Bifunctional Zinc-Molybdate/Metal-Nitrogen-Carbon Catalytic Layers with Improved Four-Electron Selectivity for Oxygen Reduction in Acidic Medium

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Abstract

Platinum-group-metal-free metal-nitrogen-carbon catalysts for the oxygen reduction reaction have demonstrated high initial activity and power performance in proton exchange membrane fuel cells. The main challenge facing this class of catalysts is their poor durability in operating acidic fuel cells. Their key *operando* degradation mechanisms involve oxidative attacks triggered by the catalysis of the oxygen reduction on atomically-dispersed 3d transition metals. Reactive oxygen species such as H₂O₂ or radical species are particularly aggressive to the carbon matrix. Minimizing H₂O₂ by-product formation during oxygen reduction or actively scavenging any formed H₂O₂ are therefore promising routes to improve their durability.

Here, we report that Zn-molybdates have low activity towards H_2O_2 decomposition and electroreduction in acid, but result in a strong synergistic effect with various M-N-C catalysts. The addition of ZnMoO₄ in cathode layers resulted in improved four-electron ORR selectivity of M-N-C catalysts (M = Fe, Co, Cr) as well as moderately improved durability in accelerated stress tests in O₂-saturated pH 1 electrolyte. The improvement in selectivity could be rationalized by hydrogen peroxide electroreduction measurements and catalase enzyme-like activity tests. Both types of measurements revealed higher reactivity towards H_2O_2 scavenging when Metal-N-C catalysts are physically mixed (in a catalytic layer) or together in a suspension in solution with Zn-molybdate, revealing a synergy effect. Control experiments with ZnCl₂ and Na₂MoO₄ salts however suggest that leached ions from ZnMoO₄ in acidic solution are likely at the root of the synergy effect.

1. Introduction

Platinum-group-metal (PGM)-free catalysts for the oxygen reduction reaction (ORR) have been intensively developed in the last decade, resulting in significant advances in their initial activity and power performance in proton exchange membrane fuel cells (PEMFC). [1-4] However, while some Fe-N-C catalysts with atomically-dispersed FeN_x sites have met intermediate targets defined by the United States Department of Energy (< 40% decrease in ORR activity after 10,000 load cycles), [5] this high stability in acidic conditions was achieved in O₂-free environments. Comparative studies with and without O₂ in the cathode environment showed that the introduction of O₂ during load-cycling (therefore triggering the ORR) resulted in dramatically enhanced ORR activity loss. [6,7] This was observed in both a rotating disk electrode (RDE) setup and in PEMFC, and implies that the stability of Fe-N-C in acidic medium is a necessary, but insufficient condition for their successful practical implementation. This recent conclusion highlights the difference between PGM-based catalysts, for which the key operando degradation mechanisms do not involve O₂ nor the ORR (platinum dissolution, particle growth via Ostwald ripening), and PGM-free acid-stable catalysts for which the key operando degradation mechanisms are directly or indirectly related to the presence of O₂ or the occurrence of the ORR. Examples for indirect relations with O_2 or ORR are the attack by H_2O_2 or other more reactive oxygen species (ROS) [8] and ROS-induced carbon corrosion. [6] The demetallation of Fe from FeN_x sites in acidic medium was shown to be triggered or strongly accelerated when Fe-N-C materials catalyze the ORR. This has been shown both with model FeN₄ sites such as Fe-phthalocyanines and with pyrolyzed Fe-N-C catalysts comprising Fe mainly as FeN_x sites formed during pyrolysis. [5, 9] The long-recognized, but not well-understood drawback of Fenton reactions in acidic medium between PGM-free catalysts comprising a 3d transition metal and H₂O₂ is a fundamental issue for durable PGMfree cathodes in PEMFC. Hitherto, this challenge has been faced by i) developing M-N-C catalysts with high selectivity for four-electron ORR, ii) replacing Fe with another metal such as Co, Mn or Sn, [10-12] or iii) adding radical scavengers onto the M-N-C catalyst or in the cathode active layer. [13] However, these approaches have hitherto not led to satisfactory durability in operating PEMFC. While some M-N-C catalysts have shown high initial selectivity in rotating ring disk electrode (RRDE) setup, a study showed that exposing such catalysts to a small amount of H₂O₂ progressively decreases their selectivity during ORR, thereby defining an evil circle of deactivation. [8] Concerning the nature of the transition metal in M-N-C, while Fe cations are commonly regarded as the most active for Fenton reactions, other 3d transition metals (either as MN_x sites or as metal species resulting from MN_x degradation) can also be active. For example, a study on 3d metal cations performed in neutral pH showed that Cu(II), Cr(III) and Co(II) can produce more ROS than Fe(III). [14] In addition, Co-N-C catalysts have generally lower four-electron ORR selectivity than Fe-N-C, [15] which would partially offset the expected benefit of the alleged lower Fenton reactivity of Co-N-C vs. Fe-N-C, due the higher production rate of H₂O₂ for a fixed current density of the PEMFC. Last, radical scavengers such as CeO₂ can decrease the concentration of ROS, but do not scavenge H₂O₂, which is the source of ROS for M-N-C systems.

