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Degradation by Hydrogen Peroxide of Metal-Nitrogen-Carbon Catalysts for Oxygen Reduction

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Fe-N-C and Co-N-C materials are promising catalysts for reducing oxygen in fuel cells. The degradation of such catalysts induced by H_2O_2 was investigated by contacting them ex situ with various amounts of H_2O_2 . The degradation increased with increasing amounts of H_2O_2 . The effect was most severe for Cr-N-C followed by Fe-N-C and last by Co-N-C. Treatment with H_2O_2 leads to diminished oxygen reduction activity at high potential and/or reduced transport properties at high current density in fuel cell. From spectroscopic characterisation, it was found that 66 and 80% of the CoN_xC_y and FeN_xC_y moieties present in pristine catalysts survived the extensive H_2O_2 treatment, respectively. In parallel, the activity for oxygen reduction was divided by ca 6–10 for Fe-N-C and by ca 3 for Co-N-C. The results suggest that the main degradation mechanism in fuel cell for such catalysts is due to a chemical reaction with H_2O_2 that is generated during operation. The super-proportional decrease of the oxygen reduction activity with loss of FeN_xC_y and CoN_xC_y moieties suggests either that only a small fraction of such moieties are initially located on the top surface, or that their turnover frequency for oxygen reduction was drastically reduced due to surface oxidation by H_2O_2 .

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27 Electrochemical energy conversion devices are expected to play an increasing role for stationary and transportation applications as well as 28 for reversible energy storage. Among the various applications already 29 in use or contemplated for electrochemical devices, the transportation 30 sector is the most demanding in terms of energy density, power density 31 and cost.1 Driving a full-size car over 100 km requires ca 13 kWh of 32 mechanical energy.² While batteries are more efficient than fuel cells, 33 the higher energy density of H₂/air fuel cells relative to the lithium-34 ion battery (1.8 kWh per kg of H2-reservoir at 690 bars; compared to 35 0.2 kWh per kg of Li-ion battery pack) is critical for designing a system 36 that can propel a car over 500 km without refuelling or recharging.² 37 Hence, the polymer electrolyte membrane fuel cell (PEMFC) is to-38 day the most promising electrochemical device for replacing internal 39 combustion engines, with uncompromised driving range and power 40 density.³ Mid-term remaining challenges for PEMFCs are increased 41 durability and reduced cost, while a longer-term challenge is the re-42 placement of the rare and expensive platinum element by catalysts 43 based on Earth-abundant metals.^{1,4-5} Today, ca 80-90% of the mass 44 of Pt in a PEMFC stack is placed at the cathode due to the sluggish 45 oxygen reduction reaction (ORR) compared to fast hydrogen oxida-46 tion at the anode. Based on projections for mass production, Pt-based 47 catalysts in today's PEMFC technology are expected to account for 48 ca 50% of the material's cost of a fuel cell stack.^{4,6} Significantly re-49 ducing the Pt loading in cathodes or eliminating the platinum metal 50 altogether from ORR catalysts are two possible approaches to tackle 51 this cost problem.³ 52

While the concept of ORR catalysis by materials or molecules 53 comprising Fe or Co, nitrogen and carbon elements is known since 54 1964,⁷ breakthroughs in the ORR activity and fuel cell performance 55 that were necessary to seriously consider Fe(Co)-N-C catalysts as po-56 tential substitutes to platinum in the acidic environment of PEMFCs were reported only after 2008.^{2,8–14} While major advances in the ORR 57 58 activity and power density of PEMFC comprising Fe(Co)-N-C based 59 cathodes have been reported, the next grand scientific challenges are 60 i) improved understanding of the structure of the active sites and 61

ii) improved understanding of the degradation mechanisms. Depend-62 ing on the latter, mitigation approaches might be very different: i) 63 system-based solutions to avoid the transient high-potential polariza-64 tion during start-up and shutdown, ii) catalyst-based approaches in 65 order to avoid the formation of H₂O₂ during ORR or to endow the 66 catalysts with a higher graphitic character, iii) novel cathode designs 67 in order to stabilize its electrical, hydrophilic and diffusion properties. 68 A restricted number of studies have focused on the degradation mech-anisms of Fe(Co)-N-C catalysts.^{15–28} Fe-N-C catalysts pyrolyzed in 69 70 NH₃ have shown the highest initial activity and power performance 71 thus far, but degrade very quickly in PEMFC, mostly due to a proto-72 nation and anion-binding effect of highly-basic surface N-groups that 73 are characteristic of NH₃ pyrolysis.¹⁶ Fe(Co)-N-C catalysts synthe-74 sized at high temperature in inert gas are initially less active but are 75 more stable in PEMFC.^{8–10,15,26} Disentangling the various degradation 76 phenomena of Fe(Co)-N-C cathodes occurring during PEMFC opera-77 tion is complex. Water flooding, changes of the hydrophilic properties 78 of the cathode, loss of a fraction of the active sites through dissolu-79 tion or via oxidative attacks generated by either the cathode potential, 80 H_2O_2 or radical oxygen species (ROS) formed between H_2O_2 and 81 the transition-metal-based active sites may all account to some de-82 gree for the overall decline of the cathode performance with time. In 83 a first attempt to disentangle these phenomena, we recently investi-84 gated on a well-defined Fe-N-C catalyst the degradation incurred by 85 high-voltage cycling.²¹ This phenomenon is the least complex one 86 to separately investigate from other degradation routes since it is not 87 linked to the ORR. It was concluded that the FeN_xC_y moieties initially 88 present in the Fe-N-C catalyst were not immediately destroyed after a 89 polarization up to 1.4-1.5 V vs. RHE. However, bulk oxidation of the 90 carbon matrix in which these active sites are embedded ultimately lead 91 to the massive dissolution of FeNxCy moieties. In addition, increased 92 electric resistance of the cathode layer after high potential cycling 93 also contributed to the decline of the cathode performance.²¹ Due to 94 the integration of $Fe(Co)N_xC_v$ moieties at the edge of graphene sheets 95 or within the sheets, these materials are intrinsically more prone to 96 degradation when subjected to high electric potential than Pt/C cat-97 alysts. More graphitic matrices may be resorted to for synthesizing 98 Fe(Co)-N-C catalysts. However, it is notoriously difficult to achieve a 99 high density of $Fe(Co)N_xC_y$ moieties in highly graphitic substrates via 100 pyrolysis.^{23,29} Mitigation of the degradation induced by high-voltage

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¹⁰² on Fe(Co)-N-C catalysts should therefore be most effective by focus-¹⁰³ ing on the causes of the high-voltage transients, namely i) the purging ¹⁰⁴ of uncombusted H_2 at the anode with air and ii) the non-selectivity of ¹⁰⁵ platinum for hydrogen oxidation.^{30–32}

The present work focuses on the oxidative attack of Me-N-C 106 catalysts by minute amount of H₂O₂ that is produced due to the 107 incomplete electro-reduction of O₂ to H₂O during fuel cell operation. 108 This degradation mechanism can probably not be dealt with at a sys-109 tem level and is therefore an inescapable issue for Me-N-C catalysts. 110 While the % H₂O₂ produced during ORR is often not much higher 111 on Fe(Co)-N-C than on Pt/C catalysts,^{25-27,33-34} the residence time 112 of H₂O₂ in the electrode may be longer with Fe(Co)-N-C catalysts 113 due to their extremely poor activity for the electro-reduction of H2O2 114 to H_2O ,^{33,35} and poor activity for the chemical disproportionation of 115 $H_2O_2^{-27,33,35-36}$ Chemical disproportionation of $H_2O_2^{-1}$ is ca two orders 116 of magnitude faster at high pH for such catalysts,³⁶ highlighting the 117 possibility to recycle H₂O₂ at high pH but impossibility to do so at 118 low pH. In contrast, platinum is an excellent catalyst for both the 119 four-electron reduction of O₂ to H₂O and the two-electron reduction 120 of H_2O_2 to H_2O .^{33,37} The important difference between the durability 121 of Pt/C and Fe(Co)-N-C catalysts in PEMFC may thus stem from their 122 different reactivity toward H₂O₂. Moreover, chemical disproportion-123 ation of H₂O₂ on Fe(Co)-N-C catalysts via Fenton chemistry might 124 produce a significant amount of radical oxygen species (ROS).³⁸⁻ 125 While suspected, the role of H₂O₂ in the degradation of Fe(Co)-N-C 126 based cathodes during steady-state operation of a PEMFC has not 127 yet been investigated in depth. Early work by Wiesener showed that 128 addition for every 100 h of a given aliquot of H₂O₂ to the electrolyte 129 lead to a faster degradation of a Co-N-C catalyst.⁴¹ Ex situ chemical 130 degradation of Fe-N-C catalysts with H2O2 was investigated in 2003 131 by Lefèvre.²⁵ A series of catalysts were immersed in H₂O₂ solutions of 132 various concentrations. The ORR activities measured with a rotating-133 disk electrode (RDE) were shown to have strongly decreased. More 134 recently, Zelenay's group applied a different ex situ H₂O₂ treatment, 135 keeping the H_2O_2 concentration fixed but varying the duration of im-136 mersion of the catalyst.²⁶ In addition to measuring the ORR activity 137 in RDE before and after the treatments, also the elemental compo-138 sition was determined by X-ray photoelectron spectroscopy (XPS). 139 A comprehensive understanding of the degradation mechanism of 140 Me-N-C catalysts upon contact with H₂O₂ solution is however still 141 lacking, especially regarding spectroscopic changes of the transition 142 metal coordination. 143

In the present work, we synthesized a Fe-N-C, a Co-N-C and a Cr-N-C catalyst and contacted them with various amounts of H_2O_2 corresponding to ratios of mol H_2O_2 per mass of catalyst ranging over ca two orders of magnitudes. The ORR activities of the catalysts before and after H_2O_2 treatment were determined with RDE and in PEMFC. The pristine and treated materials were characterized with X-ray diffraction, X-ray absorption spectroscopy, X-ray photoelectron spectroscopy and, for Fe-N-C, with ⁵⁷Fe Mössbauer spectroscopy.

