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1. Introduction

The production of ethylene (C_2H_4), which is at the core of the petrochemical industry, is generally accompanied by the production of a small amount of acetylene (C_2H_2). This is one of the key reasons for catalyst poisoning and products oligomerization, which may exert negative effects on the production of C_2H_4 . The selective hydrogenation of C_2H_2 has been considered as an effective method for removing the trace C_2H_2 due to the simple process, low energy consumption and low C_2H_4 loss.¹⁻⁴ The core mechanism of this method is the complete conversion of C_2H_2 to C_2H_4 *via* an addition reaction. However, the side reactions with oligomer production under improper control have always been the main bottleneck of C_2H_2 hydrogenation. One of the solutions to solve this problem is the development of high selective catalysts that could effectively reduce the occurrence of side reactions.

Although Fe, Cu, Zn and other non-noble metal-based catalysts have long been used in hydrogenation reactions,⁵⁻⁷ satisfactory

Anchoring ultrasmall Pd nanoparticles by bipyridine functional covalent organic frameworks for semihydrogenation of acetylene[†]

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Acetylene hydrogenation is a well-accepted solution to reduce by-products in the ethylene production process, while one of the key technical difficulties lies in developing a catalyst that can provide highly dispersed active sites. In this work, a highly crystalline layered covalent organic framework (COF) material (TbBpy) with excellent thermal stability was synthesized and firstly applied as support for ultrasmall Pd nanoparticles to catalyze acetylene hydrogenation. 100% of C_2H_2 conversion and 88.2% of C_2H_4 selectivity can be obtained at 120 °C with the space velocity of 70 000 h⁻¹. The reaction mechanism was elucidated by applying a series of characterization techniques and theoretical calculation. The results indicate that the coordination between Pd and N atom in the bipyridine functional groups of COFs successfully increased the dispersibility and stability of Pd particles, and the introduction of COFs not only improved the adsorption of acetylene and H₂ onto catalyst surface, but enhanced the electron transfer process, which can be responsible for the high selectivity and activity of catalyst. This work, for the first time, reported the excellent performance of Pd@TbBpy as a catalyst for acetylene hydrogenation and will facilitate the development and application of COFs materials in the area of petrochemicals.

activity is hard to obtain at low temperatures. In contrast, the substitution by Pd, a typical noble metal, overcomes the limit of temperature successfully.8 Pd-based catalysts with unique electronic properties often exhibit outstanding advantages in catalytic activity for selective hydrogenation of alkynes.9,10 Especially the research on the electronic structure between Pd atoms and carriers has always been a hot topic pursued by people. For instance, Yang et al. dispersed single atom Pd onto nanoscale TiO2 through self-assembly, resulting in a TOF of over 8000 h^{-1} and a selectivity of over 91% for styrene.11 At the same time, the doping of Pd atoms also contributes to the dissociation of H₂ on the catalyst surface, improving hydrogenation activity.12,13 However, the high cost and easy agglomeration during reaction still remain as major concerns in the real application. A well-recognized strategy to solve these issues is the impregnation of Pd onto high-surface-area carrier which can not only increase the dispersibility of particles, but also enhance the thermal and chemical stability. For the selective hydrogenation of C2H2, the most studied catalyst carrier for Pd was alumina due to its low cost.14-16 However, the lacking of porous structure brings certain difficulties in improving mass transfer, dispersing active sites, as well as enhancing anti-coking properties.

Recently, a new class of materials *i.e.* metal–organic frameworks (MOFs) have attracted increasing attention as the catalyst carrier in many catalytic reactions due to their unique crystalline porosity, flexible and tailorable properties.^{17,18} Metal–

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organic frameworks (MOFs) with unique regular channels are more likely to show strong adsorption capacity for C₂H₂ in mixed systems containing unsaturated hydrocarbon.19,20 Recent studies have shown that MOFs usually need to be combined with other mesoporous materials to improve the reusability and stability of catalysts. For instance, Li et al. have used mesoporous silica for stabilization Pd@ZIF-8 to achieve a conversion rate of 98% for phenylacetylene and a selectivity of 96% for styrene.²¹ It can be seen that the instability of metal coordination bonds in the structure of MOFs still makes their widespread application in industry impossible. In addition, graphene, as carbon materials with porous structures and large specific surface areas, are often used as carriers for fixing metal atoms through local coordination, effectively solving the problem of metal atom dispersion. By way of illustration, Zhou et al. successfully introduced Pd atoms onto nitrogen-doped graphene using freeze-drying assisted method. The local coordination from Pd-N hindered the aggregation of Pd particles, achieving a result of 99% acetylene conversion rate and 93.5% ethylene selectivity.22 Furthermore, Huang et al. dispersed Pd atoms onto Graphene through Pd-C bond anchoring, obtaining 100% conversion and 90% ethylene selectivity.23 However, the underdeveloped pore structure of graphene limits its mass transfer efficiency in heterogeneous reactions. Therefore, it is impressive that covalent organic frameworks (COFs), as an emerging crystalline organic porous material, not only have the advantages of the aforementioned properties,24,25 but also are more unique in terms of thermal and chemical stability. More significantly, as a well-defined and predictable functional material, COFs can be endowed with different catalytic functions by varying the means of post-synthesis. Therefore, it's reasonable to conduct that COFs might be an ideal catalyst carrying material for Pd in the reaction of C₂H₂ hydrogenation.

