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Highly dispersed Pd-based pseudo-single atoms in zeolites for hydrogen generation and pollutant disposal†

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Atomically dispersed metal catalysts with excellent activity and stability are highly desired in heterogeneous catalysis. Herein, we synthesized zeolite-encaged Pd-based pseudo-single atoms via a facile and energy-efficient ligand-protected direct H₂ reduction method. C_s-corrected scanning transmission electron microscopy, extended X-ray absorption, and pair distribution function measurements reveal that the metal species are close to atomic-level dispersion and completely confined within the intersectional channels of silicalite-1 (S-1) zeolite with the MFI framework. The Pd@S-1-H exhibits excellent activity and stability in methane combustion reactions with a complete combustion temperature of 390 °C, and no deactivation is observed even after 100 h on stream. The optimized bimetallic 0.8Pd0.2Ni(OH)₂@S-1-H catalyst exhibits an excellent H₂ generation rate from FA decomposition without any additives, affording a superhigh turnover frequency up to 9308 h⁻¹ at 333 K, which represents the top activity among all of the best heterogeneous catalysts under similar conditions. Significantly, zeolite-encaged metal catalysts are first used for Cr(vi) reduction coupled with formic acid (FA) dehydrogenation and show a superhigh turnover number of 2980 mol(Cr₂O₇²⁻) mol(Pd)⁻¹ at 323 K, surpassing all of the previously reported catalysts. This work demonstrates that zeolite-encaged pseudo-single atom catalysts are promising in efficient hydrogen storage and pollutant disposal applications.

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Introduction

Zeolites, a class of crystalline materials with orderly distributed micropores, large specific surface areas, and adjustable acidity/basicity, have been widely used in diverse industrial applications as adsorbents, separation materials, and catalysts.^{1–4} In recent decades, zeolites have also been regarded as ideal supports to immobilize metal species with small sizes and high dispersity. The rational design and fabrication of zeolite-supported metal catalysts with superior catalytic activity and

stability is always a hot topic in heterogeneous catalysis.^{5–13} Despite the excellent confinement effect of zeolites, zeolite-supported metal catalysts prepared by the conventional impregnation method still often suffer from the formation of large and uneven metal particles as well as unsatisfactory sintering and leaching during the catalytic process. This is because the metal species are inclined to be located on the outer surface of zeolite crystals during the synthetic process. To overcome these shortcomings, several innovative *in situ* synthetic strategies have been developed for the confined synthesis of ultra-small metal species within zeolite crystals.^{14–26} For example, Iglesia and coworkers developed a ligand-stabilized approach to encapsulate a series of small metal nanoparticles with a mean size of 1–2 nm into micropores of various aluminosilicate zeolites.^{23,24} Corma and coworkers developed a zeolite structure transformation approach to encage subnanometric Pt clusters within MWW zeolites.²⁵ Xiao and coworkers developed a metal-containing-seed-directed synthesis method to fix different kinds of metal nanoparticles with a diameter range of 0.8–3.6 nm within zeolite crystals.²² Our group also reported the encapsulation of subnanometric metal clusters within pure silica zeolites *via* a ligand-protected method under direct hydrothermal conditions.^{19,20,26,27} Recently, single- and pseudo-single atom catalysts with extremely high metal dispersity have attracted extensive research interest from researchers due to

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their nearly 100% atom-utilization efficiency and excellent catalytic properties.^{28–32} However, such catalysts usually suffer from poor durability and inferior reusability in catalysis. The rational design and fabrication of zeolite-supported metal species with (pseudo)single-atom dispersion while maintaining excellent stability during catalytic reactions and further expanding their catalytic applications remains a nontrivial task but is greatly challenging.

Methane, as a typical greenhouse gas, produces a 20-fold higher greenhouse effect than carbon dioxide. The release of unburned methane from exhaust gas has become an environmental issue in recent years, and catalytic methane combustion technology is an effective way to reduce its pollution of the atmosphere.^{33–36} Supported Pd-based catalysts have been considered the most active catalysts for methane combustion, but they still often suffer from either poor activity or inferior long-term stability toward the complete oxidation of methane due to the severe aggregation of metal species. The development of efficient catalysts with outstanding activity and stability is highly desired. Hydrogen (H_2) is considered a green and efficient energy carrier, but efficient and controllable hydrogen storage remains a bottleneck for the future hydrogen economy.^{37–40} Formic acid (FA, HCOOH) has emerged as a promising liquid-phase hydrogen storage material for portable H_2 storage applications.^{41–44} In addition, FA can also be used as a green and cost-effective reductant for the reduction of toxic pollutants, such as hexavalent chromium (Cr(vi)), which possesses serious mutagenicity and carcinogenicity. The *in situ* generated H_2 from FA dehydrogenation can efficiently reduce Cr(vi) to Cr(III).^{45,46} Although some heterogeneous catalysts have recently been used for this reaction, few catalysts with both excellent performance and stability have been reported.

In this work, we synthesized zeolite-encaged highly-dispersed pseudo-single Pd and bimetallic Pd–Ni(OH)₂ atoms *via* a facile and energy-saving ligand-protected direct H_2 reduction method. The location, pseudo-single atom nature, and precise structure of the metal species were determined by C_s -corrected scanning transmission electron microscopy, extended X-ray absorption, and pair distribution function (PDF) analysis. Due to the extremely high metal dispersion and almost complete exposure of active metal sites, Pd@S-1-H exhibited superior methane combustion activity with a complete combustion temperature of 390 °C, which is lower than that of Pd@S-1-C (416 °C) subjected to calcination and reduction procedures and Pd/S-1-im (513 °C) synthesized by the impregnation method. The catalytic activity of methane combustion over Pd@S-1-H remains unchanged even after 100 h on stream. Taking advantage of the synergistic effect of the bimetallic interface, the optimized bimetallic 0.8Pd0.2Ni(OH)₂@S-1-H catalyst exhibits an excellent H_2 generation rate from FA decomposition without any additives, affording a superhigh turnover frequency up to 9308 h^{-1} at 333 K, which is 33 times higher than that over Pd/S-1-im synthesized by the impregnation method, representing the top activity among all of the best heterogeneous catalysts under similar conditions. Significantly, 0.8Pd0.2Ni(OH)₂@S-1-H also acts as an efficient catalyst in Cr(vi) reduction coupling with FA dehydrogenation, showing

a record turnover number of 2980 mol(Cr₂O₇²⁻) mol(Pd)⁻¹ at 323 K.

Results and discussion

As shown in Fig. 1A, S-1 zeolite-encaged Pd-based metal samples were prepared by a facile ligand-protection method under *in situ* hydrothermal conditions at 170 °C for 3 days in reaction gels with molar compositions of 1TEOS/0.4TPAOH/35H₂O/ a [Pd(en)₂]Cl₂/ b [Ni(en)₃](NO₃)₂, (TEOS = tetraethoxysilane, TPAOH = tetrapropylammonium hydroxide, en = ethylenediamine, $a + b = 0.0045$, $a/b = 10/0, 9/1, 8/2, 7/3$, and 0/10). Next, the obtained samples were subjected to two different thermal treatment methods: one was calcinated in air at 550 °C for 6 h and then reduced in H_2 flow at 500 °C for 2 h (named x Pd(1- x)Ni(OH)₂@S-1-C, where x represented the molar ratio of Pd/(Pd + Ni) in the initial synthesis gel); the other was directly reduced in pure H_2 flow at 500 °C for 2 h (named x Pd(1- x)Ni(OH)₂@S-1-H). For comparison, the Pd/S-1-im and Pd–Ni(OH)₂/S-1-im samples were also prepared by the incipient wetness impregnation method with the same metal loadings as the Pd@S-1 and Pd–Ni(OH)₂@S-1 samples, respectively. Powder X-ray diffraction measurements confirm that the obtained zeolite-encaged Pd-based samples possess an MFI topological structure with high crystallinity (Fig. S1†). Inductively coupled plasma atomic emission spectroscopy (ICP-AES) measurements reveal that all samples possessed a similar molar amount of total metal species, giving total metal loadings of Pd@S-1 and Pd–Ni(OH)₂@S-1 samples in the range of 0.64–0.48 wt%. The molar ratios of Pd/Ni in 0.9Pd0.1Ni(OH)₂@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-H, and 0.7Pd0.3Ni(OH)₂@S-1-H are 0.89/0.11, 0.81/0.19, and 0.72/0.28, respectively, which are similar to the preset ratios in synthetic gels (Table S1†).

