Study on Methanol Oxidation at Pt and PtRu Electrodes by Combining in situ Infrared Spectroscopy and Differential Electrochemical Mass Spectrometry

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Methanol oxidation reaction (MOR) at Pt and PtRu electrodes examined at potentials from 0.3 V to 0.6 V (vs. RHE) have studied under potentiostatic conditions by in situ FTIR spectroscopy in attenuated-total-reflection configuration and differential electrochemical mass spectrometry under controlled flow conditions. Results reveal that (i) CO is the only methanol-related adsorbate observed by IR spectroscopy at all the Pt and PtRu electrodes examined at potentials from 0.3 V to 0.6 V (vs. RHE); (ii) at PtRu electrodes, two IR bands, one from CO adsorbed at Ru islands and the other from CO at Pt substrate are detected, while at other electrodes, only a single band for CO adsorbed at Pt is observed; (iii) MOR activity decreases in the order of PtRu > Pt; (iv) at 0.5 V, MOR at Pt reaches a current efficiency of 50% for CO₂ production, the turnover frequency from CH₃OH to CO₂ is ca. 0.1 molecule/(site-s⁻¹). Suggestions for further improving of PtRu catalysts for MOR are provided.

**Key words:** Differential electrochemical mass spectrometry, Electrochemical in situ infrared spectroscopy, Methanol oxidation, PtRu electrode, Current efficiency

II. EXPERIMENTS

Millipore Milli-Q water and perchloric acid (suprapure, Merck) are used to prepare 0.1 mol/L HClO₄, which is used as supporting electrolyte. Pt electrode with a thickness of ca. 50 nm on the flat reflecting face of a hemi-cylindrical Si prism formed by electroless deposition [23] is used as the working electrode (WE) or used as substrate for preparing the “PtₓRuᵧ” WE, PtRu catalysts [6]. A bifunctional mechanism and a ligand effect have been suggested to be the origin for the improvement in MOR activity [22]. Despite the great achievements mentioned above, no consensus for the optimal composition and structure for MOR catalysts has been reached so far. In order to develop more efficient MOR catalysts, further information on MOR mechanism, on the relationship of MOR activity with catalysts composition and structure are necessary.

In this work, we present results on MOR at Pt and PtRu electrodes in a flow cell by using combined EC-ATR-FTIRS and DEMS. Quantitative information on the development of both the surface absorbate (i.e., CO_ads) and the volatile product CO₂ is obtained. Suggestions for further improving of PtRu catalysts for MOR will be provided based on present results and the information from literature.
The DEMS setup used in the present study is a HidenHPR-40 DSA Bench top-membrane inlet gas analysis system, mass signals are collected 20 points/s. The mass signal for CO₂ produced has been calibrated by oxidative stripping of a saturated CO adlayer preadsorbed at 0.06 V, mass calibration constant $k=Q_T/Q_{mass}$ is found to be $3.65 \times 10^6$ mA/Torr in this study. The IR spectroscopic measurements are carried out using a Varian FTS-7000 spectrometer equipped with a mercury cadmium telluride (MCT) detector. A spectrum taken at the corresponding reaction potential in the supporting electrolyte is used as the reference spectrum. All spectra are obtained with a resolution of 4 cm⁻¹ and 1 spectrum/s. The spectra are presented in absorbance mode, i.e., $\lg(R_0/R)$, where $R_0$ and $R$ are the reflectance at reference and sample potential, respectively.

