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Design and investigation of molybdenum modified platinum surfaces for modeling of CO tolerant electrocatalysts

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Abstract

Mo overlayers were prepared on smooth polycrystalline platinum and platinized platinum electrode surfaces by *in situ* electrochemical deposition of molybdenum oxide at potential below 500 mV for modeling Mo-Pt electrocatalysts. Correlations were found between the applied potential and the amount of deposited Mo, which never exceeded a monolayer, thus Pt-Mo bonds stabilize the deposited Mo oxide. Electrochemical measurements suggested that Mo deposited from a Mo(VI) solution was reduced to the 4+ oxidation state. In line with the *ex situ* XPS findings a certain part (20-25%) of the initial Mo layer remained irreversibly adsorbed on the Pt/Pt electrode even after oxidation into the 6+ state at high potentials; this fractional monolayer cannot be dissolved even by prolonged cyclic polarization up to 1000 mV.

It has been demonstrated that the irreversibly bound Mo partial monolayer is enough to change significantly the CO poisoning properties of the Pt surface. On this Mo:Pt (1:4) surface CO oxidation is initiated at extremely low potentials (ca. 100 mV). Moreover, only Pt modified by Mo(IV) species is active in low-potential CO oxidation reaction as after oxidizing the irreversibly adsorbed Mo to the 6+ state, CO oxidation is no longer observable. Nevertheless, the catalyst can be reactivated by reduction of molybdenum into the 4+ oxidation state. However, this reduction requires clean, CO-free Pt surface. If Pt is largely covered by CO, reduction of Mo(VI) into Mo(IV) does not occur and thus the low potential CO oxidation remains hindered.

Keywords: *In situ* electrochemical deposition, Molybdenum oxide, Pt electrodes, Optimized surface composition, Anode electrocatalysts, CO-tolerance

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1 Introduction

Polymer Electrolyte Membrane Fuel Cells (PEMFCs), due to their low operation pressure and temperature, form a very attractive class of fuel cells particularly suited for low and medium power mobile applications. Key components of a PEM fuel cell are the electrocatalysts on which the anode (hydrogen oxidation) and cathode (oxygen reduction) reactions take place. The optimal electrocatalyst should be highly active both in hydrogen oxidation and oxygen reduction, possess good long term stability and have adequate electrical conductivity at an affordable price. Experience shows that still platinum supported on active carbon is the most suitable choice both at the cathode and the anode. However, the sensitivity of Pt for poisoning by contaminants in the fuel (especially CO), together with the pronounced corrosion of the active carbon support under fast load change conditions [1,2] can only be compensated by extremely high Pt loadings, which keeps the price of the PEMFCs high.

Recent investigations suggest that some of the limitations of the Pt/C system can be overcome by applying a second oxophilic metal. In general, reactive hydroxyl species form more easily on the oxophilic additive than on Pt, which not only enhances the hydrogen oxidation reaction rate [3-5] but is also beneficial in the much slower oxygen reduction process [6,7], and facilitates easy oxidation of the CO contaminant as well via the so-called bifunctional mechanism [8,9], supposed that atomic closeness between the oxophilic sites and Pt is ensured. In these systems the additive may be present as an alloying component in the carbon supported Pt nanoparticles; examples involve Pt-Sn [10,11], Pt-Mo [12-17] or Pt-Ru [18,19], which is also commercially available as the state-of-art CO-tolerant electrocatalyst recommended for PEM fuel cells. Moreover, due to the beneficial co-catalytic effect of Mo in CO and ethanol oxidation numerous studies were focused on the preparation of a well-characterized Pt-Mo bulk alloy electrodes [20-23].

Apart from alloying, the required atomic closeness of the oxophilic component and Pt may also be ensured if the former is applied as support or as a modifier of the active carbon support. Numerous works investigated the properties of such systems in which Pt is supported either by molybdenum- [24] or tungsten-oxides [25] or by active carbon doped with these oxides [26-30]. According to these studies, molybdenum- or tungsten oxides offer several advantages when they are used as components of the support for Pt electrocatalysts [31-35]. Apart from enhanced tolerance against poisoning by the bifunctional effect, the adsorption properties of the Pt particles can be tuned by changing their electronic properties by the ligand effect. In addition, hydrogen spillover from Pt towards the oxide may enhance the catalytic activity and electrooxidation of small organic molecules may be promoted. Moreover,

stabilization of the highly dispersed state of the noble metal nanoparticles can be achieved by strong metal-support interaction, which prevents their agglomeration. Finally, relative stability is expected from these oxides across the anticipated potential/pH window.

The vast majority of heterogeneous catalytic reactions occur at high temperature and elevated pressure. As a result, elucidation of structure-activity-stability relationships under *in operando* conditions is still challenging. Electrocatalysts for PEMFC applications form a notable exception, as a significant amount of information about their behavior can be obtained by simple electrochemical measurements under conditions closely resembling those found in fuel cell electrodes. Thus it is not surprising that extensive studies have been done on the development of Pt-MoO_x model electrodes, in which atomic closeness of Pt atoms and Mo ions is ensured. These catalysts can be prepared by means of versatile techniques. Most synthetic routes use either co-deposition of the Pt and Mo precursors on glassy carbon electrodes [36-38] or Mo deposition on platinum single crystals [39,40] or platinum [41,42] or platinized platinum electrodes [43,44]. The resulting systems contain molybdenum in different oxidation states depending on the synthetic procedure.

Molybdenum oxides include five Magnéli phases with compositions between MoO₂ and MoO₃ [45]. MoO₂ has a rutile-type structure and unusually high electronic conductivity [46], while MoO₃, which is more stable under oxidative conditions [47], has an orthorhombic structure and is non-conducting. However, it can be electrochemically reduced to a non-stoichiometric and electroconductive hydrogen molybdenum bronze. The formation of H_xMoO₃ endows MoO₃ with high electronic and proton conductivity [48]. Usually, molybdenum oxides do not exist in a single phase but in mixed-valence oxides (MoO_x) [49]. The lattice defect structure and self-diffusivity of oxygen are the determining factors of the electrochemical properties concomitant with the changing valence state of molybdenum [50]. In agreement with the bifunctional mechanism, the co-catalytic activity is supposed to be due to a rapid and easy change of the oxidation state of molybdenum.

Although reducible oxides of oxophilic metals are indeed valuable co-catalysts or supports for PEMFC electrocatalysts, their dissolution under the working conditions of the cell is always a concern, as dissolved metal cations can absorb in the polymer membrane, resulting in the loss of its proton conducting property [51]. To overcome this problem, a composite support was proposed, in which the oxophilic metal co-catalyst is incorporated into a TiO₂ matrix which provides long term stability, while the good electrical conductivity and high surface area of the system is ensured by an active carbon backbone. The preparation and

the thorough characterization of Pt electrocatalysts deposited onto $\text{Ti}_{0.7}\text{M}_{0.3}\text{O}_2\text{-C}$ (M= W, Mo) composites was presented in our recent studies [52-54].

