

One-step synthesis and chemical characterization of Pt-C nanowire composites by plasma sputtering

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Small size of nanoparticles (NPs) is proposed as responsible of their specific properties.^[1-3] Nanomaterials have thus found numerous applications in chemistry, electronics, optics, information storage, medical, biotechnology.^[4] Numerous processes of heterogeneous catalysis (vehicle exhaust and industrial effluent treatments, petrochemistry, sensors, energy storage and conversion...) used costly and strategic metals from the platinum group.^[5] Here, a high surface-to-volume ratio leading to large fraction of accessible atoms is required for increasing the activity of the catalyst.^[6-8] There is thus a real interest to obtain the highest catalytic surface area from a low metal weight, as for fuel cells^[9] or sensors. On the other hand, carbon nanofibers or carbon nanotubes have been proposed as catalytic support^[10-12] in place of carbon powders because of their high electrical conductivity, unique surface structure, large surface area and chemical inertia.^[13] Moreover, metal NPs deposited on carbon nanotubes or carbon nanofibers display higher electrocatalytic activity and stability.^[14,15] Therefore, numerous multistep syntheses of carbon nanowire arrays were developed, such as template methods^[16] and catalytic growths^[17].

Here we report a plasma-assisted one-step synthesis of Pt/C nanowires (Pt-CNW). Such synthesis method is convenient for controlling the metal spreading on a porous support, its localization in depth and the metal NPs size distribution.^[18,19] It is an environmental-friendly process (time, energy and atom savings, no harmful chemicals, organic solvent or precursor). The co-sputtering of platinum and carbon atoms on an E-Tek® carbon porous layer (CPL) is achieved in a magnetron sputtering reactor. Two rectangular targets, one of pure carbon and one of Pt₁C₉₉ (atomic ratio), are simultaneously DC biased at fixed power leading to a thin film growth rate of ca. 4 nm min⁻¹. Electrodes loaded at 0.010 mg cm⁻² Pt (Pt-CNW/CPL) with a Pt ratio of ca. 30 at% in the deposited film are produced. The Pt loading and localization profile inside the CPL is determined using Rutherford Backscattering Spectroscopy: 90% of Pt atoms are localized in the first 200 nm depth of the material.

The general morphology of Pt-CNW/CPL (Figure 1a) resembles that of a cauliflower with platinum clusters forming dense sponge-like film on the surface of carbon grains. The

TEM image in Figure 1b shows that a platinum film (darker zones) is deposited on the top of the CPL surface (light grey zones). Columnar structures emerge in the radial direction of the platinum film with respect to the CPL support. Higher magnification in Figure 1c reveals a Pt NPs arrangement surrounding carbon nanowires over their whole length. The carbon column enlargement from the bottom (ca. 5 nm) to the top (ca. 10 nm) is explained in terms of a shadowing effect in magnetron sputtering process.^[20] Increasing the magnification allows visualizing the platinum clusters and estimating their size to ca. 1 nm (Figure 1d).

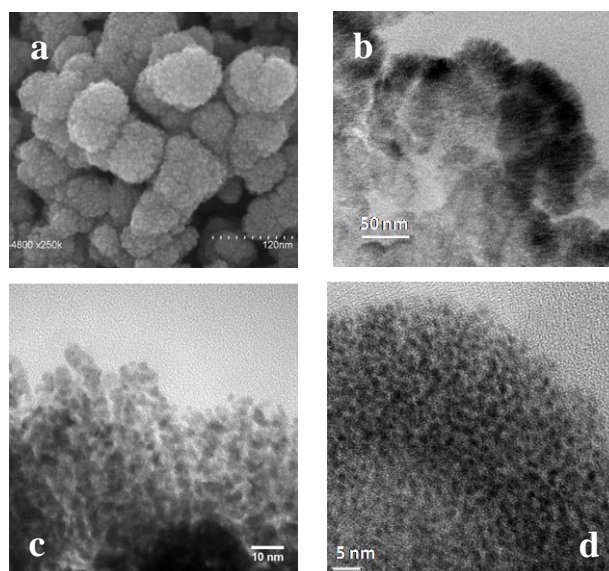


Figure 1. (a) SEM picture of a 0.010 mg cm⁻² Pt on a commercial CPL (courtesy of D. Cot, IEM); (b-d) TEM images of 0.010 mg_{Pt} cm⁻² Pt-CNW on a commercial CPL at different magnifications.

ToF SIMS was used to analyze the chemical structure of the material. Due to the complexity of the sample, a lot of mass peaks are recorded and only semi-quantitative analysis can be performed. CH_xPt and CH_xO_yPt are the main molecular ions detected, indicating that strong chemical bonds exist between Pt atoms and carbon or oxygen atoms, which is a good point for activity and durability issues.^[21] ToF SIMS and XPS (Figure 2) measurements show that Pt is mainly under oxidized states (PtO_x, PtO_xH_y). Platinum is only found under +II oxidation state by XPS. Plasma synthesis is performed under argon atmosphere, free of O₂ and water molecules. Oxygenated species could come from the targets and be sputtered with Pt and C atoms. The edges of graphene planes are likely ended by hydrogen atoms or oxygenated functions.^[22] Because ToF SIMS and XPS measurements were not conducted *in situ*, the oxidation of the Pt and C atoms could also occur as soon as the Pt-CNW/CPL is put in contact with air, forming surface oxides.^[23] XPS analysis of 2.5-3.0 nm Pt NPs synthesized by

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colloidal routes^[24,25] showed that Pt atoms on particle surface were +II oxidation state, whereas core Pt atoms remained under metallic state.^[26] For Pt particles co-deposited with carbon by plasma method, all Pt atoms are found to be oxidized, confirming that the Pt particle size is close to 1 nm. In the case of (hemi)spherical Pt particles,^[19] the dispersion D (ratio of surface N_s to total N_t atom numbers) is inversely proportional to the diameter of the particle d in nm^[27]; 5 nm diameter particle has 20% surface atoms ($N_t \approx 6\,000$ atoms), and for 1 nm, one can consider all atoms as surface atoms. The XRD pattern showed no diffraction peak corresponding to Pt fcc structure, only a broad peak at ca. 25° characteristic of carbon, several peaks assigned to the presence of PTFE in the commercial CPL and a potato-like background, which translates the very low Pt loading and a very amorphous structure. This result is coherent with the very small particle size suggested by TEM and XPS measurements.