III. RESULTS AND DISCUSSION

Figure 1 gives the typical cyclic voltammograms of Pt, Pt₀.8₃Ru₀.₁₇, Pt₀.₇₃Ru₀.₂₇, and Pt₀.₅₄Ru₀.₄₄ electrodes. It is clearly seen that after the deposition of Ru, the current in H-UPD region decreases, while that in the double layer region increases. The decrease in the current for H-UPD is due to the fact that Pt sites initially for H to adsorb is now covered by Ru and the increase in the current in the double layer region is due to the oxidation of Ru to Ru-OH. The nominal coverage of Pt and Ru at the surface (relative to the original active surface area of Pt electrode) is estimated from the decrease in the charge for H-UPD. Time-resolved IR spectra in the spectral region from 1700–2200 cm⁻¹ during MOR at Pt electrode after switching from 0.1 mol/L HClO₄ to 0.1 mol/L HClO₄+0.5 mol/L CH₂OH are given in Fig.2. The bands in the region of 2000–1900 cm⁻¹ come from the stretching vibration of bridge- and multi-bonded CO (denoted as CO₂ and CO₃) linearly. From Fig.2, it is seen that right after switching to methanol containing solution, IR bands for CO₂ and CO₃ appear, whose band intensity increases with time, and reaches maximum at ca. 10 s.

(x and $y$ in the subscripts of “PtₓRuᵧ” represent the nominal coverage of Pt and Ru at the surface which is relative to the original active surface area of Pt electrode, $x+y=1$). The geometric surface area of the Pt electrode is 0.8 cm² and the roughness factor of the Pt electrode is ca. 7 as estimated from the charge to follow the kinetics for CO ad layer preadsorbed at 0.06 V, mass calibration constant $k=Q_T/Q_{mass}$ is found to be $3.65 \times 10^6$ mA/Torr in this study. The IR spectroscopic measurements are carried out using a Varian FTS-7000 spectrometer equipped with a mercury cadmium telluride (MCT) detector. A spectrum taken at the corresponding reaction potential in the supporting electrolyte is used as the reference spectrum. All spectra are obtained with a resolution of 4 cm⁻¹ and 1 spectrum/s. The spectra are presented in absorbance mode, i.e., $\lg(R_0/R)$, where $R_0$ and $R$ are the reflectance at reference and sample potential, respectively.

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to 30 s after the solution switch. We have checked carefully the full spectral range from 1000 cm$^{-1}$ to 4000 cm$^{-1}$ (Fig. 3), except small IR signal in the region above 3300 cm$^{-1}$ from the stretching vibration O–H of interfacial water, no other bands from methanolic fragments is observed at all the reaction potentials from 0.3 V to 0.6 V. At potentials between 0.3 and 0.5 V, the steady-state CO$_L$ band intensity is nearly the same, which corresponds a CO$_{ad}$ coverage of ca. 0.5 ML as estimated from a linear relationship of CO$_L$ band intensity and CO$_{ad}$ coverage in this coverage region [19, 20].

Time-resolved IR spectra during MOR at Pt$_{0.73}$Ru$_{0.27}$ and Pt$_{0.56}$Ru$_{0.44}$ electrodes at 0.3, 0.4, 0.5, and 0.6 V recorded upon switching the solution from 0.1 mol/L HClO$_4$ to 0.1 mol/L HClO$_4$+0.5 mol/L MeOH at 0 s are given in Fig. 4 and Fig. 5. IR spectra during MOR at Pt$_{0.83}$Ru$_{0.17}$ electrode are nearly the same as that at pure Pt electrode, except that the IR band intensities of CO$_L$ and CO$_M$ are slightly smaller (spectra are not given here). From Fig.4, it is found that CO$_{ad}$ is also the only methanolic intermediates formed at the Pt$_{0.73}$Ru$_{0.27}$ electrode surface. However, compared to the case with pure Pt electrode, the IR signals for both CO$_L$ and CO$_M$ are much weaker, whose band intensity is ca. 10 times smaller than that at Pt electrode under otherwise identical condition. From the linear relationship between the CO$_L$ band intensity and CO$_{ad}$ surface coverage [19], under steady state for MOR at 0.4 V, CO$_{ad}$ coverage at Pt$_{0.73}$Ru$_{0.27}$ is estimated as ca. 0.05 ML. The IR band intensity for both CO$_L$ and CO$_M$ is found to be maximum at 0.5 V (which corresponds to ca. 0.08 ML of CO$_{ad}$ at Pt$_{0.73}$Ru$_{0.27}$), it decreases toward both higher and lower potentials. The increase in CO$_{ad}$ band intensity with potential from 0.3 V to 0.5 V is explained by the fact that at $E \leq 0.4$ V the surface is partly occupied by UPD-H atoms. As a result, the rate for methanol dehydrogenation decreases with increase in the coverage
FIG. 4 Time-resolved IR spectra of CO formed during methanol oxidation on Pt_{0.75}Ru_{0.27} electrode at 0.3, 0.4, 0.5, and 0.6 V, electrolyte is switched from 0.1 mol/L HClO$_4$ to 0.1 mol/L HClO$_4$+0.5 mol/L CH$_3$OH at 0 s.