Using $\text{Pt}/\text{Ti}_{(1-x)}\text{Mo}_x\text{O}_2\text{-C}$ ($x= 0.2\text{-}0.4$) catalytic systems it has been demonstrated [53,54] that the oxidation of CO takes place at exceptionally low potentials (the onset potential $E_{\text{CO, onset}} \leq 100$ mV) compared to the Pt/C and PtRu/C catalysts. Additionally, a characteristic feature of the cyclic voltammograms of the $\text{Pt}/\text{Ti}_{(1-x)}\text{Mo}_x\text{O}_2\text{-C}$ catalysts is the appearance of a molybdenum redox peak pair between 380 and 530 mV. The mixed oxide component increased the CO tolerance under simulated CO poisoning conditions by providing adsorbed hydroxyl species (OH_{ads}) for CO oxidation at very low electrode potentials [54,55]. After the weakly bound CO is oxidized, surface sites released in this way can adsorb some hydrogen [25,54]. Thus, in the presence of CO impurities oxophilic dopant-containing composites improved the catalytic activity in the hydrogen oxidation reaction (HOR) with respect to conventional Pt/C catalysts [53,54].

The results of the mentioned investigations suggested a possible correlation between the molybdenum redox phenomenon and the CO tolerant behaviour of the Mo-containing electrocatalysts. To further elucidate this point, model experiments were designed in which the electrochemical properties of Pt-Mo catalytic surfaces prepared by *in situ* electrochemical deposition of Mo on smooth polycrystalline platinum and platinized platinum electrodes were explored. Using electrochemical methods the amount of irreversibly deposited Mo and the change of the oxidation state of molybdenum during polarisation were assessed. The influence of the molybdenum oxidation state on the CO tolerance and stability of the Mo-Pt/Pt electrode was also investigated. The results reported here are supposed to give insight into the processes determining the activity and stability of the Mo-containing Pt-based electrocatalysts and may also give hints about the optimum surface composition and Pt/Mo ratio in real catalysts.

2 Experimental

2.1 Electrochemical experiments

Mo-Pt bimetallic model electrocatalyst surfaces were prepared by electrodeposition of Mo-species onto platinum. Ammonium heptamolybdate tetrahydrate ($(\text{NH}_4)_6\text{Mo}_7\text{O}_{24} \times 4\text{H}_2\text{O}$, Merck, 99%) was used as Mo precursor compound. Two types of Pt electrode were used: smooth polycrystalline Pt and platinized platinum with a relatively large real surface area (roughness factor – R between 200 and 250).

A special three-compartment electrochemical cell was used, in which the solution of the main compartment could be replaced with deoxygenated electrolyte with the exclusion of

air. Hydrogen electrode was used as reference and a Pt sheet as counter electrode. All potentials are given on RHE scale. All electrochemical measurements were carried out at ambient temperature. The electrolyte was 0.5 M H₂SO₄ (Merck, P.A.).

Solutions were prepared from Millipore MilliQ water and p.a. reagents. 5.5 N Ar gas was used for deoxygenation the solutions. Real surface area of Pt was estimated from the amount of charge used for oxidation of the adsorbed hydrogen.

2.2 Physical characterization

To estimate the amount of adsorbed Mo on smooth Pt electrodes, *ex situ* X-ray photoelectron spectroscopy (XPS) measurements were carried out using an EA125 electron spectrometer manufactured by OMICRON Nanotechnology GmbH (Germany). The electrodes were transferred to the electron spectrometer under water to minimize air exposure and avoid excessive oxidation. The photoelectrons were excited by nonmonochromatized MgK α (1253.6 eV) radiation. Spectra were recorded in the Constant Analyser Energy mode of the energy analyser with 30 eV pass energy resulting in a spectral resolution around 1 eV. Binding energies were referenced to the Fermi edge of the Pt foil. Data were processed using the CasaXPS software package [56] and the XPS MultiQuant software package [57].

3 Results and Discussion

3.1 Preparation of Mo modified Pt surfaces

Fig. 1 compares the cyclic voltammograms (CVs) of a smooth polycrystalline platinum electrode recorded using different cathodic potential limits in 0.5 M H₂SO₄ solution containing 5×10^{-4} M Mo(VI) precursor compound ((NH₄)₆Mo₇O₂₄ \times 4H₂O). CV of the Pt electrode recorded in pure H₂SO₄ solution is also depicted for comparison.

In the Mo-free solution, a typical CV of Pt with the classical features of the underpotentially deposited hydrogen adsorption/desorption (i, i') between 50 mV and 350 mV along with platinum oxide formation (iii) and its reduction region can be identified. Comparing the voltammograms of platinum electrode in the solutions with and without heptamolybdates it is obvious that significant changes take place on the surface in the molybdenum containing solution. The massive cathodic current below 550 mV indicates some reduction process and since the hydrogen adsorption and desorption (i, i') peaks of the platinum disappeared it can be concluded that the product of this reduction totally covered the platinum surface.

As shown in Fig. 1 upon using different potential limits during the cyclic polarization the quantity of the Mo-species electrodeposited on the surface of Pt electrode can be varied. Upon shift of the potential limit from 400 mV to 100 mV the amount of adsorbed Mo-species increases. Accordingly, as can be seen from Fig. 1, the highest peak of the oxidation of adsorbed Mo-species (ii) was detected at the potential limit of 100 mV. It also can be seen from Fig. 1, that deposition of Mo-species starts at more positive potentials comparing to the beginning of the H-adsorption on platinum.

