



Cite this: *Mater. Horiz.*, 2024, 11, 3437

Received 14th March 2024,
Accepted 25th April 2024

DOI: 10.1039/d4mh00293h
rsc.li/materials-horizons

Revisiting the phosphonium salt chemistry for P-doped carbon synthesis: toward high phosphorus contents and beyond the phosphate environment[†]

Rémi F. André, ^{*a} Christel Gervais, ^b Hannes Zschiesche, ^a Teodor Jianu, ^a Nieves López-Salas, ^{ac} Markus Antonietti ^{*a} and Mateusz Odziomek ^{*a}

The introduction of phosphorus and nitrogen atoms in carbon-catalysts is a common way to tune the electronic density, and thereby the reactivity, of the material, as well as to introduce surface reactive sites. Numerous environments are reported for the N atoms, but the P-doping chemistry is less explored and focuses on surface PO_x groups. A one-step synthesis of P/N-doped carbonaceous materials is presented here, using affordable and industrially available urea and tetrakis(hydroxymethyl)phosphonium chloride (THPC) as the N and P sources, respectively. In contrast to most of the synthetic pathways toward P-doped carbonaceous materials, the THPC precursor only displays P–C bonds along the carbon backbone. This resulted in unusual phosphorus environments for the materials obtained from direct thermal treatment of THPC–urea, presumably of type C–P–N according to ^{31}P NMR and XPS. Alternatively, the *in situ* polymerization and calcination of the precursors were run in calcium chloride hydrate, used as a combined reaction medium and porogen agent. Following this salt-templating strategy led to particularly high phosphorus contents (up to 18 wt%), associated with porosities up to $600 \text{ m}^2 \text{ g}^{-1}$. The so-formed P/N-doped porous materials were employed as metal-free catalysts for the mild oxidative dehydrogenation of N-heterocycles to N-heteroarenes at room temperature and in air.

1. Introduction

The intentional introduction of heteroatoms into carbon matrices allows the tuning of the material electronic structure

New concepts

Phosphorus-doped carbonaceous materials have garnered interest for applications in catalysis, energy storage or water treatment. However, much of the research has relied on oxidized precursors like H_3PO_4 , limiting the variety of accessible P-containing functional groups. In this study, inspired by a robust chemistry developed by the textile industry, we revisit the condensation, and calcination, of $\text{P}(\text{CH}_2\text{OH})_4^+$ (THPC) with urea to synthesize original P/N-doped carbonaceous materials. The peculiar features of THPC, such as the high reactivity of its hydroxymethyl groups and the sole presence of P–C bonds, result in unreported phosphorus environments in carbons, tentatively attributed to C–P–N domains. Besides, utilizing a metal chloride hydrate as the reaction medium for polymerization, echoing a salt-templating approach, offers leverage to adjust the composition and porosity of the materials. Benefiting from the exceptionally large P doping levels (18 wt%), the material was explored as a catalyst for the oxidative dehydrogenation (ODH) of N-heterocycles under mild conditions using air as an oxidant. This work not only presents a novel strategy for producing highly doped carbonaceous materials from simple precursors, but also offers insights into the active sites of metal-free catalysts for ODH reactions, particularly through comparison with other heteroatom-doped carbonaceous materials.

and exposed chemical groups, directly affecting the performances of the materials for catalysis,^{2–5} energy storage,^{6,7} or environmental remediation applications.^{8,9} For both chemo-catalysis and electro-catalysis, the doping with phosphorus (and possibly the co-doping with nitrogen) permits the regulation of the energies of adsorption of the reagents and chemical intermediates in order to improve the activity and/or the selectivity.^{10–17} Surface functional groups, typically phosphate or phosphinate ones, were also reported as being the active sites for Brønsted acid-catalysed reactions (e.g. esterification, dehydration, and hydrolysis),^{18–22} oxidation of alcohols under mild conditions,^{23,24} and for increasing the selectivity of carbon-based catalysts in oxidative dehydrogenation (ODH) of alkanes to alkenes.²⁷

Phosphorus atoms are usually incorporated in carbonaceous materials *via* the direct calcination of a mix of C and P-containing chemicals or *via* the post-treatment of a prefabricated carbon with a phosphorus compound at high

^a Colloid Chemistry Department, Max Planck Institute of Colloids and Interfaces (MPIKG), 14476 Potsdam, Germany. E-mail: mateusz.odziomek@mpikg.mpg.de

^b Sorbonne Université, CNRS, Laboratoire de Chimie de la Matière Condensée de Paris (LCMCP), 4 place Jussieu, 75005 Paris, France

^c Chair of Sustainable Materials Chemistry, Paderborn University, Warburger Strasse 100, 33098, Paderborn, Germany

[†] Electronic supplementary information (ESI) available: SEM-EDX sample composition for salt system optimization; pore size distribution analysis; XRD, FTIR, and XPS results; XPS fitting parameters; solid-state NMR spectra; conversion of indoline as a function of solvent polarity and catalyst loading; recycling test; NMR spectra of substrate scope. See DOI: <https://doi.org/10.1039/d4mh00293h>



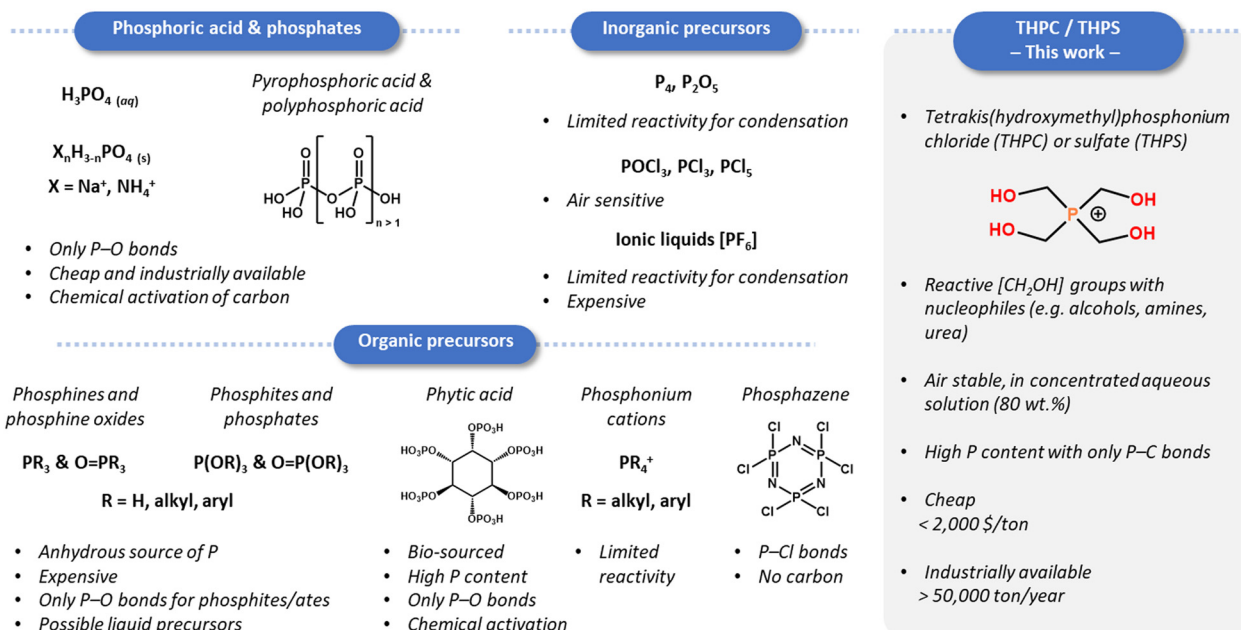


Fig. 1 Overview of the phosphorus sources used for the synthesis of P-doped carbonaceous materials.¹

temperature.¹ Various P sources were explored depending on the requirements of the synthesis (air sensitivity, absence/presence of water, and liquid/solid state), the targeted bonding environment (P–O vs. P–C bonds), the cost, the toxicity, and the potential release of phosphoric acid which then activates/etches the carbon (Fig. 1).²³ Most of these precursors present an oxidized P environment with only P–O bonds (e.g. H₃PO₄, P₂O₅, phytic acid, and O=P(OR)₃), or do not allow a straightforward condensation with the carbon precursor at low temperature (e.g. elemental phosphorus and PR₃). The quest for new P-containing groups in carbonaceous materials, beyond the usual phosphate groups [PO₄] (see Fig. S1, ESI† for nomenclature), and for higher P contents therefore goes through the exploration of different precursors.

As early as 1999, Oya *et al.* proposed to carbonize a novolac-type phenolic resin, cross-linked by tetrakis(hydroxymethyl)-phosphonium sulphate (THPS) as the P-source, to yield carbons with a P content of 5 wt%.³⁰ Such an approach looks promising to reach trivalent or reduced P sites as the phosphorus atoms are placed along the carbon backbone with only P-C bonds.^{31,32} The THP salts (THPS and the chloride equivalent, THPC) indeed condense readily with urea and amines to yield P-CH₂-NH bridges, thanks to the reactivity of the hydroxymethyl groups.³³ For decades, the corresponding polymers and resins, typically the “THPC-urea”, have been studied as a flame-retardant finishing on textiles, in both academia and industry.^{34,35} Besides, THPS and THPC are produced and commercialized on a large scale (applications in fire-retardant treatments and as microbiocides or reducing agents) and should be considered as low-cost commodities, with a price established below 2000 \$ per ton for a global market size above \$200 million in 2022.³⁶ The use of THPC/THPS was not further explored after Oya’s work for P-doped carbon synthesis, likely due to the limited surface areas obtained by the classical

calcination approach. The emergence of salt-templating and ionothermal strategies in these past two decades to synthesize porous inorganic and carbon materials appears as a promising solution to fully exploit this THPC specific chemistry.^{37,38}

We propose herein a one-step procedure to obtain porous P/N-doped carbons through the *in situ* formation of a THPC-urea resin in a metal salt hydrate, directly followed by its calcination. The as-synthesized samples display phosphorus contents among the highest reported in the literature (up to 18 wt%) while maintaining a high N content of 10 wt% and decent specific surface areas up to $600 \text{ m}^2 \text{ g}^{-1}$. In a first part, we discuss the underlying mechanism and how the used amounts of porogen salt affect the development of micro- and mesopores. We then analyse the different stages of the synthesis by means of EELS (electron energy loss spectroscopy), XPS (X-ray photoelectron spectroscopy) and solid-state ^{31}P NMR (nuclear magnetic resonance) spectroscopy. This last technique not only confirms different intermediates but also reveals unusual phosphorus environments for the samples produced in the absence of CaCl_2 , tentatively attributed to C-P-N based structures. This confirmed the interest of the initial choice of this atypical precursor to expand the library of P environments in carbonaceous materials. The so-formed material finally proved to be an efficient metal-free carbo-catalyst for the aerobic ODH of N-heterocycles to N-heteroarenes under mild conditions.

