Formation of lithiated gold and its use for the preparation of reference electrodes — an EQCM study

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Abstract
Lithiated gold wires can be used to build reference electrodes with outstanding potential stabilities over several days and even over the course of one year. These electrodes are well suited for investigations in the context of lithium-ion batteries (LIBs). In this work, a detailed procedure for the preparation of such electrodes with tailored mechanical properties, which can be fitted gastight into electrochemical cells using commercially available fittings, is given. The electrochemical lithiation process is studied using the electrochemical quartz crystal microbalance (EQCM) technique, and the differences in lithiation of wire type and thin film type gold electrodes are discussed. All experiments were carried out with two different electrolytes, namely, a LiPF6 and a lithium bis(trifluoromethane sulfonyl) imide (LiTFSI)-based electrolyte, and we conclude that for a higher lithiation rate and long-term stability, the use of LiTFSI-based electrolyte in the preparation phase is beneficial. The EQCM data provides a better insight in the analysis of film formation processes, like the buildup of the solid electrolyte interphase (SEI) during the lithiation, the rate of deposition of metallic lithium, or additional information on the kinetics of Li-Au alloy formation.

Keywords Gold-lithium alloy · SEI formation · LiPF6 · LiTFSI · Lithiation · Reference electrodes

Introduction
Lithium-ion batteries (LIBs) represent the most important achievement among modern electrochemical energy storage devices that fundamentally changed our society through portable electronics and electric vehicles. The ever-growing demand for higher energy density LIBs requires new ways and technologies to meet the needs for their future development. An important step in battery performance improvement is gaining deeper understanding of underlying electrochemical processes, accompanied by thorough electrochemical testing and characterization, as well as the development of sophisticated measurement methods. Studies in full battery cells are in that sense sometimes limiting, as they always contain overlapping information of both, anodic and cathodic, processes. In contrast, three electrode electrochemical cells, which employ a reference electrode with well-defined and stable electrode potential, can separate the characterization of cathode and anode active materials and thus provide more straightforward insights into half-reactions. However, the choice of the right reference electrodes in nonaqueous systems is not obvious. Due to the large number of applied solvents, solvent mixtures, and electrolyte salts, it is hardly possible to find a commercial reference electrode fitting all the experimental requirements. Aqueous electrochemistry in this respect is in many cases simpler, and companies offer many possibilities for commercially available, diverse reference electrodes. An excellent compilation of both aqueous and nonaqueous reference electrodes can be found in the Handbook of Reference Electrodes edited by G. Inzelt et al. [1].

In case of LIB research, the Li/Li+ system, a piece of lithium wire or foil in contact with a lithium ion containing electrolyte as half-cell, is probably the most widely used reference electrode. But in many cases, this half-cell serves
as both the counter and the reference electrode at the same time. This might be problematic even with well-established cell designs, such as coin type cells, Swagelok T cells, or pouch/coffee bag cells due to the known issue with lithium dendrite formation, especially in consecutive charge/discharge experiments. Another problem of using Li as counter and reference electrode simultaneously has been demonstrated by LaMantia et al. [2]. At high current densities, the behavior of Li becomes stochastic; i.e., the potential difference between lithium and the working electrode (WE) changes due to the polarizability of lithium. To overcome these problems, lithium might be used as the reference electrode by inserting it into the separator or in between two separator sheets [3–5]. This insertion of lithium metal can, however, be problematic due to the mechanical properties of the metal. It is soft, malleable, and easy to deform. Therefore, it is usually attached to a copper, nickel, or stainless-steel wire. The size of the so obtained reference electrodes, however, is relatively big and can disturb the electric field between the cathode and anode and falsify sensitive measurements, like impedance spectroscopy, via non-ideal positioning relative to the other electrodes [6, 7].

An alternative solution to lithium is the use of cathode materials, like Li₄Ti₅O₁₂ or LiFePO₄ [2, 8] or lithium metal alloys [9, 10] for the preparation of reference electrodes for nonaqueous electrolytes. And the use of so-called point type reference electrodes helps to significantly improve the quality and reproducibility of sensitive measurements. These reference electrodes are composed of a thin wire with an electronic insulator on its surface, being exposed to the electrolyte (the cross sectional area). Most of the remaining gold lithiation experiments reported utilize thin films, sputtered either onto a Cu substrate [20, 21] or onto a W wire [14].