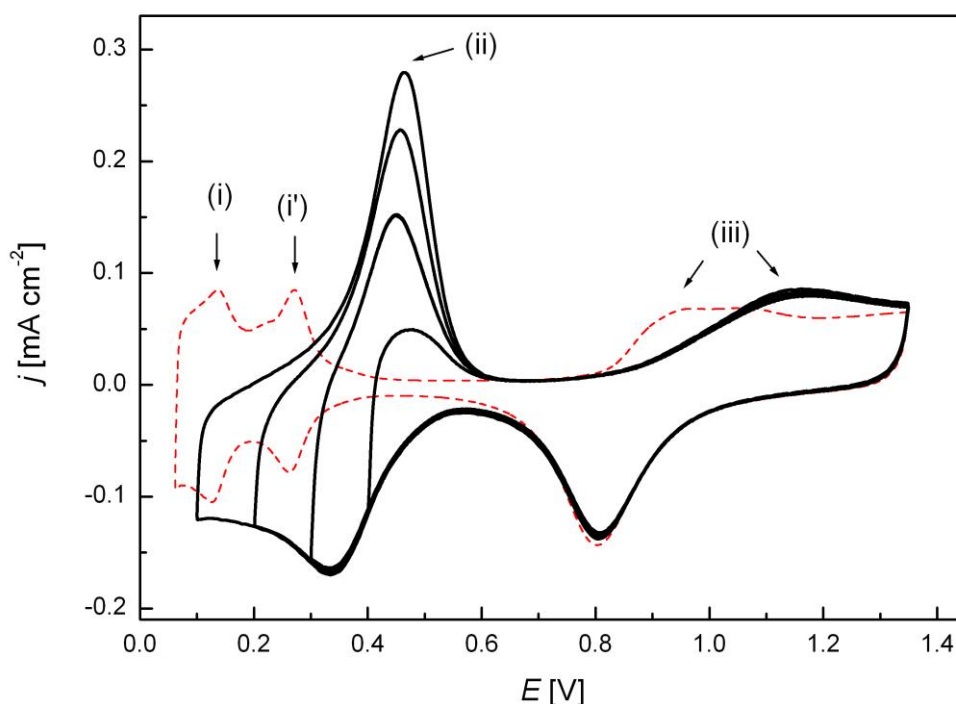


Fig. 1. Cyclic voltammograms of a smooth Pt electrode in 5×10^{-4} M Mo(VI) containing 0.5 M H_2SO_4 with different cathodic potential limits (400, 300, 200, 100 mV). Dashed line: CV of the Pt electrode in 0.5 M H_2SO_4 , sweep rate: 50 mV/s. Marks of the anodic peaks: (i), (i') - oxidation of weakly and strongly bonded hydrogen, respectively, (ii) - oxidation of Mo surface species, (iii) - oxygen adsorption on Pt.

It can be also noticed that during the electrodeposition of Mo-species onto the platinum electrode the position of the oxidation current peak usually attributed to platinum oxide formation (iii), above 800 mV, is also influenced. The onset potential of this (iii) peak is shifted to more positive potential than that observed on pure platinum electrode.

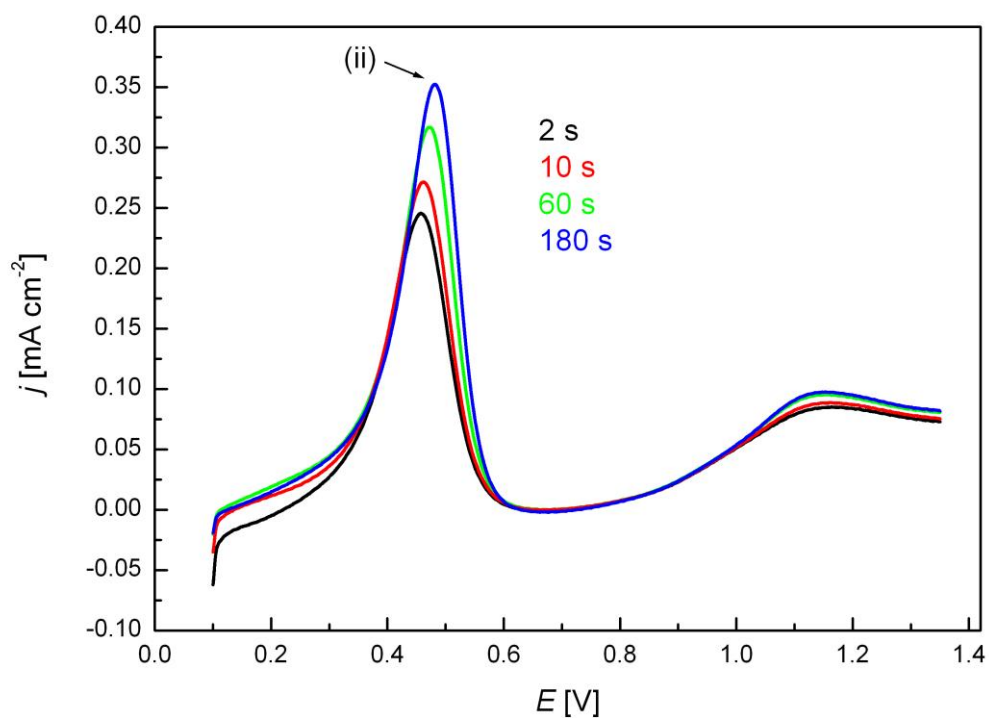


Fig. 2. Linear potential sweep of a smooth Pt electrode in 5×10^{-4} M Mo(VI) containing 0.5 M H_2SO_4 electrolyte after potentiostatic Mo-deposition at 100 mV for 2, 10, 60 and 180 s. Sweep rate: 50 mV/s. (ii) - Oxidation peak of Mo surface species.

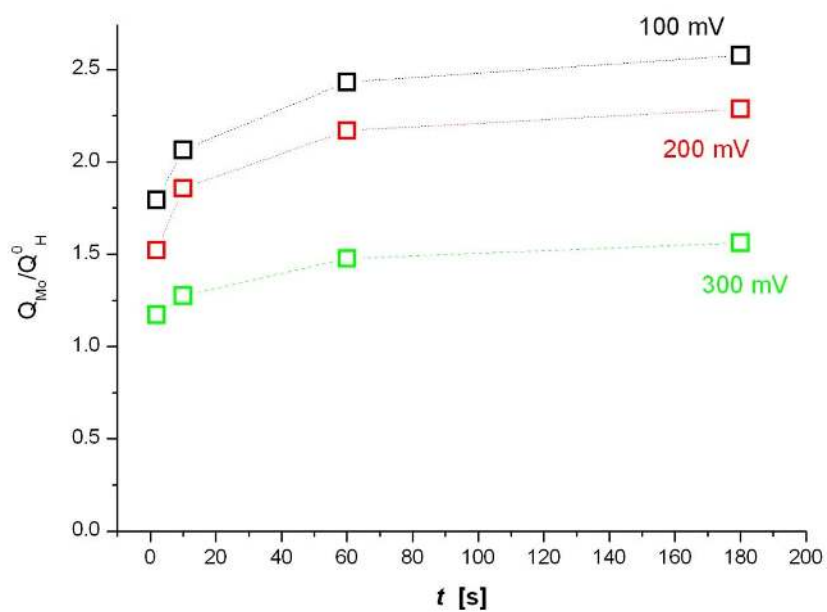


Fig. 3. $Q_{\text{Mo}}/Q_{\text{H}}^0$ ratio as a function of deposition time of Mo at different potentials.

By adding various ions (e.g. Cl^-) to the 0.5 M H_2SO_4 electrolyte solution similar shift of the above mentioned peak was observed [58,59], which could serve as a possible explanation for the observed phenomenon in our case (however, this phenomenon is out of the scope of this paper and shall not be discussed here).

As shown in Fig. 2 in the case of potentiostatic Mo-deposition at 100 mV the magnitude of the oxidation current peak (ii) observed at 450 mV depends on the duration of the deposition. Increasing the time of deposition resulted in increase of the Mo-oxidation peak (ii) at 450 mV (i.e. the amount of the deposit). However, the rate of this increase was reduced with the time of deposition, and the amount of deposit tended towards a saturation value at all investigated potentials, as can be seen in Fig 3, where the charge of the Mo-oxidation peak is normalized to the charge of the adsorbed hydrogen of the bare Pt.

