

Magnetic-field-assisted electrodeposition of metal to obtain conically structured ferromagnetic layers

Huang, M.; Eckert, K.; Mutschke, G.;

Originally published:

October 2020

Electrochimica Acta 365(2020), 137374

DOI: <https://doi.org/10.1016/j.electacta.2020.137374>

Perma-Link to Publication Repository of HZDR:

<https://www.hzdr.de/publications/Publ-31638>

Release of the secondary publication
on the basis of the German Copyright Law § 38 Section 4.

CC BY-NC-ND

Magnetic-field-assisted electrodeposition of metal to obtain conically structured ferromagnetic layers

Mengyuan Huang^{a,*}, Kerstin Eckert^b, Gerd Mutschke^{a,*}

^a*Helmholtz-Zentrum Dresden-Rossendorf (HZDR), Institute of Fluid Dynamics, 01328 Dresden, Germany*

^b*Technische Universität Dresden, Institute of Process Engineering, 01069 Dresden, Germany*

Abstract

Micro- or nano-structured ferromagnetic layers often possess superior electrocatalytic properties but are difficult to manufacture in general. The present work investigates how a magnetic field can possibly support local cone growth on a planar electrode during electrodeposition, thus simplifying fabrication. Analytical and numerical studies were performed on conical structures of mm size to elaborate the influence of the magnetic forces caused by an electrode-normal external field. It is shown that, beside the Lorentz force studied earlier in the case of single cones [1], the magnetic gradient force enabled by the field alteration near the ferromagnetic cathode significantly supports cone growth. Detailed studies performed for sharp and flat single cones allow conclusions to be drawn on the support at different stages in the evolution of conical deformations. Furthermore, the influence from neighboring cones is studied with arrays of cones at varying distances apart. Nearby neighbors generally tend to mitigate the flow driven by the magnetic forces. Here, the support for cone growth originating from the magnetic gradient force is less heavily affected than that from the Lorentz force. Our results clearly show that the magnetic field has a beneficial effect on the growth of ferromagnetic conical structures, which could also be useful on the micro- and nanometer scales.

Keywords:

metal electrodeposition, magnetic field, surface-structured electrode, Lorentz force, magnetic gradient force, numerical simulation

*Corresponding authors.

Email addresses: m.huang@hzdr.de (Mengyuan Huang), g.mutschke@hzdr.de (Gerd Mutschke)

1. Introduction

Conical micro- and nano-structures have numerous applications due to their particular physical and chemical properties which appear when length scales are reduced. Inspired by cicada wings, super-hydrophobic surfaces covered with arrays of conical protuberances were reported to have good self-cleaning properties [2, 3]. For electrochemical applications, nano-structured electrodes are of great importance due to their enhanced catalytic activity. Great efforts have been devoted to increasing the activity of noble-metal-free catalysts by surface nano-structuring in order to replace the rare, expensive Pt-based catalysts widely used in fuel cell technologies [4, 5, 6]. In this respect, conically structured ferromagnetic electrodes have also been reported to reduce the overpotential for the hydrogen evolution reaction by enhancing the active surface area and supporting the bubble detachment [7, 8].

Among the various methods of synthesizing arrays of micro- and nano-cones, electrochemical deposition techniques have been widely applied [9, 10, 11]. As the current density can be expected to be greater at cathode regions closer to the counter-electrode, according to Faraday's law, the deposition rate is higher at e.g. the tip of a cone compared to other flat electrode regions [12]. This mechanism generally supports the growth of non-uniformities during template-free deposition. In order to further enhance a possible structuring effect during deposition, magnetic fields may be utilized. In the past, magnetic fields have already proven a promising tool for controlling the mass transport of the ionic species during electrodeposition processes. A variety of effects can be achieved, among them the enhancement of the limiting current [13, 14, 15, 16], the homogenization of the deposit thickness [17, 18], the modification of the morphology of the deposited layers [19, 20, 21], the reduction of dendrite growth in Li metal batteries [22, 23], and the enhancement of hydrogen evolution in electrolysis [24, 25, 26, 27]. Structured deposits down to the micrometer scale were obtained using planar but magnetically patterned electrodes. Due to the strongly inhomogeneous magnetic field created locally, a correspondingly patterned deposit follows [28, 29]. However, the local modification of the magnetic field during growth of a ferromagnetic elevation has not yet been investigated in detail, which adds motivation for the present study.

It is well known that applying a magnetic field during electrodeposition causes magnetic forces to act on the electrolyte and to drive a flow. These forces are the Lorentz force [13, 30, 31], and possibly also the magnetic gradient force [32, 33, 34]. If the magnetic forces can be designed such as to generate an electrolyte flow which brings fresh electrolyte towards the tip of a cone, local mass transfer would be enhanced and, thus, cone growth would be supported. For the Lorentz force generated by a surface-normally oriented magnetic field, this beneficial effect has firstly been shown near single copper cones [1]. Here, simulations and measurements based on shadowgraphy and Mach-Zehnder interferometry were performed, and a

41 moderate effect could be observed in a weak magnetic field of 60 mT. The Lorentz
 42 force is given by the vector product of the current density \mathbf{j} and the magnetic flux
 43 density \mathbf{B} ,

$$\mathbf{f}_L = \mathbf{j} \times \mathbf{B} \quad (1)$$

44 Here, as the electrolyte velocity is usually low ($U \ll 1\text{m/s}$), flow-induced parts of
 45 the current density and the magnetic field may safely be neglected [35]. Figure 1(a)
 46 schematically shows the flow near a conical cathode driven by the Lorentz force \mathbf{f}_L
 47 originating from an external magnetic field directed vertically upwards (\mathbf{B}_0). For
 48 diamagnetic and paramagnetic media, there is $\mathbf{B} \approx \mathbf{B}_0$. Then, the vertical magnetic
 49 field and the radial component of the current density caused by the electric field lines
 50 bending near the cone surface create an azimuthal flow around the cone [1]. The
 51 centrifugal acceleration of the electrolyte away from the cone caused by this primary
 52 rotating flow then gives rise to a secondary downward flow which brings fresh bulk
 electrolyte to the cone tip and enriches the boundary layer [33].

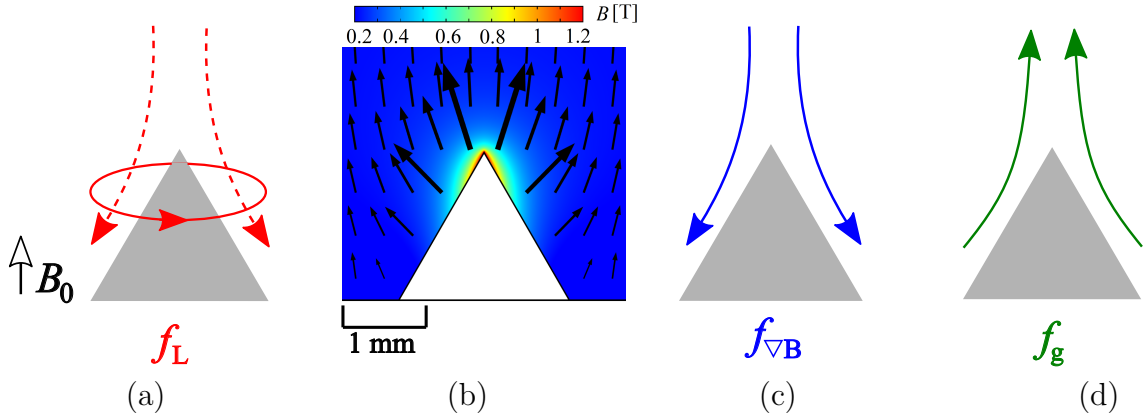


Figure 1: Sketch of the flows generated by (a) the Lorentz force \mathbf{f}_L (primary and secondary flow indicated by solid and dashed lines, respectively), (c) the magnetic gradient force $\mathbf{f}_{\nabla B}$ and (d) the buoyancy force \mathbf{f}_g near a conical cathode under the influence of an external vertical magnetic field B_0 . (b) Vectors and color contours of the magnetic flux density \mathbf{B} near a single Fe cone (diameter 2mm, tip angle 60° , $B_0 = 200$ mT; simulation result).

53

54 If the deposition is performed in an inhomogeneous magnetic field, the gradient
 55 of the magnetic field gives rise to a magnetic gradient force, which may also cause
 56 electrolyte flow and thus affect the deposition processes. If the length scale over
 57 which the magnetic field changes is small, e.g. below 1 mm, a strong field gradient
 58 is created, and the magnetic gradient force may even dominate over the Lorentz
 59 force [33]. In the case of the ferromagnetic layers studied here, the field gradient is
 60 caused by the magnetization of the cones distorting the uniform external magnetic
 61 field. Figure 1(b) shows the magnetic field distribution near a Fe cone in the vertical
 62 center plane when a uniform external vertical field of 200 mT is applied. Here, the
 63 magnetization of the Fe cone has not yet completely reached the saturation level
 64 [36], and large magnitudes of magnetic induction are visible near the tip, creating

65 strong field gradients. The magnetic gradient force is given as [33]:

$$\mathbf{f}_{\nabla\mathbf{B}} = \chi_{\text{sol}}/\mu_0(\mathbf{B} \cdot \nabla)\mathbf{B}, \quad \chi_{\text{sol}} = \sum_i \chi_i^{\text{mol}} c_i + \chi_{\text{H}_2\text{O}} \quad (2)$$

66 Here, $\mu_0 = 4\pi \cdot 10^{-7} \text{ Vs}/(\text{Am})$, c_i , χ_i^{mol} , $\chi_{\text{H}_2\text{O}}$ and χ_{sol} denote the vacuum perme-
 67 ability, the concentration and the molar magnetic susceptibility of species i , and the
 68 magnetic susceptibilities of water and the solution, respectively.

69 It has been pointed out previously that for electrodeposition performed in a
 70 closed electrochemical cell bounded by walls, the potential parts of forces are bal-
 71 anced by the wall pressure and cannot drive electrolyte flow [30]. Therefore, with
 72 respect to $\mathbf{f}_{\nabla\mathbf{B}}$ studied here, the rotational part is responsible for the resulting flow
 73 [33]:

$$\nabla \times \mathbf{f}_{\nabla\mathbf{B}} = \frac{1}{2\mu_0} \left(\sum_i \chi_i^{\text{mol}} \nabla c_i \right) \times (\nabla B^2) \quad (3)$$

74 Thus, the flow forced by $\mathbf{f}_{\nabla\mathbf{B}}$ is determined by the gradients of the species concen-
 75 trations and of the magnetic field. As the concentration gradient near the cathode
 76 develops with ongoing deposition, the magnitude and direction of $\mathbf{f}_{\nabla\mathbf{B}}$ will vary over
 77 time. Fig. 1 (c) shows a possible flow pattern caused by $\mathbf{f}_{\nabla\mathbf{B}}$.

78 Beside the magnetic forces, the buoyancy force which arises from variations in
 79 the density of the electrolyte may also cause electrolyte flow during electrodeposition
 80 [37]:

$$\mathbf{f}_{\mathbf{g}} = (\rho - \rho_0)\mathbf{g} \quad (4)$$

81 Here, ρ , ρ_0 and $\mathbf{g} = -9.81 \text{ m/s}^2 \mathbf{e}_z$ denote the local and the bulk density of the elec-
 82 trolyte and the vector of the gravitational acceleration which points in a downward
 83 direction $-\mathbf{e}_z$ (see Fig. 2 for the coordinate system). Density variations originate
 84 from a spatially varying electrolyte composition caused by electrode reactions or
 85 from temperature variations in the electrolyte caused by Ohmic heating. However,
 86 the thermal effect is often much smaller than the solutal one and can be safely
 87 neglected, see Section 2.3.

88 In the following we consider electrochemical cells where the cathode is placed
 89 at the bottom. The metal deposition at the cathode reduces the density of the
 90 electrolyte, and solutal buoyancy tends to bring upward the lighter electrolyte, as
 91 shown in Fig. 1(d). As the cathodes considered are not planar in shape, even in
 92 the case of homogeneous deposition, $\mathbf{f}_{\mathbf{g}}$ has a non-zero rotational part, which drives
 93 a horizontal flow towards the foot of the cone and an upward flow along the cone
 94 surface:

$$\nabla \times \mathbf{f}_{\mathbf{g}} = g \nabla \rho \times \mathbf{e}_z \quad (5)$$

95 As the deposition starts, this flow sets in unconditionally, which is different from
 96 solutal convection at planar horizontal electrodes; this requires a critical Rayleigh
 97 number to start [38]. The upward flow along the cone can be expected to generate a

concentration boundary layer of growing thickness. Thus, near the cone tip, the concentration gradient and diffusive mass transfer may be weakened, and cone growth supported by only the geometrical non-uniformity of the cathode may be impeded. In this respect, properly adjusted magnetic fields offer the possibility to force an opposing flow that is strong enough to compensate for the unfavorable buoyancy flow and thus to enhance the desired structuring effect.

Extending the reflections to include planar cathodes with several conical elevations (see Fig. 2), the flow generated near each single cone may interact with flow originating from its neighbors. Corresponding damping of the azimuthal flow near magnetically templated electrodes was already reported in [39]. However, the neighbor influence on the magnetization, the resulting magnetic gradient force and also the impact of buoyancy has not been studied so far.

This work aims at studying the utilization of magnetic fields for the electrodeposition of conically structured ferromagnetic deposits. Although the majority of today's practical applications are for micro- and nano-sized conical structures, here we perform first investigations on a larger scale to gain a basic understanding of the magnetic field effects. This work extends the earlier study related to Lorentz force effects with single cones [1] and firstly elaborates on the magnetic gradient force, which plays an important role at ferromagnetic layers. The study includes single cones of different shapes and also the influence of neighboring cones at varying distances from one another. Thus, a detailed understanding is gained of the contributions made by the different forces involved during electrodeposition in a vertical magnetic field.