2. Results and discussion

2.1 Calcination of THPC-urea resin: from neat reaction to salt-templating synthesis

Inspired by the robust chemistry of THPC-urea finishing treatment developed by the textile industry for combustion

resistance applications, we first envisioned the thermal treatment of a THPC–urea resin as a path to obtain high concentration of hetero-elements in a carbonaceous material, while limiting the mass loss upon calcination and achieving high yields.^{31,33,39} The chemical synthesis of the P/N-doped carbonaceous materials consisted in the mixing of the THPC, available as an 80 wt% aqueous solution, with urea in a 1:2 stoichiometry, followed by heat treatment at 600–800 °C (Fig. 2A). The condensation was first investigated by thermogravimetric analysis (TGA) coupled with mass spectrometry (MS) to characterize the produced fragments and volatile species (Fig. 2B). A first mass loss of 21% is detected at 120 °C, accompanied by a significant release of H_2O ($m/z = 18$) and, to a lower extent, of NH_3 ($m/z = 17$) and CH_3OH ($m/z = 32$) (region I: polymerization). The water originates from the initial THPC aqueous solution and from the condensation reactions between hydroxymethyl groups and urea (Fig. 3A, step 1).³³ A second mass loss starts at 236 °C, up to 450 °C, accounting for a total of 50 wt% and concomitant with the release of CH_4 ($m/z = 16$), NH_3 ($m/z = 17$), CO ($m/z = 28$), formaldehyde CH_2O ($m/z = 30$), and CO_2 ($m/z = 44$) (region II: thermal restructuration) (step 2). Accordingly, foaming clearly occurs between 200 °C and 350 °C (Fig. 1C, left) and the resin becomes brown. The formation of methane and H_2 ($m/z = 2$) at *ca.* 500 °C indicates the formation of chemical unsaturation and aromatization (step 3). Afterwards, the so-formed material is thermally stable up to 800 °C (region III) before presenting a final mass loss, accompanied by the release of CO and CO_2 at 860 °C due to a severe degradation (region IV). Part of the phosphorus-containing compounds may be degraded in volatile elemental phosphorus. The P_4 tetramers ($m/z = 124$), stable in the gas phase at high temperature, were however not detected by MS, potentially because of the decrease in

temperature in the connection between the TGA and MS devices. The synthesis was therefore performed in a tubular oven, and deposition of a red solid was indeed observed downstream of the crucible at $T > 600$ °C, attributed to the condensation of red phosphorus ($T_{\text{boiling}}(\text{red P})$ *ca.* 500 °C) (Fig. S2, ESI†).

The thermal treatment at 600 °C of THPC–urea resin leads to a P/N co-doped carbonaceous material with decent yields (48% C yield, 54% P yield, all yield data available in Table S1, ESI†) but with virtually no porosity (Table 1 and Fig. S3, ESI†). Following a salt-templating strategy, the precursors were mixed with a metal chloride hydrate to achieve an intimate liquid mixture where the metal salt acts as reaction a medium and porogen.^{40,41} We conducted preliminary tests to determine an optimal salt system and retained $\text{CaCl}_2 \cdot 4\text{H}_2\text{O}$ which enabled specific surface areas significantly larger than in the absence of salt ($> 100 \text{ m}^2 \text{ g}^{-1}$) while preserving high P contents (Table S2 and Fig. S4, ESI†). Interestingly, the addition of the metal salt not only generated porosity but also changed the phosphorus environments, as will be explained in Section 2.3 (Fig. 3B). THPC and urea form a totally liquid solution at 60 °C when mixed with $\text{CaCl}_2 \cdot 4\text{H}_2\text{O}$, *i.e.* with a total water content of only 18 wt%, favouring the homogeneity of the final material (Fig. 1C). This efficient mixing was ascribed to a large number of hydrogen bonds between the H-bond donors (H_2O , urea) and acceptors (Ca^{2+} and phosphonium cation), in a manner that echoes the formation of deep-eutectic solvents.⁴² This not only assures the high homogeneity of the condensation process but also the association of Ca^{2+} with the precursors. Upon temperature increase, the polymerization of the constituents and the evaporation of water destabilize the system and trigger the recrystallization of the calcium chloride, thus forming an entangled system resin–metal chloride akin to forming porosities at high temperature. The recorded TGA–MS curve in the presence of $\text{CaCl}_2 \cdot 4\text{H}_2\text{O}$ (Fig. S5, ESI†) presents a maximal mass loss rate at similar temperatures to those in the absence of salt (137 °C and 282 °C, and 434 °C), with similar released fragments. This suggests that the polymerization step likely occurs following similar paths as for the neat THPC/urea mixture (Fig. 2B), *i.e.* through the condensation of the urea units and hydroxymethyl ones, despite the presence of salt. The foaming of the crude at 350 °C and beyond, due to the release of gases following material fragmentation, was drastically limited (Fig. 2C).

2.2 Influence of the CaCl_2 : THPC ratio on material structure

As the introduction of CaCl_2 successfully generated porosity in the system, we evaluated the impact of the ratio of CaCl_2 vs. the carbon precursors, which can be perceived here as the dilution factor in the reaction medium, to target the minimal amount of salt required. The samples are specified as TU-ST-R where S represents the metal of the salt hydrate, T the first digit of the final temperature and R the metal:THPC molar ratio. Firstly, X-ray diffraction (XRD) measurements of the crude samples, recorded immediately after synthesis and before washing, display the characteristic pattern of calcium chlorophosphate $\text{Ca}_2(\text{PO}_4)\text{Cl}$, although always as traces and without

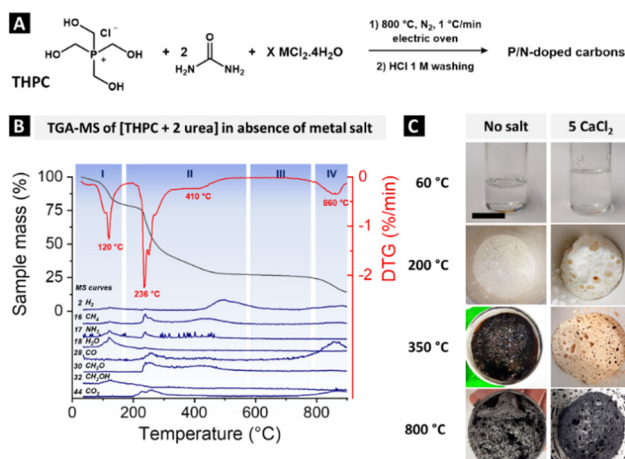


Fig. 2 Formation of P/N co-doped carbons from the THPC–urea condensate. (A) Reaction scheme. (B) TGA–MS curves of thermal treatment of [THPC + 2 urea] under helium (heating ramp of $2.5 \text{ }^{\circ}\text{C min}^{-1}$). Sample mass (black) and derivative (red), with corresponding MS curves (arbitrary units) at different m/z ratios (blue). Suggestions of fragments are given for MS. (C) Digital photographs of the materials synthesized in the presence or absence of $5 \text{ CaCl}_2 \cdot 4\text{H}_2\text{O}$ at various stages (scale bar is 1 cm).



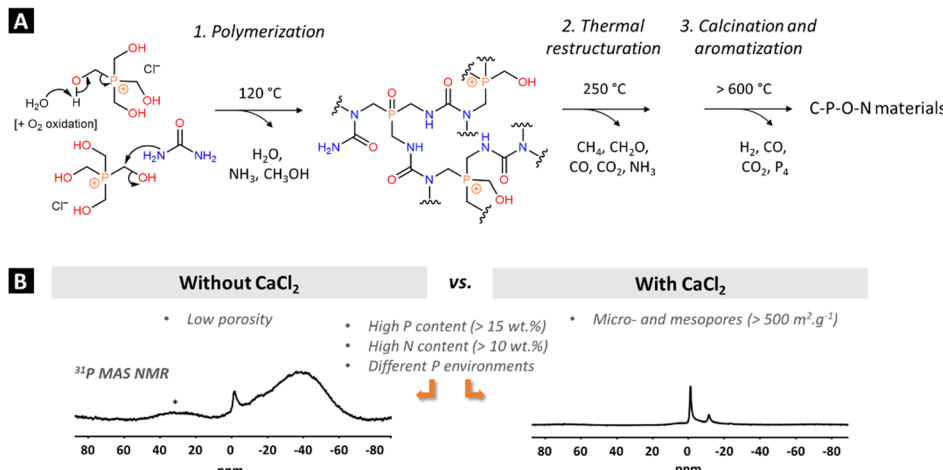


Fig. 3 (A) Mechanistic proposal for materials synthesis from THPC and urea in the absence and presence of calcium chloride. (B) Illustration of the ubiquitous use of THPC–urea precursor in the absence or in presence of a porogen agent with representative ³¹P solid-state NMR spectra (more details in Section 2.3).

