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Use of magnetic fields in electrochemistry: a selected review.

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Abstract:

Electrochemical reactions are usually thermally-activated and submitted to masstransfer effects. Although classically, enhanced kinetics of an electrochemical reaction is obtained by heating the cell and feeding the reactant by forced convection, other means can be used to improve mass- and charge-transfer. This paper shortly reviews the effects of magnetic fields in electrochemistry. Using a static or an alternating magnetic field enables to enhance electrodeposition and electrocatalysis, via improved gas and species convection, electrochemical kinetics and whole reaction efficiency. Such enhancement can mainly be related to Lorentz and Kelvin forces, magneto-hydrodynamics (MHD), chiral-induced spin selectivity (CISS) and hyperthermia, these effects being described herein.

Keyword: magnetic field; Lorentz force; Kelvin force; electrodeposition; charge-transfer; spindependent electrochemistry; electrocatalysis

Highlights:

- The Lorentz force induces convection (via MHD and micro-MHD effects).
- The Kelvin force induces convection in magnetic field gradient on para(dia)magnetic

species.

- The polarization of the spins influences reaction pathways.
- Hyperthermia, a local heating of the electrode materials, enhances reaction kinetics.

1 Introduction

Electrochemical reactions are usually thermally-activated and submitted to masstransfer effects. Although classically, enhanced kinetics of an electrochemical reaction is obtained by heating the cell and feeding the reactant by forced convection, other means can be used to improve mass- and charge-transfer. Although not trivial and beyond the usual electrochemists' expertise, magnetic fields are coupled with electrochemistry since 40 years. Researchers firstly observed its effect on mass-transfer, limiting current density and chargetransfer [1][2][3][4], and electrodeposition [5]. Many practically-interesting results were obtained in various domains, *e.g.* proton exchange membrane fuel cells (PEMFC) [6][7], electrodeposition [8], lab-on-chips [9] or enantioselectivity [10]. Whether the magnetic field is alternating (AMF) or static (SMF), and applied parallel ($B_{i/i}$) or perpendicular (B_{\perp}) to the electrode surface, several effects (forces) can be triggered, such as Lorentz and Kelvin forces [11,12]. Generally, these phenomena are concomitant, and isolating their individual influence is difficult; this selected-review aims to summarize and discuss the main results obtained recently on the theme.

2 Static Magnetic Field

2.1 The Lorentz force

The most common and earliest-studied effect [1] upon application of a SMF in electrochemistry is the Lorentz force density F_{L}^{1} (equation 1):

$$\boldsymbol{F}_{\mathrm{L}} = \boldsymbol{J} \times \boldsymbol{B} \tag{1}$$

where the current density of electrochemical process is (equation 2):

¹ In this review, vectors are written in **bold** fonts (*e.g.* vector **B**) and their modulus are in normal fonts ($||\mathbf{B}|| = B$).

$$\boldsymbol{J} = \boldsymbol{\sigma} \left(\boldsymbol{E} + \boldsymbol{v} \times \boldsymbol{B} \right) - n \, \boldsymbol{\mathcal{F}} D_{+,-} \nabla c_{+,-} \tag{2}$$

with E the electric field, σ the electrical conductivity, v the velocity of charged species and B the magnetic field. σ ($v \times B$) is the induced current, *n* the charge number of the electroactive ionic species $c_{+,-}$, $D_{+,-}$ its diffusion coefficient and \mathcal{F} the Faraday constant. The diffusion of ions in the electrolyte during electrolysis increases the concentration gradient, implying higher reaction overvoltage [13]. Expressing the diffusion current density in MHD investigation is uncommon [14], but considering this effect is nevertheless relevant. Under AMF exposure or with a gas-evolving electrode, non-static effects could involve diffusion current, which depend on local potential variations, the value of which are difficult to access.² The Lorentz force acts on moving charged species, depends on their velocity and the magnetic field amplitude; it is maximal (resp. minimal) when v and B are orthogonal (resp. parallel). Its order of magnitude ~10³ N/m³ for B = 1 T and for J = 100 mA/cm², is comparable to the buoyancy force [15] but remains inferior to mechanical-stirring. Its common effect is to thin the diffusion layer, which improves mass-transfer to the electrode [16][17][18]. In electrodeposition, the deposition rate and limiting current density are increased proportionally to $B^{1/3}$ [19][20][21]. The effect of the field-magnitude strongly depends on the distance d from the permanent magnet, decreasing as $1/d^2$, as studied by Mogi *et al.* for a $B \perp [22]$.