Scanning electron microscopy (SEM) measurements demonstrate that all zeolite-supported metal samples possess a uniformly hexagonal prism morphology with average sizes of 200–300 nm. Note that many bright dots corresponding to Pd nanoparticles (NPs) can be clearly observed in Pd/S-1-im, indicating that large amounts of Pd species with large sizes are located outside of S-1 crystals. In contrast, no metal species are detected on the outside of zeolite-encaged metal samples synthesized by the *in situ* method (Fig. S2†). To obtain precise information about the size and location of the metal species in the samples, C_s -corrected high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) and high-resolution TEM measurements were performed. As shown in Fig. 1B–I and S3, S4,† homogeneously distributed metal clusters/atoms with subnanometer sizes are completely confined within intersectional void spaces between the straight and sinusoidal channels of S-1 zeolites in the representative Pd@S-1-C, Pd@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-C, and 0.8Pd0.2Ni(OH)₂@S-1-H samples, and no metallic species could be observed on the outside of the zeolites. In contrast, the metal NPs are considerably larger in Pd/S-1-im (4–5 nm), and the great majority of metal species are located on the outer surface of zeolite crystals, which is consistent with the SEM observation. The introduction of Ni species does not affect the size of metal



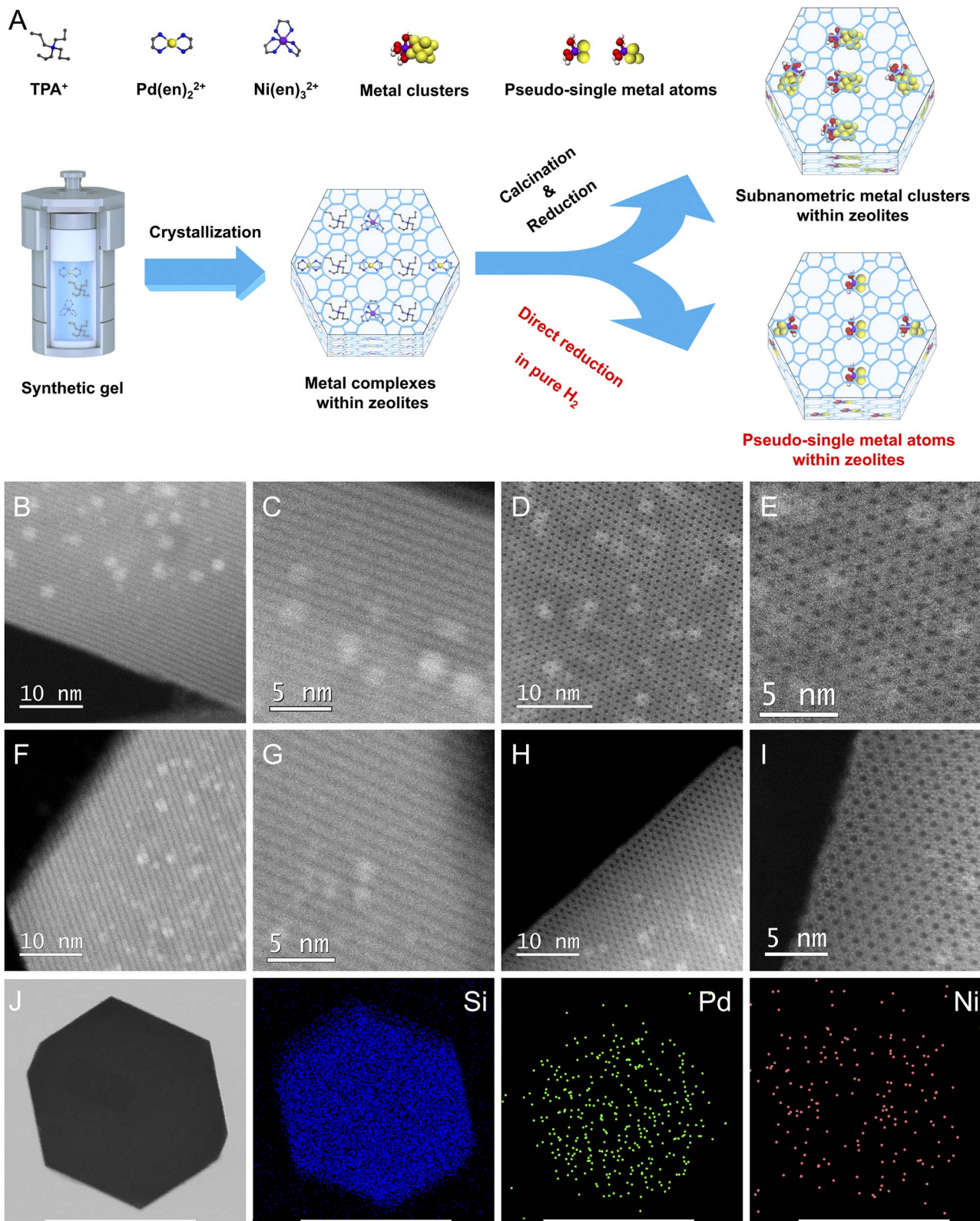


Fig. 1 (A) Schematic illustration of the preparation of zeolite-encaged Pd-based catalysts. C_s -corrected STEM images of (B and C) Pd@S-1-C; (D and E) 0.8Pd0.2Ni(OH)₂@S-1-C; (F and G) Pd@S-1-H; (H and I) 0.8Pd0.2Ni(OH)₂@S-1-H. (J) TEM image of 0.8Pd0.2Ni(OH)₂@S-1-H catalyst and corresponding EDX mapping images for Si, Pd, and Ni elements. Scale bar: 250 nm.



clusters in zeolites. Significantly, the Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H samples subjected to direct H₂ reduction possess smaller metal sizes than the corresponding Pd@S-1-C and 0.8Pd0.2Ni(OH)₂@S-1-C samples treated by the calcination-reduction process. The metal species in the Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H samples are in the form of single and pseudo-single atoms. This can be mainly attributed to the fact that during the direct H₂ reduction process, the hydrogenolysis of organic templates and ligands and reduction of metal species occur simultaneously; the ammonia gas produced by hydrolysis of organic species (confirmed by H₂-temperature-programmed reduction/mass spectrometry analysis, Fig. S5A†) can act as protective agents to restrain the aggregation of metal species, resulting in the formation of single and pseudo-single atoms with decreased size and increased dispersion.