Functionalizing M-N-C catalysts or M-N-C catalytic layers with active sites that can electrochemically reduce H_2O_2 to water or chemically decompose it into O_2 and water is therefore a promising strategy for improved durability. This bifunctionality concept was investigated for example by Mechler *et al.* by adding a small amount of platinum nanoparticles (1-2 wt%) onto Fe-N-C catalysts. [16] While this resulted in dramatically improved durability in PEMFC while keeping the same apparent activity for Fe-N-C and Pt/Fe-N-C (suggesting Pt did not contribute to the initial activity, but only impacted the durability of FeN_x sites), a subsequent study showed that the annealing step applied for reducing the Pt salt to Pt NPs resulted in core-shell particles comprising a Pt core and a Fe₂O₃ shell. [17] While such core-shell particles have lower ORR specific activity than exposed Pt NPs, their ORR specific activity is significant when the Fe₂O₃ shell is thinner than 1 nm. In addition, the Fe₂O₃ shell on top of Pt resulted in the deactivation of Pt for the electroreduction of H₂O₂. As a consequence, the concept of improved durability of FeN_x sites by the addition of a H₂O₂ decomposing catalyst could not be unambiguously demonstrated in that work. Moreover, a content of 1-2 wt % Pt onto Fe-N-C combined with loadings of 4 mg $_{Fe-N-C} \cdot cm^{-2}$ leads to cathode loadings of 40-80 $\mu g_{Pt} \cdot cm^{-2}$. Similar or even higher PEMFC performance can be obtained with Pt catalysts at such low Pt loading, without resorting to Fe-N-C. [18]

Thus, while the concept of adding a PGM-free catalyst to M-N-C for the active removal of *operando* formed H_2O_2 is conceptually attractive, there are no known PGM-free catalysts with significant activity toward H_2O_2 electroreduction or decomposition in acidic medium. Metal-carbides and metal-oxides based on W, Mo, Ti and Sn are stable in the acidic conditions of PEMFC within the typical cathode potential range, but have no or little reactivity toward H_2O_2 . Metal-molybdates comprising a divalent metal cation (general formula MMoO₄, or $M_xM'_{1-x}MoO_4$ for trimetallic systems, in which M and M' can be a 3d transition metal or an alkaline earth metal of group 2A) are an interesting class of materials with specific properties provided by M (M') for optical, [19,20] catalytic, [21-25] and electrochemical energy storage applications. [26,27] In particular, ZnMoO₄ has photocatalytic, [28] optical and antibacterial properties, [29-31], and has been investigated also as an anode material for rechargeable lithium-ion battery. [32,33]

Here, we report that Zn-molybdates supported on carbon or nitrogen-doped carbon have *per se* low activity towards H_2O_2 decomposition and H_2O_2 electro-reduction in acid, but a strong synergistic effect is observed when they are combined with M-N-C catalysts comprising single-metal-atom sites. The addition of ZnMoO₄ in cathode layers resulted in improved four-electron ORR selectivity of various M-N-C catalysts (M = Fe, Co, Cr) in RRDE setup, as well as moderately improved durability in accelerated stress tests in O₂-saturated pH 1 electrolyte. The improvement in selectivity could be rationalized by hydrogen peroxide electro-reduction measurements and with catalase enzyme-like activity tests. Both types of measurements revealed higher reactivity towards H_2O_2 scavenging when Metal-N-C catalysts are physically mixed (in a catalytic layer) or added together in a suspension in solution with Zn-molybdate, revealing a synergy effect. Control experiments with ZnCl₂ and Na₂MoO₄ salts however suggest that leached ions from ZnMoO₄ in acidic solution are likely at the root of the synergy effect between atomically dispersed Metal-N_x sites and large ZnMoO₄ particles.

2. Results and Discussion

2.1. Materials characterisation

The two ZnMoO₄ polymorphs were first studied as peroxide scavengers. The α -ZnMoO₄ material was purchased from Sigma Aldrich (99.9 % pure) and used as received, while β -ZnMoO₄ was prepared from α -ZnMoO₄ by reacting it with a 30 wt% H₂O₂ aqueous solution at room temperature (see details in the Experimental Methods). The commercial α -ZnMoO₄ polymorph was phase pure, all X-ray diffraction (XRD) peaks matching those calculated for this structure (**Figure S1a**), while the prepared β -ZnMoO₄ contains a residual amount of α -ZnMoO₄ (**Figure S1b**). Their porosity was measured with N₂ sorption (**Figure S1c**). The as-received α -ZnMoO₄ is almost non-porous, while β -ZnMoO₄ adsorbs a significant amount of N₂ across the entire range of P/P⁰, implying the coexistence of some microporosity (some volume is already adsorbed at the lowest P/P^o value), and mainly mesoporosity. The BET surface area of β -ZnMoO₄ was estimated to be 44 m²·g⁻¹. Scanning electron microscopy (SEM) also identified morphological differences between the two ZnMoO₄ materials, α -ZnMoO₄ presenting 5-8 µm large cobblestone-resembling morphology with smooth outer surface, and β -ZnMoO₄ showing a fine lamellar structure about 60 nm thick and from 300 nm up to 3-5 µm in lateral size (**Figure S2**).

Then, we investigated the effect of these two ZnMoO₄ polymorphs on the ORR selectivity of Fe-N-C, Co-N-C and Cr-N-C materials with a RRDE setup. Except for the nature of the metal, the three M-N-C materials were prepared similarly *via* a two-step pyrolysis approach, the first pyrolysis leading to the formation of N-C from ZIF-8 (Basolite[®] Z1200 from BASF), and the second pyrolysis leading to M-N-C after the addition of metal chloride onto N-C (see details in the Experimental Methods). These three M-N-C materials were characterised by XRD (**Figure 1a**) and, for Fe-N-C, also with X-ray absorption spectroscopy and ⁵⁷Fe Mössbauer spectroscopy (**Figure 1b-c**). The XRD patterns only show broad peaks associated with the presence of small and/or amorphous graphite crystallites, suggesting the presence of metal as atomically dispersed M-N_x moieties and/or as ultra-small clusters. The Fourier-transform of the Extended X-ray Absorption Fine Structure (FT-EXAFS) at the Fe K-edge prior to phase correction shows a main peak at *ca* 1.5 Å for Fe-N-C, that can be assigned to Fe-N contribution from FeN_x sites, and a minor peak at *ca* 2.5 Å that can be assigned either to Fe-C interaction from the second coordination sphere of Fe in FeN_x sites, or to Fe-Fe interaction in iron oxides. The FT-EXAFS results on Fe-N-C are supported by ⁵⁷Fe Mössbauer spectroscopy (**Fig. 1c** and **Table S1**), identifying two doublets D1 and D2 that are ubiquitous in atomically-dispersed Fe-N-C catalysts, [34-36] and recently identified by us to be high-spin Fe(III)-N₄ sites and low- or medium-spin Fe(II)-N₄ sites, respectively. [37,38] The ⁵⁷Fe Mössbauer spectrum was fitted with a third spectral component, namely a broad singlet. The latter may be assigned to magnetically disordered Fe clusters, and its isomer shift of ~0.47 mm s⁻¹ suggests some oxidized form of Fe clusters.



