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Cell voltage model for Li-Bi liquid metal batteries

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Abstract

Lithium-bismuth bimetallic cells are amongst the best explored liquid metal batteries. A simple and fast one-dimensional cell voltage model for such devices is presented. The equilibrium cell potential is obtained from a complex twodimensional fit of data drawn from multiple studies of equilibrium cell potential and rendered congruent with the phase diagram. Likewise, several analytical and fit functions for the ohmic potential drop across the electrolyte are provided for different battery geometries. Mass transport overpotentials originating from the alloying of Li into Bi are modelled by solving a diffusion equation, either analytically or numerically, and accounting for the volume change of the positive electrode. The applicability and limitations of the model are finally illustrated in three distinct experimental settings.

Keywords: liquid metal battery, diffusion, volume change, cell voltage model

1. Introduction

The ever increasing deployment of highly fluctuating renewable energies requires stationary energy storage to balance energy production and consumption. Offering extreme current densities as well as an extended lifetime at a competitive price, liquid metal batteries (LMBs) have been discussed as an ideal

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candidate for large scale energy storage [1]. While the heavy positive electrode metal (e.g. Bi, cathode at discharge) forms the bottom layer, a light negative electrode metal (e.g., Li, anode at discharge) is usually soaked into a Ni foam – as illustrated in figure 1. Both liquid electrodes are separated by an ionconducting (e.g. Li⁺) molten salt mixture. At discharge, the negative electrode metal is oxidised, crosses the electrolyte layer in ionic form, is then reduced at the interface of the lower electrode and finally alloys with the positive electrode metal.



Figure 1: Sketch (not to scale) of an LMB with a metal foam negative electrode (left) and Ni foam filled with Li (right).

Between the many possible electrode-combinations, the Li||Bi cell is one of the best explored systems, as it provides unique benefits compared to other chemistries. Table 1 gives an overview on the cells and their properties built in the past. On the one hand, Li has a low solubility in its molten salts, which ensures a very high coulombic efficiency by reducing self-discharge [2]. On the other hand, the Li||Bi cell potentially can be operated in the two-phase area as well, which distinguishes it from many other systems, as, e.g., Na||Bi or Ca||Bi [2]. When Li is alloyed into Bi during discharge, the intermetallic phase Li₃Bi will form should a certain Li-concentration be exceeded locally. The intermetallic phase would then float on top of the Bi-electrode due to its lower density [2]. As Li₃Bi is an electric semiconductor and has a very high diffusivity for Li [2, 3], the cells can be operated in a semi-solid state enabling a very high Liutilisation of more than 90%. Although the intermetallic phase dissolves slightly

electrolyte	η	$\eta_{\rm C}$	life time	$T_{\rm O}$	price	j	U	$x_{\rm Li}$	${}^{h}\mathrm{E}$	source
		%	cycles	°C	\$/kWh	A/cm^2	v		mm	
				527			0.81	0.2		[11, p. 154]
				527			0.9			[9, p. 186]
							0.931			[12, p. 119]
LiCl-KCl				489			0.93		15-33	[11, p. 155]
LiCl-KCl				489		0.8	0.93	0		[13, p. 113]
LiCl-KCl				352			0.9			[14, p. 216]
LiCl-KCl				400		1.6	0.94			[15]
LiCl-KCl				550		6				[16]
LiCl-LiF				500-850			0.96	0.05		[17]
LiCl-LiF				500-850			0.96	0.05		[11, p. 109]
LiCl-LiF	70	99.7	1000	550	220-242	1.25	0.96	0	10	[2]
LiCl-LiF				500-850			0.96	0.05		[18]
LiCl-LiF				477-880			0.96	0.05		[12, p. 118]
LiCl-LiF				550		6				[16]
LiCl-LiF				510 - 560			0.96	0.03		[19]
LiCl-LiF				500-717						[20]
LiCl-LiF-LiI				380-485		2.2	1.4	0	3.4	[21]
LiCl-LiF-LiI				485	>1.8		1.25	0	3.4	[22, p. 187]
LiCl-LiF-LiI				500		2.2	0.9			[23, p. 111]
LiCl-LiF-LiI				460						[24]

Table 1: Properties of experimental Li||Bi cells: energy efficiency η , coulombic efficiency η_c , operating temperature T_o , maximum current density j, cell potential U at a Li molar fraction x_{Li} and electrolyte layer thickness h_{E} .

in molten salts, which is well visible due to its red colour [4–6] (see image in [7]), the solubility and irreversible Bi-transport to the positive electrode is rather low [8–10].

Based on the good availability of material properties and experimental data for Li||Bi cells, a number of modelling works have been performed for the system [25]. Especially fluid dynamic phenomena, such as the sloshing instability [26–28], electro-vortex flow [29], thermal convection [30–32], Marangoni convection [32], mass transfer [33–35] and solutal convection [24, 36, 37] have been studied. In addition, a one-dimensional electrochemical model for Mg-Sb and Li-Bi cells has been developed by Newhouse [38], three-dimensional models for mass-transport overpotentials by various authors [24, 39] and a 3D cell voltage model by Weber et al. [40, 41].

The aim of the present paper is to discuss and classify the various effects determining the cell voltage of Li||Bi LMBs in order to develop a one-dimensional, simple and fast cell voltage model, which accounts for all relevant effects.

2. Model

2.1. Overview

The one-dimensional model describes the cell voltage as function of current and time by first computing the equilibrium cell potential and then subtracting the various overpotentials. Moreover, a diffusion equation for the Liconcentration in Bi is solved in the positive electrode to obtain (after a conversion) the Li molar fraction, which is needed to calculate the equilibrium cell potential. Finally, volume changes of the positive electrode and electrolyte layer are taken into account when solving the diffusion equation and when determining the ohmic losses.

2.2. Open circuit potential

The equilibrium cell potential E_{eq} can be described by the Nernst equation as [1]

$$E_{\rm eq} = -\frac{RT}{\nu_e F} \ln(a_{\rm Li(Bi)}) \tag{1}$$

with R denoting the universal gas constant, T the temperature in K, ν_e the number of exchanged electrons and F the Faraday constant. Generally, the activity $a_{\text{Li(Bi)}}$ of Li in Bi or E_{eq} itself might be fitted, using e.g. data listed in table 5. We take the latter option leading to a molar fraction and temperature dependent equilibrium cell potential in the *liquid phase* of

$$E_{\rm lq}(V) = 10^{-3}(o + p\ln(x) + T(a\ln(x) + bx + cx^2 + dx^3 + ex^4 + fx\ln(x)))$$
(2)

with T in K and

$$o = 786.66, \quad p = -6.10, \quad a = -0.07, \quad b = 4.66,$$
 (3)

$$c = -16.50, \quad d = 28.96, \quad e = -23.01, \quad f = 1.75.$$
 (4)

The obtained fit function inter- and extrapolates the measurement values of [17, 19, 20], and is valid for temperatures between $415 \,^{\circ}\text{C}$ and approximately $600 \,^{\circ}\text{C}$, with higher errors up to $900 \,^{\circ}\text{C}$.



Figure 2: Phase diagram of Li-Bi according to [42] with different cell voltage areas and the region of validity shaded in green.

Once reaching the two-phase area (figure 2), the cell potential will stay constant, when forming the intermetallic phase Li_3Bi . The modelled equilibrium cell potential E_{eq} reads therefore:

$$E_{\rm eq} = \begin{cases} E_{\rm lq} & \text{in the liquid area L,} \\ E_{\rm lq}(x=0.01) & \text{in the liquid phase L for x < 0.01,} \\ E_{\rm lq}(T_{\rm lq}) & \text{in region IV,} \\ E_{\rm lq}(T_{\rm lq}) \left(1 - \frac{x - 0.73}{0.75 - 0.73}\right) & \text{in region V.} \end{cases}$$
(5)

Table 2: Phases present in the lithium-bismuth system.						
region	phase	state				
L	Li+Bi	liquid				
Ι	Bi+L	two-phase				
II	L+LiBi (low temp.)	two-phase				
III	L+LiBi (high temp.)	two-phase				
IV	$L+Li_3Bi$	two-phase				
V	$L+Li_3Bi$ (extended intermetallic)	two-phase				
VI	Li_3Bi+L	two-phase				
VII	L+Li	two-phase				
\mathbf{SI}	Bi+LiBi (low temp.)	solid solution				
SII	LiBi (low temp.)+ Li_3Bi	solid solution				
SIII	Li ₃ Bi+Li	solid solution				
SIV	LiBi (high temp.)+ Li_3Bi	solid solution				

with T_{lq} denoting the liquidus temperature in K, obtained from the phase diagram [43]; for the different phases/regions, see table 2.

