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¹ The thermocapillary effect on gas bubbles growing on electrodes of different sizes

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6 Abstract

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Recently, the strongly inhomogeneous current density occuring near a microelectrode was identified as driving a thermocapillary electrolyte flow near gas bubbles growing during electrolysis [1]. The present paper is investigating this effect in more detail under various operating conditions. Furthermore, by simplified modeling, the question is answered of whether this effect is also of importance at large planar electrodes. The direction of the thermocapillary force on the bubble is found to change from retarding to advancing the bubble release when the size of the electrode is increased. Conclusions are drawn on how the thermocapillary effect at planar electrodes depends on the electrode coverage and the bubble departure size, also considering industrially relevant values of the current density.

7 Keywords: hydrogen evolution, electrolysis, thermocapillarity, Marangoni force, microelectrode,

8 macroelectrode

9 1. Introduction

Interfaces between media of different phases are ubiquitous in nature and play an important role in many 10 physico-chemical systems. If liquids are involved, capillary effects often influence the system behavior [2]. 11 The surface tension for a given liquid-gas pair is known to depend on physical properties of the interface, 12 namely the temperature, chemical composition and electric potential in the presence of interfacial charge [3]. 13 Thus, interfacial gradients of these physical properties cause the surface tension to vary along the interface. 14 As a result, due to unbalanced forces at the interface, fluid elements there experience a net shear stress and 15 move towards interface regions of higher interfacial tension. The resulting capillary flow is commonly called 16 Marangoni flow [4]. This Marangoni effect is known to occur in many systems of scientific and technological 17 importance. For example, thermo-capillarity is the mechanism which drives the well-known Bénard cells 18 ([5–7] and references therein), a phenomenon occurring in many engineering heat transfer applications. 19 In liquid-liquid extraction processes, soluto-capillary flows may significantly change the mass transfer rate 20

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[8]. Furthermore, Marangoni flow exists at sessile evaporating droplets due to the interfacial temperature
gradient caused by a nonuniform rate of evaporation [9]. This influences the well-known coffee-ring-like
structures [10], and may thus affect patterned deposition [11] and particle self-assembly [12].

In 1959, Young et al. [13] demonstrated the profound effect which Marangoni flow can have on air bubbles 24 in liquids by applying a positive temperature gradient in the direction of gravity to suspend them against 25 buoyancy. The temperature gradient induced a surface tension gradient along the bubble interface and 26 thus exerted a thermocapillary stress on the interface, which caused the bubble to experience a force against 27 buoyancy. At a sufficiently steep temperature gradient, the thermocapillary force can be as large as buoyancy. 28 Later, with the advent of space research, bubble actuation in micro-gravity by means of thermocapillarity 29 tracted much interest [14, 15]. Experiments conducted in the space shuttle in orbit showed that by applying 30 temperature gradient of the order of 1 K/mm, a bubble migration speed of $\sim 1 \text{mm/s}$ could be achieved а 31 for a bubble of millimeter size [16]. McGrew et al. [17] speculated that Marangoni convection might have a 32 significant contribution to heat transfer in nucleate boiling, as the observed flow structure around a pendant 33 air bubble heated from above was found to be similar to that of a vapor bubble during boiling. Later, in 34 nucleate boiling experiments conducted under microgravity conditions, the heat transfer rate obtained by 35 Straub was similar to that obtained under normal gravity conditions [18], which lent further credence to the 36 importance of Marangoni flow around the bubbles. Further experiments and numerical simulations [19–21] 37 established the significant importance of thermocapillarity-driven flow in the subcooled boiling regime and 38 showed that the resulting Marangoni force acting on the bubble slowed down their detachment from the 39 boiling surface [22]. 40

It has long been hypothesised that the Marangoni effect also influences the dynamics of bubbles grown 41 electrochemically on electrodes [23, 24]. Definitive experimental evidence of the Marangoni effect was first 42 provided by Yang et al. [25], who performed a detailed investigation of the interfacial flow around hydrogen 43 bubbles grown on a Platinum microelectrode. Then, by simultaneously measuring the electrolyte velocity 44 and temperature and correlating the results with the numerical solution obtained when considering only ther-45 mocapillarity, Massing et al. [1] were able to show that the Marangoni flow observed is primarily attributed 46 the thermocapillary effect caused by the temperature gradient along the bubble interface. Interestingly, 47 for strong local boiling at microheaters, the thermocapillary flow found near the gas bubble qualitatively 48 matches the flow structure near the electrogenerated gas bubble at a microelectrode [1], though due to strong 49 laser heating the maximum flow velocity is about 10 times higher [26]. 50

As gas bubbles are generated in many electrochemical processes such as plating, refining of metal or the chloralkali process [27-29], a better understanding of the dynamics of electrogenerated bubbles and associated thermo-fluidic phenomena is of great technological and scientific interest. This is particularly relevant for H₂ production through water electrolysis, as the bubble evolution rate directly influences both the system throughput and the process efficiency [30], which may be of increasing importance for next⁵⁶ generation green energy storage and mobility applications [31–33].

