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# Metal-organic framework-derived hierarchical porous N/Co-doped carbon-supported sponge-like Pd-SnO<sub>2</sub> nanostructures for low-temperature CO oxidation†

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Metal-organic framework-derived porous N/Co-doped carbon (MOF-PNC) nanostructures-supported metal nanoparticles (NPs) are of great importance in multidisciplinary catalytic reactions; however, their catalytic performance toward low-temperature CO oxidation ( $CO_{Oxid}$ ) is rarely reported. Herein, a MOF-PNC-supported Pd-SnO<sub>2</sub> (Pd-SnO<sub>2</sub>/MOF-PNC) was synthesized *via* a microwave-irradiation (MW-I), annealing, and chemical etching approach for thermal  $CO_{Oxid}$ . The as-prepared Pd-SnO<sub>2</sub>/MOF-PNC had hierarchical porous sponge-like nanostructures composed of porous two-dimensional ultrathin nanosheets (NSs), co-doped with N/Co, with a high specific surface area (185.40 m² g<sup>-1</sup>) and pore volume (0.045 cm³ g<sup>-1</sup>), and ornamented with Pd-SnO<sub>2</sub> NPs (7.79  $\pm$  1.42 nm). These merits endowed the Pd-SnO<sub>2</sub>/MOF-PNC with excellent thermal catalytic  $CO_{Oxid}$  activity at a low complete CO conversion temperature ( $T_{100}$  = 65.6 °C) compared to those of Pd(1%)-SnO<sub>2</sub>/MOF-PNC (165.2 °C), Pd-SnO<sub>2</sub> (199.1 °C), Pd/MOF-PNC (107.9 °C) and commercial Pd/C catalysts (201.2 °C), due to the augmented electronic interaction and synergy of Pd NPs with oxygen-rich SnO<sub>2</sub> supports and Co-N<sub>x</sub> active sites in MOF-PNC. Thus, coupling two supports (*i.e.*, SnO<sub>2</sub>/MOF-PNC) is more crucial for promoting the low-temperature  $CO_{Oxid}$  activity of Pd NPs.

#### Introduction

The global energy demand has increased substantially in the last few decades and is expected to continue to rise owing to the inevitable industrialization and civilization. <sup>1-3</sup> Limitless efforts to solve these issues lie in developing green energy technologies (*i.e.*, fuel cells, <sup>4</sup> batteries, <sup>5</sup> and water splitting <sup>6,7</sup>) and gas conversion reactions. <sup>8-10</sup> CO<sub>Oxid</sub> is formed during heterogeneous catalysis electrochemically, <sup>11,12</sup> or thermally, <sup>13-16</sup> but the latter is feasible for large-scale applications. Pd-based catalysts are among the most active catalysts for thermal CO<sub>Oxid</sub>, however, the high cost and earth-rarity of Pd are critical barriers. <sup>17</sup> Using metal oxide supports (SnO<sub>2</sub>, SiO<sub>2</sub>, Fe<sub>2</sub>O<sub>3</sub>, TiO<sub>2</sub>, and CeO<sub>2</sub>) could boost

Unlike other supports, metal–organic framework-derived porous N/Co-doped carbon (MOF-PNC) nanostructures possess outstanding features (*i.e.* thermal/chemical stability and impressive surface area), and rich metal- $N_x$  active sites,  $^{22-24}$  which are beneficial for uniform distribution and stabilization of Pd NPs. $^{25-27}$  In addition, they make Pd active sites readily accessible and maximize their utilization for thermal  $CO_{Oxid}$ .  $^{28-30}$  For instance, porous C-supported Cu/Cu<sub>2</sub>O nanojunctions derived from (Cu-BTC)-MOF had impressive  $CO_{Oxid}$  activity achieved at  $T_{100}$ 

the catalytic performance and stability, and reduce the cost. <sup>18,19</sup> Also, the electronic interaction and synergy of Pd with metal oxide supports enhances the activation/dissociation of CO/O<sub>2</sub> reactants and quickly desorbs intermediate species (*i.e.*, carbonate/formate) and products (*i.e.*, CO<sub>2</sub>). For instance, the Pd/MgO-h-BN catalyst showed a lower complete CO conversion temperature ( $T_{100}$  = 140 °C) than Pd/MgO (180 °C), but Pd/h-BN could not achieve  $T_{100}$ , due to the interaction of Pd with the MgO-h-BN support that enhanced the large amount of adsorbed O<sub>2</sub> and rapidly desorbed the intermediates. <sup>20</sup> Pd@SiO<sub>2</sub>-673-CeO<sub>2</sub> catalysts had great thermal CO<sub>Oxid</sub> at a lower  $T_{100}$  (92 °C) than Pd@SiO<sub>2</sub>-673 (130 °C), owing to the interaction of Pd with dual supports (*i.e.*, SiO<sub>2</sub> and CeO<sub>2</sub>). <sup>21</sup>

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(155–190 °C), 31 due to a high ratio of Cu metal phases (Cu<sup>0</sup>, Cu<sup>+</sup>, and Cu<sup>2+</sup>) and interaction with the porous C support. Noticeably, MOF-PNC-supported metal NPs are not emphasized enough, particularly using SnO2 as a co-support, for thermal COoxid, and their effect remains ambiguous. 24,32-34