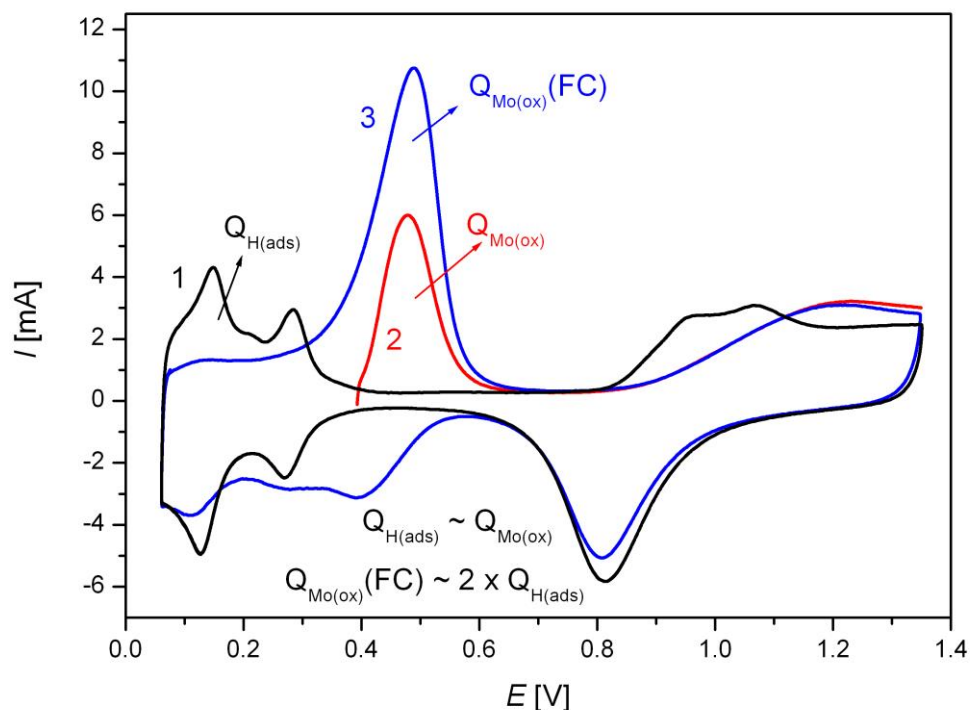


Fig. 4. Cyclic voltammogram of a Pt/Pt electrode in 0.5 M H_2SO_4 (curve 1). Linear potential sweep of the same electrode after Mo deposition via oxidation of preadsorbed hydrogen (curve 2) and after Mo-deposition at 50 mV for 5 min (curve 3). Sweep rate: 10 mV/s.

$Q_{\text{H(ads)}}$: oxidation charge of adsorbed hydrogen; $Q_{\text{Mo(ox)}}$: oxidation charge of surface Mo-species; $Q_{\text{Mo(ox)}(\text{FC})}$: oxidation charge of surface Mo-species in the case of full coverage.

In order to obtain more reliable data about the relative amount of Mo-deposit platinized platinum (Pt/Pt) electrode was used in the following experiments (in the case of larger real surface area the role of disturbing factors -for example, small amount of impurities- can be decreased).

CV of the Pt/Pt electrode and linear potential sweep after Mo deposition via oxidation of preadsorbed hydrogen are shown on Fig. 4 (curves **1** and **2**, respectively).

As a first step of this experiment the Pt/Pt electrode was saturated with adsorbed hydrogen in pure H₂SO₄ electrolyte, then Mo(VI) ions were introduced into the main compartment of the cell under open circuit conditions (potential was: 50 mV). Adsorbed hydrogen, as a reducing agent, reacted with Mo(VI) leading to deposition of Mo-species in an amount equivalent to the charge transfer from hydrogen. It can be mentioned that the potential of the electrode as a result of this process rose to 390 mV in 5 minutes. As indicated on Fig. 4 the amount of the oxidation charge of surface deposited Mo-species ($Q_{\text{Mo(ox)}}$) was equal to that of the hydrogen adsorption/desorption ($Q_{\text{H(ads)}}$). This means that the deposition of Mo species from the Mo(VI)-solution and the oxidation of the deposit take place without loss of charge, the same amount of charge was needed for the oxidation as it was available for the deposition. Thus, it has been proven that all of the product of the reduction remained on the surface and did not desorbed into the electrolyte.

Linear potential sweep of the Mo-Pt/Pt electrode after potentiostatic Mo-deposition for 5 min at 50 mV potential corresponding to full coverage can be also seen in Fig 4 (curve **3**). It is known that initiation of Mo-containing electrode at 50 mV is sufficient to reduce Mo⁶⁺ to Mo⁴⁺ [60]. At full coverage considerable increase of the peak corresponding to the Mo oxidation reaction is obvious ($Q_{\text{Mo(ox)}}(\text{FC}) \sim 2 \times Q_{\text{H(ads)}}$). Thus, the change of the oxidation state of the molybdenum during Mo-deposition at 50 mV for 5 min from 6+ to 4+ can be proposed. Oxidation of the deposited Mo monolayer requires twice as much charge than oxidation of adsorbed hydrogen, which suggests a probable Mo⁶⁺→Mo⁴⁺ transition upon deposition and a Mo⁴⁺→Mo⁶⁺ transition upon oxidation.

According to the literature [13] the comparison with Pourbaix diagram for Mo confirms that the oxidation peak around 450 mV is due to the Mo⁴⁺→Mo⁶⁺ transition. In Ref. [61] the number of electrons involved in the redox reaction calculated from the Mo redox peak pair of CVs recorded at slow sweep rates for the Pt/MoO_x/C system was also in a good agreement with assignment of this system to MoO₂/MoO₃.

3.2 Stability of the adsorbed Mo-species

In the next set of experiments the stability of the adsorbed surface Mo-species was studied.