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Experimental

Synthesis.- Catalyst precursors were prepared via a dry 153 ballmilling approach from a Zn(II) zeolitic imidazolate framework 154 (ZnN₄C₈H₁₂, Basolite Z1200 from BASF, labelled ZIF-8), Fe(II) ac-155 etate (Fe(II)Ac), Co(II)Ac, Cr(II)Ac and 1,10-phenanthroline. The 156 Fe-N-C and Co-N-C catalysts were synthesized identically: weighed 157 amounts of the dry powders of Fe(II)Ac (or Co(II)Ac), phenanthroline 158 and ZIF-8 were poured into a ZrO2 crucible. Then, 100 zirconium-159 oxide balls of 5 mm diameter were added and the crucible was sealed 160 under air and placed in a planetary ball-miller (FRITSCH Pulverisette 161 7 Premium) to undergo 4 cycles of 30 min of ballmilling at 400 rpm. 162 The resulting catalyst precursor was pyrolyzed at 1050°C in Ar for 163 h. The catalyst precursors contained 1 wt% of metal and the mass 1 164 ratio of phenanthroline to ZIF-8 was 20/80.9,21 The mass loss during 165 pyrolysis was 60-65 wt%, leading to Fe and Co contents of 2.5-2.9 166 wt%. The catalyst precursor for Cr-N-C was prepared identically, but 167 the pyrolysis was performed under flowing NH₃ at 950°C for 15 min.

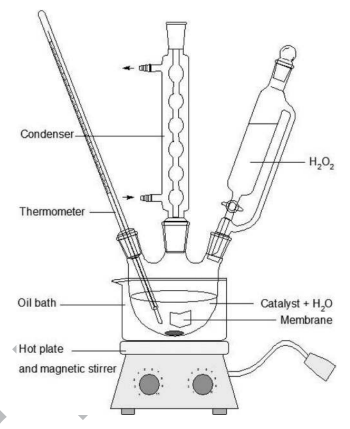


Figure 1. Scheme of the experimental setup used for ex situ H_2O_2 degradation.

When pyrolyzed in Ar, the initial ORR activity of Cr-N-C was too low to be of interest. The resulting Cr-based catalyst was then acid 170 washed in a pH 1 sulphuric acid solution and copiously washed with 171 de-ionized water. The acid washing was applied only to Cr-N-C in or-172 der to compare its ORR activity to the Ar-pyrolyzed Fe- and Co-N-C 173 catalysts. It is known that pyrolysis in NH₃ leads to a higher initial 174 ORR activity, mostly due to the presence of highly basic surface N-175 groups.¹⁶ This promotion of ORR activity due to the high basicity of 176 N-groups is usually suppressed after an acid wash, which protonates 177 basic N-groups followed by anion adsorption. The mass loss during 178 pyrolysis for the Cr-based catalyst precursor was 85–90 wt%, leading 179 to 8-10 wt% Cr in Cr-N-C. 180

Chemical treatment with hydrogen peroxide.— 100 mg of catalyst 181 was dispersed and stirred in (500 - X) ml of deionized water heated 182 to 80°C under reflux. A piece of Nafion 117 membrane with area 4.84 183 cm² was also immersed in solution in order to serve as an indicator 184 for ROS. The scalar X represent the total volume of 30 wt% H_2O_2 185 solution that was added to reach the desired number of mol H_2O_2 186 per mg of catalyst. This volume was systematically divided in four 187 aliquots of X/4 ml, each aliquot being added with a one hour time 188 interval. Once the total desired volume of H_2O_2 had been added, the solution was stirred for 24 h at 80°C under reflux and then allowed to 190 cool down to room temperature. The entire experimental setup can be 191 visualized in Fig. 1. For example, in order to reach a ratio of 10^{-2} mol 192 H₂O₂ per mg of catalyst, four aliquots of 25 ml each of 30 wt% H₂O₂ 193 solution were added every hour to a solution initially containing 100 194 mg of catalyst dispersed in 500-(4 \times 25) ml of deionized water. In 195 the present study, the ratio of mol H₂O₂ per mg catalyst was spanned 196 from 5 \times 10⁻⁴ to 2 \times 10⁻² by changing the cumulative volume of 30 197 wt% H₂O₂ from 5 to 200 ml. Each H₂O₂ treatment was terminated 198 with a one hour sonication of the solution to solubilize fluoride ions 199 originating from membrane degradation. The Nafion membrane piece 200

was removed and the solution filtrated using a Büchner setup with 201 a 220 nm pore-sized filter. The filtrate was stored for fluoride ion 202 analysis while the collected catalyst powder was copiously washed 203 with de-ionized water (ca one litre) before overnight drying at 90°C. 204 The collected mass of dry catalyst powder was then weighed while 205 the fluoride ion concentration in the filtrate was measured using an ion 206 selective electrode. To avoid any interference of the pH, the electrolyte 207 solution was adjusted with the appropriate ionic strength adjustment 208 buffer (TISAB IV from Sigma-Aldrich) in order to keep the pH in the 209 range of 5-7. TISAB solution also prevents complexation of fluoride 210 ions with any iron or other trace metals present in solution. 211

Structural characterization.- Metal K-edge X-ray absorption 212 spectra were recorded at room temperature at SAMBA beamline of 213 Synchrotron SOLEIL. The beamline is equipped with a sagittaly fo-214 cusing Si 220 monochromator and two Pd-coated collimating mirrors. 215 The mirrors have been used to remove X-rays harmonics. The catalysts 216 were pelletized as disks of 10 mm diameter using a 50/50 mass ratio 217 of Teflon powder (1 µm particle size) and catalyst. The spectra were 218 recorded in transmission mode and the energy was calibrated with 219 the corresponding metal foils to correct any change in beam energy 220 during the experiments. All experimental spectra were normalized 221 and analyzed identically with Athena software in order to derive the 222 XANES and Fourier-transform EXAFS spectra. X-ray diffraction was 223 224 conducted using a PANanalytical X'Pert Pro powder X-ray diffractometer. XPS spectra were recorded with a Physical Electronics PHI 225 5701 spectrometer. Non-monochromatic Mg K_α X-ray was used as 226 the excitation source. For the pristine and H2O2-treated Fe-N-C cata-227 lysts, ⁵⁷Fe Mössbauer spectra were measured with a source of ⁵⁷Co in 228 229 rhodium. The measurements were performed keeping both the source 230 and the absorber at room temperature. The spectrometer was operated with a triangular velocity waveform, and a NaI scintillation detector 231 was used for detecting the γ -rays. Velocity calibration was performed 232 233 with an α-Fe foil.

234 Electrochemical characterization. -- Electrochemical activity of 235 the catalysts toward the ORR was determined using the RDE technique. A catalyst ink including 10 mg of catalyst, 109 µl of a 236 5 wt% Nafion solution containing 15-20% water, 300 µl of ethanol 237 and 36 µl of de-ionized water was sonicated and mixed using a vortex. 238 An aliquot of 7 μ l was deposited on the glassy-carbon disk (0.196 cm²) 239 resulting in a catalyst loading of 800 $\mu g \cdot cm^{-2}.$ The working electrode 240 with the deposited catalyst layer was used in a three-electrode cell 241 setup connected to a potentiostat (Versastat 3, AMETEK) and rotator 242 (MSR, Pine Instruments). The counter electrode and reference elec-243 trode were a graphite rod and a reversible hydrogen electrode (RHE), 244 respectively. The electrolyte was an O₂-saturated aqueous solution of 245 0.1 MH₂SO₄. The RDE polarization curves were recorded with a scan 246 rate of $10 \text{ mV} \cdot \text{s}^{-1}$ at 1500 rpm and corrected for the background cur-247 rent measured in N2-saturated electrolyte. Additional measurements 248 were similarly performed with a rotating-ring disk electrode for de-249 termination of the % H₂O₂ released during ORR. The ring potential 250 was held at 1.2 V vs. RHE. Selected catalysts were investigated in 251 fuel cell. For the membrane electrode assembly (MEA), cathode inks 252 were prepared using the following formulation: 20 mg of catalyst, 253 652 μl of a 5 wt% Nafion solution containing 15–20% water, 326 254 µl of ethanol and 272 µl of de-ionized water. The inks were alterna-255 tively sonicated and agitated with a vortex mixer every 15 min, for a 256 total of 1 h. Then, three aliquots of 405 μ l of the catalyst ink were 257 successively deposited on the microporous layer of an uncatalysed 258 4.84 cm² gas diffusion layer (Sigracet S10-BC) to reach a catalyst 259 loading of 4 mg cm⁻². The cathode was then placed in a vacuum oven 260 at 90°C to dry for 1 h. The anode used for all PEMFC tests performed 261 in this work was 0.5 $mg_{Pt} \cdot cm^{-2}$ on Sigracet S10-BC. MEAs were 262 prepared by hot-pressing 4.84 cm² anode and cathode against either 263 side of a Nafion NRE-211 membrane at 125°C for 2 min. PEMFC 264 tests were performed with a single-cell fuel cell with serpentine flow 265 field (Fuel Cell Technologies Inc.) using an in-house fuel cell bench 266 267 and a Biologic Potentiostat with a 50 A load and EC-Lab software.

For the tests, the fuel cell temperature was 80° C, the humidifiers were set at 85° C, and the inlet pressures were set to 1 bar gauge for both anode and cathode sides. The flow rates for humidified H₂ and O₂ were ca 50–70 sccm downstream of the fuel cell. Polarization curves were recorded by scanning the cell voltage at 0.5 mV \cdot s⁻¹.