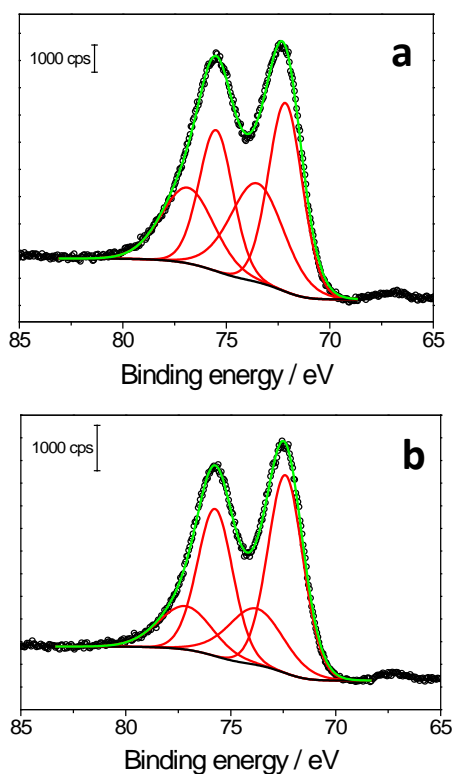


Figure 2. XPS spectra for Pt 4f orbital of $0.010 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ Pt-CNW deposited on (a) a Si wafer and (b) a commercial CPL.

Integration of the hydrogen desorption current in the voltammograms (Figure 3) gives the electrochemical active surface area (EASA)^[28,29]. Values ranging from 150 to $200 \text{ m}^2 \text{ g}_{\text{Pt}}^{-1}$ were obtained for Pt-CNW/CPL against ca. $50 \text{ m}^2 \text{ g}^{-1}$ for a reference Pt (40wt.%) /C catalyst made by a colloidal method.^[30] Assuming spherical particles of similar radius and whole Pt particle surface utilization, a mean particle size d of ca. $1.3 - 1.8 \text{ nm}$ is calculated.^[26] Values are higher than that estimated from TEM. The contact between Pt NPs in agglomerates and between Pt NPs and carbon can avoid the accessibility to Pt sites. $\text{C}_x\text{H}_y\text{O}_z$ species formed on the Pt surface can also be involved. The assumption of spherical particle could also not be valid. After numerical simulation the

equilibrium shape of small nanometric clusters could be the icosahedron,^[31] which is the most compact shape close to the spherical one, but the equilibrium shape of a supported crystal is the Wulff polyhedron for a free crystal truncated at the interface (here a truncated cuboctahedron).

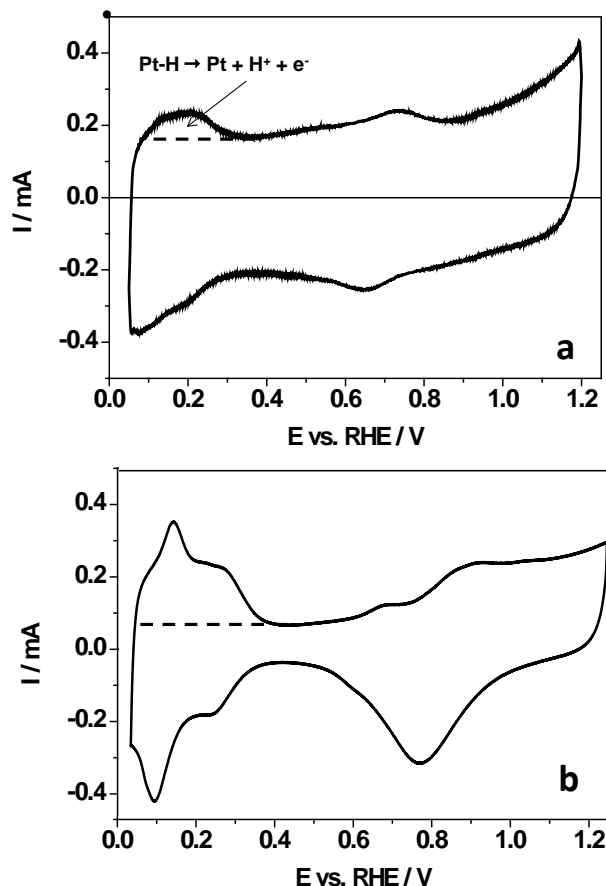


Figure 3. Cyclic voltammograms recorded on (a) $0.010 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ Pt-CNW/CPL electrode and (b) classical Pt/C electrode with 117 µg cm^{-2} of Pt for powders with metal loading of 40 wt.%. (scan rate = 50 mV s^{-1} , N_2 saturated, $0.05 \text{ mol dm}^{-3} \text{ H}_2\text{SO}_4$, $T = 20^\circ\text{C}$).