2. Methods

2.1. Problem description and simulation approach

We perform transient numerical simulations for the galvanostatic electrodeposition of metal on planar cathodes with periodic arrangements of conical elevations. The electrodes are oriented horizontally, and the electrochemical cell is exposed to a uniform magnetic field which is oriented vertically. We consider cathodes made of iron and additionally also of copper, to compare with earlier investigations at diamagnetic electrodes [1]. In order to study the basic effects of both magnetic forces, as well as the influence of neighbor effects, we start with cones of mm size, which will also facilitate the later experimental validation of our simulation results. The deposition times considered are in the range of minutes. We can therefore neglect changes of the electrode shape during deposition and treat the electrode geometry as fixed. Hence, the magnetic field near the electrodes also does not change during deposition. In order to further exclude any possible influence from the different reaction kinetics of the metals deposited, we deposit copper at both the ferromagnetic

136 and the diamagnetic cathodes. We also neglect any possible side reactions such as
137 the hydrogen evolution reaction.

138 Our simulation approach consists of three steps which are outlined in Fig. 2. In
139 the first step, we study the generic behavior at single cones and neglect any influence
140 from neighboring cones, which are assumed to be far away. Thus, in a cylindrical
141 region with the cone placed in the center of the cathode, the problem can be assumed
142 to be axisymmetrical. The computational domain is thus the vertical plane bounded
143 by the symmetry axis and the outer mantle boundary, and by the electrodes (see
144 Fig. 2(a-b)). The simulations allow for an azimuthal flow caused by the Lorentz
145 force, but neglect any dependencies on the azimuthal (θ) position.

146 In the second step, we take neighbor effects into account. We consider an in-
147 finitely extended planar cathode with a regular checkerboard arrangement of conical
148 deformations (see Fig. 2(c-d)). In the following we assume that the electrodeposition
149 proceeds identically at all cones. Global effects originating from e.g. recirculating
150 flows around electrodes of finite size [28] are beyond the scope of this study. We
151 therefore pick a cuboid computational domain between the electrodes where the
152 cone is located in the center of the cathode square. The side length of the square
153 corresponds to the cone distance, which will be varied later to study different cone
154 densities. At the opposite vertical faces of the cuboid domain, periodic boundary
155 conditions will be applied in the simulations.

156 As these simulations have to be performed in 3D and are expensive in terms
157 of computing resources, in the third step, we introduce an approximate scheme
158 for considering the neighbor influence. In order to facilitate simpler axisymmetric
159 simulations, we assume that there are neighboring cones at the same distance for all
160 azimuthal directions. Thus, the cylindrical domain shown in Fig. 2(a-b) can be used
161 again. Here, in contrast to Step 1, the radial extension is defined as half the distance
162 between the cones. At this outer domain boundary, for reasons of symmetry, the flow
163 has no radial component, and the azimuthal component originating from the Lorentz
164 force can be expected to be canceled out, as the azimuthal flow direction is opposite
165 between neighbors. This approximation allows 2D simulations to be performed
166 instead of 3D simulations as a means of considering the influence of neighboring
167 cones, and will be validated below by presenting a comparison of the two simulation
168 approaches for the case of the copper cones. The boundary conditions applied in
169 the three steps will be explained in full detail in Section 2.2.

170 *2.2. Model equations and boundary conditions*

171 The numerical model consists of a coupled system of equations for the magnetic
172 field, the electrolyte velocity, the concentration of the ionic species and the electric
173 field. The computational domains for the three steps of the simulations are outlined
174 in Fig. 2(b)(d), and the boundary conditions applied are summarized in Table 1.

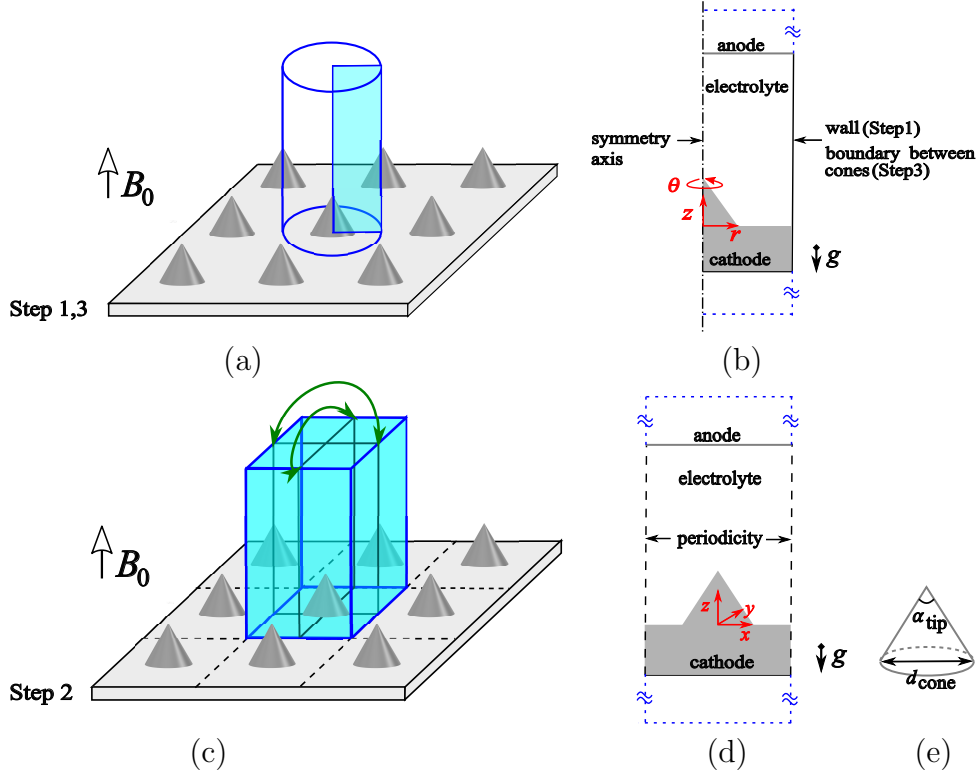


Figure 2: Sketch of the computational domains (marked in light blue), the related coordinate systems (cylindrical in (b) and Cartesian in (d)) and the specific boundary conditions applied to illustrate the three steps of the computational approach. (a-b) Step 1 and Step 3. (c-d) Step 2. At the opposite vertical faces of the cuboid, green arrows indicate the periodic boundary conditions which are applied. The dashed blue lines indicate the vertically extended domains for calculating the magnetic field. (e) Sketch of the cone geometry with α_{tip} and d_{cone} denoting the cone tip angle and the cone diameter. For more details see Sections 2.1 and 2.2.

175 The magnetic field is obtained by solving the Maxwell equations [40]:

$$\nabla \cdot \mathbf{B} = 0, \quad \nabla \times \mathbf{H} = 0 \quad (6)$$

176 The magnetic field strength \mathbf{H} is linearly related to the magnetic flux density \mathbf{B} by
 177 the vacuum permeability μ_0 for the paramagnetic and diamagnetic materials in our
 178 calculations, i.e. $\mathbf{B} = \mu_0 \mathbf{H}$. Inside the ferromagnetic cathodes, there is

$$\mathbf{B} = \mu_0(\mathbf{H} + \mathbf{M}) \quad (7)$$

179 Here, the magnetization \mathbf{M} is given by a corresponding magnetization curve
 180 [41]. \mathbf{H} is the sum of the external field strength (\mathbf{B}_0/μ_0) and the demagnetizing
 181 field, the latter being directed antiparallel to \mathbf{M} in ferromagnetic materials [42].
 182 We apply $\mathbf{B} = \mathbf{B}_0$ on the outer horizontal boundaries of the magnetic field domain
 183 for all three steps and at the vertical side wall of Step 1, because a homogeneous
 184 external magnetic field is imposed on the cell. For Steps 2 and 3, the neighboring
 185 effects imply symmetry conditions at the vertical boundaries between the cones. We
 186 therefore apply periodic conditions at the oppositely located vertical boundaries of
 187 the cuboid domain in Step 2, and $\mathbf{B} \cdot \mathbf{n} = 0$ at the outer radius in Step 3. Here \mathbf{n}

denotes the unit vector normal to the boundary. It should be mentioned that the magnetic boundary conditions applied here exclude specific deposition effects at the outer edges of the planar part of the ferromagnetic electrode found elsewhere [16].

The electrolyte velocity \mathbf{U} is obtained by solving the Navier-Stokes equations including the volume forces mentioned above and complemented by the incompressibility constraint,

$$\rho_0 \left(\frac{\partial \mathbf{U}}{\partial t} + (\mathbf{U} \cdot \nabla) \mathbf{U} \right) = -\nabla P + \mu \nabla^2 \mathbf{U} + \mathbf{f}_L + \mathbf{f}_{\nabla B} + \mathbf{f}_g, \quad \nabla \cdot \mathbf{U} = 0 \quad (8)$$

Here, P and μ denote the dynamic pressure and the dynamic viscosity of the electrolyte. A no-slip boundary condition is applied at the electrodes for all steps and at the vertical side wall for Step 1. For Step 2, periodic boundary conditions are enforced at the vertical boundaries. For Step 3, symmetry with respect to the neighbor influence allows a free-slip boundary condition to be applied ($\mathbf{U} \cdot \mathbf{n} = 0$) at the outer radius, supplemented by the constraint that the azimuthal velocity (U_θ) has to vanish.

The distribution of species in the electrolyte is obtained by solving the Nernst-Planck equations of dilute solutions:

$$\frac{\partial c_i}{\partial t} = -\nabla \cdot \mathbf{N}_i \quad (9)$$

Here, \mathbf{N}_i denotes the flux density of species i

$$\mathbf{N}_i = -z_i F \frac{D_i}{RT} c_i \nabla \phi - D_i \nabla c_i + c_i \mathbf{U} \quad (10)$$

which, beside transport by migration and diffusion, also includes convective transport by the electrolyte velocity from Eq. (8) [37]. Here, z_i , D_i , F , R , T and ϕ are the charge number and the diffusion coefficient of species i , the Faraday constant, the universal gas constant, the temperature and the electric potential, respectively. The electric current density in the solution is the net flux density of all charged species:

$$\mathbf{j} = F \sum_i z_i \mathbf{N}_i \quad (11)$$

We further assume electroneutrality in the electrolyte, i.e. $\sum z_i c_i = 0$. Combining Eqs. 10, 11 and the conservation of charge ($\nabla \cdot \mathbf{j} = 0$), the electric potential can be obtained as the solution of a Poisson equation:

$$\nabla \cdot \left(-F^2 \nabla \phi \sum_i z_i^2 \frac{D_i c_i}{RT} \right) - \nabla \cdot \left(F \sum_i z_i D_i \nabla c_i \right) = 0 \quad (12)$$

Regarding the boundary conditions applied, on electrically passive walls, due to electric insulation, the normal components of the electric current and the flux of all

214 ionic species have to vanish for Step 1. The same boundary conditions are applied
 215 for Step 3, but for reasons of symmetry. For Step 2, periodic boundary conditions
 216 are again implemented. On the electrodes, a Butler-Volmer relation is applied to
 217 describe the kinetics of the copper reaction for all three steps [37]:

$$j_n = j_0 \left(\exp \left(\frac{\alpha_a F \eta_s}{RT} \right) - \exp \left(\frac{-\alpha_c F \eta_s}{RT} \right) \right) \quad (13)$$

218 Here j_0 , α_a and α_c denote the exchange current density and the apparent transfer
 219 coefficients. The surface overpotential η_s is defined as the potential of the electrode
 220 ϕ_e relative to the solution potential ϕ minus the equilibrium electrode potential
 221 ϕ_{eq} given by the Nernst equation, i.e. $\eta_s = \phi_e - \phi - \phi_{eq}$ [37]. In the simulations,
 222 the electrode potential is adjusted at each time step to satisfy the galvanostatic
 condition $I = I(\phi_e) = \text{const.}$

Table 1: Boundary conditions applied in the different steps of the simulations. The subscripts "src" and "dst" denote the source and the destination boundaries in the case of periodic boundary conditions (see Fig. 2 (c-d)).

| | Boundary | Step 1 | Step 2 | Step 3 |
|----------------------|------------|--|---|---|
| Magnetic field | Axis | $\partial \mathbf{B} / \partial r = 0$ | - | $\partial \mathbf{B} / \partial r = 0$ |
| | Vertical | $\mathbf{B} = \mathbf{B}_0$ | $\mathbf{B}_{\text{src}} = \mathbf{B}_{\text{dst}}$ | $\mathbf{B} \cdot \mathbf{n} = 0$ |
| | Horizontal | $\mathbf{B} = \mathbf{B}_0$ | | |
| Electrolyte velocity | Axis | $\partial \mathbf{U} / \partial r = 0$ | - | $\partial \mathbf{U} / \partial r = 0$ |
| | Vertical | $\mathbf{U} = 0$ | $\mathbf{U}_{\text{src}} = \mathbf{U}_{\text{dst}}$ | $\mathbf{U} \cdot \mathbf{n} = 0, U_\theta = 0$ |
| | Electrodes | $\mathbf{U} = 0$ | | |
| Electric field | Axis | $\partial \mathbf{j} / \partial r = 0$ | - | $\partial \mathbf{j} / \partial r = 0$ |
| | Vertical | $\mathbf{j} \cdot \mathbf{n} = 0$ | $\mathbf{j}_{\text{src}} = \mathbf{j}_{\text{dst}}$ | $\mathbf{j} \cdot \mathbf{n} = 0$ |
| | Electrodes | $\mathbf{j} \cdot \mathbf{n}$ from Eq. 13 | | |
| Species | Axis | $\partial \mathbf{N}_i / \partial r = 0$ | - | $\partial \mathbf{N}_i / \partial r = 0$ |
| | Vertical | $\mathbf{N}_i \cdot \mathbf{n} = 0$ | $\mathbf{N}_{i,\text{src}} = \mathbf{N}_{i,\text{dst}}$ | $\mathbf{N}_i \cdot \mathbf{n} = 0$ |
| | Electrodes | Passive ions: $\mathbf{N}_i = 0$ Active ions: $\mathbf{N}_i \cdot \mathbf{n}$ from Eq. 11, Eq. 13 | | |

223

224 2.3. Material and simulation parameters

225 The material parameters for solving Eqs. 8-13 are listed in Table 2. The elec-
 226 trolyte is an aqueous 0.1 M CuSO_4 solution at room temperature. Because of elec-
 227 trical neutrality, it holds that $c_{\text{Cu}^{2+}} = c_{\text{SO}_4^{2-}}$, and the species index i can be omitted
 228 in the following. With respect to the reaction kinetics of copper, the exchange cur-
 229 rent density j_0 is given as $j_0 = j_{00} (c_s/c_0)^\gamma$, with c_s and c_0 denoting the surface and
 230 the bulk concentration of the electroactive Cu^{2+} species. The reference exchange
 231 current density j_{00} and the kinetic parameter γ are taken from [43]. In the simu-
 232 lations performed, we found that the influence of the concentration-related term of
 233 the Nernst equation on the surface overpotential is negligibly small. We therefore
 234 take ϕ_{eq} to be equal to the standard equilibrium potential, $\phi_{eq,0}$, which reduces the
 235 non-linearity of the boundary condition.