Lorentz force can induce flow patterns at a very small scale, where high current density can exist, which is impossible via mechanical-agitation. At the edge of an electrode with a SMF perpendicular to the electrode surface, the current density is no longer parallel to the SMF, inducing magneto-hydrodynamic (MHD) effect / micro-MHD convection; this corresponds to circular vortices around nucleation sites (F_L along u_{θ} in polar coordinates), favoring two-

 $^{^{2}}$ A quick estimation from usual gradient values [13] reaches 10^{1} - 10^{3} N/m³. However, the authors have never seen this scrutinized in the literature.

dimensional growth against three-dimensional growth (Figure 1a,b) [23][24]. Many related effects were observed during electrodeposition, depending on the applied magnetic field direction on the cathode surface. B_{II} can induce smoother deposits, more homogeneous surface with finer grains and better magnetic properties (for electroplated amorphous quaternary FeNiPGd films, Figure 1c) **Erreur ! Source du renvoi introuvable.**[25]. Besides, crystallographic growth-directions are preferred for metallic electrodeposits, which can alter the deposits shape [26].

The rest potential for (non)magnetic electrodes can be shifted with a B_{\perp} , leading to rougher deposits [27][28] or enhanced corrosion [29][30][31], these effects depending on the operating parameters. Using a rotating magnetic field or a B_{\perp} , Mogi et al. selectively-deposited an enantiomer preferentially to the other, depending on the polarity of the magnetic field [22][32]. Zou *et al.* used a rotating SMF to create dendrites of self-supported Ni_xCo_{1-x} nanowire membrane [33]. In both cases, the film thickness is generally larger, owing to enhanced mass-transfer [25].



Figure 1: Using the Lorentz force in electrodeposition. a,b) To eliminate Li dendrite growth. a) The electric field around lithium nucleii or protuberances on the current collector is larger than for flat areas, inducing Li⁺ flow towards the tips and yielding dendrite growth. b) With a magnetic field applied perpendicular to the current collector, local Li⁺ flux around the tip area cuts perpendicularly the magnetic line, inducing Lorentz force and micro-MHD vortices (swirling convection), leading to more homogeneous Li plating. Reproduced from [23] with

permission from Wiley. c) **During plating of FeNiPGd films**. Atomic force micrographs of FeNiPGd films electroplated at different magnetic fields amplitude B = (a) 0 mT, (b) 10 mT, (c) 20 mT, (d) 30 mT, (e) 40 mT; and (f) roughness value (*Ra*) of the films obtained under different magnetic field intensity (10 mT < *B* < 40 mT). Reproduced from [25] with permission from Elsevier.

SMF also influence bubbles: both B_{\perp} or B_{ll} favor bubbles removal, lured by micro-MHD convection in the clockwise or anticlockwise direction, depending on the sign of the charge adsorbed on the bubble [16]. This reduces the electrode masking by decreasing the bubbles' coverage, *e.g.* for water-splitting reaction, thereby reducing the Ohmic-loss and increasing the number of nucleation sites [34]. Bubbles removal also fosters better electrodeposits with fewer vacancies and defects [18][35]. Using a CCD camera for H₂ generation under SMF, Liu *et al.* showed a bubbles' size reduction ranging from 50 to 100 µm on a magnetized Ni-wire electrode (residual magnetic field: $B_r = 0.03$ T) [15][17][36][37].