Table 1 Composition and morphology characteristics of the samples

Sample	C yield%	P content ^a (wt%)	P yield%	S_{BET} ^b (m ² g ⁻¹)	V_{T} ^b (mL g ⁻¹)
TU-no salt-600	48	20 ^c	54	<5	<0.01
TU-Ca6-1	53	17.2	65	<5	<0.01
TU-Ca6-2	49	18.4	65	<5	<0.01
TU-Ca6-5	51	18.2	47	54	0.11
TU-Ca6-10	42	17.3	33	172	0.36
TU-no salt-800	60	19 ^c	45	70	0.03
TU-Ca8-1	59	18.4	61	50	0.03
TU-Ca8-2	67	16.3	48	570	0.48
TU-Ca8-3	70	17.3	50	606	0.51
TU-Ca8-5	59	18	42	603	0.73
TU-Ca8-10	50	14.7	28	536	0.72
TU-Ca8-20	41	13.2	20	347	0.55

^a Determined by ICP-OES with a typical error bar of 0.5 wt%. ^b Determined from the N₂ adsorption isotherm at 77 K for 0.03 < P/P₀ < 0.15 for S_{BET} , and at P/P₀ = 0.95 for total pore volume V_{T} (pores less than 40 nm in diameter). ^c Samples synthesized without salt could not be effectively digested for ICP analysis; the reported P contents correspond to the ones from EDX analysis.

further growth beyond CaCl₂:THPC ratio of 2:1 (Fig. S6, ESI[†]). A fraction of the P atoms is therefore undeniably oxidized to phosphate but the P yield only decreases from 61% to 42% for CaCl₂:THPC ratios increasing from 1:1 to 5:1, and thus stays high even in a large excess of CaCl₂. No hydroxyapatite Ca₁₀(PO₄)₆(OH)₂ was formed, and the excess of CaCl₂ was instead recovered as anhydrous CaCl₂ (Fig. S7, ESI[†]).

Upon increase of CaCl₂ content, the morphology observed by scanning electron microscopy (SEM) evolved from non-rough particles (>100 μm) to smaller aggregated flake-like structures (<1 μm) (Fig. 4A). This morphology reflects the dual role of CaCl₂ as a porogen agent and a reaction medium for the polymerization process, leading to polymeric, and then carbonaceous, particles. At the lowest ratios, the system corresponds to a dispersion of calcium chloride clusters in the resin material, ultimately resulting in the cracking of a bulk into chunks, rather than to a polymer embedded in a salt matrix. According to energy dispersive X-ray spectroscopy (EDX), all

samples contain large amounts of heteroatoms with N, O and P contents in the 9–15 wt%, 13–26 wt% and 13–20 wt% ranges, respectively (Fig. 4B). The overall composition was confirmed by XPS for TU-Ca8-5, with similar values. The phosphorus contents were also determined by inductively coupled plasma-optical emission spectroscopy (ICP-OES), presenting here the best accuracy, and appeared consistent with EDX: from 18.4 wt% for the 1:1 ratio to 13.2 wt% for 20:1 (Table 1).⁴³ The highest P contents reported in the literature are obtained via phosphorylation of a porous carbonaceous material with concentrated phosphoric acid and are *ca.* 12 wt%.¹ The unusually high phosphorus contents reached here are due to the use of THPC as a simultaneous C and P-source with a high P:C molar ratio of 1:4.

All samples calcined at 800 °C display similar XRD patterns, irrespective of the amount of CaCl₂ used, with two broad peaks at 24° and 43° (Fig. S8, ESI[†]). The shift toward low angles, as compared with typical peaks of turbostratic structures (2θ = 26°,



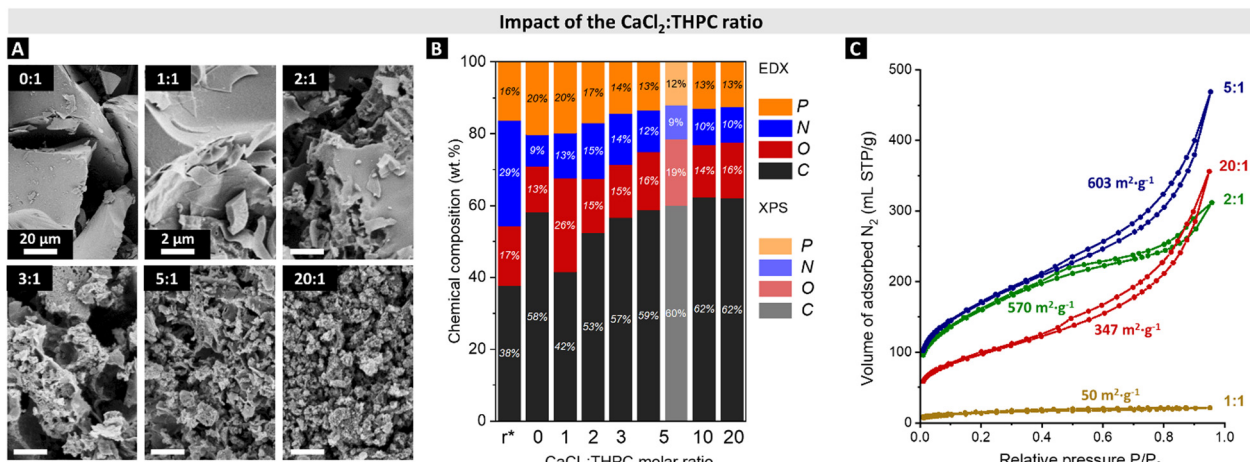


Fig. 4 (A) SEM images of the samples calcined at 800 °C with different CaCl₂:THPC ratios, indicated in the images (scale bar of 2 μm, except for TU-Ca8-0). (B) Chemical composition according to SEM-EDX, and XPS for TU-Ca8-5 (faded). The theoretical composition of the resin according to stoichiometry is indicated at r^* . (C) N₂ physisorption experiments performed at 77 K (adsorption and desorption).

$d = 3.35 \text{ \AA}$), is attributed to a local geometric distortion of the interlayer packing due to the high concentration of phosphorus. Fourier transform infrared spectroscopy (FTIR) measurements display numerous absorption bands in the 1600–800 cm⁻¹ region, typical for N and P doped carbonaceous materials and absent in non-doped carbon materials (*e.g.* pure graphite), even after calcination at 800 °C (Fig. S9, ESI[†]). This corroborates the presence of numerous N- and P-containing pendant functional groups and the absence of extensive graphitization (further discussion in the ESI[†]).

The porous structure of the synthesized materials was analysed by means of N₂ physisorption at 77 K (Fig. 4C). In the absence or at a low concentration of CaCl₂, the material has a low porosity with specific surface areas of 50–70 m² g⁻¹ (Table 1). Upon increasing the amount of salt, samples develop accessible micro- and mesopores, as indicated by the measured type IV sorption isotherms, with typical BET (Brunauer–Emmett–Teller) specific surface areas ranging from 300 to 600 m² g⁻¹ (Table 1C). The H3/H4 hysteresis loops suggest small “slit-like” mesopores or groove pores formed by flaky particles.⁴⁴ While the adsorption behaviour stays similar for molar ratios of 2:1 and 5:1 for the low pressure region ($P/P_0 < 0.4$), the volume of adsorbed N₂ decreases upon further dilution with salt at 20:1, indicating lower volume of micropores. This decrease may be due to the formation of a higher proportion of isolated polymer branches upon porogen content increase. These branches are not connected to the bulk of the material, or too thin to resist calcination. As such, they totally decompose or are lost upon washing without generating microporous regions, in coherence with the decreasing of the overall micro-porosity. The pore size distribution reveals essentially small mesopores below 8 nm in diameter for TU-Ca8-2, and a wider distribution up to more than 20 nm for TU-Ca8-5 and TU-Ca8-20 (Fig. S10, ESI[†]). The porosity mainly developed upon increase of the CaCl₂:THPC ratio from 1:1 ($S_{\text{BET}} = 50 \text{ m}^2 \text{ g}^{-1}$) to 2:1 ($S_{\text{BET}} = 570 \text{ m}^2 \text{ g}^{-1}$), without being associated with a

significant drop of carbon yield (from 61% to 48%), ruling out an etching mechanism as the main one. Instead, the calcium and chloride ions, initially miscible with the precursors, crystallize to clusters and nanoparticles at higher temperatures upon water evaporation, and thereby act as a hard template to create pores, as already discussed by Fechler *et al.* on the salt-melt templating effect.⁴¹ Accordingly, the increase of pore volume is concomitant with the recovery of CaCl₂ at the end of the reaction, on addition of Ca₂(PO₄)Cl. Visually, no macroscopic phase separation of the salt and the carbonaceous material is observed after the synthesis. Nonetheless, the melting point of CaCl₂ (772 °C) is in between the two studied calcination temperatures, which could explain part of the porosity development: the salt melts and favours the opening of pores while leaving the small pockets it was entrapped in, thus giving access to a higher microporosity (in the case of < 2 nm CaCl₂ clusters). Note the minimal molar ratio required to develop porosity (2:1) corresponds to a mass ratio between CaCl₂ and the two carbon precursors (urea + THPC) of 0.7:1, significantly lower than the values generally used for salt-melt templating, up to 10:1.³⁷