Regions not included in the cell voltage model are the phases with high lithium fractions (x > 0.75) and the liquid and solid phases below 415 °C. While sufficient datapoints were available for higher temperatures ($450 \degree C$ to $900 \degree C$), only few datasets exist down to $380 \degree C$ for the two-phase region [44]. The lowtemperature liquid and bismuth-rich phases are not represented in available measurement data. Therefore, we model the electromotive force as follows in these phases:

$$E = 0$$
 in regions I-III, SI-SIV, VI and VII. (6)

The resulting equilibrium cell potential as function of the Li molar fraction and temperature is exemplarily illustrated in figure 3.



Figure 3: Equilibrium cell potential fitted with experimental data from [17, 19, 20].

2.3. Overpotentials

2.3.1. Activation losses

The activation overpotential η_{act} can be described by the Butler-Volmer equation as [45–47]

$$j = j_0 \cdot \left(\exp\left(\frac{\alpha n F}{RT} \eta_{\text{act}}\right) - \exp\left(-\frac{(1-\alpha) n F}{RT} \eta_{\text{act}}\right) \right)$$
(7)

with j denoting the current density, j_0 the exchange current density, α the charge transfer coefficient and n the reaction order of the rate limiting step. For a small overpotential, i.e. if

$$\eta_{\rm act} < {\rm R}T/{\rm F} \approx 66 {\rm mV} \quad ({\rm at} \ 500^{\circ}{\rm C}),$$
(8)

the Butler-Volmer equation can be linearised by approximating $\exp(y)$ as 1 + y, which leads to [45]

$$j = j_0 \frac{nF}{RT} \eta_{\text{act}}.$$
(9)

Newhouse measured concentration dependent exchange current densities at the positive electrode between 4 and 50 A/cm^2 [38, 48]. Using both limiting values, we find for a typical temperature of 500 °C an activation loss of

$$\eta_{\rm act} = \begin{cases} 0.0170 \cdot j \ (A/cm^2) & \text{for} \quad j_0 = 4A/cm^2 \\ 0.0013 \cdot j \ (A/cm^2) & \text{for} \quad j_0 = 50A/cm^2 \end{cases},$$
(10)

which would result in a voltage loss of 0.4...5 mV for a typical current density of 0.3 A/cm^2 and to 1.3...17 mV for a high current density of 1 A/cm^2 . Assuming that the exchange current densities are similar at the negative electrode, the activation losses appear to be minuscule compared to the ohmic overpotential. In line with Newhouse' conclusion [48], the activation losses are neglected in the present model.

2.3.2. Ohmic overpotential

The ohmic loss represents typically the most important overpotential of an LMB. In general, its value can either be measured or calculated. As the electrolyte conductivity is four orders of magnitude lower than those of the metals, only ohmic losses in the molten salt will be accounted for. In the most simple case, if positive and negative electrode have the same cross section area S, the voltage loss is simply

$$\eta_{\Omega} = IR = I \frac{H}{S\sigma} \tag{11}$$

with I denoting the cell current, R the electrolyte resistance, H its thickness, σ the electrical conductivity and S the surface area.

However, in real cells, often a metal foam is used to contain the molten Li. As illustrated in figure 4, the negative electrode has then a smaller diameter than the positive electrode. For a cylindrical cell or square cell, as shown in figure 4,



Figure 4: Sketch of a cylindrical (left) and rectangular (right) LMB using a metal-foam negative electrode current collector.

a simple Laplace equation can be solved in the electrolyte to obtain an estimate of the ohmic potential drop. Considering the comparably low conductivity of the salt, iso-potential boundary conditions are expected to provide the most realistic results. Solving this equation for a rotational-symmetric cylindrical geometry as well as a square cell with OpenFOAM [49] for a large parameter set and subsequent fitting, we obtain the ohmic overpotential as function of current, radii and layer height – as outlined in Appendix A.2 and Appendix A.3.

Alternatively, the Laplace equation can be solved analytically for a cylindrical cell using certain assumptions and simplifications – as explained in Appendix A.1. This leads to an explicit expression in terms of an infinite Fourier series:

$$\eta_{\Omega} = \frac{IH}{\sigma \pi R_{1}^{2}} + \sum_{n=1}^{\infty} \frac{IJ_{1}(\kappa_{0n})}{\sigma \pi \epsilon_{0n}^{2} R_{2}} \tanh\left(\frac{\epsilon_{0n}}{R_{1}}\frac{H}{2}\right) \frac{\pi H_{0}(\kappa_{0n}) J_{1}(\kappa_{0n}) + (2 - \pi H_{1}(\kappa_{0n})) J_{0}(\kappa_{0n})}{J_{0}^{2}(\epsilon_{0n})}$$
(12)

where J_0 and J_1 denote zero- and first-order Bessel functions of the first kind, H_0 and H_1 are zero- and first-order Struve functions and the numbers κ_{0n} are given by $\kappa_{0n} = \epsilon_{0n} R_2/R_1$, with ϵ_{0n} being the *n* roots of the first derivative of the zero-order Bessel function, see Appendix A.1 for more details. The solution appears to be rather intricate at first glance, but it allows for very fast calculations making it suitable for future integration into optimisation routines. Both the fitted and the analytical overpotential and corresponding relative errors are shown in figure 5 for different heights of the salt layer H and electrode sizes R_2/R_1 . The fitting functions lead to only very tiny errors, while the analytical formula deviates by up to 30% from the OpenFOAM simulations for very small H and R_2 . This was to be expected because iso-current boundary conditions had to be applied for the solution (12), whereas complementary iso-potential boundary conditions were used in the simulation. However, the introduced error is fairly low in a large parameter range such that the analytical solution still provides a good and fast estimate of the voltage loss. The comparison further reveals that the voltage drop is not very sensitive to the boundary conditions such that the exact current distribution in the metal foam plays only a minor role and does not necessarily have to be taken into account for the calculation of the overpotential.



Figure 5: Fitted ohmic overpotential for a cylindrical cell (a) and corresponding error (b), analytically computed ohmic loss (c) with error (d) and fitted ohmic overpotential for a square cell (e) with error (f). In the examples (a, c, e), $\sigma = 1$ S/cm, I = 1 A, $R_1 = 5$ cm and $L_1 = 10$ cm.

Finally, it should be noted that the salt layer thickness will vary considerably during operation, if the negative electrode metal is contained in a fixed metalfoam. This volume change needs to be accounted for as explained in section 2.6.

2.4. Diffusion equation

2.4.1. General considerations

The cell voltage of the LMB depends on the Li molar fraction at the positive electrode-electrolyte interface. In order to obtain the latter, a diffusion equation for the Li-concentration in Bi needs to be solved. According to Newhouse [38], the finite geometry of the positive electrode, its volume change and the concentration dependent diffusion coefficient need to be taken into account.

The general diffusion equation reads [50]

$$\frac{\partial c}{\partial t} = \nabla \cdot D \nabla c, \tag{13}$$

with D denoting the diffusion coefficient, t the time and c the concentration. It can either be solved for the molar (mol/m³) or for the mass concentration (kg/m³). However, when including convection in the cell voltage model, only the mass concentration is applicable any more, because the Navier-Stokes equations are written for a mass-averaged velocity. The density is strongly dependent on the concentration of Li in Bi. Therefore, the molar and mass averaged velocities are not equal – for a detailed explanation, see Levicky [51]. Finally, it is not admissible to solve directly for mass or molar fraction. The transport equation of the latter is derived by dividing Fick's law (equation 13) by the density. As the density changes continuously, when alloying Li into Bi, this operation is not permitted [52].

After solving the diffusion equation, the equilibrium cell potential can be obtained by converting the concentration to molar fraction as explained in section 2.5.