236 The simulation parameters are summarized in Table 3. For the single cone
 237 studies of Step 1, a cone is considered with a base diameter of $d_{\text{cone}} = 2$ mm and a
 238 tip angle of $\alpha_{\text{tip}} = 60^\circ$. These dimensions are varied only in the cone shape studies,
 239 see Section 3.2. The chosen cell height of 30 mm is large enough to avoid any
 240 influence from the anodic mass transfer at the cathode. The radial extension of the
 241 electrochemical cell is chosen to be $5 d_{\text{cone}}$, which is large enough to ensure that the
 242 flow forced near the cone is not affected by the side wall. The current density used
 243 for the simulations in the three steps varies between 8 and 16 mA/cm². In order to
 244 obtain comparable amplitudes of the Lorentz force, the magnetic field amplitude is
 245 accordingly adjusted between 200 and 400 mT.

246 We now come back to estimating the relative significance of thermal and solutal
 247 buoyancy. For the maximum current density considered, of 16 mA/cm², when the
 248 Joule heat is assumed to be completely transferred to the electrolyte, the local tem-
 249 perature rises by about 0.2 K within one minute. The corresponding thermal density
 250 variation is less than 0.01% [44, 45]. However, assuming there is a concentration
 251 variation of 0.1 M, the solutal density variation is more than 100 times larger and
 252 reaches about 1.6% [45, 46]. Thermal buoyancy can therefore safely be neglected
 253 in the following. As the solutal density variation is still small, we may apply the
 254 Boussinesq approximation [38], and the buoyancy force expressed by Eq. 4 can be
 255 simplified to:

$$\mathbf{f}_g = \rho_0 \mathbf{g} \beta_{\text{CuSO}_4} (c - c_0) \quad (14)$$

with β_{CuSO_4} denoting the volume expansion coefficient of the electrolyte (Table 2).

Table 2: Material parameters for 0.1 M CuSO₄ at room temperature.

| | | |
|--|------------------------|----------|
| $\chi_{\text{Cu}^{2+}}^{\text{mol}}$ (m ³ /mol) | $1.57 \cdot 10^{-8}$ | [32] |
| $\chi_{\text{H}_2\text{O}}$ | $-9.0 \cdot 10^{-6}$ | |
| ρ_0 (kg/m ³) | 1014 | [46] |
| β_{CuSO_4} (m ³ /mol) | $1.6 \cdot 10^{-4}$ | |
| μ (Pa · s) | $1.04 \cdot 10^{-3}$ | |
| $D_{\text{Cu}^{2+}}$ (m ² /s) | $5.6 \cdot 10^{-10}$ | [43] |
| $D_{\text{SO}_4^{2-}}$ (m ² /s) | $10.04 \cdot 10^{-10}$ | |
| $z_{\text{Cu}^{2+}}$ | 2 | |
| $z_{\text{SO}_4^{2-}}$ | -2 | [37, 43] |
| j_{00} (mA/cm ²) | 10 | |
| $\phi_{\text{eq},0}$ (V) | 0.337 | |
| γ | 0.42 | [37, 43] |
| α_a | 1.5 | |
| α_c | 0.5 | |

256

257 2.4. Numerical details and validations

258 The simulations were performed using the Finite Element software package Com-
 259 sol V.5.5 [41]. Second-order shape functions are used for the magnetic field \mathbf{B} , the

Table 3: Simulation parameters.

| | Step 1 | | | Step 2 | Step 3 |
|--|--------|-----|-----|--------------|--------------|
| α_{tip} | 30° | 60° | 90° | 60° | 60° |
| d_{cone} (mm) | 1.4 | 2 | 2.4 | 2 | 2 |
| Cone surface area (mm ²) | 6.28 | | | 6.28 | 6.28 |
| Cell height (mm) | 30 | | | 30 | 30 |
| Height of magnetic field domain (mm) | 100 | | | 100 | 100 |
| Cell width (mm) | 10 | | | - | - |
| Cone distance (d_{cone}) | - | | | 1.5, 2, 3, 5 | 1.5, 2, 3, 5 |
| $j_{\text{cathode,avg}}$ (mA/cm ²) | 16 | | | 8 | 8 |
| B_0 (mT) | 200 | | | 400 | 400 |
| c_0 (M) | 0.1 | | | 0.1 | 0.1 |

260 velocity \mathbf{U} , the concentration c and the electric potential ϕ , and first-order shape
261 functions are used for the pressure P . The initial conditions for the transient sim-
262 ulations are an electrolyte with a homogeneous bulk concentration c_0 at rest, i.e.
263 $\mathbf{U} = 0$. Time integration was carried out using an implicit backward differentiation
264 formula method of up to the second order. The time step is adjusted by the solver
265 based on error estimation during the calculations, with an upper limit of 0.1 s which,
266 according to a time-step study, ensures the accuracy of the results.

267 Unstructured triangular (2D) or prismatic (3D) meshes were generated and re-
268 fined near the boundaries so as to sufficiently resolve the spatial gradients of the
269 concentration, the velocity and the magnetic field. A mesh study has shown that
270 the steep magnetic field gradients near the Fe cones are most demanding. Thus,
271 the mesh size near the cone surface was chosen to be $0.0005 d_{\text{cone}}$ for the Fe cones
272 and $0.001 d_{\text{cone}}$ for the Cu cones, respectively. This choice also ensures the sufficient
273 resolution of the concentration and velocity boundary layers.

274 For the magnetic field, a domain size study was performed to ensure that the
275 proximity of the top and bottom boundary does not affect the field distribution inside
276 the electrochemical cell. The magnetic field distributions obtained numerically have
277 further been validated with experimental data for different magnet geometries [1].
278 As a result, a domain height of 100 mm was chosen.

279 The full numerical model described above was validated by experimental data
280 from a similar deposition process on a single Cu cone [1] and also by deposition
281 problems at magnetically templated electrodes, where the magnetic gradient force
282 is of importance [33, 34]. As a result, the numerical model is able to deliver accu-
283 rate, reliable information on the magnetic field distribution, the electrolyte flow, the
284 species distribution and the current density distribution in the cell.

285 3. Results and Discussion

286 3.1. Single cones

287 In the following, the generic case of single cones (Step 1) is considered in order to
 288 obtain a basic understanding of the influence of the different volume forces involved
 289 in the deposition process. An intermediate tip angle of 60° is chosen for the copper
 290 and iron cones investigated. Fig. 3 shows the azimuthal flow driven by the Lorentz
 291 force \mathbf{f}_L after a deposition time of 10 s. Surprisingly, an opposite direction of rotation
 292 is observed for the iron cone compared to the copper cone. Moreover, the azimuthal
 293 flow for the iron cone extends over a smaller area than for the copper cone. This
 294 can be understood by considering the Lorentz force of the axisymmetric problem,
 295 which consists of an azimuthal component only:

$$\mathbf{f}_L = (j_z B_r - j_r B_z) \mathbf{e}_\theta \quad (15)$$

296 Here, \mathbf{e}_θ denotes the unity vector in the azimuthal direction (Fig. 2(b)). In general,
 297 when approaching the cone, the vertical current density vectors bend towards the
 298 surface-normal direction of the cone. Thus, $j_r < 0$, $j_z < 0$. For the Cu case, the
 299 magnetic field is not modified by the diamagnetic cathode, and the magnetic field
 300 equals the external vertical field applied, i.e. $B_r = 0$, $B_z = B_0$. Therefore, as shown
 301 on the right-hand side of Fig. 3, $j_r B_0 < 0$ is responsible for the anticlockwise
 rotation seen from the top of the cell.

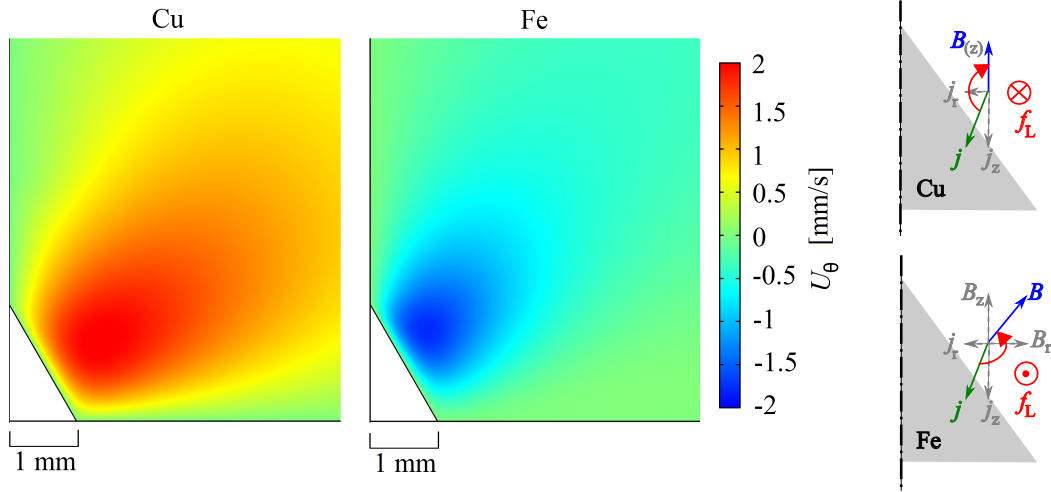


Figure 3: Color surface of the azimuthal velocity after 10 s deposition time near the Cu and the Fe cones ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$). The direction of azimuthal forcing by the Lorentz force is outlined on the far right.

302

303 For the Fe case, the magnetic field is modified in the vicinity of the cone due to its
 304 magnetization, as shown in Fig. 1. Unlike the Cu case, radial components of \mathbf{B} give
 305 rise to negative force amplitudes $j_z B_r$ which may exceed the positive part $(-j_r B_z)$.
 306 The resulting clockwise forcing is shown on the right-hand side of Fig. 3. As large
 307 radial components of the magnetic field are only found close to the surface of the

308 magnetized cone, the region of rotation is smaller compared to the Cu case. Similar
 309 clockwise rotation was also found for the steeper and flatter iron cones investigated
 310 in Section 3.2. However, it should be noted that, irrespective of the direction of the
 311 azimuthal flow, the secondary flow created by the centrifugal force is always directed
 312 downwards towards the cone tip [33, 39].

313 Fig. 4 compares the meridional velocity near the Cu and Fe cones with and
 314 without a magnetic field after 5 s and 10 s of deposition time. In the absence of the
 315 magnetic field, the behavior is identical for Cu and Fe cones. An upward-directed
 316 buoyant flow develops with ongoing deposition, reaching a velocity of about 1 mm/s
 317 above the cone after 10 s of deposition. Similar flow pattern were found in an earlier
 318 study of copper deposition at a copper cone in a weak vertical magnetic field of
 319 60 mT. There, the downward acceleration generated by the Lorentz force was not
 320 strong enough to reverse the direction of flow, and the concentration boundary layer
 was advected upwards and detached at the cone tip [1]. In the following, a stronger

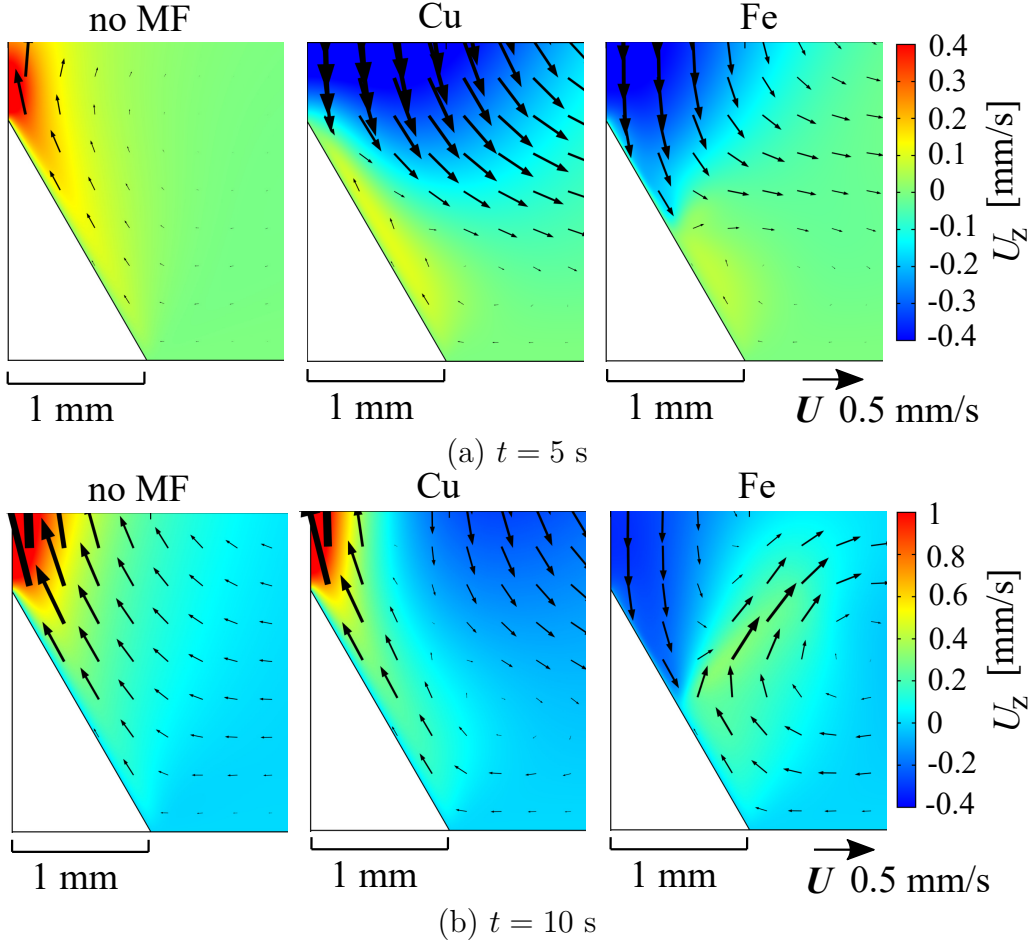


Figure 4: Color surface of the vertical velocity after 5 s and 10 s deposition time near a cone without a magnetic field (left: no MF), near the Cu cone with a magnetic field (middle) and the Fe cone with a magnetic field (right). $\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 16$ mA/cm², $B_0 = 200$ mT. Black arrows represent the meridional velocity vectors.