Figure 1. Structural characterisation of M-N-C materials. a) XRD patterns of Fe-N-C, Co-N-C, and Cr-N-C, b) FT-EXAFS of Fe-N-C, c) ⁵⁷Fe Mössbauer spectrum of Fe-N-C. The Mössbauer spectrum was acquired at 5 K and the velocity scale was calibrated versus a metallic iron foil.

2.2. Activity and selectivity towards ORR

The peroxide yield and Faradaic ORR current density were measured as a function of the electrochemical potential in a RRDE setup for each M-N-C catalyst, and then for each M-N-C material with the addition of ZnMoO₄ in a mass ratio for ZnMoO₄/M-N-C of 1/10 at the ink preparation stage. The results for β -ZnMoO₄/Fe-N-C and α -ZnMoO₄/Fe-N-C show that both zinc molybdate polymorphs contribute to the decrease of the peroxide yield across the entire potential range compared to Fe-N-C alone (Figure 2a and Figure 2b, respectively). The ORR activities of the bifunctional ZnMoO₄/Fe-N-C layers are identical to that of Fe-N-C alone, as shown by superimposed polarisation curves at E > 0.8 V, indicating that the addition of ZnMoO₄ did no block Fe-based active sites. This is further supported by the cyclic voltammograms (CV) acquired in N₂-saturated electrolyte for the Fe-N-C layer and the β -ZnMoO₄/Fe-N-C bifunctional layer (Figure S3). The signals are generally superimposed, indicating similar electrochemical surface area of carbon. However, for β -ZnMoO₄/Fe-N-C, an additional reduction peak is seen at ca 0.37 V vs. RHE and the oxidation peak at 0.62-0.64 V vs. RHE has a higher intensity compared to Fe-N-C, and its position on the potential scale is negatively shifted by ca 30 mV. This can be interpreted as a redox peak contribution from β -ZnMoO₄, with broad separation between its reduction and oxidation peaks. The ORR polarisation curve of β -ZnMoO₄/Fe-N-C shows a higher and better-defined diffusion-limited current density than Fe-N-C (Figure 2a), while the polarisation curve of α -ZnMoO₄/Fe-N-C is superimposed with that of Fe-N-C across the entire potential range (**Figure 2b**). Since the peroxide yield on Fe-N-C is at maximum ca 6 % only (at 0.0 V vs. RHE), the maximum increase in current density that might be observed in the diffusion-limited region is only ca 3% if no peroxide at all were produced with ZnMoO₄/Fe-N-C layers. However, the gain in current density for β -ZnMoO₄/FeN-C vs. Fe-N-C at E < 0.7 V vs. RHE significantly exceeds this value (*ca* 7 and 21% higher current density at 0.0 and 0.5 V vs. RHE, respectively). Therefore, the higher current density observed at low potentials for β -ZnMoO₄/Fe-N-C vs. Fe-N-C is the combined result of decreased peroxide yield and improved O₂ mass-transport or layer uniformity in the β -ZnMoO₄/Fe-N-C layer. In contrast, the nearly superimposed polarisation curves for α -ZnMoO₄/Fe-N-C and Fe-N-C (**Figure 2b**) suggest that the addition of α -ZnMoO₄ did not improve O₂ mass-transport nor layer uniformity. This difference can be assigned to the contrasted morphologies of the two ZnMoO₄ polymorphs that were described previously.

On the basis of slightly further decreased peroxide yield with β -ZnMoO₄ compared with α -ZnMoO₄ and the peroxide-induced phase transformation from α - to β -form (used to prepare β -ZnMoO₄, but which may transform also α -ZnMoO₄ into β -ZnMoO₄ in *operando*, as a result of H₂O₂ production during ORR), we selected β -ZnMoO₄ to investigate its effect on other M-N-C materials. The results for β -ZnMoO₄/Co-N-C and β -ZnMoO₄/Cr-N-C (Figure 2c and Figure 2d, respectively) show strongly reduced peroxide yields compared to the measurements with Co-N-C and Cr-N-C. The trends are similar to those observed with the β -ZnMoO₄/Fe-N-C bifunctional layer, with a reduced peroxide yield across the entire potential range, similar ORR activity at 0.8 V vs. RHE with and without Zn molybdate, and increased current density with β -ZnMoO₄ in the layer at E < 0.70-0.75 V vs. RHE. The positive effect of β -ZnMoO₄ addition is dramatic with Co-N-C, due to the high peroxide yield with that catalyst alone (12-19 % at E < 0.7 V vs. RHE). As for β -ZnMoO₄/Fe-N-C, the increase in current density with β -ZnMoO₄ in the diffusion-limited region exceeds the value that would be expected from the decreased peroxide yield. Even for Co-N-C, the maximum increase in current density at low potential expected from peroxide yield measurements would be ca 11 %. However, 35 and 63 % increase in current density is experimentally observed at 0.0 and 0.5 V vs. RHE, respectively. Therefore it may be concluded that the addition of β-ZnMoO₄ in the catalytic layers of Fe-N-C, Co-N-C and Cr-N-C results in i) decreased peroxide yield and ii) improved layer uniformity or O₂ mass-transport. In summary, the results show that the addition of α -ZnMoO₄ and β -ZnMoO₄ strongly reduces the peroxide yield of M-N-C catalysts without negatively affecting the ORR activity of the layers and that this effect is applicable to M-N-C catalysts with different active metal centers (Fe, Co, Cr).