In this work, a novel bipyridine-functionalized COFs (TbBpy) was developed via the reaction of 1,3,5-triformylbenzene (Tb) with 2,2'-bipyridine-5,5'-diamine (Bpy), applying a solvothermal synthesis method.26 Pd supported on TbBpy (Pd@TbBpy) was then obtained by a simple impregnation process and applied as catalyst for the selective C2H2 hydrogenation. A series of techniques were carried out to characterize the structure of Pd@TbBpy. The catalytic activity and long-term stability were systematically investigated under different conditions. To further elucidate the reaction mechanism and the function of TbBpy, in situ diffusion reflectance infrared Fourier transform spectra (DRIFTS) and the quantum chemical calculations were used to probe the surface processes. This work for the first time reported the excellent properties of TbBpy as carrier for Pd catalyst, which will facilitate the development and application of Pd in industrial C₂H₂ hydrogenation process.

2. Experimental section

2.1 Chemical and materials

All the chemicals used in the experiments were at A.R. grade. 1,3,5-Triformylbenzene and 2,2'-bipyridyl-5,5'-diamine were obtained from Jilin Chinese Academy of Sciences-Yanshen Technology Co., Ltd. Ethanol, mesitylene, acetic acid, *N*,*N*-dimethylformamide, tetrahydrofuran, and acetone were obtained from Aladdin. Palladium acetate $(Pd(OAc)_2)$ and NaBH₄ were purchased from Tianjing Fuchen Chemical Reagents Factory. And the water used in the experiment was deionized water.

2.2 Catalyst preparation

The preparation process of Pd@TbBpy was shown in Fig. 1.

2.2.1 Synthesis of TbBpy. TbBpy was synthesize *via* the Schiff-base reaction of 1,3,5-triformylbenzene (0.2 mmol) with 2,2'-bipyridyl-5,5'-diamine (0.3 mmol). Two organic monomers were dissolved in the mixture of ethanol and mesitylene (with volume ratio of 1 : 4) and transferred into a glass ampule, then aqueous acetic acid (AcOH) of 6.0 mol L⁻¹ was added, and the glass ampoule was quickly frozen in a liquid nitrogen bath. The internal pressure of the bottle is evacuated and flame-sealed, and after defrosting at room temperature, it was further heated to 120 °C. After three days, the yellow solid in the ampule was collected and washed with excess *N*,*N*-dimethylformamide (DMF), tetrahydrofuran (THF), and acetone, and then dried under vacuum at 80 °C for 24 h to yield the as-prepared TbBpy. The details about TbBpy synthesis are provided in the ESI (Tables S1–S3†).

2.2.2 Synthesis of Pd@TbBpy. In order to improve the dispersion and anchor of Pd particles based on TbBpy, achieving the goal of high ethylene selectivity, Pd based catalysts were prepared using traditional impregnation methods.^{27,28} Here, two reduction methods are provided to prepare catalysts to obtain high selectivity catalysts:

The method using NaBH₄ as reducing agent is as follows: TbBpy (100 mg) was dispersed in 10 mL of ethanol, and then 0.76 mL of Pd(OAc)₂ (1 mg mL⁻¹) was added slowly under continuous stirring. After 24 h, NaBH₄ (2 mol L⁻¹, 2 mL) was added dropwise into the mixture and stirred for 2 h. Then, the residue was collected through centrifugation, washed three times with ethanol (10 mL), and dried at 60 °C under vacuum for 24 h.

The method using H_2 as reducing agent is as follows: TbBpy (100 mg) was dispersed in 10 mL of ethanol, and then 0.76 mL of Pd(OAc)₂ (1 mg mL⁻¹) was added slowly under continuous stirring. After 24 h, the residue was collected through centrifugation, washed three times with ethanol (10 mL), and dried at 60 °C under vacuum for 24 h. The catalyst was reduced in H_2/N_2 (20% H_2 , 50 mL min⁻¹) gas flow at 80 °C for 1 h, and then Pd@TbBpy was obtained.