Herein, MOF-PNC-supported Pd-SnO<sub>2</sub> (Pd-SnO<sub>2</sub>/MOF-PNC) exhibited hierarchical porous sponge-like nanostructures comprising porous 2D ultrathin MOF-PNC NSs with abundant N/Co dopants, large specific surface area (185.40 m<sup>2</sup> g<sup>-1</sup>), and encapsulated spherical-like Pd-SnO<sub>2</sub> NPs (7.79  $\pm$  1.42 nm). Coupling the properties of SnO<sub>2</sub> (i.e., ease of adsorption and activation/ dissociation of CO/O2) and the merits of MOF-PNC (i.e., high surface area and rich metal-N<sub>x</sub> active sites) can ease CO oxidation on Pd at a low temperature and enhanced durability. The thermal CO<sub>Oxid</sub> activities of Pd-SnO<sub>2</sub>/MOF-PNC, Pd(1%)-SnO<sub>2</sub>/MOF-PNC, Pd-SnO<sub>2</sub>, SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC are benchmarked to commercial Pd/C catalysts in order to estimate the effect of the supports.

#### Materials and methods

#### **Materials**

Potassium tetrachloropalladate( $\Pi$ ) ( $K_2$ PdCl<sub>4</sub>  $\geq$  98%), tin( $\Pi$ ) chloride dihydrate (SnCl<sub>2</sub>·2H<sub>2</sub>O ≥ 98%), cobalt(II) nitrate hexahydrate  $(Co(NO_3)_2 \cdot 6H_2O \ge 94.5\%)$ , biphenyl-4,4'-dicarboxylic acid (BPDC  $\geq$  97%), ethylene glycol (EG  $\geq$  99.8%), triethyleneamine (TEA  $\geq$  99.5%), dimethylformamide (DMF  $\geq$  99.8%), and commercial Pd/C catalyst (20 wt%) were purchased from Sigma-Aldrich Chemie GmbH (Munich, Germany).

# Preparation of MOF-derived porous Co/N-doped carbon (MOF-

MOF-PNC was synthesized by mixing Co(NO<sub>3</sub>)·6H<sub>2</sub>O (0.44 g), BPDC (0.36 g), TEA (1.5 mL) and DMF (50 mL) under magnetic stirring at 25 °C, then microwave-irradiated at 600 W for 30 min.33 The obtained precipitates were washed and dried in a vacuum oven at 60  $^{\circ}$ C, followed by annealing at 800  $^{\circ}$ C for 5 h. The obtained powder was soaked in an aqueous solution of HCl (3 M) for 24 h, washed, and dried to give MOF-PNC.

#### Preparation of SnO<sub>2</sub>/MOF-PNC

SnO<sub>2</sub>/MOF-PNC was prepared by magnetically stirring SnCl<sub>2</sub>· 2H<sub>2</sub>O (47.8 mg) in a mixture of EG and water (4:1 by volume); then, MOF-PNC (100 mg) was added and the mixture was microwave-irradiated at 600 W for 1 h. The resulting product (SnO<sub>2</sub>/MOF-PNC) was washed and dried at 80 °C for 4 h under vacuum.

#### Preparation of Pd-SnO<sub>2</sub>/MOF-PNC

The as-prepared SnO<sub>2</sub>/MOF-PNC (100 mg) was mixed with K<sub>2</sub>PdCl<sub>4</sub> (61.35 mg) in EG (50 mL) at a pH of 12 with NaOH under magnetic stirring at 25 °C for 30 min, followed by microwave-irradiation at 600 W for 1 h (Anton Paar - Multiwave 3000).35 Then, the mixture's pH was lowered to 3 using 0.1 M HNO<sub>3</sub>, and it was washed thoroughly with deionized H<sub>2</sub>O, and

dried at 80 °C for 4 h under a vacuum to afford Pd-SnO<sub>2</sub>/MOF-PNC. A similar method was used for the preparation of Pd(1%)-SnO<sub>2</sub>/MOF-PNC (1 wt% Pd loading) and Pd-SnO<sub>2</sub>.

#### Preparation of Pd/MOF-PNC

Pd/MOF-PNC was prepared by mixing MOF-PNC (100 mg) with K<sub>2</sub>PdCl<sub>4</sub> (61.35 mg) in EG (50 mL) under magnetic stirring at 25 °C for 30 min and the pH was adjusted to 12 using NaOH solution (1 M). Then, the solution was placed in a microwave and irradiated at 600 W for 1 h. 35 Finally, the mixture's pH was decreased to 3 using 0.1 M HNO3, washed with deionized H2O, and dried at 80 °C for 4 h under vacuum.

#### Characterisation

The morphology and composition analysis were conducted on a scanning electron microscope (SEM, Hitachi S-4800, Hitachi, Tokyo, Japan) and transmission electron microscope (TEM, TecnaiG220, FEI, Hillsboro, OR, USA) equipped with an energy dispersive spectrometer (EDS). The electronic structure and surface composition were carried out by X-ray photoelectron spectroscopy ((XPS) Ultra DLD XPS Kratos, Manchester, UK). The powder X-ray diffraction pattern (XRD) was measured on an X-ray diffractometer (X'Pert-Pro MPD, PANalytical Co., Almelo, Netherlands). The N<sub>2</sub>-physisorption isotherms were measured on a Quanta chrome Autosorb-1 analyzer (Quanta chrome Instrument Corporation). The Fourier transform infrared spectra (FT-IR) were recorded on a Thermo Nicolet Nexus 670 FT-IR spectrometer (Thermo Scientific).