Thermogravimetry analyses reveal that no noteworthy weightlessness can be observed in either Pd@S-1-C or Pd@S-1-H when the temperature is above 300 °C, indicating that almost all organic species were removed by the calcination-reduction and direct H₂ reduction processes (Fig. S5B†). N₂ adsorption/desorption measurements show that the micropore volume of zeolite-encaged metal samples decreased by approximately 10% compared with that of pure S-1 zeolite ($\sim 0.115 \text{ cm}^3 \text{ g}^{-1}$) because metal species confined within the zeolite matrix caused a slight decrease ($\sim 0.012 \text{ cm}^3 \text{ g}^{-1}$), but large microporous volumes ($0.103\text{--}0.105 \text{ cm}^3 \text{ g}^{-1}$) and abundant surface areas ($399\text{--}415 \text{ m}^2 \text{ g}^{-1}$) were still preserved in Pd@S-1-C, Pd@S-1-H, and 0.8Pd0.2Ni(OH)₂@S-1-H, respectively (Fig. S6 and Table S1†).

To further identify the electronic and local structure information of metal species encapsulated within S-1 zeolites, the X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) spectra of representative Pd@S-1-C, Pd@S-1-H, and 0.8Pd0.2Ni(OH)₂@S-1-H samples are measured. The detailed structural parameters according to EXAFS fittings are summarized in Table S2.† As shown in Fig. 2A and S7,† the Pd K-edge XANES spectra of Pd@S-1-C, Pd@S-1-H, and 0.8Pd0.2Ni(OH)₂@S-1-H samples are different from Pd foil and PdO, showing a higher oxidation state than the Pd foil. It can be attributed to the formation of ultrasmall Pd species, which are easy to coordinate with oxygen atoms of zeolite frameworks. A small shift to the higher absorption edge energy and increased intensity of the white line can be observed for Pd@S-1-H in comparison with Pd@S-1-C, suggesting the higher oxidation state of Pd in the Pd@S-1-H sample. Note that a small edge shift to lower energy and slightly decreased intensity of the white line is found in 0.8Pd0.2Ni(OH)₂@S-1-H as compared with Pd@S-1, indicating that the Pd species in 0.8Pd0.2Ni(OH)₂@S-1-H possess a higher electron density than that in Pd@S-1-H. The Ni K-edge XANES spectrum of 0.8Pd0.2Ni(OH)₂@S-1-H is nearly the same as that of the Ni(OH)₂ reference spectrum, revealing that the Ni species in the bimetallic clusters exist in the form of a Ni(OH)₂ structure (Fig. 2B). In the Pd K-edge Fourier-transformed EXAFS spectra, the significant peaks at ~ 2.01 and 2.72 \AA were identified as the Pd–O and Pd–Pd metallic bonds, respectively (Fig. 2C). Based on the EXAFS fitting results, the average coordination number (CN)

of Pd–O shell for Pd@S-1-H is 2.9, which is higher than that of Pd@S-1-C (1.9) and 0.8Pd0.2Ni(OH)₂@S-1-H (2.4), further confirming the higher oxidation state of Pd species in Pd@S-1-H sample. The average CN of the Pd–Pd shell for Pd@S-1-H is just 1.9, which is much smaller than that for Pd@S-1-C (3.5). It indicates that Pd clusters in both Pd@S-1-H and Pd@S-1-C are subnanometer in size, and the average size of Pd clusters in Pd@S-1-H is smaller than that in Pd@S-1-C, in line with the observation of C_s-corrected STEM measurements. Notably, the average CN of the Pd–Pd shell for 0.8Pd0.2Ni(OH)₂@S-1-H is as low as 0.9, indicating the Pd species are dimer- or single-atomic dispersion. A small peak at 2.45 \AA corresponding to Pd–Ni metallic bond (CN = 0.13) can be also detected, which reveals the partial formation of Pd–Ni alloy structure, and some electrons may transfer from Ni to Pd and form electron-enriched Pd surfaces in 0.8Pd0.2Ni(OH)₂@S-1-H. The average valence states of Pd sites in different samples could be disclosed by establishing a linear correlation between the edge energy and the valence state of reference samples. The Pd@S-1-H shows a valence state of Pd (+1.83), which is slightly higher than Pd@S-1-C (+1.67). Notably, the valence state of Pd species in 0.8Pd0.2Ni(OH)₂@S-1-H (+1.17) is lower than those of two Pd samples, indicating the higher electron density in 0.8Pd0.2Ni(OH)₂@S-1-H (Fig. S8†). In the Ni K-edge EXAFS spectra of 0.8Pd0.2Ni(OH)₂@S-1-H is consistent with the Ni(OH)₂ reference spectrum, further confirming the formation of the bimetallic Pd–Ni(OH)₂ structure (Fig. 2D).

The interatomic structure of the encapsulated metallic clusters in zeolite was further investigated by pair distribution function (PDF) analyses. The experimental PDFs of different S-1 zeolite-encaged metal catalysts and pure S-1 zeolite are shown in Fig. 2E. The peaks at *ca.* 1.6, 2.6, and 3.1 \AA can be attributed to the Si–O, O–O, and Si–Si distances in S-1 zeolites, respectively. The above signals are very similar in Pd@S-1-C, Pd@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-H, and pure S-1 zeolite samples, indicating the introduction of metal species inside the zeolites rarely affect the structure of S-1 zeolites. The differential PDF (d-PDF) analysis is a method based on the difference between two PDFs and can be used for obtaining structural information of guest species.⁴⁷ Therefore, we can extrapolate the structure of Pd species in S-1 zeolites corresponding to the calculated difference in the experimental PDFs of the Pd@S-1 and pure S-1 zeolite. Fig. 2F shows the calculated d-PDF of encapsulated metal species in zeolite-encaged metal catalysts. The peaks at approximately 2.7 and 4.7 \AA , assigned to the first and second-shell Pd–Pd bonds can be observed in Pd@S-1-C, indicating the formation of small Pd clusters. In contrast, the first-shell Pd–Pd bond in Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H are nearly neglected, reflecting the formation of atomically dispersed Pd atoms in the samples subjected to the direct H₂ reduction treatment, in accordance with the observation of C_s-corrected STEM and EXAFS results. Notably, a unique peak at about 2.0 \AA can be observed in 0.8Pd0.2Ni(OH)₂@S-1-H, which is attributed to the formation of the Ni–O bond.

The thermal stabilities at high temperatures under various atmospheres are also investigated over the representative 0.8Pd0.2Ni(OH)₂@S-1-H sample. The average particle size of