As bubbles attached to the electrode surface disturb a homogeneous surface reaction rate, reduce the 57 active reaction area and increase the electrical resistance of the cell, the speedy removal of generated bubbles 58 from the electrode surface is highly desirable with respect to both deposit quality and electrical efficiency. 59 The instant of bubble departure and the departing diameter is determined by the equilibrium of forces acting 60 on the bubble [34]. However, the often-used Fritz equation [35] does not provide an accurate prediction of 61 the departure diameter [36]. For the special case of oxygen bubbles grown photocatalytically on TiO_2 62 nanorods where a temperature gradient is caused by light irradiation, Chen et al. have recently shown that 63 the inclusion of both thermocapillary and solutal Marangoni forces in the force balance of the bubble could 64 considerably improve the estimate of the bubble departure [37]. However, beside the aforementioned study 65 by Massing et al. [1], where a retarding influence of the thermocapillary effect on the bubble departure was 66 found, no systematic study has been carried out for electrogenerated bubbles. 67 For studying bubble evolution during electrolysis, microscale electrodes have been widely used, as the 68 nucleation area is limited, and single bubble growth can be more easily observed compared to the spatially 69 random nucleation occuring at large planar electrodes [38–41]. The recent work which was the first to 70 confirm the existence of thermocapillary flow around electrogenerated bubbles [1] was also carried out at 71 microelectrodes. However, there remains a need for more detailed investigations of this phenomenon. In 72

⁷³ particular, the question has not yet been addressed of whether this effect is also of importance at the larger ⁷⁴ electrodes, which are more relevant to industrial applications. Therefore, in this study we perform detailed ⁷⁵ numerical simulations and analyses of the thermocapillary effect at electrogenerated bubbles on electrodes ⁷⁶ of varying size. Different operating conditions are investigated, and the important qualitative differences ⁷⁷ found between microelectrodes and macroelectrodes will be emphasized.

78 2. Simulation setup

Microelectrode. To study the thermocapillary effect around a hydrogen bubble at a microelectrode of vary-79 ing size, our simulation setup closely follows the methodology used by Massing et al. [1]. A cylindrical 80 electrochemical cell with a radius $R_c = 5 \text{ mm}$ and a height $R_h = 5 \text{ mm}$ is considered. It is filled with an 81 aqueous solution of $1 \text{ M H}_2\text{SO}_4$. The computational domain is sketched in Figure 1a. At the bottom of 82 the cell, a bubble with a radius $R_b = 560 \ \mu \text{m}$ sits on a Pt microelectrode (radius R_e), which works as the 83 cathode. The contact angle between the bubble and electrode is $\theta_c = 4.2^{\circ}$ [1]. It should be noted here that 84 the bubble radius is much smaller than the size of the cell. The Pt microelectrode is embedded in a glass 85 bottom. The top of the cell is completely covered by a counter-electrode made of platinum. The electrodes 86 and the glass bottom have a height of 5 mm each. The inclusion of the glass bottom and both electrode 87 domains is important for an accurate calculation of the temperature field in the electrolyte due to non-zero 88



Figure 1: Computational domain (not to scale). a) microelectrode, b) macroelectrode.

⁸⁹ and non-uniform heat flux through these domains.

We follow Massing et al. [1] and consider a bubble of fixed size to resemble a late stage of the bubble growth cycle. As shown in Figure 1, we utilize the rotational symmetry of the cell and hence perform axisymmetrical simulations. The leftmost edge in Figure 1 is the axis of symmetry. The following equations to be solved describe the spatial and temporal distribution of the electric potential (ϕ), velocity (**u**) and temperature (*T*) in the electrolyte:

$$\nabla^2 \phi = 0 \tag{1}$$

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$$\nabla \cdot \mathbf{u} = 0 \tag{2}$$

$$\rho\left(\frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \cdot \nabla)\mathbf{u}\right) = -\nabla p + \mu \nabla^2 \mathbf{u}$$
(3)

$$\rho C_p \left(\frac{\partial T}{\partial t} + \left(\mathbf{u} \cdot \nabla \right) T \right) = k \nabla^2 T + \frac{|\mathbf{j}|^2}{\sigma}$$
(4)

Here, the primary current density $\mathbf{j} = -\sigma \nabla \phi$ is obtained by solving Eq. (1), where σ denotes the electrical conductivity of the electrolyte, which is assumed to be constant. The electrolyte velocity is obtained by solving the incompressible Navier-Stokes equation (3) complemented by the incompressibility constraint (2), where p, ρ and μ denote the pressure field and the material properties of density and viscosity. Thermal and solutal buoyancy effects may safely be neglected, as discussed in [1]. The electrolyte temperature is obtained by solving Equation (4), where C_p and k denote the material properties of the specific heat capacity and thermal conductivity. The latter term denotes Joule heating due to electric current passing through the electrolyte. It should be noted here that this term may be neglected in the electrodes, where the electrical conductivity σ is several orders of magnitude larger than in the electrolyte. Therefore, in all other domains (both electrodes, glass bottom and gas bubble), only heat diffusion is solved:

$$\rho C_p \frac{\partial T}{\partial t} = k \nabla^2 T \tag{5}$$

¹⁰⁸ Here, convective heat transport in the gas bubble may safely be neglected, as discussed in [1].

The boundary conditions applied to these equations are as follows: for Equation (1), the electrode surfaces exposed to electrolyte are kept at a fixed potential, i.e. $\phi = 0$ for the working electrode and $\phi = \phi_0$ for the counter electrode. The outer boundary at R_c is electrically insulating, i.e. $\frac{\partial \phi}{\partial \mathbf{n}} = 0$, where **n** is the normal unit vector. For Equation (3), a shear stress balance is applied at the electrolyte–bubble interface, i.e. $\tau_H = \tau_M$, where the hydrodynamic shear stress is given by $\tau_H = \mu \left(\nabla \mathbf{u} + (\nabla \mathbf{u})^T \right)$. **n**, and the thermal Marangoni stress is given by

$$\tau_M = \frac{\partial \gamma}{\partial T} \cdot \nabla_s T \tag{6}$$

Here, γ denotes the interfacial tension, which is assumed to depend on the temperature only, and 115 $\nabla_s \equiv \nabla - (\nabla \mathbf{n}) \mathbf{n}$ denotes the gradient along the surface of the interface. As the temperature field is 116 solved in all domains by Eqns. (4) and (5), boundary conditions are required only at the outermost surfaces. 117 The top and the bottom of the computational domain are kept at an ambient temperature $T_{amb} = 20^{\circ}$ C, 118 while at the outer surface at R_c far from the bubble an adiabatic condition $\frac{\partial T}{\partial \mathbf{n}} = 0$ is applied. Finally, at the 119 symmetry axis at r = 0, a zero radial gradient condition is applied on all variables, i.e. $\frac{\partial}{\partial r}(\phi, \mathbf{u}, T) = 0$. The 120 temperature coefficient of the surface tension at the gas-liquid interface is taken to be $\partial \gamma / \partial T = -1.6 \cdot 10^{-4}$ 121 N/m· K [1]. The value of the surface tension γ does not appear directly in our model as the bubble is 122 assumed to be of fixed shape and size. All other material properties are assumed to be constant over the 123 electric potential and temperature range solved for and are shown in Tables 1 and 2. 124