During the first gold wire lithiations with a LiPF₆ containing electrolyte, we experienced many peculiarities with huge discrepancies between the reported current–time curves and our own results. Switching from LiPF₆ to lithium bis(trifluoromethane sulfonyl) imide (LiTFSI) just partially solved the problem, so we decided to investigate the phenomena deeper, utilizing the electrochemical quartz crystal microbalance (EQCM) technique. It has the advantage that it employs thin Au films deposited onto quartz, thereby conforming to previous studies, while also being a very sensitive technique for surface mass changes. Due to big differences in the construction and composition of the SEI in the two applied electrolytes, this latter aspect seemed to be especially beneficial.

The EQCM is mainly a surface mass sensitive method, widely used in investigations including metal and alloy deposition, underpotential deposition (UPD) or electroplating just to mention a few. A book chapter written by G. Inzelt gives a good overview of the method through selected applications by the author [22]. The mass changes occurring during deposition processes together with the simultaneously measured electrochemical data can be very helpful for the elucidation of reaction mechanisms via identification of the species accumulated on the surface. The method has also been used for investigations of LIB systems. In the 1990s, it was used to study the behavior of battery electrolyte solutions on Au electrodes [23], for the study of the deposition and stability of SEI forming Li compounds in the UPD region of lithium [24] or to monitor the mass change of electrodeposited Li on Ni-coated crystals [25]. Nowadays, it is mainly used to investigate intercalation processes at both the cathode and anode or formation of SEI on graphite, usually together with dissipation monitoring taking into account the porous structure of the layers [26, 27]. However, to the best of our knowledge, it has so far never been used to monitor alloy formation between Li and Au during long-term galvanostatic Li deposition that is of interest in relation with the preparation of our PEEK embedded lithiated gold wire reference electrodes (AuLi-Ref).

As it has been very well described in the preface of [1]: “In order to produce reliable experimental data, reference electrodes have to fulfill many, partly contradictory conditions, which practically never can be met to full satisfaction. For the choice and application of a particular reference electrode it is of utmost importance to know its properties as detailed as possible.” Therefore, we decided to dedicate a detailed study to the preparation of macroscopic sized Au/
Li reference electrodes that can be used in a huge variety of electrochemical cells to study nonaqueous systems, especially LIBs.

**Experimental**

**Materials**

The battery electrolytes LiPF₆ (1 mol·L⁻¹ in EC:EMC, 3:7 by wt.) and LiTFSI (1 mol·L⁻¹ in EC:EMC, 3:7 by wt.) used for the experiments were bought from E-Lyte Innovations GmbH (Münster, Germany). The silver salt AgClO₄ (anhydrous) which was used for the calibration of the EQCM was obtained from Alfa Aesar (by Thermo Fisher GmbH, Kandel, Germany). Silver (99.9%; 0.25 mm outside diameter, OD), gold (99.9%; 0.25 mm OD), and platinum (99.99%; 0.25 mm OD) wires as well as lithium foil (99.8%; 0.5 mm) were from MaTecK (MaTecK Material-Technologie & Kristalle GmbH, Jülich, Germany). The glassy carbon rods (SIGRADUR® G; 1 mm OD) used for potential stability measurements of the AuLi-Ref were purchased from HTW Hochtemperatur-Werkstoffe GmbH (Thierhaupten, Germany). All crystals for the EQCM, namely, Au-Ti (5 MHz, 100 nm Au on thin Ti adhesion layer) and Au-Cr (5 MHz, 100 nm Au on thin Cr adhesion layer), were obtained from MicroVacuum Ltd. (Budapest, Hungary). The PEEK electrode bodies (LF-1.6; Leak-Free Reference Electrode Junction, 1.6 mm OD, 1.0 mm ID), which were used for the assembly of the reference electrodes, were ordered from Innovative Instruments, Inc. (Tampa, FL, USA).

**Instruments**

All experiments were performed inside an Ar filled glovebox (M. BRAUN Inertgas-Systeme GmbH, Garching, Germany) with water and oxygen levels < 1 ppm. Electrochemical measurements were conducted with either an SP-300 or a VSP-300 potentiostat by Biologic Sciences Instruments (Seyssinet-Pariset, France). The EQCM device (eQCM-I Mini) was obtained from MicroVacuum Ltd. (Budapest, Hungary).