2.4.2. Analytical solutions

The diffusion equation (13) can be solved analytically for a semi-infinite geometry with the concentration flux q at the electrolyte interface as [38, p.83]

$$q = \nabla c \cdot \boldsymbol{n} = \frac{j}{\nu_e \mathrm{F} D},\tag{14}$$

where c denotes the molar concentration, \boldsymbol{n} the normal vector and ν_e the number of exchanged electrons. Using the approach of Carslaw [53, p. 75] with $f = F_0 =$ $Dq = j/(\nu_e F), K = D, \kappa = D$, we obtain the concentration at the interface as

$$c = c_0 + 2q \left(\frac{Dt}{\pi}\right)^{1/2} \tag{15}$$

with the initial concentration c_0 . The concentration inside the layer reads [53]

$$c = c_0 + 2q \left(\left(\frac{Dt}{\pi}\right)^{1/2} \exp(-z^2/(4Dt)) - \frac{z}{2} \operatorname{erfc} \frac{z}{2\sqrt{Dt}} \right),$$
(16)

with z denoting the vertical coordinate. By defining the integral of the error function ierfc as

$$\operatorname{ierfc}(z) = \frac{1}{\sqrt{\pi}} e^{-z^2} - z \cdot \operatorname{erfc}(z) \tag{17}$$

we obtain

$$c = c_0 + 2q\sqrt{Dt} \left(\text{ierfc} \frac{z}{2\sqrt{Dt}} \right), \tag{18}$$

and replacing z by H - z with H denoting the layer thickness finally leads to

$$c = c_0 + \frac{2j\sqrt{Dt}}{\nu_e FD} \left(\text{ierfc} \frac{H-z}{2\sqrt{Dt}} \right).$$
(19)

Here, the coordinate z runs from the interface downwards. This approach has already been used for LMBs by Newhouse [38] and Personnettaz et al. [24]. Of course, the simplification of a semi-infinite layer is only valid for thick layers or short charge/discharge times.

Alternatively, the diffusion equation can be solved analytically for a finite layer with the same boundary condition at the electrolyte interface as above, and $\nabla c \cdot \mathbf{n} = 0$ at the bottom [38]. Using the original solution for heat transfer [53, p. 112] and setting $\rho c_p = 1$, K = D, $\kappa = K/c_p = D$ and $F_0 = Dq = \frac{j}{\nu_e F}$ we obtain [54, p. 61][39]

$$c = c_0 + \frac{jH}{\nu_e FD} \left(\frac{Dt}{H^2} + \frac{3z^2 - H^2}{6H^2} - \frac{2}{\pi^2} \sum_{i=1}^{\infty} \frac{(-1)^i}{i^2} \exp(-Di^2 \pi^2 t/H^2) \cos\frac{i\pi z}{H} \right).$$
(20)

This approach has already been used for LMBs by Newhouse [38]; for alternative formulations of the same equation, see [38, 53].

2.4.3. Numerical solution

The analytical solutions of the diffusion equation are fast to compute, but neither account for the concentration dependent diffusion coefficient nor for volume change. Therefore, equation (13) is discretised using finite differences and finite volumes following Ferziger & Perić [55] as explained in detail in Appendix B.1 and Appendix B.2. The resulting equation system is solved in Python.

Volume change is accounted for by adjusting the cell volume or distance between the discretisation points in each time step. For this purpose, the initial masses in each control volume (CV) or at each point are computed, and then updated in every iteration. After computing the density using equation (35), the volume of each CV is obtained. The latter is used to compute a correction factor to increase either the size of the CV (finite volumes) or the distance between the points (finite differences). As the amount of moles of Li needs to stay constant during this operation, the concentration needs to be divided by the same factor, when increasing the volume.

2.5. Conversion between concentration and molar fraction

The equilibrium cell potential is given as function of the molar fraction (section 2.2). However, it is not possible to solve for the molar fraction directly – instead, the equations need to be written for molar or mass concentration (section 2.4). Hence, a conversion between concentration and fraction is required in both directions: from fraction to concentration to obtain the initial condition for the diffusion equation (13) and from concentration to molar fraction to obtain the cell voltage by equation (2).

The molar concentration of Li can be described as

$$c = \frac{n_{\rm Li}}{V} = \frac{x \cdot n}{V} \tag{21}$$

with $n_{\rm Li}$ denoting the amount of Li in mol, n denoting the total amount of Li and

Bi and V the volume. This gives with $n = m/M, V = m/\rho$ and $M = \sum_i x_i M_i$

$$c = \frac{\rho(x)x}{xM_{\rm Li} + (1-x)M_{\rm Bi}},$$
(22)

$$x = \frac{cM_{\rm Bi}}{\rho(c) - cM_{\rm Li} + cM_{\rm Bi}}.$$
(23)

As the density is usually only given as function of the molar fraction $\rho(x)$ (section 3.2), the second formula cannot simply be used. Converting concentration to molar fraction is possible using one of the following simplifications:

- Using a linear density law allows an arbitrary conversion between concentration and fraction without further simplifications. As the linear approximation holds only near a working point, it should be used for short charge/discharge times only. A good domain of application are complex three-dimensional simulations [41] together with the Oberbeck-Boussinesq approximation [56, 57].
- 2. The concentration of Bi might be assumed to be constant during the time of simulation. Then, the mole fraction is given as $x = c_{\text{Li}}/(c_{\text{Li}} + c_{\text{Bi}})$. Again, this simplification should be limited to short discharge times.
- 3. The density might be fitted twice once for concentration and once for mole fraction, as explained in section 3.2. Even though both fits will not be perfectly equivalent, this allows an easy conversion between c and x and back using formulae 22 and 23.
- 4. Assuming the density to be constant is a very strong simplification, limited to very short operation times.
- 5. The equilibrium cell potential might be directly fitted as function of the concentration. Although this approach seems easy, as it eliminates the molar fraction from all equations, it has one drawback: the phase diagram needs to be converted from x to c. In this step, it gets an additional dimension, because c depends on temperature, while x does not.

Within this work, we will use simplification (3) as it is the most accurate one for long discharge times.

2.6. Volume change

When the cell is discharged, and Li alloys into Bi, the positive electrode thickness will increase. This volume change needs to be accounted for, when solving the diffusion equation [38]. In most of the LMBs built in the past, the Lithium was contained in a metal-foam current collector, mounted at a fixed position. If in these cells the positive electrode layer changes its thickness, the electrolyte layer height will change at the same time. Considering the high ohmic resistance of the molten salt, the volume change of the electrolyte might be a very important effect.

The change of height of the positive electrode layer during operation can be estimated as

$$\Delta H = \frac{x \frac{n_{\rm Bi}}{1-x} M_{\rm Li} + m_{\rm Bi}}{S\rho(x)} - \frac{x_0 \frac{n_{\rm Bi}}{1-x_0} M_{\rm Li} + m_{\rm Bi}}{S\rho(x_0)} \tag{24}$$

with x_0 denoting the initial Li molar fraction, m the mass and S the surface area. Figure 6 gives an illustrative example for the large Li-Bi cell experiment, described in section 4.3. There, the positive electrode layer changes between charged and discharged state between a thickness of 2.7 and 6.5 mm, i.e. roughly by a factor of two. The volume change is considerable – and needs to be accounted for when solving the diffusion equation, but also when computing the ohmic overpotential.



Figure 6: Thickness of the positive electrode layer as function of the Li molar fraction in Bi for the experiment described in section 4.3.

An elegant way to account for volume change in LMBs when solving the diffusion equation has been introduced by Newhouse [38]. She solved the diffu-

sion equation in a solvent-fixed reference frame, which means that no net flow of Bi (the solvent) occurs over the single control volumes [58–60]. This approach involves introducing a new spatial unit ξ , which describes the distance between two solvent atoms, and to convert the diffusion coefficient to this reference frame. After solving the modified diffusion equation, and obtaining the Li-flux relative to the Bi-atoms, the solvent-fixed concentration needs to be reconverted to the "true" concentration. This approach is based on the original article by Hartley & Crank [61] (available also in [54]), which is very well explained by Gekas & Lamberg [62] and has been used for other applications, as well [63, 64]. Apart from the difficult derivation, the biggest drawback of this approach is the assumption of constant partial volumes – i.e. the assumption that the alloy density can be described as that of an ideal solution (equation 30).

An alternative approach is to compute every single cell volume in each time step, when solving the diffusion equation. Then, the size of the control volume can be updated in real-time. When solving the diffusion equation in 1D, as explained in Appendix B.1 and Appendix B.2, the distances between two nodes need to be scaled, respectively.

3. Material properties

3.1. Conductivity

The electrical conductivity of the molten salt is required to compute the ohmic losses. In the past, mostly mixtures of LiCl-LiF [2, 11, 12, 17, 18] and LiCl-LiF-LiI [21–23] have been used for Li-Bi cells. LiCl-KCl was employed as well [11, 13–15], although it is known that Li reduces KCl at temperatures above 500 °C [17, 65]. Table 3 gives an overview on the melting temperature and conductivity of the salts.