2.3 Catalyst characterization

The crystalline phase of materials was analyzed by Powder X-ray diffraction (PXRD, X'Pert PRO MPD, PANalytical) with Cu K α radiation ($\lambda = 1.54$ Å). The structure of materials can be characterized by The Fourier transform infrared (FT-IR, Nicolet IS 10, Thermo Scientific) spectroscopy. The specific surface area and pore-size distribution were calculated using the Brunauer-Emmett-Teller (BET) method and non-local density functional theory (NLDFT) modelling based on N₂ adsorption data. The nitrogen adsorption and desorption isotherms were tested at 77 K on a Micromeritics TriStar II system. The samples were



degassed under vacuum at 150 °C for 6 h before the measurements. The surface morphology of TbBpy and Pd@TbBpy was analyzed by high-resolution transmission electron microscopy (TEM, JEM-2100, JEOL). The sample should be pre-treated before the test. Firstly, it was ground and sonicated, and then the sample was dispersed and casted on a thin carbon-coated TEM grid (copper), and finally, it can be transferred to the filter and the solvent (i.e. ethanol) will evIaporate. The STEM image was recorded with a JEOL JEM-2100 F high-resolution transmission electron microscope equipped with energy dispersive X-ray spectroscopy (EDS) operating at 200 kV. The chemical properties of Pd(II)@TbBpy and Pd@TbBpy were analyzed by X-ray photoelectron spectra (XPS, K-Alpha, Thermo Scientific) equipped with Mono Al Ka. The binding energy calibration based on surface contamination carbon C1s (284.8 eV). The thermal properties of TbBpy and Pd@TbBpy can be analyzed by Thermogravimetric Analyzer (TGA, 5500, TA). The temperature was raised from 26 °C to 800 °C in N2 atmosphere at a heating rate of 10 $^{\circ}$ C min⁻¹. The quantitative elemental analysis for Pd was performed by inductively coupled plasma spectrometry (ICP-AES) on an Agilent 720ES ICP-OES/MS. Temperature programmed desorption of H₂ (H₂-TPD) experiments were carried out on xianguan TP-5080. Typically, 100 mg catalyst was pretreated in 10% H2-90% Ar at 120 °C for 2 h and then returned to room temperature. The temperature programmed desorption experiment was carried out in pure Ar atmosphere (flow rate of 30 ml min⁻¹) at a heating rate of 10 $^{\circ}$ C min⁻¹ to 600 °C.

The material transformation mechanism of C₂H₂ hydrogenation on Pd@TbBpy was explored by in situ diffusion reflectance infrared Fourier transform spectra (DRIFTS, Nicolet iS50, Thermo Scientific) equipped with a high-temperature in situ diffuse reflection cell and a liquid nitrogen cooled MCT detector. The small porcelain crucible is filled with Pd@TbBpy powder and carefully leveled, and the solid powder were first pretreated in 20% H₂/N₂ at 150 °C for 2 h, and cooled to 120 °C in pure He at a flow rate of 10 mL min⁻¹ for 30 min to ensure a stable condition and the system purged of H₂, then the background spectrum was recorded. Thereafter, the Pd@TbBpy powder was exposed to reactant stream (20% H₂/N₂, 20% C₂H₂/ N2 and pure He) flow for 100 min. During the process, a series of spectral data in the range of 4000–650 cm⁻¹ at different minutes were obtained.

All density functional theory (DFT) calculations have been performed by using the Perdew-Burke-Ernzerhof (PBE)29 formula in generalized gradient approximation (GGA) using VASP.^{30,31} The ion nucleus was described by selecting the projection enhanced wave (PAW) potential.32 Take valence electrons into account using a plane wave basis set with a kinetic energy cutoff of 450 eV. Partial occupancies of the Kohn-Sham orbitals were allowed using the Gaussian smearing method and a width of 0.05 eV. The electronic energy was considered self-consistent when the energy change was smaller than 10⁻⁵ eV. A geometry optimization was considered convergent when the energy change was smaller than 0.05 eV $Å^{-1}$. The Brillouin zone is sampled with $1 \times 1 \times 1$ Gamma *k*-points.³³