The sample TU-Ca8-5, presenting the highest porosity, was further examined by scanning transmission electron microscopy (STEM) (Fig. 5A–C). The material appears homogeneous without embedded nanoparticles of elemental phosphorus, contrary to Imamura *et al.*'s report of their presence upon thermal treatment of P-doped carbon at 1000 °C.³⁰ The presence of *ca.* 10–100 nm wide “holes”, originating from the formation of crystals (CaCl₂, Ca₂(PO₄)Cl) upon reaction, agrees with the porogenesis mechanism discussed earlier (Fig. 5A, orange arrow). The elemental mapping by using STEM-EDX reveals a homogeneous distribution of C, N, O and P in the carbonaceous material (Fig. 5A). At the atomic scale, high resolution annular dark-field (ADF)-STEM, where a brighter contrast refers to the heavier P atoms, nonetheless revealed a superior density of phosphorus at the edges of carbon ribbons



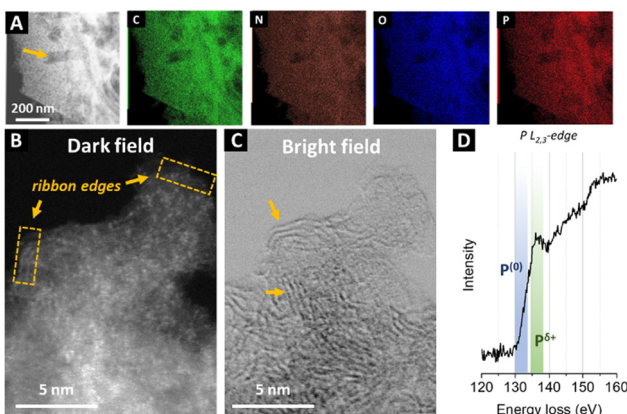


Fig. 5 Scanning transmission electron microscopy study of sample TU-Ca8-5. (A) Bright-field STEM image and the corresponding EDX elemental maps. (B) Annular dark-field and (C) bright field STEM image of the same region. (D) EELS spectra of P $L_{2,3}$ -edge.

with typical distance between two P atoms of 4 Å, *i.e.* separated by only 2–3 chemical bonds (Fig. 5B). Such a close vicinity of the P atoms is expected to exacerbate the reactivity of the surrounding carbon and nitrogen atoms by combining their electronic effects. Complementary to the elemental information in ADF-STEM, bright-field (BF)-STEM carries structural information, which show diffraction fringes, though not on extended domains (orange arrows) (Fig. 5C). The partly ordered structure (STEM limited here to 2D projection) is coherent with the large concentration of heteroatoms, preventing a long-range ordering. Such a structure is expected to lead to numerous defects

which may be as many catalytic active sites. The P $L_{2,3}$ -edge shape observed by EELS coincides with the P $L_{2,3}$ -edge recorded in X-ray absorption spectroscopy for phosphate-like structures.⁴⁵ Although the analysis of P structures by this means is not specific enough to distinguish $R_2(RO)PO$, $R(PO)_2PO$ and $(RO)_3PO$ groups, it unambiguously confirmed an oxidized state for P with the main feature at 138 eV (Fig. 5D), instead of 130 eV for elemental phosphorus.⁴⁶

2.3 Nature of the phosphorus-containing groups

^{31}P solid state-NMR experiments were run to investigate the nature of the phosphorus environments at the different stages of the synthesis. At first, a mix [THPC + 2 urea] without $CaCl_2$ was heated for 1 h at 200 °C, 350 °C, 600 °C or 800 °C. The ^{31}P MAS NMR spectrum of the resin at 200 °C displays mainly one broad peak centred at 30 ppm (A) with a shoulder at 45 ppm (B), attributed to R_4P^+ and R_3PO species, respectively (Fig. 6A and C). These attributions are in line with liquid state NMR spectra of the corresponding oligomers, tabulated at 29 ppm and 44 ppm, respectively (Table S3, ESI†).³³ Besides, $^{31}P\{-^1H\}$ HETCOR (heteronuclear correlation) experiments indicate a correlation of the ^{31}P signal at 30 ppm with a 1H peak at 4.5 ppm, typical of $P-CH_2-O/P-CH_2-N$ environments (Fig. S11A, ESI†).³³ At 350 °C, R_4P^+ and R_3PO species are still present, though shifted to 23 ppm and 50 ppm, respectively, and a broad peak C centred at –10 ppm appears. The simultaneous presence of aliphatic-like (2–4 ppm) and aromatic-like (8–10 ppm) protons already attests of the formation of conjugated structures at this temperature (Fig. S11B, ESI†). Upon

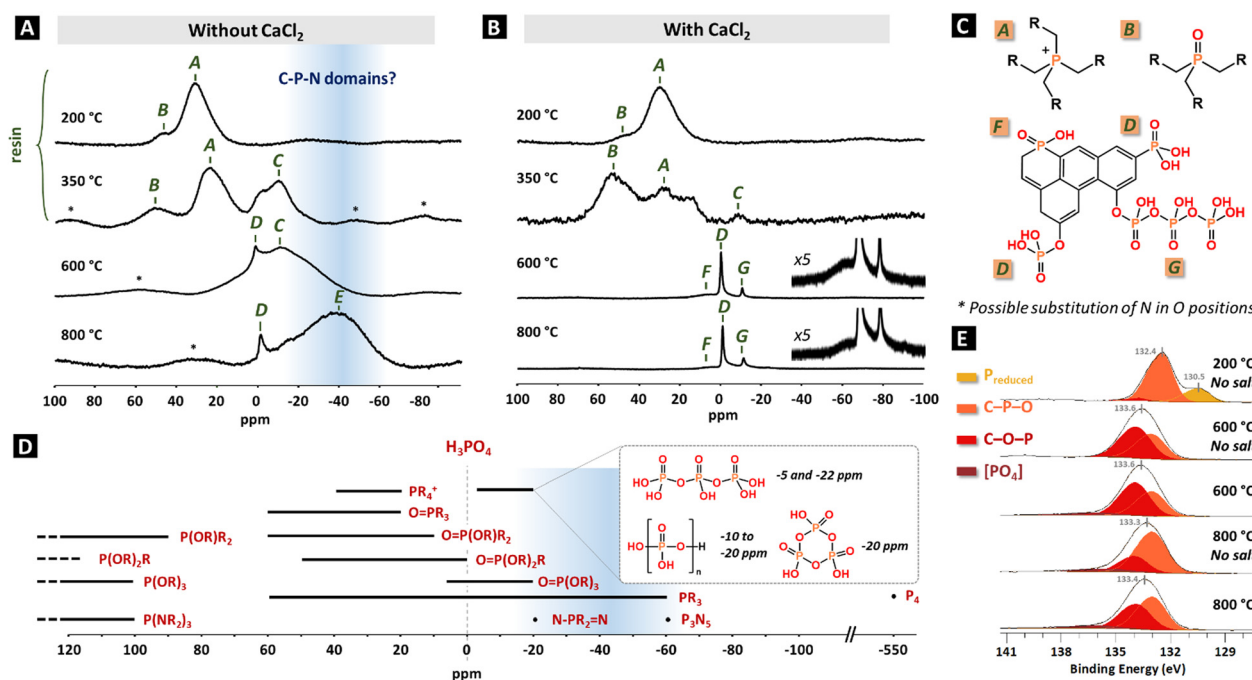


Fig. 6 Nature of the phosphorus-containing groups. ^{31}P MAS NMR spectra of materials synthesized in absence (A) and in the presence (B) of $CaCl_2$ salt (molar ratio of 5 : 1). Asterisks denote the presence of side bands. (C) Suggested attribution of several components. (D) Representative positions of P-containing groups in liquid-state ^{31}P NMR (except P_3N_5 and P_4 from solid state NMR). (E) X-ray photoelectron spectra (P 2p) of the resin and the carbonaceous materials, with proposed deconvolution.



thermal treatment at 600 °C and 800 °C, the phosphonium and phosphine oxides species completely vanished and a peak D appeared at *ca.* 0 ppm, ascribed to $(RO)_3PO/(RO)_2RPO$ groups (Fig. 6A and C). At the same time the peak C at –10 ppm significantly grew, dominating spectra at 600 °C, while the main one at 800 °C, denoted E and equally broad, is centred at –40 ppm. At 600 °C, ^{13}C CP MAS NMR spectrum mainly consists in “aromatic” carbons (130 ppm) indicating an important redistribution of the C–N and C–P bonds to create C–C/ C=C bonds with possible integration of the phosphorus atoms directly into the aromatic system (Fig. S12, ESI†). The presence of a contribution at 14 ppm for 1H is typical of P–OH groups, supposedly at the surface of the material as the sharpness of the peak suggests a certain mobility (Fig. S13, ESI†).