321 magnetic field of 200 mT is considered. As shown in Fig. 4, near the copper cone
 322 after a deposition time of 5 s a strong downward-directed flow is clearly visible,
 323 which can be attributed to the dominant secondary flow caused by the Lorentz
 324

force. However, the flow direction very close to the cone surface has already turned upwards due to the buoyancy of the concentration boundary layer, indicated by the yellow region. As the deposition continues, the buoyant flow accelerates with time. As shown after 10 s, the direction of flow near the surface of the cone is dominated by buoyancy, which causes the concentration boundary layer to detach from the tip of the cone. A weak downward flow remains at a larger distance from the cone.

For the Fe cone in the magnetic field, the magnetic gradient force $\mathbf{f}_{\nabla\mathbf{B}}$ also comes into play. In comparison to the copper case, the downward flow close to the cone surface seems to be stronger. After 5 s, the downward flow along the upper cone region is clearly visible, whereas the flow velocity near the lower part of the cone is low. After 10 s of deposition, the downward flow along the upper cone region is still maintained, but an upward flow caused by buoyancy is clearly visible in the lower part. This results in a jet-like flow leaving the cone surface in an approximately normal direction at about half the height of the cone. Although the influence of $\mathbf{f}_{\mathbf{L}}$ in the case of the iron cone is slightly weaker than in the case of the copper cone, $\mathbf{f}_{\nabla\mathbf{B}}$ is effectively counteracting $\mathbf{f}_{\mathbf{g}}$ in the vicinity of the cone surface.

In order to further analyze the influence of $\mathbf{f}_{\nabla\mathbf{B}}$ on the deposition process, Figs. 5(a-b) show the concentration of the copper ions close to the iron cone after a deposition time of 5 s and 10 s. The images are superimposed with the meridional velocity vectors. The downward flow along the surface of the upper part of the cone continuously brings fresh bulk electrolyte to the tip, keeping the local concentration boundary layer thin. By contrast, the upward buoyant flow already originates at the horizontal part of the cathode, with a horizontal flow towards the cone, and the thickness of the concentration boundary layer there grows with time. The aforementioned jet-like flow about halfway up the cone is accompanied by the corresponding detachment and advection of the concentration boundary layer in an approximately electrode-normal direction. A similar effect of boundary layer separation caused by the opposing action of $\mathbf{f}_{\mathbf{L}}$ and $\mathbf{f}_{\mathbf{g}}$ during electrodeposition was already reported elsewhere [47].

The concentration boundary layer near the cone, in turn, affects $\mathbf{f}_{\nabla\mathbf{B}}$ and thus the flow field. Applying Eq. 2, based on the material parameters given in Table 2, the magnetic susceptibility of the bulk solution is negative, as the diamagnetic contribution of the water molecules dominates [32]:

$$\chi_{\text{sol}} = \chi_{\text{Cu}^{2+}}^{\text{mol}} c_0 + \chi_{\text{H}_2\text{O}} = -7.43 \cdot 10^{-6} \quad (16)$$

This also holds inside the concentration boundary layer, where $c_{\text{Cu}^{2+}} \leq c_0$ can be assumed. Therefore, $\mathbf{f}_{\nabla\mathbf{B}}$ is acting in the opposite direction to that of the magnetic gradient. As the strongest magnetic field is found near the tip of the cone, $\mathbf{f}_{\nabla\mathbf{B}}$ points away from the tip. However, this contradicts the simulation result, which indicates that the electrolyte flow is directed towards the tip. This issue can easily

be resolved by only considering the rotational parts of the magnetic gradient force and the buoyancy force, as discussed in the introduction. As axisymmetry applies, the curl of both forces defined in Eqs. 3 and 5 in cylindrical coordinates reads:

$$\nabla \times \mathbf{f}_{\nabla B} = \frac{\chi^{\text{mol}}}{\mu_0} \left(\frac{\partial c}{\partial z} B \frac{\partial B}{\partial r} - \frac{\partial c}{\partial r} B \frac{\partial B}{\partial z} \right) \mathbf{e}_\theta \quad (17)$$

$$\nabla \times \mathbf{f}_g = \beta_c g \rho_0 \left(\frac{\partial c}{\partial r} \right) \mathbf{e}_\theta \quad (18)$$

Near the cone surface, the concentration gradient vector essentially points away from the surface into the electrolyte volume, i.e. $\frac{\partial c}{\partial r} > 0$, $\frac{\partial c}{\partial z} > 0$ (see Fig. 5(b)). As the maximal magnetic field exists near the cone tip, along the cone surface $\frac{\partial B}{\partial r} < 0$, $\frac{\partial B}{\partial z} > 0$. According to Eq. 17, the resulting $\nabla \times \mathbf{f}_{\nabla B}$ has a negative azimuthal amplitude and forces the flow to rotate in an anticlockwise direction in the meridional plane, i.e. downwards along the cone surface. In contrast, $\nabla \times \mathbf{f}_g$ has a positive azimuthal amplitude along the cone surface because $\frac{\partial c}{\partial r} > 0$. As the final flow is determined by the curl of both forces, in Fig. 5(c) the sum of $\nabla \times \mathbf{f}_{\nabla B}$ and $\nabla \times \mathbf{f}_g$ is shown after 10 s of deposition time. It can be clearly seen that the upper part of the cone boundary layer is dominated by the negative curl of $\mathbf{f}_{\nabla B}$, whereas the lower part is dominated by the positive curl of \mathbf{f}_g . This means that a downward flow is forced along the surface of the upper cone part, counteracting the upward buoyant flow. These flows force the jet-like departure of the concentration boundary layer from the cone surface at about mid-height. Thus, the azimuthal components of $\nabla \times \mathbf{f}_{\nabla B}$ and $\nabla \times \mathbf{f}_g$ deliver a vivid interpretation of the flow pattern observed.

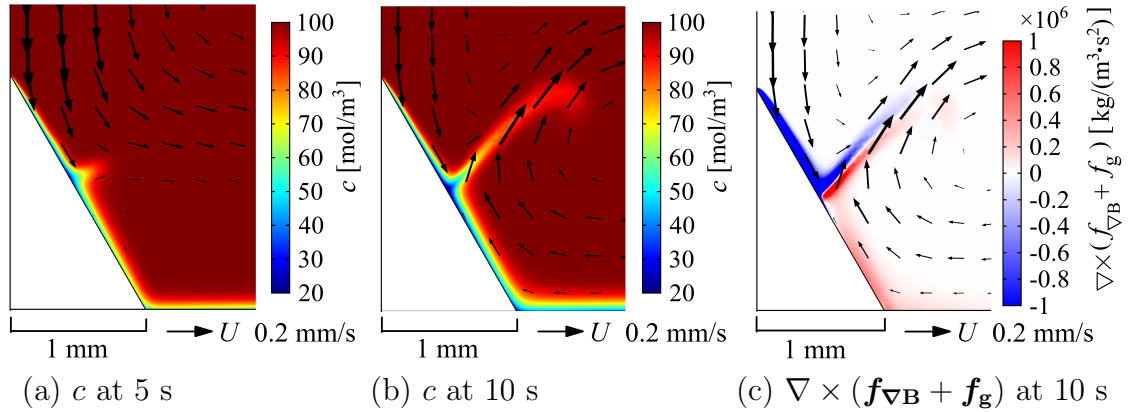


Figure 5: Color surface of the concentration after (a) 5 s and (b) 10 s deposition time and (c) the sum of the curl of the magnetic gradient force and the buoyancy force after 10 s deposition near the Fe cone ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$). Black arrows represent meridional velocity vectors.

381

382 The influence of the magnetic forces on the growth dynamics of conical elevations
383 on the cathode can be evaluated by considering the deposit thickness along the cone
384 surface [37]:

$$d(r, T) = \frac{V_m}{zF} \int_0^T j_n(r, t) dt \quad (19)$$

Here, $V_m = 7.11 \times 10^{-6} \text{ m}^3/\text{mol}$ denotes the molar volume of copper [48], and T denotes the deposition time. Fig. 6 shows the initial height contour of the cone (left) and the deposit thickness along the surface of the Cu and Fe cones after 10 s of deposition time (right). In the case without a magnetic field, the deposit thickness increases monotonically from the foot to the tip of the cone, as a result of the previously mentioned support for cone growth for geometrical reasons. Despite the counteraction of the upward buoyant flow discussed above, in total, cone growth would be supported.

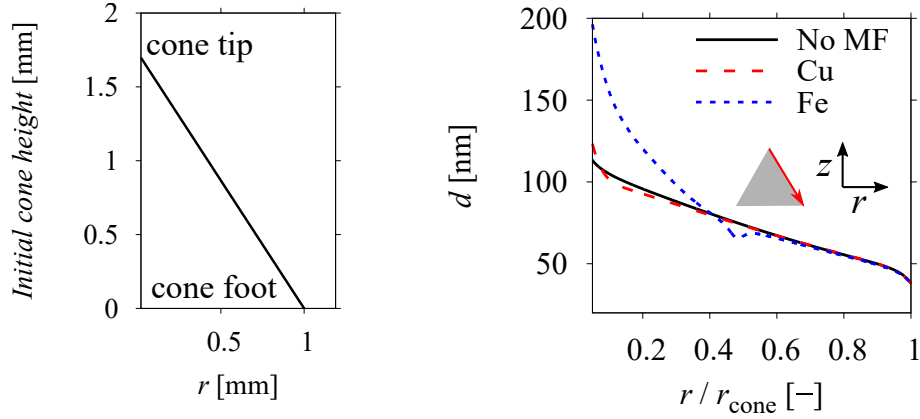


Figure 6: Left: Initial height contour of the cone. Right: Deposit thickness along the cone surface for the cases without magnetic field (No MF) and for Cu and Fe cones in a vertical magnetic field after 10 s deposition time ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$). The horizontal axis is the radial surface coordinate normalized by the cone radius r_{cone} .

392

393 In the case of the copper cone in the vertical magnetic field, small differences
 394 are found with respect to the case without a magnetic field. Close to the tip of the
 395 cone, the thickness of the deposit is increased. This is a result of the downward
 396 secondary flow caused by \mathbf{f}_L , which initially supersedes the upward buoyant flow.
 397 This reversal of the flow direction in the tip region in the first few seconds of the
 398 deposition enriches the boundary layer and thus enhances mass transfer, see Fig. S1
 399 in the Supplementary Information (SI). In the region below, down to about half
 400 the height of the cone, the deposit is slightly thinner in comparison to the case
 401 without a magnetic field. This suggests that during the deposition period of 10 s,
 402 the counteraction between the opposing secondary flow and the buoyant flow may
 403 have briefly led to a thicker boundary layer in the region slightly below the cone tip.
 404 For a more detailed discussion we refer to section 3.2 and also to Fig. S1 in SI.

405 For the case of the iron cone in the magnetic field, stronger support for cone
 406 growth is found compared to the case of copper. As can be seen in Fig. 6, the
 407 deposit thickness in the upper half of the cone is considerably increased and almost
 408 doubled at the tip compared to the case without a magnetic field. This results
 409 from a stronger downward flow caused by the additional action of the magnetic
 410 gradient force $\mathbf{f}_{\nabla B}$ compared to the case of copper. The local minimum of the
 411 deposit thickness found at about $r/r_{\text{cone}} = 0.5$ is related to the detachment of the

concentration boundary layer in a jet-like flow, as shown in Fig. 5(b).

As a means of further assessing the impact of the different volume forces on the deposition process, the ratios of the magnetic forces to the buoyancy force are introduced as follows [33]:

$$R_{\text{MHD}} = \frac{\int_V |\mathbf{f}_{\text{L}}| dV}{\int_V |\mathbf{f}_{\text{g}}| dV}, \quad R_{\nabla\text{B}} = \frac{\int_V |\mathbf{f}_{\nabla\text{B}}| dV}{\int_V |\mathbf{f}_{\text{g}}| dV} \quad (20)$$

Here, the integration over volume V is restricted to the main flow region ($0 < r < 2.5 d_{\text{cone}}, 0 < z < 2.5 d_{\text{cone}}, 0 < \theta < 2\pi$). The temporal behavior of both quantities during the deposition process in the magnetic field is presented in Fig. 7.