2.4 Acetylene hydrogenation

The gas phase selective hydrogenation of C_2H_2 was performed in a heated fixed-bed microreactor under continuous flow conditions. After volume controlling by gas pressure reducer and flowmeter, the gases (*i.e.*, 20% C_2H_2/N_2 acetylene, 20% H_2/N_2 N_2 hydrogen and pure He with a H_2/C_2H_2 -ratio of 2:1) have been fed into the reaction pipe (Φ 8 mm \times 200 mm) loaded with Pd@TbBpy. In a typical experiment, 20 mg of the catalyst was diluted with 100 mg crushed quartz sand and filled in the middle of the reactor pipe. Before the reaction, Pd@TbBpy was activated for 2 h at 20% H_2/N_2 with a flow rate of 50 mL min⁻¹ and 150 °C, and then cooled to room temperature at pure He with a flow rate of 10 mL min⁻¹. The outlet gas stream has been collected for analysis by a gas chromatography (GC7890A, Agilent, USA) equipped with a thermal conductivity detector and a long silica packed column (Φ 0.25 mm \times 50 mm, Al₂O₃/ Na₂SO₄ coating). The conversion of C₂H₂ and selectivity of C₂H₄ for the catalytic reactions can be calculated as bellow, *i.e.*,

$$\text{Conversion}_{C_{2}H_{2}}(\%) = \frac{C_{C_{2}H_{2_{\text{inlet}}}} - C_{C_{2}H_{2_{\text{outlet}}}}}{C_{C_{2}H_{2_{\text{inlet}}}}} \times 100 \quad (1a)$$

Selectivity_{C₂H₄}(%) =
$$\frac{C_{C_2H_{4_{outlet}}}}{C_{C_2H_{2_{inlet}}} - C_{C_2H_{2_{outlet}}}} \times 100$$
 (1b)

3. Results

3.1 Characterization of Pd@TbBpy catalysts

The long-range order of TbBpy can be characterized by XRD. At first, the effect of solvent species on the crystal structure of TbBpy were explored (Fig. S1[†]), it's clear that no obvious characteristic peaks of TbBpy can be observed when 1,4-dioxane and n-butanol were applied as solvents. While the replacement by mesitylene and ethanol clearly promoted the production of TbBpy, with the characteristic peak showing at $2\theta = 24.8^{\circ}$, which can be assigned to the (001) plane of TbBpy, corresponding to the layered stacking structure in the COFs. It might be resulted from the fact that the solvents would determine the solubility as well as the condensation reaction rate of the two organic monomers.³⁴ Besides, the ratio of solvent components would influence the reaction balance between skeleton formation and crystallization,35 and a much better crystal structure has been observed in Fig. S2[†] at volume ratio of ethanol/mesitylene of 1:4. Further, the concentration of acetic acid aqueous solution was adjusted to improve the crystallinity of TbBpy. As can be seen in Fig. S3,† when the concentration was 6 mol L^{-1} , the crystal morphology is relatively complete. While as it further increased, the crystallinity of TbBpy decreased instead. This is probably because the high concentration of acetic acid disrupted the rate of reversible polycondensation in the process of linking organic monomers through polycondensation and orderly assembly of organic monomers, which led to a incomplete self-healing of the structure.35

The XRD patterns of TbBpy and Pd@TbBpy synthesized under the optimal condition are shown in Fig. 2a, the typical diffraction peaks of TbBpy at 2θ of 3.53° and 24.3° can be clearly seen, which can be attributed to the crystal planes of (100) and (001), respectively. For Pd@TbBpy, only diffraction peak of Pd corresponding to (111) plane at 2θ of 39.94° can be observed,³⁶ suggesting the successful loading of Pd particles onto the TbBpy. While the disappearance of peaks of (100) and (001) planes might be ascribed to the partial collapse of internal structure of TbBpy after the loading of Pd.³⁷

To further confirm that TbBpy is composed of 1,3,5-triphenylcarbaldehyde and 5,5-diamino-2,2-bipyridine connecting to each other through Schiff base reaction, the functional group structure of TbBpy was determined by FT-IR (Fig. S4†). The FT-IR spectrum of TbBpy shows a C=N stretching vibration peak at 1618 cm⁻¹, which means that the polycondensation reaction has occurred. When Pd nanoparticles were embedded in the pores of TbBpy, as shown in Fig. 2b, a strong vibrational peak (C=N stretching vibration) was still observed at a wavelength of 1618 cm⁻¹, and the intensity was weakened, indicating that the immersion of Pd may cause the collapse of layered structure inside TbBpy, which is in accordance with the results of XRD analysis.