The oxygen reduction reaction (ORR) at the cathode of Proton Exchange Membrane fuel cells is very challenging and is activated by platinum-based catalysts for reaching acceptable kinetics.^[32-34] Pt-CNW/CPL materials were then evaluated as cathode catalyst under the harsh fuel cell working conditions. The voltage/power density vs current density curve in figure 4 was obtained with two $0.010 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ Pt-CNW/CPL electrode pressed on a Nafion 212 membrane. At 80°C , a maximum power density of 0.40 W cm^{-2} is achieved, i.e. $20 \text{ kW g}_{\text{Pt}}^{-1}$ (highest platinum utilization ever obtained). Fuel cells performances obtained with the reference Pt(40 wt.%) /C catalyst were previously found higher, with a maximum power density of ca. 1.2 W cm^{-2} , but with Pt loading of ca. 0.35 mg cm^{-2} in electrodes, leading to a Pt utilization rate of only $1.7 \text{ kW g}_{\text{Pt}}^{-1}$.^[30]

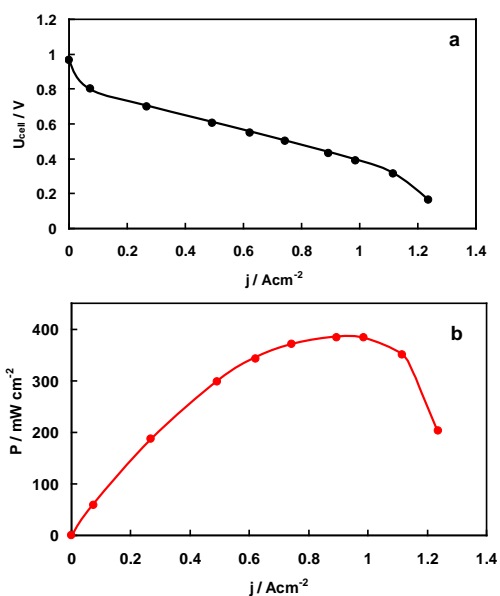


Figure 4. (a) $E(j)$ and (b) $P(j)$ curves obtained at 70°C with a MEA of Nafion 212 membrane sandwiched between symmetrical 0.010 mg_{Pt} cm⁻² Pt-CNW/CPL electrodes (Flow rates: O₂ = 350 sccm, H₂ = 500 sccm; T_{O₂} = 40° C, T_{H₂} = 80° C, P_{O₂} = 4bars, P_{H₂} = 3 bars).

The controlled co-sputtering of platinum and carbon on a porous carbon layer led to a one step formation of carbon nanowires decorated by platinum NPs with diameters from 1 to 2 nm. Strong chemical bonds exist between platinum atoms and carbon or oxygen atoms, and platinum is observed mainly under oxidized states. Very high active surface area and catalytic activity towards ORR were obtained. The film structure, consisting in very small Pt particles decorating carbon columns with strong interactions, is responsible of the very high metal utilization rate, as exemplified by the electrochemical performances of the material as fuel cell electrodes.

Migration/aggregation of Pt crystallites, Ostwald ripening, dissolution of metal and carbon support corrosion are the main degradation processes of Pt/C classical electrodes. The strong interaction between the carbon support and the platinum atoms is according to Zhou et al.²¹ a good point for activity and durability issues. According to the Gibbs-Thompson relation the thin film layer structure composed of agglomerated metallic nanoclusters is expected to cause an increase in their chemical potential, and to translate into lower Pt corrosion kinetics.^[35] The relatively dense thin film of Pt cluster surrounding the carbon nanowires (at the nanoscale level) and the carbon CPL (at a higher scale) is expected to lower metal/carbon/oxygen contact, which could decrease the support corrosion rate. Moreover, the procedure for fuel cell tests imposed the condition that, before fuel cell experiments for several days, the cell had to be conditioned for 2 days at 80 °C with humidified hydrogen and oxygen fed to the anode and the cathode, respectively. Therefore, because the cell performance was maintained for several days, the nanocomposite structure could be considered to be stable on the FC experimental timescale.

Although the electrode fabrication process is relatively expensive (targets and high vacuum conditions) and fuel cell performances remain too low for automotive applications, where the electric kW cost has to be low (EU FP7 and US DoE targeted 50 € / kW for electric vehicle), such technology of

electrode fabrication is quite applicable for stationary and portable applications. Moreover, physical deposition methods under high vacuum conditions are routinely used in many sectors of industry, especially in metallurgy and microelectronics domains, where the cost reduction is primordial. At last, the very high Pt utilization makes this material convenient for sensor applications, where the environmental (temperature, pressure, corrosive medium) and electrochemical conditions are smoother.

Experimental Section

SEM and TEM measurements are carried out with a Hitachi S-4800 and a JEOL JEM 2010 (equipped with a LaB6 filament and with a resolution of 0.35 nm), respectively.

XPS measurements are performed with an Escalab MKII (VG scientific) set-up using the Magnesium monochromatic beam (1253.6 eV), at room temperature below 8 10⁻⁹ Torr pressure in the analysis chamber. Core level spectra are recorded with a 50 meV resolution. Binding energies are lined up with respect to the C 1s peak at 284.6 eV.

TOF SIMS measurements are carried out with an IONTOF SIMS IV apparatus on different zones (analyzed surface = 1600 μm²) of a Pt-CNW/CPL to check the homogeneity of the sample.

Electrochemical measurements are performed with a computer controlled Voltalab PGZ 402 potentiostat, at 20°C in N₂-saturated (U quality from l'Air Liquide) 1.0 M NaOH (suprapur from Aldrich) electrolyte. A glassy carbon plate as counter electrode and a reversible hydrogen electrode as reference are used; all potentials are referred to the reversible hydrogen electrode (RHE).