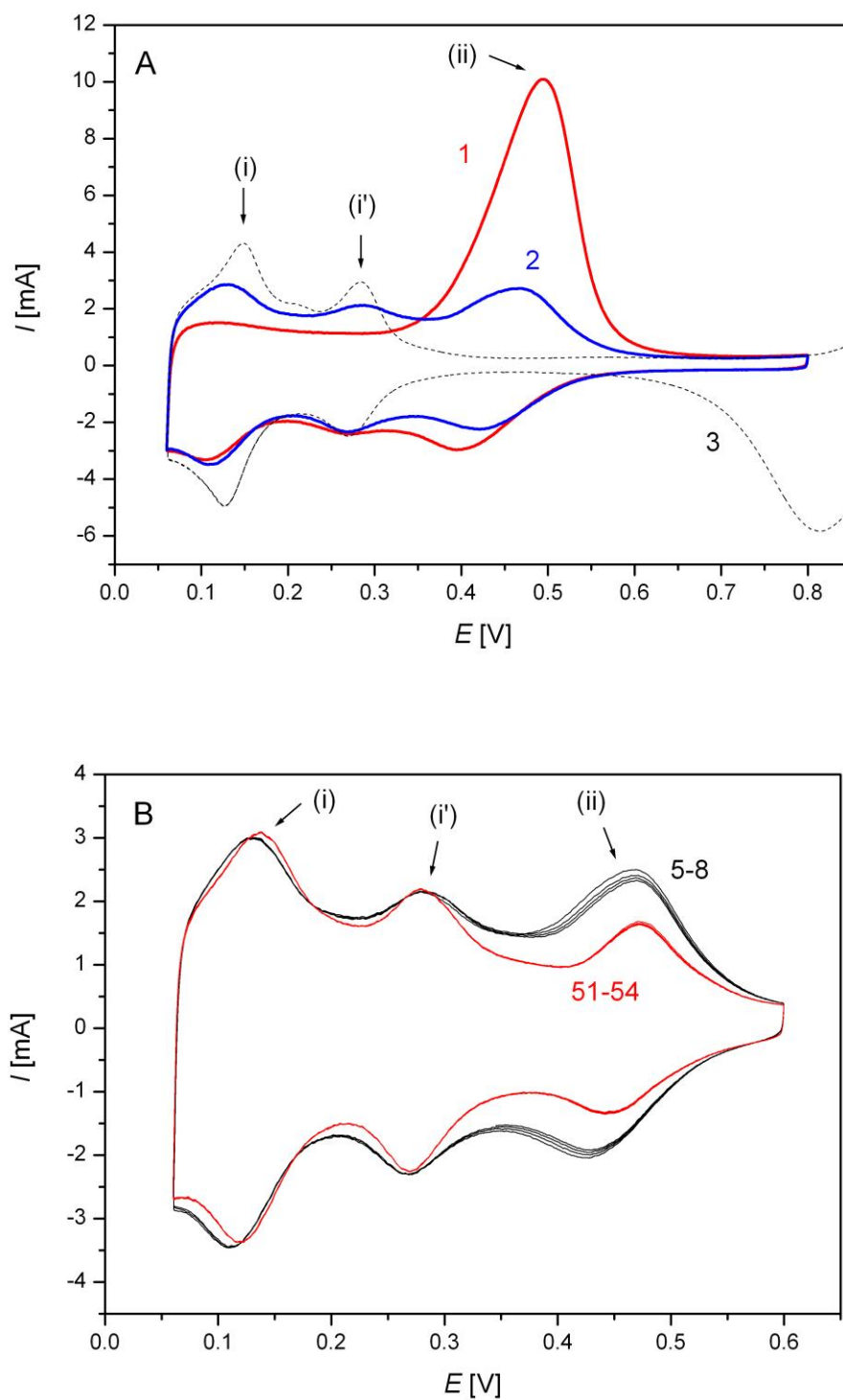


Fig. 5. (A) Cyclic voltammograms of a Mo-Pt/Pt electrode after Mo-deposition at 50 mV for 5 min; CVs were measured in pure Mo-free 0.5 M H₂SO₄. Curve 1: 1st cycle, curve 2: 3rd

cycle, curve 3 (dashed line): CV of the bare Pt/Pt electrode in 0.5 M H₂SO₄. (B) Continuation of the cyclic polarization of the Mo-Pt/Pt electrode in 0.5 M H₂SO₄. Numbers of the cycles are indicated. Marks of the anodic peaks: (i), (i') - oxidation of weakly and strongly bonded hydrogen, respectively, (ii) - oxidation of Mo surface species.

After Mo deposition at 50 mV the electrolyte was changed and CVs were recorded in pure Mo-free H₂SO₄ electrolyte. As shown in Fig. 5.A after the first cycle a significant decrease of the current peak (ii) at 450 mV could be observed (see curve 2 in Fig. 5.A). This behavior was accompanied with some restoration of the hydrogen adsorption/desorption peaks (i, i'), indicating that certain part of the monolayer Mo-deposit has been desorbed during the first potential cycle. However the 25-30 % of the Pt surface remained covered after the first cycle, and the cathodic pair of the Mo oxidation peak (ii) indicates that the oxidized Mo can be reduced. Continuing the cyclic polarization this strongly adsorbed part of the Mo-oxide proved to be quite stable, as demonstrated by the 5-8 and 51-54 cycles on Fig. 5.B. As seen from Fig. 5.B prolonged polarization cycling results in only small decrease of the Mo deposit (ii). Thus, even after oxidation into the 6+ state a certain part (20-25%) of the initial Mo monolayer irreversibly deposited on the Pt/Pt electrode remains stable on the surface even after prolonged cyclic polarization; this Mo surface species can be reduced back into the 4+ state even after polarization up to 1000 mV. It is necessary to mention that the amount of deposited Mo depends on the applied potential but never exceeds a monolayer, thus Pt-Mo bonds stabilize the deposited Mo.

The stability of Mo-containing Pt electrocatalysts is still under debate in the existing literature. Irreversible Mo loss from the electrode surface has been previously reported in the literature [22,60,62], which was attributed to the formation of Mo species soluble in acid solution. In Ref. [63] the fact of Mo dissolution has been confirmed in a test with a smooth Pt electrode immersed into the working electrolyte. A peak at ca. 450 mV, very characteristic to the presence of Mo species in the system, appears on the CV of the Pt test electrode after a few cycles of the working Pt-Mo/C electrode. At the same time, much inhibited dissolution of Mo from the Pt-Mo alloys was attributed to the stabilization effect of the alloying with Pt [13]. Our own observations also suggest that Mo species adsorbed at certain locations are particularly strongly bound and cannot be easily oxidized into dissolvable species. In this aspect it may be interesting to note that theoretical calculations of Wang *et al.* [64] predicted a non-uniform distribution of Pt on Mo atoms in the outermost atomic layer in Pt₈₀Mo₂₀

nanoparticles: the Pt atoms segregate preferentially to the facet sites, while Mo atoms favor the low coordination edge and vertex sites, alternating with Pt whenever possible.

3.3 Behaviour of the Mo-Pt/Pt electrode in the presence of CO

In our recent study it has been demonstrated [54] that the oxidation of CO on the Mo-containing Pt-based catalyst commences at exceptionally low potential values (ca. 100 mV). We demonstrated [54] that the CO stripping voltammograms of the Pt/Ti_(1-x)Mo_xO₂-C (x= 0.2-0.4) catalysts display two clear oxidation features: (i) the main anodic peak at 705 mV, and (ii) two small overlapping “pre-peaks” centered at ca. 215 and 400 mV, which may be attributed to partial oxidation of weakly adsorbed CO and oxidation of Mo surface species, respectively.

Behaviour of the model catalytic surfaces prepared by *in situ* electrochemical deposition of molybdenum oxide on platinized platinum electrodes in the presence of CO is demonstrated in Fig. 6.A. Four consecutive cyclic voltammograms of Mo-Pt/Pt electrode with Mo coverage of about 25% in CO saturated and CO-purged H₂SO₄ electrolyte were recorded between 50 and 550 mV with low sweep rate (0.5 mV/s). Considering the very low sweep rate the current values can be regarded as representation of quasi-stationer states.