#### Thermal CO oxidation reaction

The thermal CO oxidation reaction was carried out in a fixed bed quartz tubular reactor connected to an online gas analyzer (Mass spectra, HIDEN ANALYTICAL the HPR-20 System) using 50 mg of each catalyst at the same Pd loading amount (20 wt%).<sup>36</sup> The catalyst was packed with quartz wool and fixed in the reactor and then pretreated at 200 °C (5 °C min<sup>-1</sup> heating ramp) under O<sub>2</sub> (5% in Ar) with a flow rate of 20 mL min<sup>-1</sup> for 1 h and subsequently under H<sub>2</sub> (5% in Ar) with a flow rate of 20 mL min<sup>-1</sup> for 1 h. After cooling to 25 °C, the catalyst was exposed to the gas mixture (O2 (20%) + CO (4%) + Ar (76%)) at a flow rate of 20 mL min<sup>-1</sup> under heating to 300  $^{\circ}$ C (5 $^{\circ}$  min<sup>-1</sup> ramping rate).<sup>36</sup> All the catalysts were subjected to the same treatment before the thermal CO oxidation. The percentage of CO conversion (%CO) was calculated using the following (eqn (1)):

$$^{\circ}$$
/CO =  $\left(\frac{\text{CO}_{\text{in}} - \text{CO}_{\text{out}}}{\text{CO}_{\text{in}}}\right) \times 100$  (1)

where COin is the input amount of CO and COout is the output amount of CO estimated from the mass spectra.

The CO temperature-programmed desorption (CO-TPD) was conducted by the initial pretreatment of each catalyst (50 mg) under Ar (50 mL min<sup>-1</sup> flow rate) at 300  $^{\circ}$ C (5  $^{\circ}$ C min<sup>-1</sup>) for 1 h in a Micromeritics ChemiSorb 2750 analyzer equipped with a thermal conductivity detector (TCD). After cooling to room temperature, the catalysts were exposed to (4% CO + 96% Ar) at a flow rate of 30 mL min  $^{-1}$  for 30 min under heating to 350  $^{\circ}$ C (5  $^{\circ}$ C min  $^{-1}$ ).  $^{36}$ 

The  $\rm H_2$  temperature-programmed reduction ( $\rm H_2$ -TPR) was measured by an initial pretreatment under Ar (50 mL min<sup>-1</sup>) at 300 °C (10 °C min<sup>-1</sup>) for 1 h and then exposed to (5%  $\rm H_2$  + 95% Ar) at a flow rate of 30 mL min<sup>-1</sup> under heating to 350 °C (5 °C min<sup>-1</sup>).<sup>36</sup>

The  $O_2$  temperature-programmed oxidation ( $O_2$ -TPO) was studied by initial treatment under Ar at a flow rate of 50 mL min<sup>-1</sup> at 300 °C (5 °C min<sup>-1</sup>) for 1 h and then exposed to (20%  $O_2$  + 80% Ar) at a flow rate of 30 mL min<sup>-1</sup> under heating to 350 °C (20 °C min<sup>-1</sup>).

To get more insights into the thermal CO oxidation, the CO oxidation rate ( $r_{\rm CO}$ ) of the catalysts was calculated at different CO conversion temperatures (10, 20, 50, and 80 °C) based on total gas flow ( $V_{\rm Gas}$ ), initial concentration of CO ( $X_{\rm CO}$ ), and molar weight of catalyst ( $M_{\rm Cat}$ ) using eqn (2).

$$r_{\rm CO} = \left[ \frac{X_{\rm CO} \times V_{\rm Gas}}{M_{\rm Cat}} \right] \tag{2}$$

The apparent activation energy ( $E_a$ ) was calculated at varied CO conversion (10–50%), based on the Arrhenius equation (eqn (3)) and the relationship between  $\ln r_{\rm CO}$  and 1/T.

$$\ln r = \ln A - \frac{E_{a}}{RT} + a \ln[CO] + b \ln[O_{2}]$$
 (3)

#### Results and discussion

The Pd–SnO<sub>2</sub>/MOF-PNC was synthesized  $\nu ia$  the MW-I of Co(NO<sub>3</sub>)·6H<sub>2</sub>O with biphenyl-4,4′-dicarboxylic acid (BPDC) and triethyleneamine (TEA) in dimethylformamide (DMF) solution to form a MOF, which was annealed and etched in HCl solution to afford MOF-PNC NSs (Fig. 1a).<sup>33</sup> This was followed by MW-I with the Sn precursor and then K<sub>2</sub>PdCl<sub>4</sub> in ethylene glycol (EG) to give Pd–SnO<sub>2</sub>/MOF-PNC.<sup>35</sup> The SEM of Pd–SnO<sub>2</sub>/MOF-PNC shows hierarchical porous sponge-like nanostructures (Fig. 1b), composed of porous 2D sheet-supported Pd NPs, with mean size (7.79  $\pm$  1.42 nm), proved by TEM (Fig. 1c and d), which is important for stabilizing the Pd NPs against aggregation during the CO<sub>Oxid</sub>. The lattice fringe (0.229 nm) of the Pd NPs is assigned to the {111} facet of face-center-cubic (fcc) Pd (Fig. 1e),<sup>35</sup> and the selected area electron diffraction pattern (SAED) reveals the typical rings of Pd (Fig. 1f).<sup>35</sup>