Table 3: Electric conductivity σ and melting temperature ϑ_m of the molten salts.

salt	composition in mol%	ϑ_{m} in $^{\rm o}{\rm C}$	σ in S/cm	source
LiCl-LiF	70:30	501		[2]
			6.78 at 927 $^{\rm o}{\rm C}$	[66]
	various			[67]
LiCl-LiF-LiI	eutectic	340.9	3 at 475 $^{\rm o}{\rm C}$	[21]
	eutectic	340.9	2.3 at 475 $^{\rm o}{\rm C}$	[68, p. 97]
	29.1:11.7:59.2	340.9	2.3 at 375 $^{\rm o}{\rm C}$	[21]
	29.1:11.7:59.2	341	2.3 at 375 $^{\rm o}{\rm C}$	[22, p. 167]
	$29.1{:}11.7{:}59.1$	341	$8.895 \exp(-872.6/T)$	[69]
LiCl-KCl		354	1.83 at 500 $^{\rm O}{\rm C}$	[70]
	40.45:59.55		$13.21 \exp(-13995.71415/RT)$	[71, p. 205]
	41:59	353	1.7 at 476 $^{\rm o}{\rm C}$	[1]
	58.8:41.2	353	1.57 at 450 $^{\rm O}{\rm C}$	[22, p. 167]
	58.8:42.2		$18.7876 \exp(-1800.6/T)$	[69]
	58.8:41.2		$23.021 \exp(-16204.90311/RT)$	[71, p. 205]
	70.36:29.64	-1101.08 -	$+2.7461T - 13.2471 \cdot 10^{-4}T^{2}$	[72, p. 1045]
	81.77:18.23		$13.886 \exp(-9982.35421/RT)$	[71, p. 205]
	various			[72, p. 1045]
	various			[73]

3.2. Density

The densities of molten Li and Bi are given in kg/m^3 as [74, p. 14-10]

$$\rho_{\rm Li} = 518 - 0.1(T - 453.5) \tag{25}$$

$$\rho_{\rm Bi} = 10050 - 1.18(T - 544) \tag{26}$$

with T denoting the temperature in K. Densities of binary alloys are sometimes estimated based on Vegard's law [75]. The latter predicts a linear relation of the mean lattice distance in a solid solution, if two different components are mixed. Applied to fluid mixtures, it is assumed that the volume V can be computed as a linear weight of the amounts of the substances n_i and the third roots of their molar volumes v_i as [76]

$$V = \sum \left(n_i \sqrt[3]{v_i} \right)^3, \tag{27}$$

which leads for a binary alloy of component 1 and 2 to [76]

$$\rho = \frac{x_1 M_1 + (1 - x_1) M_2}{\left(x_1 \sqrt[3]{M_1/\rho_1} + (1 - x_1) \sqrt[3]{M_2/\rho_2}\right)^3}.$$
(28)

Alternatively, the volume of an *ideal solution* can be described as the sum of its components as

$$V = \sum n_i v_i, \tag{29}$$

which leads to the very similar equation [77]

$$\rho = \frac{x_1 M_1 + (1 - x_1) M_2}{(x_1 M_1 / \rho_1 + (1 - x_1) M_2 / \rho_2)}.$$
(30)

Both density laws are not well suited for Li-Bi, as can be seen in figure 7. The deviation from Vegard's law perhaps can be explained by electronic structure. While Li is strongly electropositive ($\chi = 0.98$), Bi is considerably more electronegative ($\chi = 1.9$). This difference might cause a deviation from ideality for the liquid solution – similar as the Hume-Rothery Rules predict for solid state solutions a deviation from ideality for large electronegativity differences.

Only few density values for the Li-Bi alloy are available in the literature [78–82]. These data are used to fit the density as

$$\rho = \frac{xM_{\rm Li} + (1-x)M_{\rm Bi}}{(x^{0.94}M_{\rm Li}/\rho_{\rm Li} + (1-x)^{1.64}M_{\rm Bi}/\rho_{\rm Bi})}.$$
(31)

As illustrated in figure 7, this fitted function is describing the Li-Bi density best, and is therefore used in the following.

In order to allow an easy conversion between concentrations and molar fractions, the density is fitted as function of the concentration, as well. For this purpose, the molar fractions belonging to the measured density values are first converted to the Li molar concentrations as

$$c = \frac{\rho x}{x M_{\rm Li} + (1 - x) M_{\rm Bi}} \tag{32}$$

with ρ denoting the mixture density. The resulting fit function fulfils the boundary conditions

$$\rho(c=0) = \rho_{\rm Bi} \tag{33}$$

$$\rho(c = c_{\infty}) = \rho_{\rm Li} \tag{34}$$

and reads

$$\rho = \frac{\alpha \cdot c^{\gamma} + \beta \cdot (c_{\infty} - c)^{\delta}}{\alpha \cdot \frac{c^{\gamma}_{\infty}}{\rho_{\mathrm{Li}}} \left(\frac{c}{c_{\infty}}\right)^{\varepsilon} + \beta \cdot \frac{c^{\delta}_{\infty}}{\rho_{\mathrm{Bi}}} \left(\frac{c_{\infty} - c}{c_{\infty}}\right)^{\varphi}}$$
(35)

with the concentration of pure Li defined as

$$c_{\infty} = \frac{\rho(x=1)}{M_{\rm Li}} = \frac{\rho_{\rm Li}}{M_{\rm Li}}$$
(36)



$$\varepsilon = 1.45, \ \varphi = 0.67, \ \alpha = 8.1 \cdot 10^{-9}, \ \beta = 37, \ \gamma = 2.37 \text{ and } \delta = 0.7.$$
 (37)

Figure 7: Density of Li-Bi depending on the Li molar fraction (a) and Li concentration (b).

3.3. Diffusion coefficient

and

Diffusion coefficients of Li in Bi have been measured by Temnogorova, Weppner, van Norman and Newhouse, as shown in table 4 and figure 8. Due to the available concentration dependence, we use the formula of Newhouse.

Table 4: Diffusivity of Li in Bi.					
$D \text{ in } \text{cm}^2/\text{s}$	ϑ in $^{\circ}\mathrm{C}$	source			
$2.2 \cdot 10^{-5}$	450	[83]			
$1.56 \pm 0.15 \cdot 10^{-5}$	500	[15]			
$3.25 \pm 0.53 \cdot 10^{-5}$	550	[15]			
$3.27 \pm 0.3 \cdot 10^{-5}$	550	[15]			
various (Li_3Bi)	360-600	[3]			
$\exp\left(\frac{-4.081c - 0.01315}{c^2 + 0.3742c + 0.001572}\right)$	450	[38, p. 174]			

3.4. Thermodynamic properties

The thermodynamics of the Li-Bi system have been studied several times – table 5 gives an overview on literature sources for the activity $a_{\text{Li(Bi)}}$, the Gibbs free energy G, the equilibrium cell potential (E_{eq}) , the enthalpy H and the entropy S.

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Figure 8: Concentration dependent diffusivity according to various authors and fit by Newhouse [38]; x has been converted to c using equations (31) and (32).

4. Application of the model

4.1. Small Li-Bi cell

The first use case for the developed model is illustrated in figure 9 – a small Li-Bi cell with a diameter of approximately 3 cm. The Li-metal is contained in spirally rolled Ni-sheet serving as current collector and separated from the positive electrode by an effectively 3-4 mm thick eutectic LiCl-LiF-LiI electrolyte.



Figure 9: Setup and dimensions (in mm) of the small Li-Bi laboratory cell (not to scale). The thicknesses of positive electrode and electrolyte are given for the start of the charge phase (figure 10). As the electrolyte thickness can not easily be measured, it has been derived from the known geometry and conductivity.