For fuel cell measurements two 5 cm² surface area 0.010 mg_{Pt} cm⁻² Pt-CNW/CPL electrodes are mechanically pressed on a Nafion 212 membrane (Quintech, Germany) at 2 Nm torque in the cell hardware (Electrochem). Measurements are performed using a ECL150/MTS 150/HSA unit (Electrochem. Inc.).

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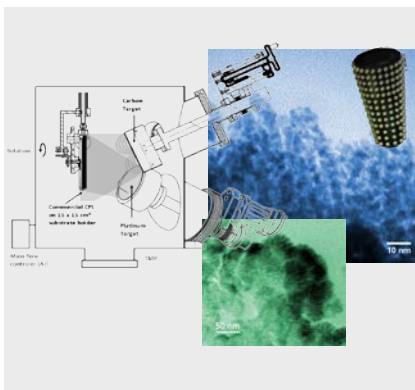
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COMMUNICATION

A one-step synthesis of Pt-C nanowire composites using a plasma co-deposition method is reported. Electrodes with a Pt loading as low as 0.010 mg cm^{-2} are obtained. Pt particles with sizes from 1 to 2 nm are decorating with strong interactions columnar carbon nanostructures. The composite microstructure is responsible of the very high metal utilization rate as exemplified from reactions involved in fuel cell electrodes ($20 \text{ kW g}_{\text{Pt}}^{-1}$).



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One-step synthesis and chemical characterization of Pt C nanowire composites by plasma sputtering

Supporting informations

One-step synthesis of Pt-C nanowire composite materials by plasma sputtering

C. Coutanceau, P. Brault, A. Caillard, S. Baranton, M. Mougnot, S. Cuyenet

Plasma co-sputtering method for synthesis

Pt-CNW are synthesised on a commercial CPL (Elat from E-Tek) in a plasma sputtering apparatus. Figure 1 shows the TEM image of the CPL.

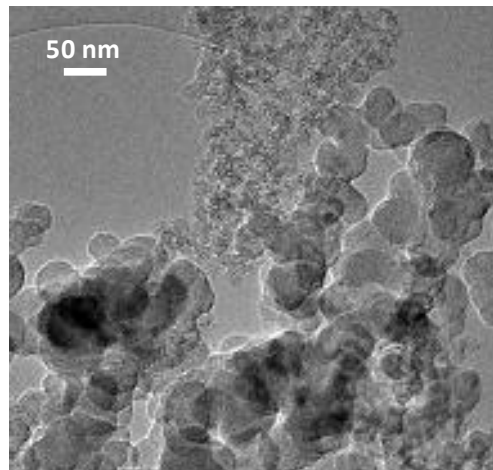


Figure 1: TEM image of the CPL used for Pt-C co-deposition by plasma sputtering.

Before deposition process, the 304L cylindrical stainless steel chamber is pumped to $5 \cdot 10^{-6}$ mbar using a 500 L s^{-1} turbomolecular pump (TV-551 Navigator™ from Varian®). The CPL is introduced into the deposition chamber through a load lock unit and placed on a $20 \times 20 \text{ cm}^2$ substrate holder. The argon pressure is adjusted to $75 \text{ }\mu\text{bar}$. Two high purity planar targets (one of pure carbon the other of Pt₁C₉₉ atomic ratio) are simultaneously DC biased at fixed power of 530 W (target current 0.78 A , target voltage 700 V). The targets are placed at a distance of 10 mm from the substrate holder at an angle of 45° with respect to the substrate holder axis. The deposition rate of each material, the Pt loadings and the distributions into the commercial CPL were measured using Rutherford Backscattering Spectroscopy (RBS)^[1] in the laboratory CEMHTI (Orleans, France). A 2 MeV α particle beam ($1 \times 1 \text{ mm}^2$) produced in a Van de Graaf accelerator impinges onto the Pt-CNW material. The α particles are backscattered after colliding with a carbon atom or a Pt atom and they are collected by a 25 mm^2 α detector (placed at 70 mm from the sample with a scattering angle of 165°). Their energy depends on the mass and on the in-depth position of the collision. The sticking coefficients of Pt and C atoms on Si were found to be equal to the one on commercial CPL. The deposition time was adjusting to reach a Pt loading of 0.010 mg cm^{-2} and a ratio of $30 \text{ wt}\%$ of Pt in the deposited film. Figure 2 shows the RBS spectrum of the $0.01 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ loaded CPL. On the right side of the spectrum, the asymmetric peak corresponds to the Pt atoms. The area under this Pt peak is related to the number of Pt atoms inside the Pt-CNW material. The tail on the left side of the Pt peak indicates that the Pt atoms have diffused into the porous CPL. Pt distribution in the depth of the material can be deduced^[2].

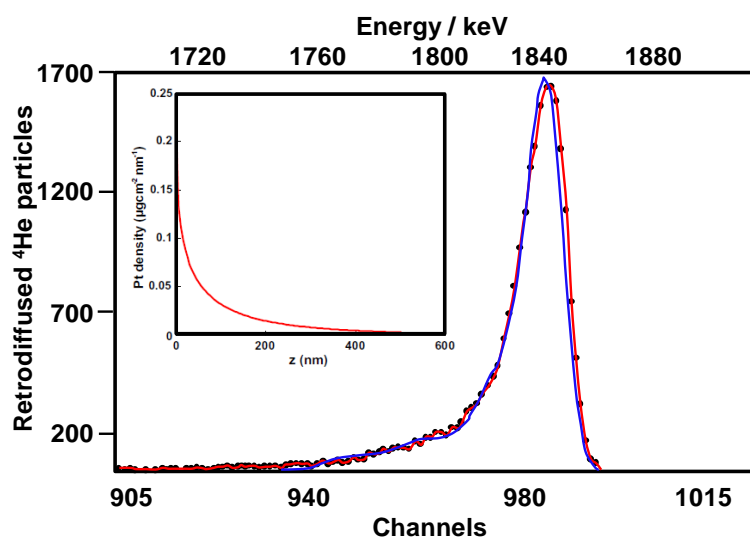


Figure 2: RBS spectrum of Pt from which the platinum density profile (inset) in the composite Pt-CNW/CPL material was determined; experimental data are represented by the red curve, the blue curve is the simulated one to access the density profile.