As shown in Fig. 6.A during the first sweep in CO saturated H₂SO₄ significant oxidation current in the anodic scan was observed. In this case the oxidation of CO takes place at very low potential (the onset potential $E_{\text{CO,onset}} < 100$ mV). Thus the irreversibly adsorbed Mo partial monolayer is enough to change significantly the CO poisoning properties of the Pt surface. However, in the second and during the subsequent cycles in this potential range pronounced decrease of the anodic current was observed. The pre-oxidation peak completely disappeared in the fourth cycle (see curve 4 in Fig. 6.A).

Thus, the CO tolerance of the Mo-Pt/Pt electrode was totally lost after the third sweep of the cyclic polarization from 50 mV to 550 mV. Since the upper potential limit 550 mV is higher than the potential needed for the Mo(IV)→Mo(VI) transition, this observation suggests that only Pt modified by Mo(IV) species is active in the low-potential CO oxidation reaction (LPCOR). Despite the fact that after oxidizing to the 6+ state Mo remains on the Pt surface, LPCOR is not observable. This behavior is in analogy with that found previously for Pt/Ti_(1-x)Mo_xO₂-C mixed oxide-carbon composite supported catalysts, which also lost their activity in LPCOR when Mo had become fully oxidized (above ca. 400 mV) and re-reduction of these oxidized species was hindered when strongly adsorbed carbonyl species were still present.

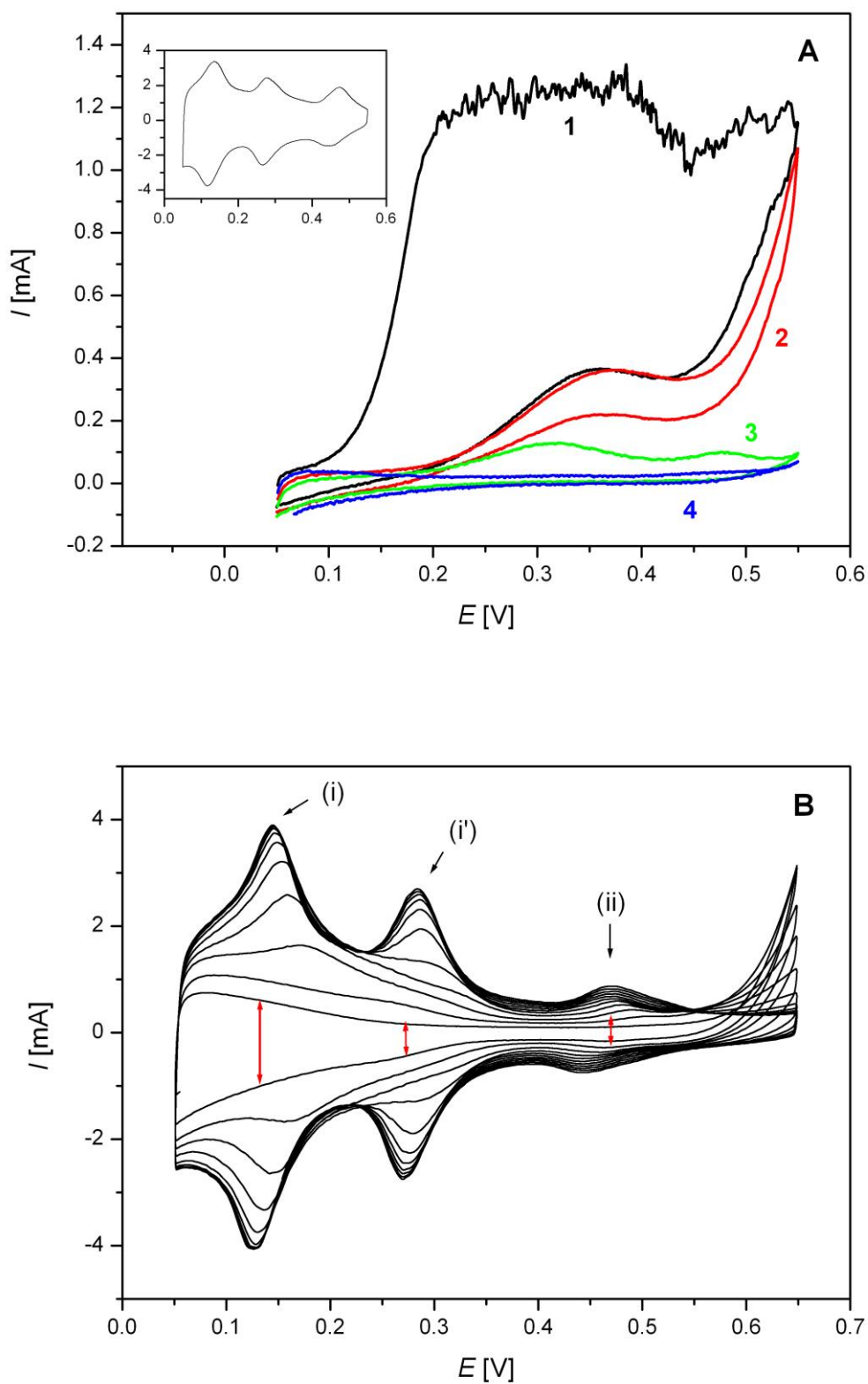


Fig. 6. (A) Cyclic voltammograms of Mo-Pt/Pt electrode (Mo coverage is about 25%) with low sweep rate (0.5 mV/s) in CO saturated and CO-purged 0.5 M H_2SO_4 . Numbers indicate the number of cycles. Insert: CV of the same electrode in Ar-purged H_2SO_4 with 10 mV/s

sweep rate. (B) Cyclic voltammograms of the CO-covered Mo-Pt/Pt electrode in pure Ar-purged 0.5 M H₂SO₄. Red colored arrows indicate the direction of changes. Marks of the anodic peaks: (i), (i') - oxidation of weakly and strongly bonded hydrogen, respectively, (ii) - oxidation of Mo surface species.

In the case of the composite supported electrocatalysts oxidized Mo species can be re-reduced and the activity in the LPCOR can be restored only if the platinum surface is CO-free. If Pt is largely covered by CO, reduction of Mo(VI) into Mo(IV) does not occur. It means that reduction of these Mo(VI) species requires an abundant source of hydrogen in the form of hydrogen spillover and/or charge transfer from Pt particles released from CO [65]. Note that, CO_{ads} species can be completely removed only above 550 mV (above the so called pre-peak potential).

In order to explore the relation between the presence of strongly bound CO adsorbates and the reduction behavior of the mentioned Mo(VI) species in case of the Mo-Pt model system, in the next set of experiments the upper potential limit of the polarization was set to 650 mV, where oxidation of a fraction of the CO species left on the Pt surface after the previous scans limited to 550 mV is expected. Fig. 6.B illustrates the influence of the number of the polarization cycles to 650 mV on the change of the electrochemical behaviour of the Mo-Pt/Pt electrode in pure CO-free solution.

