



Cite as
Nano-Micro Lett.
(2023) 15:149

Received: 2 February 2023
Accepted: 20 April 2023
Published online: 7 June 2023
© The Author(s) 2023

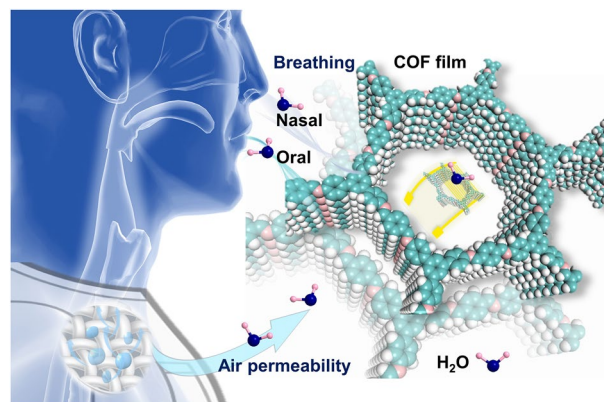
Outstanding Humidity Chemiresistors Based on Imine-Linked Covalent Organic Framework Films for Human Respiration Monitoring

Xiyu Chen¹, Lingwei Kong¹, Jaafar Abdul-Aziz Mehrez¹, Chao Fan¹, Wenjing Quan¹, Yongwei Zhang¹, Min Zeng¹ ✉, Jianhua Yang¹, Nantao Hu¹, Yanjie Su¹, Hao Wei¹, Zhi Yang¹ ✉

HIGHLIGHTS

- Imine groups in covalent organic framework (COF) films act as dual-active sites for humidity sensing, inducing an intrinsic enhanced mechanism of reversible protonated tautomerism via water molecule-induced hydrogen bonding.
- The *cis*-ketoimine reciprocal isomerization induces a stretching vibration effect for the ordered conjugated conductive frame of COF films, realizing fast response, wide range, and high sensitivity characteristics for humidity detection.
- Resistance changes of COF film-based sensors keep a strong linear relationship with low-range relative humidity, reflecting the quantitative sensing mechanism at the molecular level.

ABSTRACT Human metabolite moisture detection is important in health monitoring and non-invasive diagnosis. However, ultra-sensitive quantitative extraction of respiration information in real-time remains a great challenge. Herein, chemiresistors based on imine-linked covalent organic framework (COF) films with dual-active sites are fabricated to address this issue, which demonstrates an amplified humidity-sensing signal performance. By regulation of monomers and functional groups, these COF films can be pre-engineered to achieve high response, wide detection range, fast response, and recovery time. Under the condition of relative humidity ranging from 13 to 98%, the COF_{TAPB-DHTA} film-based humidity sensor exhibits outstanding humidity sensing performance with an expanded response value of 390 times. Furthermore, the response values of the COF film-based sensor are highly linear to the relative humidity in the range below 60%, reflecting a quantitative sensing mechanism at the molecular level. Based on the dual-site adsorption of the (–C=N–) and (C–N) stretching vibrations, the reversible tautomerism induced by hydrogen bonding with water molecules is demonstrated to be the main intrinsic mechanism for this effective humidity detection. In addition, the synthesized COF films can be further exploited to effectively detect human nasal and oral breathing as well as fabric permeability, which will inspire novel designs for effective humidity-detection devices.



KEYWORDS Covalent organic frameworks; Humidity sensors; Reversible tautomerism; Non-invasive diagnosis; Health monitoring

✉ Min Zeng, minzeng@sjtu.edu.cn; Zhi Yang, zhiyang@sjtu.edu.cn

¹ Key Laboratory of Thin Film and Microfabrication (Ministry of Education), Department of Micro/Nano Electronics, School of Electronic Information and Electrical Engineering, Shanghai Jiao Tong University, Shanghai 200240, People's Republic of China



1 Introduction

Human metabolism is rich in information and involves water molecules, ranging from skin moisturization to more complex internal respiration that can shed light on the internal health of the human body. Particularly, data from the respiratory tract can represent the physiological and psychological state of the individual, and even the early diagnosis of some diseases (such as lung cancer, epidemic diseases, and apnea syndrome) [1, 2]. Recent studies have demonstrated that respiratory viruses may enhance aerosol transmission risks and are susceptible to large outbreaks, even in lower-humidity environments [3]. Additionally, doctors can use perspiration to diagnose some diseases, uncover drugs, and analyze the performance of athletes [4, 5]. Eventually, to satisfy the quest for practical application, developing a humidity sensor with ultra-sensitivity, real-time quantitative analysis, and a wide detection range is necessary.

Two-dimensional (2D) materials with distinctive microstructures and unique electrical characteristics have attracted great interest in the field of humidity sensing. For example, nanocarbon materials [6, 7], conducting polymers [8], metal–organic frameworks [9], transition metal dichalcogenides [10], and transition metal carbides/nitrides/carbonylides [11, 12] were investigated as sensing materials. In general, the microstructure of these humidity-sensitive materials can not be precisely designed at the molecular level to obtain optimized sensitivity. To address this issue, a class of porous covalent organic frameworks (COFs) is explored with distinct advantages in the sensing field [13], profiting from the structural stability in complex environments and abundant active sites [14]. The pre-designed COFs can form single or hetero-porous frameworks by selecting monomer linkers and functional groups according to sensing requirements [15].

The chemical affinity between COFs and target analyte is generated by electron-rich/deficient building units to a periodic lattice of donor–acceptor pairs that lead to the stimulated response. Replicating multiple identical binding sites through the whole extended topology shows that the signal can be effectively transduced and amplified through the COFs to achieve a high sensitivity [16]. The benefits of COFs for sensing applications include not only molecule identification but also charge transfer rate acceleration. Furthermore, COFs manufactured into films with continuous pores can expose the greater interior surface area to the

analytes, resulting in enhanced sensing performance [17]. It is noteworthy that the strong non-covalent interactions between COF materials and target molecules make them highly selective [18], allowing for reversible colorimetric sensing *via* the inherent linking chemistries as the sensing site, employing complex electron transfer and reversible protonation of the linkage groups [19]. However, it is critical to develop COF-based humidity chemiresistors, especially COF films with (1) higher sensitivity and easier quantification ability; (2) better conductivity, and (3) electronic equipment compatibility for the increasing needs of wearable electronics and health monitoring [20]. For example, the BTA-TAPT COF film-based interdigitated electrodes (IDEs) capacitive sensor exhibits high sensitivity toward benzene vapor [21]. Similarly, since water and benzene molecules can have host–guest interactions with COFs, the researchers developed COF-based electrical humidity sensors. The response and recovery times of the COF-TXDBA-based %RH sensor were 37 and 42 s, respectively [22]. Nevertheless, the response/recovery time of dozens of seconds reported in previous studies remains insufficient for overcoming the limitation of the COFs-based sensor response to human metabolism-related humidity detection.