The EDX reveals the presence of Pd/Sn/Co/N/C/O with atomic contents (2.44/4.80/2.74/9.14/66.76/14.11 at%), indicating the successful formation of Pd/SnO<sub>2</sub> over MOF-PNC; however, Pd/MOF-PNC shows the existence of Pd/Co/N/C/O with contents (2.12/4.39/20.54/49.81/23.24 at%) (Fig. 1g) and uniform distribution mapped (Fig. S1, ESI†). Bulk and actual metal contents (Pd/Sn/Co (15.90/18.12/3.16 wt%)) in Pd-SnO<sub>2</sub>/MOF-PNC, (Pd/Co (17.70/11.24 wt%)) in Pd/MOF-PNC and (Pd (19.78 wt%)) in Pd/C are confirmed by ICP-OES (Table S1, ESI†). Meanwhile, the existence of Co in both catalysts is attributed to the partial etching by HCl, which is particularly important to coordinate with Pd-SnO<sub>2</sub> and provide additional active sites for thermal CO<sub>Oxid</sub>. The SEM of Pd/MOF-PNC shows its porous sponge-like structure (Fig. S2a, ESI†), but lower porosity than Pd-SnO<sub>2</sub>/MOF-PNC, owing

to the possible gas release during the MW-I in the absence of the Sn precursor.

The distribution of Pd NPs (9.07  $\pm$  1.75 nm), the lattice fringe (0.225 nm) for Pd{111} and SAED, but only Pd/Co/C distributed as mapped (Fig. S2b-f, ESI†). The Raman reveals the D- (1358.2 cm<sup>-1</sup>) and G-bands (1592.0 cm<sup>-1</sup>), but SnO<sub>2</sub> incorporation induced more defects in Pd-SnO<sub>2</sub>/MOF-PNC, proved by its higher  $I_D/I_G$  (2.01) than Pd/MOF-PNC (1.77) (Fig. S3a, ESI†).

The XRD of Pd–SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC at  $40^{\circ}$ ,  $46^{\circ}$  and  $68^{\circ}$  is assigned to  $\{111\}$ ,  $\{200\}$  and  $\{220\}$  of fcc Pd along with the  $\{002\}$  facet of amorphous C at  $26^{\circ}$ , but Pd–SnO<sub>2</sub>/MOF-PNC has additional peaks of  $\{101\}$ ,  $\{200\}$ ,  $\{211\}$ , and  $\{110\}$  attributable to the tetragonal SnO<sub>2</sub> (Fig. 2a), compared to Pd/C (Fig. S4, ESI†). The peak assigned to the C $\{002\}$  facet in Pd–SnO<sub>2</sub>/MOF-PNC is broadened with higher full width at half maximum (FWHM) than Pd/MOF-PNC, due to the possible coordination of Pd–SnO<sub>2</sub> with Co–N<sub>x</sub> in MOF-PNC. This is also evidenced by the slight positive shifts of fcc Pd in Pd–SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC relative to Pd/C, implying lattice contraction of Pd. Crystallite sizes (2.8 and 3.0 nm) from the Scherrer equation for Pd–SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC, respectively.

The XPS survey of the catalysts displays the valence state of Pd 3d/Co 2p/C 1s/O 1s/N 1s, but Pd–SnO<sub>2</sub>/MOF-PNC showed additional spectra of Sn 3d (Fig. S3b, ESI†). The atomic contents of Pd (2.16 at%) in Pd–SnO<sub>2</sub>/MOF-PNC and 2.71 at% in Pd/MOF-PNC imply the coherent distribution of Pd on the surface, but the bulk metal contents are given by the ICP-OES (Table S1, ESI†), which is critical for providing enough active sites for thermal CO<sub>Oxid</sub>. Pd 3d spectra of Pd–SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC display the phases of Pd<sup>0</sup>, Pd<sup>2+</sup>, and Pd<sup>4+</sup> (Fig. 2b).<sup>35</sup> The ratio of Pd<sup>0</sup> to Pd<sup>2+</sup> in Pd–SnO<sub>2</sub>/MOF-PNC (0.62) was lower than Pd/MOF-PNC (0.77), due to the incorporation of metal oxide (*i.e.*, SnO<sub>2</sub>) signifying more Pd<sup>2+</sup> in the Pd–SnO<sub>2</sub>/MOF-PNC.<sup>35</sup>