**Cells**

EQCM measurements were carried out in a temperature controlled cell of the type “QSHE-open” at 25 °C (MicroVacuum Ltd.) using a modified lid. Lithiation experiments of gold wires were performed in homemade 2-electrode cells of either polypropylene or glass. These cells were built by modification of chromatography sample vials of varying volumes depending on the desired immersed length of the wires into the electrolyte. In all cells, a piece of lithium foil fixed to a copper wire served as the counter electrode (CE), while the gold wires or EQCM-crystals were used as the WEs.

**Calibration**

The calibration of the EQCM was carried out using a spiral Ag wire as the counter and reference electrode and 0.1 mol·L⁻¹ AgClO₄ added into the different electrolytes. Silver was chosen as the calibration system because its redox potential lies within the stability windows of the electrolytes, while also not being able to form alloys with the used Au crystals. During the calibration, constant current densities of 200, 20, and 2 µA·cm⁻² were applied consecutively between the Au quartz and the Ag wire for 2 min each while monitoring the frequency change. First negative currents were applied to deposit silver, then positive currents were used to redissolve the deposited Ag atoms. The thereby calculated calibration factor is obtained as the mean mass per Hz change of all applied current densities (ng·Hz⁻¹) and is used for the conversion of frequency changes to mass values in the subsequent experiments.

**Results and discussions**

**Preparation and testing of the PEEK embedded lithiated gold wire reference electrodes**

Based on previous lithiation experiments from the literature [11, 14], 200 µA·cm⁻² current density was used to lithiate a gold wire with 250 µm outer diameter. The first experiments were carried out in a commercial battery electrolyte (1 mol·L⁻¹ LiPF₆ in EC:EMC, 3:7 by wt.). The lithiation time has been chosen to be 72 h. However, the typical $E-t$ curves, containing the characteristic plateaus, reported in the literature for thin film electrodes could not be reproduced. Switching to a LiTFSI based electrolyte, which is widely used for lithiation experiments [14, 20, 21], and trying different current densities only slightly changed the shapes of the $E-t$ curves, and well-pronounced plateaus could not be observed. The equilibrium electrode potentials (equilibrium open circuit voltages, OCV) of the wires, however, were very close to the potentials reported in [13]. Figure 1a shows typical chronopotentiometric (CP) curves obtained with gold wires in LiTFSI-based electrolyte and the subsequent OCV values for 48 h. Even if the lithiation curves slightly differed, the equilibrium potentials of the lithiated gold wires converged to a stable value in all cases (Fig. 1b). For a gold wire having a radius of $r$ and a length of $l$ immersed into the solution, the ratio of the amounts of lithium and gold ($n_{Li}/n_{Au}$) after a lithiation time $t$ with a constant current density $i$, assuming 100% faradaic efficiency (FE) and neglecting the
cross sectional area of the thin wire, can be calculated by the following formula:

$$\frac{n_{\text{Li}}}{n_{\text{Au}}} = \frac{i \times 2 \pi \times l \times t \times M_{\text{Au}}}{F} \times \frac{r^2 \pi \times l \times \rho_{\text{Au}}}{2 \times i \times M_{\text{Au}} \times F \times r \times \rho_{\text{Au}}} \times t = k \times t$$

where $F$ is the Faraday constant and $M_{\text{Au}}$ and $\rho_{\text{Au}}$ are the molar mass and density of gold, respectively. In our case, $k = 3.4 \times 10^{-6}$s$^{-1}$, i.e., after 72 h lithiation, the ratio of the two metals in the wire is close to 1 (0.9). Nevertheless, this ratio is enough to keep the potential of the AuLi-Ref created from such a wire constant for an extended period of time, as will be demonstrated below. However, its potential is still ca. +310 mV positive to Li, which will be different for thin-film electrodes with a much higher $n_{\text{Li}}/n_{\text{Au}}$ ratio.