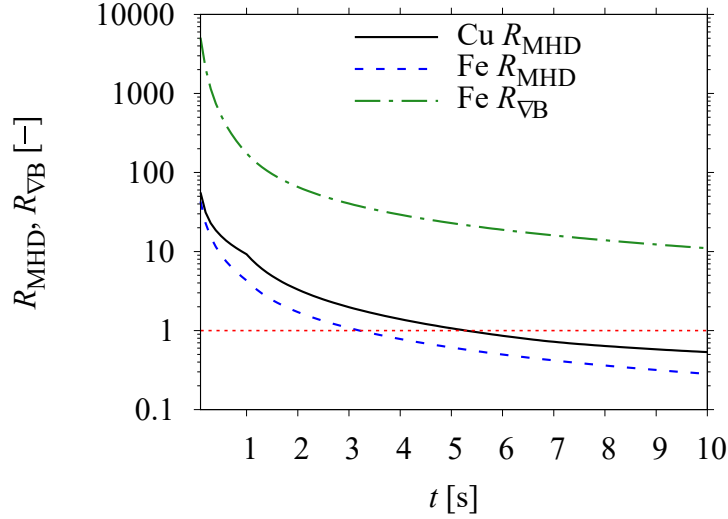


Figure 7: Temporal evolution of the force ratios R_{MHD} and $R_{\nabla\text{B}}$ defined in Eq. (20) for deposition on single copper and iron cones ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$). The vertical axis is shown on a logarithmic scale for better visibility, the red dashed line indicates the critical value of the force ratio of 1.

418

For both copper and iron cones, R_{MHD} shows a strong decrease with time. This is caused by the different temporal behavior of \mathbf{f}_{L} and \mathbf{f}_{g} . As a galvanostatic process is considered, the current density distribution can be expected to change only slowly with ongoing deposition [1]. Therefore, \mathbf{f}_{L} is nearly constant, whereas \mathbf{f}_{g} grows considerably with time due to the development of the concentration boundary layer. In the case of copper, R_{MHD} sinks below the critical value of unity at $t \approx 5 \text{ s}$. This is in line with the above discussion of the corresponding electrolyte flow pattern in Fig. 4. In comparison, in the case of iron R_{MHD} is slightly lower and already drops below unity at $t \approx 3 \text{ s}$. This is in accordance with the primary flow shown in Fig. 3, which is only forced in a comparably smaller region, and therefore also results in a correspondingly weaker secondary flow.

The ratio of the magnetic gradient force to the buoyancy force $R_{\nabla\text{B}}$ for the iron cone also shrinks with the deposition time. However, the values are much larger than in the two cases considered before. At the final instant considered, $t = 10 \text{ s}$, $R_{\text{MHD}} \approx 10$. This indicates that $\mathbf{f}_{\nabla\text{B}}$ has a much stronger influence on the process

433

than \mathbf{f}_L and also effectively counteracts the influence of \mathbf{f}_g , which is in line with the discussion of the flow pattern shown in Fig. 4. It should be noted that the region where \mathbf{f}_g acts is extended with the buildup and the advection of the concentration boundary layer, while $\mathbf{f}_{\nabla B}$ is only of importance close to the cone where large field gradients are found. This easily explains the decrease in $R_{\nabla B}$ with the deposition time.

3.2. Flat and sharp cones

To enable conclusions to be drawn on how a vertical magnetic field affects the growth of surface elevations at the different stages of development, we now include flat and steep cones in the investigation. The cone tip angles additionally considered are 30° and 90° , with the surface area of the cones remaining constant. More details of the cone dimensions are given in Table 3.

The shape of the cone can be expected to influence the distribution of the current density near the cone and also the magnetic field near the iron cone. We first investigate the influence of the current density. For geometrical reasons, steep cones are expected to be characterized by a less uniform current density distribution along the cone surface compared to flat cones [12]. This is also confirmed by the analytical solution of the primary current density distribution along an infinitely extended cone as derived in SI. Fig. 8(a) compares the numerically obtained primary current density distribution along the surface of the cones with different tip angles studied here, exhibiting good qualitative agreement with the analytical solution, as shown in Fig. S3 in SI.

In order to additionally account for kinetic effects at the electrode, which might mitigate the purely geometric effect considered in the primary current density [49, 50], Fig. 8(b) shows the numerical results of the current density at cones with different tip angles at $t = 10$ s. In these calculations, the electrode kinetics according to Eq. 13 are included, but the electrolyte flow is excluded. Although the inhomogeneity in general is reduced compared to the primary current density shown in Fig. 8(a), the numerical results indicate that the steep cone still has the largest value for the current density near the tip of the cone. This indicates a stronger Lorentz force effect for steeper cones.

A variation in the shape of the iron cones also affects the distribution of the magnetic field nearby. Fig. 9 shows the magnitudes of B and $B\nabla B$ along a nearby surface-parallel line for cones of different tip angles. In the vicinity of the cone tip, as already mentioned above, the sharp edge of the magnetized surface generally leads to a high magnetic flux density [40]. This implies that when the cone becomes sharper, a stronger stray field and correspondingly a stronger field gradient is found near its tip. However, for steeper cones, the cone surface grows more parallel to the external vertical field. This reduces the demagnetization of the Fe cones, which also reduces the stray field and thus the magnetic gradient near the cones. For more

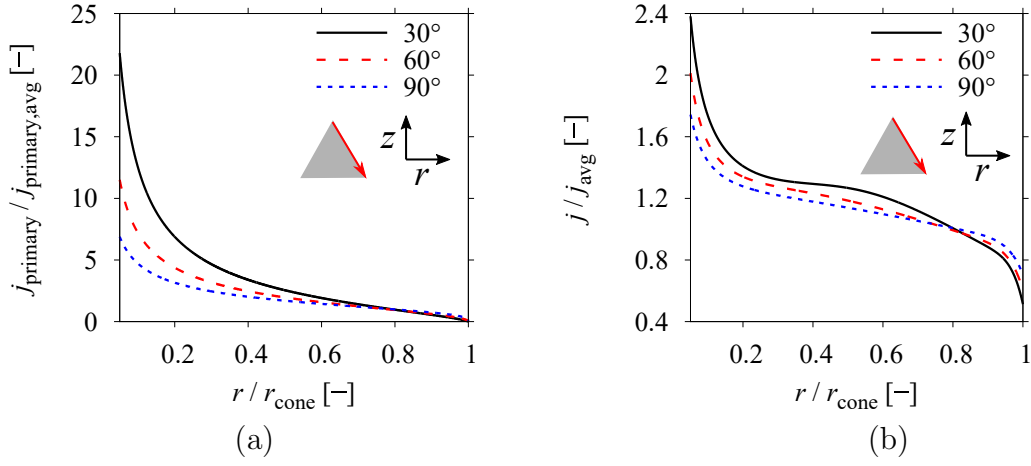


Figure 8: (a) Normalized primary current density and (b) normalized current density (No magnetic field, no convection) at cones of different tip angles α_{tip} after 10 s of deposition. $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$. The horizontal axis shows the radial surface coordinate normalized by the cone radius r_{cone} .

474 details on the magnetization of ferromagnetic surfaces in external magnetic fields,
 475 we refer to [40]. Due to the two opposite effects mentioned, the intermediate cone tip
 476 angle of 60° presented in Fig. 9 yields the strongest amplitudes of B and $B\nabla B$ near
 477 the tip. The strongest influence of the magnetic gradient force on the deposition
 might further be expected to be found at intermediate cone tip angles.

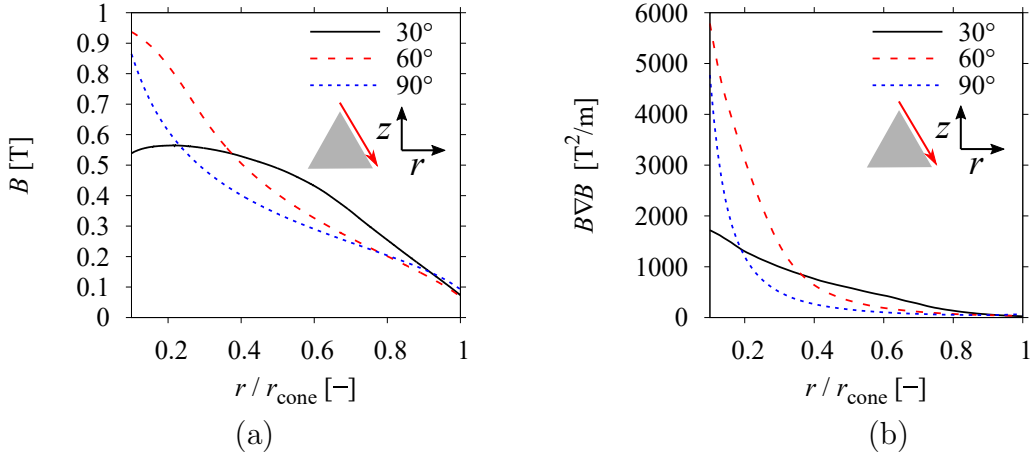


Figure 9: Magnitudes of (a) the magnetic flux density B and (b) the magnetic gradient term $B\nabla B$ along a monitoring line parallel and near (distance $35 \mu\text{m}$) to the slanted surface of Fe cones of different tip angles. $B_0 = 200 \text{ mT}$. The horizontal axis shows the radial surface coordinate normalized by the cone radius r_{cone} .

478
 479 Eventually, the simulation of the deposition process at copper and iron cones
 480 of different shapes, Fig. 10, shows the distribution of the copper ion species and
 481 the meridional velocity vectors obtained after a deposition time of 10 s. For the
 482 copper cones, the downward secondary flow driven by \mathbf{f}_{L} is strongest for the steep
 483 cone of $\alpha_{\text{tip}} = 30^\circ$ and weakest for the flat cone of $\alpha_{\text{tip}} = 90^\circ$. As the magnetic field
 484 remains unchanged, this is in line with the behavior of the current densities shown in
 485 Fig. 8. Furthermore, although not shown in detail, a larger region with horizontal
 486 components of the current density exists near steeper cones, as the vectors must

487 become normal to the cone surface. All this compensates for the closer proximity
 488 of the cone surface to the symmetry axis, which reduces the azimuthal momentum
 489 delivered by \mathbf{f}_L . For the steep cone, the concentration boundary layer rising from
 490 below is forced to separate from the cone surface slightly below the tip due to the
 491 strong downward flow, locally reducing the thickness of the boundary layer. The
 492 flatter the cone becomes, the weaker the downward flow caused by \mathbf{f}_L , while the
 493 concentration boundary layer passes the cone tip and continues to rise freely. This
 494 leads to a thicker concentration boundary layer near the tip of the cone, as observed
 495 also experimentally for an intermediate tip angle in [1]. The plume-like shape of the
 496 tip of the separated concentration boundary layer is similar to the behavior of the
 temperature in free thermal convection [51].

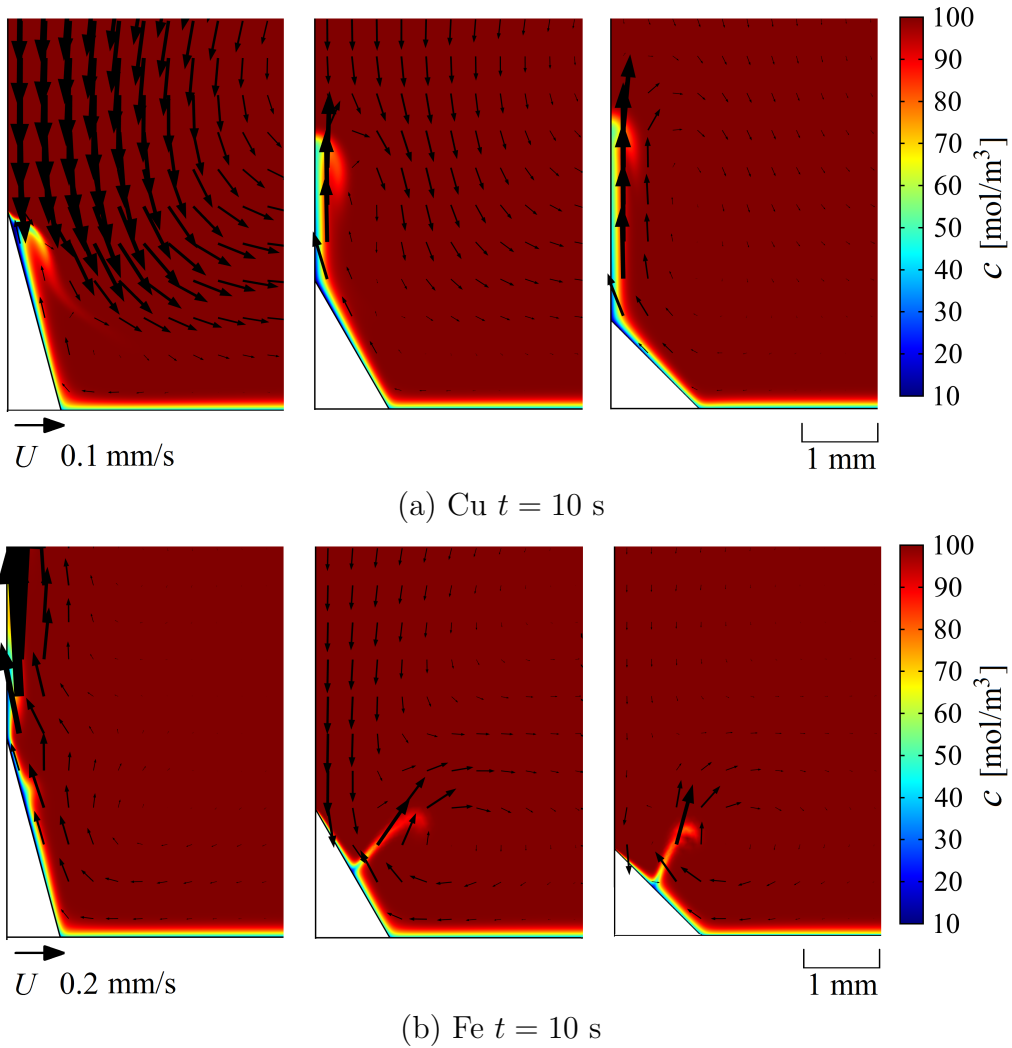


Figure 10: Color surface of the concentration near (a) Cu and (b) Fe cones after 10 s deposition time ($j_{\text{cathode,avg}} = 16$ mA/cm², $B_0 = 200$ mT). Black arrows represent the meridional velocity vectors. Cone tip angles from left to right: 30°, 60°, 90°.