The Pd-SnO<sub>2</sub>/MOF-PNC had a higher ratio of Pd<sup>2+</sup> than Pd/MOF-PNC, owing to its possible interaction with SnO2 during the reduction process, which led to the partial oxidation of Pd to generate more active PdO<sub>r</sub> species and slightly decreased the dband center of Pd, evidenced by the slight positive shift of Pd binding energies of Pd-SnO<sub>2</sub>/MOF-PNC than Pd/MOF-PNC (Table S2, ESI†). Norskov et al. suggested that a d-band center and dbandwidth slightly below the Fermi level is optimal for CO chemisorption, so the reduced d-band center of Pd may allow strong interaction and possible poisoning and deactivation. 37,38 However, this did not happen in the case of Pd-SnO2/MOF-PNC, due to the promotional effect of Pd<sup>0</sup>/Pd-O<sub>x</sub> and their interfacial interaction with SnO2 and Co/N-doped porous MOF-derived C. Notably, Norskov and co-workers reported the ability of promoters to balance the CO chemisorption on the metal surface and enhance its activity.<sup>37,38</sup> The C 1s spectra are assigned to sp<sup>3</sup>/sp<sup>2</sup> (C-C/C=C) and the C-N bond (Fig. 2c), but shifted positively in Pd-SnO<sub>2</sub>/MOF-PNC than Pd/MOF-PNC, due to the reduced electron density on C by the interaction with Pd-SnO2. The Sn 3d spectra display major Sn2+  $(3d_{5/2} \text{ and } 3d_{3/2})$  and minor Sn<sup>0</sup> (Fig. 2d). Meanwhile, Co 2p spectra show  $Co^{2+}$  (3d<sub>3/2</sub> and 3d<sub>1/2</sub>) and  $Co^{0}$  (Fig. 2e). The N 1s spectra are attributed to pyridinic, pyrrolic, and graphitic (Fig. 2f).

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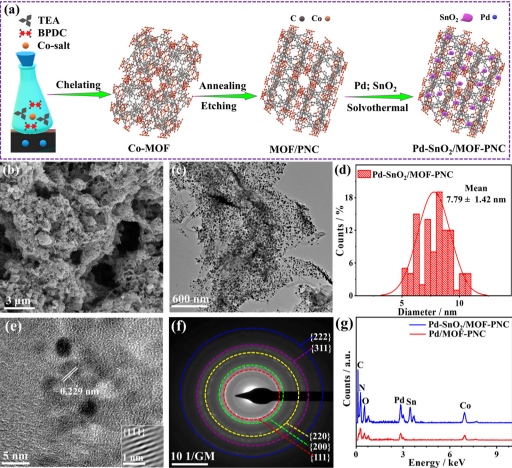


Fig. 1 (a) Schematic synthesis, (b) SEM, (c) TEM, (d) NPs size distribution, (e) HRTEM, (f) SAED of Pd-SnO<sub>2</sub>/MOF-PNC and (g) EDX of Pd-SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC.

The BET-specific surface area of Pd-SnO<sub>2</sub>/MOF-PNC (185.40 m<sup>2</sup> g<sup>-1</sup>) is slightly higher than that of Pd/MOF-PNC  $(152.83 \text{ m}^2 \text{ g}^{-1})$ , and Pd/C  $(107.91 \text{ m}^2 \text{ g}^{-1})$  (Fig. S5a-c, ESI†); meanwhile, the pore volume of Pd-SnO<sub>2</sub>/MOF-PNC (0.045 cm<sup>3</sup> g<sup>-1</sup>) was slightly higher than that of Pd/MOF-PNC (0.030 cm<sup>3</sup> g<sup>-1</sup>), and Pd/C  $(0.012 \text{ m}^3 \text{ g}^{-1})$ , in addition to their multiple pore size range (2-110 nm) with mean pore sizes of 66.78, 56.54, and 12.31 nm for Pd-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC, and Pd/C, respectively (Fig. S5d-f, ESI†).

The thermal CO<sub>Oxid</sub> shows typical light-off curves for the conversion of CO to CO<sub>2</sub> at heating temperatures (25–300 °C) and atmospheric pressure, but a superior activity on Pd-SnO<sub>2</sub>/ MOF-PNC than Pd/MOF-PNC, and Pd/C (Fig. 3a).  $^{39,40}$  The  $T_{100}$ of Pd-SnO<sub>2</sub>/MOF-PNC (65.6 °C) is lower than that of Pd/MOF-PNC (107.9  $^{\circ}$ C) by 42.3  $^{\circ}$ C and Pd/C (201.2  $^{\circ}$ C) by 135.6  $^{\circ}$ C. This is due to the electronic and synergistic interaction of Pd<sup>0</sup>/Pd-O<sub>x</sub> active sites with oxygen-enriched SnO<sub>2</sub>, Co-N<sub>x</sub> and MOF-PNC, which optimizes CO + O<sub>2</sub> adsorption/activation and desorption of CO<sub>2</sub> at low temperatures.