ToF SIMS analysis

The ToF SIMS surface analysis technique consists in impinging the surface of a sample with a primary ion beam (in the present case $^{69}\text{Ga}^+$) and to analyze the ejected secondary ions. By measuring the ratio masse/charge (m/z) of secondary ions and their time of flight between the sample and the detector, the elemental composition and the chemical structure of the material upper layers can be determined. The depth resolution is 1-3 monolayers, the atomic resolution is lower than 5×10^{11} at cm^{-2} . Three different zones of the samples were measured in order to check the homogeneity of the material. The analysis chamber pressure is set at 5×10^{-9} Torr. The primary ions source of $^{69}\text{Ga}^+$ is powered at potential of 15 kV leading to 0.67 pA pulsed ionic current (100 ns). The Molecular ions with mass from 1 to 400 containing C, O, H and Pt were searched in the spectra corresponding to positive and negative polarity of ions (Table 1). Several molecular ions could correspond to the same m/z value; however, the DtMass value may help to discriminate the molecular ions: the lower is its absolute value, the closer is the measured mass to the calculated one (the more probable molecular ions are in bold characters). Moreover, due to dependence of the ionic yield of secondary ions on the matrix, the ion intensity is not directly proportional to their concentration; however, while the same matrix is involved for all secondary ions, semi-quantitative analysis could be performed, and the main molecular fragments, in terms of amount, could be pointed out.

Molecular peak from ToF SIMS

Positive molecular ions	Sample 1			Sample 2			Sample 3		
	Ct Mass	Dt Mass	CI	Ct Mass	Dt Mass	CI	Ct Mass	Dt Mass	CI
PtO	210,9819	105,5	63,38	210,9819	100,8	70,16	210,9819	98,1	64,96
CH₄Pt	210,9819	-67,1	63,38	210,9819	-71,8	70,16	210,9819	-74,4	64,96
PtOH	211,9839	77,2	70,64	211,9839	75,2	75,37	211,9839	77,7	75,71
CH ₃ Pt	211,9839	-94,6	70,64	211,9839	-96,5	75,37	211,9839	-94	75,71
PtH₂O	212,9861	50,7	48,44	212,9861	43,3	56,55	212,9861	44,6	60,65
CH ₂ Pt	212,9861	-120,2	48,44	212,9861	-127,6	56,55	212,9861	-126,3	60,65
PtH₄O	214,9866	-20,4	14,26	214,9866	-14,8	18,02	214,9866	-31,4	15,55
PtO ₂	227,0004	201,5	13,33	227,0004	203,4	14,82	227,0004	198,8	13,68
CH₄OPt	227,0004	41,2	13,33	227,0004	43	14,82	227,0004	38,4	13,68
PtO ₂ H	228,0083	201,5	29,96	228,0083	198,9	37,81	228,0083	194,8	40,62
CH₅OPt	228,0083	41,9	29,96	228,0083	39,3	37,81	228,0083	35,1	40,62
PtH ₂ O ₂	229,0104	175,3	35,95	229,0104	176,7	39,97	229,0104	176	40,7
CH₆OPt	229,0104	16,8	35,95	229,0104	17,7	39,97	229,0104	17,1	40,7
PtH₃O₂	230,0132	152,4	23,21	230,0132	148,9	28,67	230,0132	151	27,03
C₃H₂Pt	232,9772	-13,9	17,98	232,9772	-19	19,14	232,9772	-16,1	15,39
C₃H₃Pt	233,979	-39,8	18,57	233,979	-37,8	21,15	233,979	-29,5	23,2
C ₂ OPt	234,9865	114,2	33,51	234,9865	115,2	33,88	234,9865	118,5	30,77
C₃H₄Pt	234,9865	-40,7	33,51	234,9865	-39,8	33,88	234,9865	-36,4	30,77
C ₂ HOPt	235,9938	111,5	23,8	235,9938	99,7	28,35	235,9938	104,7	25,97
C₃H₅Pt	235,9938	-42,8	23,8	235,9938	-54,5	28,35	235,9938	-49,5	25,97
C₂H₂OPt	236,9905	64,3	23,88	236,9905	68,4	24,19	236,9905	68,2	24,83
C ₃ H ₆ Pt	236,9905	-89,3	23,88	236,9905	-85,2	24,19	236,9905	-85,4	24,83
CHO ₂ Pt	239,9919	122,9	15,44	239,9919	105,3	17,62	239,9919	103,5	14,82