All equations are solved using the FEM-based simulation software COMSOL 5.4. The initial conditions applied are zero potential, electrolyte at rest and ambient temperature. As discussed extensively in [1], a time integration period of one second was chosen at which nearly stationary and realistic values of the temperature and velocity distribution and the resulting Marangoni force may be expected. To study the effect of the size of the working microelectrode, R_e is varied while R_b is kept constant. We introduce a

	$\mu = 10^{-3}$ Pa·s
	$\sigma = 40 \text{ S/m}$
Electrolyte	$\rho = 10^3 \ \rm kg/m^3$
	$C_p = 4.182 \text{ kJ/kg·K}$
	$k = 0.58 \text{ W/m} \cdot \text{K}$

Table 1: Material properties of the electrolyte for Equations 3 and 4

Material	$\rho (\rm kg/m^3)$	$C_p(kJ/kg\cdot K)$	$k(W/m \cdot K)$
Platinum	21450	0.13	72
Glass	2201	1.052	1.38
Hydrogen	0.09	14.32	0.186

Table 2: Material properties for Equation 5

¹³⁰ non-dimensional length parameter as the ratio of the bubble radius and the microelectrode radius,

$$\alpha = \frac{R_b}{R_e} \tag{7}$$

As we are here primarily interested in studying how the increasing size of the microelectrode affects the behavior in the vicinity of the gas bubble, unless otherwise mentioned, the radius of the microelectrode is varied from 50 μ m to 400 μ m such that $R_e < R_b$, i.e. $\alpha > 1$.

Macroelectrode. To study the thermocapillary effect on a large, flat electrode where multiple gas bubbles 134 are growing simultaneously, we take a simplified approach as follows: we first zoom into a small part of the 135 electrode in order to focus only on a single bubble. A cylindrical domain around this bubble is considered, 136 where the radial extension R_c of the domain is half the distance to the next neighboring bubble. We 137 assume that the cell with large electrodes is essentially composed of this periodically repeating domain. 138 Though there are time shifts between the evolution cycles of neighboring bubbles, our simplified approach 139 therefore assumes that all bubbles develop synchronously and also ignores possible coalescence phenomena 140 of neighboring bubbles during growth. A further approximation consists in neglecting azimuthal variations 141 of potential, temperature and velocity which might arise from the finite number of neighboring bubbles 142 sitting at unknown azimuthal positions. This approach treats neighboring effects in an approximate manner 143 averaged over the azimuthal direction and allows the problem to be simulated axisymmetrically. We can 144 therefore make use of the axisymmetric computational domain shown in Figure 1b, which is similar to the 145 one used previously for the microelectrode shown in Figure 1a, and which only needs minor modifications: 146 at the bottom of the cell, the outer glass part is removed, and the electrode is now enlarged over the full 147 radial extent of the cell R_c . At the outer radial boundary, the conditions of electrical insulation for the 148

¹⁴⁹ potential and an adiabatic condition for the temperature remain unchanged, but in the case of the velocity,
¹⁵⁰ vertical slip is applied. The governing equations to be solved for remain as before. In order to quantify the
¹⁵¹ ratio between the bubble size and the size of the cell, we introduce

$$\Theta = \left(\frac{R_b}{R_c}\right)^2 \tag{8}$$

In the framework of our approach, this ratio describes the bubble coverage of the electrode which is defined as the ratio of the bubble area projected on the electrode and the electrode surface area. We can later simply change the bubble coverage of the electrode (which is known to depend on e.g. the mean current density), for example by varying the radial extension of the cell while retaining a fixed bubble size.

156 3. Results and Discussion

157 3.1. Microelectrode

158 3.1.1. General overview

It is evident from Equation 4 that the electrolyte is heated because of the electric current passing through 159 the cell. This causes a temperature gradient along the bubble interface which gives rise to Marangoni 160 flow. Therefore, the current distribution near the interface strongly influences the temperature and flow 161 distribution in the electrolyte. Figure 2(a) shows numerically obtained current lines near the interface for 162 $\alpha = 11.2$. As the gas bubble is electrically insulating, current must pass around it to converge at the 163 microelectrode. Therefore, as the entire cell current must squeeze through a narrow wedge-like region at 164 the bubble foot, the local current density there is very high. In addition, the presence of the bubble makes 165 the outer region of the electrode more accessible to the electric current. This causes an inhomogeneous 166 distribution of the current density over the electrode surface wetted by the electrolyte, as shown in Figure 2(b)167 for different microelectrode sizes. Here, the cell voltage was kept constant at $\phi_0 = 4.5$ V, and the radial 168 position on the microelectrode is presented normalized by the electrode radius, i.e. $r^* = r/R_e$. As the 169 current density is shown on a logarithmic scale, it clearly can be seen that it varies quite strongly over the 170 radial electrode position and is largest at the electrode periphery. The smaller the microelectrode becomes, 171 the stronger the radial variation of the current density and also its maximum value at the electrode periphery. 172 As the rate of heat generation scales with the current density squared $(|\mathbf{j}|^2)$, the electrolyte experiences 173 strong heating near the periphery of the electrode. Figure 3 shows a zoomed view of the distribution of the 174 temperature rise $(\Delta T = T - T_{amb})$ in the vicinity of the outer electrode edge for $R_e = 200 \,\mu\text{m}$ ($\alpha = 2.8$). It 175 can be seen that the temperature maximum occurs just above the microelectrode periphery at $r^* = 1$. As 176 the surface tension of the bubble interface decreases with increasing temperature, the electrolyte is pulled 177 towards colder parts of the interface, thus establishing an interfacial flow away from the point of maximum 178 temperature. As this bidirectional interfacial flow must be replenished from the bulk, a double-vortex 179