Herein, we present imine-linked COF films for ultra-sensitive chemiresistive humidity sensors with a wide detection range. We choose the 1,3,5-tris(4-aminophenyl) benzene (TAPB) as the electron-rich structure in order to induce charge exchange between water molecules. On the other hand, the existence of 2,5-dihydroxyterephthaldehyde (DHTA) compared to other monomers such as 4,4'-biphenyldicarboxaldehyde (BPDA) and 1,4-phthalaldehyde (PDA), with strong intramolecular O–H...N=C hydrogen bonding can protect the imine bond from any nucleophilic attacks or the construction of an intact crystalline film. The pre-designed COF films can powerfully facilitate the diffusion of water molecules into and out of the porous framework, thus increasing the response to changes in human breath [23, 24]. Besides, the flexible and free-standing COF films can be easily integrated into humidity detection devices with higher stability. The results indicate that the current amplitude of the humidity sensor based on COF film has a 390-fold increase in a humidity range of 13–98% and a fast response/recovery time of 0.4/1.0 s. Notably, the imine groups of COF films acted as dual-site, and the intrinsic sensing mechanism of hydrogen bonding caused reversible protonated tautomerism for this excellent humidity sensing

performance. The humidity sensing mechanism is confirmed by in situ Raman spectroscopy and density functional theory (DFT) calculations. Furthermore, we apply COF film-based sensors to monitor human breathing and the breathability of certain fabrics. The precise molecular structure regulation and continuous porous properties of COF films can accurately capture moisture perturbations. Therefore, the COF films developed in this work hold great promise for ultra-sensitive human-related humidity detection.

2 Experimental and Calculation

2.1 Materials and Apparatus

Materials, COF synthesis route and apparatus involved in this work are all represented in the Supporting Information.

2.2 COF Films Design

Electron-rich structures, such as TAPB, were selected as the basis for humidity-induced electrical signals via electron exchange with water molecules. Designed active sensing sites are represented by the N atoms of DHTA which is responsible for the inducement of tautomerism [25]. Water molecules combined with electron-withdrawing groups are supposed to produce group dissociation leading to the formation of a charge transfer complex across the conjugated imine bonds. Moreover, the existence of DHTA with strong intramolecular O–H···N=C hydrogen bonding can protect the imine bond from any nucleophilic attacks or the construction of an intact crystalline film [26]. Given the above considerations, we chose three increasingly electron-rich TAPB, namely 1,3,5-tris-(4-aminophenyl)triazine (TAPT) and 1,3,6,8-tetrakis(4-aminophenyl)pyrene (TAPPy) to provide the polarized electron-accepting imine-linker blocks. These three films referred to as COF_{X-DHTA} are shown in Fig. S1, with their corresponding typologies and synthesis methods.

2.3 Growth Process of COF Films

The growth of COF into continuous porous films allows large-scale preparation, and the films are more convenient

for processing electronic devices. Inspired by the COFs separation membrane preparation [27], the connecting monomers undergo an imine condensation process at the two-phase interface, resulting in the self-assembly of functional COF films at room temperature (RT). Figure 1 depicts the COF powder- and film-based sensor methods to illustrate the characteristics of the two types of COF sensors. Following a washing procedure, the COF films were directly transferred to the hydrophilic-treated IDEs and assembled into a chemiresistive sensor for subsequent sensing applications. It is critical to ensure that oxygen is isolated throughout the whole process to obtain highly crystalline COF films. The similar-structured monomers TAPT or TAPPy were also successfully synthesized into COF films, demonstrating the generality of the large-scale preparation method. Figure S2 shows the optical photos of COF films prepared by interfacial synthesis and self-assembly. Due to the disorder-to-order transition of the grain boundary, the as-prepared COF_{X-DHTA} films have significantly higher carrier transport and mobilities than powders, resulting in a remarkable humidity-sensing ability. Simultaneously, to speculate the hypothesis that DHTA is the active functional group in humidity sensing, comparative COF films containing BPDA and PDA were prepared (Figs. S3 and S4). Moreover, to determine whether the thickness of the COF film affects sensing capability, a series of COF_{TAPB-DHTA} films of varying thicknesses were prepared by regulating the condensation reaction of a constant amount of TAPB with varied masses of DHTA monomer.

2.4 Humidity-Sensing Measurements

The advantages of COF films will be demonstrated in the following humidity performance tests. The preparation of the IDEs-COF powder- and film-based sensors and the humidity testing are elaborated in Figs. S5–S8. However, the COF_{X-DHTA} powder-based humidity sensors have a low signal-to-noise ratio and a small signal amplitude over a wide RH range (0–74.9% RH). As a result, COF powders are incompatible with applications requiring ultra-sensitive humidity detection. The performance comparison between COF_{X-DHTA} powder- and film-based sensors can further elaborate the advantages of COF_{X-DHTA} films in sensing. Furthermore, we investigate the relationship between COF film