Better efficiency of polymerization [38], organic reactions [39] and water reduction/oxidation were also recorded [34][40], these examples further demonstrating the impact of the Lorentz force in electrochemistry.

2.2 The Kelvin force

"Kelvin effects are ubiquitous because any single-electron-transfer redox process generally involves paramagnetic species. Magnetic species in an electrolyte are usually paramagnetic cations, free-radicals or molecular species with unpaired spin such as O_2 . An applied field H induces a magnetization $M = \chi H$, where, $\chi (= \chi_m c^3)$ is the dimensionless Curie-

³ Some authors propose [77] to write the susceptibility as the sum of two contributions: solvent and electrochemical reactants ($\chi = \chi_s + \chi_m c$), where χ_m is the molar susceptibility of the electroactive species *c* and χ_s the susceptibility of the solvent. Nevertheless, χ_s is generally far below $\chi_m c$.

law magnetic susceptibility", proportional to the spin of the species in aqueous solution [12]. The Kelvin force density (also denominated magnetic field-gradient force or magnetophoretic force) is given by equation 3:

$$\boldsymbol{F}_{\mathbf{K}} = 1/(2\mu_0) \ c \ \boldsymbol{\chi}_{\mathrm{m}} \ \boldsymbol{\nabla}(\boldsymbol{B}^2) \tag{3}$$

with μ_0 the magnetic constant in free space, *c* the concentration and χ_m the molar magnetic susceptibility. Independent on the magnetic field direction, this force creates convection, which drives paramagnetic species ($\chi > 0$) towards high magnetic field gradient and diamagnetic species ($\chi < 0$) towards low magnetic field gradient. Order of magnitude from 10³ to 10⁶⁻⁷ N/m³ can be reached at the vicinity of micrometer-size ferromagnetic (FM) element, the strength increasing for smaller FM objects [12]. Okada *et al.* used this effect for the ORR in a PEMFC, using a magnetized [20 wt.% Pt/C + Nd/Fe/B nanoparticles (NPs)] electrode under a magnetic field B = 4 T: O_{2 (g)} ($\chi_m = 3.449 \times 10^{-6} \text{ cm}^3/\text{mol} >> \chi_m = -12.96 \times 10^{-6} \text{ cm}^3/\text{mol}$, for liquid water at T = 273 K [41]) was attracted toward the cathode surface, limiting the mass-transfer issues [7]. This force being non-conservative ($\nabla \times F_K = (1/2 \mu_0)\chi_m$ [$\nabla(c) \times \nabla(B^2)$], convection should arise in region of non-uniform concentration if the concentration gradient is perpendicular to the magnetic field gradient, thereby thinning the diffusion layer and enhancing mass-transfer, as for the MHD effect [30]. Using field gradient near the working electrode (WE) surface with magnetic elements (3d-elements), direct and inverse patterning could be obtained upon electrodeposition [12][42][43].

2.3 Spin-dependent electrochemistry

Another domain of growing interest is the spin-polarization of exchanged electrons and open-shell (valence) electrons of catalysts. Naaman *et al.* observed that chiral compounds act as spin-filter: one polarization of the electron's spin is favoured against the other when transmitted through a chiral molecule, yielding a current of polarized-spins. This "chiral-