FIG. 5 Electrochemical in situ IR spectra recorded during the methanol oxidation on Pt$_{0.56}$Ru$_{0.44}$ at 0.3, 0.4, 0.5, and 0.6 V, the electrolyte was switched to 0.1 mol/L HClO$_4$+0.5 mol/L MeOH at 0 s.
of UPD-H atoms. Results at pure Pt electrode reveal that the rate for methanol dehydrogenation increases with potential increase from 0.3 V to 0.7 V [20], the decrease in CO$_{ad}$ band intensity at Pt$_{0.73}$Ru$_{0.27}$ above 0.5 V is explained by the enhanced oxidation rate of CO$_{ad}$ by the addition of Ru.

From the IR spectra given in Fig.5 recorded during MOR on Pt$_{0.56}$Ru$_{0.44}$ electrode surface, it is seen that at 0.3 and 0.4 V another band in the region 1970–2020 cm$^{-1}$ appears in addition to the IR spectra for CO$_L$ and CO$_M$ adsorbed at Pt as similar to the case given in Fig.4. Furthermore, it is found that the intensity of this band decreases with increase of electrode potential from 0.3 V to 0.4 V, and it disappears at higher potentials. This band is assigned to CO$_{ad}$ adsorbed at Ru islands, which will only appear when Ru adatoms at Pt electrodes becomes islands with large enough size ($\geq$1 nm). Such result confirms that Ru adatoms on Pt surface tends to form clusters or islands instead of epitaxial growth. These behaviors mentioned above can be explained by the faster CO$_{ad}$ oxidation rate other than that for its formation from methanol dehydrogenation at such Ru sites.

The Faradaic currents, the mass signals of $m/z$=44 from CO$_2$ and $m/z$=60 from CH$_3$OOCH as a function of reaction time recorded during methanol oxidation on Pt, Pt$_{0.83}$Ru$_{0.17}$, Pt$_{0.73}$Ru$_{0.27}$ and Pt$_{0.56}$Ru$_{0.44}$ at 0.4 and 0.5 V are presented in Fig.6. When the applied potential is 0.3 V, no mass signal CO$_2$ and CH$_3$OOCH from MOR has been detected and after the initial current spike right after switching to methanol containing solution, the Faradaic currents drops immediately to zero (results are not shown here). From Fig.6 it is seen that immediately after switching from 0.1 mol/L HClO$_4$ to 0.1 mol/L HClO$_4$+0.5 mol/L CH$_3$OH at 0.4 V, MOR current at Pt film electrode increases sharply and reaches peak current at ca. 1.2 s after the solution switch. Then, it drops quickly to zero after ca. 15 s of the solution switch. At Pt electrode, no mass signal for CO$_2$ and HCOOCH$_3$ are detected at 0.4 V. At Pt$_x$Ru$_y$, both the magnitude of the initial current spike right after the solution switch and the rate of current drop after the current spike decreases with increase in Ru coverage. MOR currents on Pt$_{0.83}$Ru$_{0.17}$ at 0.4 V drops to zero after ca. 4 min exposing to methanol containing solution, while on Pt$_{0.56}$Ru$_{0.44}$, Pt$_{0.73}$Ru$_{0.27}$, and Pt$_{0.56}$Ru$_{0.44}$ electrode, a steady state current of ca. 0.03 mA can be maintained at 0.4 V.