Figure 2. ORR activity and selectivity of M-N-C catalyst and ZnMoO₄/M-N-C bifunctional catalytic layers. Peroxide yield and ORR polarisation curve for a) Fe-N-C (black) and β -ZnMoO₄/Fe-N-C (red), b) Fe-N-C (black) and α -ZnMoO₄/Fe-N-C (blue), c) Co-N-C (black) and β -ZnMoO₄/Co-N-C (red), d) Cr-N-C (black) and β -ZnMoO₄/Cr-N-C (red). The M-N-C and ZnMoO₄ loadings are 800 and 80 µg·cm⁻², respectively. The electrolyte is 0.5 M H₂SO₄, scan rate 2 mV·s⁻¹ and rotation rate 1600 rpm. Note: nonidentical curves for Fe-N-C in a) and b) are due to the fact that two different batches of Fe-N-C were used.

2.3. Reactivity of Zn-molybdates and bifunctional layers towards hydrogen peroxide

Then we designed experiments to identify the underlying mechanism through which β -ZnMoO₄ and α -ZnMoO₄ decrease the peroxide yield during ORR catalysis with M-N-C catalysts. For this purpose, we used a reference Fe-N-C catalyst previously developed at our laboratory and that does not contain any Fe particles (or in such low amount that they cannot be detected by the present techniques). [34] The catalyst is labelled Fe_{0.5}, as in our previous studies. [8, 34, 37, 38] We first investigated the activity of β -ZnMoO₄ for the hydrogen peroxide reduction reaction (PRR). A layer of β -ZnMoO₄ and Ketjenblack (a high surface area carbon black) with a mass ratio β -ZnMoO₄/carbon of 1/4 shows no PRR activity down to ca 0.3 V vs. RHE (Figure S4, black curve). Under the assumption that PRR activity may arise from a synergy between M-N-C and β -ZnMoO₄, we then measured the PRR activity of β -ZnMoO₄/Fe_{0.5} and α -ZnMoO₄/Fe_{0.5} layers (1/10 mass ratio) and compared it to that of Fe_{0.5}. The results show strongly enhanced PRR activity for the bifunctional layers with similar enhancement for β -ZnMoO₄/Fe_{0.5} and α -ZnMoO₄/Fe_{0.5} layers relative to Fe_{0.5} (Figure 3). The results explain the increased four-electron selectivity for the bifunctional layers reported in Figure 2. However, the positive effects of β -ZnMoO₄ and α -ZnMoO₄ in increasing the PRR activity are similar (Figure 3), which is in slight contrast with the stronger reduction in % H_2O_2 with β -ZnMoO₄ than with α -ZnMoO₄ seen in **Figure 2**. The decreased % H_2O_2 with bifunctional layers in **Figure 2** is seen at E < 0.8 V vs. RHE, in line with the onset potential for PRR at *ca* 0.84 V vs. RHE seen in **Figure 3**. A control experiment with β -ZnMoO₄ mixed with an ironfree N-doped carbon prepared otherwise similarly as Fe_{0.5} (labelled as N-C) shows that the synergy at play is between ZnMoO₄ and the Fe-N₄ centers, not between ZnMoO₄ and metal-free nitrogen moieties or Zn-N₄ moieties present in N-C (dotted red curve in Figure 3). Also supporting this, the $%H_2O_2$ produced during ORR on N-C is high, and unmodified by the addition of β -ZnMoO₄ (Figure S5).



Figure 3. Activity toward H₂O₂ electroreduction of α - or β -ZnMoO₄/Fe_{0.5}, β -ZnMoO₄/N-C, Fe_{0.5} and N-C. H₂O₂ electroreduction was measured in 10 mM H₂O₂ in 0.5 M H₂SO₄ electrolyte at 1600 rpm with a scan rate of 2 mV s⁻¹. The loading of Fe_{0.5} and ZnMoO₄ was 800 and 80 µg·cm⁻², respectively.

To investigate the suspected activity of ZnMoO₄ in catalyzing heterogeneous reactions with hydrogen peroxide, we also applied methods previously developed for investigating the catalase-like activity of nanozymes, [39,40]. Nanozymes are generally defined *as* inorganic nanoparticle with the ability to mimick the catalytic activity of natural biomolecules. [41] The activity of the enzyme catalase or a nanozyme enzyme-mimetic is therefore the ability to accelerate the disproportionation of H_2O_2 into O_2 and water, without the need for an electron donor. The catalase-like activity of β -ZnMoO₄ and α -ZnMoO₄ was investigated in the presence of N-C or Fe_{0.5} by measuring the rate of O_2 evolved from H_2O_2 as a function of time (see Methods). Control experiments were also performed with N-C or Fe_{0.5} alone. For each material (or combination), the O_2 evolution rate was measured for four different H_2O_2 concentrations (3, 15, 30, 60 mM). The plots for oxygen evolution at all four concentrations are shown for N-C, Fe_{0.5}, β -ZnMoO₄/N-C and α -ZnMoO₄/Fe_{0.5} in **Figure S6**, and for β -ZnMoO₄/Fe_{0.5} in **Figure 4a**. For catalase-like activity tests, the ZnMoO₄ content was always kept fixed at 5 wt % relative to Fe_{0.5} or N-