Such lithiated Au wires offer many advantages for the preparation of REs: (i) their size can be flexibly varied and with that the size of the RE; (ii) they can be placed in a Luggin-capillary, as demonstrated below, in an easy manner; (iii) they offer good mechanical stability unlike thin Li wires; (iv) gold has excellent electrical conductivity and can be directly connected to the cables of the potentiostat at the free end of the wire; and (v) after proper lithiation its potential stays very stable even for long experimental times. In the following part, we discuss the preparation of PEEK housed AuLi-Refs from the wires.

For the lithiation, the gold wire is placed in a simple homemade cell filled with LiTFSI electrolyte (1 mol·L$^{-1}$; EC:EMC (3:7 by wt.)) with 200 µA·cm$^{-2}$ current density applied over 72 h and b OCV curves of five Au wires lithiated as in a all converging to a stable equilibrium electrode potential with an aqueous solution containing H$_2$SO$_4$ to maintain a wet junction. In order to avoid cross reactions of the acidic aqueous contents of the body with the later introduced electrolyte and lithiated wire, a thorough cleaning procedure was developed. This procedure consists of three stages with three different solvents. In the first stage, the body is repeatedly purged with water using a syringe and cleaned inside a water-filled beaker in an ultrasonication bath outside the glovebox; after that, the same steps are repeated in the second stage, this time with EtOH. After the second stage, the body is set to rest in a EtOH filled vial. In the third stage, the body is again repeatedly purged and cleaned in the sonication bath, this time with target solvent mix of EC:EMC (3:7 by wt.). After the final cleaning, the body is placed inside a sealed vial filled with EC:EMC and later introduced into the glovebox for assembly of the final electrode. Before the lithiated gold wire is introduced into the body, the body is filled with the target electrolyte depending on the electrode’s desired field of use (1 mol·L$^{-1}$ LiTFSI or 1 mol·L$^{-1}$ LiPF$_6$ in EC:EMC, 3:7 by wt.) (Fig. 2a). After that, the wire is introduced into the body (Fig. 2b). Finally, a wire end ferrule is placed on top of the body and wire. The metal part of the ferrule is crimped onto the wire to ensure electric contact, while the cavities between PEEK body and wire ferrule are sealed using a two-component epoxy resin to prevent evaporation of the electrolyte and ensure mechanical stability of the electrode (Fig. 2c). The electrode is then placed in a 1.5-mL vial filled with the target electrolyte to prevent the junction from drying out. After complete hardening of the epoxy resin (ca. 24 h), the electrode can be used.

In order to test the long-term stability of the reference electrode, it has been applied in a three electrode cell configuration containing a glassy carbon WE, a Pt wire CE and 10 mmol·L$^{-1}$ ferrocene (ferrocene is bis(η-cyclopentadienyl)iron(II)) dissolved in 1 mol·L$^{-1}$ LiTFSI containing EC:EMC (3:7 by wt.). The reason of using ferrocene is first that the ferrocene (Fc)/ferrocenium (Fc$^+$) couple as a reference redox system is recommended for
reporting electrode potentials in nonaqueous media [28]; therefore, the value of the equilibrium electrode potential of the Fc/Fc⁺ couple should be reported for the newly created RE. Second and luckily, this compound demonstrates good solubility, invariant redox potentials, and excellent chemical and electrochemical reversibility in the studied battery electrolytes. It has been shown to be a potentially useful redox reagent for the chemical overcharge protection of rechargeable lithium and lithium-ion batteries and therefore has been studied in detail by Laoire et al. [29]. The authors studied the kinetics of oxidation of Fc to Fc⁺ in the non-aqueous lithium ion conducting electrolyte composed of a solution of 1 mol·L⁻¹ LiPF₆ in EC:EMC (1:1 by vol.) solvent mixture and concluded that the Fc/ Fc⁺ redox couple is reversible even in this highly concentrated electrolyte.

Figure 3a demonstrates this excellent reversibility with a cyclic voltammogram obtained in 10 mmol·L⁻¹ Fc containing LiTFSI-based electrolyte. The long-term stability has been checked by carrying out cyclic voltammetric experiments every 1 h in the above-described cell. The half wave potential \( E_{1/2} \) has been determined from each CV curve from the anodic \( E_{pa} \) and cathodic \( E_{pc} \) peak potentials using a homemade software with regard to the huge number of CVs. The \( E_{1/2} \) values were estimated as \( \frac{1}{2}(E_{pa} + E_{pc}) \). Figure 3b shows the change of \( E_{1/2} \) directly after preparation.