Figure 2: (a) Current lines near the bubble interface as obtained from a simulation for $\phi_0 = 4.5$ V and $\alpha = 11.2$. (b) Distribution of the magnitude of the current density ($|\mathbf{j}|$) on microelectrodes of different size versus the radial position on the wetted part of the electrode normalized with the electrode radius ($r^* = r/R_e$). $\phi_0 = 4.5$ V. $|\mathbf{j}|$ is shown in log scale.

structure is created, as shown for the first time in Figure 3. As will be elaborated in the following, this double-vortex structure is a characteristic feature of the thermocapillary flow driven at the interface of an electrogenerated bubble. In the case of microelectrodes, for geometric reasons, the temperature hotspot appears above the electrode periphery, thus confining the left vortex, and the thermocapillary effect is dominated by the large right vortex. The small size of the left vortex explains, why it was not observed during earlier work at microelectrodes [1].

186 3.1.2. Interfacial temperature and velocity profile

¹⁸⁷ We now discuss the interfacial temperature and velocity profiles at electrodes of different sizes R_e while ¹⁸⁸ the cell voltage is kept constant at $\phi_0 = 4.5$ V. It is to be noted here that, at constant voltage, the electrical ¹⁸⁹ resistance of the cell decreases and hence the cell current increases when the microelectrode is enlarged. The ¹⁹⁰ obtained profiles of the temperature increment ΔT and tangential velocity ($u_t = \mathbf{u} \cdot \mathbf{t}$) along the bubble ¹⁹¹ interface are shown in Figure 4, with the definition of the angular position θ based on Figure 1. Starting ¹⁹² from the point of contact between the bubble and electrode ($\theta_c = 4.2^\circ$), an initial temperature increment ¹⁹³ is seen in all cases. The smaller the microelectrode is, i.e. the larger α is, the steeper the increase in



Figure 3: Color contours of the temperature distribution ($\Delta T = T - T_{amb}$) and related electrolyte flow pattern (black: velocity vectors) near the temperature hotspot for a microelectrode of $R_e = 200 \,\mu\text{m}$ ($\alpha = 2.8$) at potential $\phi_0 = 4.5$ V. The radial coordinate is $r^* = r/R_e$.

temperature becomes. At the same time, the angle θ_m at which the maximum temperature increase ΔT_m 194 is observed reduces. This behavior can qualitatively be understood from the discussion on the temperature 195 hotspot in the section above: this was found to be located approximately above $r^* = 1$. Therefore, when the 196 microelectrode grows in size, θ_m also increases. A more detailed discussion on θ_m is given in Section 3.1.3. 197 Moving further along the interface away from θ_m , ΔT gradually decreases towards the bubble north pole at 198 $\theta = 180^{\circ}$. This general trend in the interfacial temperature profile is seen for all cathode sizes investigated. 199 It is also to be noted here that the variation of ΔT_m with α is not monotonic. This is because increasing 200 R_e while keeping ϕ_0 constant increases the cell current. Thus the total heat generation in the cell increases, 201 which tends to increase the peak interfacial temperature. As the electrode gets bigger, more heat is carried 202 away by the Pt microelectrode. Greater heat advection from the hotspot by the electrolyte (cf. Figure 4(b)) 203 also contributes similarly. These effects tend to reduce ΔT_m . As a result of the combined action of greater 204 heat generation and higher heat transfer, ΔT_m varies non-monotonically with R_e and thus α . Hence we can 205 see from Figure 4(a) that ΔT_m first increases and then decreases to an almost constant value with a larger 206 electrode i.e. smaller α . 207

The non-uniform interfacial temperature profile causes thermocapillary stress as shown in Equation 6. Because of the stress balance at the interface, $\tau_M = \tau_H \sim \frac{\partial u_t}{\partial \theta}$, where u_t denotes the tangential velocity at the interface, a Marangoni flow is driven, as already mentioned in Section 3.1.1. The interfacial flow profiles for different microelectrode sizes are shown in Figure 4(b) and can be understood by applying the following sequence of reasoning: According to Equation 6, when the negative temperature coefficient of the surface tension of the electrolyte mentioned in Section 2 is taken into account, positive or negative slopes

of the temperature profile cause negative or positive thermocapillary stress, respectively. Hence, because of 214 the stress balance, the tangential velocity of the Marangoni flow becomes negative or positive, respectively. 215 Consequently, as the flow is purely driven by thermocapillarity, the temperature hotspot separates a region 216 of negative tangential velocity at small angles from a region of positive tangential velocity at larger angles. 217 At the temperature hotspot θ_m itself, the tangential velocity must vanish, i.e. $u_t = 0$. In Figure 4 and later 218 also in Figure 5, an exemple of this is emphasized for the case $\alpha = 5.6$ by a vertical black line. The two 219 regions of velocity of opposite sign are the interfacial parts of the two counter-rotating vortices already seen 220 in Figure 3. As the microelectrode increases in size, the region of negative velocity also increases, yielding 221 a larger vortex above the microelectrode. At the same time, the extremum positions of maximum negative 222 and positive interfacial velocity are both shifted towards larger angles. During this process, the maximum 223 negative velocity decreases monotonically in amplitude, with the initially growing maximum velocity later 224 seeming to level out near a value of about 15 cm/s for $\alpha \sim 1$. However, the total kinetic energy at the 225 interface reaches a maximum at the largest microelectrode size, a straightforward example of the above 226 argument of maximum heat generation at smallest α . 227