Accompanying with the initial current spike, there are also significant mass signals for CO$_2$ produced from MOR on all three Pt$_x$Ru$_y$ electrodes at 0.4 V. After the current spike, mass signals for CO$_2$ drops sharply. At Pt$_{0.73}$Ru$_{0.27}$ a CO$_2$ current efficiencies of ca. 5% is attained under steady state, while at Pt$_{0.83}$Ru$_{0.17}$ and Pt$_{0.56}$Ru$_{0.44}$ the current efficiency for CO$_2$ production at 0.4 V are less than 1%. At all three Pt$_x$Ru$_y$ electrodes, the mass signal for CH$_3$OOCH is two orders of magnitude smaller than that for CO$_2$, it is not sure whether this is due to the slow kinetics for the reaction between HCOOH and CH$_3$OH, or just because

![FIG. 6 Time course for (a) reaction current, mass signal of (b) $m/z$=44 from CO$_2$ and (c) $m/z$=60 from CH$_3$OOCH for methanol oxidation at Pt, Pt$_{0.83}$Ru$_{0.17}$, Pt$_{0.73}$Ru$_{0.27}$ and Pt$_{0.56}$Ru$_{0.44}$ electrode at 0.4 V (left) and 0.5 V (right) after solution switching from 0.1 mol/L HClO$_4$ to 0.1 mol/L HClO$_4$+0.5 mol/L CH$_3$OH at 0 s.](image-url)
HCOOH produced at the potential is rather small. Further studies are underway to clarify this issue. In this work, we focus on analysis of the mass signal of CO$_2$ together with the IR signal of CO$_{ad}$. The higher rate for CO$_2$ production on the three Pt$_x$Ru$_y$ electrodes at 0.4 V right in methanol containing solution, is probably due to the fact that methanol is decomposed at Pt sites neighboring to Ru to CO$_{ad}$, and the CO$_{ad}$ can be oxidized easily to CO$_2$ with OH$_{ad}$ previously adsorbed at neighboring Ru edges. After the consumption of OH$_{ad}$ species at Ru, the rate for the formation of OH$_{ad}$ at Ru sites are smaller than that for methanol dehydrogenation to CO$_{ad}$ at Pt. Such CO$_{ad}$ may also diffuse fast to the neighboring Ru sites. Once such sites become poisoned, the CO$_2$ production rate becomes very small. This is in agreement with the fact that Ru islands deposited at clean Pt catalysts surface is not good for methanol dehydrogenation, however, they are good for CO$_{ad}$ oxidation.

At 0.5 V, at Pt and Pt$_{0.83}$Ru$_{0.17}$ electrodes, MOR current immediately drops to zero as similar to the case at 0.4 V. In contrast for the cases with Pt$_{0.73}$Ru$_{0.27}$ and Pt$_{0.56}$Ru$_{0.44}$, although the initial current right after switch to methanol containing solution much smaller than that for the cases at Pt and Pt$_{0.83}$Ru$_{0.17}$ electrodes and they are comparable to those Pt$_{0.73}$Ru$_{0.27}$ and Pt$_{0.56}$Ru$_{0.44}$ at 0.4 V, however the decay of the Faradic current with reaction time becomes much slower than the other two cases (Fig.6(a), right). Furthermore, high mass signals for CO$_2$ and CH$_3$OOCH have also been detected. The Faradic current and the mass signals from MOR at Pt$_{0.73}$Ru$_{0.27}$ electrode are ca. 1.5 times higher than the corresponding values at Pt$_{0.56}$Ru$_{0.44}$ electrode. After correcting the CO$_2$ mass signal with the mass calibration constant, the current efficiency for CO$_2$ produced from MOR on Pt$_{0.73}$Ru$_{0.27}$ electrode at 0.5 V is estimated to be ca. 50%, this indicates that there are still more than half of methanol molecules incompletely oxidized to HCOOH or HCHO. Additionally, both the Faradic current and the mass signals from MOR on Pt$_{0.73}$Ru$_{0.27}$ electrode at 0.5 V display a continuous decay. Considering that CO$_{ad}$ coverage at Pt$_x$Ru$_y$ with y>0.27 is rather small (ca. 0.08 ML) and its band intensity reaches maximum at ca. 30 s after solution switch, the current decay is the most probably due to the fact that the ensemble sites for methanol dehydrogenation are not optimized.