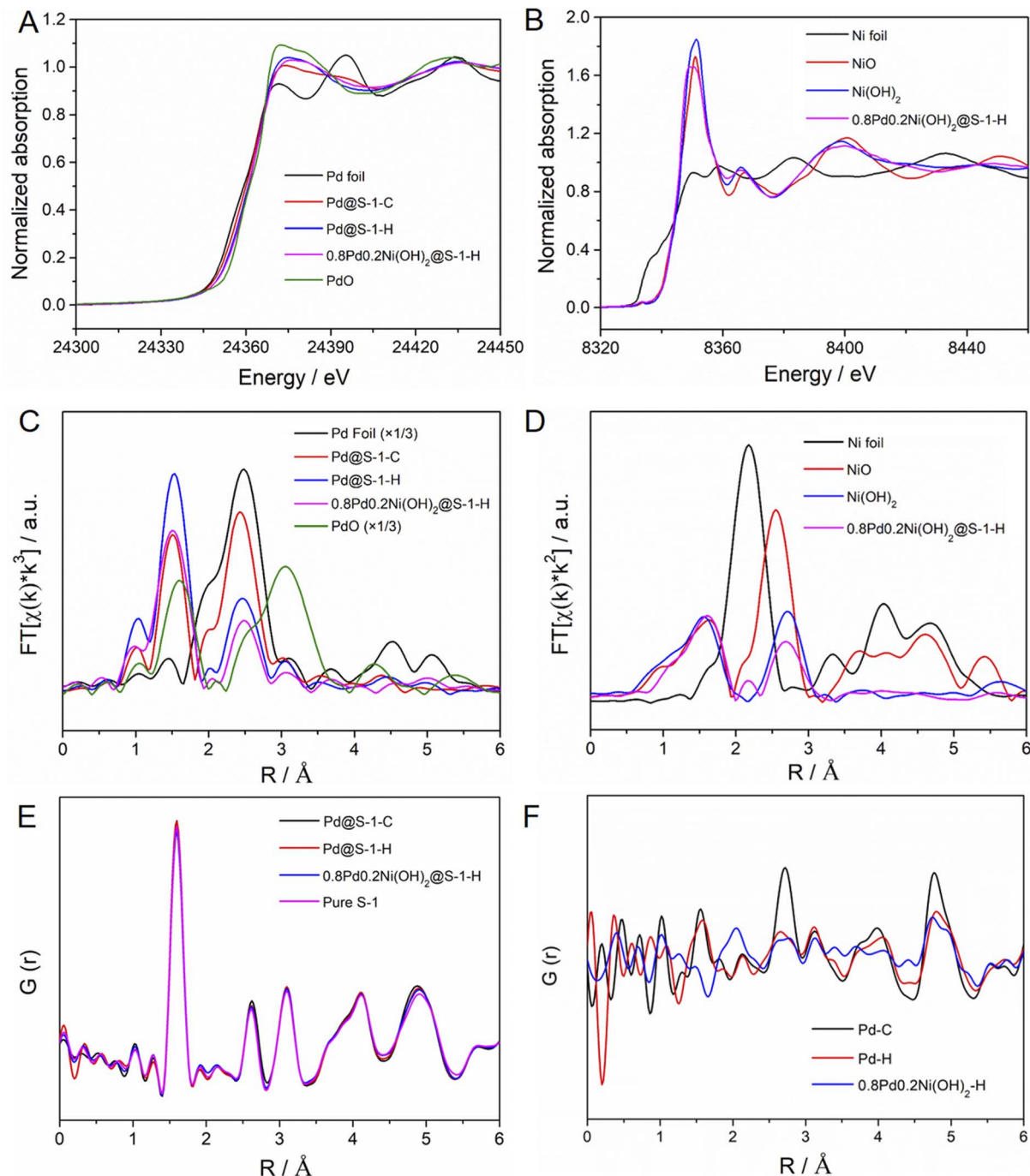


Fig. 2 (A) Pd K-edge XANES spectra of Pd@S-1-C, Pd@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-H and Pd foil, PdO standards. (B) Ni K-edge XANES spectra of 0.8Pd0.2Ni(OH)₂@S-1-H and Ni foil, NiO, Ni(OH)₂ standards. (C) Fourier transform of k^3 -weighted EXAFS spectra of Pd@S-1-C, Pd@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-H and Pd foil, PdO standards. (D) Fourier transform of k^2 -weighted EXAFS spectra of 0.8Pd0.2Ni(OH)₂@S-1-H and Ni foil, NiO, Ni(OH)₂ standards. (E) PDFs of Pd@S-1-C, Pd@S-1-H, 0.8Pd0.2Ni(OH)₂@S-1-H, and pure S-1 zeolite samples. (F) d-PDFs of Pd-C, Pd-H, and 0.8Pd0.2Ni(OH)₂-H species.

metal species is only 1.6 nm after calcination at 700 °C for 2 h under N₂ atmosphere and all metal species remain to be confined within zeolite crystals. In contrast, the mean particle size of metal species in 0.8Pd0.2Ni(OH)₂/S-1-im dramatically increases to 6.2 nm and most of the metal NPs are located on the outside of zeolites under the same thermal treatment

(Fig. S9†). Moreover, the particle sizes of metal species in 0.8Pd0.2Ni(OH)₂@S-1-H are 1.0, 1.8, and 2.2 nm after calcination in H₂, O₂, and water-vapor treatment at 600 °C, respectively (Fig. S10†). Significantly, all metal species are still encapsulated inside the zeolite crystals after high-temperature thermal treatments. The above results demonstrate that the

0.8Pd0.2Ni(OH)₂@S-1-H possesses superior thermal and hydrothermal stability under high temperatures, suggesting its promising applications under practical harsh conditions.

The catalytic performances of Pd@S-1-H, Pd@S-1-C, and Pd/S-1-im catalysts in methane combustion reactions at a GHSV of 36 000 mL g_{cat}⁻¹ h⁻¹ are shown in Fig. 3A. The methane conversion improves with the increase of the reaction temperature. Among all of the catalysts, the Pd@S-1-H exhibits the highest methane combustion activity with a complete combustion temperature of 390 °C, which is lower than those of Pd@S-1-C (416 °C) and Pd/S-1-im (513 °C), respectively. The significantly improved activity can be attributed to the Pd@S-1-H possessing decreased sizes of Pd species compared with the Pd@S-1-C and Pd/S-1-im, thus exposing more accessible active sites (especially the corner site) for the reactants. In addition, the unique microenvironment of Pd species and highly dispersed Pd^{δ+} ($0 < \delta < 2$) species due to the confinement effect of zeolites may also be a key factor affecting the catalytic performance of methane oxidation reactions. The water resistance ability of Pd@S-1-H is further studied through on-stream tests in the presence of 4.5 vol% water vapor. As shown in Fig. S11,† although the catalytic activity of Pd@S-1-H is slightly decreased under the humidity condition as compared to the dry condition, the Pd@S-1-H catalyst still gives an excellent activity with the complete methane conversion temperature at 415 °C under humidity condition. Such a temperature is close to that under dry condition and much lower than that of Pd/S-1-im, indicating the excellent water vapor resistance of the Pd@S-1-H in methane oxidation reactions. Significantly, the Pd@S-1-H exhibits superhigh long-term stability, and 100% of methane conversion remains unchanged even after 100 h on stream at 470 °C. In contrast, the methane conversion of Pd/S-1-im decreases rapidly from 88.0% to 68.7% within 65 h under the same condition (Fig. 3B). When the reaction temperature decreases to 365 °C, the initial methane conversion of Pd@S-1-H is about 81% and the methane conversion increases gradually as the reaction goes on. Note that the methane conversion can

be stable at around 89% even after 100 h on stream. The above results indicate that the encapsulation of metal species inside zeolites by using the ligand-protected direct H₂ reduction method, can not only enhance the utilization of metal species and catalytic activity, but also remarkably improved the durability during methane combustion reactions.

Catalytic activities of H₂ generation from FA dehydrogenation over various zeolite-supported metal catalysts were evaluated at 333 K without any additive. As shown in Fig. 4A, the Pd@S-1-H exhibited an excellent H₂ generation rate (TOF = 1936 h⁻¹), which is nearly 3 and 7-fold higher than that of Pd@S-1-C (TOF = 651 h⁻¹) and Pd/S-1-im synthesized by impregnation method (TOF = 281 h⁻¹). The activity improvement of Pd@S-1-H can be mainly attributed to the decreased size of Pd species, resulting in more accessible active Pd sites for FA dehydrogenation. The introduction of Ni(OH)₂ species into Pd clusters can further enhance the catalytic activities of H₂ generation from FA dehydrogenation (Fig. 4B). Among all catalysts, the 0.8Pd0.2Ni(OH)₂@S-1-H exhibited the highest H₂ generation rate: 147 mL of CO₂ and H₂ without CO impurity (<10 ppm, Fig. S12†), corresponding to 100% FA conversion (HCOOH = H₂ + CO₂), could be released within 1.67 min at 333 K without any additive, affording a superhigh initial TOF value of 9308 h⁻¹, which is 4.8 and 33 times higher than those over Pd@S-1-H and Pd/S-1-im, respectively. Given that Ni(OH)₂@S-1-H shows no activity for the FA dehydrogenation, the Pd species are responsible for the H₂ generation from FA dehydrogenation. As expected, the H₂ generation rate of 0.8Pd0.2Ni(OH)₂@S-1-H reduction is also about 1.3-fold improvement in comparison with that over 0.8Pd0.2Ni(OH)₂@S-1-C. The TOF value of representative 0.8Pd0.2Ni(OH)₂@S-1-H represents the top level among all of the previously reported heterogeneous catalysts and is even higher than many results obtained with additives under similar catalytic conditions (Table S3†).^{27,48,49} In comparison, the H₂ generation rate over the physical mixture of Pd@S-1-H and Ni(OH)₂@S-1-H are identical to the single Pd@S-1-H, but much lower than that over 0.8Pd0.2Ni(OH)₂@S-1-H.