The surface areas (BET) of TbBpy and Pd@TbBpy have been calculated by the BET method, which were 76.10 and $36.32 \text{ m}^2 \text{ g}^{-1}$, respectively (Fig. 2c). The pore size distribution embedded in Fig. 2c shows that the average pore size of TbBpy and Pd@TbBpy are 5.8 nm and 5.12 nm, respectively. The decrease in pore size may resulted from the block by Pd particles, which further demonstrates the successful loading onto TbBpy.²⁶ Comparing with other COFs, such as COF-LZU1 (410 m² g⁻¹)²⁶ and TpPa-1 (484 m² g⁻¹),³⁸ TbBpy has a much lower BET, which might be resulted from its unique structure with inhomogeneous mesopores and lower long-range ordered.

The morphological structures of the Pd@TbBpy were characterized by TEM and STEM scanning. The random distribution of Pd particles can be clearly observed in Fig. 2d, and the average diameter of the Pd loaded is about 2.4 nm (Fig. 2f). Moreover, a 0.211 nm of interplanar crystalline space was observed in HRTEM image (Fig. 2e), which matched well with (111) crystal plane of Pd. The EDS elemental mapping of Pd@TbBpy were also analyzed to reveal the distribution of Pd, C and N. The EDS mapping clearly showed that Pd homogeneously distributed on the TbBpy surface, which further suggested the well dispersion of Pd nanoparticles. In addition, inductively coupled plasma spectroscopy (ICP) analysis indicated that the Pd content of Pd@TbBpy is 0.72 wt%, which deviates slightly from the theoretical Pd content (0.75 wt%) we applied.

The thermal stability of TbBpy and Pd@TbBpy was evaluated by TGA. As shown in Fig. S5,† the mass loss of the material was relatively small when the temperature was lower than 400 °C. Then, the heat cracks started to appear in TbBpy and Pd@TbBpy at 400–550 °C, with the mass loss reaching to 20– 30%. And the further rising of temperature to 800 °C gradually led to the continuous increase in ash content. These phenomenon show that TbBpy can maintain good thermal stability below 400 °C, and the load of Pd will not affect the thermal stability.

The XPS analysis was conducted to elucidate the interactions between Pd and N groups in the TbBpy. At first, from the total survey scan of XPS (Fig. 3a), the peaks of O, C, N, and Pd can be found. Specifically, in the N 1s spectrum shown in Fig. 3b, the peaks at 399.08 eV and 398.53 eV can be ascribed to pyrrole-N and pyridine-N, respectively, which are basic units of N groups in TbBpy. Besides, in the Pd 3d spectrum (Fig. 3c), the peaks at binding energy of 335.78 eV and 340.88 eV corresponded to Pd 3d_{5/2} and Pd 3d_{3/2}, respectively. The results indicate that Pd(n) has been successfully reduced to Pd(0), which was then loaded onto TbBpy.39 Notably, the binding energy peak at 338.0 eV can be attributed to the formation of PdO₂ caused by Pd@TbBpy contacting with air, which is consistent with the result of PdO/Cel-xPdxO₂-δ reported in the literature.40 For comparison, the XPS analysis of un-reduced $Pd(\pi)$, *i.e.* $Pd(\pi)$ TbBpy was also conducted, with the Pd 3d spectrum showing in Fig. 3d, with a binding energy of 337.58 eV assigning to the Pd 3d_{5/2} peak and a binding energy of 342.81 eV assigned to the Pd 3d_{3/2} peak. Compared with the characteristic peak of $Pd(OAc)_2$ (338.4 eV), the binding energy peak of Pd(II)(a)TbBpy at 337.58 eV is reduced by about 0.62 eV, which indicates that the Pd(II) species accepts the lone pair of electrons from the N atom in the imine group to form a strong coordination bond.26



Fig. 2 The (a) XRD profile, (b) FT-IR spectra, and (c) N₂ adsorption–desorption isotherms of TbBpy and Pd@TbBpy; the (d) TEM image and (e) HR-TEM image of Pd@TbBpy. (f) The Pd particle size frequency distribution histogram; (g) the STEM image and (h) corresponding elemental mapping image of Pd@TbBpy.

3.2 Catalytic performance over Pd@TbBpy catalysts

Normally, an ideal C_2H_2 hydride Pd-based catalyst should meet the following requirements: (1) the lower Pd loading; (2) the complete conversion of C_2H_2 at lower temperature and higher C_2H_4 selectivity; (3) the stable catalytic activity. With these indicators as reference, we systematically investigated the catalytic performance of the catalyst for C_2H_2 hydrogenation.