induced spin-selectivity" (CISS) [44] was applied in several domains [28,45-47]; spinpolarization was observed using FM or antiferromagnetic (AFM) WE [45], and used to reach giant-magnetoresistance [48]. By polarizing the spin through a chiral compound or by applying a magnetic field on a ferromagnetic WE, enhanced electron-transmission was observed for O₂ evolution, while the side reaction of H₂O₂ production was reduced during water-splitting [30][46][49][50][51]. The effect is explained by the more favourable energetic path of the transferred electrons upon polarization: when the OH· radicals reacting into O₂ have their electronic spin co-aligned in the laboratory frame, they interact on a triplet potential, yielding triplet oxygen, whereas peroxide formation is symmetry-forbidden (Pauli principle, Figure 2) [46]. One can imagine that by polarizing the electrons, the overlap-integral of the radicals' molecular orbitals is higher than without polarization, leading to the formation of more kinetically-favourable species, even though the thermodynamics is not favoured. Density Functional Theory (DFT) calculations indicated that the energy difference between the triplet and singlet state of O_2 is ca. 1 eV [9][52][53]. By simply approaching a permanent magnet (B = 450 mT) to a NiZnFe₄O_x/Ni-foam FM WE, Garcès-Pineda et al. enhanced the oxygen evolution reaction (OER) current from 24 to 40 mA/cm² at 1.65 V_{RHE}. Knowing that 3delements have a maximum spin-polarization of 40 and 23% for Fe and Ni respectively [45][54], and that a 450 mT magnet is probably insufficient to reach the maximum catalyst spinpolarization, superior performances might be reached [53]. This enhanced electrontransmission was explained by a decreased energy-barrier under magnetic field, following the Zeeman effect, even if this perturbation might be very small [31][55]. Forslund et al. studied the overlap-integral between the Ni and Fe 3d bands and the O 2p band for Ruddlesden-Popper oxides: cross-gap hybridization between $e_g(Ni)$, p(O) and $e_g(Fe)$ bands across the Fermi level were predicted (DFT calculations) to enhance both charge-transfer interactions across Fe-O-Ni bridges and the bandwidth available for electrode-adsorbate electron transfer. The sufficient Fe content favoured the 4-electron pathway (yielding HO·) over the 2-electron pathway (yielding HOO·) [56]. Spin-dependent electrochemistry (SDE) induced enantioselectivity too, as observed for oxalate oxidation [57].



Figure 2: Spin-dependent electrochemistry. In a) the support is achiral so the spins are not polarized, allowing the interaction on the singlet potential, yielding the thermodynamically-favourable hydrogen peroxide. In b) the hydrogen peroxide is forbidden because the photo-anode induces polarized spin that interacts on the triplet potential, favouring O₂ evolution. Reproduced from [46] with permission from Wiley.

Others studied the binding of reaction intermediates on FM or AFM WE, notably for the OER and ORR. Using DFT calculations, assuming quantum spin-exchange interaction (QSEI) for 3d-elements, they showed that the binding-strength can be controlled through spinpolarization. Dominant AFM QSEI enhance the interatomic Coulomb potential and pairlocalization versus a nonmagnetic covalent structure, while dominant FM QSEI lowers electronic repulsion and reduces pair-localization. The FM intra-atomic QSEI decreases the adsorption enthalpies of reactants, yielding milder oxidant intermediates adsorption and enhanced ORR activity. On the contrary, dominant interatomic AFM pair-localization lowers the conductivity, leading to decreased ORR efficiency. Pair-localization and strong couplings in fully AFM-orderings may strongly decrease some reaction intermediates chemisorption, detrimentally-affecting reactivity [58][59][60][61][62]. According to Barut and Kraus [63], magnetic interactions play the dominant role at small distances, likely inducing additional effects than the Coulomb interaction of e^+e^- system (two-body bound state), *e.g.* symmetry of the potential, leading to such local energy modification.

Although spin-polarization is thrilling, it faces limitations: it is constrained by the lifetime and distance over which the electrons' spin is conserved. Bullard *et al.* used magnetic-conducting atomic-force microscopy to show that the polarized current created by the CISS effect can flow through achiral molecules over ≈ 35 Å, which exceeds the length of chiral moiety (≈ 15 Å) [64], while Mishra *et al.* observed CISS effects through extracellular conduits, over distances above 10 nm [65]. Kumar *et al.* recorded a long-lifetime (> 10 ms) in an AlGaN/GaN device [66]. Furthermore, the spin-selectivity may be pH–dependent for certain reactions, as Garcès-Pineda *et al.* did not observe spin-restriction for water splitting at pH 11 using Ni-foam-supported 3d-element oxides in 1 M KOH electrolyte [53], while Tassinari *et al.* observed enhanced HER with TiO₂ NPs coated with a chiral poly(fluorene-co-thiophene) and decorated with CdSe quantum dots in 0.35 M Na₂SO₃ + 0.25 M Na₂S aqueous electrolyte solutions (pH 9.5) [67]. These results indicate that the rate-limiting step is not spin-restricted under different pH conditions, or with different catalysts.