The interfacial temperature distribution at the bubble is determined by the interplay between the gener-228 ation of heat and the diffusion and advection of heat. For the case of a constant cell voltage (ϕ_0) considered 229 above, all these quantities vary with varying electrode size (α). In order to partly decouple these effects, in 230 the following we study the case of a constant electric cell power which mainly (apart from kinetic losses) 231 determines the generation of heat by Joule dissipation. The total electric power of the cell is given by 232 $P = \phi_0 I_c$, where I_c is the cell current, which in the simulations was determined by integrating the normal 233 current density over the counter-electrode surface $(I_c = \int j_n dA)$. In the following, the electrical power is 234 kept constant at P = 13.4 mW and α is varied. This power value chosen is the same as the constant cell 235 voltage case at $\alpha = 11.2$, i.e. for the smallest electrode. Hence, the constant cell potential and constant 236 cell power cases are identical for $\alpha = 11.2$. This way, a major remaining factor determining $\Delta T(\theta)$ along 237 the bubble interface is the heat carried by the Pt microlelectrode, which increases with decreasing α . The 238 simulation results are shown in Figure 5. The interfacial temperature profile still maintains the character-239 istics seen in Figure 4(a). However, because the heat generation is kept constant and the heat carried away 240 from the electrolyte increases along with the electrode size, ΔT_m decreases monotonically with decreasing 241 α , as seen in Figure 5(a). Accordingly, the extrema of maximum negative and positive interfacial velocity 242 eventually decrease monotonically with decreasing α , as seen in Figure 5(b). The same holds for the total 243 kinetic energy at the interface. 244

245 3.1.3. Position of the temperature hotspot

As mentioned in Section 3.1.2, the reversal of the direction of interfacial flow coincides with the location of the temperature maximum ΔT_m at the angular position θ_m . It can be further seen in Figure 3 that



Figure 4: Constant cell voltage case: interfacial temperature (a) and tangential velocity (b) profile as a function of angular position for different electrode sizes expressed by $\alpha = R_b/R_e$

for $\alpha = 2.8$ the position of ΔT_m coincides with the upward projection of the microelectrode's outer edge ($r^* = 1$) onto the bubble interface, as shown in Figure 1. Therefore, we intend to study ΔT_m closer to see whether this geometrical interpretation also holds true for other microelectrode sizes. The angular position of the projection of the microelectrode's outer edge ($r^* = 1$) on the bubble interface is given as

$$\theta_e = \sin^{-1}(R_e/R_b) = \sin^{-1}(1/\alpha) \,. \tag{9}$$

Figure 6 displays the angular position of the interfacial temperature maximum θ_m versus the inverse relative electrode size $\alpha = R_b/R_e$ as obtained from simulations for the cases of constant cell voltage (Figure 4a) and of constant cell power (Figure 5a) in comparison with the values θ_e obtained from the geometrical approach by Equation 9. At small relative electrode size, i.e. at large α , the simulation results for both cases perfectly match with the geometrical relation from Equation 9, which provides a very accurate estimate of the location of the temperature maximum at the interface. As the microelectrode increases in size, only small differences



Figure 5: Constant cell electric power (P = 13.4 mW) case: interfacial temperature (a) and velocity (b) profile as a function of the angular position for different electrode sizes expressed by $\alpha = R_b/R_e$

are seen between the two simulation cases, besides an overall good match being maintained with Equation 9. 258 Eventually, at $\alpha = 1.2$, both cases seem to deviate from the analytical curve and deliver smaller angular 259 positions of the temperature hotspot than Equation 9. This is caused by the fact that as the microelectrode 260 increases in size, as seen from Figure 2, the non-uniformity of the current density, i.e. the ratio of the peak 261 value at the electrode edge in comparison to the mean value, decreases. Therefore, as α approaches unity, 262 inner parts of the electrode surface also contribute to the generation of the temperature hotspot, which leads 263 to the observed slight shift in the position of the temperature maximum towards smaller angular positions. 264 In summary, the geometrically estimated angular position θ_e of the temperature maximum θ_m based on the 265 inhomogeneous distribution of the current density at the microelectrode provides a useful estimation of the 266 position of the temperature maximum θ_m and the resulting interfacial flow characteristics at the bubble 267 interface at the microelectrode without the need to perform detailed simulations or experimentation. We 268 would also like to point out here that Equation 9 is only valid for $R_e \leq R_b$, which coincides with the range 269



Figure 6: Angular position of the interfacial temperature maximum (θ_m) versus $\alpha = R_b/R_e$ at fixed $R_b = 560 \,\mu\text{m}$ for the cases of constant cell voltage (red circles) and of constant cell power (blue squares). The dashed curve denotes the geometrical relation from Equation 9.

²⁷⁰ of studies for the microelectrode case presented here.

271 3.1.4. Marangoni force



Figure 7: Marangoni force acting on the bubble, shown as normalized by the buoyancy force $(F_M^* = F_M/F_B)$, at different values of $\alpha = R_b/R_e$. The *x*-axis is shown in logarithmic scale to emphasize the behavior at low α . In the case of constant cell voltage, $\phi_0 = 4.5$ V. In the case of constant cell power, P = 13.4 mW.