After measuring the **1-4** curves presented in Fig. 6.A, Ar was bubbled through the main compartment of the cell and 20 CVs were recorded again in the CO-free electrolyte solution. Already in the 2nd-3rd cycles the appearance of the peaks corresponding to the adsorption/desorption of weakly bonded hydrogen (i) on the Pt surface and appearance of the Mo redox peaks (ii) were observed. In the 4th cycle the peaks of the adsorption/desorption of strongly bonded hydrogen (i') were also recovered. This experiment reveals that partial oxidation of strongly adsorbed CO starts via slight increase of the potential limit up to 650 mV and subsequently the strongly oxidized surface Mo species (ii) can be re-reduced.

Fig. 7.A compares the CVs of a Pt/Pt electrode recorded in 0.5 M H₂SO₄ solutions with and without 5 x 10⁻⁴ M Mo(VI) precursor compound. As described earlier and shown in Fig. 7.A a characteristic feature of the CV of the Mo-Pt/Pt electrode (see curve **2**) is the appearance of the Mo redox peak pair (ii) in the double layer region compared to the CV of Pt/Pt. From this comparison it is obvious that after deposition of Mo-species slight decrease of the hydrogen adsorption/desorption peaks (i) and (i') was also observed.

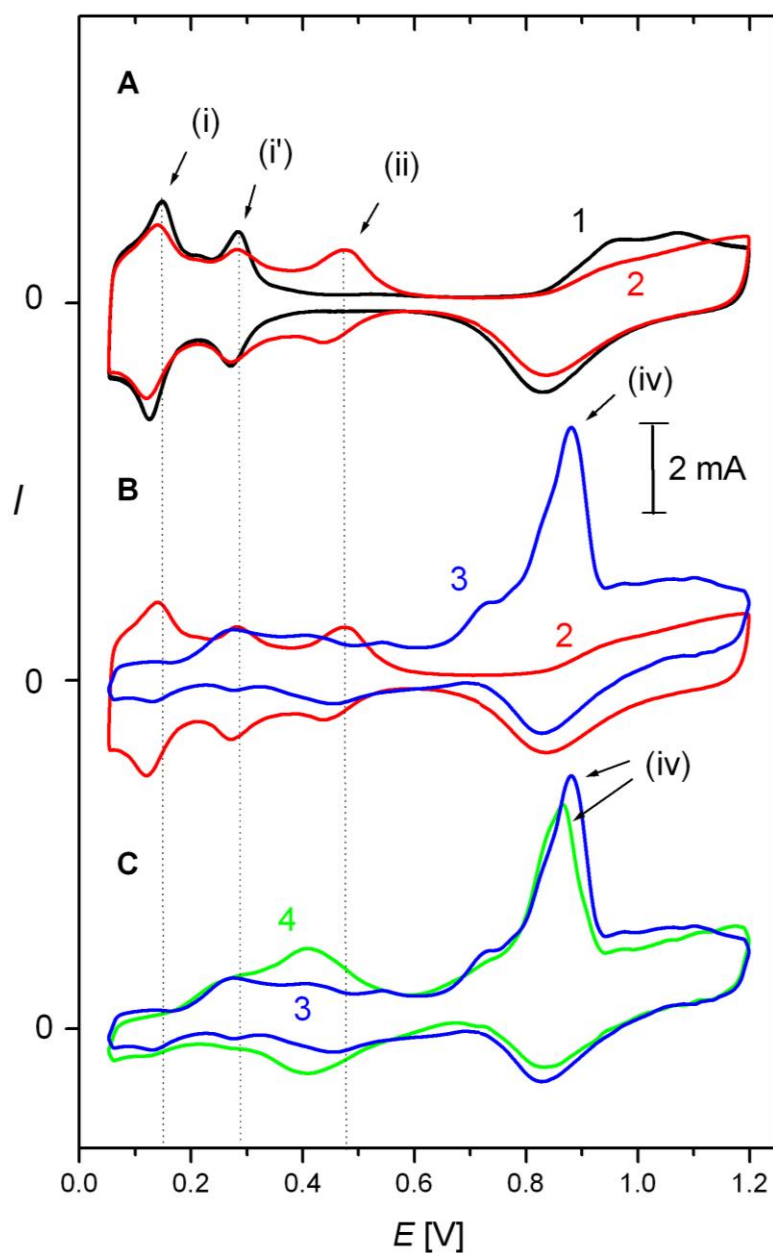


Fig. 7. Cyclic voltammogram of a Pt/Pt electrode recorded in 0.5 M H_2SO_4 (curve **1**), in 5×10^{-4} M Mo(VI)-containing 0.5 M H_2SO_4 (curve **2**), in 5×10^{-4} M Mo(VI)-containing and CO-saturated H_2SO_4 (curve **3**), in 10^{-3} M Mo(VI)-containing and CO-saturated 0.5 M H_2SO_4 (curve **4**). Sweep rate 10 mV/s. Marks of the anodic peaks: (i), (i') - oxidation of weakly and strongly bonded hydrogen, respectively, (ii) - oxidation of Mo surface species, (iv) - oxidation of strongly adsorbed CO.

This behaviour is in accordance with the presence of some amount of Mo irreversibly deposited on the Pt/Pt electrode surface.

Fig. 7.B reveals the changes observed in the shape of the CVs of the Mo-Pt/Pt electrode before (curve **2**) and after saturation the electrolyte with CO (curve **3**). Since in CO-saturated electrolyte CO blocks the platinum catalytic sites the hydrogen adsorption/desorption peaks (i) and (i') are not observed (see curve **3** in Fig. 7.B). The voltammogram of the Mo-Pt/Pt electrode recorded in CO-saturated electrolyte displays usual oxidation features corresponding to the Mo-containing Pt catalysts: the main anodic peak (iv) and several small overlapping “pre-peaks” between 200 and 600 mV (curve **3**). In our recent study [54] these pre-oxidation peaks were attributed to oxidation of both weakly adsorbed CO and Mo surface species (ii).

However, it is necessary to mention that the position of the main anodic peak (iv) observed on the model Mo-Pt/Pt electrode is shifted by ca. 150 mV toward more positive potentials comparing to the Pt/Ti_(1-x)Mo_xO₂-C (x= 0.2-0.4) catalysts [54]. According to the literature and our recent studies [39,54,65] this peak can be assigned to the electrooxidation of strongly bound CO.

Comparison of the CO_{ads} electrooxidation behaviors of the Mo-Pt/Pt electrode with different amount of surface deposited Mo species (compare curves **3** and **4** in Fig. 7.C) demonstrates an increased CO tolerance of the Mo-Pt/Pt electrode with higher Mo content. As shown in Fig. 7.C the position of the main anodic peak (iv) corresponding to the electrooxidation of strongly bounded CO for Mo-Pt/Pt electrode with increased Mo content shifts toward less positive potential values in comparison with that electrode obtained by using 5×10^{-4} M Mo precursor compound. Moreover, it is necessary to mention that the behaviors of the Mo-Pt/Pt electrode with higher Mo content strongly resembles to that observed in our earlier study over Mo-containing composite supported Pt catalysts [54].