497

498 For the steep iron cone, unlike the case of copper, the concentration boundary
 499 layer quickly rises vertically; its tip has already crossed the upper boundary of
 500 the region shown. This can be understood from Fig. 3, which shows a smaller
 501 region of azimuthal flow driven by \mathbf{f}_L compared to a copper cone. This results in

502 a correspondingly weaker secondary downward flow. The magnetic gradient force is
 503 also weakest for steep cones, as can be inferred from Fig. 9. Thus, the magnetic forces
 504 here are not strong enough to generate a downward flow. However, the interplay
 505 with buoyancy results in slight variations in the thickness of the rising boundary
 506 layer. For flatter iron cones, a downward flow towards the cone can be observed,
 507 which is stronger at $\alpha_{\text{tip}} = 60^\circ$ compared to $\alpha_{\text{tip}} = 90^\circ$, in full agreement with the
 508 discussion on the magnetic gradient term shown in Fig. 9. Compared to the copper
 509 cones discussed above, it can be concluded that this flow is mainly forced by $\mathbf{f}_{\nabla\mathbf{B}}$. As
 510 already discussed above, for $\alpha_{\text{tip}} = 60^\circ$ the concentration boundary layer separates
 511 from the cone at half its height. This is also observed in the case of the flat cone,
 512 where the weaker influence of $\mathbf{f}_{\nabla\mathbf{B}}$ is balanced by the comparably weaker influence
 513 of \mathbf{f}_g .

514 In order to evaluate the cumulative influence of the magnetic forces on the de-
 515 position, in Fig. 11 we compare the deposit thickness obtained after a deposition
 516 period of 10 s at the copper and iron cones of different shapes. In the absence of a
 517 magnetic field, as discussed above (see Fig. 8), at steep cones the natural support for
 518 further cone growth is most pronounced. When a magnetic field is applied, at steep
 519 cones the Lorentz force considerably enhances growth near the tip of the copper
 520 cone, whereas at the iron cone the support of the magnetic field for cone growth is
 521 comparably small. The intermediate tip angle of 60° was already discussed above,
 522 see Fig. 6. For the flat cone, the situation is opposite to the steep cone. The in-
 523 fluence of \mathbf{f}_L at the copper cone is negligible, whereas a profound enhancement of
 cone growth at the iron cone can be reported, mainly due to $\mathbf{f}_{\nabla\mathbf{B}}$.

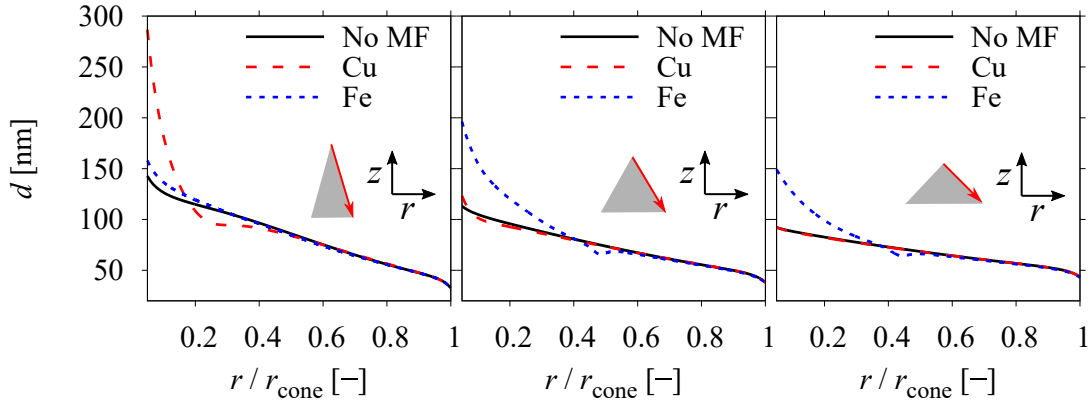


Figure 11: Deposit thickness on cones of tip angles 30° (left), 60° (middle) and 90° (right) after 10 s of deposition for the case without a magnetic field (no MF) and for Cu and Fe cones with a magnetic field ($j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$). The horizontal axis shows the radial surface coordinate normalized by the cone radius r_{cone} . The case of 60° is reprinted from Fig. 6 for comparison.

524
 525 Table 4 summarizes the relative enhancement of the deposit thickness at the
 526 tip of the Cu and Fe cones compared to the case without a magnetic field after a
 527 deposition time of 10 s. It can be concluded that cone growth at diamagnetic cones
 528 supported by the action of \mathbf{f}_L works best for cones of a sharp shape. However, as

the support at flat cones seems to be negligible, the benefit of magnetic fields for the further development of early flat surface elevations seems rather limited.

For ferromagnetic cones, a promising enhancement of the deposit thickness was found for all the cone shapes studied here, mainly caused by the magnetic gradient force. The supporting effect was found to be the greatest with intermediate tip angles and to be moderate only in the case of steep cones. As can be seen from Fig. 11, when starting from a flat surface elevation, cone growth accelerates during deposition as the cone tip angle becomes smaller. The cone continues to grow and to become sharper. However, as soon as the optimum intermediate tip angle associated with the strongest support from the magnetic gradient force is passed, this sharpening slows down.

Table 4: Enhancement of the deposit thickness at the Cu and Fe cone tips compared to the case without a magnetic field after 10 s deposition time ($\alpha_{\text{tip}} = 30^\circ, 60^\circ, 90^\circ$, $j_{\text{cathode,avg}} = 16 \text{ mA/cm}^2$, $B_0 = 200 \text{ mT}$).

| Cone tip angle | Cu | Fe |
|----------------|-------------|------|
| 30° | +93% | +10% |
| 60° | +9% | +73% |
| 90° | ≈ 0 | +62% |

3.3. Influence of neighboring cones

Depending on the distance between adjacent cones, the flow forced at each single cone may be affected by the flow originating from its neighbors. Thus, mass transfer might also change. We therefore now extend our investigation to take into consideration the influence of neighboring cones. We first consider regular arrangements of cones on a quadratic lattice as shown in Fig. 2(c). These investigations correspond to Step 2 of the simulation methodology mentioned earlier.

As a result of the 3D simulations performed, Fig. 12 shows the horizontal components of the electrolyte velocity in a horizontal plane above the cathode touching the tips of copper cones. A square with four neighboring cones is shown, and the distance between the cone centers is varied from 5 to 1.5 cone diameters d_{cone} (see Fig. 2). At the large cone distance, a strong azimuthal flow in an anticlockwise direction is clearly visible, driven by \mathbf{f}_L . As known from the single cone studies, this flow extends in a radial direction far beyond the cone radius (see Fig. 3). In the outer region close to half the cone distance, the neighbor influence becomes visible as a deformation of the circular shape of the azimuthal velocity contours towards a square. At half of the distance between the cones, the horizontal velocity is considerably reduced, as the two azimuthal flows are in opposite directions and hamper each other. When the distance between the cones is reduced, this damping influence increases, as seen from the lower amplitude of the azimuthal flow. At the same time, small clockwise-rotating vortices are formed at the four vertical edges of the

561 computational domain (most clearly visible in the center of the squares shown in
 562 Fig. 12), the amplitude of which is weak in comparison to the primary flow.

563 Fig. 13 shows the corresponding vertical velocity components in the vertical
 564 plane across the center of the copper cone. Because of symmetry, the results in x
 565 or y direction are identical. As the primary rotational flow is slowed down when
 566 the cones grow closer to each other, the secondary downward flow forced by \mathbf{f}_L also
 567 gets correspondingly weaker and has nearly disappeared at the small cone distance
 568 of $1.5 d_{\text{cone}}$. Thus, the support of cone growth in a magnetic field is weakened as the
 area density of the cones grows.

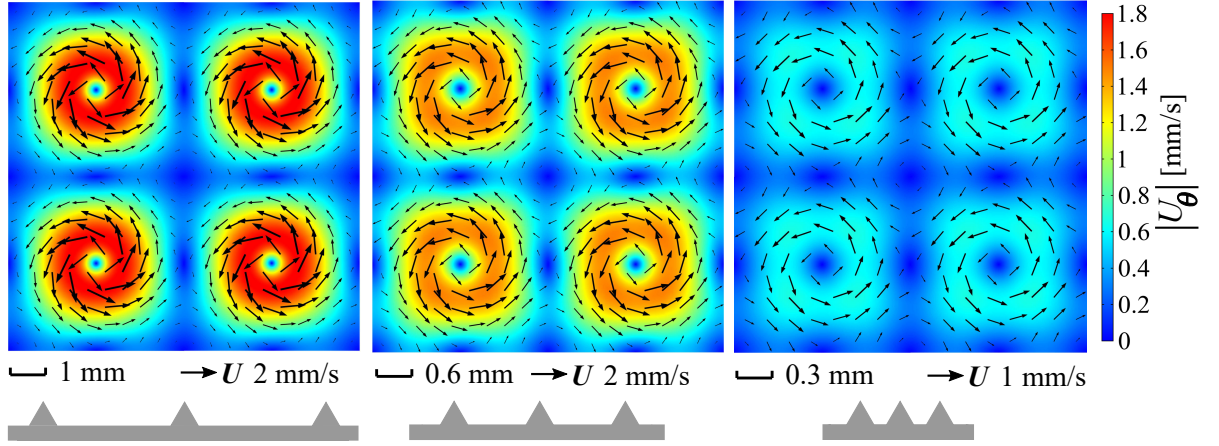


Figure 12: Horizontal velocity in a horizontal plane touching the tips of the neighboring copper cones obtained from 3D simulations after a deposition time of 5 s ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 8$ mA/cm², $B_0 = 400$ mT). Black arrows of the horizontal velocity vectors and color contours of the azimuthal velocity component. The cone distance in units of d_{cone} shrinks from left to right (5, 3, 1.5). Note that the scale of length and of the velocity vectors changes from left to right.

569

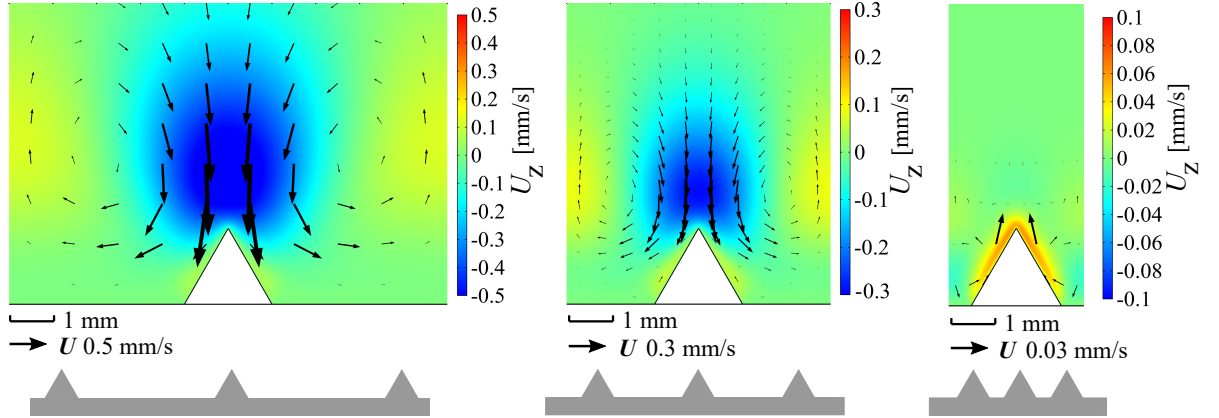


Figure 13: Vertical velocity on the vertical plane across the center of the copper cone in x direction obtained from 3D simulations after a deposition time of 5 s ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 8$ mA/cm², $B_0 = 400$ mT). The cone distance in units of d_{cone} shrinks from left to right (5, 3, 1.5). Black arrows represent the velocity vectors, the scale of which changes from left to right.

570 It should be mentioned that the computational effort for these 3D simulations
 571 is great and would be even greater for the iron cones, where steep gradients in the
 572 magnetic field near the surface of the cones additionally need to be accurately re-
 573 solved. A validation of the axisymmetric 2D approach denoted as Step 3 in Section

2.1 could allow the computational effort for these simulations to be reduced considerably. This is motivated by the persistence of strong axisymmetry at a shrinking cone distance, as shown in Fig. 12. We therefore next present a comparison of results obtained by the 3D method and by the axisymmetric 2D approach for Cu cones. Fig. 14(a) shows the magnitude of the maximum azimuthal velocity of the primary flow versus the cone distance after a deposition time of 5 s. The results of the 3D and 2D approaches for the Cu case are close and follow the trend mentioned above of the azimuthal rotational flow becoming weaker if the cone distance is reduced. Fig. 14(b) shows the vertical velocity measured slightly above the cone tip. Again, both 3D and 2D results in the case of Cu are close and show that the downward flow velocity weakens. In both cases, the 2D axisymmetric results slightly overstate the damping influence compared to the 3D results, which is reasonable, as immediately neighboring cones are assumed to be found at every angular position compared to only four immediately neighboring cones in the 3D case. Nevertheless, the axisymmetric 2D approach proves to be an accurate and effective means of studying the neighbor influence. The intrinsic slight overestimation of the neighbor influence allows upper boundaries for the damping neighbor effect to be found quickly. All results presented in the following, including the cases of the iron cones, are therefore obtained by 2D simulations.