We now focus on investigating the resulting thermocapillary force on the bubble at different microelectrode sizes. As already mentioned in the introduction, the moment of bubble detachement is determined by the equilibrium of forces acting on the bubble. Therefore, a comprehensive treatment of all the forces acting on a bubble is of importance. A number of different forces may act on the bubble depending on the predominant physical conditions i.e. growth rate, imposed flow condition, orientation relative to gravity, etc. [34] The relevant static forces acting on an electrogenerated bubble are in general the buoyancy force, surface tension force, contact pressure force and hydrodynamic force originating from the flow of the electrolyte [42, 43]. Recently it was shown that beside the thermocapillary force [1], an electrostatic force may also play an important role [44].

The thermocapillary force on the bubble is a hydrodynamical force by nature and results from the interfacial flow around the bubble driven by temperature gradients. In the following, we will denote the force as the Marangoni force, which can be calculated by integrating the Marangoni shear stress over the bubble interface (Σ),

$$F_M = -\int_{\Sigma} \tau_M dA. \tag{10}$$

Here, dA denotes the related surface differential. Because of the axial symmetry around the bubble, the above integration results in a force which is acting parallel to the z-axis, i.e. vertically upwards or downwards, thus influencing the moment of detachment. In the following, we consider the Marangoni force F_M in relation to the buoyancy force $F_B = gV\Delta\rho$ where g, V and $\Delta\rho$ denote the gravitational acceleration, the bubble volume and the density difference between the electrolyte and gas, respectively. Thus, we introduce

$$F_M^* = \frac{F_M}{F_B} \tag{11}$$

where positive values of F_M^* imply a Marangoni force acting in the direction of buoyancy.

Figure 7 shows the dependence of F_M^* on the inverse relative electrode size (α) at a constant cell voltage 291 and constant cell electric power. Note that the differential local Marangoni force experienced by any part 292 of the interface is opposite to the direction of local interfacial velocity. According to the velocity profiles 293 shown in Figure 4(b) and 5(b), the lower part of the bubble ($\theta < \theta_m$) experiences a net Marangoni force in 294 the upward direction and the upper part of the bubble experiences a net Marangoni force in the downward 295 direction. As for the microelectrode, the extent of the flow vortex to the left of temperature hotspot is 296 smaller than that to the right of the temperature hotspot. The integration of Equation 10 gives a net F_M 297 in the downward direction, hence F_M^* is negative at large α . For the potentiostatic case, as with decreasing 298 α , the mean current density and thus the heat generation in the electrolyte grows, F_M^* also increases in 299 magnitude. When decreasing α further, the growing contribution from the left vortex becomes more and 300 more important, and F_M^* starts to decrease in magnitude near $\alpha \sim 1$. We further see that at $\alpha \sim 1$, due 301 to very strong heating, the Marangoni force has reached a magnitude of roughly 55% of buoyancy, which 302 corresponds to a very large mean current density value of 34.8 A/cm^2 . 303

As the electrode size is increased while keeping the total heat generation constant, the temperature nonuniformity along the interface reduces and the contribution from the two vortices becomes comparable. Thus, F_M^* decreases with decreasing α , becoming approximately zero at $\alpha \sim 1$.

307 3.2. Macroelectrode



Figure 8: Sketch depicting the qualitative differences in current distribution, position of temperature hotspot (red circle) and resulting thermocapillary flow pattern when the size of the electrode is increased from micro (left) to macro (right). The flow inside the gas bubble and the movement of the interface are not shown.

So far we have presented a detailed characterization of the thermocapillary effect on gas bubbles growing at microelectrodes of different sizes. As larger electrodes are prevalent in industrial processes, an in-depth understanding of similar phenomena at macroelectrodes is of technological importance. An attempt is made here to gain an understanding of this kind for the first time. With respect to the modeling, we follow the simple approach described in detail in Section 2, based on considering single bubble phenomena, and originating from the previous study of microelectrodes after accounting for differences in length scales and boundary conditions.

Figure 8 schematically displays important qualitative differences of the thermocapillary effect when the size 315 of the electrode is increased from micro (left) to macro (right) with respect to the bubble size. For the 316 sake of simplicity, at the macroelectrode only a small part of the electrode is shown, above which a single 317 bubble evolves. Looking at the distribution of the current density, shown by red dotted lines, important 318 differences become obvious. At the microelectrode, as discussed before, the current has to squeeze through 319 a narrow region at the bubble foot, and large current densities occur above the electrode, with a maximum 320 near its outer edge. In the center of the figure, the intermediate case of a large microelectrode is shown, 321 where, as discussed before, the temperature hotspot has already moved upward slightly towards a larger 322 value of θ . At the macroelectrode, however, the situation is changed, as the main geometrical obstacle for 323 a uniform current flow is now the bubble itself and not the electrode. Here, the current has to pass the 324 narrower equatorial space between neighboring bubbles. Hence, the maximum of the current density appears 325 at the bubble's equator. As the position of the temperature hotspot correlates directly with the location of 326 the maximum current density, the temperature maximum moves from the bubble foot towards the bubble 327 equator when the size of the electrode is increased from micro to macro. 328

The thermocapillary flow patterns driven at the bubble interface from hot towards cold regions are sketched in the right half of the subfigures for all three cases. In line with the changing position of the temperature hotspot, the size and the position of the two counter-rotating vortices driven by thermocapillary stress change accordingly. At the microelectrode, a dominating large vortex brings electrolyte upward along the interface, and the small vortex above the electrode does not contribute much to the thermocapillary effect. This small vortex grows in size when the size of the microelectrode is increased and starts contributing to the Marangoni force on the bubble, as already discussed in the previous section. At the macroelectrode, eventually, two counter-rotating vortices of almost equal size are therefore expected which advect electrolyte away from the temperature hotspot at the equator. The ramifications of these qualitative changes are investigated quantitatively in the following.