C ₂ H ₅ OPt	239,9919	-28,8	15,44	239,9919	-46,3	17,62	239,9919	-48,1	14,82
CH ₂ O ₂ Pt	240,9862	66,1	12,66	240,9862	69,8	11,21	240,9862	74,1	10,75
C ₂ H ₆ OPt	240,9862	-84,6	12,66	240,9862	-81,3	11,21	240,9862	-76,9	10,75
C ₃ H ₃ OPt	250,0039	82,9	18,82	250,0039	76,4	20,1	250,0039	77	15,96
C ₂ O ₂ Pt	251,0009	184,4	14,35	251,0009	187,8	15,62	251,0009	178,7	15,47
C ₃ H ₄ OPt	251,0009	39,4	14,35	251,0009	42,7	15,62	251,0009	33,7	15,47
CH ₅ O ₃ Pt	259,9946	22,7	23,13	259,9946	17,3	22,75	259,9946	17	21,9
CH ₆ O ₃ Pt	260,9981	6,2	27,77	260,9981	1,6	28,03	260,9981	3,1	28,33
C ₃ O ₂ Pt	263,0022	181,1	17,89	263,0022	172,6	21,47	263,0022	171	21,41
C ₃ HO ₂ Pt	264,006	165,3	12,41	264,006	165	12,9	264,006	154,4	12,37
C ₂ H ₄ O ₃ Pt	270,9894	31,4	13,33	270,9894	38,6	15,86	270,9894	38	13,76
C ₂ H ₅ O ₃ Pt	271,9958	26,2	23,21	271,9958	17,7	23,47	271,9958	23,2	23,04
C ₂ H ₆ O ₃ Pt	273,0012	17,2	23,46	273,0012	13,2	26,19	273,0012	9,1	24,67
C ₂ H ₇ O ₃ Pt	274,0088	16,3	16,12	274,0088	0,1	19,94	274,0088	6,1	18,89
C ₃ H ₅ O ₃ Pt	283,9921	12	15,44	283,9921	15,7	14,34	283,9921	12	14,82
C ₃ H ₆ O ₃ Pt	285,0017	18,2	16,04	285,0017	23,1	15,38	285,0017	16,7	16,28
C ₃ H ₇ O ₃ Pt	286,006	5,9	13,17	286,006	17,3	15,62	286,006	10,9	13,76

Negative molecular ions	Sample 1			Sample 2			Sample 3		
	Ct Mass	Dt Mass	CI	Ct Mass	Dt Mass	CI	Ct Mass	Dt Mass	CI
¹⁹⁴ Pt	193,9519	-55,6	23,17	193,9519	-59,3	20,25	193,9519	-31,2	20,36
Pt	194,9584	-32,6	40,24	194,9584	-37	41,63	194,9584	-11,8	35,48
¹⁹⁶ Pt	195,9613	-57,5	44,03	195,9613	-58,5	44,08	195,9613	-43,6	33,95
PtH ₂	196,9708	-48,8	27,63	196,9708	-49,6	27,56	196,9708	-48,5	15,45
¹⁹⁷ Pt	197,974	-72,3	15,64	197,974	-72,3	15,75	197,974	-87,8	8,16
CH ₃ Pt	209,963	-120,2	13,73	209,963	-121,3	15,75	209,963	-122,7	10,28
CH ₄ Pt	210,9575	-183	17,59	210,9575	-179,4	16,88	210,9575	-159,2	12
CH ₅ Pt	211,9627	-194,4	20,1	211,9627	-196,9	21,82	211,9627	-194	12,93
C ₂ Pt	218,9599	-22,3	19,66	218,9599	-17,8	19,77	218,9599	-1,7	10,88
C ₂ HPt	219,9616	-50,3	64,64	219,9616	-57,8	64,21	219,9616	-34,2	28,18
C ₂ H ₂ Pt	220,9654	-68	87,33	220,9654	-71,7	83,38	220,9654	-46,5	34,35
C ₂ H ₃ Pt	221,9666	-97,5	73	221,9666	-103,6	71,68	221,9666	-79,4	28,78
C ₂ H ₄ Pt	222,9678	-127	60,11	222,9678	-131,6	61,6	222,9678	-127,9	19,1
COPt	222,9678	36,2	60,11	222,9678	31,7	61,6	222,9678	35,3	19,1
C ₂ H ₅ Pt	223,9632	-181,8	44,18	223,9632	-181,2	47,17	223,9632	-151,3	15,32
CHOPt	223,9632	-19,3	44,18	223,9632	-18,6	47,17	223,9632	11,2	15,32
C ₂ H ₆ Pt	224,9671	-198,5	31,29	224,9671	-204,9	30,74	224,9671	-172,3	8,69
CH ₂ OPt	224,9675	-34,8	30,65	224,9675	-41,1	29,98	224,9675	-10,5	8,69
CH ₃ OPt	225,959	-107,2	16,52	225,959	-106,4	17,32	225,959	-78,6	6,57
CH ₄ OPt	226,9535	-165,4	16,64	226,9535	-168,2	19,41	226,9535	-155,3	6,83
PtO ₂	226,9535	-5	16,64	226,9535	-7,8	19,41	226,9535	5,1	6,83
PtH ₃ O ₂	229,9371	-178,2	20,34	229,9371	-178,8	18,24	229,9371	-180,3	11,34
C ₃ Pt	230,9408	-104,2	23,88	230,9408	-112,4	24,91	230,9408	-101,5	11,67
PtH ₄ O ₂	230,9408	-195,6	23,88	230,9408	-203,8	24,91	230,9408	-192,8	11,67
C ₃ HPt	231,9498	-98,7	24,36	231,9498	-110,8	24,91	231,9498	-86,3	10,48
C ₃ H ₂ Pt	232,9636	-72,6	21,61	232,9636	-82,9	22,62	232,9636	-76,3	8,75
C ₃ H ₃ Pt	233,9668	-92	22,37	233,9668	-97,7	21,74	233,9668	-77,8	7,76
C ₂ HOPt	235,9669	-2,6	24,76	235,9669	-6,4	25,03	235,9669	29,6	9,28
C ₃ H ₅ Pt	235,9669	-156,8	24,76	235,9669	-160,6	25,03	235,9669	-124,6	9,28
C ₂ H ₂ OPt	236,9668	-36	28,82	236,9668	-35,4	33,15	236,9668	-7	9,75
C ₃ H ₆ Pt	236,9668	-189,6	28,82	236,9668	-189	33,15	236,9668	-160,7	9,75
C ₂ H ₃ OPt	237,9685	-61,8	27,67	237,9685	-67,5	30,22	237,9685	-42,9	7,69
C ₃ H ₇ Pt	237,9685	-214,7	27,67	237,9685	-220,4	30,22	237,9685	-195,9	7,69
CO ₂ Pt	238,9625	33,1	30,41	238,9625	25,4	31,5	238,9625	55,8	7,89
C ₂ H ₄ OPt	238,9625	-119,2	30,41	238,9625	-126,9	31,5	238,9625	-96,6	7,89
CHO ₂ Pt	239,9595	-12,1	20,94	239,9595	-20,1	22,14	239,9595	20,4	6,37
C ₂ H ₅ OPt	239,9595	-163,8	20,94	239,9595	-171,8	22,14	239,9595	-131,3	6,37
CH ₂ O ₂ Pt	240,9574	-53,5	16,76	240,9574	-58,5	15,79	240,9574	-16,7	5,04
C ₂ H ₆ OPt	240,9574	-204,5	16,76	240,9574	-209,5	15,79	240,9574	-167,7	5,04
PtO ₃	242,9563	28	16,44	242,9563	24,3	16,39	242,9563	59,6	5,97