C. The O₂ evolution vs. time curves can be compared for all materials or material combinations at the fixed concentration of 60 mM H₂O₂, as shown in **Figure 4b**. At short times, the trend of the O₂ evolution rate for the different materials or material combinations is β -ZnMoO₄/Fe_{0.5} $\geq \alpha$ -ZnMoO₄/Fe_{0.5} >> Fe_{0.5} $>> \beta$ -ZnMoO₄/N-C > N-C $\geq \beta$ -ZnMoO₄. This trend is in line with the results of peroxide yield observed in RRDE setup (**Figure 2**). While the coexistence in solution of β -ZnMoO₄ and Fe_{0.5} initially leads to the highest O₂ evolution rate, a higher conversion rate of peroxide into O₂ is observed after around 2 h with the combination of α -ZnMoO₄ and Fe_{0.5}, and a higher saturation conversion is also achieved after 3 h (96% vs. 85%, **Figure 4b**). However, at lower peroxide concentrations, both β -ZnMoO₄/Fe_{0.5} and α -ZnMoO₄/Fe_{0.5} catalyze an almost complete conversion of H₂O₂ to O₂ within 2 h (94-97%) with initial H₂O₂ concentrations of 3 and 15 mM (**Table S2**). This suggests that the decomposition channel is fast and selective. Such rapid and almost complete conversion of H₂O₂, even at low peroxide concentration, is an important property in the context of PEMFC application.



Figure 4. a) Produced O_2 amount versus time for β -ZnMoO₄/Fe_{0.5} with different concentrations of H₂O₂. b) Conversion of H₂O₂ into O₂ on the different materials at pH 4, (sodium citrate buffer solution 0.1 M) starting from the same initial concentration of H₂O₂ (60 mM).

Notably, the catalytic activities observed with iron-free materials or material combinations, *i.e.* β -ZnMoO₄, N-C and β -ZnMoO₄/N-C are modest, all iron-free combinations showing similar and low initial reaction rates. This suggests that iron species might be the catalytic centers, and that a synergy between zinc molybdates and FeN_x centers further accelerates the peroxide disproportionation by Febased sites. In contrast, no synergistic effects can be observed between the zinc molybdates and the nitrogen moieties in N-C, also in line with the electrochemical investigation in RRDE.

We then attempted to analyze the reaction profiles of peroxide heterogeneous disproportionation within the theoretical framework of a Michaelis-Menten kinetic model, which is commonly used for enzymes and nanozymes. However only in the case of β -ZnMoO₄/Fe_{0.5}, α -ZnMoO₄/Fe_{0.5}, and Fe_{0.5} were straight lines observed in the Lineweaver–Burk double reciprocal plots (Figure S7). The results of the Michaelis-Menten analysis for Fe_{0.5}, β -ZnMoO₄/Fe_{0.5} and α -ZnMoO₄/Fe_{0.5} are reported in **Table S3**. The data clearly confirm the hypothesis that the combination of zinc molybdate with the iron catalyst results in an increase in the catalase-like activity that is higher than the simple sum of the single catalytic activities. The general trend deduced from the direct comparison of the reaction profiles is quantitatively confirmed, with β -ZnMoO₄/ Fe_{0.5} being more active than α -ZnMoO₄/ Fe_{0.5}, especially in terms of V_{max} , and both of them being more active than Fe_{0.5.} Notably, the trend of the K_m values indicates that H₂O₂ has higher affinity for the iron centers than the molybdates. Overall, the catalytic performance of β -ZnMoO₄/Fe_{0.5} is very good compared with that of other transition-metal-based nanozymes developed to mimic catalase (Table S4), such as ferritin-platinum nanoparticles, [42] graphene Fe_2O_3 nanohybrids, [40] Co_3O_4 , [43-44] and capped Fe_2O_3/Fe_3O_4 nanoparticles. [45] Given the intended application of ZnMoO₄/Fe-N-C materials in the acidic environment of PEMFC, a series of experiments was performed in more acidic buffered solutions, *i.e.* phosphate buffer solution (PBS) with a pH of 2.42, and KCI/HCI with a pH of 0.9 (Figure S8). As expected, the catalytic activity is strongly suppressed as the pH is reduced, although peroxide disproportionation is still clearly detectable for 5 wt% β-ZnMoO₄/Fe_{0.5}, whereas other groups previously reported that other nanozymes in similar conditions are completely inactive. [42]

Further insights on the mechanism of hydrogen peroxide decomposition were obtained from XPS measurements (Figure 5). The β -ZnMoO₄ and its physical mixture with either Fe_{0.5} or N-C were suspended in 60 mM H_2O_2 solution buffered to pH 4 by citrate, incubated for 30 min and then drop cast on a solid support and dried under vacuum. The resulting dried films were investigated by XPS and the spectra were compared to the data acquired on samples that were not exposed to H₂O₂. For all samples, the Zn 2p_{3/2} photoemission line did not undergo any apparent change after the exposure to H₂O₂, showing a single component peak centered at about 1022 eV, which corresponds to Zn²⁺. On the other hand, the Mo 3d photoemission line exhibits an interesting behavior. In the pristine β -ZnMoO₄ sample, the Mo 3d photoemission line consists of a single doublet component with the Mo $3d_{5/2}$ peak centered at 232.2 eV, which can be ascribed to Mo⁶⁺ ions. After the incubation with H₂O₂ an additional component can be identified at 230 eV, which indicates the formation of Mo⁴⁺ species. [46, 47] Whereas in the pure β -ZnMoO₄ sample, such Mo⁴⁺ represents only 22 % of the whole Mo species after H_2O_2 incubation, in the physical mixture of β -ZnMoO₄ with Fe_{0.5}, it is possible to observe the complete disappearance of Mo^{6+} and its complete conversion to Mo^{4+} after the same H_2O_2 incubation. Moreover, when β -ZnMoO₄ is mixed with N-C, no reduction of molybdenum is observed after H₂O₂ incubation, ruling out that the reduction can be due to a direct solid-state reaction between the metal oxide and the carbon support. The body of these spectroscopic data therefore indicates a strong synergy between iron species and the metal oxide. Iron is key to activate the H₂O₂ molecule, which therefore is able to reduce Mo⁶⁺ to Mo⁴⁺. This explains why the catalytic activity towards the decomposition of H_2O_2 is poor on the zinc molybdate alone (*i.e.* H_2O_2 is not activated), while is boosted in the β -ZnMoO₄/Fe_{0.5} composite system (*i.e.* the redox active Mo species may sustain a catalytic cycle as promoters in synergy with FeN_x sites).