![Fig. 3](https://example.com/fig3.png)

**Fig. 3**  a Cyclic voltammetric response of ferrocene (10 mmol·L⁻¹ in 1 mol·L⁻¹ LiTFSI containing EC:EMC (3:7 by wt.) with 100 mV·s⁻¹ scan rate and b, c stability of potential over time
of the AuLi-Ref (dark green curve) and after 1 year of use (dark red curve). The value of $E_{1/2}$ shifted ca. 100 mV in one year. The change of $E_{1/2}$ within the timeframe of 50 h can be estimated by deriving the $E_{1/2}(t)$ function and is displayed in Fig. 3c. The values are in the lower mV/h range and therefore exhibit better potential stability than common Li-foil REs (see Fig. S1). Here we would like to note that we used the original $E_{1/2}(t)$ function for the derivation without smoothing the curve in advance. To be able to account even for these minimal changes in the potential of the AuLi-Ref, we always check its potential with the Fe/Fe$^+$ redox couple both before and after the actual measurements.

**Lithiation of Au films — insights into potential stability**

The potential stability of lithiated Au electrodes can be investigated using thin film (100 nm) Au deposited onto quartz. The CP lithiation curves showed the characteristic shape, known from the literature in both electrolytes. Figure 4 shows the results of measurement sequences consisting of a CP part with 200 µA·cm$^{-2}$ for 4 h followed by an OCV period for 40 h. The potential values are IR corrected values, using the current interruption method for 50 ms, after the CP part. The first two potential plateaus shown in the insets of Fig. 4a and b belong to phase transformations of the Li$_x$Au$_y$ alloys. In the first step of alloying, Bach et al. found a hexagonal structure with composition Li$_5$Au$_3$ [21]. The second plateau corresponds to the thermodynamically stable Li$_3$Au phase with face centered cubic structure. The second plateau ends at ca. 4000 s and is followed by the deposition of metallic lithium onto the formed alloy. In case of the thin Au film the $n_{Li}/n_{Au}$ ratio as function of lithiation time can be determined by the formula:

$$\frac{n_{Li}}{n_{Au}} = \frac{i \times r^2 \times \pi \times t}{F} \times \frac{M_{Au}}{r^2 \times \pi \times h \times \rho_{Au}} \times \frac{i \times M_{Au}}{F \times h \times \rho_{Au}} \times \frac{t}{k'} = k' \times t$$

**Fig. 4** Lithiation of 100-nm-thick Au EQCM electrodes for 4 h in a 1 mol·L$^{-1}$ LiPF$_6$ containing EC:EMC (3:7) and b 1 mol·L$^{-1}$ LiTFSI containing EC:EMC (3:7) with 200 µA·cm$^{-2}$ current density with the subsequent OCV periods (purple curves) and simultaneously measured frequency change data (green curves). Apparent molar masses calculated from the frequency change data during the galvanostatic lithiation experiments for c LiPF$_6$ and d LiTFSI-based electrolytes.
where \( h \) is the height of the gold film (\( h = 100 \text{ nm} \)) and the side area of the cylinder has been neglected. In this case \( k' = 2.1 \times 10^{-3} \text{s}^{-1} \); i.e., after ca. 473 s, the \( n_{\text{Li}}/n_{\text{Au}} = 1 \) ratio is achieved. Although the length of the plates is not well correlated to this value (formation of \( \text{Li}_{2}\text{Au} \) would need ca. 1418 s), it can be seen in Fig. 4 that for both electrolytes, a stable potential of the lithiated Au electrode is achieved under these conditions. The reason for the huge discrepancy between the calculated lithiation times and observed length of plateaus is most probably the invalidity of the assumption that lithium deposition occurs with 100% FE. At this point, the analysis of the simultaneously measured EQCM data greatly helps in the explanation.