Results and Discussion

Structural characterisation of pristine catalysts.— The X-ray 274 diffractograms for the pristine Fe-N-C, Co-N-C and Cr-N-C cata-275 lysts are shown in Figure 2. While the Fe-N-C and Co-N-C catalysts 276 only show two diffraction peaks that are assigned to the 002 and 101 277 reflections of graphite, the Cr-N-C catalyst shows additional intense 278 peaks that are assigned to the CrN structure. In addition, secondary 279 peaks visible in the diffraction pattern of Cr-N-C are assigned to the 280 Cr₂N structure and to a chromium carbonitride structure. The major 281 presence of CrN over Cr₂N was expected after pyrolysis in the N-282 rich atmosphere, NH₃.⁴² The superimposition of the broad peaks at 283 23–25° that is assigned to graphite highlights a similar low degree of 284 graphitization. This is understandable since the metallic cobalt and 285 iron crystalline structures known to catalyze graphitisation are not 286 detected by XRD in the pristine Fe-N-C and Co-N-C catalysts while 287 chromium nitrides do not catalyze graphitization.^{29,43–44}

The Fourier-transforms of the extended X-ray absorption fine 289 structure (EXAFS) spectra of the Fe-N-C and Co-N-C catalysts are 290 shown in Fig. 3a–3b (thick solid lines), while the corresponding 291 X-ray absorption near edge structure (XANES) spectra are shown 292 in Fig. 3d–3e. The radial distribution function of Fe-N-C shows a first 293 peak at 1.3–1.5 Å arising from Fe-N interactions and a second peak 294 at 2.2 Å mostly assigned to backscattering from carbon atoms. Both 295 the EXAFS and XANES spectra for pristine Fe-N-C resemble those 296 of Fe(II) phthalocyanine (Fe^{II}Pc) (thin red line in Figs. 3a and 3d). 297 The main difference is the absence of a pre-edge peak at 7117 eV 298 in the XANES spectrum of Fe-N-C, while it is present in the spec-299 trum of Fe^{II}Pc. The absence of the pre-edge feature in Fe-N-C can 300 be explained on the basis that the square-planar D_{4h} local symmetry 301 is broken down in the Fe-N-C catalyst due to the additional coordi-302 nation of dioxygen along the axial direction.⁴⁵ The pre-edge feature 303 at 7117 eV is indeed observed for square-planar Fe^{II}N₄ compounds 304 but disappears for a square-pyramid coordination resulting from the 305 coordination by a fifth ligand.^{17,46} Such is the case for Hemin, a chloro-306 Fe(III)-porphyrin compound (green line in Fig. 3d). Thus, the XANES 307 and EXAFS data for Fe-N-C show that i) the active-site structure in 308 Fe-N-C is similar to that in Fe^{II}Pc and ii) no or almost no Fe-Fe 309 bonds exist in that catalyst. Similarly, the EXAFS data for Co-N-C 310 suggest that only few Co-Co bonds are present in the pristine cata-311 lyst due to the small signal at 2.3–2.5 and 4–5 Å corresponding to 312

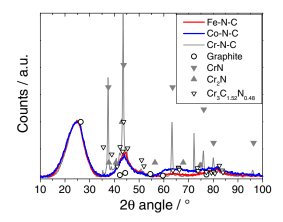


Figure 2. X-ray diffractograms of the pristine Fe-N-C, Co-N-C and Cr-N-C catalysts and reference X-ray diffraction lines for graphite (JCPDS file 75-1621), CrN (JCPDS file 11-0065), Cr_2N (JCPDS file 01-1232) and $Cr_3C_{1.52}N_{0.48}$ (JCPDS file 76-1037).

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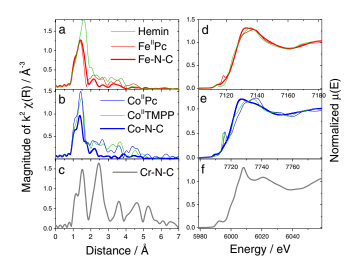


Figure 3. Left handside: Magnitude of the k^2 -weighed Fourier transform of the EXAFS signal for the pristine Fe-N-C, Co-N-C and Cr-N-C catalysts and for reference compounds. The radial distance is not corrected for phase shift. Right handside: the corresponding normalized XANES spectra, at energies between -10 and +70 eV around the corresponding Metal K-edge.

Co-Co bond distances in metallic cobalt (thick blue line in Fig. 3b). 313 On the basis of the relative intensity of the Fourier transform at 1.4 Å 314 (Co-N interactions) for the present Co-N-C catalyst compared to that 315 of a Co-based catalyst free of metallic cobalt particles (lower cobalt 316 content), it was estimated that ca 79-83% of the cobalt atoms in the 317 present Co-N-C catalyst are involved in CoNxCy moieties. Similar to 318 the case for iron, the pre-edge feature is observed for the square-planar 319 Co^{II}N₄ compounds Co^{II}Pc and Co(II)-tetramethoxyphenylporphyrin 320 (Co^{II}TMPP) but not observed for Co-N-C (Fig. 3e). The active site 321 structure in Co-N-C is also different from that in Co^{II}Pc or Co^{II}TMPP 322 as revealed by the lower intensity of its Fourier transform at 1.4 Å, 323 corresponding to Co-N interactions (Fig. 3b). This suggests a lower 324 coordination of cobalt in Co-N-C than in CoN4 macrocycles. In con-325 clusion, the restricted amount of metal-based crystalline structures in 326 the pristine Co- and Fe-N-C catalysts is important as it allows an eas-327 ier tracking with X-ray absorption spectroscopy (XAS) of the fate of 328 the CoN_xC_y and FeN_xC_y moieties after H₂O₂ treatment. This aspect 329 had already proved important for post mortem Mössbauer analysis of 330 Fe-N-C cathodes.²¹ Last, the Fourier transform function for Cr-N-C 331 shows a long-range ordering up to 6 Å, in agreement with the CrN 332 structure. The first peak at 1.5 Å is assigned to Cr-N interactions with 333 N atoms while the second and third peaks at 2.5 and 3.6 Å are assigned 334 to Cr-Cr interactions. 335

Electrochemical characterization of pristine and degraded 336 catalysts.- Figure 4 shows the polarization curves measured with 337 RDE for the pristine Fe-N-C, Co-N-C and Cr-N-C catalysts (thick 338 black curves) as well for the three series of catalysts obtained by 339 treating these catalysts with increasing amounts of H₂O₂. The po-340 larization curves for the pristine Fe- and Co-N-C catalysts show a 341 gradual \transition from a kinetic control to a diffusion control ex-342 tending over 150 mV. This is typical for catalytic films with poor 343 diffusion properties. These experimental curves are similar in shape 344 to those calculated for films including an O₂ concentration gradient 345 at high current density.47 This does however not impede the reading 346 of the ORR activity at high potential where the current density is 347 low. For both Fe- and Co-N-C, the ORR activity at 0.8 V vs. RHE 348 gradually decreases with increasing amounts of H₂O₂ used during the 349 treatment, while the diffusion-limited current density is practically 350 unmodified. This suggests that the ORR mechanism was unchanged 351 while the number of FeN_xC_y and CoN_xC_y active sites on the surface 352 gradually decreased. The general trend of decreasing ORR activity 353 354 with increased amount of H2O2 used in the treatment is however more

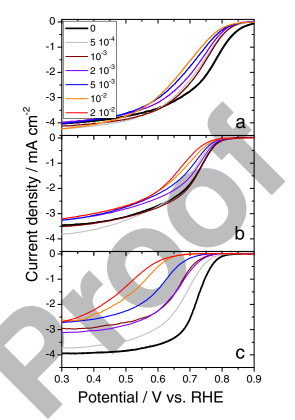


Figure 4. Polarization curves measured with RDE for a) Fe-N-C, b) Co-N-C and c) Cr-N-C pristine catalysts and after ex situ treatment with various amounts of H_2O_2 . The legend indicates the number of moles H_2O_2 per mg of catalyst used for treating Me-N-C catalysts.

complex at low peroxide amounts. For Fe-N-C, the ORR activities 355 after treatment with $5 \cdot 10^{-4}$ and 10^{-3} mol H₂O₂ per mg of catalyst 356 are identical, suggesting that the degradation stalled. For Co-N-C, the 357 ORR activity first decreased after treatment with $5 \cdot 10^{-4}$ mol H₂O₂ 358 per mg of catalyst, but rose back to the activity of pristine Co-N-C 359 after treatment with 10⁻³ mol H₂O₂ mg⁻¹. For higher amounts of 360 H₂O₂, the ORR activity then decreased gradually for both Fe- and 361 Co-N-C. The H_2O_2 treatment seems to result in the superimposition 362 of a minor positive effect at low H₂O₂ amount and of a major negative 363 effect at high H₂O₂ amount. Following a restricted surface oxidation 364 with H_2O_2 , the top surface of Co- or Fe-N-C might have been cleaned, thereby exposing to the electrolyte some active sites that were pre-366 viously buried under a thin carbon layer. Alternatively, the grafting 367 of oxygen groups on the surface might have positively affected the 368 ORR turnover of the active sites via electronic effects.⁴⁸ Next, the 369 pattern for Cr-N-C is different. The polarization curve of the pristine 370 catalyst is steeper in the kinetic region. The diffusion-limited current 371 density is also better defined. The decrease of the ORR activity after 372 H₂O₂ treatment is much larger than that observed for Fe- and Co-373 N-C. This suggests that the nature of the active sites is completely 374 different, which is corroborated by XRD, XANES and EXAFS char-375 acterisations (Figs. 2–3). As will be seen later, the chromium nitride 376 structures resist the peroxide treatment, but they are ORR inactive. 377 The ORR activity of pristine Cr-N-C is assigned to highly basic N-378 groups that are formed during pyrolysis in NH₃. The assignment is 379 based on the poor ORR activity of a Cr-based sample pyrolyzed in Ar 380 and also on the high activity of a metal-free N-C sample pyrolyzed in 381 NH₃ (not shown). Regarding the diffusion-limited current density, the 382 pattern is also very different from that observed for Fe- and Co-N-C. 383 The ORR mechanism seems to gradually shift from a four-electron 384 to a two-electron reduction mechanism (Fig. 4c). This is understand-385 able if the top-surface N-groups present in pristine Cr-N-C were re-386 moved from the surface during the treatment with H₂O₂, or chemically 387