Figure 5. Mo 3d (a) and Zn 2p (b) photoemission spectra acquired on the different materials or material combinations after incubation in citrate buffer solution with or without H_2O_2 (concentration 60 mM).

Given the application of these catalysts at low pH conditions (pH ~ 1) in PEMFC, we explored the possibility that the Zn-molybdates can partially dissolve and provide ions that can act as homogeneous catalysts. We therefore evaluated the catalase-like activity of soluble salts (0.05 mg/mL) of Zn²⁺ (ZnCl₂) and MoO₄²⁻ (Na₂MoO₄), either with or without Fe_{0.5} suspended in solution, in order to identify the possible existence of synergistic effects between Fe_{0.5} and these ions. The kinetic profiles of O₂ evolution as a function of time clearly indicate the molybdates anions have *per se* a low activity, whereas Zn²⁺ cations are significantly active with an overall performance for ZnCl₂ alone that is comparable to that of Fe_{0.5} (**Figure S9**). As in the previous cases, strong synergistic effects are revealed when Fe_{0.5} and the metal salts are simultaneously present in solution (**Figure S9**), and this is particularly visible in the case of i) Na₂MoO₄ and Fe_{0.5} combination, and ii) ZnCl₂ and Fe_{0.5} combination, which achieve a higher performance than β -ZnMoO₄/Fe_{0.5}. This agrees with the XPS data, indicating that only in the presence of Fe_{0.5}, the molybdate phase was activated and able to react with H₂O₂. These data help also to explain why systems obtained by a physical mixture of micrometric nanoparticles

unexpectedly exhibit such a strong (electro)catalytic synergy, which on the other hand implies a nanoscale interaction. In fact, we can assume that the high activity observed in the heterogeneous $ZnMoO_4/M-N-C$ composites results from the interaction of dissolved ions, either Zn(II) or $MoO_4^{2^-}$, with Fe-N_x centers. Such a mechanism is particularly appealing because it may activate Fe-N_x catalytic centers within micro- and meso- pores that normally would not have contact with large $ZnMoO_4$ particles.

2.4. Accelerated stress tests in RDE setup

Based on the foregoing results and identification of β -ZnMoO₄ as a promising peroxide scavenger working in synergy with Fe-N_x sites to decrease the peroxide yield during ORR, we performed loadcycling accelerated stress tests (AST) in RRDE setup on the bifunctional layer of β -ZnMoO₄/Fe_{0.5} and compared the results to those with Fe_{0.5} alone. Figure 6 shows comparable increase in peroxide yield at end-of-test (EoT) relative to beginning-of-test (BoT) for $Fe_{0.5}$ (Figure 6a) and β -ZnMoO₄/Fe_{0.5} (Figure 6c), and also comparable decrease in ORR activity (Figure 6b and 6d). The electrochemical potential at -1 mA cm⁻² shifted negatively by 36 mV for Fe_{0.5} and only 28 mV for β -ZnMoO₄/Fe_{0.5}. The data therefore suggests a partial stabilisation of the ORR activity of $Fe_{0.5}$ by β -ZnMoO₄ in acidic medium. The incomplete stabilisation may be due to the low concentration in the catalytic layer of the ions resulting from ZnMoO₄ dissolution, or to a still insufficient rate of peroxide scavenging. The peroxide produced during ORR may undergo three main routes: a) removal from the catalytic layer by diffusion/convection, b) reaction with Fe-N_x sites leading to formation of ROS, c) reaction with $ZnMoO_4$ modified Fe-N_x sites, leading to peroxide disproportionation or electroreduction. For an efficient protection of Fe-N_x sites, the rate of the route c) should be higher than route b), which is likely not yet achieved with the present systems. Future studies will be needed to further improve our fundamental understanding of the synergy between ZnMoO₄ or its constituting metal ions and Metal-N_x sites, and practical approaches must be developed to efficiently utilize this effect in catalytic layers, especially in the PEMFC environment.



Figure 6. Stability of β-ZnMoO₄/**Fe**_{0.5} **bifunctional catalytic layer compared to Fe**_{0.5}. a) Peroxide yield at BoT and EoT for Fe_{0.5}; b) ORR polarisation curve at BoT and EoT for Fe_{0.5}; c) Peroxide yield at BoT and EoT for β-ZnMoO₄/Fe_{0.5}; d) ORR polarisation curve at BoT and EoT for β-ZnMoO₄/Fe_{0.5}. The solid and dashed curves correspond to BoT and EoT results, respectively. The Fe_{0.5} and β-ZnMoO₄ loadings are 800 and 80 µg·cm⁻², respectively. Electrolyte is 0.5 M H₂SO₄, scan rate 2 mV·s⁻¹ and rotation rate 1600 rpm for ORR polarisation curve acquisition. The AST consisted of 10,000 cycles between 0.6 and 0.925 V vs. RHE in O₂-saturated electrolyte, at a scan rate of 100 mV s⁻¹.

3. Experimental Methods

3.1. Material synthesis

3.1.1. Synthesis of β -ZnMoO₄

The α -ZnMoO₄ was purchased from Sigma Aldrich (99.9 %) and was converted to β -ZnMoO₄ as follows. Approximately 4 g of α -ZnMoO₄ was dispersed in 10 mL of deionized water and placed in an ultrasonic bath for 5 minutes. 1 mL of 30 % (v/v) H₂O₂ solution (Sigma Aldrich) was then added to the α -ZnMoO₄ dispersion and left at room temperature under continuous stirring for 24 h. After 24 h, the dispersion was filtered and rinsed several times with deionized water then dried in air at 80° C overnight.