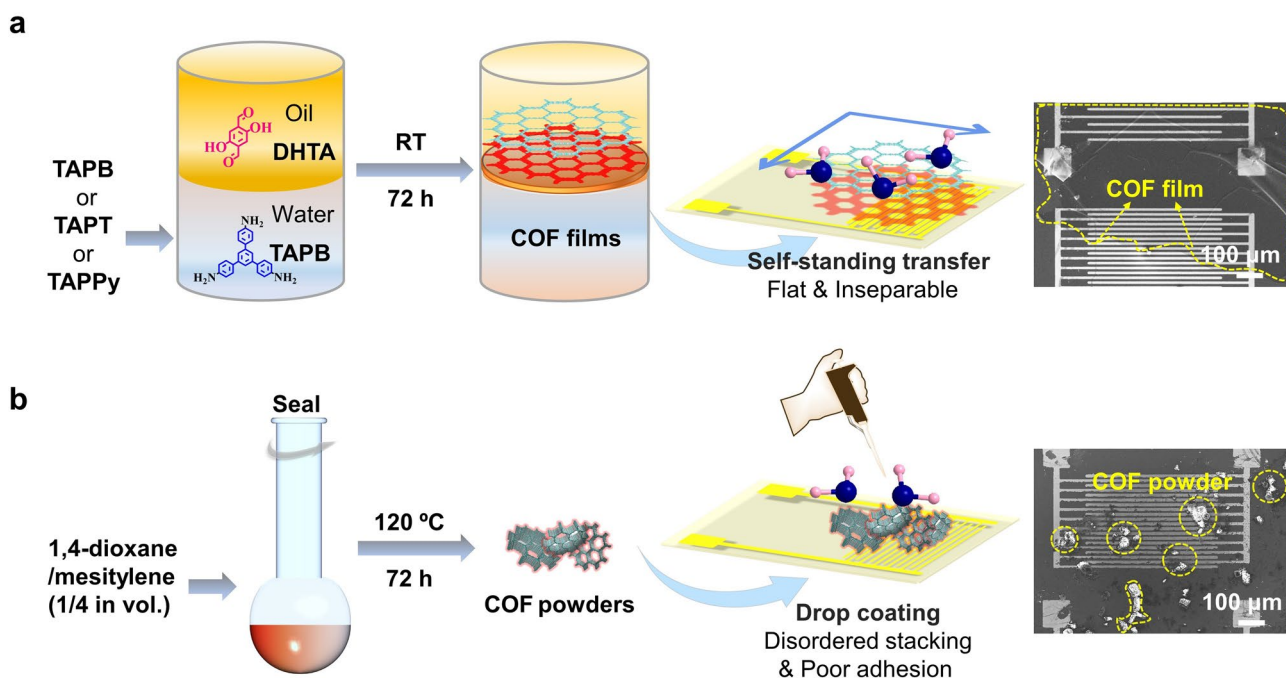


Fig. 1 Schematic illustration of the preparation process of the humidity sensor based on the COF **a** films (tightly adhered) and **b** powders (randomly dispersed)

thickness and sensing performance. The 1.5-COF_{TAPB-DHTA} film (108.9 μm) from the X-COF_{TAPB-DHTA} series with the μm-level exhibited the highest response values (Figs. S9 and S10). To exclude the interference of the complex ambient atmosphere, selectivity tests for different vapors (including water) were also performed. The COF_{X-DHTA} films have an intense response to water vapor, and the most outstanding selectivity is found in the case of the COF_{TAPT-DHTA} film with a strongly polar triazine structure (Fig. S11) [28]. On the other hand, the COF_{TAPB-BPDA} and COF_{TAPB-PDA} film displayed a negligible humidity response and a low signal-to-noise ratio. Therefore, it is confirmed that the more electron-rich structural functional monomer of the COF films generated a higher humidity sensing response value. The humidity response is defined here using Eq. (1):

$$\text{Response \%} = \left(\frac{I_{\text{humidity}} - I_{\text{dry}}}{I_{\text{dry}}} \right) \times 100\% \quad (1)$$

where I_{humidity} and I_{dry} represent the current of the sensor under analyte vapors and dry compressed air, respectively.

3 Results and Discussion

3.1 Characterizations of COF Powders and Films

The powder X-ray diffraction (PXRD) patterns of COF films depicted in Fig. 2a–c contain some distinct reflections, indicating the acquisition of the predicted structure of COF films with high crystallinity. The features at 2.75°, 4.93°, 5.62°, 7.59°, and 9.90° are denoted as (100), (110), (200), (210), and (220) facets, respectively. Notably, there is a weak peak at the angle of 26.02°, corresponding to the (001) reflection of COF_{TAPB-DHTA} [17]. Comparing the experimental and simulated results of PXRD, we can determine that COF_{TAPB-DHTA} is in the AA stacking mode (Fig. S12). The COF_{TAPT-DHTA} and COF_{TAPPy-DHTA} also exhibit AA stacking similar to previously reported COF structures [26, 29]. Excluding the amorphous state in the contrast films that affect the sensing results, the sharp reflections of PXRD patterns were also obtained (Fig. S13). Furthermore, the stretching characteristic peak of C=N produced by imine condensation confirms the existing topology of COF films, which is about 1620 cm⁻¹ in the fourier transform infrared

spectroscopy (FT-IR) spectra. A new characteristic peak that appears at $3300\text{--}3500\text{ cm}^{-1}$ can be ascribed to the N–H stretching vibration of amine groups and the hydroxyl on the DHTA monomer (Figs. 2d–f and S14) [30]. To evaluate the chemical components of the prepared $\text{COF}_{\text{X-DHTA}}$ films, XPS analysis was performed. The full XPS spectra of the $\text{COF}_{\text{X-DHTA}}$ films indicate that only C, N, and O elements exist, and the peak positions of each element are very similar (Fig. S15). The prepared ultrathin COF films with nm-level thickness can facilitate clearer observation of the microscopic morphology of COF films. According to the atomic force microscopy (AFM) images, the thickness of

the prepared ultrathin $\text{COF}_{\text{TAPB-DHTA}}$ film with layer-by-layer self-assembly stacking at the interface is 143.8 nm (Fig. 2g) [31]. The stabilized nm-level $\text{COF}_{\text{TAPB-DHTA}}$ film can also be delaminated to homogenous thinner sheets. The average thickness is approximately 3.56 nm by ultrasonication for 3 h in alcohol. Comparing the 3D topography and AFM height of the nm-level $\text{COF}_{\text{X-DHTA}}$ films, it was found that the degree of uniform dispersion of $\text{COF}_{\text{TAPB-DHTA}}$, $\text{COF}_{\text{TAPT-DHTA}}$ and $\text{COF}_{\text{TAPPy-DHTA}}$ films gradually decreases. As a result, there are variations in the stacking microstructure of the three samples, which might affect their capacity to sensors detect humidity (Figs. 2h and S16) [32]. The average thickness is