The current density for MOR at Pt$_{0.73}$Ru$_{0.27}$ after normalizing to the active surface area of the Pt electrode before deposition of Ru is ca. 35 µA/cm$^2$ it is slightly higher than Ru deposited on Pt(111) single crystal, however it is ca. one fourth of the MOR activity reported for Pt$_{0.73}$Ru$_{0.25}$ alloys prepared under ultrahigh vacuum conditions through high temperature annealing [3, 29]. We have tried to use many different ways to deposit Ru onto Pt film, but we found that MOR activity at such PtRu electrocatalysts are rather small (typically <20 µA/cm$^2$ at 0.5 V) [30], which are similar to what has been reported typically for the PtRu system [31]. The small MOR activity are probably due to the following two factors: (i) Pt ensembles exposed at the surface are not good enough for the dehydrogenation of methanol,- (ii) Ru atoms are much less uniformly distributed in PtRu electrode in which Ru is electrochemically deposited on top of the Pt substrate than the in Pt$_{0.75}$Ru$_{0.25}$ alloy prepared by high temperature annealing under UHV.

The turn-over frequency (TOF) from CH$_3$OH to CO$_2$ on Pt$_{0.73}$Ru$_{0.27}$ is ca. 0.1 molecule/(site-sec), this number is still one order of magnitude smaller than what is required for the operation of practical DMFCs. Further optimization of PtRu nanocatalysts with ensemble structure good for methanol adsorption and its subsequent dehydrogenation, together with active Ru adatoms in its neighborhood which could accelerate oxidation of CO$_{ad}$ and other methanolic-fragments formed at the Pt sites are necessary in order to further enhance the MOR activity. On the other hand, as mentioned above, we do not observe appreciable steady state MOR current at potentials lower than 0.3 V, which is similar to what was found for the PtRu system [32]. Further studies are underway by addition of a third element which may produce more active oxygen species for the oxidation of CO$_{ad}$ while does not significantly inhibit methanol adsorption and dehydrogenation, in order to further reduce MOR overpotential down to 0.3 V.

IV. CONCLUSION

Methanol oxidation at Pt and Pt film electrode deposited with various amount of Ru in 0.1 mol/L HClO$_4$+0.5 mol/L MeOH has been studied under potentiostatic conditions by combining in situ FTIR spectroscopy and differential electrochemical mass spectrometry. We found that on all Pt and PtRu electrodes examined at potentials from 0.3 V to 0.6 V, CO is the only methanol-related adsorbate observed by IR spectroscopy. Ru deposited may only form very small clusters at Pt surface for the cases with Ru coverage below 0.3 ML, since only at Pt$_{0.56}$Ru$_{0.44}$, distinct IR band for CO$_2$ adsorbed at Ru islands is detected. The overall MOR activity and CO$_2$ current efficiency decrease in the order of Pt$_{0.73}$Ru$_{0.27}$>Pt$_{0.56}$Ru$_{0.44}$>Pt$_{0.83}$Ru$_{0.17}$>Pt$_x$, the current density for MOR on Pt$_{0.73}$Ru$_{0.27}$ electrode at 0.5 V is ca. 35 µA/cm$^2$ after normalizing to the active surface area of the Pt film electrode before deposition of Ru, and the current efficiency for CO$_2$ production is ca. 50%, this corresponds to a turn over frequency from CH$_3$OH to CO$_2$ of ca. 0.1 molecule/(site-sec). Possible reasons for the low MOR activity of the as prepared Pt$_x$Ru$_y$ electrocatalysts and strategies for further improvement of MOR electrocatalysts are briefly discussed.

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