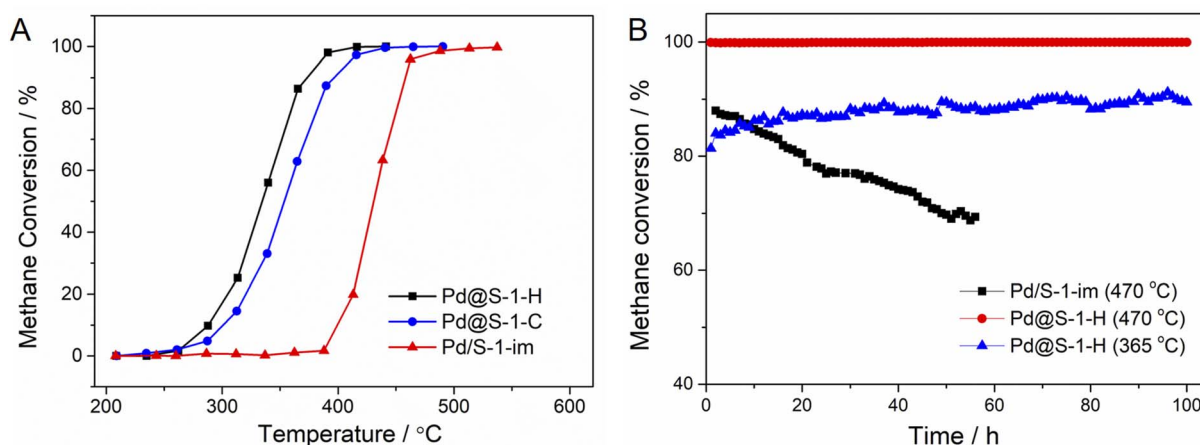


Fig. 3 (A) Catalytic performance of methane combustions over Pd@S-1-H, Pd@S-1-C, and Pd/S-1-im catalysts. Reaction condition: 2% CH₄–20% O₂–78% N₂, gas hourly space velocity (GHSV) = 36 000 mL g_{cat}⁻¹ h⁻¹. (B) Long-term stability of methane combustions over Pd@S-1-H and Pd/S-1-im catalysts with a GHSV of 36 000 mL g_{cat}⁻¹ h⁻¹.



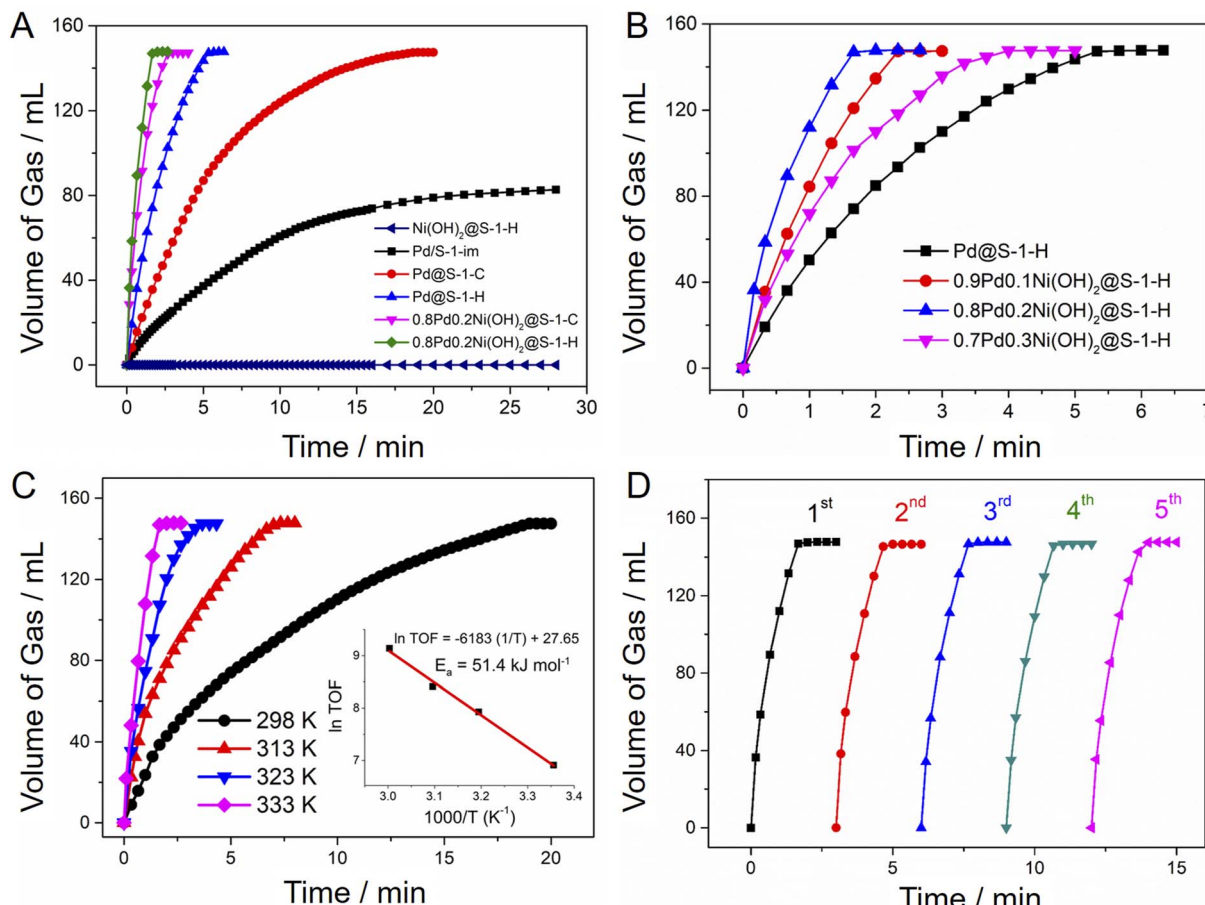


Fig. 4 (A) Volume of the generated gas (CO₂ + H₂) versus time for FA dehydrogenation over (A) Pd@S-1-C, 0.2Pd0.8Ni(OH)₂@S-1-C, Pd@S-1-H, Ni(OH)₂@S-1-H, 0.2Pd0.8Ni(OH)₂@S-1-H, and Pd/S-1-im; (B) Pd@S-1-H and Pd-Ni(OH)₂@S-1-H with various Pd/Ni ratios at 333 K (2 M FA, 3 mmol, $n_{\text{metal}}/n_{\text{FA}} = 0.012$). (C) Volume of the generated gas versus time for FA dehydrogenation at different temperatures over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst (2 M FA, 3 mmol, $n_{\text{metal}}/n_{\text{FA}} = 0.012$). Inset of (C): Arrhenius plot (ln TOF vs. 1000/T). (D) Recycling tests for FA dehydrogenation over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst at 333 K ($n_{\text{metal}}/n_{\text{FA}} = 0.012$). The TOF values of the first run to five runs are 9308, 9789, 8741, 8945, and 9074 h⁻¹, respectively.