3.4 *Ex situ* photoelectron spectroscopy analysis

In order to obtain further information on the properties of the Mo deposits on the Pt electrodes, *ex situ* X-ray photoelectron spectroscopy experiments were carried out on samples prepared by saturating of the smooth polycrystalline platinum electrode surface with Mo at 50 mV. After the Mo deposition the platinum foil was taken out from the electrochemical cell at 50 mV (see curve (a) on Fig. 8) or a part of the Mo overlayer was removed by polarizing to 600 mV (curve (b) on Fig. 8) or 1000 mV before removal (curve (c) on Fig. 8). In order to

avoid undesired oxidation, the samples were carried to the photoelectron spectrometer under water; nevertheless, a few minutes exposure to ambient conditions was unavoidable.

Fig. 8.A shows the XPS spectra of the Mo 3d region at the beginning of the data collection. Irrespective to the preparation conditions, in all three samples Mo⁶⁺ and Mo⁵⁺ ionic states were observed with leading contributions around 232.5 and 231.1 eV binding energies [66]. Assuming that some oxidation may easily occur during sample transfer, the presence of a strong Mo⁵⁺ contribution in the spectrum of the Mo-saturated sample is in agreement with the proposed reductive deposition as described in the electrochemical results. The XPS data confirm that a certain fraction of the adsorbed Mo species remain on the Pt surface even after polarization to 1000 mV. Somewhat surprisingly, in both cases of oxidative removal of the dissolvable Mo, the *ex situ* XPS measurement reveals the presence of a mixture of Mo⁶⁺ and Mo⁵⁺ states.

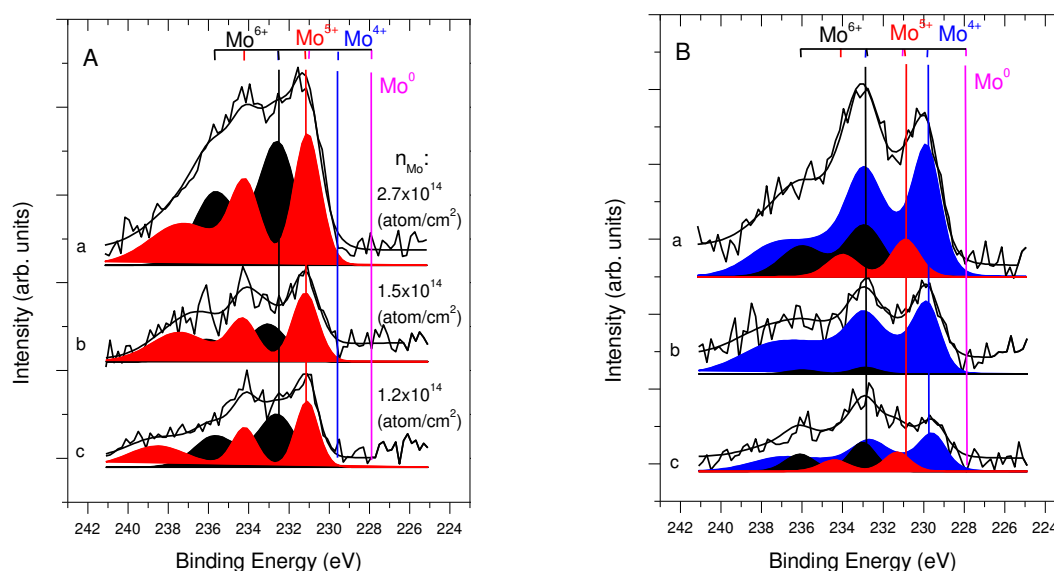


Fig. 8. Mo 3d spectra of the Mo/Pt model samples at the beginning of the data collection (A) and after 1 h exposure to the X-ray radiation in the electron spectrometer (B). Curve (a): after deposition of Mo up to the saturation coverage at 50 mV; (b) after removing of dissolvable Mo at 600 mV; (c) after removing of dissolvable Mo at 1000 mV. Numbers in panel A indicate the estimated surface concentration of Mo.

The easily reversible oxidation of the adsorbed Mo species is indicated by the very pronounced radiation sensitivity of the Mo spectra. As shown in Fig. 8.B 1 h exposition to the X-ray radiation transforms a significant part of the higher Mo oxides into the Mo⁴⁺ state,

which has a leading peak slightly below 230 eV binding energy. The Mo⁴⁺ state is characterized by a complex line shape due to differently screened final states [67,68]; the line shape used here was derived from measurements on MoO₂ [54]. According to our observations, the sample obtained by polarization to 1000 mV resists somewhat more to reduction by X-ray exposure.

Quantitative evaluation of the Mo and Pt peak intensity data gives an estimation on the surface concentration of Mo ions. According to these data, the saturation amount of deposited Mo is around 0.2 monolayer (1 monolayer is around 1.3×10^{15} atom/cm², corresponding to the surface atomic density of polycrystalline Pt). The polarization treatment indeed significantly decreases the surface concentration of Mo and after oxidation at 1000 mV less than 0.1 ML Mo remains on the Pt.

4 Conclusions

Model Pt-Mo catalytic surfaces were prepared using electrochemical and electroless deposition of molybdenum at potential below 500 mV onto smooth polycrystalline Pt and Pt/Pt electrodes. According to the results of the electrochemical measurements there was correlation between the applied potential and the amount of deposited Mo, which never exceeded a monolayer, thus Pt-Mo bonds stabilize the deposited Mo oxide. Since the hydrogen adsorption and desorption peaks of the platinum disappear it has been concluded that the deposited product totally covered the platinum surface. Electrochemical data indicated that at full coverage oxidation of the deposited Mo monolayer requires twice as much charge as needed for oxidation of adsorbed hydrogen, which suggests a probable Mo⁶⁺→Mo⁴⁺ transition upon deposition and a Mo⁴⁺→Mo⁶⁺ transition upon oxidation. In line with the *ex situ* XPS findings even after oxidation into the 6+ state a certain part (20-25%) of the initial Mo monolayer remains irreversibly bound to the Pt surface; this fractional layer remains stable on the surface even after prolonged cyclic polarization. The irreversibly adsorbed Mo surface species can be reduced back into the 4+ state even after polarization up to 1000 mV.

It has been demonstrated that this irreversibly deposited Mo partial monolayer is enough to change significantly the CO poisoning properties of the Pt surface. On the Mo:Pt (1:4) surface CO oxidation is initiated at extremely low potentials (ca. 100 mV). Moreover, only Pt modified by Mo(IV) species is active in low-potential CO oxidation reaction. Despite the fact that after oxidizing to the 6+ state Mo remains on the Pt surface, low potential CO oxidation becomes hindered in the presence of completely oxidized Mo. Thus, the Pt-Mo

catalytic surface lost their activity in LPCOR when Mo had become fully oxidized (above ca. 400 mV). However, Mo(VI)-oxide remaining on the surface can be reactivated by reduction of molybdenum into the 4+ oxidation state, but this reduction requires clean, CO-free Pt surface. If Pt is largely covered by CO, reduction of Mo(VI) into Mo(IV) does not occur.

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