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Table 5: Sources for thermodynamic properties of the Li-Bi system: activity $(a_{\text{Li(Bi)}})$, Gibbs free energy (G), equilibrium cell potential (E_{eq}) , entropy of mixing (S) and enthalpy of mixing (H).

phase diagram	^a Li(Bi)	G	E_{eq}	S	H	literature
•	•	•		•	•	[84]
			•			[5, p. 152]
		•				[12, p. 118]
			•			[14, p. 216]
•						[2]
			•			[18, p. 226]
•		•	•			[17]
•		•	•	•	•	[20]
			•			[85]
•	•	•		•	•	[86]
	•	•	•	•	•	[19]
			•			[11, p. 109f]
			•			[87, p. 143f]
			•			[38, p. 110f]
			•			[44]
•	•				•	[42]
	•					[83]
	•		•			[88]

The metals (Li: Alfa Aesar, 99.9%; Bi: Alfa Aesar, 99.998%) were first cleaned mechanically (oil removal and skin cut off for Li), before melting them and skimming floating pollutants. Additionally, a small amount of salt was added to bind remaining impurities. After freezing the contaminated salt, the clean metal was poured into a second crucible. To prepare the electrolyte, LiCl (Alfa Aesar, 99.995%) and LiF (Beantown, 99.99%) were mixed and allowed to rest for several hours at 550 °C. LiI (Beantown, 99.95%) was added thereafter, and the complete mixture filtered though a glass frit.

The tantalum crucible was filled with 0.1 mol Bi and 18.5 g electrolyte. Finally, molten Li was soaked into the negative electrode current collectors (Ni alloy 200). The cell was operated on a ceramic heating plate (BACH Resistor Ceramics) in a glove box filled with argon gas (H₂O < 0.1 ppm, O₂ < 0.1 ppm) at a heating plate temperature of 460°C [24].

The modelling work starts with the initial condition: the known mass of Bi, and the measured equilibrium cell potential lead to the initial Li-concentration in Bi. Then, the amount of Li, which leaves the positive electrode during charge, is calculated. Considering the strong solutal convection during charging, perfect



Figure 10: Measured charge-discharge curve at 1 A and modelled curves using the analytical solution for a finite and semi-infinite layer and the numerical solution with variable diffusivity and volume change.

mixing is assumed [24]. The ohmic loss for charging is obtained for each timestep using equation (A.11) and taking the change of the electrolyte layer height into account.

Switching to discharging, the current direction is reversed and Li transferred back to the positive electrode. For comparison, the diffusion equation is solved once by the analytical solutions for a finite and semi-infinite layer and once numerically. The latter accounts for volume change of positive electrode and electrolyte as well as the variable diffusivity, while the analytical solutions neglect volume change at all and use a constant diffusion coefficient of $4 \cdot 10^{-5} \text{ cm}^2/\text{s}$.

Comparing the measured and modelled curves, it becomes apparent that an approximate reproduction of experimental curves is easy. Already Personnettaz et al. [24] reproduced the same experiment using an estimated diffusion coefficient of $7 \cdot 10^{-5}$ cm²/s and a different formulation for the mixture density. Obtaining a perfect match is, however, challenging. For example, the small deviation of the equilibrium cell potential before charge/discharge is probably caused by the fit function (equation 2). As no measured electromotive force values were available for 460 °C, the fit had to be slightly extrapolated. The small mismatch of the cell potential during charge is surely related to the ohmic loss. While the positive electrode layer changes its height by 5% during cycling, it is not easy to predict the exact shape of the negative electrode. The Li might be soaked fully into the Ni-sheet or form a large droplet below it. The shape of this droplet might vary due to the Lorentz forces at charge/discharge and also due to the transfer of Li-metal [89]. Finally, the slight deviation of the discharge curves might be related to the 3-dimensionality of the diffusion problem. While the model assumes only downward diffusion, Li will in reality also diffuse sidewards in the positive electrode – which then increases the cell voltage. Moreover, any very small flow effects, caused, e.g. by Marangoni convection or electro-vortex flow, might lead to a higher cell potential, too. Anyway, the numerical solution is extremely close to the measured cell potential.

4.2. Large Li-Bi cell

Highlighting the broad applicability of the developed model, a second, larger Li-Bi cell will be studied. Now, the cell diameter is 9 cm, with the negative electrode metal being contained in a Ni foam as shown in figure 11.



Figure 11: Setup and dimensions (in mm) of the large Li-Bi cell (not to scale). The layer thicknesses are given for the start of discharge in figure 12; as the electrolyte thickness was not accessible to a direct measurement, it has been derived from the known geometry and conductivity.

Metals and salts were cleaned as for the small cell (section 4.1). The battery was set up by pouring 1 mol Bi into the tantalum vessel and adding 80.28 g eutectic LiCl-LiF-LiI electrolyte. The negative current collector (Ni foam, Recemat BV) was prepared by heating 1.5 mol Li in a stainless steel vessel to 450 °C and letting the foam rest for 2 h in the bath. Finally, the cell was operated such that the heating plate obtained a temperature of 500 °C, while the salt reached only 420 °C.



Figure 12: Experimental and numerical charge-discharge curve for the large Li-Bi cell at 5A.

The charge/discharge curve is modelled exactly like for the previous experiment: the initial values are obtained from the measured electromotive force and the positive electrode is assumed to be well mixed at charge. Volume change is accounted for during charge, but during discharge only when solving the diffusion equation numerically. As the cell is operated with a Li-concentration of 1 to 5 kmol/m³, a diffusion coefficient of $7 \cdot 10^{-5}$ cm²/s is assumed when using the analytical solutions of the diffusion equation.

The model reproduces the experimental data very well – with a certain deviation during the discharge phase. Compared to the small Li-Bi cell experiment described in section 4.1 the charge phase is reproduced better. The reason is most likely that the negative electrode shape is now defined much better, as the Li is contained in a metal foam instead of a rolled Ni-sheet. The slight deviation during cell discharge can most probably be attributed to the threedimensionality of the problem. As the positive electrode diameter is three times larger than the negative electrode, substantial lateral diffusion will occur, which surely influences the cell voltage. Nevertheless, the experimental curve is reproduced reasonably well.

4.3. Li-Bi cell by Ning et al.

The last test case is a large Li-Bi cell built at Massachusetts Institute of Technology and published by Ning et al. [2]. The setup is shown in figure 13: the cell is 15 cm in diameter, containing 455 g of Bi, which results in a 2.7 mm thick positive electrode layer. The Li-negative electrode is soaked into a Ni foam, and is assumed to be 10 cm in diameter. The cell operates at $550 \,^{\circ}\text{C}$ using an eutectic LiCl-LiF (70:30 mol%) electrolyte with an interelectrode distance of 10 mm in charged state. Having a capacity of 175 Ah, 3 h and 20 min are theoretically needed to discharge the complete cell at a current density of 300 mA/cm².



Figure 13: Setup and dimensions (in mm) of the Li-Bi cell investigated by Ning et al. [2] (not to scale).

Figure 14 illustrates the discharge curves of this cell using different models. The equilibrium cell potential (equation 5) – which would be obtained at very small current densities – clearly exhibits a long plateau in the two-phase area. When reaching a Li molar fraction of about 0.73, there is a predicted extended intermetallic phase region [90]. In our model, the potential in this region decreases linearly to zero.

Discharging with a current density of 300 mA/cm^2 , the numerical solution of the diffusion equation gives the most accurate results as it accounts for volume change as well as the concentration dependent diffusivity. As expected, the

ohmic overpotential shifts the cell potential to lower values. Moreover, the usable cell capacity is reduced by roughly 30%, as well. The reason is simply that due to the high current, a layer of saturated Li₃Bi forms at the electrolyte-positive electrode interface, which causes the cell potential to drop to 0V. The "missing" capacity is located just below of this intermetallic layer, where a stoichiometric composition of Li₃Bi is not reached. The usable cell capacity depends therefore not only on the mass of the active materials and side reactions, but also on the discharge current.

Modelling the discharge curve with the analytical solution of the diffusion equation for a finite layer (equation 20) using an average diffusivity of $D = 8 \cdot 10^{-5} \text{ cm}^2/\text{s}$ [41], the predicted capacity is only 50% of that found by the numerical solution. This considerable discrepancy is simply caused by the fact that volume changes are not considered. As the analytical solution of the diffusion equation neglects the fact that the positive electrode thickness changes approximately by a factor of two (see figure 6) between charge and discharge, it predicts a much lower capacity.

The analytical solution of the diffusion equation for a semi-infinite layer does the opposite: it predicts a much larger cell capacity than theoretically possible. As it ignores the lower boundary of the positive electrode, it assumes that much more Li can be alloyed into Bi than practically possible.

These examples highlight that the analytical solutions for the diffusion equation should not be used when considering full charge-discharge cycles as they induce considerable errors due to the negligence of volume-change effects.

5. Limitations of the model and possible improvements

Modelling the cell voltage of LMBs is a complex task. In the opinion of the authors, the presented model and its portability to other geometries and cell chemistries is mainly limited by the following points:

1. Material properties: measured densities of typical LMB electrode couples are usually scarce. Moreover, diffusivities are not always available with



Figure 14: Discharge curves for 0.3 A/cm² using different numerical and analytical models.

their concentration dependence, and the measurement error may be considerable.