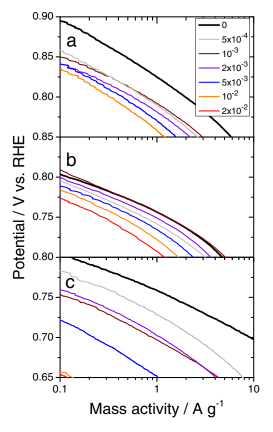


Figure 5. Tafel plots derived from RDE polarization curves for a) Fe-N-C, b) Co-N-C and c) Cr-N-C catalysts initially and after ex situ treatment with various amounts of H_2O_2 . The legend indicates the number of moles H_2O_2 per mg of catalyst used to treat Me-N-C catalysts.

modified. The low ORR activity remaining after extensive degrada-388 tion with H₂O₂ possibly comes from less active N-groups or from 389 oxygen groups that formed during the peroxide treatment. In order to 390 391 quantify the ORR activity, the polarization curves were corrected for diffusion limitation with the Koutecky-Levich equation and normal-392 ized per mass of catalyst. Figure 5 shows the Tafel plots corresponding 393 to Figure 4. Within a given catalyst series, the Tafel slopes remained 394 constant with H₂O₂ treatment, except for pristine Fe-N-C for which 395 the Tafel slope is slightly higher than that for treated catalysts (77 396 vs. 62–65 mV dec⁻¹). The ORR activity after treatment with 10^{-2} 397 mol H₂O₂ mg⁻¹ was divided by 6.6 for Fe-N-C, 2.9 for Co-N-C and 398 1400 for Cr-N-C (Fig. 5). The activity was read at 0.8 V vs. RHE 399 for Fe- and Co-N-C, and at 0.75 V vs. RHE for Cr-N-C. Thus, the 400 ORR activity decay following H₂O₂ treatment strongly depends on 401 the nature of the metal in Me-N-C catalysts. This in turn strongly sug-402 gests that the oxidizing species degrading the catalysts is not H₂O₂ 403 but rather the ROS generated via a Fenton reaction between transition 404 metal atoms and H₂O₂. Direct evidence for the formation of ROS is 405 reported in Collection yield and radical oxygen species after ex situ 406 H_2O_2 treatment section. 407

The ORR mechanism on pristine catalysts was then investigated 408 with a rotating ring disk electrode (RRDE). Measured at a loading 409 of 0.8 mg cm⁻², the % H₂O₂ detected at the ring is shown in Fig. 6a 410 and the polarization curves measured at the disk are shown in Fig. 6b. 411 The % H₂O₂ is far lower for ORR when catalyzed by Fe-N-C than 412 catalyzed by Co-N-C and Cr-N-C. The curves of % H₂O₂ vs. potential 413 for Fe-N-C and Co-N-C have a similar shape, but Co-N-C produces ca 414 20 times more H₂O₂ at any given potential. For Cr-N-C, the % H₂O₂ is 415 quite high and almost independent of potential. In order to investigate 416 whether the low % H2O2 measured on Fe-N-C indicates a true four 417 418 electron reduction mechanism, a 2 + 2 electron mechanism or a

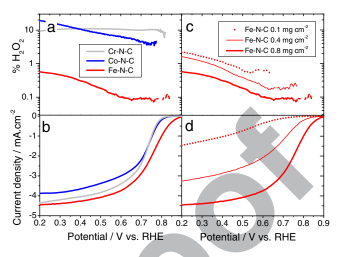


Figure 6. Rotating ring disk electrode measurements for Fe-N-C, Co-N-C and Cr-N-C (left handside) at a fixed loading of 0.8 mg cm⁻² and rotating ring disk electrode measurements for Fe-N-C at different loading (right handside). In c and d, the loading at the disk was 0.8 mg cm⁻² (thick solid line), 0.4 mg cm⁻² (thin solid line) or 0.1 mg cm⁻² (dashed line).

two-electron mechanism followed by chemical disproportionation, 419 lower Fe-N-C loadings were investigated (Fig. 6c-6d).⁴⁹ At lower 420 loadings, the diffusion-limited current density was drastically reduced 421 (Fig. 6d), due to incomplete coverage of the disk, or due to a low 422 density of active sites per geometric area of the disk. This drastic de-423 crease does not reflect the small change in % H2O2 that is detected at 424 the ring. The % H₂O₂ increased with decreased loading, but remained 425 in the range 3.3-4.5% at potentials of 0.2-0.6 V vs. RHE even at 426 0.1 mg_{Fe-N-C} cm⁻² (dashed curve in Fig. 6d). This shows that ORR 427 on Fe-N-C proceeds mostly according to a direct four electron 428 mechanism. The minute amount of H₂O₂ formed on the catalyst 429 surface may however have formed ROS through a chemical reaction 430 on FeN_xC_y moieties. 431

While the RDE and RRDE techniques are useful for investigating 432 the ORR activity and ORR mechanism, they do not allow predicting 433 the behavior of Me-N-C cathodes at high current density in PEMFC. 434 We selected the pristine catalysts and catalysts treated with $2 \cdot 10^{-3}$ 435 and 10^{-2} mol H_2O_2 mg⁻¹ for beginning-of-life (BoL) PEMFC tests. Figure 7 shows PEMFC polarization curves recorded with 4 mg cm⁻² 436 437 of Me-N-C catalyst at the cathode. Analyzing the extent of degrada-438 tion after ex situ H_2O_2 treatment should a priori be carried out by 439 comparing the BoL polarisation curve of H₂O₂-treated catalysts to 440 the BoL polarization curves of the corresponding pristine catalysts. 441 However, for the pristine Fe- and Co-N-C catalysts, this simple ap-442 proach is impeded by the fact that the BoL polarization curve does 443 not correspond to the best performance of the pristine catalysts. For 444 pristine Fe-N-C, the polarization curve continuously improves during 445 the first 6 h of operation, and the same occurs for pristine Co-N-C 446 during the first 15 h of operation. The improvement is most visible 447 in the high current density region, while the ORR activity at high 448 potential slightly decreases during this break-in period (Fig. 7a–7b). 449 The break-in phenomenon is assigned to changes in the hydrophilic 450 properties of the catalytic surface that had never been in contact with 451 water previously. This break-in period is neither observed for pristine 452 Cr-N-C nor for any H₂O₂-treated catalyst. When the high current den-453 sity region is considered, the reference curves for the pristine Fe-N-C 454 and Co-N-C catalysts will therefore be the polarisation curves after 455 break-in (dotted black lines in Fig. 7a–7b). When the ORR activity 456 at 0.8 V is considered, the reference curves for the pristine Fe-N-457 C and Co-N-C catalysts will be the BoL polarization curves (solid 458 black lines in Fig. 7a-7b). The current densities measured at 0.8 and 459 0.5 V in fuel cell as extracted from Fig. 7 for pristine and H₂O₂-treated 460 catalysts are summarized in Figure 8. 461

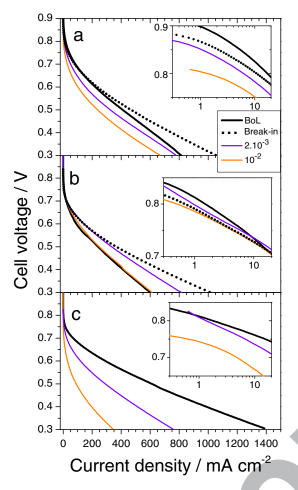


Figure 7. Effect of ex situ H_2O_2 treatment on the fuel cell polarization curves of MEAs with cathodes comprising a) Fe-N-C, b) Co-N-C and c) Cr-N-C catalysts. The insets show the high-potential region of the polarization curves in semi-logarithmic plot. The break-in involved potentiostatic control at 0.5 V for 6 h (pristine Fe-N-C) or 15 h (pristine Co-N-C).

As previously observed with RDE, the degradation of the cathode 462 performance after treating the catalysts with 10^{-2} mol H₂O₂ mg⁻¹ is 463 most severe for Cr-N-C, followed by Fe-N-C and then Co-N-C. After 464 a treatment with 10^{-2} mol H₂O₂ mg⁻¹, the ORR activity measured in 465 PEMFC at 0.8 V was divided by ca 45 for Cr-N-C, by 9 for Fe-N-C 466 and by 3 for Co-N-C (Fig. 8b). These numbers are comparable to those 467 measured with RDE, except for Cr-N-C for which the decay in fuel 468 cell is much smaller than the one measured in RDE (Fig. 5c and 7c). At 469 0.5 V, the decrease in current density from the BoL polarization curve 470 (Cr-N-C), or from the polarisation curve after break-in (for Co- and 471 Fe-N-C) to the BoL polarisation curve measured for catalysts treated 472 with 10^{-2} mol H₂O₂ mg⁻¹ is highest for Cr-N-C (decrease from 606) 473 to 85 mA cm⁻²), followed by Fe-N-C (470 to 238 mA cm⁻²) and 474 smallest for Co-N-C (366 to 240 mA cm^{-2}). Noteworthy, the ORR 475 activity at 0.8 V is almost identical for the pristine Co-N-C catalyst 476 after break-in and for the Co-N-C catalyst after treatment with 10^{-2} 477 mol H_2O_2 mg⁻¹ (Fig. 8b). In spite of this, a large difference is observed 478 in the current density at 0.5 V (Fig. 8a). Thus, for Co-N-C, the loss of 479 performance at high current density induced by the peroxide treatment 480 is mostly due to decreased transport properties of the cathode layer. 481 The story is different for Fe-N-C and Cr-N-C. The Fe-N-C catalyst 482 shows a stronger decay of its ORR activity after peroxide treatment 483 (Fig. 8b) but the transport properties of the cathode layer are also 484 negatively impacted by the ex situ peroxide treatment of the catalyst, 485 as can be deduced from the steeper slope of the polarization curve 486 487 at high current density (Fig. 7a). The Cr-N-C based cathode shows a