At first, H_2 and NaBH₄ were applied as Pd(II) reductants to investigate the effect of reductant phase on the catalytic activity (Fig. S6†). It turns out that both of the C₂H₂ conversion rates reached 100% with two reductants. While the C₂H₄ selectivity of NaBH₄-reduced catalyst was 52.2%, being higher than that of H₂-reduced catalyst, which was only 43.5%. This might be explained by the fact that NaBH₄ could penetrate into the pores of the TbBpy *via* capillary force more easily, and reduced the Pd precursor with higher efficiency.⁴¹ Meanwhile, Pd particles can be restricted by the bipyridine functional group and would not overflow again, which may guarantee the uniform distribution of Pd. On the other hand, the reduction by H_2 has to be conducted under high temperature (120 °C), which can be inclined to result in aggregation of Pd particles.^{42,43}

Moreover, the effect of Pd loading on the catalytic activity of Pd@TbBpy was explored. As shown in Fig. 4a, when the loading of Pd nanoparticles increased from 0.25 wt% to 1 wt%, the conversion remained at 100%, while the C_2H_4 selectivity increased first and then decreased, which were 67.8%, 69.3%, and 75.1%, and 52.2%, respectively. This trend implies that a lower Pd loading may not guarantee enough active sites for the reactants, while an exorbitant Pd loading may lead to the agglomeration of Pd nanoparticles. The Pd particle size analysis (Fig. S7†) after the reaction also demonstrates that the Pd particle size is much larger than before the reaction. Therefore,



Fig. 3 (a) XPS total spectrum, (b) N 1s spectrum, and (c) Pd 3d spectrum of Pd@TbBpy; (d) Pd 3d XPS spectrum of Pd(II)@TbBpy.

based on the comparative results, a Pd content of 0.75 wt% was selected for subsequent experiments.

Besides, the effect of space velocity on the catalytic performance of Pd@TbBpy was also investigated (Fig. 4b). Over 120 °C, the C_2H_2 conversion remained at 100% at all selected space velocities, which were 40 000 h⁻¹, 50 000 h⁻¹, 60 000 h⁻¹, 70 000 h⁻¹, and 80 000 h⁻¹. As for the C_2H_4 selectivity, it was 73.4%, 78.5%, 81.02%, 88.2%, and 82.3%, respectively. The highest selectivity was obtained at velocity of 70 000 h⁻¹. Apparently, when the space velocity is too low, the products may not be blown away from the catalyst surface in time, resulting in an excessive hydrogenation or formation of other intermediates.⁴⁴ On the other hand, it would also be hard for the reactants to steadily adsorb on the active sites for reaction at higher space velocities. In addition, as shown in Fig. S8,† the C₂H₂ conversion of the Pd@TbBpy catalyst achieves complete conversion at 100 °C, which is about 40% higher than that of the Pd precursor. The C₂H₄ selectivity of Pd@TbBpy catalyst reached 88.2% at 120 °C and the efficiency of



Fig. 4 Catalytic activities of Pd@TbBpy (a) with different Pd contents and (b) at different space velocities.

C₂H₄ selectivity was twice that of Pd precursor at the same temperature. Therefore, the difference in reaction efficiency is very remarkable between Pd@TbBpy and Pd precursor. Based on the above results, the Pd loading of 0.75 wt% and the space velocity of 70 000 h^{-1} was selected as the optimal condition for following activity tests. At a low reaction temperature of around 120 °C, The catalytic performance of Pd@TbBpy is comparable to other highly selective Pd based catalysts supported on carbon materials listed in Table S4.† In fact, compared to other single atom Pd based catalysts, our developed catalyst has more advantages in experimental results and Pd distribution. By comparison, it was found that in this work, the Pd content we used was slightly higher than other catalytic groups, and the experimental parameters were not strict and harsh. In addition, it is important that the high ethylene selectivity is due to the layered stacking structure and the N-site anchoring of COFs, which enables the uniform distribution and stability of Pd nanoparticles.

The reactions involved in industrial production usually need to be carried out for a long time, which puts forward strict requirements on the life of the catalyst. Therefore, the longterm stability of the Pd@TbBpy was evaluated under the optimal reaction condition. As shown in Fig. 5, during the 1500 min of continuous reaction, the catalytic activity kept excellent with the C_2H_2 conversion being 100% and the C_2H_4 selectivity being ~90%. Compared with other reported Pd-based catalysts,⁴⁵⁻⁴⁸ Pd@TbBpy shows more remarkable long-term stability. However, a gradual formation of green oil, which was resulted from the deposit of by-products during the continuous hydrogenation, has been observed. It might has exerted some negative effects on the process activity as well as the catalyst stability.^{44,49}

3.3 Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) analysis

To further analyze the reaction mechanism, in this section, the dynamic adsorption–desorption processes of C_2H_2 and C_2H_2/H_2 over Pd@TbBpy were investigated by *in situ* FT-IR spectra. Notably, before the reactions, Pd@TbBpy has been reduced by