- 2. Complexity of intermetallic phases: the conductivity of the Li-Bi alloy changes by almost two orders of magnitude during alloying, reaching the same order of magnitude as for the molten salt, when forming the intermetallic phase [78, 91, 92]. The corresponding ohmic losses might be included in improved models. Likewise, the diffusivity of Li in Bi changes by a factor of up to 3, when forming the intermetallic. This makes accurate predictions of mass transport challenging.
- 3. Ohmic overpotential: the ohmic losses in the salt can usually not be approximated by a one-dimensional model. The fit equation, used here, might be replaced alternatively by solving a Laplace equation for the electric potential in 2 or 3D.
- 4. One-dimensionality of mass transport: mass transport has been modelled as a one-dimensional effect. This simplification is often appropriate – as long as the negative electrode and positive electrode diameters are similar (see section 4.2 for an example, where this was not the case).
- 5. Simplification of flow effects: any type of flow can increase the diffusion

coefficient "artificially" [41]. This effect has been neglected. The positive electrode is simply be assumed to be perfectly mixed by solutal convection at charge [24], while mass transfer is assumed to be controlled by diffusion only at discharge. One example, where this does not work well, is shown in figure 15(a). When charging the large Li-Bi cell described in section 4.2 with a very large current (here 20 A), a Li-depleted concentration layer will form in the positive electrode despite most of the volume is well mixed by solutal convection. The measured cell potential will rise therefore steeply, while the modelled potential is lower.

6. Memory effect: the cell potential is not only a function of the state of charge, but is history-dependent as well. Figure 15(b) shows one example: the measured cell potential of the first cycle is larger than of the two subsequent ones. At the beginning of cycling, the positive electrode was at rest and perfectly mixed. Therefore, solutal convection sets in immediately at charge, mixing the alloy well, and leading to a high cell voltage. After discharge, a stable density stratification formed. When now charging again, solutal convection takes longer to destroy the stable density stratification and to mix the positive electrode again – therefore, the potential of the second and third cycle is initially lower. Such "memory effects" can generally be accounted for using the developed model; however, care needs to be taken not to ignore them.

6. Summary

The Lithium-Bismuth system is one of the best explored liquid metal batteries (LMBs). After discussing the peculiarities of this chemistry and summarising the previous experimental studies, we gave a short overview of the most relevant modelling work of LMBs. Further, we briefly described our motivation to develop a simple one-dimensional cell voltage model.

The equilibrium cell potential is obtained by a complex two-dimensional fit of experimental data as function of temperature and Li-molar fraction. While



Figure 15: Experimental and modelled charge/discharge curve for the large Li-Bi cell presented in section 4.2 at 20 A (a) and overlay of the first three experimental cycles at 10 A (b).

the very small activation losses have been neglected, different sub-models for the ohmic loss have been proposed. It was shown that the voltage drop in the salt can be solved analytically, but only under certain assumptions. As alternative, two fit functions for cylindrical and rectangular cells are provided, which describe the ohmic losses well.

Special care was taken to obtain the correct mass transfer overpotential. After discussing the possible formulations of the diffusion equation to obtain the Li-content in Bi, different possibilities to convert molar and mass fraction to molar or mass concentration were proposed. Two analytical solutions for the diffusion equation were presented: one for a semi-infinite, and one for a finite positive electrode. Finally, a numerical scheme was implemented in the finite difference and finite volume method, which accounts for volume change of the positive electrode during operation.

The developed model was finally applied to three different Li||Bi cells. Although the results generally match well with the experimental data, the comparison revealed certain limitations of the model. Most importantly, the onedimensional model fails in describing three-dimensional mass transfer effects, which become important if the negative electrode and positive electrode diameter deviate strongly from each other. Moreover, the influence of flow effects on the cell voltage has been simplified to a large extend. The modelling of the three experiments further highlighted that for short-time cycling the analytical solutions of the diffusion equation are well suited. However, in order to describe the full discharge of an LMB, volume change is of highest importance, and needs to be included as well by solving the mass transport equation numerically.

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Appendix A. Ohmic overpotential

Appendix A.1. Analytical solution for cylindrical geometries

We calculate the potential distribution in the electrolyte for the left asymmetric cylindrical cell shown in Figure 4. Since the electrical current can be uniquely described by a potential through Ohm's law $\boldsymbol{j} = \sigma \nabla \phi$ and must be closed $(\nabla \cdot \boldsymbol{j})$, we need to solve the two-dimensional Laplace equation

$$\Delta \phi = \frac{\partial^2 \phi}{\partial r^2} + \frac{1}{r} \frac{\partial \phi}{\partial r} + \frac{\partial^2 \phi}{\partial z^2} = 0 \tag{A.1}$$

in the electrolyte's domain, where we have applied cylindrical coordinates (r, φ, z) with the origin located in the centre of the salt-negative electrode interface. For the boundary condition, we assume that the sidewalls are perfectly insulating and that we have a homogeneous vertical current distributions in the positive electrode, yielding

$$\left. \frac{\partial \phi}{\partial r} \right|_{r=R_1} = 0, \tag{A.2}$$

$$\left. \frac{\partial \phi}{\partial z} \right|_{z=-H} = \frac{I}{\sigma \pi R_1^2},\tag{A.3}$$

$$\left. \frac{\partial \phi}{\partial z} \right|_{z=0} = \frac{I}{\sigma \pi R_2^2} \Theta(R_2 - r),$$
 (A.4)

where I denotes the total cell current and Θ is the Heaviside function ensuring that no current bypasses the negative electrode in the region $r > R_2$. Please note that the current distribution is usually not homogeneous, in particular not in the negative electrode foam if its radius R_2 is considerably smaller than the cylinder radius R_1 . However, we seek for an approximate solution of the cell voltage to be expressed analytically, requiring to keep the boundary conditions as simple as possible. This approximation is *a posteriori* justified in figure 5, where we discuss the relative error following from this approximation. The cylindrical boundary value problem (A.1) - (A.4) is solved by expanding a Fourier-Bessel series. We take the ansatz

$$\phi = -\frac{I}{\sigma \pi R_1^2} z + \sum_{n=1}^{\infty} a_n \cosh\left(\frac{\epsilon_{0n}}{R_1}(z+H)\right) J_0\left(\epsilon_{0n}\frac{r}{R_1}\right), \qquad (A.5)$$

with J_0 being the zero-order Bessel function of the first kind and the numbers ϵ_{0n} given as the *n* roots of the first derivative of the zero-order Bessel function

$$J_0^{\prime}(\epsilon_{0n}) = 0, \tag{A.6}$$

which can be easily determined numerically or found in, e.g., [93]. The coefficients a_n are to be determined by applying condition (A.4); ansatz (A.5) already fulfils the simple boundary conditions (A.2) and (A.3). Inserting ansatz (A.5) into (A.4), multiplying both sides of the equation by $rJ_0(\epsilon_{0m}r/R_1)$ and eventually integrating the equation over the cylinder radius R_1 , we find explicit expressions for the coefficient a_n as

$$a_n = \frac{2IJ_1\left(\epsilon_{0n}\frac{R_2}{R_1}\right)}{\sigma\pi R_2 \sinh\left(\frac{\epsilon_{0n}}{R_1}H\right)\epsilon_{0n}^2 J_0^2(\epsilon_{0n})}.$$
(A.7)

For the calculation, we have exploited the orthogonality condition

$$\int_0^{R_1} r J_0\left(\epsilon_{0m} \frac{r}{R_1}\right) J_0\left(\epsilon_{0n} \frac{r}{R_1}\right) \mathrm{d}r = \frac{1}{2} R_1^2 J_0^2(\epsilon_{0n}) \delta_m n.$$
(A.8)

Inserting (A.7) into (A.5) finally yields the potential solution

$$\phi = -\frac{I}{\sigma \pi R_1^2} z - \sum_{n=1}^{\infty} \frac{2I J_1\left(\epsilon_{0n} \frac{R_2}{R_1}\right)}{\sigma \pi \epsilon_{0n}^2 R_2} \frac{\cosh\left(\frac{\epsilon_{0n}}{R_1}(z+H)\right)}{\sinh\left(\frac{\epsilon_{0n}}{R_1}H\right)} \frac{J_0\left(\epsilon_{0n} \frac{r}{R_1}\right)}{J_0^2\left(\epsilon_{0n}\right)}.$$
 (A.9)