3 Alternating magnetic field (AMF)

To our knowledge, only one work reported the effect of an AMF on water electrolysis processes [68]: Niether *et al.* used carbon-felt-supported core-shell magnetic and catalytic NPs as electrode in AMF-enhanced alkaline water-splitting. Under radiofrequency (300 kHz) AMF,

the NPs magnetization changes direction with the AMF direction, the induced work releasing heat, which heightens the NPs temperature. This principle, hyperthermia, was first used for cancer therapy [69]. The heating of the NPs, characterized by the Specific Absorption Rate (SAR) value (the area A of the hysteresis-loop multiplied by the frequency f, equation 4) [70], enhances the reaction kinetics (Figure 3).

$$SAR = \int_{-H\max}^{+H\max} \mu_0 M(H) dH f$$
 (4)

OER and HER chronopotentiometric measurements showed decreased overpotential by 250 and 150 mV, respectively, for B = 48 mT, J = 18 mA/cm², although heating may not be the only effect at stake (Kelvin of Lorentz forces may help releasing O₂ gases).

Other works used switchable magnetic field to micro-manipulate single Fe₃O₄ microparticles, and take benefit of their electrochemical activity by attracting or removing the particles by switching on or off a magnetic field [71]. Manipulating particles individually may be of interest for several domains such as engineering, biology, chemistry and physics.

Besides, AMF has been used to magnetized sea water (0.3 T at 25 Hz), which then shows notably limited corrosion effect on Cu (corrosion current reduced by 10), and fewer corrosion products [72].



Figure 3: Water electrolysis activated by the local heating of MNPs in an AMF (**Hyperthermia**). a) General representation of the AWE cell operating inside the coil that generates the AMF, the cell being, ideally, powered with renewable electricity. b) Schematics of the electrolysis AWE flow-cell set-up. Reproduced from [68] with permission from Springer-Nature.

4 Concluding remarks

Many different puzzling effects at several length scales are observed when coupling alternating or static magnetic field with electrochemistry. The most important are the (i) Lorentz force, inducing convection (MHD and micro-MHD effects), (ii) Kelvin force, inducing convection in magnetic field-gradient on para(dia)magnetic species (iii) spin-polarization, orienting reaction pathways and (iv) hyperthermia, a local heating at the electrode surface, that enhances reaction kinetics. Although qualitative explanations are proposed, these phenomena are not fully-investigated yet and no uniform/universal model exists to explain them, though numerical calculations tried to explain MHD convection [73]. A permanent magnet of $B \approx 1$ T or Helmholtz coils are usually used to trigger the effects, and specially-designed cells are necessary to scrutinize them, a hurdle for industrial applications. Table 1 summarizes some recorded enhancements of water oxidation/reduction reaction, thanks to the application of a magnetic field.

Other effects were reported, like liquid-metal droplet actuation induced by a magnetic field [74][75][76], that are not presented here. Neither are exposed the techniques used to measure spin-polarization [9][51], and the force acting on the double layer or the one proportional to $\nabla(c) * B^2$ [8][28], although Coey *et al.* found it negligible [77]. Anyway, these discoveries are the "tip of the iceberg" for many applications, especially for spin-dependent electrochemistry, and they could have major impacts in several domains, notably for water-splitting at lower voltage, hence lower prices.

Table 1: selected summary of the main "magnetic effects" mentioned for

electrochemical reactions (results with currents above c.a. mA/cm² have been recorded).