Fig. 5 Life test of Pd@TbBpy for the hydrogenation of C_2H_2 (temperature = 120 °C; space velocity = 70 000 h⁻¹; time \ge 1500 min; 20% C_2H_2/N_2 , 20% H_2/N_2 and He).

the mix of H₂ (20%)/N₂ (80%) at 120 °C for 60 min, and then purged with pure He to remove all adsorbates on the surface. Firstly, the adsorption behaviors of C2H2 onto Pd@TbBpy were analyzed under 80 °C, 100 °C, 120 °C, 150 °C, and 200 °C. As can be seen in Fig. 6a, the intensity of peaks didn't vary much as the temperature changing, demonstrating that the saturation adsorption of C₂H₂ could be obtained at low temperature. Specifically, the characteristic peaks of C_2H_2 at 3308–3264 cm⁻¹, which are attributed to the \equiv CH bending mode ($R(\delta J = +1)$, $P(\delta J = +1)$) (= -1)), and the central Q branch peaks ($\delta J = 0$) of \equiv CH $(\sim 730 \text{ cm}^{-1})$, as well as the IR-active \equiv CH bending vibrations $(717 \text{ and } 756 \text{ cm}^{-1})$ can be clearly observed, indicating a typical π -bond adsorption.³³ The C₂H₂ hydrogenation over zeolitesupported bimetallic catalyst has also been reported in Huang's work, in which the π -bond adsorption was confirmed as the main adsorption mode, and it was the dominant factor affecting the higher hydrogenation activity.⁵⁰

Besides, the C₂H₂ adsorption over Pd@TbBpy under 120 °C for 100 min was also continuously monitored by FT-IR spectroscopy (Fig. S9a[†]). It's clear that the characteristic peaks of C₂H₂ at 3308-3264 cm⁻¹ remained almost same during the whole period, which indicated that the C₂H₂ adsorption reached saturation at the very beginning and the adsorbed C2H2 has stayed at stable form without transforming into oligomeric state. In addition, the products transformation process of C₂H₂ hydrogenation over Pd@TbBpy was also investigated under 80 ° C, 100 °C, 120 °C, 150 °C, and 200 °C, as shown in Fig. 6b. During the adsorption-desorption processes, the peaks of C_2H_2 gradually diminished as the temperature increased, while some characteristic peaks at 3100-2750 cm⁻¹ and 1700-800 cm⁻¹, which can be attributed to the vibration of related functional groups of C₂H₄ and saturated alkanes, gradually showed up. The detailed dynamic adsorption process of feed gas has been evaluated by the real-time in situ DRIFTS spectra, in which, different from FT-IR spectra, the structural changes of functional groups during the reaction can be recorded, as shown in Fig. 6c and d.

The absorption peaks from 3123 cm^{-1} to 3012 cm^{-1} can be assigned to the stretching vibration of the C-H bond on the olefinic double bond (C=C-H). Thereinto, the characteristic peak at 3076 cm⁻¹ was assigned to the stretching vibration of the terminal double bond hydrogen ()C=CH₂). Besides, the peaks at 1670 cm⁻¹, 1616 cm⁻¹ and 1582 cm⁻¹ were corresponded to the stretching vibration of alkene C=C. Meanwhile, the peaks around 986 cm^{-1} , 965–949 cm^{-1} , and 908 cm^{-1} were attributed to the torsional vibration of alkene CH₂, the out-ofplane bending vibrations of trans-configuration CH (R1-HC= $CH-R_2$), and the out-of-plane rocking vibrations of alkene CH_2 in alkyl alkenes, respectively. These results confirmed the alkene production during the C₂H₂ hydrogenation over Pd@TbBpy. However, the typical peaks belonging to alkane can also been observed, including the asymmetric stretching vibration of $-CH_3$ (2972 cm⁻¹) and $-CH_2$ (2943 cm⁻¹), and the symmetric stretching vibrations of $-CH_3$ (2905 cm⁻¹) and $-CH_2$ (2852 cm^{-1}) . It suggested that the saturated alkanes were also generated inevitably during the hydrogenation process.⁵⁰

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Fig. 6 (a) In situ DRIFTS spectra of C_2H_2 adsorption on Pd@TbBpy. (b) In situ DRIFTS spectra of C_2H_2 hydrogenation on Pd@TbBpy. The real-time in situ DRIFTS spectra (c) 3500–2500 cm⁻¹ and (d) 2000–650 cm⁻¹ of C_2H_2 hydrogenation on Pd@TbBpy (temperature = 120 °C; space velocity = 70 000 h⁻¹; 20% $C_2H_2/80\%$ N₂, 20% $H_2/80\%$ N₂).