CH ₄ O ₂ Pt	242,9563	-121,8	16,44	242,9563	-125,5	16,39	242,9563	-90,2	5,97
PtO₃H	243,9604	12,7	15,96	243,9604	-0,1	16,96	243,9604	55,3	4,64
CH ₅ O ₂ Pt	243,9604	-136,5	15,96	243,9604	-149,4	16,96	243,9604	-93,9	4,64
PtH₂O₃	244,9636	-6,6	26,87	244,9636	-16,5	25,6	244,9636	1,5	7,83
CH ₆ O ₂ Pt	244,9636	-155,2	26,87	244,9636	-165,1	25,6	244,9636	-147,1	7,83
PtH₃O₃	245,9632	-39,8	32,6	245,9632	-41,8	32,35	245,9632	-18,7	9,42
C₃OPt	246,9663	26,5	37,66	246,9663	22,4	38,86	246,9663	45,6	12
PtH ₄ O ₃	246,9663	-58,9	37,66	246,9663	-63	38,86	246,9663	-39,8	12
C₃HOPt	247,9603	-29,3	39,69	247,9603	-25,3	38,25	247,9603	-3,7	10,08
C₃H₂OPt	248,9639	-46,3	33,04	248,9639	-53,4	32,51	248,9639	-12,6	7,23
C₃H₃OPt	249,9621	-84,3	19,7	249,9621	-80,2	19,81	249,9621	-61,5	5,31
C₂HO₂Pt	251,9595	-11,8	11,54	251,9595	-14,5	11,93	251,9595	-2	2,85
C ₃ H ₅ OPt	251,9595	-156,2	11,54	251,9595	-159	11,93	251,9595	-146,5	2,85
C₂H₂O₂Pt	252,9538	-65,4	11,86	252,9538	-59,4	12,13	252,9538	-37,8	3,12
C ₃ H ₆ OPt	252,9538	-209,3	11,86	252,9538	-203,3	12,13	252,9538	-181,7	3,12
C₂H₃O₂Pt	253,9492	-113,9	13,45	253,9492	-104	12,94	253,9492	-109,8	4,18
CO₃Pt	254,9366	-50,6	20,82	254,9366	-48,6	22,5	254,9366	-21,9	5,84
C ₂ H ₄ O ₂ Pt	254,9366	-193,4	20,82	254,9366	-191,4	22,5	254,9366	-164,7	5,84
C₁6H₃1O₂	255,2263	-23,7	36,78	255,2263	-34,6	37,89	255,2263	10,6	10,15
CHO₃Pt	255,9389	-72,3	26,15	255,9389	-71,6	23,91	255,9389	-46,1	8,09
C ₂ H ₆ O ₂ Pt	256,9319	-271,4	35,39	256,9319	-278,2	33,79	256,9319	-246	9,15
CH₂O₃Pt	256,9319	-129,8	35,39	256,9319	-136,5	33,79	256,9319	-104,4	9,15
C ₂ H ₇ O ₂ Pt	257,9323	-298,8	29,62	257,9323	-302,6	27,12	257,9323	-258,7	8,02
CH₃O₃Pt	257,9323	-157,7	29,62	257,9323	-161,5	27,12	257,9323	-117,6	8,02
CH₄O₃Pt	258,9272	-207,2	25,64	258,9272	-209	23,71	258,9272	-177	7,23
C₃O₂Pt	262,9635	33,5	19,78	262,9635	29,5	17,52	262,9635	21,9	4,97
C₃HO₂Pt	263,9576	-18,4	17,2	263,9576	-21	18,04	263,9576	-3,8	4,18
C₃H₂O₂Pt	264,9558	-54,8	16,2	264,9558	-58,5	16,88	264,9558	-19	4,11
C₃H₃O₂Pt	265,9479	-113,5	13,77	265,9479	-98,5	14,02	265,9479	-105,5	3,05
C ₃ H ₄ O ₂ Pt	266,9536	-121,4	12,7	266,9536	-113,8	12,09	266,9536	-69,9	2,79
C₂O₃Pt	266,9536	15	12,7	266,9536	22,5	12,09	266,9536	66,4	2,79
C ₃ H ₅ O ₂ Pt	267,9562	-140,3	11,58	267,9562	-145,2	11,29	267,9562	-123,1	3,38
C₂HO₃Pt	267,9562	-4,5	11,58	267,9562	-9,4	11,29	267,9562	12,7	3,38
C ₃ H ₆ O ₂ Pt	268,9605	-152,7	14,01	268,9605	-153,1	12,09	268,9605	-101,7	3,18
C₂H₂O₃Pt	268,9605	-17,4	14,01	268,9605	-17,8	12,09	268,9605	33,6	3,18
C₂H₃O₃Pt	269,9638	-34,1	12,06	269,9638	-37,9	11,21	269,9638	-17,2	3,18
C₂H₄O₃Pt	270,9621	-69,1	12,78	270,9621	-63,6	12,62	270,9621	-19	3,91