In the presence of $CaCl_2$, the spectrum is identical for the heating at 200 °C, confirming that the presence of salt does not drastically interfere with the polymerization mechanisms (Fig. 6B and C). At 350 °C, the proportion of phosphine oxides is higher, potentially due to a higher exposed surface. Upon thermal treatment at 600 °C and 800 °C, three main contributions are noted: a sharp phosphate peak D at –1 ppm, a broad one F at 5 ppm, attributed to $(RO)_2RPO$ units, and a small sharp one G at –10 ppm. This last one may correspond to inner phosphorus atoms in pyrophosphate/polyphosphates ($-P(=O)OH-O-$)_n, as observed in the case of activation of carbon fibres by H_3PO_4 .⁴⁷ Note that as the nitrogen content is high in the material, and the molar O/P ratio limited, the substitution of some of the O atoms by N ones is expected. However, ^{31}P NMR analysis does not allow the unambiguous distinction between equivalent groups bearing P–O and P–N bonds as they are tabulated in the same regions.⁴⁸ The heat treatment of carbon precursors (typically lignocellulosic sources) in the presence of phosphoric acid is the most widespread way to access P-doped carbon materials, with a maximum phosphorus content obtained following a calcination at 800 °C. Upon heating, both XPS and ^{31}P solid-state NMR data suggest an evolution of the C–O–P linkages into C–P–O ones, and eventually elemental phosphorus due to the carbothermal reduction of phosphate groups, explaining the decrease of the overall P content. In these cases, the reported NMR peaks correspond to the components F and D at respectively *ca.* 10 and 0 ppm.^{49,50}

Rationalizing ^{31}P NMR chemical shifts in P-doped carbonaceous materials is challenging due to the lack of referenced data in the literature (most of samples display PO_4 groups) and to the complex effects ruling the peak positions (*e.g.* electronic density surrounding the phosphorus, P valence). As a consequence, the predicted ranges for the different groups mostly overlap (Fig. 6D). Nonetheless, the chemical shifts for C and E (–10 and –40 ppm) in the ^{31}P NMR spectra are rather uncommon for P-doped carbons in the literature. The C contribution most likely does not correspond to a polyphosphate species due to the too low temperature for its first apparition (350 °C), its broadness and the absence of a larger peak at 0 ppm for the terminal phosphate group. Both C and E features are extremely broad, even though acquired at a high field (16.4 T). This is the

sign either of a large number of closed chemical environments, though not equivalent, or of a limited mobility of these chemical groups: both reasons point at phosphorus atoms buried in the core of the carbonaceous particles. They are therefore preserved from oxidation during acid washing and further storage in air. In contrast, the sharpness of the phosphate peak D suggests that the species are located at the surface. In liquid-state NMR, species with a trivalent phosphorus $P(OR)_x(NR_2)_yR_{z \leq 2}$ usually present a signal beyond 80 ppm and are thereby excluded to explain the nature of C and E components.⁴⁸ On the other side, pentavalent phosphorus presenting $-N-PR_2=N-$ bonding, for instance derived from tricyclopophosphazene condensation, are reported *ca.* –20 ppm,⁵¹ and phosphorus nitride (P_3N_5) also displays a down-shifted signal at –60 ppm in solid-state MAS NMR.⁵² Noticeably, Schmidpeter *et al.* reported that signals stemming from pentavalent phosphorus span from 45 ppm to –20 ppm for PO_4 environments, but are extended to –40 ppm for PN_4 ones.⁵³ Overall, the unattributed features likely correspond to a phosphorus connected to nitrogen atoms through single and/or double bonds, eventually forming C–P–N domains; or to a phosphine-like environment generated by the cleavage of a phosphonium ($P(CH_2OH)_3$) is reported at –24 ppm).⁵⁴ Although X-ray absorption spectroscopy is a common tool to investigate and confirm the chemical environments of specific atoms, the analysis of the K-edge of phosphorus is not expected here to bring further information than NMR considering the poor discrimination of phosphorus-containing compounds based on spectral features on one hand,⁴⁵ and the very close P–O and P–N distances (analysis of the extended X-ray absorption fine structure oscillations) on the other hand.

In addition to NMR, and in order to characterize the chemical states of the elements at the surface of the material, XPS was performed on the resin obtained at 200 °C without metal salt and on the carbonaceous materials synthesized at 600 °C and 800 °C, both in the absence and in the presence of $CaCl_2$ (Table S4, ESI† for fitting parameters and Fig. S14 (ESI†) for high resolution C 1s, N 1s, and O 1s XPS spectra). The analysis of the P 2p spectra reveals in the case of the resin (200 °C no salt in Fig. 6E), a first component at 132.4 eV, assigned to $R_3P=O$ units originating from the condensation of THPC followed by the loss of a substituent and an aerobic oxidation.⁵⁵ Another well-defined component is observed at lower binding energy (130.5 eV), in the region usually ascribed to reduced phosphorus, as in phosphines or elemental P. Here, this component may be assigned to phosphonium centres R_4P^+ considering the NMR analysis.^{55–57} Alternatively, it might correspond to surface localized phosphine species created upon beam irradiation on the R_4P^+ groups. The spectra of the calcined samples are very similar and comprise a large peak centred at *ca.* 133.5 eV, deconvoluted in an electron-poor component C–O–P and an electron-rich one O–C–P. These two components rather reflect a continuum of environments between phosphate ester $(RO)_3PO$ and phosphine oxide R_3PO rather than precise functional groups. The non-negligible O–C–P component indicates the partial incorporation of the



phosphorus atoms in the carbon matrix *via* C-P bonds. In addition, there is a minor contribution at 136.5 eV of fully oxidized phosphorus component [PO₄], assigned to (poly)phosphate. The strong similarities of the XPS spectra of all carbonaceous materials obtained at 600 °C and 800 °C, even though the ³¹P NMR revealed important differences, support that the surface phosphorus atoms are mostly oxidized (C–O–P/C–P–O type) and that the previously discussed reduced [C–P–N] and PR₃ species are either indiscernible by XPS or absent from the surface.

2.4 Carbo-catalyst for the oxidative dehydrogenation of N-heterocycles

One notable success of carbon materials as catalysts lies in their industrial use for oxidative dehydrogenation (ODH) of alkanes to alkenes, and of alcohols to aldehydes.^{58,59} The reactivity is generally attributed to carbonyl/quinone groups at their surface. Interestingly, an improved selectivity was reported after insertion of phosphorus, as phosphate groups, in the material.⁶⁰ A similar ODH reaction starting from N-heterocycles generates N-heteroarenes, essential building blocks in fine chemistry. N-doped carbo-catalysts in particular proved to efficiently activate O₂ as primary oxidant for the ODH of indole and quinoline derivatives.^{25,26,28,29,61} On the basis of these works, we evaluated the performance of the synthesized P/N co-doped carbonaceous material for the oxidation of indole to indole, as a model reaction. As these reactions are usually run on N-heterocycles bearing other reactive groups, we targeted mild conditions in terms of temperature (<100 °C) and oxidant (air). Considering the catalytic context, a sample with a large exposed area is required, we therefore chose the TU-Ca8-5 one, presenting the optimal balance of high porosity and high P and N contents, *ca.* 18 and 10 wt%, respectively. Note that this sample was prepared in the presence of CaCl₂, leading to phosphonic acid-like surface groups.

We first evaluated the influence of the solvent at room temperature in air with a catalyst loading of 100 wt%, with respect to the indole reagent (Table 2, entries 1–8). The polarity of the solvent did not correlate with the conversion, suggesting that the limiting step of the reaction does not involve ionic or polarized species (Fig. S15A, ESI†). The best conversion (93%) was obtained in a 1:1 volume mixture of H₂O/EtOH. As this nontoxic solvent solubilized the indole and efficiently dispersed the catalyst, it was used in the following experiments. A blank reaction without a catalyst and the use of various metal-free materials such as carbon black, graphite or carbon nitride led to a null conversion, revealing the catalytic activity is indeed due to specific heteroatom-containing surface groups (entries 9–12). The absence of a reaction in presence of phosphoric acid indicates that the sole Brønsted acid groups present at the catalyst surface, if involved in the mechanism at all, are not sufficient for the dehydrogenation (entry 13).

Replacing the air by pure O₂ afforded a quantitative conversion at 100 wt% at r.t. (entries 14–16). In many cases and for less active systems, carbo-catalysis requests rather high catalyst loadings.² In the present case, a 91% conversion was obtained

Table 2 Optimization of the reaction conditions for the oxidative dehydrogenation of indole by the P/N co-doped carbo-catalyst

Entry	Catalyst	Solvent	Loading (wt%)	T (°C)	
				Conv. (%)	
1 ^a	P/N carbon	H ₂ O	100	20	64
2	P/N carbon	MeOH	100	20	46
3	P/N carbon	EtOH	100	20	51
4	P/N carbon	ACN	100	20	58
5	P/N carbon	EtOAc	100	20	62
6	P/N carbon	Dioxane	100	20	22
7	P/N carbon	Toluene	100	20	75
8	P/N carbon	H ₂ O/EtOH	100	20	93
9	—	H ₂ O/EtOH	—	20	0
10	Carbon black	H ₂ O/EtOH	100	20	0
11	Graphite	H ₂ O/EtOH	100	20	0
12	C ₃ N ₄	H ₂ O/EtOH	100	20	0
13 ^b	H ₃ PO ₄ (50 mol%)	H ₂ O/EtOH	—	20	0
14 ^c	P/N carbon	H ₂ O/EtOH	25	20	44
15 ^c	P/N carbon	H ₂ O/EtOH	50	20	73
16 ^c	P/N carbon	H ₂ O/EtOH	100	20	>98
17 ^d	P/N carbon	H ₂ O/EtOH	10	80	56
18 ^d	P/N carbon	H ₂ O/EtOH	25	80	91
19 ^d	P/N carbon	H ₂ O/EtOH	50	80	>98
20 ^e	P/N carbon	H ₂ O/EtOH	100	80	>98
21 ^f	NPCH	H ₂ O	21	120	91
22 ^{gc}	NCC-800	H ₂ O	67	80	>99
23 ^h	N-HPC-800	t-Amyl alcohol	134	110	87
24 ^{ic}	NC(242-800)	EtOH	67	60	96

Reaction conditions: indole (28 μ L, 30 mg, 0.25 mmol), solvent (2.5 mL), catalyst (30 mg), 1 bar, 18 h. Conversions determined by ¹H NMR.

^a Limited solubility of the indole in water. ^b Similar P content to that introduced with 30 mg of P/N carbon. ^c Reaction performed in O₂.