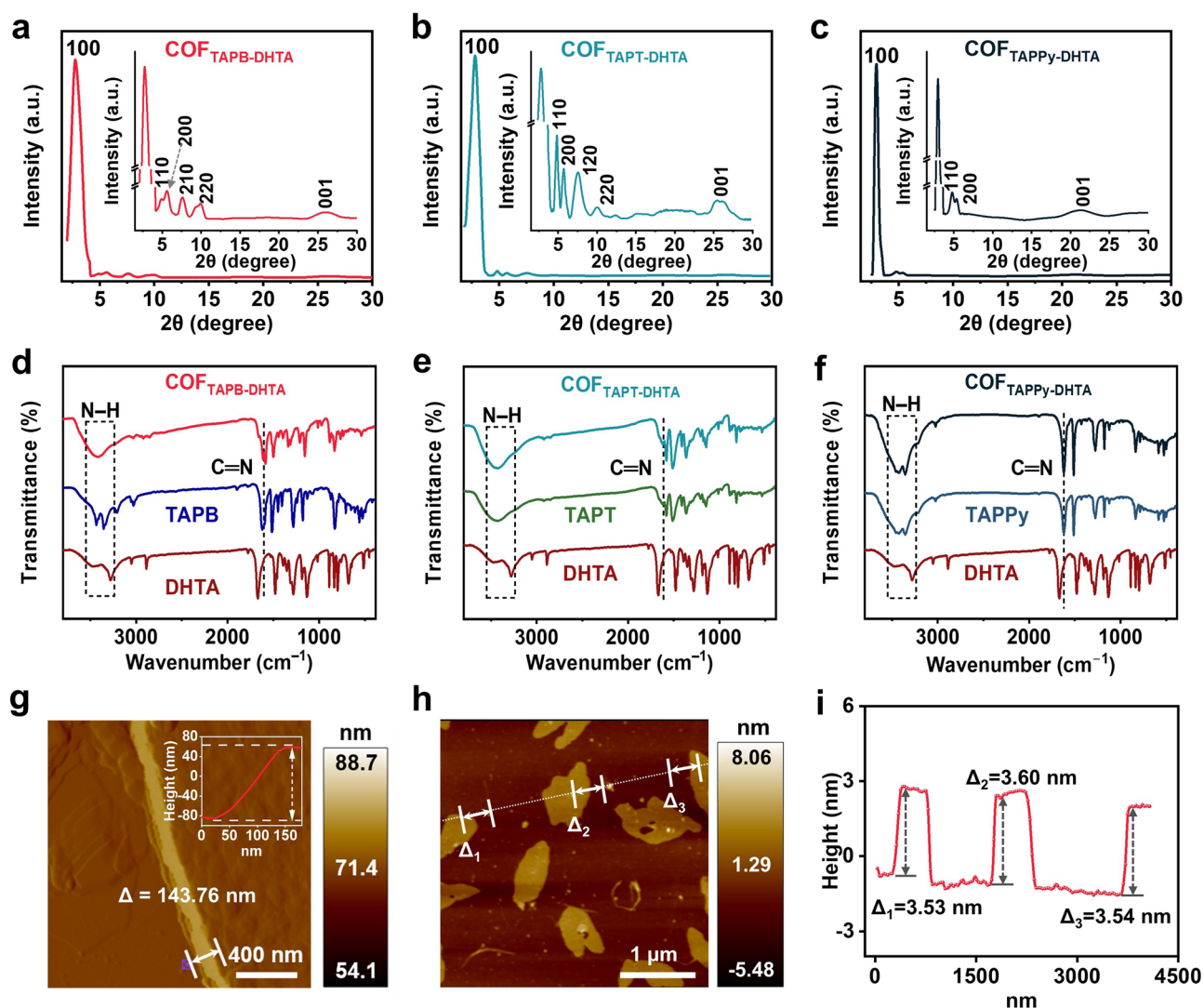


Fig. 2 PXRD patterns of **a** $\text{COF}_{\text{TAPB-DHTA}}$, **b** $\text{COF}_{\text{TAPT-DHTA}}$, and **c** $\text{COF}_{\text{TAPPy-DHTA}}$ film. FT-IR spectra of **d** $\text{COF}_{\text{TAPB-DHTA}}$, **e** $\text{COF}_{\text{TAPT-DHTA}}$, and **f** $\text{COF}_{\text{TAPPy-DHTA}}$ film. **g** AFM height image of $\text{COF}_{\text{TAPB-DHTA}}$ film. **h** AFM height image of $\text{COF}_{\text{TAPB-DHTA}}$ film after ultrasound for 3 h. **i** Height profile along the white line in **(h)**

approximately 3.56 nm by ultrasonication for 3 h in alcohol. Comparing the 3D topography and AFM height of the nm-level $\text{COF}_{\text{X-DHTA}}$ films, it was found that the degree of uniform dispersion of $\text{COF}_{\text{TAPB-DHTA}}$, $\text{COF}_{\text{TAPT-DHTA}}$, and $\text{COF}_{\text{TAPPY-DHTA}}$ films gradually decreases. As a result, there are variations in the stacking microstructure of the three samples, which might affect their capacity to sensors detect humidity (Figs. 2h and S16) [32].

Scanning electron microscopy (SEM) images of COF films demonstrate the continuous plane accompanied by porous morphology on the surface (Figs. 3a–c and S17–S18). In contrast, the COF powders prepared by the solvothermal method possess a uniform spherical shape with disorderly stacked blocks (Fig. 3d–f). The clear lattice fringes can be observed in high-resolution transmission electron microscopy (HR-TEM) images of $\text{COF}_{\text{TAPB-DHTA}}$ film, where the spacing is about 2.98 Å, corresponding to (100) facets (Fig. 3h) [33]. HR-TEM images and selected area electron diffraction (SAED) patterns of the $\text{COF}_{\text{TAPB-DHTA}}$ film are consistent with the crystal structure provided by PXRD results (Fig. S19) [34]. The stacking, crystallinity and grain size of COFs will seriously interfere with the transition energies required for electron transfer during the sensing process [35]. Therefore, the long-range continuous porous COF films will accelerate charge transfer and provide a solid basis for favorable sensing performance [36].