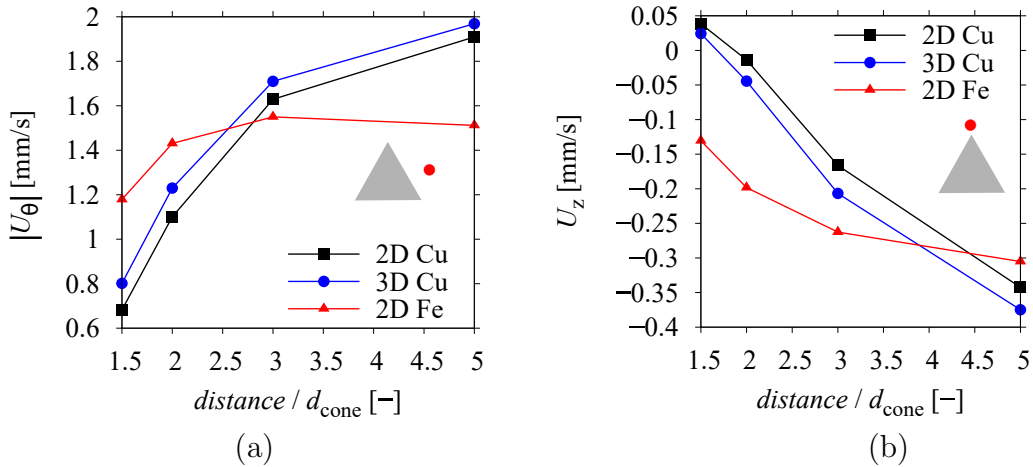


Figure 14: (a) Magnitude of the maximal azimuthal velocity, $|U_\theta|$, and (b) the vertical velocity U_z at 0.3 mm above the cone tip versus cone distance for Cu (2D and 3D results) and Fe cones (2D results) after 5 s of deposition ($\alpha_{tip} = 60^\circ$, $j_{cathode,avg} = 8 \text{ mA/cm}^2$, $B_0 = 400 \text{ mT}$).

For the iron cones, unlike the copper cones, varying the distance between the cones can change their magnetization and thus the magnetic field. Fig. 15 shows the magnitudes of B and $B\nabla B$ near the cone surface for different cone distances. When the neighboring cones grow closer, the magnetic field near a magnetized Fe cone becomes slightly weaker. Regarding the magnetic field gradient term, reducing the distance to the neighbor cones has a small influence near the lower half of the iron cone only.

As shown in Fig. 14, shrinking the cone distance has a much weaker effect on the

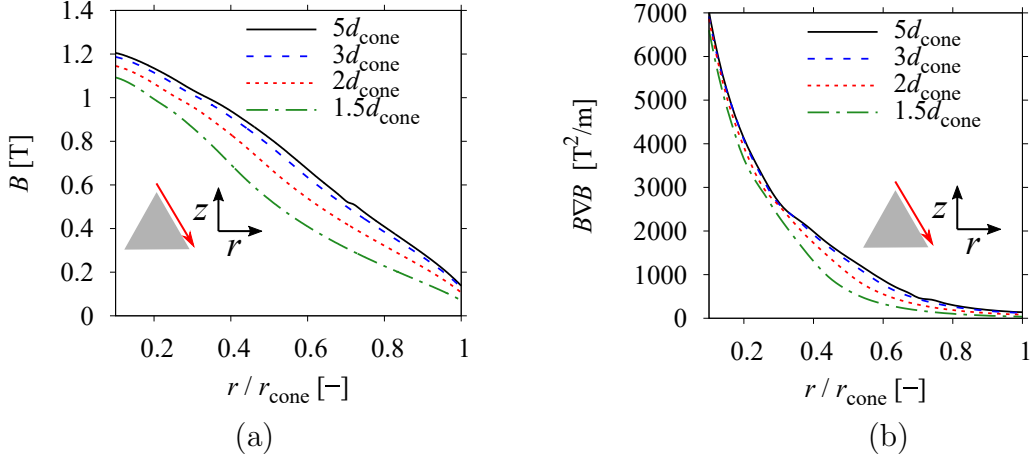


Figure 15: Magnitudes of (a) the magnetic flux density, B , and (b) the magnetic gradient term, $B\nabla B$, along a monitoring line parallel to the slanted cone surface (distance to the cone surface: $35\ \mu\text{m}$) for different distances between the Fe cones ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 8\ \text{mA}/\text{cm}^2$, $B_0 = 400\ \text{mT}$). The horizontal axis shows the radial surface coordinate normalized by the cone radius r_{cone} .

electrolyte flow near the iron cones compared to the copper cones. The maximum azimuthal velocity shown on the left (a) remains nearly constant and eventually decreases only slightly towards a distance of $1.5\ d_{\text{cone}}$. The comparably weak neighbor effects on the azimuthal flow are related to the bending of the magnetic field near the iron cones. More details are given in SI. The vertical velocity shown on the right (b) and depicted slightly above the cone tip, contrary to the Cu cones, remains negative even at the shortest distance of $1.5\ d_{\text{cone}}$ shown. As the cone distance shrinks, this downward flow is also weakened due to the continuity of the flow, as the vertical backflow in the narrow gap between the cones is in the opposite direction. But the slowing of this flow is moderate only. The obvious reason is that the downward flow caused by $\mathbf{f}_{\nabla B}$ is driven near the cones. This is unlike the downward flow caused by $\mathbf{f}_{\mathbf{L}}$, which is a result of the primary azimuthal flow driven in a radially extended region. Thus, the supporting influence of $\mathbf{f}_{\nabla B}$ for cone growth can be expected to be less affected by neighboring cones compared to $\mathbf{f}_{\mathbf{L}}$.

Fig. 16(a) shows the concentration and meridional flow for the Cu case at a later deposition time (13 s), at which the downward secondary flow caused by $\mathbf{f}_{\mathbf{L}}$ is only visible at the largest cone distance. As the distance between the cones shrinks and the secondary flow of the Lorentz force becomes weaker, the buoyant upward convection of the concentration boundary layer is enhanced. The only exception is that, as the cone distance decreases from $2\ d_{\text{cone}}$ to $1.5\ d_{\text{cone}}$, the buoyant flow seems to be slightly weaker again, as seen from the height of the buoyancy plume. This might be due to stronger damping between the upward buoyant flow and the downward backflow, as the radial distance between these two flow regions also shrinks with the cone distance.

In the case of the iron cones, as shown in Fig. 16(b), unlike the Cu cones, the downward flow resulting from the two magnetic forces seems to be only slightly weakened as the cone distance shrinks. Therefore, support for cone growth by the

628 magnetic field can be expected in all cases. The weakening of the downward flow
 629 results in a slight increase in the height at which the concentration boundary layer
 630 leaves the cone, as discussed above in the case of single iron cones. In all cases,
 631 the concentration boundary layer at the Fe cones exhibits further periodic thickness
 632 variations along the cone at a length scale much smaller than the cone diameter.
 633 These are mainly caused by the action of the magnetic gradient force $\mathbf{f}_{\nabla\mathbf{B}}$. The
 634 position of the departing jet of depleted electrolyte can also be understood as the
 most unstable location of the boundary layer [38, 52].

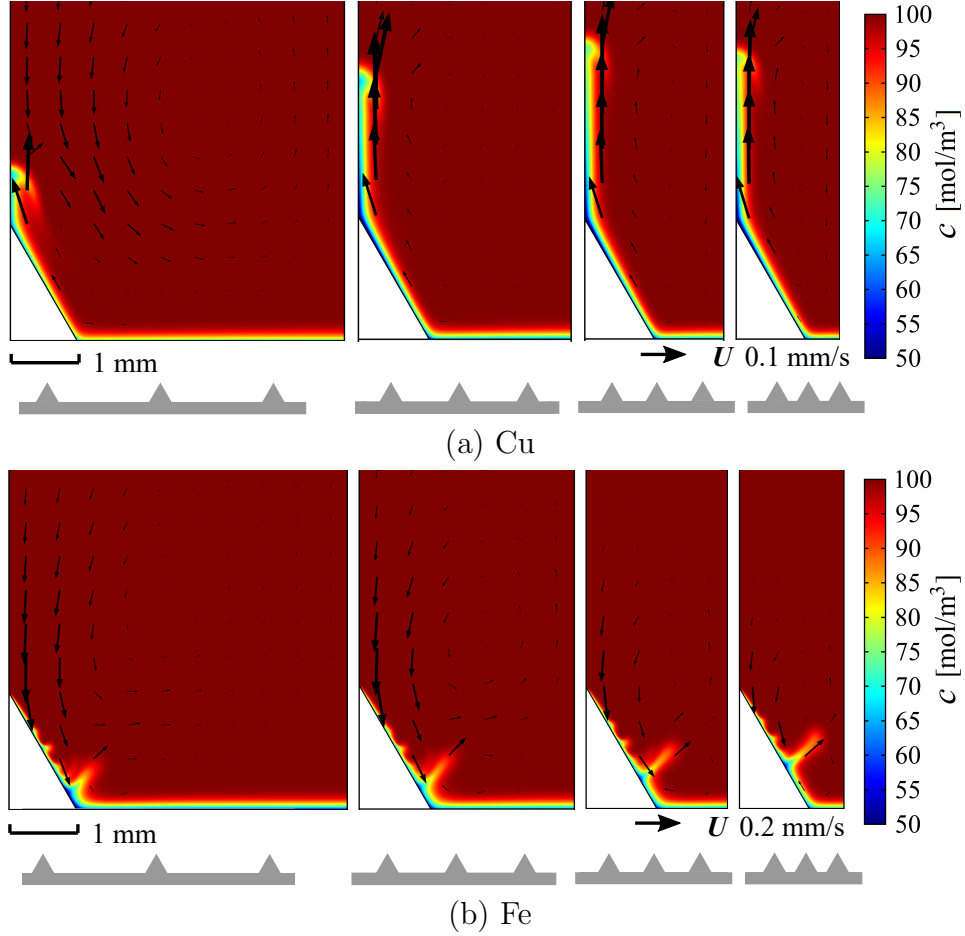


Figure 16: Species concentration (color surface) and meridional velocity vectors (black arrows) for
 (a) Cu and (b) Fe cones at different distances after 13 s of deposition. ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 8$
 mA/cm², $B_0 = 400$ mT). The cone distance in unit of d_{cone} shrinks from left to right (5, 3, 2, 1.5).
 Note that the scales of the velocity vectors for Cu and Fe cones are different.

635

636 Finally, Fig. 17 shows the deposit thickness obtained for the smallest cone dis-
 637 tance of $1.5 d_{\text{cone}}$ investigated. A longer deposition time of 50 s is also considered.
 638 As can be seen, for the Cu cones, the structuring effect due to $\mathbf{f}_{\mathbf{L}}$ is negligible when
 639 compared to the case without a magnetic field, which is in accordance with the
 640 buoyancy-dominated flow pattern shown in Fig. 16(a). For the iron cones, however,
 641 the support for cone growth by the magnetic field, i.e. mainly by $\mathbf{f}_{\nabla\mathbf{B}}$, is clearly
 642 visible.

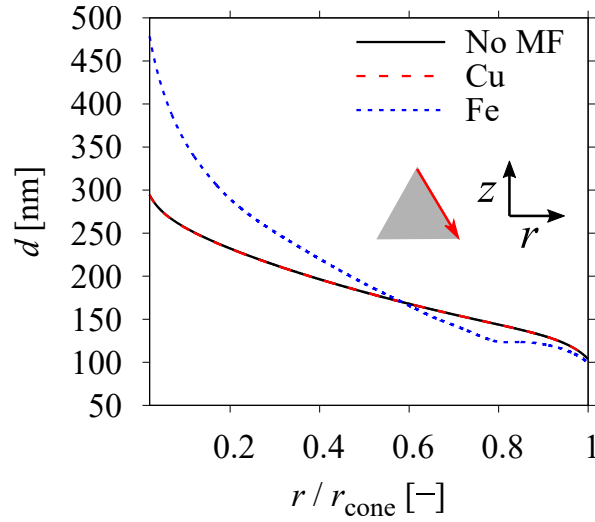


Figure 17: Deposit thickness along the cone surface for the case without a magnetic field (No MF) and for Cu and Fe cones in a vertical magnetic field after 50 s of deposition ($\alpha_{\text{tip}} = 60^\circ$, $j_{\text{cathode,avg}} = 8 \text{ mA/cm}^2$, $B_0 = 400 \text{ mT}$). The distance to neighboring cones is $1.5 d_{\text{cone}}$, the horizontal axis shows the radial surface coordinate normalized by the cone radius r_{cone} .

643 4. Conclusions

644 The analytical and numerical results of this work offer an insight into the basic
 645 effects of a vertical magnetic field on electrodeposition at conically shaped metal
 646 electrodes. The focus is on ferromagnetic conical structures of mm size. We show
 647 that cone growth can be supported by the flow driven by the Lorentz force and
 648 the magnetic gradient force that enriches the electrolyte near the cone, and thus
 649 enhances the local mass transfer.

650 As the cathode is assumed to be placed at the bottom of the electrochemical
 651 cell, the beneficial magnetic effects are counteracted by solutal buoyancy arising
 652 from the electrode reaction. The Lorentz force is surpassed by the buoyancy force
 653 after the first few seconds of the deposition. In comparison, the magnetic gradient
 654 force enabled by the magnetization of the ferromagnetic cones provides stronger
 655 support for cone growth, thereby often dominating over buoyancy. In general, the
 656 flow is caused by the rotational parts of both magnetic gradient force and buoyancy
 657 force.