(Fig. S13†). The above results revealed that the superior catalytic activity of FA dehydrogenation over 0.8Pd0.2Ni(OH)₂@S-1-H can be mainly ascribed to the formation of ultrasmall and highly-dispersed metal species coupled with the synergistic effect of the Pd–Ni(OH)₂ interface that can remarkably lower the energy barrier of FA dehydrogenation, which is also confirmed by theoretical calculations in our previous work.¹⁹ In addition, a more electron-rich Pd surface favors the C–H activation of the Pd-formate intermediate to produce H₂ and CO₂, accelerating the H₂ generation from FA decomposition.

The catalytic activities of catalysts are highly dependent on the reaction temperature, and the H₂ generation rates improved along with the increase in catalytic temperatures (Fig. 4C and S14, S15†). The apparent activation energy (E_a) of 0.8Pd0.2Ni(OH)₂@S-1-H is calculated as 51.4 kJ mol⁻¹, which is lower than that of Pd@S-1-H catalysts (66.5 kJ mol⁻¹), further indicating the higher activity of H₂ generation over the zeolite-encapsulated bimetallic catalytic system than the corresponding monometallic counterpart. The zeolite-encaged Pd-based metal catalysts also possess excellent recycling stability for H₂

generation from FA dehydrogenation. No degradation of catalytic activities for FA dehydrogenation can be observed over Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H after five cycles at 333 K (Fig. 4D and S16, S17†). Significantly, there is no variation in the morphology and crystallinity of zeolites as well as the particle size and distributions of metal species in spent catalysts after recycling reactions, further confirming the excellent stabilities of the zeolite-encaged metallic catalyst during FA dehydrogenation (Fig. S18–S20†).

Given the zeolite-encaged Pd-based catalysts exhibit excellent catalytic activity and stability in FA dehydrogenation, such catalyst can be expected to have high activity in the Cr(vi) reduction coupled with FA dehydrogenation. Catalytic activities of Cr(vi) reduction in aqueous solution over the representative Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H catalysts are evaluated at 323 K by using K₂Cr₂O₇ and FA as a representative Cr(vi) compound and reducing agent, respectively. As shown in Fig. 5A, B and S21,† the peak at ~350 nm corresponding to Cr(vi) in UV-vis absorption spectroscopy decreases continually as the reaction time is prolonged, and the color of the reaction

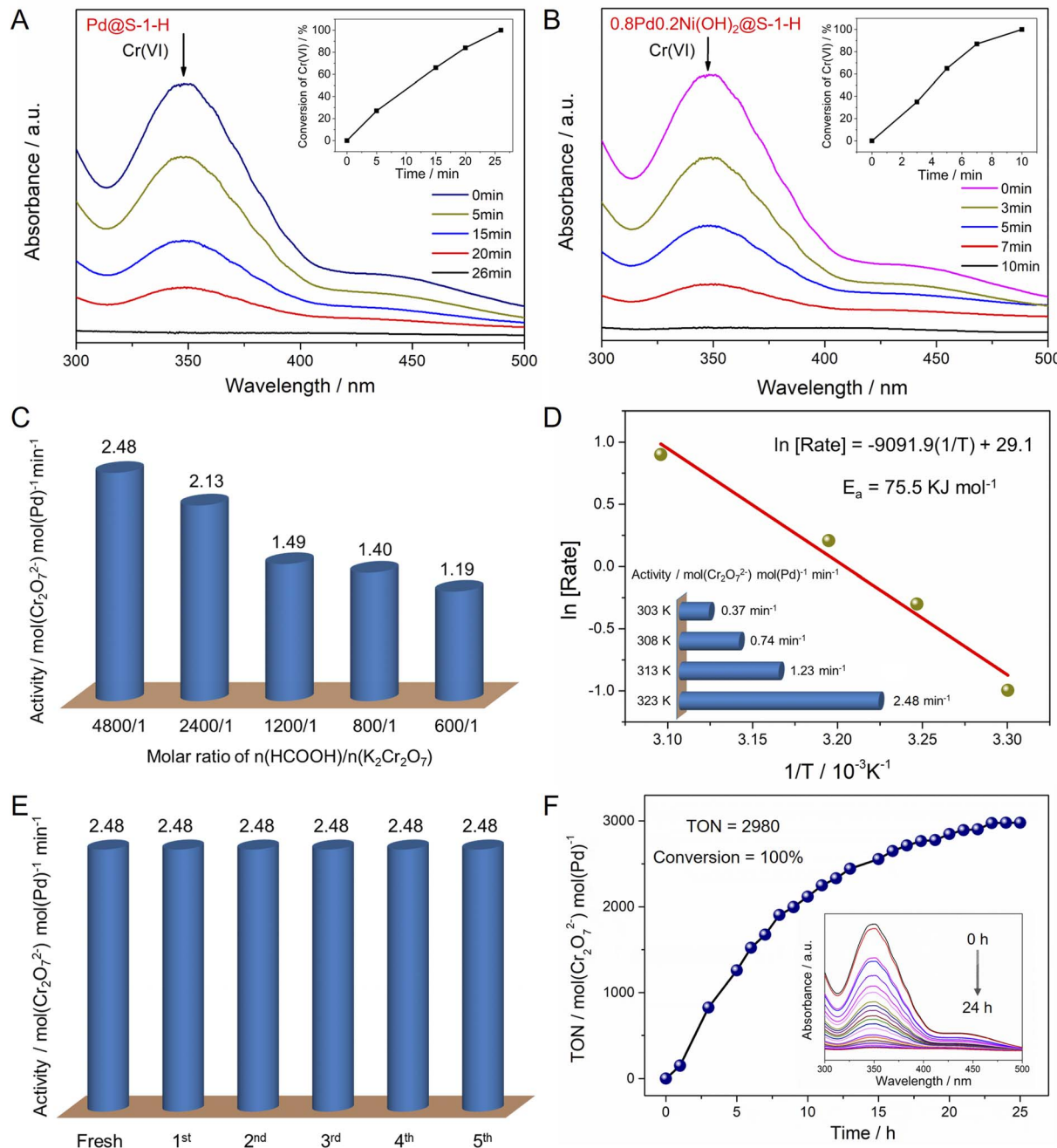


Fig. 5 (A and B) UV-vis absorption spectra for the Cr(VI) reduction by FA over Pd@S-1-H and 0.8Pd0.2Ni(OH)₂@S-1-H catalysts. Catalytic condition: 28 mg catalyst, 0.2 mL K₂Cr₂O₇ (0.1 M), 4 mL HCOOH (6 M) (*n*(HCOOH)/*n*(K₂Cr₂O₇) = 1200/1), 5.8 mL H₂O, *T* = 323 K. (C) Catalytic activity of Cr(VI) reduction over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst with different *n*(HCOOH)/*n*(K₂Cr₂O₇) ratios. Catalytic condition: 28 mg catalysts, 0.05–0.4 mL K₂Cr₂O₇ (0.1 M), 4 mL HCOOH (6 M) (*n*(HCOOH)/*n*(K₂Cr₂O₇) = 4800/1–600/1), 5.95–5.6 mL H₂O, *T* = 323 K. (D) Catalytic activity of Cr(VI) reduction over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst at different temperatures. Catalytic condition: 28 mg catalyst, 0.05 mL K₂Cr₂O₇ (0.1 M), 4 mL HCOOH (6 M), 5.95 mL H₂O, *T* = 303–323 K. (E) Recycling stability of Cr(VI) reduction over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst. Catalytic condition: 28 mg catalyst, 0.05 mL K₂Cr₂O₇ (0.1 M), 4 mL HCOOH (6 M), 5.95 mL H₂O, *T* = 323 K. (F) Durability test of Cr(VI) reduction over 0.8Pd0.2Ni(OH)₂@S-1-H catalyst. Catalytic condition: 28 mg catalyst, 40 mL K₂Cr₂O₇ (0.1 M), 200 mL HCOOH (6 M), 260 mL H₂O, *T* = 323 K.