The Brunauer–Emmett–Teller (BET) measurement result of μm -level COF films is shown in Fig. S20. The nitrogen sorption isotherm exhibits a type IVb isotherm with a hysteresis loop, confirming the mesoporous character of the $\text{COF}_{\text{TAPB-DHTA}}$ and $\text{COF}_{\text{TAPT-DHTA}}$ films (Fig. S20a, b). The equilibrium model of quenched solid density functional theory (QSDFT) analyzes $\text{COF}_{\text{TAPB-DHTA}}$ film and yields a very narrow pore size distribution with a maximum of 2.28 nm (Fig. S20a) which agrees with the structural model in SEM and TEM images. $\text{COF}_{\text{TAPB-DHTA}}$ film has the largest BET surface of $1143.18 \pm 38.66 \text{ m}^2 \text{ g}^{-1}$, with a cumulative pore volume of $0.63 \pm 0.05 \text{ cm}^3 \text{ g}^{-1}$. There are two peaks in the cumulative pore size volume of $\text{COF}_{\text{TAPPY-DHTA}}$ film, which play an adsorption function at 2.34 and 32.23 nm, indicating the existence of a hierarchically porous structure (Fig. S20f). The alternating stacking of $\text{COF}_{\text{TAPPY-DHTA}}$ film layers containing heteroporous frameworks may cause an even more pore volume reduction. DHTA hydrogen bonding can prevent a nucleophilic attack during the condensation reaction, resulting in $\text{COF}_{\text{X-TAPB}}$ films with a controllable pore

size (Figs. S20d, e and S21). In contrast, the pore size distribution of $\text{COF}_{\text{TAPB-BPDA}}$ and $\text{COF}_{\text{TAPB-PDA}}$ films without DHTA is more dispersed. $\text{COF}_{\text{TAPB-BPDA}}$ and $\text{COF}_{\text{TAPB-PDA}}$ films only have BET surface results of 666.26 ± 13.31 and $611.16 \pm 6.49 \text{ m}^2 \text{ g}^{-1}$, respectively (Fig. S21a, b).

3.2 Humidity Detection and Applications Based on the IDEs-COF Film Sensors

Within the $\text{COF}_{\text{X-DHTA}}$ films, the most sensitive film is $\text{COF}_{\text{TAPB-DHTA}}$ with the lowest detection limit when using the normalizing control of water vapor exposure time. The real-time current response curve can still be observed when the relative humidity is under 50%. Within a 3.2% RH change (from 13.1 to 16.3%, the low RH), the response value increased by as much as 108.7% (inset in Fig. 4a). The IDEs- $\text{COF}_{\text{TAPB-DHTA}}$ film-based sensor has a current amplitude enhancement of 390 times when the humidity ranges from 13.1 to 98.2% RH (Fig. 4a). Such an ultra-high sensitivity performance of the COF film-based sensor is critical for the perturbation of human-related humidity detection [37].

As contrasted with $\text{COF}_{\text{TAPB-DHTA}}$, $\text{COF}_{\text{TAPT-DHTA}}$ and $\text{COF}_{\text{TAPPY-DHTA}}$ films were less sensitive with higher detection limits (Fig. 4b, c). A logarithmic fit for the response and RH from the lowest detection capability of the $\text{COF}_{\text{X-DHTA}}$ film-based sensor to 60% RH (atmospheric environment in sunny weather) shows a linear relationship for all films (Fig. 4d–f insets). The fact that COF films can be designed at the molecular level, shows an excellent linear response at low humidity. As for $\text{COF}_{\text{TAPB-DHTA}}$ film, the theoretical detection limit can reach 7%, which is far better than the detection ability of traditional humidity sensors at RT (Fig. 4d inset) [38]. It is clarified that by fine-tuning the ligand monomers and functional groups of the COF topology, we can achieve the precise design of functional structural units adapted to multiple application scenarios with high sensitivity, wide range, fast response, and high linear relationship. To evaluate the humidity monitoring stability of the COF films, we performed the cycle stability test in the atmosphere of ambient humidity. Even after 12 cycles, there is a negligible change in the current output characteristic curve, suggesting the $\text{COF}_{\text{X-DHTA}}$ films are stable (Fig. 4g–i). When the IDEs-COF film-based sensor is exposed to a high-humidity atmosphere (76.2% RH) for 60 s, it recovers only in 3.4 s on