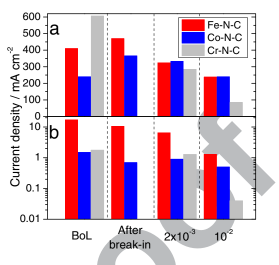


Figure 8. Current density measured in PEMFC at 0.5 V (a) and 0.8 V (b) for the pristine catalysts at BoL or after break-in, and for the catalysts after treatment with $2 \cdot 10^{-3}$ or 10^{-2} mol H₂O₂ mg⁻¹. The break-in involved potentiostatic control at 0.5 V for 6 h (pristine Fe-N-C) or 15 h (pristine Co-N-C). No break-in was applied to Cr-N-C. The activity of Cr-N-C at 0.8 V after 10^{-2} mol H₂O₂ mg⁻¹ needed an extrapolation.

dramatic decrease of both its ORR activity and transport properties $_{488}$ due to the H_2O_2 ex situ treatment of the Cr-N-C catalyst (Fig. 8, gray columns). $_{490}$

Collection yield and radical oxygen species after ex situ H_2O_2 491 treatment.— The weight percentage of catalyst powder collected after each H_2O_2 treatment is reported in Fig. 9a. After a treatment with $5 \cdot 10^{-4}$ to $5 \cdot 10^{-3}$ mol H_2O_2 mg⁻¹, the collection yields are 87–90% for Fe-N-C (red bars) and 77–85% for Cr-N-C (gray bars). Collection yields lower than 100% can be due to i) handling loss, ii) chemical loss due to the oxidative attack by peroxide and ii) filtration loss. The latter occurs if, at the end of the treatment, some catalytic particles are smaller than the pore size of the filter used to separate the treated 499

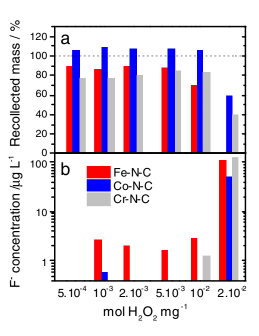


Figure 9. Extent of the H_2O_2 ex situ degradation assessed a) with the relative mass of catalyst recollected after each H_2O_2 treatment + filtration + drying, or b) with the concentration in the filtrate of fluoride ions released by a Nafion membrane.

catalyst from the solution. The non-collected mass of catalyst for the 500 treated Fe-N-C and Cr-N-C samples is mostly assigned to chemical 501 and filtration losses. The loss due to material's handling is believed to 502 be negligible since for Co-N-C samples handled similarly, a weight 503 gain was observed after treatment up to 10^{-2} mol H₂O₂ mg⁻¹ (blue 504 bars in Fig. 9a). This weight gain is assigned to the grafting of oxygen 505 functionalities on the surface of Co-N-C, without any loss of carbon 506 atoms from Co-N-C. This explanation is supported by XPS elemen-507 tal analysis (III.6). The lower collection yields for Cr-N-C samples 508 than for Fe-N-C samples is assigned to the higher porosity and higher 509 specific area of the pristine Cr-N-C catalyst due to the ammonia gas 510 used during pyrolysis. A significant decrease of the collection yield 511 is observed at and above 10^{-2} mol H₂O₂ mg⁻¹. After treatment with 512 $2 \cdot 10^{-2}$ mol H₂O₂ mg⁻¹, only 60 and 40% of the initial Co-N-C 513 and Cr-N-C mass was collected whereas no Fe-N-C catalyst could 514 be collected on the filter. A large fraction of the missing mass of 515 catalyst is suspected to have passed through the 220 nm sized pores 516 of the filter as a consequence of finer powders following the H₂O₂ 517 treatment. This hypothesis is supported by the greyish color of the 518 filtrates after treatment with large amounts of peroxide. The lower 519 granulometry of the powders after extended treatment with perox-520 ide may explain the poorer transport properties of fuel cell cathodes 521 (Fig. 7). The cathode macroporosity may have decreased due to a 522 lower 3D branching character and lower inter-particle connections for 523 524 H_2O_2 -treated catalysts. In conclusion, after treatment with 10^{-2} mol H_2O_2 mg⁻¹, the extent of catalyst modification clearly depends on the 525 nature of the transition metal in Me-N-C catalysts, in the order Fe > Cr 526 > Co. This, in turn, might be related to the production of ROS ensuing 527 from the reaction between H_2O_2 and metal species. The production 528 529 of ROS during the peroxide treatment was quantified with the con-530 centration of fluoride ions measured in the filtrate at the end of each treatment. These fluoride ions originate from the simultaneous attack 531 by ROS of the Nafion membrane immersed in the solution during the 532 entire process. While the fluoride concentration remained low after 533 treatment up to 5.10^{-3} mol H₂O₂ mg⁻¹, it dramatically increased by 534 ca two orders of magnitude for all three catalysts after a treatment with 535 $2 \cdot 10^{-2}$ mol H₂O₂ mg⁻¹ (Fig. 9b). A correlation is observed between 536 a high collection yield and a low concentration of fluoride, suggesting 537 that Co-N-C produces less ROS than Fe-N-C when it is contacted 538 by H₂O₂. This explains the restricted decrease of the electrochemical 539 performance of Co-N-C after H₂O₂ treatment. 540

X-ray absorption spectroscopy on pristine and degraded 541 catalysts.— The XANES spectra for the three series of H₂O₂ treated 542 catalysts are shown in Fig. 10. For Fe-N-C, there is practically no 543 change of the XANES spectra from the pristine Fe-N-C catalyst up to 544 the highly degraded catalyst after treatment with 10^{-2} mol H₂O₂ mg⁻¹ 545 (Fig. 10a). Also the Fourier-transforms of the EXAFS spectra of the 546 Fe-N-C samples did not reveal any changes and were identical to that 547 of pristine Fe-N-C shown in Fig. 3a. The XANES and EXAFS spec-548 tra of the Fe-based series therefore demonstrate that the coordination 549 chemistry of the Fe atoms that remained in the samples after H_2O_2 550 treatment is identical to the one before treatment. It is paramount to 551 realize that this observation does not preclude the possibility that the 552 FeN_xC_y moieties located on the top-surface in pristine Fe-N-C were 553 partially or fully removed during H_2O_2 treatment. If similar FeN_xC_y 554 moieties are present both on the top surface and in the bulk of pristine 555 Fe-N-C, the normalized XANES and EXAFS spectra will remain un-556 changed even after the complete removal of top surface sites due to 557 the bulk-averaging nature of XANES and EXAFS. Hence, identical 558 spectra before and after H2O2 treatment is not paradoxical with the 559 decreased ORR activity before and after H₂O₂ treatment. It strongly 560 suggests, however, that only a fraction of the FeN_xC_y moieties existing 561 in pristine Fe-N-C were located on the top surface. Only the latter may 562 contribute to the ORR activity. For the series of H2O2-treated samples 563 derived from pristine Co-N-C, the XANES spectra changed notice-564 ably with H_2O_2 treatment (Fig. 10b). The inset in Fig. 10b shows 565 the initial increase of the white line intensity at ca 7726 eV after 566 treatment with $5 \cdot 10^{-4}$ and 10^{-3} mol H₂O₂ mg⁻¹, followed by a con-567

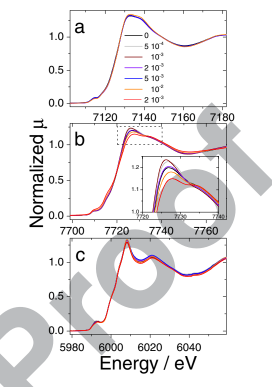


Figure 10. Normalized XANES spectra for pristine and H_2O_2 treated catalysts. a) Fe-N-C, b) Co-N-C, c) Cr-N-C. The spectra are shown in the energy range -10 to +70 eV around the corresponding Metal K-edge.

tinuous decrease from 10^{-3} to $2 \cdot 10^{-2}$ mol H₂O₂ mg⁻¹. Noteworthy, 568 after treatment with 10^{-3} mol H₂O₂ mg⁻¹, the ORR activity measured 569 in RDE increased back to the ORR activity of pristine Co-N-C, and 570 these two observations might be interrelated. The two XANES spectra 571 corresponding to treatments with 10^{-2} and $2 \cdot 10^{-2}$ mol H₂O₂ mg⁻¹ are 572 significantly different at 7726 eV and are also negatively shifted by 573 1.5-2.5 eV at the beginning of the absorption edge (7715 eV). This in-574 dicates a decreased average oxidation state of cobalt. This observation 575 is further supported by the Fourier-transforms of the EXAFS signal, 576 showing an increased peak at ca 2.2, 3.8 and 4.5 Å (Fig. 11). These 577 distances (not corrected for phase shift) correspond to back-scattering 578 from cobalt atoms in metallic cobalt. Extensive treatment with H₂O₂ 579 probably resulted in the etching of a significant fraction of the CoN_xC_y 580 moieties present in pristine Co-N-C, thereby increasing the fraction of 581

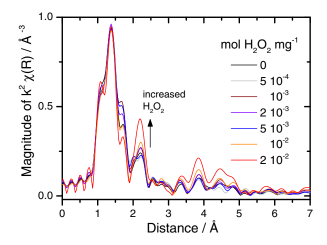


Figure 11. Magnitude of the k^2 -weighed Fourier transform of the EXAFS signal for pristine Co-N-C and for H_2O_2 treated Co-N-C. The radial distance is not corrected for phase shift.