658 We studied cones with different tip angles and found that the sharpest cone
 659 yields the strongest Lorentz force, while an intermediate cone tip angle generates
 660 the highest magnetic gradient force. As the evolution of the conical structures
 661 usually develops from flat surface elevations, it could be anticipated that the growth
 662 of ferromagnetic cones may be slow at the beginning, followed by an acceleration
 663 of the growth speed until an optimum shape is approached with respect to the
 664 supporting effect. Later on, growth will continue at a lower speed.

665 When the neighboring cones come closer to each other, the azimuthal flow caused
 666 by the Lorentz force is strongly damped. In comparison, the meridional flow caused
 667 by the magnetic gradient force is less affected, as it is driven in the close vicinity of

668 the cone surface.

669 Our results demonstrate the superiority of the magnetic gradient force in terms
670 of supporting the structured electrodeposition in a magnetic field. We expect this
671 superiority to enhance further when extending the investigation towards the micro-
672 and nanometer scale, where larger field gradients are to be expected. This may en-
673 courage further research on magnetic field assisted electrodeposition as a simple and
674 efficient method for synthesizing micro- and nano-structured ferromagnetic surfaces.

675 **Acknowledgments**

676 We would like to thank Xuegeng Yang, Margitta Uhlemann, Piotr Zabinski
677 and Katarzyna Skibinska for fruitful discussions. Financial support by Deutsche
678 Forschungsgemeinschaft, grant no. 381712986 is gratefully acknowledged.

679 References

- 680 [1] M. Huang, G. Marinaro, X. Yang, B. Fritzsche, Z. Lei, M. Uhlemann, K. Eckert,
681 G. Mutschke, Mass transfer and electrolyte flow during electrodeposition on a
682 conically shaped electrode under the influence of a magnetic field, *Journal of*
683 *Electroanalytical Chemistry* 842 (2019) 203–213.
- 684 [2] T. Hang, A. Hu, H. Ling, M. Li, D. Mao, Super-hydrophobic nickel films with
685 micro-nano hierarchical structure prepared by electrodeposition, *Applied Sur-*
686 *face Science* 256 (2010) 2400–2404.
- 687 [3] T. Mouterde, G. Lehoucq, S. Xavier, A. Checco, C. T. Black, A. Rahman,
688 T. Midavaine, C. Clanet, D. Quéré, Antifogging abilities of model nanotextures,
689 *Nature materials* 16 (2017) 658–663.
- 690 [4] H. Zhang, S. Hwang, M. Wang, Z. Feng, S. Karakalos, L. Luo, Z. Qiao, X. Xie,
691 C. Wang, D. Su, et al., Single atomic iron catalysts for oxygen reduction in
692 acidic media: particle size control and thermal activation, *Journal of the Amer-*
693 *ican Chemical Society* 139 (40) (2017) 14143–14149.
- 694 [5] L. Liu, A. Corma, Metal catalysts for heterogeneous catalysis: from single
695 atoms to nanoclusters and nanoparticles, *Chemical reviews* 118 (10) (2018)
696 4981–5079.
- 697 [6] G. B. Darband, M. Aliofkhazraei, S. Shanmugam, Recent advances in methods
698 and technologies for enhancing bubble detachment during electrochemical water
699 splitting, *Renewable and Sustainable Energy Reviews* 114 (2019) 109300.
- 700 [7] N. Wang, T. Hang, D. Chu, M. Li, Three-dimensional hierarchical nanostruc-
701 tured Cu/Ni–Co coating electrode for hydrogen evolution reaction in alkaline
702 media, *Nano-micro letters* 7 (2015) 347–352.
- 703 [8] G. B. Darband, M. Aliofkhazraei, S. Hyun, A. S. Rouhaghdam, S. Shan-
704 mugam, Electrodeposition of Ni–Co–Fe mixed sulfide ultrathin nanosheets on
705 ni nanocones: a low-cost, durable and high performance catalyst for electro-
706 chemical water splitting, *Nanoscale* 11 (35) (2019) 16621–16634.
- 707 [9] S. Arai, M. Ozawa, M. Shimizu, Communication micro-scale columnar archi-
708 tecture composed of copper nano sheets by electrodeposition technique, *Journal*
709 *of The Electrochemical Society* 164 (2) (2016) D72–D74.
- 710 [10] G. B. Darband, M. Aliofkhazraei, A. S. Rouhaghdam, Three-dimensional
711 porous Ni-CNT composite nanocones as high performance electrocatalysts for
712 hydrogen evolution reaction, *Journal of Electroanalytical Chemistry* 829 (2018)
713 194–207.

- [11] A. K. Singh, D. Sarkar, Enhanced light absorption and charge carrier management in core-shell Fe_2O_3 @Nickel nanocone photoanodes for photoelectrochemical water splitting, *ChemCatChem* 11 (24) (2019) 6355–6363.
- [12] K. Aoki, Theory of current distribution at a conical electrode under diffusion control with time dependence, *Journal of electroanalytical chemistry and interfacial electrochemistry* 281 (1990) 29–40.
- [13] R. Aogaki, K. Fueki, T. Mukaibo, Application of magnetohydrodynamic effect to the analysis of electrochemical reactions 1. mhd flow of an electrolyte solution in an electrode-cell with a short rectangular channel, *Denki Kagaku oyobi Kogyo Butsuri Kagaku* 43 (9) (1975) 504–508.
- [14] A. Bund, S. Koehler, H. Kuehnlein, W. Plieth, Magnetic field effects in electrochemical reactions, *Electrochimica Acta* 49 (1) (2003) 147–152.
- [15] G. Mutschke, A. Hess, A. Bund, J. Fröhlich, On the origin of horizontal counter-rotating electrolyte flow during copper magnetoelectrolysis, *Electrochimica Acta* 55 (5) (2010) 1543–1547.
- [16] Y. Zhang, B. Yuan, L. Li, C. Wang, Edge electrodeposition effect of cobalt under an external magnetic field, *Journal of Electroanalytical Chemistry* (2020) 114143.
- [17] S. Mühlenhoff, G. Mutschke, M. Uhlemann, X. Yang, S. Odenbach, J. Fröhlich, K. Eckert, On the homogenization of the thickness of cu deposits by means of mhd convection within small dimension cells, *Electrochemistry Communications* 36 (2013) 80–83.
- [18] M. Najminoori, A. Mohebbi, K. Afrooz, B. G. Arabi, The effect of magnetic field and operating parameters on cathodic copper winning in electrowinning process, *Chemical Engineering Science* 199 (2019) 1–19.
- [19] K. Nishikawa, T. Saito, H. Matsushima, M. Ueda, Holographic interferometric microscopy for measuring Cu^{2+} concentration profile during cu electrodeposition in a magnetic field, *Electrochimica Acta* 297 (2019) 1104–1108.
- [20] H. A. Murdoch, D. Yin, E. Hernández-Rivera, A. K. Giri, Effect of applied magnetic field on microstructure of electrodeposited copper, *Electrochemistry Communications* 97 (2018) 11–15.
- [21] P. Żabiński, K. Mech, R. Kowalik, Electrocatalytically active Co–W and Co–W–C alloys electrodeposited in a magnetic field, *Electrochimica Acta* 104 (2013) 542–548.

- [22] K. Shen, Z. Wang, X. Bi, Y. Ying, D. Zhang, C. Jin, G. Hou, H. Cao, L. Wu, G. Zheng, et al., Magnetic field-suppressed lithium dendrite growth for stable lithium-metal batteries, *Advanced Energy Materials* 9 (20) (2019) 1900260.
- [23] A. Wang, Q. Deng, L. Deng, X. Guan, J. Luo, Eliminating tip dendrite growth by Lorentz force for stable lithium metal anodes, *Advanced Functional Materials* 29 (25) (2019) 1902630.
- [24] J. A. Koza, M. Uhlemann, A. Gebert, L. Schultz, Desorption of hydrogen from the electrode surface under influence of an external magnetic field, *Electrochemistry communications* 10 (9) (2008) 1330–1333.
- [25] J. A. Koza, S. Mühlenhoff, P. Żabiński, P. A. Nikrityuk, K. Eckert, M. Uhlemann, A. Gebert, T. Weier, L. Schultz, S. Odenbach, Hydrogen evolution under the influence of a magnetic field, *Electrochimica Acta* 56 (6) (2011) 2665–2675.
- [26] D. Fernández, Z. Diao, P. Dunne, J. Coey, Influence of magnetic field on hydrogen reduction and co-reduction in the Cu/CuSO₄ system, *Electrochimica acta* 55 (28) (2010) 8664–8672.
- [27] H. Liu, Q. Hu, L. Pan, R. Wu, Y. Liu, D. Zhong, Electrode-normal magnetic field facilitating neighbouring electrochemical bubble release from hydrophobic islets, *Electrochimica Acta* 306 (2019) 350–359.
- [28] K. Tschulik, R. Sueptitz, J. Koza, M. Uhlemann, G. Mutschke, T. Weier, A. Gebert, L. Schultz, Studies on the patterning effect of copper deposits in magnetic gradient fields, *Electrochimica Acta* 56 (1) (2010) 297–304.
- [29] P. Dunne, J. Coey, Patterning metallic electrodeposits with magnet arrays, *Physical Review B* 85 (22) (2012) 224411.
- [30] G. Mutschke, A. Bund, On the 3D character of the magnetohydrodynamic effect during metal electrodeposition in cuboid cells, *Electrochemistry Communications* 10 (4) (2008) 597–601.
- [31] L. M. Monzon, J. M. D. Coey, Magnetic fields in electrochemistry: The Lorentz force. a mini-review, *Electrochemistry Communications* 42 (2014) 38–41.
- [32] J. Coey, F. Rhen, P. Dunne, S. McMurtry, The magnetic concentration gradient force - is it real?, *Journal of solid state electrochemistry* 11 (6) (2007) 711–717.
- [33] G. Mutschke, K. Tschulik, T. Weier, M. Uhlemann, A. Bund, J. Fröhlich, On the action of magnetic gradient forces in micro-structured copper deposition, *Electrochimica Acta* 55 (28) (2010) 9060–9066.

- [34] G. Mutschke, K. Tschulik, M. Uhlemann, A. Bund, J. Fröhlich, Comment on magnetic structuring of electrodeposits, *Physical review letters* 109 (2012) 229401.
- [35] M. Uhlemann, K. Tschulik, A. Gebert, G. Mutschke, J. Fröhlich, A. Bund, X. Yang, K. Eckert, Structured electrodeposition in magnetic gradient fields, *The European Physical Journal Special Topics* 220 (2013) 287–302.
- [36] H. Danan, A. Herr, A. Meyer, New determinations of the saturation magnetization of nickel and iron, *Journal of Applied Physics* 39 (2) (1968) 669–670.
- [37] J. Newman, K. E. Thomas-Alyea, *Electrochemical systems*, 3rd Edition, John Wiley & Sons, 2012.
- [38] A. Bejan, *Convection heat transfer*, 3rd Edition, John Wiley & Sons, 2013.
- [39] T. Weier, D. Baczyzmalski, J. Massing, S. Landgraf, C. Cierpka, The effect of a Lorentz-force-driven rotating flow on the detachment of gas bubbles from the electrode surface, *International Journal of Hydrogen Energy* 42 (33) (2017) 20923–20933.
- [40] J. D. Jackson, *Classical electrodynamics*, 3rd Edition, John Wiley and Sons, 1999.
- [41] COMSOL Multiphysics Documentation Suite V 5.5, COMSOL Inc., Burlington, MA 01803, USA (2019).
- [42] J. Coey, New permanent magnets; manganese compounds, *Journal of Physics: Condensed Matter* 26 (6) (2014) 064211.
- [43] D. Koschichow, G. Mutschke, X. Yang, A. Bund, J. Fröhlich, Numerical simulation of the onset of mass transfer and convection in copper electrolysis subjected to a magnetic field, *Russian Journal of Electrochemistry* 48 (7) (2012) 682–691.
- [44] C. Akilan, T. Chen, T. Vielma, P. M. May, G. Senanayake, G. Hefter, Volumes and heat capacities of cobalt(II), nickel(II), and copper(II) sulfates in aqueous solution, *Journal of Chemical & Engineering Data* 65 (9) (2020) 4575–4581.
- [45] P. Novotny, O. Sohnel, Densities of binary aqueous solutions of 306 inorganic substances, *Journal of Chemical and Engineering Data* 33 (1) (1988) 49–55.
- [46] D. R. Lide, *CRC handbook of chemistry and physics*, Vol. 85, CRC press, 2004.
- [47] X. Yang, S. Mühlenhoff, P. A. Nikrityuk, K. Eckert, The initial transient of natural convection during copper electrolysis in the presence of an opposing Lorentz force: current dependence, *The European Physical Journal Special Topics* 220 (1) (2013) 303–312.

- 815 [48] C. N. Singman, Atomic volume and allotropy of the elements, Journal of Chem-
816 ical Education 61 (2) (1984) 137–142.
- 817 [49] M. Matlosz, C. Creton, C. Clerc, D. Landolt, Secondary current distribution
818 in a hull cell boundary element and finite element simulation and experimental
819 verification, Journal of the Electrochemical Society 134 (12) (1987) 3015–3021.
- 820 [50] C. Low, E. Roberts, F. Walsh, Numerical simulation of the current, potential
821 and concentration distributions along the cathode of a rotating cylinder hull
822 cell, Electrochimica Acta 52 (11) (2007) 3831–3840.
- 823 [51] E. Sparrow, R. Husar, R. Goldstein, Observations and other characteristics of
824 thermals, Journal of Fluid Mechanics 41 (4) (1970) 793–800.
- 825 [52] R. Nelson, A. Bejan, Constructal optimization of internal flow geometry in
826 convection, Journal of heat transfer 120 (2) (1998) 357–364.