Thus, Pd-SnO<sub>2</sub>/MOF-PNC has higher CO<sub>Oxid</sub> kinetics than Pd/MOF-PNC, and Pd/C, owing to its capacity to oxidize CO at all applied temperatures (Fig. 3b), i.e., 50% of CO to CO<sub>2</sub>

 $(T_{50} = 58.4 \, ^{\circ}\text{C})$  on Pd-SnO<sub>2</sub>/MOF-PNC was lower than Pd/MOF-PNC (99.9 °C) and Pd/C (186 °C). So, Pd-SnO<sub>2</sub>/MOF-PNC completely oxidizes CO within only 13.12 min compared to Pd/MOF-PNC (21.58 min) and Pd/C (40.24 min) (Fig. 3c). The CO<sub>Oxid</sub> activity of Pd-SnO<sub>2</sub>/MOF-PNC was superior to previously reported Pd-based catalysts, i.e., Pd/CeSn, Pd@SiO2/TiO2, Pd@CeO<sub>2</sub>, Pd/MgO, (Pd@SiO<sub>2</sub>-673-CeO<sub>2</sub> (92 °C), <sup>21</sup> Pd/MgO-h-BN (140 °C), <sup>20</sup> Cu/Cu<sub>2</sub>O-500 nanojunctions (155 °C), <sup>31</sup> Pd/Cu/gC<sub>3</sub>N<sub>4</sub>NTs (154 °C),<sup>41</sup> Pd-Cu/gC<sub>3</sub>N<sub>4</sub>NWs (149 °C),<sup>42</sup> and Au/Pd/gC<sub>3</sub>N<sub>4</sub>NFs (149 °C)<sup>43</sup>) (Table S3, ESI†). The  $T_{100}$  of Pd–SnO<sub>2</sub>/MOF-PNC (65.6 °C) is among the lowest values reported for Pd-based catalysts as far as we found. The  $CO_{Oxid}$  rate  $(r_{CO})$  of Pd-SnO<sub>2</sub>/MOF-PNC was 1.95 and 2.15 times that of Pd/MOF-PNC and Pd/C, respectively (Fig. 3d), indicating maximum utilization of Pd active sites in Pd-SnO<sub>2</sub>/MOF-PNC, due to its greater porosity, which makes Pd active sites more accessible during CO<sub>Oxid</sub>.

This is further seen in the lower activation energy ( $E_a$  = 69.5 kJ mol<sup>-1</sup>) of Pd-SnO<sub>2</sub>/MOF-PNC than Pd/MOF-PNC  $(74.1 \text{ kJ mol}^{-1})$  and Pd/C  $(89.2 \text{ kJ mol}^{-1})$  (Fig. 3e). The CO<sub>Oxid</sub> stability of Pd-SnO2/MOF-PNC at 65.6 °C was shown by a timeon-stream (TOS) for 108 h (Fig. 3f), which reveals excellent durability with insignificant loss in  $T_{100}$ . The stability of

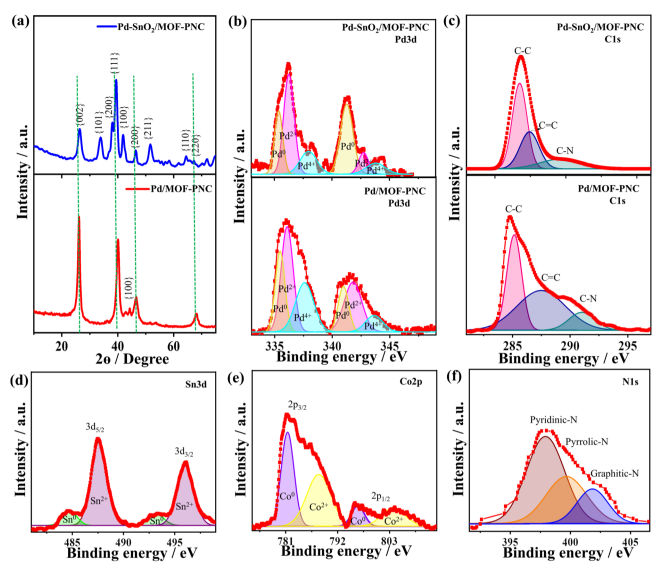


Fig. 2 (a) XRD, and high-resolution XPS (b) Pd 3d, (c) C 1s (d) Sn 3d, (e) Co 2p and (f) N 1s of Pd-SnO<sub>2</sub>/MOF-PNC and Pd/MOF-PNC. The green lines in (a) refer to the positions of pure Pd NPs on C, which were taken from the JCPDS database.

Pd–SnO<sub>2</sub>/MOF-PNC is further provided by TEM, which displayed the good dispersion of Pd nanoparticles over SnO<sub>2</sub>/MOF-PNC without any obvious aggregation, indicating the architecture durability (Fig. S6a, ESI†). The XPS full-scan showed the presence of C 1s, Pd 3d, Co 2p, O 1s, N 1s, and Sn 3d without significant degradation for Pd (2.13 at%), implying compositional stability (Fig. S6b, ESI†). The BET maintained the same isotherm adsorption/desorption features before stability with an inferior loss in the surface area and pore volume (Fig. S6c and d, ESI†). This implies the reservation of the physiochemical properties of Pd–SnO<sub>2</sub>/MOF-PNC after the durability test, as also proved by measuring the CO<sub>Oxid</sub> after the stability test, which displayed a minimal loss in the  $T_{100}$  (only 5 °C) (Fig. S4b, ESI†).