Very interestingly, the frequency change curves (green lines in Fig. 4) also reflect the formation of the different alloy phases: the rate of frequency (i.e., mass) change decreases towards the end of the potential plateaus, i.e., towards the completeness of the stable phase and remains constant for the rest of the lithiation time (from ca. 4000 s to 4 h). It is possible to calculate apparent molar masses from the measurements after calibration and assuming the validity of the Sauerbrey-equation [22], which states that the frequency change is directly proportional to the mass change (\( \Delta f \propto -\Delta m \)). It should be noted that especially due to the formation of a possibly non-rigidly attached SEI at the crystal’s surface, the applicability of the Sauerbrey-equation is not given without restrictions. The good agreement of theoretical and calculated mass change values, however, validates its use in this situation and also provides a clear basis for following calculations. We have carried out the calibration of the EQCM cell by using 0.1 mol·L\(^{-1}\) AgClO\(_4\) dissolved in the battery electrolytes. AgClO\(_4\) dissolves very slowly, and to reach a concentration of 0.1 mol·L\(^{-1}\), solutions were left overnight to dissolve. Surprisingly, the value of the integral sensitivity of the crystals was very close to the theoretical one for LiTFSI-based electrolytes (17.3 ng·Hz\(^{-1}\)·cm\(^{-2}\) calculated vs. 17.7 ng·Hz\(^{-1}\)·cm\(^{-2}\) theoretical), while the difference for the LiPF\(_6\)-based electrolyte was somewhat bigger (14.9 ng·Hz\(^{-1}\)·cm\(^{-2}\) calculated). Although PF\(_6^-\) is supposed to be a weakly coordinating anion, it seems to interact strongly to the Au surface to hinder the deposition of Ag (we can rule out competing electrode reactions at the potential of Ag deposition, which is ca. 3.8 V vs. Li). Apparent molar masses or \( M_{\text{pe}} \) (mass per electron) values have been calculated for the lithiation experiments and are shown in Fig. 4c and d. The curves are the derivatives of the corresponding \( m = m(Q/F) \) functions and are smoothed by adjacent averaging taking 50 adjacent points for the sake of better visibility.

It is evident from the change of the \( M_{\text{pe}} \) values, which decrease asymptotically after applying the constant lithiation current that first an SEI is being built on the electrode resulting in \( M_{\text{pe}} \gg 7 \text{ g·mol}^{-1} \). At the end of the potential plateaus, the \( M_{\text{pe}} \) is close to zero, i.e., no mass change, neither related to SEI formation nor to Li intercalation accompanies these phase transformations. In the last phase of the 4-h long deposition starting at ca. 4000 s and for the rest of the CP period, the \( M_{\text{pe}} \) is very close to 7 g·mol\(^{-1}\) (the horizontal line at the figure is a constant function at 7 g·mol\(^{-1}\)) for the LiTFSI-based electrolyte. The total frequency change (green line) is ca. half as much for the LiPF\(_6\)-based electrolyte as for the LiTFSI-based electrolyte (ca. 95.9 µg for LiPF\(_6\) vs. ca. 216.2 µg for LiTFSI) at same applied current density until the end of the lithiation. The reason for this is most probably the different SEI structure for the two electrolytes. Therefore, we can say that while both Au electrodes — lithiated in the two different electrolytes — will exhibit stable potentials over a prolonged period of time, it seems that Li deposition happens more readily through an SEI built up in LiTFSI. This difference in SEI build up has also a huge influence on the long-term potential stability of lithiated Au electrodes prepared in the two electrolytes with shorter lithiation times as will be demonstrated below.

**Short-term lithiation — insights into the properties of SEI**

In order to see how the potential stability of lithiated Au electrodes is influenced by lithiation times, experiments with CPs for only 1000 s instead of 4 h have been conducted (Fig. 5). Although after 1000 s lithiation, a theoretical \( n_{\text{Li}}/n_{\text{Au}} \) ratio of ca. 2 is achieved; in both cases, the \( E \) vs \( t \) curves do not stay constant and rapidly change most probably after the spontaneous delithiation of gold under OCV conditions. This is less surprising if considering that completion of the lithium alloy formation is only achieved after ca. 1400 s. Interestingly, however, the mass of the electrode in LiPF\(_6\) decreases first right after the CP period and stays more or less constant for LiTFSI. It increases for both electrolytes as the potential starts to rapidly increase. Figure 5 demonstrates an important difference between the two electrolytes: the potential stays stable for more than 30 h in LiTFSI and for only ca. 10 h in LiPF\(_6\). We speculate that the initial mass loss for LiPF\(_6\) is not related to lithium loss, since as we described above, the LiPF\(_6\) type SEI does not facilitate Li transfer as much as the LiTFSI type and the potential also stays in this phase rather constant. Maybe the inner inorganic SEI layer [30] is being dissolved in lack of a thicker and protective organic SEI in this region. However, this problem needs further investigations to be able to convincingly prove our speculation. The delithiation happens as the potential increases. In this phase, Li is being oxidized and electrolyte reduced and deposited onto the electrode, causing the mass
increase in both cases. These results demonstrate that lithiated gold electrodes will suffer potential instability on the long term if not enough Li is deposited onto the electrode.