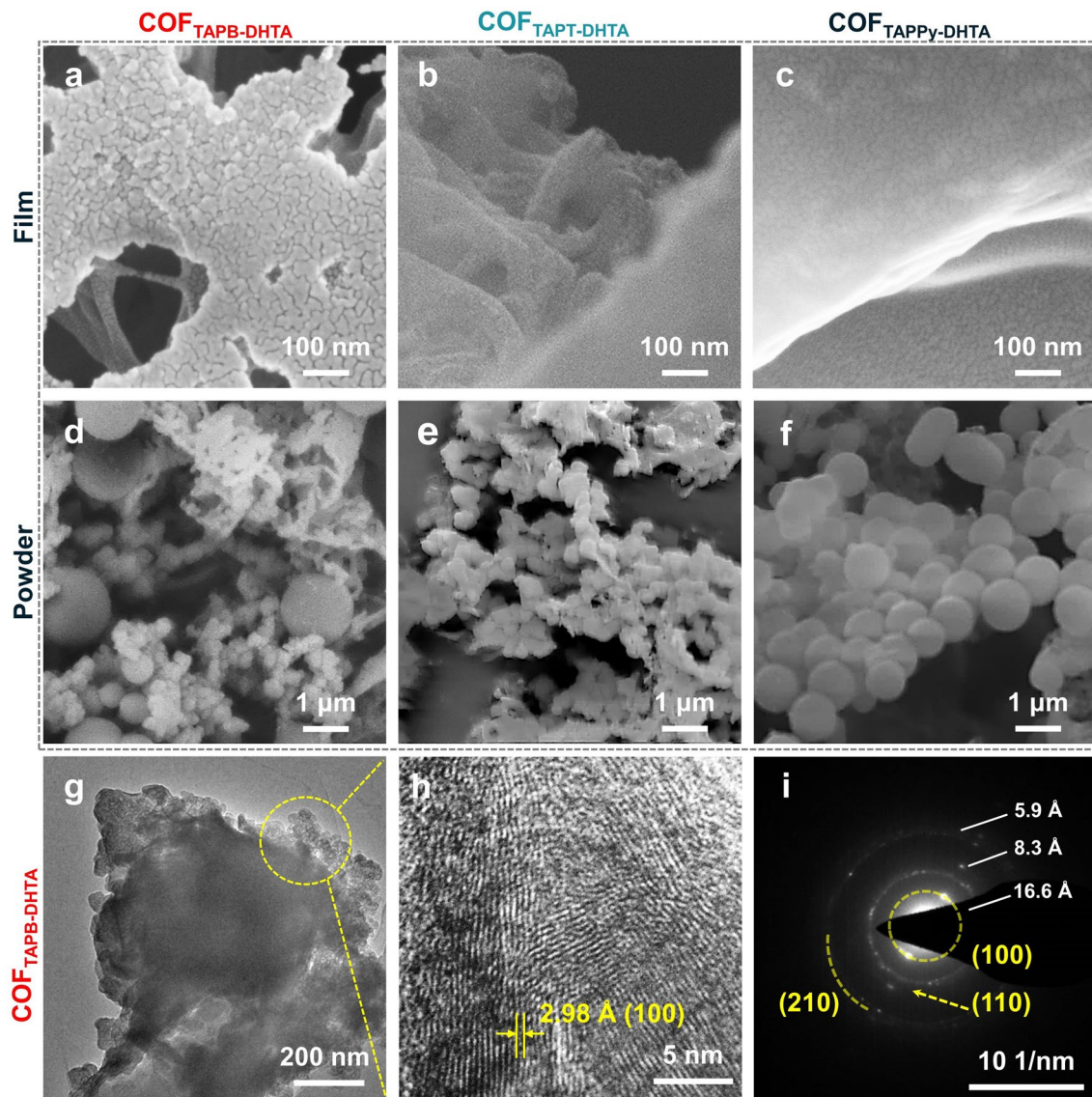


Fig. 3 SEM images of **a** $\text{COF}_{\text{TAPB-DHTA}}$, **b** $\text{COF}_{\text{TAPT-DHTA}}$, and **c** $\text{COF}_{\text{TAPPy-DHTA}}$ film. SEM images of **d** $\text{COF}_{\text{TAPB-DHTA}}$, **e** $\text{COF}_{\text{TAPT-DHTA}}$, and **f** $\text{COF}_{\text{TAPPy-DHTA}}$ powder. **g** TEM, **h** HR-TEM and **i** SAED patterns of $\text{COF}_{\text{TAPB-DHTA}}$

average (Fig. 4i). Long-term sensing response tests demonstrate the structural stability of the $\text{COF}_{\text{TAPB-DHTA}}$ film (Fig. S22). The humidity test for the same $\text{COF}_{\text{TAPB-DHTA}}$ film was repeated every two months under the same conditions. After ten months, the COF film-based humidity sensor retained 87.7% of its original response value.

Benefiting from the fast response and short interval periods of the COF film-based humidity sensor, we investigated its potential application for monitoring human breath rate. The IDEs- $\text{COF}_{\text{X-DHTA}}$ was used to detect two of these breathing patterns of volunteer-I, namely oral and nasal breathing.

Previous humidity monitoring reports indicate that the RH of oral breathing is higher than that of nasal due to the presence of saliva in the mouth [39]. As shown in Fig. 5a–f, our detection yields the same result, the oral breathing monitoring signal current value is greater than the nasal. Volunteer-I must rest for 1 min before each test and wear a mask with a breathing valve (Fig. S23). The IDEs- $\text{COF}_{\text{X-DHTA}}$ film-based humidity sensor is embedded in the mask to ensure that the data of each test is relatively objective and accurate. The nasal breathing monitored for 2 min shows that the response and recovery times of the IDEs- $\text{COF}_{\text{TAPB-DHTA}}$ film-based