339 3.2.1. Temperature and velocity distribution

We now study in detail the temperature and velocity distribution near a gas bubble of constant size $(R_b = 560 \,\mu\text{m})$ at different values of the bubble coverage Θ of the macroelectrode. In the following, a potential difference of $\phi_0 = 1 V$ is applied between the electrodes. We note that, e.g. for a bubble coverage of $\Theta = 0.87$, an average current density of 4.9 kA/m² is obtained, which is close to typical values of 1–3 kA/m² in related industrial applications [30]. Figure 9(a) shows the temperature distribution in and around



Figure 9: ($\phi_0 = 1$ V and $\Theta = 0.87$) (a) Temperature contours and field lines of the electrical current density. Temperature is maximum at the location of maximum current density near bubble equator. (b) Contours of the electrolyte flow velocity distribution along with velocity streamlines. The electrolyte at the bubble interface flows away from the equator towards both poles.

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the gas bubble at a bubble coverage of $\Theta = 0.87$. As can be seen from the current lines in the electrolyte, the electric current must pass around the gas bubble and hence is highly concentrated in the narrow inter-bubble region near the bubble equator, where the largest current density occurs, in agreement with the expectations formulated in the previous section. The temperature maximum also occurs at the same spot, as can be seen from the temperature distribution. Figure 9(b) shows the velocity distribution and the streamlines in the electrolyte. The thermocapillary flow is driven away from the temperature hotspot, and the electrolyte flows away from the equator towards either pole. This gives rise to two vortices in the northern and southern hemisphere of the bubble and causes two interfacial velocity maxima, visible as the red regions in Figure 9(b). It is also to be noted here that the vortices are not fully symmetric with respect to the equatorial plane. The lower vortex is constrained by the electrode surface, the bubble interface and the outer boundary of the domain, whereas the upper vortex extends upward above the top of the bubble.



Figure 10: Interfacial (a) temperature and (b) velocity distribution as a function of θ as defined in Figure 1 for different electrode coverage Θ at $\phi_0 = 1$ V. The vertical dashed line marks $\theta = 90^{\circ}$.

356

³⁵⁷ Next we study in detail the influence of the bubble coverage on the interfacial temperature and flow ³⁵⁸ profiles. R_c is varied from 0.6 mm to 1 mm, yielding a range of Θ from 0.87 to 0.31, which, according ³⁵⁹ to earlier work, provides a realistic range of gas coverage values [45]. Figure 10(a) shows the interfacial ³⁶⁰ temperature profile as a function of the angular position θ . The occurrence of a temperature maximum ³⁶¹ close to the bubble equator ($\theta_m \sim 90^\circ$) is clearly visible in all cases. The temperature maximum also ³⁶² increases with increasing bubble coverage, as the electric current lines are more densely placed in the inter-³⁶³ bubble space, causing larger local Joule heating. Figure 10(b) shows the related interfacial velocity profile.

Similarly to the microelectrode case discussed in Section 3.1.2, the tangential velocity u_t vanishes at the 364 position of the temperature maximum at θ_m . According to the slopes of opposite sign in the temperature 365 profile left and right of θ_m , interfacial flow in the southern or northern hemisphere is directed downward 366 and upward, respectively, as also shown in Figure 10(b). In this figure, the value of zero tangential velocity 367 = 0 and the angular position of the equator $\theta = 90^{\circ}$ are additionally marked by lines to allow the more u_t 36 accurate identification of the location of the flow reversal (θ_m) in relation to the bubble equator. As can 369 be seen, the temperature hotspot is slightly shifted above the equator at smaller bubble coverage values. 370 Furthermore, it is to be noted that the magnitude of the velocity minimum in the southern hemisphere 371 is larger than the magnitude of the velocity maximum in the northern hemisphere in general. The latter 372 can easily be explained. As the metal electrode at the bottom is a good conductor of heat compared to 373 the electrolyte, the related downward heat flux tends to increase the temperature gradient in the southern 374 hemisphere, leading to a larger velocity magnitude. This larger velocity magnitude also causes enhanced 375 advection of heat at the interface, which leads to a shift in the temperature hotspot above the equator. 376 As increasing the bubble coverage narrows the gap between the bubble equator and domain boundary, the 377 temperature hotspot at the equator increases, and the relative influence of cooling and advection reduces. 378 This is also seen in the two extrema of the velocity profiles, which grow closer to each other in terms of 379 magnitude. 380

381 3.2.2. Marangoni force on the bubble

It is well known from measurements on large planar electrodes that the bubble evolution characteristics depend on the operating current density [45, 46]. At a higher current density, more nucleation sites are activated at the electrode, and thus the bubble coverage increases. Increasing the average current density also reduces the departure diameter and causes early detachment from the electrode surface [47]. In order to discuss the possible influence of thermocapillary effects on these phenomena, we carried out parametric studies to elaborate the influence of varying bubble coverage and bubble size on the Marangoni force under different operating conditions. The results are shown in Figure 11.

³⁸⁹ Constant bubble radius. In Figure 11(a) the Marangoni force normalized with the buoyancy force F_M^* (as ³⁹⁰ defined in Section 3.1.4) is shown versus the bubble coverage of the electrode Θ for a gas bubble of fixed ³⁹¹ size ($R_b = 560 \,\mu\text{m}$) at a constant cell voltage of $\phi_0 = 1 \, V$. It should be noted first that unlike the case of ³⁹² the microelectrode considered earlier, the Marangoni force is directed upwards, thus advancing the bubble ³⁹³ departure. The Marangoni force F_M^* is found to amount to about 2% of the buoyancy force in the range of ³⁹⁴ parameters considered and increases almost linearly with the bubble coverage Θ .

