrium, surface roughness on larger length scales might lead to interactions of...
bulk transport an layer charging causing deviations from ideal behavior of perfectly plain surfaces.

References