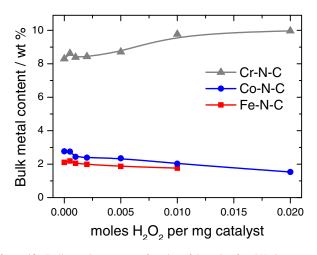


Figure 12. Bulk metal content as a function of the ratio of mol H_2O_2 per mass catalyst used for ex situ degradation.

metallic cobalt in the treated Co-N-C samples. Pristine Co-N-C seems 582 to contain a small fraction of metallic cobalt (Fig. 11), although it was 583 not detected by XRD (Fig. 2). The better resistance of metallic cobalt 584 particles to the peroxide treatment is normal since such particles are 585 usually surrounded by a continuous layer of graphitic carbon after 586 pyrolysis. Next, the XANES spectra for the pristine and treated Cr-N-587 C catalysts show little change with H_2O_2 treatments (Fig. 10c). This 588 agrees with the known stability of chromium nitride structures. 589

In addition to providing knowledge on the oxidation state and co-590 ordination chemistry of transition metals, XAS allows quantifying the 591 bulk metal content from the absolute height of the absorption step 592 combined with the exact mass of each pellet used for measurements 593 in transmission mode. Figure 12 shows the bulk content of Fe, Co and 594 595 Cr determined from XAS as a function of the number of mol H_2O_2 596 used to treat the catalysts. The bulk contents of Fe and Co decreased with increasing amounts of H2O2. However, the loss is moderate. Af-597 ter a treatment with 10^{-2} mol H₂O₂ mg⁻¹, the relative decrease of 598 Fe and Co content was only 17 and 27%, respectively. For Cr-N-C, 599 the bulk metal content even increased by 18%, highlighting the better 600 resistance to the peroxide treatment of CrN particles than the N-doped 601 carbon matrix. Dodelet's group also reported in 2003 a relative loss 602 of Fe of only 30-40% after H₂O₂ treatment. In parallel, the half-wave 603 potential for ORR in RDE decreased by 150-200 mV, corresponding 604 to ca two-order-of-magnitude decrease in ORR activity.²⁵ For another 605 Fe-N-C catalyst, Zelenay's group observed a decrease of Fe content 606 from 0.9 to 0.6 at. % (33% relative los) after H₂O₂ treatment for 607 72 h, which was accompanied by a 150 mV downshift of the half-608 wave potential for ORR when measured in RDE.²⁶ A XANES analysis 609 with linear combination fitting of reference spectra identified that only 610 20-25% of the iron atoms in their pristine catalyst were involved in a 611 Fe^{II}Pc-like coordination, the remainder being iron sulfide, iron oxide 612 and metallic iron structures. This complicated the identification of a 613 correlation, or lack of correlation, between the decrease in bulk con-614 tent of FeN_xC_y moieties during H₂O₂ treatment and the decrease of 615 ORR activity. Here in contrast, ca 80 and 90% of the metal atoms in the 616 pristine Co-N-C and Fe-N-C catalysts are assigned to MeN_xC_y moi-617 eties, respectively (see Structural characterisation of pristine catalysts 618 section and ⁵⁷Fe Mössbauer spectroscopy on pristine and degraded 619 Fe-N-C catalysts section). The relative loss of only 27 and 17% of 620 Co and Fe compared to the ORR activity loss measured in RDE after 621 treatment with 10^{-2} mol H₂O₂ mg⁻¹ (activity divided by a factor a 622 6.6 for Fe-N-C and 2.9 for Co-N-C) suggests that only a fraction of 623 the Fe and Co atoms in the pristine catalysts are located on the top 624 surface and participate in the ORR. Alternatively, if a major fraction 625 of MeN_xC_y moieties is located on the top surface initially, the results 626 suggest that surface oxidation due to H2O2 drastically reduced the 627 628 turnover frequency for ORR of the remaining MeN_xC_y moieties. In

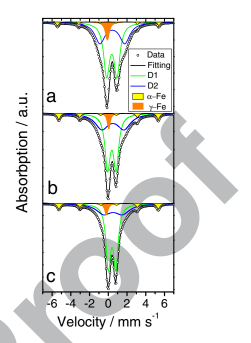


Figure 13. Mössbauer absorption spectra measured for pristine Fe-N-C and after ex situ treatment with H_2O_2 . a) pristine, b) after $2 \cdot 10^{-3}$ mol H_2O_2 mg⁻¹, c) after 10^{-2} mol H_2O_2 mg⁻¹.

this second hypothesis, the fact that little or no modification of the 629 XANES spectra is observed after H₂O₂ treatment, especially for Fe-630 N-C, suggests that the oxidation would have mostly occurred on the 631 N-doped carbon matrix. Preferential oxidation at the edge of graphene 632 sheets could have modified the electronic properties. The latter might in turn have modified the access of electrons to the MeN_xC_y moieties. 634 The latter are believed to be covalently integrated near the edge, or at 635 the edge, of graphene sheets.^{48,50–55} Decreased electronic properties 636 and conductivity at the edge would have profound implications in 637 particular for MeN_xC_y moieties located in micropores.^{56,5} 638

⁵⁷Fe Mössbauer spectroscopy on pristine and degraded Fe-N-C 639 catalysts.- Mössbauer spectroscopy is a powerful technique to iden-640 tify the presence of different iron coordinations in multiphase mate-641 rials. It is therefore ideally suited to be used alone or in combina-642 tion with XAS.^{58–60} With EXAFS, the secondary peak observed at ca 2.1–2.3 Å in the Fourier-transform may be assigned to backscatter-644 ing from either carbon atoms in the second coordination sphere of 645 FeN_xC_y moieties, or from iron atoms in the first or second coordina-646 tion sphere of metallic iron or iron carbide (Fig. 3a). The detection 647 of a minor fraction of iron atoms involved in Fe-based crystalline 648 structures is therefore difficult with EXAFS. Figure 13 shows the ⁵⁷Fe 649 Mössbauer spectra for pristine Fe-N-C and for samples obtained after 650 its treatment with $2 \cdot 10^{-3}$ and 10^{-2} mol H₂O₂ mg⁻¹. The Mössbauer 651 spectrum of pristine Fe-N-C shows that the major components are the 652 doublets D1 and D2 (Fig. 13a). They have previously been assigned 653 to a square-planar $Fe^{II}N_4$ coordination with Fe^{II} in low and medium 654 spin, respectively.⁵⁸⁻⁶¹ Doublets D1 has Mössbauer parameters sim-655 ilar to those of unpyrolyzed Fe^{II}Pc adsorbed on carbon, while D2 has parameters similar to those of crystalline Fe^{II}Pc in which stacked 657 molecules interact with each other (Table I).⁶² A sextet and a singlet 658 component are also detected and unambiguously assigned to α -Fe and 659 γ -Fe, respectively (Fig. 13a).⁶³ The spectral areas of these two com-660 ponents represent only 10% (Table I), demonstrating that the major 661 fraction of iron atoms in pristine Fe-N-C is engaged in molecular-like 662 FeN_xC_y moieties. Figures 13b and 13c show the spectra after perox-663 ide treatment with $2 \cdot 10^{-3}$ and 10^{-2} mol H₂O₂ mg⁻¹, respectively. No 664 major changes are observed. With increasing amount of H₂O₂, a trend 665 of increased relative fraction of D1 and decreased relative fraction 666 of D2 is noticed. In parallel, the relative fraction summed for α -Fe

Table I. Mössbauer parameters derived from the fitting with four components of the spectrum for pristine Fe-N-C and for Fe-N-C after ex situ treatment with $2 \cdot 10^{-3}$ and 10^{-2} mol H₂O₂ mg⁻¹. The hyperfine field of the sextet was 33.6, 33.0 and 32.9 T for pristine Fe-N-C, $2 \cdot 10^{-3}$ and 10^{-2} treated Fe-N-C samples, respectively.

	Relative Area/%			Iso	omer Shift/mm	s^{-1}	Quadrupole Splitting/mm s ⁻¹			
	Pristine	$2 \cdot 10^{-3}$	10 ⁻²	Pristine	$2 \cdot 10^{-3}$	10^{-2}	Pristine	$2 \cdot 10^{-3}$	10^{-2}	Assignment
D1	53 ± 5	51 ± 2	59 ± 3	0.38 ± 0.01	0.38 ± 0.00	0.40 ± 0.00	1.10 ± 0.02	0.90 ± 0.01	0.80 ± 0.01	Fe ^{II} N ₄ /C, LS
D2	38 ± 3	37 ± 2	29 ± 2	0.43 ± 0.02	0.49 ± 0.01	0.51 ± 0.02	2.65 ± 0.08	2.36 ± 0.06	2.01 ± 0.13	Fe ^{II} N ₄ /C, MS
Sextet	3 ± 1	7 ± 0	7 ± 0	-0.02 ± 0.03	0.03 ± 0.01	0.04 ± 0.01	_	-	-	α-Fe
Singlet	6 ± 1	4 ± 0	4 ± 0	-0.10 fixed	0.05 ± 0.01	-0.20 ± 0.01	-	-	-	γ-Fe

and γ -Fe increases only from 9 to 11%. Such metallic particles are 668 embedded in a graphitic shell and resist oxidative attacks, as previ-669 ously shown in a high-voltage degradation study of the same catalyst.²¹ 670 If a large amount of FeN_xC_y moieties had been removed from the cata-671 lyst during H_2O_2 treatment, the relative fraction of α -Fe and γ -Fe after 672 treatment would have dramatically increased. The fact that it only in-673 creased from 9 to 11% (+22% relative increase) confirms that most of 674 the FeN_xC_y moieties existing in pristine Fe-N-C are still present after 675 treatment with 10^{-2} mol H₂O₂ mg⁻¹. This agrees with the moderate 676 decrease in bulk Fe content, only 17% relative decrease after treat-677 ment with 10^{-2} mol H₂O₂ mg⁻¹ (Fig. 12). Due to the trends of relative 678 area of D1 and D2, it can be concluded that the peroxide treatment 679 preferentially removed FeNxCy moieties assigned to D2. The relative 680 loss of 17% Fe after treatment with 10⁻² mol H₂O₂ mg⁻¹ can however 681 not entirely be assigned to the removal of D2 sites. If this were true, 682 683 the decrease in the relative area of D2 would have been commensurate with that number. However, it only decreased from 38 to 29% 684 (Table I). Hence, the H₂O₂ treatment etched a minor fraction of 685 both the D2 and D1 sites that were initially present in pristine Fe-686 N-C. These fractions can be estimated mathematically. If one con-687 siders 100 Fe atoms in pristine Fe-N-C, then 53 Fe atoms are in D1 688 sites, 38 atoms in D2 sites and the remainder in metallic structures 689 (Table I). The H_2O_2 treatment with 10^{-2} mol H_2O_2 mg⁻¹ removed 17 690 out of those 100 atoms (17% loss). After trial-and-error calculations. 691 it is found that when, among those 17 Fe atoms, 4 come from D1 692 sites, 13 from D2 sites and none from the metallic structures, then 693 the final relative % of D1 and D2 in the H_2O_2 treated sample are 694 $100 \times (53-4)/(100-17) = 59\%$, and $100 \times (38-13)/(100-17) = 30\%$. 695 These percentages match those experimentally observed for D1 and 696 D2 after treatment with 10^{-2} mol H₂O₂ mg⁻¹ (Table I, column 4). The 697 decrease in the absolute content of Fe atoms involved in D1 and D2 698 coordinations upon treatment with 10^{-2} mol H₂O₂ mg⁻¹ can now be 699 calculated. It is ca $100 \times 4/53 = 7.5\%$ for D1, and $100 \times 13/38 = 34\%$ 700 for D2. This information is more precise than the overall loss of 17% 701 Fe obtained from XAS. Nevertheless, no proportional relationship can 702 be drawn with the decrease in ORR activity for Fe-N-C. After treat-703 ment with 10^{-2} mol H₂O₂ mg⁻¹, the latter was divided by 6.6 and by 9 in RDE and PEMFC measurements, respectively (Electrochemical 704 705 characterization of pristine and degraded catalysts section). 706