3.1.2. Synthesis of Fe-N-C, Co-N-C and Cr-N-C

Nitrogen doped carbon was synthesized from zeolitic imidazole framework ZIF-8 (Basolite® Z1200 Sigma Aldrich). First, 1.6 g of ZIF-8 was added to a 45 mL zirconia grinding bowl with 100 zirconia grinding balls with diameter 5 mm. The precursor was then ball milled at a rotation of 400 RPM for 4 cycles each of 30 min. Next, the precursor was inserted in a horizontal quartz reactor pre-heated to 1050 °C and purged in Ar. A flash pyrolysis was performed for 1 hour and the quartz reactor was then quenched to room temperature. The N-doped carbon product is labelled N-C. Metal coordination on N-C was performed by first mixing 0.5 g of N-C with MeCl₂ (98% anhydrous Sigma Aldrich) to give a total metal loading on NC of 1.0 wt. %. The N-C-flash and MeCl₂ precursor were added to a 45 mL zirconia grinding bowl with hundred 5-mm diameter zirconia grinding balls and then ball milled at a rotation of 500 RPM for 4 cycles each at 30 min. After milling, the catalyst precursor was inserted in a horizontal quartz reactor pre-heated to 800 °C and purged in Ar. A flash pyrolysis was performed in Ar with a dwell time of 1 hour, after which the quartz reactor was quenched to room temperature.

3.1.3. Synthesis of Fe_{0.5}

Fe_{0.5} was prepared by mixing zeolitic imidazole framework ZIF-8 (Basolite® Z1200 Sigma Aldrich), 1,10phenanthroline (Sigma Aldrich), and anhydrous Fe(II) acetate (Sigma Aldrich) with a mass ratio of ZIF- 8:1,10-phenanthroline of 4:1. Fe(II) acetate was added to give a total iron content of 0.5 wt. % in the precursor mixture. 1.0 g of the precursor mixture was added to a 45 mL zirconia grinding bowl with 100 5-mm zirconia grinding balls then ball milled at a rotation of 400 RPM for 4 cycles each at 30 minutes. Next, the precursor was inserted in a horizontal quartz reactor pre-heated to 1050 °C and purged in Ar. A flash pyrolysis was performed for 1 hour and the quartz reactor was quenched to room temperature

3.2. Electrochemical measurements in RRDE and RDE setups

3.2.1. ORR activity and peroxide yield measurement in RRDE

Catalyst inks were prepared using an ionomer to carbon mass ratio of 0.5. Nafion 1100 EW perfluorosulfonic acid solution (5 wt. % in aliphatic alcohols containing 15- 20 % H₂O, Sigma Aldrich) was added to a suspension of catalyst (either carbon, N-C or M-N-C or a physical mixture of 90 wt % carbon, N-C or M-N-C with 10 wt% α - or β -ZnMoO₄) in 1-propanol with a carbon concentration of 2 wt. %. The inks were sonicated in an ultrasonic bath at 20° C for 1 h prior to being drop-cast on the working electrode, to give a total carbon loading of 800 µg·cm², unless otherwise stated. Electrochemical measurements were performed in a 3-electrode cell using a graphite counter electrode and reversible hydrogen reference electrode. ORR polarization curves were performed using a glassy carbon working electrode with platinum ring held at a constant potential of 1.3 V vs. RHE. Polarization curves were measured in O₂ saturated 0.5 M H₂SO₄ (prepared from Merck 98% Merckmillipore) at a scan rate of 2 mV·s⁻¹ starting from OCP then scanned from 0 to 1 V vs. RHE with a rotation speed of 1600 RPM. Prior to ORR measurements a catalyst break-in procedure was performed where the working electrode was cycled at 20 mV·s⁻¹ in N₂ saturated electrolyte until negligible change in capacitance was observed.

3.2.2. PRR activity measurement in RDE

Identical catalyst ink formulation and RDE setup was used for PRR measurements with the exception that the working electrode used was a glassy carbon RDE without a platinum ring. After the catalyst break-in procedure in N_2 saturated electrolyte, an aliquot of 30 % (v/v) H_2O_2 solution (Sigma Aldrich)

was added to the electrolyte to give a concentration of 10 mM H_2O_2 in N_2 saturated 0.5 M H_2SO_4 . PRR polarization curves were measured at 2 mV·s⁻¹ starting from OCP then scanned from 0 to 1 V vs. RHE with a rotation speed of 900 RPM.

3.3. Catalase measurements

The measurements for the evaluation of the catalase-like activity were performed in a three-neck round-bottomed flask: one for the O₂ probe (NEOFOX –KIT PROBE from Ocean Optics) used to measure continuously in time the produced oxygen, one neck for the septum for the injection of H₂O₂, and the last one for the expansion chamber to allow constant pressure in the system. The catalytic tests were performed with 16.5 mL of buffer solution (i.e. sodium citrate 0.1 M pH 4.02, or PBS pH 2.4 or HCl/KCl pH 0.9) and 0.51 mL of a catalyst suspension made up of either 1 mg/mL Fe_{0.5} or 1 mg/mL NC, in which 0.05 mg mL⁻¹ of $\alpha(\beta)$ -ZnMoO₄ or other soluble salts (ZnCl₂ or Na₂MoO₄) was further added. The oxygen produced by the H₂O₂ disproportionation was removed from the head space of the flask (head space volume 51.4 mL), whereas the unreacted H₂O₂ remaining in solution was independently measured to verify the overall yield of the catalase-like reaction. The detection of H₂O₂ was carried out using a titanium oxysulfate solution (~ 5 % Ti basis, in H₂SO₄ 27–31 %, Sigma-Aldrich), which forms a complex with H₂O₂ with a characteristic absorption peak at 409 nm. An optical absorbance vs concentration calibration curve in the range from 1.2 μ M to 2 mM of H₂O₂ was used to quantify the H₂O₂ concentration remaining at the end of the catalytic tests.