The solution contains the voltage loss expected to occur in the electrolyte. It is defined as the potential difference between negative and positive electrode, but we must take into account that the potential is not constant along the negative electrode foam. To approximate the global cell voltage, we consider the mean potential in the negative electrode (the positive electrode potential is approximately constant), so that the voltage drop is given as

$$\eta_{\Omega} := \frac{1}{R_2} \int_0^{R_2} \phi(r, z = -H) - \phi(r, z = 0) dr$$

$$= \frac{IH}{\sigma \pi R_1^2} + \sum_{n=1}^{\infty} \frac{IJ_1(\kappa_{0n})}{\sigma \pi \epsilon_{0n}^2 R_2} \tanh\left(\frac{\epsilon_{0n}}{R_1} \frac{H}{2}\right) \frac{\pi H_0(\kappa_{0n}) J_1(\kappa_{0n}) + (2 - \pi H_1(\kappa_{0n})) J_0(\kappa_{0n})}{J_0^2(\epsilon_{0n})}$$

(A.10)

with $\kappa_{0n} = \epsilon_{0n}R_2/R_1$. For the calculation we have applied the hyperbolic identity $\operatorname{coth}(x) - \sinh(x)^{-1} = \tanh(x/2)$. H_0 and H_1 refer here to the Struve functions of zero and first order, see [93], which result from the integration of J_0 . The first term in (A.10) describes the homogeneous voltage for the trivial case $R_1 = R_2$. The following Fourier sums correct this solution to account for the inhomogeneities caused by the asymmetry of positive and negative electrode $R_2 < R_1$. The convergence behaviour of the series strongly depends on the ratio R_2/R_1 . For point-like negative electrodes $R_2 \ll R_1$ the solution converges very slowly but correspondingly fast if the electrodes are in the same order. For all the practical cases shown in figure 5 it was sufficient to keep the first 400 terms allowing for a very fast calculation.

Appendix A.2. Numerical solution for cylindrical geometries

The voltage loss in the electrolyte of a cylindrical cell, as illustrated in figure 4 is

$$\eta_{\Omega} = I \frac{0.05^2}{\sigma R_1^2} (a + a_1 A + a_2 A^2 + a_3 A^3 + a_4 A^4 + a_5 A^5 + a_6 A^6 + b_1 H + b_2 H^2 + b_3 H^3 + b_4 H^4 + b_5 H^5 + b_6 H^6 + d_1 A H + d_2 (A H)^2 + d_3 (A H)^3 + d_4 (A H)^4 + d_5 (A H)^5 + e_1 H / A + e_2 H^2 / A + e_3 H A^2 + e_4 A H^2 + e_5 A H^3),$$
(A.11)

with the ratio of the radii defined as $A = R_2/R_1$ and

$$a = 12.5, \ a_1 = -105, \ a_2 = 358, \ a_3 = -652 \ a_4 = 676, \ a_5 = -378, \ a_6 = 89,$$

$$b_1 = 214, \ b_2 = -5.8 \cdot 10^4, \ b_3 = 3.3 \cdot 10^6, \ b_4 = -8 \cdot 10^7, \ b_5 = 1.3 \cdot 10^9, \ b_6 = -2.8 \cdot 10^9,$$

$$d_1 = -1277, \ d_2 = -4507, \ d_3 = -4.3 \cdot 10^6, \ d_4 = 2.4 \cdot 10^8, \ d_5 = -4.5 \cdot 10^9,$$

$$e_1 = 419, \ e_2 = -1.1 \cdot 10^4, \ e_3 = 691, \ e_4 = 9.6 \cdot 10^4, \ e_5 = -2 \cdot 10^7.$$

(A.12)

The formula is valid for A > 0.3 and electrolyte layers thicker than 2.5 mm.

Appendix A.3. Numerical solution for rectangular geometries

The voltage loss in the electrolyte of a square cell, as illustrated in figure 4 is

$$\eta_{\Omega} = I \frac{0.05^2}{\sigma L_1^2} (a + a_1 A + a_2 A^2 + a_3 A^3 + a_4 A^4 + a_5 A^5 + a_6 A^6 + b_1 H + b_2 H^2 + b_3 H^3 + b_4 H^4 + b_5 H^5 + b_6 H^6 + d_1 A H + d_2 (A H)^2 + d_3 (A H)^3 + d_4 (A H)^4 + d_5 (A H)^5 + e_1 H / A + e_2 H^2 / A + e_3 H A^2 + e_4 A H^2 + e_5 A H^3),$$
(A.13)

with the ratio of the side lengths defined as $A = L_2/L_1$ and

$$a = 45.5, \ a_1 = -383, \ a_2 = 1312, \ a_3 = -2395 \ a_4 = 2485, \ a_5 = -1390, \ a_6 = 326,$$

$$b_1 = 354, \ b_2 = -1.6 \cdot 10^5, \ b_3 = 1 \cdot 10^7, \ b_4 = -2.6 \cdot 10^8, \ b_5 = 4.2 \cdot 10^9, \ b_6 = -1.3 \cdot 10^{10},$$

$$d_1 = -3384, \ d_2 = 3.9 \cdot 10^4, \ d_3 = -1.4 \cdot 10^7, \ d_4 = 7.5 \cdot 10^8, \ d_5 = -1.3 \cdot 10^{10},$$

$$e_1 = 1393, \ e_2 = -3.5 \cdot 10^4, \ e_3 = 1746, \ e_4 = 2.3 \cdot 10^5, \ e_5 = -5.1 \cdot 10^6.$$

(A.14)

The formula is valid for A > 0.2 and electrolyte layers, which are at least 2 mm thick.

Appendix B. Discretisation of the diffusion equation

Appendix B.1. Finite difference method

Appendix B.1.1. Equation

The diffusion equation

$$\frac{\partial c}{\partial t} = \nabla \cdot D \nabla c \tag{B.1}$$

is discretised using central differencing and the implicit Euler method as [55] (see also [94])

$$\frac{c^t - c^{t-1}}{\Delta t} = \frac{\left(D\frac{\partial c}{\partial z}\right)_{i+0.5} - \left(D\frac{\partial c}{\partial z}\right)_{i-0.5}}{0.5(z_{i+1} - z_{i-1})}$$
(B.2)

with t denoting the time, i the point-index and z the coordinate running from bottom to top. The first derivatives are discretised as

$$\left(D\frac{\partial c}{\partial z}\right)_{i+0.5} = \frac{D_i + D_{i+1}}{2} \cdot \frac{c_{i+1} - c_i}{z_{i+1} - z_i},\tag{B.3}$$

and

$$\left(D\frac{\partial c}{\partial z}\right)_{i=0.5} = \frac{D_{i-1} + D_i}{2} \cdot \frac{c_i - c_{i-1}}{z_i - z_{i-1}}.$$
(B.4)

Combining these equations leads to

$$c^{t} = \frac{\frac{D_{i} + D_{i+1}}{2} \cdot \frac{c_{i+1} - c_{i}}{z_{i+1} - z_{i}} - \frac{D_{i-1} + D_{i}}{2} \cdot \frac{c_{i} - c_{i-1}}{z_{i} - z_{i-1}}}{0.5(z_{i+1} - z_{i-1})} \Delta t + c^{t-1}$$
(B.5)

and after further simplification to

$$c^{t} = \frac{\frac{D_{i}+D_{i+1}}{2} \cdot (c_{i+1}-c_{i}) \cdot (z_{i}-z_{i-1}) - \frac{D_{i-1}+D_{i}}{2} \cdot (c_{i}-c_{i-1}) \cdot (z_{i+1}-z_{i})}{0.5(z_{i+1}-z_{i-1}) \cdot (z_{i+1}-z_{i}) \cdot (z_{i}-z_{i-1})} \Delta t + c^{t-1}$$
(B.6)

and finally to

$$c^{t} = c_{i-1} \cdot \frac{\frac{D_{i-1}+D_{i}}{2} \cdot (z_{i+1}-z_{i})}{0.5(z_{i+1}-z_{i-1}) \cdot (z_{i+1}-z_{i}) \cdot (z_{i}-z_{i-1})} \Delta t$$