solution changes from yellow to colorless, indicating the gradual reduction of Cr(VI) in the presence of catalysts (Fig. S22†). 0.02 mmol Cr₂O₇²⁻ can be converted completely to Cr³⁺ within 10 min over 0.8Pd0.2Ni(OH)₂@S-1-H, which is 2.6-fold faster than Pd@S-1-H. In contrast, pure S-1 zeolite was

inactive for the Cr(VI) reduction even with the addition of FA, indicating that the Pd species were responsible for the Cr(VI) reduction (Fig. S23†). In addition, neglectable Cr(VI) conversion can be observed over 0.8Pd0.2Ni(OH)₂@S-1-H when the FA was replaced by H₂ (1 bar), indicating that the *in situ* generated H₂

from FA dehydrogenation is more favorable to the $\text{Cr}_2\text{O}_7^{2-}$ reduction to Cr^{3+} via an H_2 transfer pathway ($\text{Cr}_2\text{O}_7^{2-} + 8\text{H}^+ + 3\text{H}_2 \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O}$) (Fig. S24†). Because the bimetallic Pd-Ni(OH)_2 species is more active for the FA dehydrogenation than monometallic Pd species, not unexpectedly, the $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ exhibits the significantly enhanced activity of cascade $\text{Cr}(\text{vi})$ reduction coupled with FA dehydrogenation than $\text{Pd}@\text{S-1-H}$.

The molar ratio of $n(\text{HCOOH})/n(\text{K}_2\text{Cr}_2\text{O}_7)$ can affect the catalytic activity of $\text{Cr}(\text{vi})$ reduction. The reaction rate of $\text{Cr}(\text{vi})$ reduction doubled with the increase of $n(\text{HCOOH})/n(\text{K}_2\text{Cr}_2\text{O}_7)$ ratio from 600/1 to 4800/1 (1.19 vs. 2.48 min^{-1}) (Fig. 5C). The catalytic activity of $\text{Cr}(\text{vi})$ reduction over $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ is dependent on the reaction temperature, and the reduction rate of $\text{Cr}(\text{vi})$ enhances with the increase of reaction temperatures, affording apparent activation energy of 75.5 kJ mol^{-1} (Fig. 5D and S25†). Moreover, the catalytic activity of $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ remains unchanged after five consecutive cycles, reflecting its excellent recycling stability during the $\text{Cr}(\text{vi})$ reductions (Fig. 5E and S26†). The Pd atoms in the spent $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ catalyst are uniformly dispersed throughout the S-1 crystals, which is much similar to that in a fresh one, indicating the high stability of $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ during $\text{Cr}(\text{vi})$ reduction reactions (Fig. S27†). Significantly, the $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ also possesses superior durability for the cascade reduction of $\text{Cr}(\text{vi})$ species, 4 mmol $\text{K}_2\text{Cr}_2\text{O}_7$ can be completely converted to Cr^{3+} within 24 h over only 28 mg of $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ catalyst, affording a high turnover number of 2980 $\text{mol}(\text{Cr}_2\text{O}_7^{2-}) \text{mol}(\text{Pd})^{-1}$ at 323 K (Fig. 5F), which represents the best stability among all of the previously-reported heterogeneous catalysts (Table S4†).^{50,51}

Conclusions

In summary, we prepared a series of zeolite-encaged Pd -based pseudo-single atoms via a facile and energy-saving ligand-protected direct H_2 reduction method. In comparison with the conventional calcination-reduction treatment, the direct H_2 reduction process is more energy-saving and able to further reduce the size and improve the dispersity of Pd -based metal species within zeolite micropores. Thanks to the extremely high metal dispersion and confinement effect of zeolites, the $\text{Pd}@\text{S-1-H}$ exhibited superior activity and durability in methane combustion with a complete combustion temperature of 390 °C that is much lower than Pd/S-1-im (513 °C). By virtue of the synergy between the interface of Pd and Ni(OH)_2 species, the $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ catalyst exhibited a superhigh H_2 generation rate from FA decomposition, affording a TOF value up to 9308 h^{-1} at 333 K, which is 33 times higher than that of Pd/S-1-im . Significantly, the zeolite-encaged metal catalysts are for the first time used for the $\text{Cr}(\text{vi})$ reduction coupling with FA dehydrogenation reaction, and the $0.8\text{Pd}0.2\text{Ni(OH)}_2@\text{S-1-H}$ gave a record turnover number of 2980 $\text{mol}(\text{Cr}_2\text{O}_7^{2-}) \text{mol}(\text{Pd})^{-1}$ at 323 K. This work provides references for the preparation of high-performance zeolite-encaged atomically dispersed metal catalysts. The superhigh metal utilization,

superior catalytic activity and excellent stability of such catalyst suggest important prospects for practical applications in the field of efficient hydrogen storage and pollutant disposal in the future.

Author contributions

Q. S., J. Y. and N. W. conceived the idea, designed the experiments, and supervised this work. K. Z., N. W., Y. M., T. Z., P. Y., and Q. S. performed experiments and analysed the data. Q. S. wrote the manuscript, Q. S. and J. Y. revised the manuscript. All the authors discussed the results and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 L.-H. Chen, M.-H. Sun, Z. Wang, W. Yang, Z. Xie and B.-L. Su, *Chem. Rev.*, 2020, **120**, 11194–11294.
- 2 M. Dusselier and M. E. Davis, *Chem. Rev.*, 2018, **118**, 5265–5329.
- 3 Y. Li and J. Yu, *Nat. Rev. Mater.*, 2021, **6**, 1156–1174.
- 4 B. M. Weckhuysen and J. Yu, *Chem. Soc. Rev.*, 2015, **44**, 7022–7024.
- 5 Y. Chai, W. Dai, G. Wu, N. Guan and L. Li, *Acc. Chem. Res.*, 2021, **54**, 2894–2904.
- 6 L. Liu and A. Corma, *Chem. Rev.*, 2018, **118**, 4981–5079.
- 7 L. Liu and A. Corma, *Nat. Rev. Mater.*, 2021, **6**, 244–263.
- 8 Z. Qu and Q. Sun, *Inorg. Chem. Front.*, 2022, **9**, 3095–3115.
- 9 Q. Sun, N. Wang and J. Yu, *Adv. Mater.*, 2021, **33**, 2104442.
- 10 H. Wang, L. Wang and F.-S. Xiao, *ACS Cent. Sci.*, 2020, **6**, 1685–1697.
- 11 S.-M. Wu, X.-Y. Yang and C. Janiak, *Angew. Chem., Int. Ed.*, 2019, **58**, 12340–12354.
- 12 D. Farrusseng and A. Tuel, *New J. Chem.*, 2016, **40**, 3933–3949.
- 13 N. Wang, Q. Sun and J. Yu, *Adv. Mater.*, 2019, **31**, 1803966.
- 14 Y. Chai, G. Wu, X. Liu, Y. Ren, W. Dai, C. Wang, Z. Xie, N. Guan and L. Li, *J. Am. Chem. Soc.*, 2019, **141**, 9920–9927.
- 15 S. Goel, S. I. Zones and E. Iglesia, *J. Am. Chem. Soc.*, 2014, **136**, 15280–15290.
- 16 L. Liu, M. Lopez-Haro, C. W. Lopes, C. Li, P. Concepcion, L. Simonelli, J. J. Calvino and A. Corma, *Nat. Mater.*, 2020, **3**, 628–638.