As the Marangoni force (F_M) results from the integration of the interfacial temperature gradient along the interface (see Equation 10), the slight asymmetry of the interfacial temperature profile with respect to

³⁹⁷ the equator seen in Figure 10(a) gives rise to a remaining nonzero force F_M . As the temperature gradient



Figure 11: Marangoni force normalized with buoyancy force F_M^* versus (a) bubble coverage Θ at a constant bubble radius $R_b = 560 \,\mu\text{m}$ and cell voltage $\phi_0 = 1$ V and versus (b) bubble radius R_b at a constant bubble coverage $\Theta = 0.87$ and mean cell current density $\bar{j} = 4.9 \text{ kA/m}^2$.

is negative in the region below the equator and larger in magnitude than the positive gradient above the equator, eventually an upward Marangoni force appears. The almost linear increase in the Marangoni force as the bubble coverage increases is related to the strong temperature increase of the hotspot near the equator in conjunction with the heatflux through the electrode, thus retaining a steep temperature profile below the equator.

⁴⁰³ Constant bubble coverage. In order to extend our study to include smaller gas bubble departure sizes, ⁴⁰⁴ Figure 11(b) summarizes the Marangoni force F_M^* obtained for different bubble radii R_b at a constant ⁴⁰⁵ bubble coverage of $\Theta = 0.87$ and at mean cell current density of $\overline{j} = 4.9 \text{ kA/m}^2$. We would like to point out ⁴⁰⁶ here that smaller bubbles have shorter residence times [47]. This was taken into account by adapting the ⁴⁰⁷ integration time in the simulations accordingly. The simulation details are summarized in Table 3. Starting out from an integration time of 1 s at $R_b = 560 \,\mu\text{m}$ in accordance with the constant bubble radius case considered earlier, the integration time is then linearly decreased along with the bubble size such that for a bubble of 200 μ m radius it matches the bubble lifetime of about 0.4 s reported in [48]. Please also note that for smaller bubbles the contact angle between the bubble and electrode was slightly increased. As can be seen in Figure 11(b), the Marangoni force F_M^* strongly increases when the size of the bubble reduces. This behavior is mainly caused by the corresponding variation in the buoyancy force, which scales with the cube

of the bubble size. For a small bubble of $R_b = 50 \,\mu\text{m}$, the Marangoni force is found to amount to about 20% of the buoyancy force.

$R_b \ (\mu { m m})$	$ heta_c$	Integration Time (s)
50	14.1°	0.15
100	9.9°	0.23
200	7.0°	0.4
300	5.7°	0.57
400	5.0°	0.73
560	4.2°	1.0

Table 3: Bubble radius, contact angle and integration time in the simulation

415

416 **4. Conclusions**

In this study we characterized in detail the thermocapillary flow around electrogenerated bubbles at 417 electrodes of different sizes, ranging from microelectrodes to large planar electrodes. At microelectrodes, a 418 detailed analysis of the current density and temperature profiles revealed a clear correlation between the 419 position of the temperature hotspot at the bubble interface and the upward projection of the outer edge 420 of the microelectrode. As a result, a double vortex structure of the thermocapillary flow was unveiled 421 which was overlooked previously [1] because the lower vortex is small at the microelectrodes. This double 422 vortex structure is also found at larger electrodes. Increasing the electrode size changes the position of the 423 temperature hotspot, thus increasing the size of the lower vortex and affecting the amplitude and also the 424 direction of the Marangoni force. 425

At large planar electrodes, multiple bubbles grow simultaneously at random nucleation sites where the electrical current must pass through the inter-bubble space. The simplified approach we take in order to gather initial results assumes that the bubbles grow synchronously at a given uniform nucleation density, meaning that a single bubble can only be considered in the simulations. As a result, the maximum current density and thus the temperature hotspot are located close to the equatorial inter-bubble region, and a nearly symmetric double vortex structure is generated near the bubble interface. However, mainly because

of the cooling effect of the metal electrode, the interfacial flow of the lower vortex is stronger than that of 432 the upper vortex. Consequently, an upward Marangoni force acts on the bubble, assisting its release, which 433 is the opposite of what is found at the microelectrodes. This clearly helps explain why larger bubbles can 434 be grown on macroelectrodes compared to microelectrodes. At a constant cell voltage, the Marangoni force 435 is found to grow as the bubble coverage of the electrode increases due to higher temperatures occurring in 436 the smaller inter-bubble gaps. For a fixed current density and related large bubble coverage, the force on 437 the bubble is found to get larger when the bubble size is reduced, reaching about 20% of the buoyancy force 438 at $\Theta = 0.87$, $\bar{j} = 4.9 \text{ kA/m}^2$. 439

We note here that the quantitative results obtained for the macroelectrode are based on the assumption of 440 uniform bubble distribution, i.e. simultaneous in-phase bubble evolution, and that azimuthal dependencies 441 of quantities originating from neighboring bubbles growing at certain angular positions have so far been 442 neglected. The Marangoni force values obtained for reasonable integration times in our model may further 443 depend on the specific value of the contact angle and neglect more complex aspects such as microbubbles 444 partly covering the cathode. It should also be noted that our results for all electrode sizes naturally depend 445 in quantity on the details of the thermal boundary conditions of the cell. This especially holds true for 446 the metal electrode where the bubble grows, which takes heat out of the electrolyte and thereby shapes the 447 thermocapillary effect. Nevertheless, as this is a common feature, we believe the qualitative findings of the 448 thermocapillary effect are valid in general. 449

In summary, the thermocapillary effect is found to be important for the dynamics of bubbles generated at electrodes under various conditions. A proper inclusion of this effect in future studies may lead to an improved understanding of the instant of the bubble departure and the departure diameter. Finally, appropriate temperature management of the electrodes may be a useful means of improving the efficiency of electrolyzers.

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