The active sites and  $CO_2$  production rate of Pd-SnO<sub>2</sub>/MOF-PNC are proved by its higher turnover number (TON = 14.8  $\mu$ mol  $g_{Cat}^{-1}$ ) and turnover frequency (TOF = 61.97 h<sup>-1</sup>) relative to Pd/MOF-PNC (11.5  $\mu$ mol  $g_{Cat}^{-1}$ ; 45.75 h<sup>-1</sup>) and Pd/C

 $(7.86~\mu mol~g_{Cat}^{-1}; 31.49~h^{-1})$  (Fig. 4a and b). The recyclability of Pd–SnO<sub>2</sub>/MOF-PNC reveals that the catalyst remained active for 5 consecutive cycles with no degradation. The CO-TPD gave sharp peaks for Pd–SnO<sub>2</sub>/MOF-PNC (74.5 °C) and Pd/MOF-PNC (72.6 °C), compared to a broad peak for Pd/C (174.2 °C) (Fig. 5a). This implies more accessible Pd active sites in Pd–SnO<sub>2</sub>/MOF-PNC and its capacity to adsorb/oxidize CO at a lower temperature due to the presence of an oxygen-rich SnO<sub>2</sub> support, higher surface area, and abundant active sites.<sup>36</sup>

The O<sub>2</sub>-TPO displays a sharp oxygen-uptake peak on Pd–SnO<sub>2</sub>/MOF-PNC (66.8 °C) relative to broad peaks on Pd/MOF-PNC (116.3 °C) and Pd/C (197.6 °C) (Fig. 5b), implying ease of O<sub>2</sub> adsorption (O<sub>ads</sub>) on Pd–SnO<sub>2</sub>/MOF-PNC, due to the interfacial interaction of Pd with SnO<sub>2</sub> and MOF-PNC. This leads to rapid turnover of the adsorbed CO/O<sub>2</sub>, which is important for inducing a reaction between the active O<sub>ads</sub> and CO<sub>ad</sub> on the Pd surface to allow quick CO<sub>Oxid</sub> kinetics.<sup>36</sup> Also, the oxygenated species (*i.e.*, SnO<sub>2</sub>) in Pd–SnO<sub>2</sub>/MOF-PNC enables a lower energy barrier for

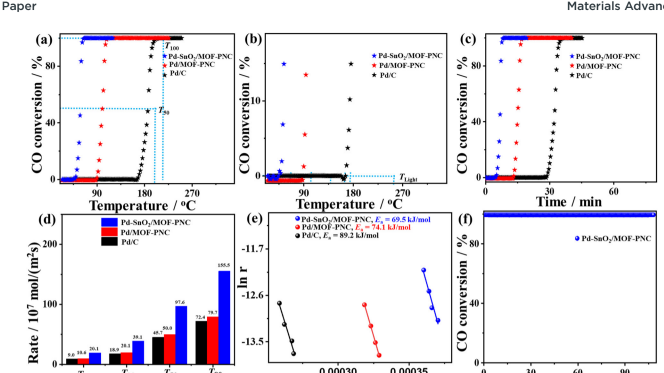


Fig. 3 (a, b) Temperature-dependent CO conversion, (c) time-dependent CO conversion, (d) rate at different CO conversion temperatures, (e) Arrhenius plots, and (f) time on stream (TOS) of Pd-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC, and Pd/C.

0.00030

0.00035

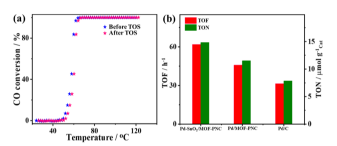


Fig. 4 (a) Temperature of CO conversion before and after the TOS test and repetitive CO conversion after H2-TPR and O2-TPO of Pd-SnO2/ MOF-PNC. (b) Turnover number (TON) and turnover frequency (TOF) of Pd-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC and Pd/C

CO/O2 uptake/activation, thereby accelerating the complete CO<sub>Oxid</sub> kinetics.<sup>24,33</sup> The H<sub>2</sub>-TPR confirms the reducibility of the catalysts, 36 where Pd-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC, and Pd/C show a broad H<sub>2</sub> adsorption at 62.3, 184.1, and 199.9 °C, respectively, which implies that Pd-SnO<sub>2</sub>/MOF-PNC got reduced fast, owing to the interaction of SnO<sub>2</sub>, Pd<sup>2+</sup>, and Co<sup>2+</sup> (Fig. 5c). This may serve as evidence for its exposed metal active sites.

The H<sub>2</sub>-TPR, O<sub>2</sub>-TPO, and CO-TPD reveal that coupled Pd, SnO<sub>2</sub>, and MOF-PNC enhance the CO redox properties and CO<sub>Oxid</sub> activity of Pd-SnO<sub>2</sub>/MOF-PNC. Hence, the CO<sub>Oxid</sub> mechanism on Pd-SnO<sub>2</sub>/MOF-PNC could follow Langmuir-Hinshelwood, 36 i.e., co-adsorption of CO/O2 on Pd-SnO2/ MOF-PNC, followed by dissociation of O2 to form O lattice and O adsorbed (O<sub>ads</sub>) (eqn (R1) and (R2)), which then oxidizes CO<sub>ads</sub> to CO<sub>2ads</sub> (i.e., the rate determining step (eqn (R3))) and CO<sub>2</sub> desorbed from Pd-SnO<sub>2</sub>/MOF-PNC (eqn (R4)).

$$O_{2g} + Pd-SnO_2/MOF-PNC \rightarrow 2O_{ads}-Pd-SnO_2/MOF-PNC$$
(R1)