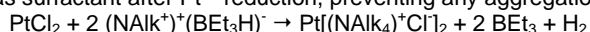
Table 1 : Molecular ions with mass from 1 to 400 containing C, O, H and Pt obtained in the spectra corresponding to positive and negative polarity of ions (CtMass = centre of mass; DtMass = mass deviation; Cl = signal intensity). The more probable molecular ions are given in bold characters.

XPS measurements

Oxidation states of Pt were evaluated by X-ray photoelectron spectroscopy (XPS). The deconvolutions of the XPS spectra are carried out using a Shirley function for the background. The separation between peaks corresponding to Pt4f7/2 and Pt4f5/2 is fixed to 3.35 eV, and the ratio was fixed at 3/2. 1 x 1 cm silicon 100 is used as second substrate for the XPS measurement on Pt/(C nanowires)/Si material. The peaks corresponding to Pt 4f7/2 are located at binding energies of 72.2 eV (Si wafer) / 72.4 eV (CPL), and 73.5 eV (Si wafer) / 73.2 eV (CPL), corresponding to metallic Pt(OH)₂ and PtO, the Pt 4f7/2 peak related to metallic Pt⁰ species being located at binding energy lower than 72 eV.

Electrochemical measurements

The procedure described by Bönnemann and co-workers^[3] was slightly modified and adapted for the preparation of the reference Pt/C catalyst material.^[4] Syntheses were carried out under atmosphere free of oxygen and water, using non-hydrated PtCl₂ (99.9% from Alfa Aesar). The first step consisted in the preparation of a tetraalkyltriethylborohydride reducing agent (NAlk₄)⁺(BEt₃H)⁻, which also acts as surfactant after Pt²⁺ reduction, preventing any aggregation of the metallic particles:



The colloidal precursor was then dispersed on a carbon support (Vulcan XC72) and treated at 573 K for 90 min under air to entirely remove the organic surfactant, as shown by thermo-gravimetric analyses (TGA) [24]. Platinum loadings of 40 wt.% was expected by adjusting the amount of Vulcan XC72 added to the platinum colloid before heat treatment (Pt(40 wt.%)/C).

The working electrode from the reference Pt/C material was prepared by deposition of a catalytic ink on a 0.071 cm² gold disc. The catalytic powder was added to a mixture of 25 wt.% (based on the powder content) Nafion solution (5 wt.% in aliphatic alcohols from Aldrich) and ultra-pure water (MilliQ, Millipore, 18.2 MX cm). After ultrasonic homogenization of the Pt/Vulcan XC72-Nafion ink, a given volume was deposited from a syringe onto a fresh polished gold substrate yielding a catalytic powder loading of 293 μg cm⁻²,

i.e. 117 lg cm⁻² of platinum for powders with metal loading of 40 wt.%. The solvent was then evaporated under pure nitrogen flow at room temperature.

The electrochemical active surface area (EASA) of the catalysts is evaluated from the cyclic voltammograms recorded at 20 mV s⁻¹ in support electrolyte (H₂SO₄ 0.5 M), by integration of the charge involved in the adsorption–desorption region of hydrogen between 0.05 and 0.4 V after correction of the capacitive contribution of the carbon support^[5] and assuming a charge of 210 μC cm⁻² for the adsorption of a monolayer of atomic hydrogen on a smooth polycrystalline platinum surface, according to the following equation:^[6]

$$EASA = \frac{1/v \int IdV}{210 \times 10^4 \times m_{Pt}}$$

where EASA is the electrochemical active surface area (m² g⁻¹), v the linear potential scan rate (V s⁻¹), i the current (μCs⁻¹), V the electrode potential (V) and m_{Pt} the mass of platinum deposited on the electrode (g).

Fuel Cell measurements

For Membrane electrode assembly (MEA) fabrication, a membrane (a Nafion 212 membrane, purchased from Quintech, Germany, noted further in the text N212) was sandwiched without pre-humidification between two electrodes (an anode and a cathode of 5 cm² geometric surface each) used as prepared by plasma sputtering (without Nafion solution addition). The MEA is prepared by simply tightening the membrane and symmetric electrodes between two graphite bipolar plates at a 2 Nm torque. No preliminary hot pressing process of electrodes against the Nafion 212 membrane (used as received from Electrochem. Inc.) was performed. No Nafion solution was added in the electrode catalytic layers before use. Under our testing conditions, the pressure is manually regulated with the output valves and hydrogen and oxygen are humidified by bubbling in water at 80°C and 40°C, respectively. Measurements in 5 cm² PEMFC using pure H₂ and O₂ gases are performed using a ECL150/MTS 150/HSA unit (Electrochem. Inc.).

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