3.4. Structural and morphological characterizations

3.4.1. X-ray diffraction measurements

X-ray diffraction patterns of the M-N-C materials and the Zn-molybdates were recorded using a PANalytical X'Pert Pro powder X-ray diffractometer equipped with a Cu Kα radiation source and 1-D strip detector in Bragg-Brentano focusing geometry. A receiving slit filter was used to remove Cu Kβ. Powder specimens were loaded into zero-background Si single crystal sample holders mounted on a

spinning sample stage to improve particle statistics. A continuous scan rate of 1.5 °/min was used and intensity was averaged over a 0.05° step size.

3.4.2. ⁵⁷Fe Mössbauer spectroscopy measurement

The ⁵⁷Fe Mössbauer spectrum measured for the Fe-N-C prepared via a two-step synthesis in the present study was acquired in identical conditions to those applied in our earlier studies to measure the spectrum of Fe_{0.5}. The source was ⁵⁷Co in rhodium and the spectrometer was from Wissel (Germany). The spectrum was acquired in transmission mode. The velocity driver was operated in constant acceleration mode with a triangular velocity waveform. The velocity scale was calibrated at room temperature with the magnetically split sextet of a high-purity α -Fe foil. The measured spectrum was fitted to appropriate combinations of Lorentzian profiles representing quadrupole doublets, sextets by least-squares methods. The fittings were performed with unconstrained parameters (relative area, IS, QS, LW, H) for each spectral component. The fitted isomer shift (IS) values are reported relative to α -Fe at room temperature. Fe-N-C powder was mounted in a 2 cm² holder. Mössbauer measurements at 5 K were performed in a helium flow cryostat (SHI-850 Series from Janis, USA).

3.4.3. X-ray absorption spectroscopy

Fe K-edge X-ray absorption spectroscopy (XAS) was conducted in transmission mode on the Fe-N-C material at room temperature at the SAMBA beamline (Synchrotron SOLEIL). The beamline is equipped with a sagittally focusing Si 220 monochromator and two Pd-coated mirrors that were used to remove X-ray harmonics. The catalyst was pelletized as a disk of 10 mm diameter using Teflon powder (1 μm particle size) as a binder.

3.4.4. Scanning electron microscopy

The morphology of the Zn molybdates was investigated with SEM (Hitachi S-4800, Hitachi, Tokyo, Japan) operating at 2 kV. Zn molybdate samples were dispersed on electrically conductive adhesive tape followed by gold metallization to improve sample conductivity.

3.4.5. Nitrogen physisorption measurements

The Brunauer–Emmett–Teller (BET) surface area and pore volume of the Zn-molybdates were measured with N_2 sorption at 77 K using a Micromeritics ASAP 2020 instrument. Before the measurement, the samples were degassed at 200 °C for 12 h under vacuum.

3.4.6. XPS measurements

X-ray photoemission spectra were acquired in a custom build Ultra High Vacuum chamber with a base pressure of 5×10^{-10} mBar coupled with a non-monochromatized double-anode X ray source (Omicron DAR-400, Scienta-Omicron GmbH, Uppsala, Sweden), a hemispherical electron analyzer (Omicron EA-125, Scienta-Omicron GmbH, Uppsala, Sweden) and a 5-channeltron detection assembly. The electron analyzer had an acceptance angle of $\pm 4^{\circ}$ and the diameter of the analyzed area was 3mm. The spectra were collected with Al-k α radiation, for the high-resolution spectra was used 20 eV as pass energy, a step of 0.1 eV and a dwell time of 0.25 s.

4. Conclusions

The addition of zinc molybdates to M-N-C catalysts was shown to improve the selectivity toward the four-electron reaction path in the ORR in acidic conditions. RRDE measurements suggest that both ZnMnO₄ polymorphs, and especially the β phase, when combined with M-N-C catalysts exhibit excellent electrocatalytic peroxide reduction activity, confirming their potential to improve cathode layer durability by keeping the H_2O_2 and ROS concentration low. The high activity in H_2O_2 conversion was also confirmed by a series of catalase-like assays. The systematic investigation of different combinations of materials and model compounds allowed us to conclude that the FeN_x sites are the active catalase-like sites, but that their efficiency in significantly improved in the presence of Znmolybdates. The study of the catalase-like activity of soluble metal salts, such as $ZnCl_2$ and Na_2MOO_4 , which represent the homogenous catalyst counterparts of ZnMoO₄, suggest that these metal ions when leached from ZnMoO₄ particles at low pH conditions can be responsible for the synergistic interactions between ZnMoO₄ and FeN_x sites. Despite the excellent activity in both the chemical disproportionation and the electrochemical reduction of hydrogen peroxide, the results of accelerated stress tests in O₂-saturated acidic conditions showed only a modest improvement in the durability of catalytic layers comprising a Fe-N-C catalyst and β-ZnMoO₄. This could be explained by the still insufficiently effective mixing between the M-N-C layer and the zinc molybdate or by the need for even faster abatement of the local H_2O_2 concentration in order to fully suppress the possible formation of other ROS.

Acknowledgements

This work was funded by the Fuel Cells and Hydrogen 2 Joint Undertaking (now the Clean Hydrogen Joint Undertaking) under grant agreement no. 779366 (CRESCENDO project). This Joint Undertaking receives support from the European Union's Horizon 2020 research and innovation program, Hydrogen Europe and Hydrogen Europe Research. We are grateful to Bernard Fraisse for help with XRD

measurements, to Nicolas Donzel for help with N_2 sorption measurements, and to Andrea Zitolo for help with XAS measurements.

Declaration of competing interest

The authors have no competing interests to declare.

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