^d Reaction conditions: indole (112 μ L, 120 mg, 1 mmol), solvent (10 mL), catalyst (12/30/60 mg). ^e Gram-scale experiment: indole (1.12 mL, 1.2 g, 10 mmol), solvent (100 mL), catalyst (1200 mg), air, reflux, 18 h. ^f Beller work.²⁵ ^g Liu work.²⁶ ^h Dai work.²⁸ ⁱ Quan work.²⁹

in air with only 25 wt% loading upon increase of the reaction temperature to 80 °C (entries 17–19, and Fig. S15B, ESI†). A catalyst loading of 10 wt%, equivalent to 58 μ mol of P per mmol of indole, led to a conversion of 56%, which definitely proves the catalytic effect of the carbon materials (equivalent turnover number *vs.* P atoms TON > 9.6). The upscaling of the reaction to 10 mmol (equiv. to 1.19 g of indole) proved to be successful (>98% conversion, entry 20) and the recovered catalyst was still active after ethanol washing and drying. The conversion decreased over the first recycling tests before stabilizing *ca.* 60% (Fig. S16, ESI†). It is noteworthy, the developed catalyst presents at least equivalent performances to the currently best performing metal-free N-doped carbo-catalysts (entries 21–24).²⁵

We then investigated the substrate scope for dehydrogenation in a range of substituted indolines and tetrahydroquinolines (Fig. 7 and Fig. S18–31, ESI†). High conversions were obtained in the presence of different groups in the 5-position of the indoline (–methyl, –methoxy, –Br, and –NO₂), though slightly lower in the case of the strong electron-attractor nitro group. The only substrate obtained with a low yield (6%) was

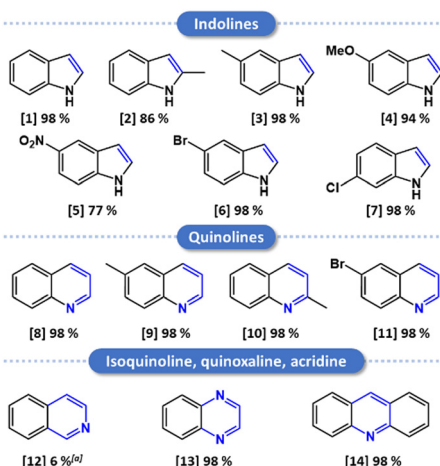


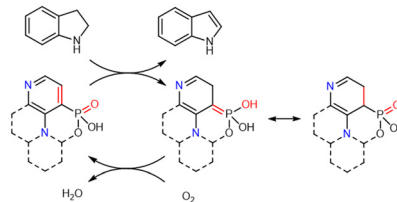
Fig. 7 Substrate scope. Reaction conditions: substrate (0.25 mmol), $\text{H}_2\text{O}/\text{EtOH}$ (1:1) (2.5 mL), catalyst (30 mg), air, 80 °C, 18 h. Yields determined by ^1H NMR. (a) Total conversion of the tetrahydroisoquinoline to 6% of isoquinoline and 94% of 3,4-dihydroisoquinoline.

isoquinoline, for which mainly the half-dehydrogenated product was obtained, a limitation also reported by Mata *et al.*²⁹ A total conversion was recorded for both the quinoxaline and the acridine, in line with previous works.^{26,28} All of these results point toward the possible use of the developed material for aerobic ODH of N-heterocycles in N-heteroarenes under mild conditions.

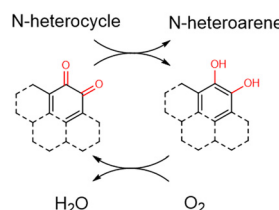
The mechanism of the ODH was assessed by control experiments changing the gas phase, or in the presence of radical scavengers (Table 3). Run in N_2 , the reaction yield falls from 73% to 12%, proving the role of O_2 as the primary oxidant (entries 1 and 2). The residual conversion is non-negligible and suggests that some functional groups at the surface of the catalyst are directly responsible for the dehydrogenation. Therefore, the molecular oxygen is thought to regenerate the oxidative groups consumed upon reaction, thus closing a catalytic loop.²³ The limited variation of yield in the presence of 2 equivalents of BHT (67%), TEMPO (80%) or $^1\text{PrOH}$ (61%) is

• Phosphorus-containing carbon materials (this work)

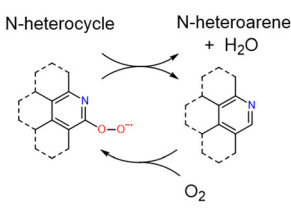
- ✓ reactivity maintained in presence of radical scavengers
- ✓ not affected by solvent polarity
- ✓ poor activity in absence of P–C bond



• Air-oxidized carbons



• N-doped carbons



Scheme 1 Proposed mechanism for N-heterocycle dehydrogenation using a P/N-doped carbonaceous material.

within the reproducibility of the experiments, indicating the main pathway does not imply free radical species (entries 3–5). Such a result is in line with the works of Beller *et al.* (N/P-doped carbon)²⁵ and Dai *et al.* (N-doped carbon),²⁸ but contrasts with the one of Liu *et al.* (N-doped carbon).²⁶ Finally, the absence of deep brown coloration, characteristic of the triiodide ion I_3^- , upon the addition of a drop of concentrated NaI, ruled out the generation of a molecular oxidant, such as H_2O_2 , as a reaction intermediate (Fig. S17, ESI†).

Recently, Patel *et al.* reported the ODH of benzyl alcohol to benzaldehyde catalysed by P-doped carbons and suggested a shuttle of the active site from $\text{P}^{(\text{III})}$ to $\text{P}^{(\text{V})}$.²³ Such a mechanism, however, involves the formation of an alkoxide directly linked to the phosphorus atom, and can less easily be transposed to the ODH of N-heterocycles. Instead, the classical mechanism proposed for ODH on air-oxidized carbon materials consists in the transfer hydrogenation of a surface quinone group (two adjacent $\text{C}=\text{O}$) to a diphenol, continuously regenerated through oxidation by O_2 (Scheme 1).^{59,60} In our case, the number of $\text{C}=\text{O}$ groups is limited as no absorption band is detected at *ca.* 1600 cm^{-1} by FTIR, most of the oxygen atoms are most likely linked to phosphorus atoms. However, as the material exhibits P–C bonds at high concentration, we suggest a modified version of the classical transfer hydrogenation step leading to an ylide structure, susceptible to rearranging rapidly to phosphonic acid ($\text{RO}(\text{RP}(\text{O})=\text{O})\text{OH}$) (Scheme 1). The activation of O_2 on another part of the material, followed by its diffusion to the so-formed unsaturated site then allows oxidation back to an aromatic structure, energetically more favourable, and generates a water molecule. The insertion of N in the aromatic structure may further favour the activation of O_2 at the surface of the material and facilitates the re-oxidation process of the P-containing groups to their resting state, as suggested by the

Table 3 Radical inhibition and control experiments

Entry	Carbo-catalyst	Additive (2 equiv.)	Gas	T (°C)	t (h)	Conv. (%)
1	P/N-doped	—	Air	20	8	73
2 ^a	P/N-doped	—	N_2	20	8	12
3	P/N-doped	BHT	Air	20	8	67
4	P/N-doped	TEMPO	Air	20	8	80
5	P/N-doped	$^1\text{PrOH}$	Air	20	8	61
6	P/N-doped	—	Air	20	16	93
7	CB-air	—	Air	20	16	6
8	CB-air	—	Air	80	16	92
9	N-doped	—	Air	20	16	46
10	P-doped	—	Air	20	16	75
11	H_3PO_4 -carbon	—	Air	20	16	9

Reaction conditions: indoline (28 μL , 30 mg, 0.25 mmol), $\text{H}_2\text{O}/\text{EtOH}$ (1:1) (2.5 mL), catalyst (30 mg), 1 bar gas. Conversions determined by ^1H NMR. BHT = butylated hydroxytoluene, TEMPO = (2,2,6,6-tetramethylpiperidin-1-yl)oxidanyl. ^a The solvent was degassed before-hand by N_2 bubbling.



good activity of N-doped carbon materials for oxygen reduction reaction (Scheme 1).^{4,13} The simultaneous presence of P and N elements in the matrix therefore allows a higher number of catalytically active sites as both are supposedly active (see below), and a more efficient catalytic loop as to the P sites thanks to the activation of O₂ as superoxide by the N sites in the vicinity. The progressive deactivation of the catalyst may be due to the leaching or hydrolysis in the phosphate groups of some of the depicted active P-sites.

The hypothesis on the mechanism was tested *via* different model materials (see Fig. S32–S35, ESI† for materials syntheses and characterization studies). At first, in our hands, an air-oxidized carbon black sample, a well-known carbo-catalyst for ODH,⁶² led to a 6% conversion at r.t. (Table 3, entries 9 and 10), pointing at the increased activity upon P-doping. Then, a highly N-doped carbonaceous sample (N content of 24 wt%) also proved to be active, but with a significantly lower conversion of 46% compared with P/N-doped materials, suggesting that most of the activity of this last one indeed originates from the phosphorus doping. Besides, a P-doped sample obtained from phytic acid calcination (similar to Patel's work²³) also displayed a non-negligible activity for ODH at r.t. (Table 3, entry 7), confirming that the sole P-doping is sufficient. However, a carbon phosphorylated by reacting phosphoric acid at low temperature (180 °C) was poorly active, even though containing 3.9 wt% of P (Table 3, entry 8). Such a sample supposedly contains only P–O bonds as no reduction of the phosphate groups occurs in this temperature range. This suggests that a conjugation of P with the aromatic system is required, likely to form the P=C bond of the intermediate species, in line with the absence of conversion with H₃PO₄ (Table 2, entry 13). Overall, the ODH of N-heterocycles catalysed by P-doped carbo-catalyst apparently proceeds *via* a mechanism differing both from the reported ones for surface oxidized carbons and N-doped carbons, while achieving higher conversions under equivalent conditions.