	Effect/equation advanced	<i>B</i> applied	Overpotential and current density	Catalysts	Medium (alkaline/acid)	References
SMF	Spin-polarization of O ₂ electrons	≤ 450 mT during experiment	$24 \rightarrow 40 \text{ mA/cm}^2$ at 1.65 V _{RHE} (iR- compensated)	NiZnFe4Ox/Ni- foam electrode	alkaline pH 14 (1 M KOH)	[53]
	large surface areas (incomplete Ostwald ripening), fast charge/mass transportation ability (metallic phase) and possible lattice defect-enhancement effect	Only used to create the nanowires	-150 mA/cm ² at - 0.2 V _{RHE}	Ni _{0.5} Co _{0.5} nanowire membrane	alkaline pH 14 (1 M KOH)	[33]
	Lorentz force which triggers gas elimination and the reduction of average gas fraction at electrode surface	0.9 T, B//	Cell voltage 3.95 V at 1.25 A/cm ²	two Platinum sheets (4×4 mm) are used as anode and cathode	4.24 M KOH	[17]
	Lorentz force which drives away H ₂ bubbles	0.9 T, B//	Cell voltage of 3.3 V at 200 mA/cm ²	foam copper was cut as the cathode and a nickel one as the anode	0.2 M KOH	[25]
	Lorentz force (MHD) reducing hydrogen bubble	magnetized electrode (B _r = 30 mT)	0.4 mA/cm ²	Ni wire as cathode	alkaline pH 14 (1 M KOH)	[37]

	diameter and bubble coverage					
	at the electrode surface					
	Zeeman energy, MHD convection	125 mT	56 mV gain at 20 mA/cm²	urchin-like Co ₃ O ₄ spheres on the nickel foam	alkaline pH 14 (1 M KOH)	[34]
	Kelvin force	150 mT	5.8 mA/cm \rightarrow 8.1 mA/cm at +0.2 V _{RHE} for ORR	Fe-N-C/S carbon gels	O ₂ -saturated 0.02 M PB buffer with 0.15 M K ₂ SO ₄ of pH 5.5	[30]
	Smaller adsorption energies and bond creation energies	magnetic catalyst	$-0.49 V_{RHE}$ at 100 mA/cm ² for HER	FeB@Fe	alkaline pH 14 (1 M KOH)	[40]
AMF	Hyperthermia	48 mT at 300 kHz during experiment	 18 mA/cm² at 1.60 V vs. RHE (iR-compensated) - 250 mV gain for the OER and 150 mV gain for the HER at 18 mA/cm² 	FeC@Ni	alkaline pH 14 (1 M KOH)	[68]

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Author contributions

JD and MC have conceived and designed the study; all authors have equally analyzed and interpreted the data. VG did draft the first version of the article, which has been critically-revised and finally-approved by all authors equally.

Lexicon

B: magnetic flux density vector, its modulus is in Tesla. We abusively call it magnetic field.

H: magnetic field vector, its modulus is in A/m

 $B_{\prime\prime}$: magnetic field applied parallel to the electrode surface

 $B\perp$: magnetic field applied perpendicular to the electrode surface

M: magnetization in A/m

 $F_{\rm L}$: Lorentz force density vector, its modulus is in N/m³

 $F_{\rm K}$: Kelvin force density vector, its modulus is in N/m³

J: current density vector, its modulus is in A/m²

E: electric field vector, its modulus is in N/C

v: species velocity vector, its modulus is in m/s

 σ : electric conductivity in S/m

d: distance from the permanent magnet

 u_{θ} : second coordinate in the polar coordinates, u_{r} and u_{z} being the first and third coordinates, respectively.

 χ : dimensionless magnetic susceptibility,

 χ_m : molar magnetic susceptibility in m³/mol

- c: concentration of the species in mol/m^3
- $\mu_0:$ magnetic permeability in free space, in kg.m.A^{-2}.s^{-2}
- ∇ : gradient operator
- $\nabla \times$: rotational operator
- *f*: frequency in Hz

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