In conclusion, Mössbauer spectroscopy reveals that most of the 707 FeN_xC_y moieties present in pristine Fe-N-C survived the peroxide 708 treatment. The Mössbauer parameters of D1 and D2 were only slightly 709 modified, especially for D1. This suggests that the active sites remain-710 ing in the peroxide-treated catalysts have an electronic structure and 711 coordination chemistry very similar to those in the pristine catalyst. 712 The apparent continuous decrease of the QS values for D1 and D2 713 with increased mol H₂O₂ mg⁻¹ used for ex situ degradation might 714 be due to electronic effects of the surrounding carbon support, which 715 becomes more oxidized and contains fewer delocalized π electrons. 716 Hence, either the ORR activity in pristine Fe-N-C is due to a mi-717 nor fraction of D1 or D2 sites located on the top surface, or the 718 top surface of the catalyst was highly oxidized during H₂O₂ treat-719 ment, leading to a loss of surface conductivity or partial exfoliation 720 of the graphene sheets. The latter transformations may be localized 721 around the surface FeN_xC_y moieties, where H_2O_2 reacted to form 722 ROS. 723

X-ray photoelectron spectroscopy on pristine and degraded 724 catalysts.— The nitrogen and oxygen content and speciation were in-725 vestigated with XPS on pristine catalysts as well as on catalysts after 726 treatment with 10^{-2} mol H₂O₂ mg⁻¹. Figure 14 shows as an exam-727 ple the N_{1s} narrow scan spectra for the pristine and degraded Fe-N-C 728 catalyst. Both experimental spectra were fitted with four components 729 (Table II). They are assigned in the order of increasing binding energy 730 (BE) to pyridinic, pyrrolic, graphitic and oxidized nitrogen.^{64–67} The 731 pyridinic component (BE 398.5-398.7), primarily assigned to N atoms 732 contributing with one electron to the π system, may also comprise N 733 atoms ligating the Fe or Co ions.^{66–67} Next, most pyrrolic N atoms are 734 expected to be located on the graphene edges, and bonded to two C 735 atoms and 1 H atom in six-membered rings. Third, the graphitic N 736 component comprises all N atoms bonded with three C-atoms, either 737 within a graphene sheet or at the edge of a graphene sheet.^{64,68} 738

From Fig. 14, it can be seen that no novel N-species arose during 739 the treatment of Fe-N-C with H₂O₂. This observation holds for Co-740 N-C and Cr-N-C. The modified shape of the overall N_{1s} spectrum in 741 Fe-N-C after degradation is due to a larger relative fraction of pyrrolic-742 N. After quantification of the atomic % of C, N and O, it is found that 743 the absolute content of pyrrolic N increased from 1.07 to 1.97 at % 744 during degradation, (Table II). This absolute increase in pyrrolic N 745 is also observed with Co-N-C and Cr-N-C (Table II). In contrast, the 746 absolute content of all other N species in Fe-N-C, Co-N-C or Cr-N-C 747 was unmodified or increased only slightly after H₂O₂ treatment. The 748 exception is the significant decrease of pyridinic N for Cr-N-C after 749 H_2O_2 degradation (ca -1 at %). The changes observed in the nitrogen 750

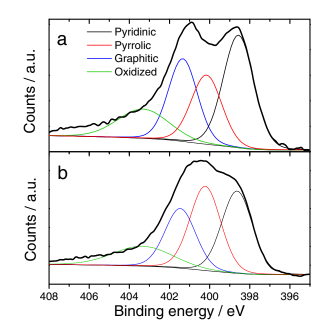


Figure 14. N_{1s} narrow scan spectra measured with XPS for a) the pristine Fe-N-C catalyst and b) the Fe-N-C catalyst treated with 10^{-2} mol H₂O₂ mg⁻¹. The experimental spectra were fitted with four nitrogen components.

Table II. X-ray photoelectron spectroscopy analysis of the N_{1s} and O_{1s} narrow scans for pristine catalysts and catalysts degraded with 10^{-2} mol H_2O_2 mg⁻¹. For Cr-N-C, a fifth N-component at 396.5–396.7 eV assigned to CrN was present with 0.15 and 0.33 atom % in the pristine and degraded catalysts, respectively.

		Oxygen								
	Pyridinic or MeN _x	Pyrrolic	Graphitc	Oxidized	Total	01	02	O3	O4	Total
BE range/eV	398.5-398.7	400.0-400.4	401.3-401.5	403.3-403.7		530.8-531.2	532.5-532.8	534.2-534.3	536.2-536.7	
Fe-N-C pristine	1.75	1.07	1.21	0.76	4.8	1.14	2.64	0.70	0.12	4.6
Fe-N-C degraded	1.93	1.97	1.40	0.90	6.2	4.20	6.55	1.50	0.55	12.8
Co-N-C pristine	1.76	0.84	1.55	0.85	5.0	1.81	2.79	0.47	0.13	5.2
Co-N-C degraded	2.20	1.46	1.54	0.90	6.1	2.79	5.40	1.00	0.32	9.5
Cr-N-C pristine	3.08	1.25	1.43	0.79	6.7	3.54	2.79	0.48	0.39	7.2
Cr-N-C degraded	2.02	2.10	1.28	0.57	6.3	5.11	6.22	1.23	0.93	13.5

speciation for Fe-N-C and Co-N-C are now discussed in parallel, due 751 to their similar active site structure. As a result of increased pyrrolic 752 N and unmodified content of other N-species, the absolute N content 753 increased by 1.1-1.4 at % after the peroxide treatment of Fe-N-C and 754 Co-N-C (Table II). This is counter-intuitive. The least stable carbon 755 atoms must have been substituted by nitrogen atoms from N₂ in air, a 756 substitution that might have been catalyzed by ROS. Increased content 757 of nitrogen was previously observed by Zelenay's group after ex situ 758 H₂O₂ treatment of another Fe-N-C catalyst.²⁶ In that study, after a 759 sharp drop from 6 to 3 at % N after 24 h of immersion in a 10% H₂O₂ 760 761 solution (unmodified ORR activity), the N-content steadily increased from 3.0 to 3.5 at % after 24-100 h of immersion (negative shift of 762 200 mV for the ORR half-wave potential measured with RDE). In 763 that study, the H_2O_2 treatment corresponds to $1.5-10^{-2}$ mol H_2O_2 764 mg⁻¹ and was also performed at 80°C.^{26,69} The N-speciation after ex 765 situ H₂O₂ treatment was not discussed, but after fuel cell testing for 766 500 h a relative increase in pyrrolic N was also observed on the N_{1s} 767 spectrum.26 768

Changes in the N content and speciation of the Cr-N-C catalysts 769 are now discussed. For Cr-N-C, a fifth N component was necessary 770 for fitting the N_{1s} spectrum. Its BE of 396.5–396.7 eV corresponds 771 to CrN.⁷⁰ The small N content assigned to CrN is explained on the 772 basis of the smaller penetration depth of X-rays in CrN (Table II). 773 Thus, the N_{1s} spectrum of Cr-N-C is practically unmodified by the 774 presence of CrN and largely reflects N-moieties integrated in the 775 carbon matrix. While pristine Cr-N-C shows contents of pyrrolic, 776 graphitic and oxidized N similar to those of pristine Fe-N-C and Co-777 N-C, its content of pyridinic N is significantly higher, 3.08 vs. 1.75 778 at % for Fe- and Co-N-C (Table III). Similar to Fe- and Co-N-C, 779 the content of pyrrolic N in Cr-N-C increased after H2O2 treatment 780 (+0.85 at %). In contrast to Fe- and Co-N-C, the content of pyridinic 781 N decreased strongly (-1.06 at %). Since CrN particles are not known 782 to be active toward ORR, the activity of Cr-N-C is assigned to highly 783 basic N-groups formed in the carbon matrix during NH₃ pyrolysis.^{16,58} 784 The strong decrease of pyridinic N in Cr-N-C with H₂O₂ treatment 785 might explain the strong decrease in ORR activity of that catalyst after 786 H_2O_2 treatment (Fig. 4c). 787