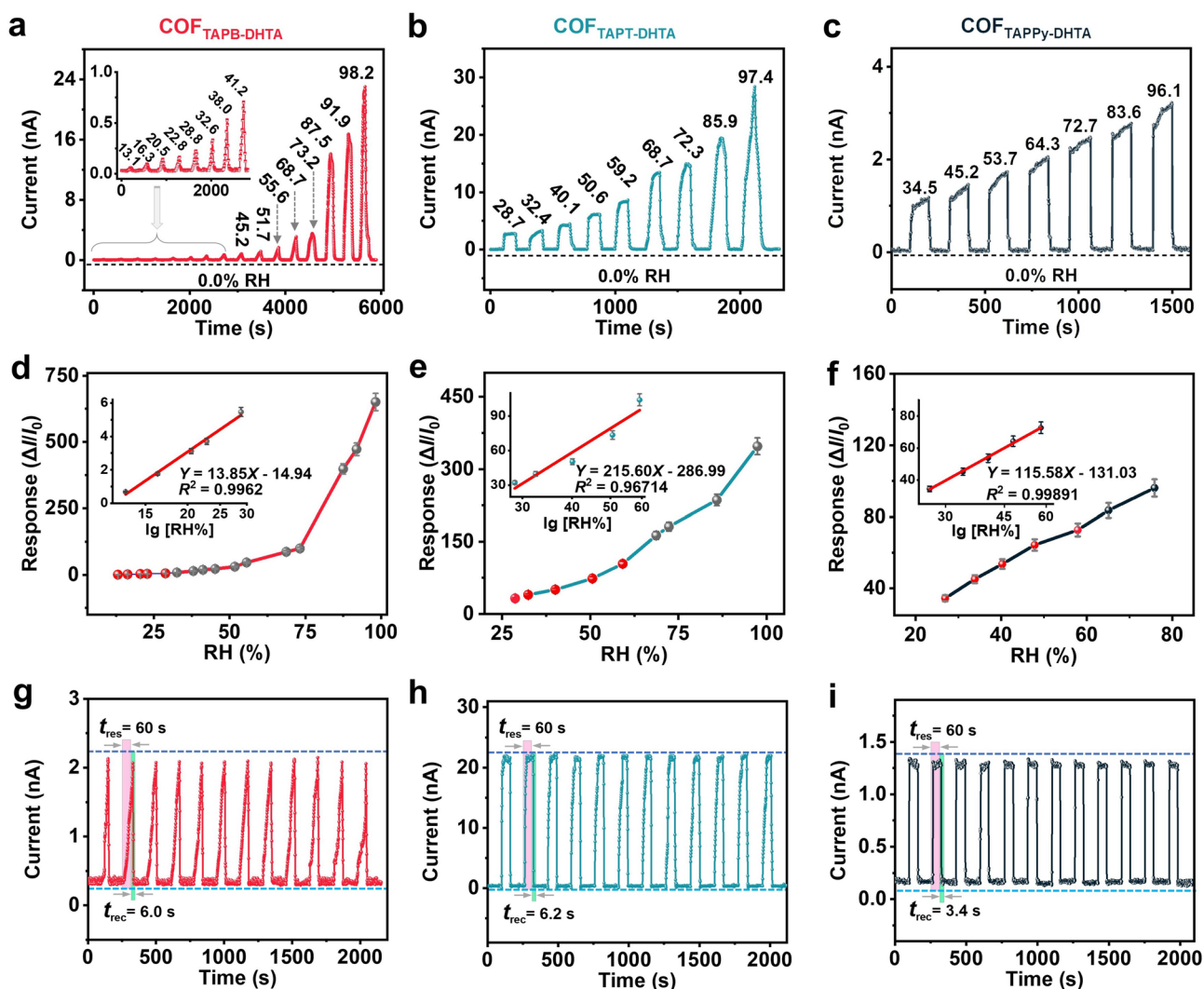


Fig. 4 Dynamic response characteristic curves of **a** IDES-COF_{TAPB-DHTA}, **b** COF_{TAPT-DHTA}, and **c** COF_{TAPPY-DHTA} film-based humidity sensors at different RH (Detection range: from dry air 0.0% to different RH). Response versus RH curve of **d** COF_{TAPB-DHTA}, **e** COF_{TAPT-DHTA}, and **f** COF_{TAPPY-DHTA} film (the inset shows the linear fit curve of response versus the RH logarithm). Response and recovery curves for 12 cycles of **g** COF_{TAPB-DHTA}, **h** COF_{TAPT-DHTA}, and **i** COF_{TAPPY-DHTA} film (Detection range: 41.7–76.2% RH)

sensor are 0.4 and 1 s, respectively (Fig. 5a). Likewise, the real-time current signal curve increased significantly in the oral breathing across 25 cycles performed by volunteer-I with a stable amplitude as shown in Fig. 5d–f. As we know, the average humidity of the transient airflow carried by human breath is generally over 90% RH [39]. The sensing results from Fig. 5a–f indicated that the COF_{X-DHTA} film-based sensor maintains good sensing performance even at a high RH of over 90%, highlighting its potential for human respiration monitoring with good stability.

Remarkably, it is meaningful to evaluate the breathability of clothing in direct contact with the skin, because the skin

plays a key role in human life activities, such as the detection of cotton and polyester fabrics' air permeability [41]. Evaporating normal saline at 36 ± 0.5 °C is used as a simulation of sweat on the surface of human skin, and the distance between the container neck and the IDES-COF_{TAPB-DHTA} film-based sensor is set at 20 mm (Fig. S24). After stabilizing for 400 s, the conical flask was completely covered with fabrics, and real-time dynamic current curves were observed (Fig. 5g). The ventilation effect of pure cotton fiber is found to be significantly better than that of polyester and tends to stabilize after about 350 s (red curve in Fig. 5g). As shown in the chemical structural formula of cellulose with abundant

hydroxyl groups that can form hydrogen bonds and assist in transporting water molecules. Interestingly, polyester fibers with a highly crystalline molecular chain structure quickly hindered the evaporation of “sweat”. Due to the accumulation of water molecules on the surface (after about 200 s), the capillary core transfer occurs on the polyester fiber, allowing a small number of water molecules to overflow the surface. Hence, the detection current increases slightly (blue curve in Fig. 5g) before stabilizing. In addition, a series of interference tests on the IDE-COF-based sensor, the conditions

of the experiments including a mixed gas atmosphere (Fig. S25), light illumination under different wavelengths (Fig. S26), variable RT (Fig. S27a–c), and the hysteresis response (Figs. S28 and S29) were conducted. And all testing results reveal that the sensitivity and stability of the COF_{X-DHTA} film-based sensor are excellent for humidity.

Moreover, we conducted a humidity sensing test using the Macro-COF_{TAPB-DHTA} film (*d* = 20 mm) under the glass substrate. The dynamic response curves of the Macro-COF_{TAPB-DHTA} film-based sensor at RT with a detection