+ $c_{i} \cdot \frac{-\frac{D_{i}+D_{i+1}}{2} \cdot (z_{i}-z_{i-1}) - \frac{D_{i-1}+D_{i}}{2} \cdot (z_{i+1}-z_{i})}{0.5(z_{i+1}-z_{i-1}) \cdot (z_{i+1}-z_{i}) \cdot (z_{i}-z_{i-1})} \Delta t$
+ $c_{i+1} \cdot \frac{\frac{D_{i}+D_{i+1}}{2} \cdot (z_{i}-z_{i-1})}{0.5(z_{i+1}-z_{i-1}) \cdot (z_{i+1}-z_{i}) \cdot (z_{i}-z_{i-1})} \Delta t$
+ c^{t-1} . (B.7)

The equation is simplified by setting $c^t = c_i$, defining

$$a = \frac{\Delta t}{(x_{i+1} - x_{i-1}) \cdot (x_{i+1} - x_i) \cdot (x_i - x_{i-1})}$$
(B.8)

and rearranging to fit to the matrix equation

$$A \cdot c = b, \tag{B.9}$$

which leads to

$$c_{i-1} \cdot -(D_{i-1} + D_i) \cdot (z_{i+1} - z_i) \cdot a$$

+ $c_i (1 + \cdot ((D_i + D_{i+1}) \cdot (z_i - z_{i-1}) + (D_{i-1} + D_i) \cdot (z_{i+1} - z_i)) \cdot a)$
+ $c_{i+1} \cdot -(D_i + D_{i+1}) \cdot (z_i - z_{i-1}) \cdot a$
= c^{t-1} . (B.10)

We now define the upper distance

$$dz_u = z_{i+1} - z_i, (B.11)$$

and lower distance

$$dz_l = z_i - z_{i-1} (B.12)$$

and

$$dz_u + dz_l = z_{i+1} - z_{i-1}.$$
 (B.13)

This gives

$$a = \frac{\Delta t}{(dz_u + dz_l)dz_u dz_l},\tag{B.14}$$

which leads to

$$c_{i-1} \cdot -(D_{i-1} + D_i) \cdot dx_u \cdot a$$

+ $c_i \cdot (1 + ((D_i + D_{i+1}) \cdot dx_l + (D_{i-1} + D_i) \cdot dx_u) \cdot a)$
+ $c_{i+1} \cdot -(D_i + D_{i+1}) \cdot dx_l \cdot a$
= c^{t-1} . (B.15)

Appendix B.1.2. Boundary conditions

The boundary condition at the lower interface reads

$$\nabla c \cdot \boldsymbol{n} = 0 = -c_1 + c_0 \tag{B.16}$$

which leads to the coefficients A[0,0] = 1, A[0,1] = -1 and b = 0.

The boundary condition at the upper interface reads

$$\nabla c \cdot \boldsymbol{n} = \frac{j}{\nu_e F D}.\tag{B.17}$$

This Neumann boundary condition is discretised as [55]

$$\nabla c = \frac{-c_3(z_2 - z_1)^2 + c_2(z_3 - z_1)^2 - c_1((z_3 - z_1)^2 - (z_2 - z_1)^2)}{(z_2 - z_1)(z_3 - z_1)(z_3 - z_2)}.$$
 (B.18)

As *i* counts from the bottom, we denote the last point by -1, the second last point by -2 and so on, and find

$$\nabla c = \frac{-c_{-3}(z_{-2} - z_{-1})^2 + c_{-2}(z_{-3} - z_{-1})^2 - c_{-1}((z_{-3} - z_{-1})^2 - (z_{-2} - z_{-1})^2)}{(z_{-2} - z_{-1})(z_{-3} - z_{-1})(z_{-3} - z_{-2})}.$$
(B.19)

We obtain the matrix coefficients as

$$b = \frac{j}{\nu_e FD} \tag{B.20}$$

and

$$A[-1,-3] = -(z_{-2} - z_{-1})^2 / a$$
(B.21)

$$A[-1,-2] = (z_{-3} - z_{-1})^2 / a$$
(B.22)

$$A[-1,-1] = -((z_{-3} - z_{-1})^2 - (z_{-2} - z_{-1})^2)/a$$
(B.23)

with

$$a = (z_{-2} - z_{-1})(z_{-3} - z_{-1})(z_{-3} - z_{-2}).$$
(B.24)

Appendix B.1.3. Volume change

Volume change is accounted for by adjusting in each time step the distance between the points dz (representing the volume) and rescaling the concentration appropriately. As the concentration is defined at the discretisation points, but the volume in between these points, two differently located scaling factors will be used.

As the first step, the initial masses m are computed both, at and between the discretisation points. After solving the diffusion equation, the mass fluxes of Li, \dot{m} , are calculated at and between the points. This way, the exact masses are obtained in each time step as

$$m = m_{t-1} + \dot{m}\Delta t, \tag{B.25}$$

with t - 1 denoting the previous time step. The masses are used to calculate the volume as

$$V = m/\rho \tag{B.26}$$

with the concentration-dependent density obtained from equation (35). The volumetric correction factor is then simply defined as

$$f = V/V_{t-1} \tag{B.27}$$

and is used to scale the distances between the discretisation points as

$$dz = f \cdot dz_{t-1}.\tag{B.28}$$

In order to the fact that the amount of moles of Li stays constant during this increase in volume, the molar concentration c needs to be divided by the same factor. As the concentration is defined at the discretisation points, a similar factor is computed there. To stabilise the solution, it is sometimes useful to perform the volume change operation only every nth iteration.

Appendix B.2. Finite volume method

Appendix B.2.1. Equation

The diffusion equation

$$\frac{\partial c}{\partial t} = \nabla \cdot D\nabla c \tag{B.29}$$

is discretised using the finite volume method as [55, 95–97]

$$\frac{\partial}{\partial t} \int_{V} c dV = \int_{V} \nabla \cdot (D\nabla c) \, dV. \tag{B.30}$$

Using the implicit Euler method, we obtain

$$\frac{(cV)^t - (cV)^{t-1}}{\Delta t} = \sum_f D_f \boldsymbol{S}_f \cdot (\nabla c)_f = \sum_f D_f |S_f| \frac{c_N - c_P}{d}$$
(B.31)

with V denoting the volume of the control volume, D_f the diffusivity on the faces, S_f the surface area vector, d the distance between two cell centres, c_P the concentration in the parent cell and c_N the concentration in the neighbour cell. By dividing the equation by the surface area and introducing the cell height dz we obtain

$$c^{t} = \sum_{f} D_{f} \frac{c_{N} - c_{P}}{d} \frac{\Delta t}{dz^{t}} + \frac{c^{t-1} dz^{t-1}}{dz^{t}}.$$
 (B.32)

Denoting the concentration in the lower cell by c_{i-1} and the one in the upper cell by c_{i+1} , c_N is replaced. Further, we denote the distance of the parent cell centre to the lower cell by d_l and the distance to the upper cell as d_u and set $c^t = c_i$. We obtain the discretised equation in the matrix-form

$$A \cdot c = b, \tag{B.33}$$

 as

$$c_{i-1}\left(D_f\frac{\Delta t}{d_l dz^t}\right) + c_i\left(-1 - \frac{D\Delta t}{d_l dz^t} - \frac{D_f\Delta t}{d_u dz^t}\right) + c_{i+1}\left(\frac{D_f\Delta t}{d_u dz^t}\right) = -\frac{c^{t-1} dz^{t-1}}{dz^t}$$
(B.34)

Assuming no volume change at small time steps, we can assume $dz^t \approx dz^{t-1}$.

Appendix B.2.2. Boundary conditions

The boundary condition at the lower interface reads

$$\nabla c \cdot \boldsymbol{n} = 0 = -c_1 + c_0, \tag{B.35}$$

which means basically that the flux over the lower interface needs to be set to zero. This leads to a modification of the coefficients as $A[0,0] = -1 - \frac{D_f \Delta t}{d_u dz^t}$. Similarly, the boundary condition on the top interface

$$\nabla c \cdot \boldsymbol{n} = \frac{j}{\nu_e F D} \tag{B.36}$$

requires a constant (given) flux through the upper interface, leading to a modified source term as

$$b[-1] = -\frac{c^{t-1}dz^{t-1}}{dz^t} - \frac{j\Delta t}{\nu_e F dz^t}$$
(B.37)

and a modified diagonal coefficient as

$$A[-1, -1] = -1 - \frac{D_f \Delta t}{d_l dz^t}.$$
 (B.38)

Appendix B.2.3. Volume change

Likewise to the finite difference method, the cell volume and concentration are scaled in each time step to account for volume change. However, here only one scaling factor is needed as concentration and volume are both defined in the cell centre of the control volumes.