Finally, after 100 min of reaction, the desorption experiment was carried out with pure He purging over Pd@TbBpy surface. As shown in Fig. S9c,† the characteristic peaks of C_2H_2 (3308– 3264 cm⁻¹, 756–717 cm⁻¹) disappear very quickly, while the peaks belonging to the products (C_2H_4 and alkanes) still existed, which indicated that C_2H_2 was more easily desorbed from the Pd@TbBpy surface than the products. However, the presence of C_2H_4 may lead to the further hydrogenation, resulting in the formation of carbon layers depositing on the catalyst surface.⁵¹ The similar result has also been reported in Zhang's work,⁵² in which, a carbonaceous layer was observed on the surface of Pd/ Al_2O_3 catalyst after the hydrogenation of C_2H_2 .

3.4 DFT analysis

To further elucidate the role of TbBpy in the reaction and the interaction between Pd and TbBpy, DFT calculation was further conducted. As can be seen in Fig. 7, after the loading of Pd onto



Fig. 7 (a) Adsorption energy diagrams of Pd and Pd@TbBpy in C_2H_2 and H_2 absorption process. Charge transfer of the adsorbed (b) C_2H_2 and (c) H_2 around the Pd and Pd@TbBpy.

(00)

TbBpy, the adsorption energies (E_{ads}) of C_2H_2 and H_2 increased from -2.12 eV and -0.73 eV to -2.92 eV and -0.88 eV, respectively. The higher E_{ads} indicated that the adsorption of C₂H₂ and H₂ onto Pd were both easier with the presence of TbBpy, which is in accordance with the results of DRIFTS. In addition, there are two peaks displayed in H₂-TPD (Fig. S10[†]) at 135 °C and 482 °C, which are attributed to the desorption of chemically adsorbed hydrogen on the Pd phase and the desorption of overflow hydrogen, respectively.53,54 Therefore, it can be seen that H₂ can be well adsorbed on the Pd@TbBpy when the temperature is below 135 °C, which provides strong evidence for obtaining good reaction activity. Clearly, the interaction of Pd and TbBpy benefited the adsorption of reactants, which is the very first step to initiate the whole reaction. Besides, the charge transfer value of C varied from -0.11e to -0.15e from the unsupported to the supported Pd, indicating the electron-grabbing power of C in C₂H₂ was enhanced with the presence of TbBpy. Meanwhile, the charge transfer value of H changed from 0.027e to 0.41e, implying that the electron donating ability of H₂ was also greatly improved. It indicated that the electron transfer process between the reactants was obviously enhanced with the presence of TbBpy. Above results show that the interaction between Pd and COFs material helped to increase the catalytic activity from the atomic level.

4. Conclusions

In this study, a novel 2D covalent organic framework (TbBpy) was synthesized via the Schiff base reaction of 1,3,5-triformylbenzene and 2,2'-bipyridyl-5,5'-diamine, accompany with the formation of C=N bond. Pd nanoparticles were loaded onto the surface and into the pores of TbBpy with uniform particle sizes (*i.e.* \sim 2.4 nm), which coordinated with N atoms in the bipyridine functional group of TbBpy. The 100% of C2H2 conversion and 88.2% of C2H4 selectivity can be obtained at the temperature of 120 °C and the space velocity of 70 000 h⁻¹. Under the this optimal condition, Pd@TbBpy showed remarkable stability as well as life (>1500 min) in the durability experiments. Over 150 °C, the over-hydrogenation of intermediate products was more inclined to happen with the production of green oil, which was composed of hydrocarbon derivatives and might exerted negative effects on catalyst activity. In addition, based on the results of DRIFTS, it was found that C₂H₂ can quickly achieve equilibrium of adsorption on the surface of Pd@TbBpy without transformation into oligomer. And a typical form of π -bond adsorption has been observed, which can be beneficial for the catalytic reactivity. However, the intermediate products were found out to be more difficult to desorb from the Pd@TbBpy surface than C₂H₂, thus leading to the generation of saturated alkanes inevitably, which further explain the production of green oil. The DFT analysis revealed that, after the loading of Pd onto TbBpy, the adsorption of C₂H₂ and H₂ onto the catalyst were both improved, and the electron transfer between C and Pd was also enhanced, which might further demonstrate the promotion effects of TbBpy as support toward the catalytic activity of Pd. The above results demonstrate and elucidate the high activity and selectivity of Pd

catalyst supported on TbBpy in the C_2H_2 hydrogenation, promoting the application of COFs materials in this area.

Conflicts of interest

The authors declare no competing financial interest.

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