The oxygen content and speciation is now discussed (Table II). 788 The total O-content increased dramatically after H₂O₂ treatment for 789 Fe-N-C (+8.2 at %), Co-N-C (+4.3 at %) and Cr-N-C (+6.3 at %). 790 The larger increase of O-content in Fe-N-C vs. Co-N-C is positively 791 correlated with a larger decrease in ORR activity (Figs. 5 & 7). The 792 O_{1s} spectra were fitted with four components, labelled O1 to O4 793 (Table II). The O1 component is assigned to oxygen doubly bonded 794 with C as in C=O or O=C-OH, O2 is assigned to C-O-H or C-O-C 795 groups and O3 to oxygen singly-bonded with C as in O=C-OH or 796 O=C-OR.⁷¹ The increase of the O1 and O2 components in Fe-N-C, 797 Co-N-C and Cr-N-C accounts for the major part of the total increase 798 in O-content after H₂O₂ treatment. The largest increase is observed 799 for C-O-H (hydroxyl) and C-O-C (epoxide). Those groups are known 800 to result in strong distortion of the graphene layers due to sp³ hy-801 bridization of carbon atoms.⁷² In graphene oxide, this distortion leads 802 to a interlayer spacing of 6.7 Å, ca twice that in graphite.⁷³ The in-803

crease in total O-content on Fe-N-C after treatment with 10⁻² mol $H_2O_2 \text{ mg}^{-1}$ (+8.4 at %) is very similar to the value of +8.0 at % 805 oxygen reported for a PANI-Fe-C catalyst over a 500 h PEMFC test 806 at 0.4 V.²⁶ Surface oxidation of carbon thus seems to be the main 807 degradation route during reaction with H₂O₂. The inclusion of large 808 amounts of oxygen on the edge of graphene sheets or on the basal 809 planes may have dramatically changed the electronic properties of 810 the surface, as reported for graphene nano-ribbons.^{74–75} The Co-N-C 811 catalyst was however less oxidized than Fe-N-C and Cr-N-C after 812 treatment with 10^{-2} mol H₂O₂ mg⁻¹. This agrees with the trends 813 in fluoride measurement and collection yield previously discussed 814 (Fig. 9). For Co-N-C, the weight gain of 6% observed after treat-815 ment with 10^{-2} mol H₂O₂ mg⁻¹ is commensurate with the increase in 816 O-content measured by XPS (+4.3 at. %). Hence, it seems that no car-817 bon atoms initially present in Co-N-C were lost during the treatment 818 with 10^{-2} mol H_2O_2 mg⁻¹, and oxygen atoms grafted on the catalyst 819 surface. 820

Comparison between ex situ H_2O_2 degradation and operando fuel 821 *cell degradation.*— In this section, we focus on the Fe-N-C catalyst. 822 The polarization curves of a PEMFC with a loading of 4 mg cm⁻² of 823 pristine Fe-N-C at the cathode have been recorded at BoL and after 824 different duration of galvanostatic operation at 0.5 A $\rm cm^{-2}$ (Fig. 15, 825 solid lines). A galvanostatic control was chosen in order to easily 826 calculate the electric charge that passed through the cathode. The 827 polarization curves measured at BoL of cathodes comprising Fe-N-828 C catalysts after treatment with H_2O_2 are also shown (dashed and 829 dashed-dot curves). As can be seen, the BoL curve for Fe-N-C treated with 10^{-2} mol H₂O₂ per mg corresponds to ca 76–100 h of operation

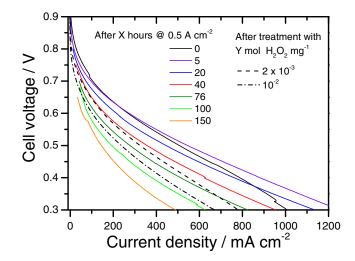


Figure 15. PEMFC polarization curves for cathodes comprising the pristine Fe-N-C and measured after X hours of operation in fuel cell at 0.5 A cm⁻² (solid lines) and for cathodes comprising Fe-N-C catalysts treated with Y mol H_2O_2 per mg and measured at BoL (dashed or dash-dot lines).

in fuel cell at 0.5 A cm⁻². If H₂O₂ released during ORR in fuel cell is 832 the most important cause of degradation under steady state operation, 833 the cumulative amount of H₂O₂ produced after 76-100 h at 0.5 A 834 cm⁻² and normalized by the mass of catalyst in the cathode should be 835 similar to the amount of 10^{-2} mol H₂O₂ per mg used for the ex situ 836 degradation. The calculation of that scalar for operando conditions 837 now only requires a hypothesis on the % H2O2 produced during ORR 838 on the Fe-N-C catalyst. This % H2O2 can be assumed on the basis of 839 the RRDE measurements on pristine Fe-N-C (Fig. 6). 840

To calculate the cumulative number of mol H₂O₂ produced in fuel 841 cell after X hours, we now derive its expression as a function of the 842 steady-state current density of the fuel cell, the duration of operation 843 and the % H₂O₂ during ORR. When the PEMFC operates at 0.5 A 844 cm^{-2} with a cathode catalyst loading of 4 mg cm⁻², then the electric 845 current produced by every mg of catalyst is 0.5/4 = 0.125 A. After 846 e.g. 100 h of operation, the electric charge passed per mg of catalyst is 847 $100 \times 3600 \times 0.125 = 4.5 \times 10^4 \text{ C mg}^{-1}$, equivalent to 0.466 mol of 848 electrons per mg of catalyst. If the % H2O2 produced as a by-product 849 of the ORR is 5% (5 mol H_2O_2 produced for 100 mol of reacted O_2), 850 then it ensues that 5 mol H₂O₂ are produced for 390 mol of electrons. 851 The latter scalar comes from 5 mol H_2O_2 x2 electrons/ H_2O_2 + 95 852 mol H₂O x4 electrons/H₂O. The scalar of 0.466 mol electrons per 853 mg can now be converted to the scalar mol H2O2 per mg catalyst by 854 multiplying it by 5/390. This yields 5.97×10^{-3} mol H₂O₂ per mg. It 855 represents the number of mol H2O2 that 1 mg of Fe-N-C catalyst has 856 produced over 100 h of operation at 0.5 A cm⁻². Based on the above 857 reasoning, the general equation relating the % H₂O₂ (x) during ORR 858 with the ratio mole H₂O₂ per mol of electrons is: 859

nol H₂O₂/mol electrons =
$$R1 = x/(400 - 2x)$$
 [1]

Also, the general relation between the current density, I, duration of 860 PEMFC operation at that current density, Δt , and the ratio of mol 861 electrons per mg catalyst is: 862

mol electrons/mg catalyst =
$$R2 = I \Delta t / (FL)$$
 [2]

Where I is the current density in A cm⁻², Δt is the duration in seconds, 863 F Faraday's constant in S.I. units and L the cathode catalyst loading 864 in mg cm⁻². From equations 1 and 2, the ratio of mol H₂O₂ per mg 865 catalyst can be expressed as: 866

$$mol H_2O_2 per mg catalyst = R1 \cdot R2 = x I \Delta t / [FL(400 - 2x)]$$
[3]

Applying this equation with x = 5, I = 0.5 A cm⁻², $\Delta t = 76$ h 867 or 100 h and L = 4 mg cm⁻², the scalars of $4.5 \cdot 10^{-3}$ and $6 \cdot 10^{-3}$ 868 mol H₂O₂ per mg catalyst are calculated. The fair agreement between 869 polarisation curves measured after 100 h of operation and that for the 870 Fe-N-C catalyst treated with 10⁻² mol H₂O₂ per mg catalyst suggests 871 that the formation of 5% H_2O_2 as a by-product of ORR is a possible 872 explanation for the degradation of the Fe-N-C catalyst under relatively 873 high load (0.5 A cm⁻²). The hypothesis of 5% H_2O_2 during ORR in 874 PEMFC is not contradicting the RRDE measurements performed on 875 pristine Fe-N-C (Fig. 6). The % H₂O₂ measured at low loading in 876 RRDE varied between 2.5 and 3.3% in the range of 0.2-0.5 V vs. 877 RHE. In conclusion, for Fe-N-C, the scalar mol H₂O₂ per mg catalyst 878 predicts both the ex situ and the operando degradation. The underlying 879 cause of the degradation is however believed to be the formation of 880 881 ROS from the decomposition of H_2O_2 on FeN_xC_y moieties.

Conclusions

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Ex situ degradation with H2O2 of a Fe-N-C, a Co-N-C and a Cr-N-883 C catalyst was investigated. The negative effect on the ORR activity 884 885 and fuel cell performance was worse for Cr-N-C, followed by Fe-N-C and least for Co-N-C. The initial activity of Cr-N-C is assigned 886 to surface N-groups while the initial activity of Fe-N-C and Co-N-887 C is assigned to MeN_xC_y moieties integrated in the carbon matrix. 888 These moieties accounted for ca 80 and 90% of the metal atoms in the 889 pristine Co- and Fe-N-C catalysts, respectively. The changes observed 890 891 with X-ray absorption and Mössbauer spectroscopy before and after degradation with H_2O_2 were minor. Most of the MeN_xC_y moieties 892

present in the pristine catalysts were still present in the Fe- and Co-N-893 C catalysts after degradation with 10^{-2} mol H₂O₂ mg⁻¹. The relative 894 loss of Fe and Co content was 17 and 27%. In parallel, the ORR activity 895 at 0.8 V vs. RHE was divided by ca 6-10 for Fe-N-C and 3 for Co-N-896 C. The super-proportional decrease of ORR activity with decreased 897 metal content can be interpreted in two ways: either only a fraction of 898 the MeN_xC_y moieties are on the top surface in pristine catalysts, or the 899 turnover frequency for ORR of the MeN_xC_y moieties remaining after 900 degradation with H₂O₂ was drastically decreased. A technique able to 901 distinguish between MeN_xC_y moieties situated in the bulk and those 902 located on the surface is needed to draw final a conclusion. The H₂O₂ 903 chemical treatment oxidized the surface of the catalysts, resulting 904 in +4 to +8 atom % oxygen. The oxidizing species is believed to 905 be radical oxygen species (ROS) generated via a Fenton reaction 906 between transition metal centres and H_2O_2 . This is supported by the 907 lower fluoride concentration and lower degradation measured for Co-908 N-C than for Fe-N-C after peroxide treatment. From a comparison 909 between ex situ and operando degradation, the chemical attack of 910 Fe-N-C by H₂O₂ seems to be the main degradation mechanism in 911 steady-state operation. To mitigate this main degradation mechanism, 912 four approaches can be envisaged for Me-N-C catalysts: i) synthesis 913 of catalysts with minimized % H₂O₂ during oxygen reduction, ii) 914 synthesis of catalysts or cathodes with a more graphitic structure, iii) 915 synthesis of catalysts that do not form ROS in the presence of H_2O_2 , 916 iv) addition of radical scavengers. 917

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