Before the first plateau, as shown in Fig. 5c and d, a potential dip can be observed between 310 and 330 s. This is in accordance with previous observations [21]. The use of EQCM provides a better insight into the initial reactions. Right after the application of the constant current, the rate of frequency change is less than half of that registered for the first plateau and showed by the fitted linear curves. Therefore, we speculate that this initial phase showing potential shoulders and a dip is rather related to the reductive decomposition of the electrolyte, i.e., the formation of the SEI and maybe to the initial nucleation of Li on Au. Zhang et al. [31] showed that the SEI formation takes place through two major stages in the case of graphite anodes. The first stage takes place before lithiation of graphite, during which a loose and highly resistive film is formed. The second stage occurs at a narrow voltage range of 0.25–0.04 V, which proceeds simultaneously with lithiation of graphite electrode. The authors found that “in the second stage, a stable, compact, and highly conductive SEI film is produced.” When looking at the frequency change curves between 300 and 320 s also two kinds of stages can be identified. However, whereas this first phase can be characterized in LiPF₆-based electrolyte with a concave function, in LiTFSI-based electrolyte, a convex function is observed. After this ca. 20–30 s, the lithiation rate becomes constant and will decrease again only towards the end of the first potential plateau (see Fig. 4). It is also apparent that in the initial two stages of lithiation (alloy formation), the differences between the deposited mass for LiPF₆ and LiTFSI are marginal. Only during the deposition of lithium, i.e., after the alloy formation is finished, differ the frequency change rates substantially.

We carried out lithiation with wire type Au electrodes, too, that also showed the instability of the potential after an extended OCV period. In the delithiation rate, the surface area over volume ratio might play an important role, if the
charge transfer is the rate determining step. In case of the thin films, this is proportional to the inverse of the layer thickness, whereas for wires to the inverse of the wire’s diameter. In our case, there are at least three orders of magnitude differences between these two ratios for the thin films and wires that reflect in the slower delithiation rate of the wires. It can be concluded that the SEI facilitates the deposition/insertion of Li in general.

During cyclic voltammetric experiments (CVs), the aforementioned differences in the SEI forming properties of the two electrolytes can be investigated in more detail. In both cases, a new gold crystal is taken for the experiments and the electrodes are polarized with 5 mV·s⁻¹ scan rate from their OCV to cathodic directions. In the LiPF₆-based electrolyte, first, the frequency increases slightly (Fig. 6a), maybe due to the repulsion of the adsorbed PF₆⁻ anions from the surface as the potential decreases. However, the frequency starts to decrease already at around 2.1 V and a sharp cathodic peak appears at this potential. This cathodic peak corresponds to the reduction of dissolved oxygen in the electrolyte and disappears in the second cycle [32]. The frequency change is in good agreement with the work of Yamamoto et al. [24], who investigated the deposition processes in LiClO₄ containing tetraethylammonium perchlorate dissolved in propylene carbonate. The authors did not observe any substantial mass change without the addition of LiClO₄ to the solution; therefore, they concluded that Li containing species are deposited onto the surface. The rate of frequency decrease, i.e., mass increase, is the highest in the following potential region that refers to the decomposition of the electrolyte. Interestingly, the deposited layer retards the rate of deposition at lower potentials that suggests that an electrically resistive layer is forming. As we have already seen in the course of the galvanostatic measurements, this SEI layer also progressively decelerates the rate of Li deposition onto Au compared to LiTFSI (see the frequency change curves in Fig. 4a and b). The second cathodic peak corresponds to the reduction of trace amounts of water in the electrolyte. After this peak, the rate of frequency change becomes smaller. The potential has been reversed at 0.4 V vs. Li to avoid the underpotential deposition of Li onto Au [23]. Upon potential reversal, the mass increase continues in the anodic scan until the OCV potential is reached again and the current remains in the cathodic region at around ~50 nA·cm⁻². The frequency change is irreversible and finally accounts to ca. 180 Hz mass increase. Since the measured currents are in the nA range, the charge passed through the electrode is also minimal compared to the large frequency changes. After converting the frequency changes to mass changes using the coefficient determined from calibration, apparent molar masses are in the range of >1000 g·mol⁻¹, referring to mainly chemical reactions causing the deposit on the surface rather than electrochemical.