17 Y. Liu, Z. Li, Q. Yu, Y. Chen, Z. Chai, G. Zhao, S. Liu, W.-C. Cheong, Y. Pan, Q. Zhang, L. Gu, L. Zheng, Y. Wang, Y. Lu, D. Wang, C. Chen, Q. Peng, Y. Liu, L. Liu, J. Chen and Y. Li, *J. Am. Chem. Soc.*, 2019, **141**, 9305–9311.

18 Y. Ma, S. Song, C. Liu, L. Liu, L. Zhang, Y. Zhao, X. Wang, H. Xu, Y. Guan, J. Jiang, W. Song, Y. Han, J. Zhang and P. Wu, *Nat. Catal.*, 2023, **6**, 506–518.

19 Q. Sun, N. Wang, Q. Bing, R. Si, J. Liu, R. Bai, P. Zhang, M. Jia and J. Yu, *Chem.*, 2017, **3**, 477–493.

20 Q. Sun, N. Wang, Q. Fan, L. Zeng, A. Mayoral, S. Miao, R. Yang, Z. Jiang, W. Zhou, J. Zhang, T. Zhang, J. Xu, P. Zhang, J. Cheng, D.-C. Yang, R. Jia, L. Li, Q. Zhang, Y. Wang, O. Terasaki and J. Yu, *Angew. Chem., Int. Ed.*, 2020, **59**, 19450–19459.

21 C. Wang, E. Guan, L. Wang, X. Chu, Z. Wu, J. Zhang, Z. Yang, Y. Jiang, L. Zhang, X. Meng, B. C. Gates and F.-S. Xiao, *J. Am. Chem. Soc.*, 2019, **141**, 8482–8488.

22 J. Zhang, L. Wang, B. Zhang, H. Zhao, U. Kolb, Y. Zhu, L. Liu, Y. Han, G. Wang, C. Wang, D. S. Su, B. C. Gates and F.-S. Xiao, *Nat. Catal.*, 2018, **1**, 540–546.

23 M. Choi, Z. Wu and E. Iglesia, *J. Am. Chem. Soc.*, 2010, **132**, 9129–9137.

24 S. Goel, Z. J. Wu, S. I. Zones and E. Iglesia, *J. Am. Chem. Soc.*, 2012, **134**, 17688–17695.

25 L. Liu, U. Diaz, R. Arenal, G. Agostini, P. Concepcion and A. Corma, *Nat. Mater.*, 2017, **16**, 132–138.

26 N. Wang, Q. Sun, R. Bai, X. Li, G. Guo and J. Yu, *J. Am. Chem. Soc.*, 2016, **138**, 7484–7487.

27 Q. Sun, B. W. J. Chen, N. Wang, Q. He, A. Chang, C.-M. Yang, H. Asakura, T. Tanaka, M. J. Hulse, C.-H. Wang, J. Yu and N. Yan, *Angew. Chem., Int. Ed.*, 2020, **59**, 20183–20191.

28 Y. Chen, S. Ji, C. Chen, Q. Peng, D. Wang and Y. Li, *Joule*, 2018, **2**, 1242–1264.

29 M. Flytzani-Stephanopoulos and B. C. Gates, *Annu. Rev. Chem. Biomol. Eng.*, 2012, **3**, 545–574.

30 B. Qiao, A. Wang, X. Yang, L. F. Allard, Z. Jiang, Y. Cui, J. Liu, J. Li and T. Zhang, *Nat. Chem.*, 2011, **3**, 634–641.

31 Q. Sun, N. Wang, T. Zhang, R. Bai, A. Mayoral, P. Zhang, Q. Zhang, O. Terasaki and J. Yu, *Angew. Chem., Int. Ed.*, 2019, **58**, 18570–18576.

32 X. Yang, A. Wang, B. Qiao, J. Li, J. Liu and T. Zhang, *Acc. Chem. Res.*, 2013, **46**, 1740–1748.

33 Z. Hou, L. Dai, J. Deng, G. Zhao, L. Jing, Y. Wang, X. Yu, R. Gao, X. Tian, H. Dai, D. Wang and Y. Liu, *Angew. Chem., Int. Ed.*, 2022, **61**, e202201655.

34 T. Li, Y. Yao, Z. Huang, P. Xie, Z. Liu, M. Yang, J. Gao, K. Zeng, A. H. Brozena, G. Pastel, M. Jiao, Q. Dong, J. Dai, S. Li, H. Zong, M. Chi, J. Luo, Y. Mo, G. Wang, C. Wang, R. Shahbazian-Yassar and L. Hu, *Nat. Catal.*, 2021, **4**, 62–70.

35 A. W. Petrov, D. Ferri, F. Krumeich, M. Nachtegaal, J. A. van Bokhoven and O. Kröcher, *Nat. Commun.*, 2018, **9**, 2545.

36 W. Wang, W. Zhou, W. Li, X. Xiong, Y. Wang, K. Cheng, J. Kang, Q. Zhang and Y. Wang, *Appl. Catal., B*, 2020, **276**, 119142.

37 S. Guan, Y. Liu, H. Zhang, R. Shen, H. Wen, N. Kang, J. Zhou, B. Liu, Y. Fan, J. Jiang and B. Li, *Adv. Sci.*, 2023, **10**, 2300726.

38 C. Lang, Y. Jia and X. Yao, *Energy Storage Mater.*, 2020, **26**, 290–312.

39 M. Navlani-García, K. Mori, Y. Kuwahara and H. Yamashita, *NPG Asia Mater.*, 2018, **10**, 277–292.

40 Q. Sun, N. Wang, Q. Xu and J. Yu, *Adv. Mater.*, 2020, **32**, 2001818.

41 J. Eppinger and K.-W. Huang, *ACS Energy Lett.*, 2017, **2**, 188–195.

42 X. Gu, Z.-H. Lu, H.-L. Jiang, T. Akita and Q. Xu, *J. Am. Chem. Soc.*, 2011, **133**, 11822–11825.

43 Z. Li and Q. Xu, *Acc. Chem. Res.*, 2017, **50**, 1449–1458.

44 A. Zhang, J. Xia, Q. Yao and Z.-H. Lu, *Appl. Catal., B*, 2022, **309**, 121278.

45 Z. H. Farooqi, M. W. Akram, R. Begum, W. Wu and A. Irfan, *J. Hazard. Mater.*, 2021, **402**, 123535.

46 P. Veerakumar and K.-C. Lin, *Chemosphere*, 2020, **253**, 126750.

47 T. Iida, D. Zanchet, K. Ohara, T. Wakihara and Y. Román-Leshkov, *Angew. Chem., Int. Ed.*, 2018, **57**, 6454–6458.

48 S.-J. Li, Y.-T. Zhou, X. Kang, D.-X. Liu, L. Gu, Q.-H. Zhang, J.-M. Yan and Q. Jiang, *Adv. Mater.*, 2019, **31**, 1806781.

49 F.-Z. Song, Q.-L. Zhu, N. Tsumori and Q. Xu, *ACS Catal.*, 2015, **5**, 5141–5144.

50 S. Dai, X. Wu, J. Zhang, Y. Fu and W. Li, *Chem. Eng. J.*, 2018, **351**, 959–966.

51 L. Tan, T. Ray Jones, J. Poitras, J. Xie, X. Liu and G. Southam, *J. Hazard. Mater.*, 2020, **398**, 122945.