3. Conclusions and outlook

We demonstrated that the use of a tetrakis(hydroxymethyl)-phosphonium salt as a P precursor for carbonaceous materials syntheses, in combination with urea, opens the way to high phosphorus doping contents up to 18 wt%. To introduce porosity, a salt-templating strategy, based on the direct polymerization of the precursors in a calcium chloride hydrate, was successfully employed. This work thus expands the already explored condensation reactions of ionic liquids, carbohydrates and nucleobases in molten salts/salt hydrates to lower the temperature of polymerization of formaldehyde-like compounds (<120 °C). Moreover, unusual P-containing functional groups were reported for the calcined samples, beyond the commonly reported phosphate/phosphonate groups. These structures likely correspond to trivalent phosphine environments or pentavalent ones of type C–P–N. We believe that exploring such a chemistry would in particular be beneficial

to tune the electronic configuration of bulk carbon materials. The samples with the highest P contents proved to be efficient carbo-catalysts for the oxidative dehydrogenation of N-heterocycles to N-heteroarenes under mild conditions, likely following a transfer hydrogenation mechanism.

4. Experimental procedures

Chemicals

All chemicals were used as received without further purification. Tetrakis(hydroxymethyl)phosphonium chloride (THPC) 80 wt% in water, urea (99%), MgCl₂·6H₂O (99%), CaCl₂·2H₂O (99%), ZnCl₂ (98%), HCl (37%), indoline (98%) and tetrahydroquinoline (98%) were purchased from Merck. All the other N-heterocycles were purchased from A2B Chem (>98%). Note that CaCl₂·4H₂O and ZnCl₂·4H₂O were obtained *in situ* by mixing CaCl₂·2H₂O and ZnCl₂ with the corresponding amounts of water.

Materials synthesis

Phosphorus-doped carbonaceous materials were prepared typically by mixing CaCl₂·2H₂O (3.67 g, 25 mmol, 5 equiv.), urea (600 mg, 10 mmol, 2 equiv.), water (900 μL, 50 mmol, cautious: exothermic), and finally THPC 80% (889 μL, 1.191 g, 5 mmol, 1 equiv.) in a ceramic crucible. The precursor-salt mixture was thoroughly mixed with a spatula, covered with a lid, transferred to a muffle furnace and subjected to heat treatment under constant N₂ flow with a heating ramp of 1 °C min⁻¹ (the oven was first purged for 1 h with N₂). The final temperature was kept for one hour, before cooling down to room temperature. The black samples were then ground, washed three times with 400 mL of 1 M HCl (8 h, r.t.) and once with 400 mL of water before being dried at 60 °C overnight to yield a black powder (360 mg, equiv. to 59% C yield and 42% P yield).

Materials characterization

ICP measurements were performed using a PerkinElmer ICP-OES Optima 8000 after digestion of the dried samples in 3 mL H₂SO₄ 95% + 3 mL H₂O₂ 50% for 12 h at 80 °C. Standard operating conditions were used (Plasma power 1300 W, plasma flux 12 L min⁻¹, nebulizer gas flux 0.55 L min⁻¹, auxiliary gas flux 0.2 L min⁻¹, sample uptake 1.7 mL min⁻¹, Ar purging of the optical part of the instrument). Fourier transformed infrared (FTIR) spectroscopy was performed using a Thermo Scientific Nicolet iS5 FT-IR spectrometer, using an attenuated total reflection iD5 accessory. Powder XRD measurements were performed on a Rigaku SmartLab diffractometer, using Cu K α radiation at 1.5406 Å, with steps of 0.1° (0.03° for samples containing CaCl₂) and a scanning rate of 0.5° min⁻¹. Physisorption measurements were performed on a Quantachrome Quadrasorb SI apparatus with N₂ adsorption at 77 K after degassing at 150 °C for 15 h. The specific surface area (S_{BET}) of each material was obtained from the N₂ adsorption data (P/P₀ < 0.2) using the Brunauer–Emmett–Teller (BET) method. The total pore volume was calculated from the amount of nitrogen



adsorbed at $P/P_0 = 0.95$ (pores of diameter less than 40 nm). The N_2 pore size distribution was obtained using a Quenched Solid Density Functional Theory (QSDFT) model with a slit/cylindrical pore shape using the N_2 adsorption branch. SEM was performed on a LEO 1550 Gemini Zeiss microscope at 5 keV beam energy after sputtering of 5 nm of carbon on the top of the samples. EDX was performed using an Oxford Instruments EDX analyser. ^{31}P MAS, ^{13}C CP MAS, 2D HETCOR $^{13}C-\{^1H\}$ and $^{31}P-\{^1H\}$ CP MAS solid-state NMR spectra were recorded on a Bruker Avance 700 spectrometer (field = 16.4 T, $\nu_0(^1H)$ = 300.39 MHz, $\nu_0(^{13}C)$ = 176.08 MHz, $\nu_0(^{31}P)$ = 283.29 MHz) using a 3.2 mm Bruker MAS probe spinning at 20 kHz and on a Bruker Avance 300 spectrometer (field = 7.0 T, $\nu_0(^1H)$ = 700.14 MHz, $\nu_0(^{13}C)$ = 75.51 MHz, $\nu_0(^{31}P)$ = 121.56 MHz) using a 4 mm Bruker MAS probe spinning at 14 kHz. CP MAS experiments were recorded with ramped-amplitude cross-polarization in the 1H channel to transfer magnetization from 1H to ^{13}C or ^{31}P (recycle delay = 3 s, CP contact time = 1 ms, optimized 1H spinal-64 decoupling). Single pulse ^{31}P MAS NMR spectra were recorded with a recycle delay of 30 s. Chemical shift values were referenced to tetramethylsilane (TMS) for ^{13}C and H_3PO_4 (85 wt%) for ^{31}P . XPS measurements were performed using a Thermo Fisher Scientific K-Alpha, and data were treated using CasaXPS software. Monochromatic X-ray radiation of 1486.6 eV (Al $K\alpha$) was used to analyse each material. The powders were pressed on a carbon tape deposited on the sample holder in order to have a homogeneous layer. Survey scans were obtained using a pass energy of 200 eV and step size of 1 eV, with a spot size of approximately 100 μm . High resolution spectra of C 1s (280–298 eV), N 1s (394–404 eV), O 1s (526–545 eV) and P 2p (125–144 eV) were recorded using a pass energy of 50 eV and a step size of 0.1 eV. The binding energy was calibrated against the 284.8 eV peak of adventitious carbon (C-C).⁶³ Relative sensitivity factors (RSF) of 1, 1.8, 2.93 and 1.19 were used for C, N, O and P, respectively. Unless otherwise stated, a U3 Tougaard background was used and peaks were deconvoluted using mixed Gauss–Lorentz sum function line shape GL(30). STEM was performed using a double-Cs corrected JEOL JEM-ARM200F, operated at 80 kV using a probe semi convergence angle of 23 mrad, equipped with a cold-field emission gun, an energy dispersive X-ray detector and a Gatan Imaging Filter (GIF) Quantum. Samples were prepared on standard TEM Cu grids with lacey carbon support by dispersing particles in ethanol, drop-casting the solution on the grid and letting it dry under ambient conditions. ADF-STEM images were recorded within an angular collection semi-angle range from 50 mrad to 180 mrad. BF-STEM images were taken from scattering signal to a maximal collection semi-angle of 3.8 mrad. EELS data were acquired using a dispersion of 0.1 eV per channel. All data were acquired and analysed using Gatan's microscopy suite (GMS) software version 3.4.

Catalytic dehydrogenation

The reactions at 80 °C were conducted in a round-bottom flask connected to a condenser and heated in an oil bath, while the

reactions at r.t. were conducted in 5 mL glass vials. In a typical reaction, the catalyst (30 mg) was dispersed in the solvent (2.5 mL) by brief sonication, followed by the addition of the indoline (0.25 mmol, 28 μL). The reaction is allowed to react under stirring at the given temperature for 18 h and cooled down naturally to r.t. To avoid solvent evaporation, the vials of the reactions run at r.t. were closed with a septum and connected to an air balloon of *ca.* 1 L in volume to provide enough molecular oxygen (8 mmol of O_2). After the reaction, an aliquot was diluted in $CDCl_3$ and directly analysed by 1H NMR (400 MHz, 298 K). For the recycling test, the catalyst was recovered by centrifugation, washed with 100 mL of EtOH and dried at 60 °C overnight before the next cycle.

Author contributions

Rémi André: conceptualization, investigation, writing – original draft. Christel Gervais: investigation (ss-NMR), writing – review & editing. Hannes Zschiesche: investigation (STEM), writing – review & editing. Teodor Jianu: investigation (STEM). Nieves López-Salas: writing – review & editing. Markus Antonietti: supervision, writing – review & editing, funding acquisition. Mateusz Odziomek: supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The Max Planck Society is gratefully acknowledged for financial support, as well as the Deutsche Forschungsgemeinschaft (DFG) for the “Germany’s Excellence Strategy-EXC 2008-390540038-UniSysCat” funding and the French Région Ile de France – SESAME program (700 MHz NMR spectrometer) funding. N. Tarakina is acknowledged for the fruitful discussions, J. Brandt for ICP measurements, A. Völkel for TGA-MS measurements, Johannes Schmidt for XPS measurements, and B. Badamdorj and H. Runge for SEM-EDX measurements. Open Access funding provided by the Max Planck Society.

References

- 1 A. M. Puziy, O. I. Poddubnaya, B. Gawdzik and J. M. D. Tascón, *Carbon*, 2020, **157**, 796–846.
- 2 M. Antonietti, N. Lopez-Salas and A. Primo, *Adv. Mater.*, 2019, **31**, e1805719.
- 3 L. H. Zhang, Y. Shi, Y. Wang and N. R. Shiju, *Adv. Sci.*, 2020, **7**, 1902126.
- 4 C. Hu and L. Dai, *Adv. Mater.*, 2019, **31**, e1804672.
- 5 X. Rao, S. Zhang and J. Zhang, *Curr. Opin. Electrochem.*, 2023, **42**, 101416.
- 6 J. P. Paraknowitsch and A. Thomas, *Energy Environ. Sci.*, 2013, **6**, 2839–2855.
- 7 Y. Yang, D.-M. Tang, C. Zhang, Y. Zhang, Q. Liang, S. Chen, Q. Weng, M. Zhou, Y. Xue, J. Liu, J. Wu, Q. H. Cui, C. Lian,



G. Hou, F. Yuan, Y. Bando, D. Golberg and X. Wang, *Energy Environ. Sci.*, 2017, **10**, 979–986.

8 H. Wang, Y. Shao, S. Mei, Y. Lu, M. Zhang, J. K. Sun, K. Matyjaszewski, M. Antonietti and J. Yuan, *Chem. Rev.*, 2020, **120**, 9363–9419.