Contrary to LiPF₆, no oxygen reduction peak can be observed in the LiTFSI-based electrolyte (Fig. 6b). The absence of this peak could either originate from varying oxygen levels of the electrolytes caused by the manufacturer or differences (i.e., inhibition) in the oxygen reduction mechanism in the presence of TFSI⁻ anions. The overall frequency change for this LiTFSI electrolyte is very moderate down
to 0.5 V compared to LiPF$_6$. Another difference is that the rate of frequency change never decreases until the potential reversal of 0.15 V, which refers to an SEI with good electrical conductivity. This potential limit was chosen because of the very steep increase in current, which starts at around ca. 0.17 V vs Li and is related to the underpotential deposition (UPD) of Li [32] (simultaneously to ongoing SEI formation as the frequency change curve shows). Most probably the deposited Li initiates a faster buildup of the SEI. Upon potential reversal, two anodic peaks appear at around 0.4 V and 0.5 V, that are related to the oxidation of Li. Parallel to these processes, the frequency increases and stabilizes after the peaks. We speculate that the two peaks are the result of oxidation of energetically two different kinds of Li: Li deposited onto Au and already alloyed with Au. From the frequency change curve, it seems that the irreversible change corresponds to the formed SEI (around 40 Hz) and the rest of the reversible change to the electrochemical oxidation of Li. Compared to LiPF$_6$, it seems that LiTFSI forms less SEI in the same potential range; however, the SEI formed from LiTFSI-based electrolyte has beneficial properties towards the deposition of Li onto Au (see also Fig. 4). This might have some implications also for graphite electrodes and on the intercalation/deintercalation kinetics of Li, which needs, however, further studies.

**Conclusions**

We have examined the electrochemical lithiation of gold in two different kind of electrolytes in form of wires and thin films with the aim to build stable reference electrodes with well-defined mechanical properties. Gold wires after 72 h of lithiation with a current density of 200 µA·cm$^{-2}$ showed excellent electrochemical stability over a prolonged test period and are highly suitable to build PEEK embodied reference electrodes with variable mechanical properties. These electrodes also showed outstanding potential stability not only over the course of several days but also over an entire year. Experiments with thin film Au electrodes in form of coated EQCM crystals provided insights into the effects of varying lithiation times and electrolytes. While too short lithiation times will result in unstable potentials, also, the use of LiPF$_6$ will result in lower potential stability compared to LiTFSI-based electrolytes. The EQCM measurements also revealed differences in the kinetics of lithiation of thin film Au compared to wired type Au electrodes. Well-pronounced potential plateaus appeared on the chropopotentiometric curves, and the simultaneously measured frequency change curves also show the changing lithiation kinetics towards the ends of the potential plateaus. These frequency change curves could also explain the large differences between the two electrolytes. Whereas the total mass change until the completion of formation of the stoichiometric Li-Au alloy is roughly the same for the two electrolytes, compared to LiPF$_6$, the amount of deposited lithium after 4 h is more than double in LiTFSI-based electrolyte. The calculation of apparent molar mass values reflects a relatively fast buildup of the SEI that is followed by a period in which nearly exclusively Li is deposited. The differences of the two types of SEI layers that are already revealed in the frequency change curves and also reflected during first CV cycles are the most probable reason for this huge difference in the lithiation behavior of the different electrolytes.

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**Availability of data and material** All experimental data is available upon request from the authors.

**Declarations**

**Conflict of interest** The authors declare no competing interests.

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