30

60

Time / h

90

$$CO_g + Pd-SnO_2/MOF-PNC \rightarrow CO_{ads}-Pd-SnO_2/MOF-PNC$$
(R2)

$$O_{ads}$$
-Pd-SnO<sub>2</sub>/MOF-PNC +  $CO_{ads}$ -Pd-SnO<sub>2</sub>/MOF-PNC  
 $\rightarrow CO_{2ads}$ -Pd-SnO<sub>2</sub>/MOF-PNC (R3)

$$CO_{2ads}$$
-Pd-SnO<sub>2</sub>/MOF-PNC  $\rightarrow CO_2 + Pd$ -SnO<sub>2</sub>/MOF-PNC (R4)

To investigate the effect of Pd nanoparticles loaded with a lower content (i.e., 1 wt%) decorated on SnO<sub>2</sub>/MOF-PNC, Pd(1%)-SnO<sub>2</sub>/MOF-PNC was prepared and tested for thermal CO oxidation, which showed significantly higher  $T_{100}$  (165.2 °C) than Pd-SnO<sub>2</sub>/MOF-PNC (65.6 °C) (Fig. S7a, ESI†). Meanwhile, in the absence of Pd nanoparticles, SnO<sub>2</sub>/MOF-PNC could not attain  $T_{100}$  even at 300 °C, implying that Pd is the main active site for the thermal CO oxidation. Also, to get more insights into the effect of the support, Pd-SnO<sub>2</sub> was examined for CO<sub>Oxid</sub> and achieved  $T_{100}$ (199.1 °C) which was greater than Pd-SnO<sub>2</sub>/MOF-PNC (65.6 °C) and even Pd/MOF-PNC (107.9 °C) (Fig. S7a-c, ESI†), which indicates that using a co-support of SnO<sub>2</sub>/MOF-PNC is crucial for promoting the CO<sub>Oxid</sub> activity and kinetics as further seen in the lower rate  $(r_{CO})$  of Pd-SnO<sub>2</sub> than that of Pd-SnO<sub>2</sub>/MOF-PNC, and Pd(1%)-SnO<sub>2</sub>/MOF-PNC (Fig. S7d, ESI†). Also, the estimated  $E_a$  of Pd-SnO<sub>2</sub> was greater than that of Pd(1%)-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC, and Pd-SnO<sub>2</sub>/MOF-PNC (Fig. S7e, ESI†). These results clarify the importance of combining mixed Pd phases (Pd<sup>0</sup>/Pd-O<sub>x</sub>) and an oxygen-rich SnO2 support for excellent COoxid activity as

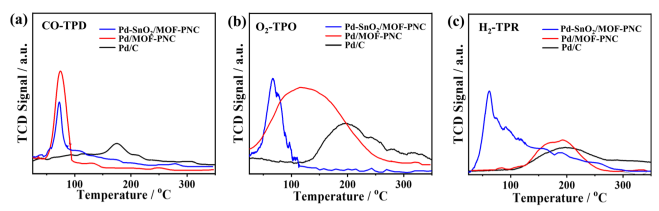


Fig. 5 (a) CO-temperature-programmed desorption (CO-TPD), (b)  $O_2$ -temperature-programmed oxidation ( $O_2$ -TPO), and (c)  $H_2$ -temperature-programmed reduction ( $H_2$ -TPR) of Pd-SnO<sub>2</sub>/MOF-PNC, Pd/MOF-PNC and Pd/C.

shown by low  $T_{100}$ , but high kinetics, TON and TOF of Pd–SnO<sub>2</sub>/MOF-PNC. Thus, coupling Pd with a metal oxide support is preferred for promoting the  $CO_{Oxid}$ , due to the optimal  $CO/O_2$  adsorption and ease of activation/dissociation at low temperatures.

#### Conclusion

In brief, hierarchical porous sponge-like Pd–SnO<sub>2</sub>/MOF-PNC was prepared by the MW-I, annealing, and chemical etching approach to initially form MOF-PNC, mixed with SnO<sub>2</sub> and Pd and then MW-I. Pd–SnO<sub>2</sub>/MOF-PNC comprises porous 2D ultrathin MOF-PNC NSs with monodispersed Pd–SnO<sub>2</sub> NPs (7.79  $\pm$  1.42 nm), a large specific surface area (185.40  $\rm m^2~g^{-1}$ ), and pore volume (0.045  $\rm cm^3~g^{-1}$ ). Thus, the CO<sub>Oxid</sub> at  $T_{100}$  of Pd–SnO<sub>2</sub>/MOF-PNC (65.6 °C) was lower than those of Pd/MOF-PNC (107.9 °C), Pd(1%)–SnO<sub>2</sub>/MOF-PNC (165.2 °C), Pd–SnO<sub>2</sub> (199.1 °C), and Pd/C (201.2 °C), and also superior to most previously reported Pd-based catalysts. This originated from the electronic interaction and synergism of Pd NPs with oxygen-rich SnO<sub>2</sub> supports and Co-N<sub>x</sub> active sites in MOF-PNC to deliver low energy barriers and high kinetics. These results indicate that using two supports, SnO<sub>2</sub>/MOF-PNC is preferred for promoting the thermal CO<sub>Oxid</sub> activity of Pd NPs.

#### Conflicts of interest

We declare no conflicts of interest.

### Acknowledgements

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