9 L. Zhang, X. Li and M. Antonietti, *Angew. Chem.*, 2021, **60**, 24257–24265.

10 N. Yang, X. Zheng, L. Li, J. Li and Z. Wei, *J. Phys. Chem. C*, 2017, **121**, 19321–19328.

11 D. S. Yang, D. Bhattacharjya, S. Inamdar, J. Park and J. S. Yu, *J. Am. Chem. Soc.*, 2012, **134**, 16127–16130.

12 D. Yu, Y. Xue and L. Dai, *J. Phys. Chem. Lett.*, 2012, **3**, 2863–2870.

13 J. Zhang, Z. Zhao, Z. Xia and L. Dai, *Nat. Nanotechnol.*, 2015, **10**, 444–452.

14 T. Najam, S. S. A. Shah, W. Ding, J. Jiang, L. Jia, W. Yao, L. Li and Z. Wei, *Angew. Chem.*, 2018, **130**, 15321–15326.

15 C. Chen, X. Sun, X. Yan, Y. Wu, H. Liu, Q. Zhu, B. B. A. Bediako and B. Han, *Angew. Chem.*, 2020, **59**, 11123–11129.

16 C. H. Choi, S. H. Park and S. I. Woo, *J. Mater. Chem.*, 2012, **22**, 12107–12115.

17 L. Zhang, H. Zhang, K. Liu, J. Hou, B. Badamdjorj, N. V. Tarakina, M. Wang, Q. Wang, X. Wang and M. Antonietti, *Adv. Mater.*, 2023, **35**, e2209310.

18 R. T. Mayes, P. F. Fulvio, Z. Ma and S. Dai, *Phys. Chem. Chem. Phys.*, 2011, **13**, 2492–2494.

19 L. Wang, X. Dong, H. Jiang, G. Li and M. Zhang, *Catal. Commun.*, 2014, **56**, 164–167.

20 K. M. Eblagon, A. Arenillas, A. Malaika, M. Fernando, R. Pereira and J. L. Figueiredo, *Fuel*, 2023, **334**, 126610.

21 M. J. Valero-Romero, E. M. Calvo-Muñoz, R. Ruiz-Rosas, J. Rodríguez-Mirasol and T. Cordero, *Ind. Eng. Chem. Res.*, 2019, **58**, 4042–4053.

22 J. Bedia, J. M. Rosas, J. Márquez, J. Rodríguez-Mirasol and T. Cordero, *Carbon*, 2009, **47**, 286–294.

23 M. A. Patel, F. Luo, M. R. Khoshi, E. Rabie, Q. Zhang, C. R. Flach, R. Mendelsohn, E. Garfunkel, M. Szostak and H. He, *ACS Nano*, 2016, **10**, 2305–2315.

24 X. Hu, M. Fan, Y. Zhu, Q. Zhu, Q. Song and Z. Dong, *Green Chem.*, 2019, **21**, 5274–5283.

25 K. Sun, H. Shan, R. Ma, P. Wang, H. Neumann, G. P. Lu and M. Beller, *Chem. Sci.*, 2022, **13**, 6865–6872.

26 X. Li, Z. Yuan, Y. Liu, H. Yang, J. Nie, G. Wang and B. Liu, *ChemSusChem*, 2022, **15**, e202200753.

27 J. Zhang, X. Liu, R. Blume, A. Zhang, R. Schlogl and D. S. Su, *Science*, 2008, **322**, 73–77.

28 S. Shang, Y. Li, Y. Lv and W. Dai, *Asian J. Org. Chem.*, 2022, **11**.

29 A. Mollar-Cuni, D. Ventura-Espinosa, S. Martin, H. Garcia and J. A. Mata, *ACS Catal.*, 2021, **11**, 14688–14693.

30 R. Imamura, K. Matsui, S. Takeda, J. Ozaki and A. Oya, *Carbon*, 1999, **37**, 261–267.

31 D. V. Moiseev and B. R. James, *Phosphorus, Sulfur Silicon Relat. Elem.*, 2019, **195**, 263–279.

32 D. V. Moiseev and B. R. James, *Phosphorus, Sulfur Silicon Relat. Elem.*, 2020, **195**, 687–712.

33 A. Granzow, *J. Am. Chem. Soc.*, 1977, **99**, 2648–2652.

34 A. Basch, B. Zvilichovsky, B. Hirshmann and M. Lewin, *J. Polym. Sci., Polym. Chem. Ed.*, 1979, **17**, 27–37.

35 W. A. Reeves and J. D. Guthrie, *Ind. Eng. Chem.*, 1956, **48**, 64–67.

36 Global Tetrakis (Hydroxymethyl) Phosphonium Sulfate Market Research Report 2022, Market Reports World.

37 X. Liu, N. Fechler and M. Antonietti, *Chem. Soc. Rev.*, 2013, **42**, 8237–8265.

38 N. Díez, A. B. Fuertes and M. Sevilla, *Energy Storage Mater.*, 2021, **38**, 50–69.

39 D. V. Moiseev and B. R. James, *Inorg. Chim. Acta*, 2011, **379**, 23–27.

40 J. Pampel, A. Mehmood, M. Antonietti and T. P. Fellinger, *Mater. Horiz.*, 2017, **4**, 493–501.

41 N. Fechler, T. P. Fellinger and M. Antonietti, *Adv. Mater.*, 2013, **25**, 75–79.

42 E. L. Smith, A. P. Abbott and K. S. Ryder, *Chem. Rev.*, 2014, **114**, 11060–11082.

43 R. F. André, J. Brandt, J. Schmidt, N. López-Salas, M. Odziomek and M. Antonietti, *Carbon*, 2024, **223**, 118946.

44 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. W. Sing, *Pure Appl. Chem.*, 2015, **87**, 1051–1069.

45 J. Kruse, P. Leinweber, K. U. Eckhardt, F. Godlinski, Y. Hu and L. Zuin, *J. Synchrotron Radiat.*, 2009, **16**, 247–259.

46 A. Favron, E. Gaufres, F. Fossard, A. L. Phaneuf-L'Heureux, N. Y. Tang, P. L. Levesque, A. Loiseau, R. Leonelli, S. Francoeur and R. Martel, *Nat. Mater.*, 2015, **14**, 826–832.

47 R. Fu, L. Liu, W. Huang and P. Sun, *J. Appl. Polym. Sci.*, 2003, **87**, 2253–2261.

48 Handbook of Phosphorus-31 Nuclear Magnetic Resonance Data (1990), 2017.

49 A. M. Puziy, O. I. Poddubnaya, R. P. Socha, J. Gurgul and M. Wisniewski, *Carbon*, 2008, **46**, 2113–2123.

50 Y. Wang, S. Zuo, J. Yang and S. H. Yoon, *Langmuir*, 2017, **33**, 3112–3122.

51 W. Zhang, J. Barrio, C. Gervais, A. Kocjan, A. Yu, X. Wang and M. Shalom, *Angew. Chem.*, 2018, **57**, 9764–9769.

52 W. Schnick, J. Lücke and F. Krumeich, *Chem. Mater.*, 1996, **8**, 281–286.

53 A. Schmidpeter and K. Schumann, *Z. Naturforsch., B: Chem. Sci.*, 1970, **25**, 1364–1370.

54 J. M. Mateo, A. de la Hoz, L. Uson, M. Arruebo, V. Sebastian and M. V. Gomez, *Nanoscale Adv.*, 2020, **2**, 3954–3962.

55 G. Biagiotti, V. Lange, C. Ligi, S. Caporali, M. Muniz-Miranda, A. Flis, K. M. Pietrusiewicz, G. Ghini, A. Brandi and S. Cicchi, *Beilstein J. Nanotechnol.*, 2017, **8**, 485–493.

56 J. Chastain and R. C. KingJr, *PerkinElmer Corp.*, 1992, **40**, 221.

57 F. G. d A. Dias, L. S. G. Souza, A. G. Veiga, A. K. Andreopoulou, J. K. Kallitsis and M. L. M. Rocco, *Surf. Interface Anal.*, 2023, **55**, 677–682.

58 W. Qi and D. Su, *ACS Catal.*, 2014, **4**, 3212–3218.



59 J. Sheng, B. Yan, W. D. Lu, B. Qiu, X. Q. Gao, D. Wang and A. H. Lu, *Chem. Soc. Rev.*, 2021, **50**, 1438–1468.

60 W. Qi, P. Yan and D. S. Su, *Acc. Chem. Res.*, 2018, **51**, 640–648.

61 H. Hu, Y. Nie, Y. Tao, W. Huang, L. Qi and R. Nie, *Sci. Adv.*, 2022, **8**, eabl9478.

62 L. Enders, D. S. Casadio, S. Aikonen, A. Lenarda, T. Wirtanen, T. Hu, S. Hietala, L. S. Ribeiro, M. F. R. Pereira and J. Helaja, *Catal. Sci. Technol.*, 2021, **11**, 5962–5972.

63 T. R. Gengenbach, G. H. Major, M. R. Linford and C. D. Easton, *J. Vac. Sci. Technol., A*, 2021, **39**, 013204.