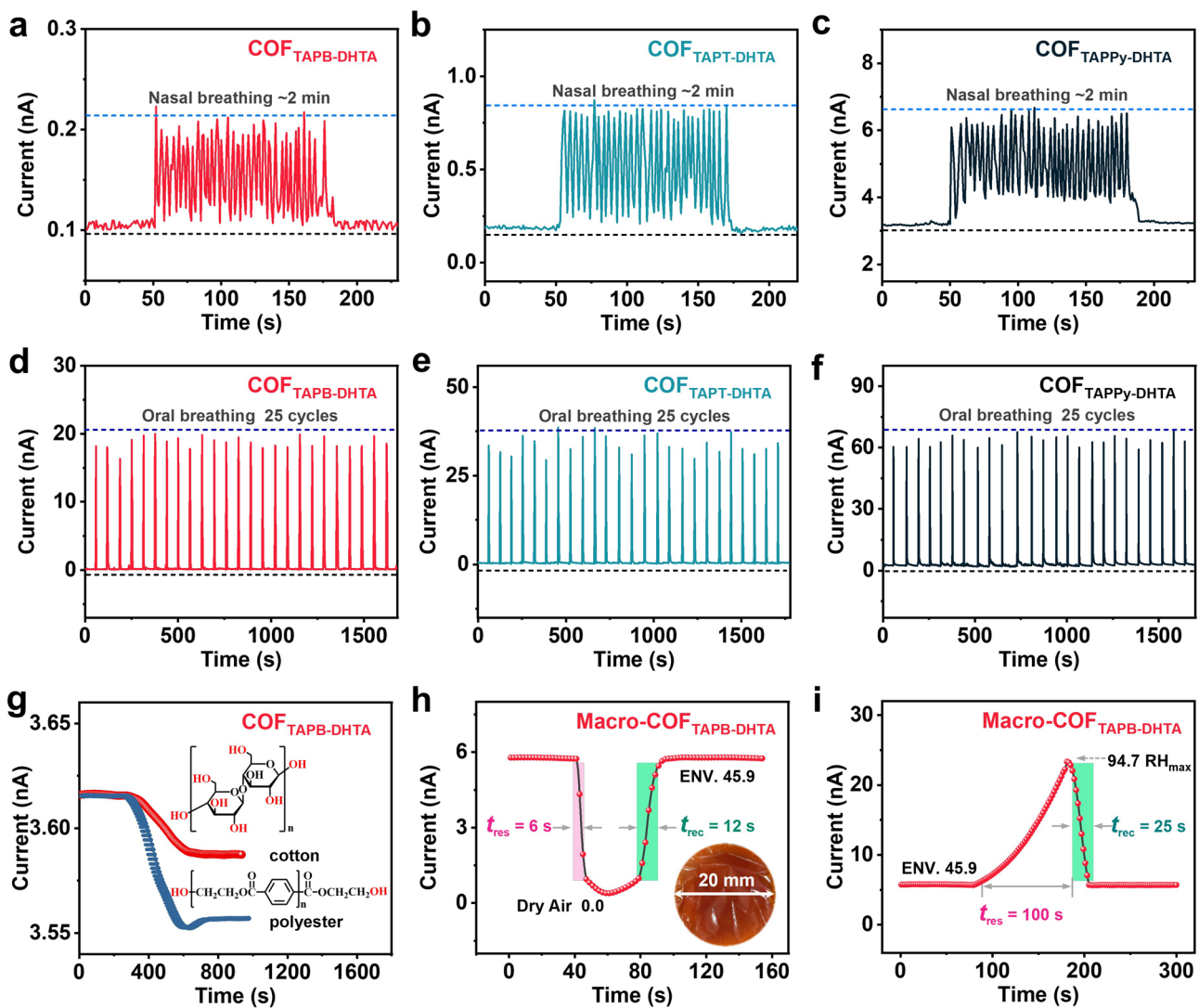


Fig. 5 Nasal breathing monitoring curves of **a** IDEs-COF_{TAPB-DHTA}, **b** COF_{TAPT-DHTA}, and **c** COF_{TAPPy-DHTA} film-based humidity sensor (the environment RH performed at 52.3%). Oral breathing monitoring curves of **d** IDEs-COF_{TAPB-DHTA}, **e** COF_{TAPT-DHTA}, and **f** COF_{TAPPy-DHTA} film-based humidity sensor (the environment RH performed at 31.7%). **g** Real-time current curve of the IDEs-COF_{TAPB-DHTA} film-based humidity sensor simulates the air permeability of clothing fabrics. Real-time dynamic humidity monitoring directly using a Macro-COF_{TAPB-DHTA} film-based humidity sensor on the glass slide detection range from **h** dry air to environment RH (0.0–45.9%) and **i** environment to maximum RH (45.9–94.7%)

range from dry air to environment RH (0.0–45.9%) and then to max RH (45.9–94.7%), respectively (Fig. 5h, i). The recovery time is a little longer compared to the IDEs-COF_{X-DHTA} film-based sensors. The main reason for this phenomenon is the accumulation of water molecules on the macroscopic film, which leads to a longer recovery time under the same atmospheric environment. A comparison of the humidity detection performance of the COF_{TAPB-DHTA} film-based sensor with previously reported studies are shown in Table S1. It can be seen that our COF film material exhibits a comprehensive sensing capability. The humidity sensor based on COF_{TAPB-DHTA} film's superior sensing performance is expected to spark new ideas for monitoring RH in human metabolite detection.

3.3 Humidity Sensing Mechanism

Numerous hydroxyl functional groups bring in the hydrophilic properties of the COF films measured by the contact angle, which is approximately 58° (Fig. S30). The higher contact area facilitates the charge transfer between COF_{X-DHTA} films and water molecules, resulting in a faster response [42]. Based on DFT calculations and in situ Raman spectra, the sensing mechanism of COF films for humidity was discussed. The 4.2.1 ORCA software was used for all calculations, and the basis was B3LYP/6–31 + G(d) with the D3BJ correction function. The data processing and image rendering was executed by Multiwfn 3.8.0 and VMD 1.9.3 [43–45]. The average local ionization energy (ALIE) distribution provides accurate and reliable analysis regarding the reaction sites and binding mode (Figs. 6 and S31). It can be seen that the ALIE minimum points of both COF_{TAPB-PDA} and COF_{TAPB-DHTA} are distributed near the nitrogen atoms of the imine bond. When water molecules are present, the N atom acts as the active site, providing the least amount of electronic constraints to form the imino/*cis*-ketoenamine isomers. Inversely, the ALIE minimum site of COF_{TAPT-DHTA} is located at 0.33 a.u. position on the benzene ring. It was attributed to the COF_{TAPT-DHTA} with an N-rich triazine structure providing strong electron-absorbing ability, and framework equilibrium electron rearrangement of the neighboring benzene ring as a more suitable active site [46]. In particular, the COF_{TAPB-DHTA} has the lowest ALIE

minimum of 0.309 a.u. compared to the other COFs. In terms of thermodynamics, it is highly favorable for water molecules to attack the imine bond in COF_{TAPB-DHTA} and lead to reciprocal isomerization (Fig. 6a, f). To further reveal the humidity sensing mechanism, the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) of COF_{TAPB-PDA}, and COF_{X-DHTA} which have demonstrated humidity-sensitive, as well as those of Water-COF_{TAPB-PDA} and Water-COF_{X-DHTA} upon water adsorption were also investigated. The energy arrangement of the donors/acceptors is pivotal for achieving an efficient driving force for separating initially generated excitons. Following this principle, the HOMO and LUMO of COF films should be delocalized and located on different nodes that serve as donors/acceptors [25].

Furthermore, the hydroxyl group of DHTA supplies additional electrons, increasing the reactivity of the COF_{TAPB-DHTA} film. The polarity of the hydroxyl group induces enhanced van der Waals force interactions between the layers, and the denser stacking of COF_{TAPB-DHTA} can contribute to the transport of water molecules (Fig. S32). The W-HOMO of COF_{TAPB-PDA} is distributed around the benzene ring and cannot induce structural heterogeneity (Fig. 6d). However, the W-HOMO and W-LUMO configurations of Water-COF_{TAPB-DHTA} film show a more delocalized level on the whole framework, and it is beneficial for improving carrier mobility (Fig. 6i, j) [47]. The entry of the water molecule leads to the iminol/*cis*-ketoenamine isomerization and induces a HOMO destabilization via conjugation along the imine linkers, which strengthens the donating ability of the imine bonds and reduces the energy gap E_g . Accordingly, the narrower E_g is obtained for Water-COF_{X-DHTA}, suggesting the enhanced electron conductivity, which conforms to the collected real-time current value being positively correlated with the RH [48]. Comparing the E_g of the COF_{X-DHTA} film and the difference after adsorbing water molecule, where COF_{TAPB-DHTA} has the largest ΔE_g is -0.19 eV, followed by COF_{TAPPY-DHTA} with -0.13 eV (Table S2). Moreover, the weak interaction allows the COF_{TAPB-DHTA} film to rapidly perform the adsorption/desorption process at RT to achieve highly sensitive and reversible humidity sensing [49]. It is also worth mentioning that the triazine structure of COF_{TAPT-DHTA} makes the ΔE_g exhibit only -0.044 eV. The extremely low ΔE_g allows COF_{TAPT-DHTA} to have a fast response value boost, but it also shows a slight perturbation

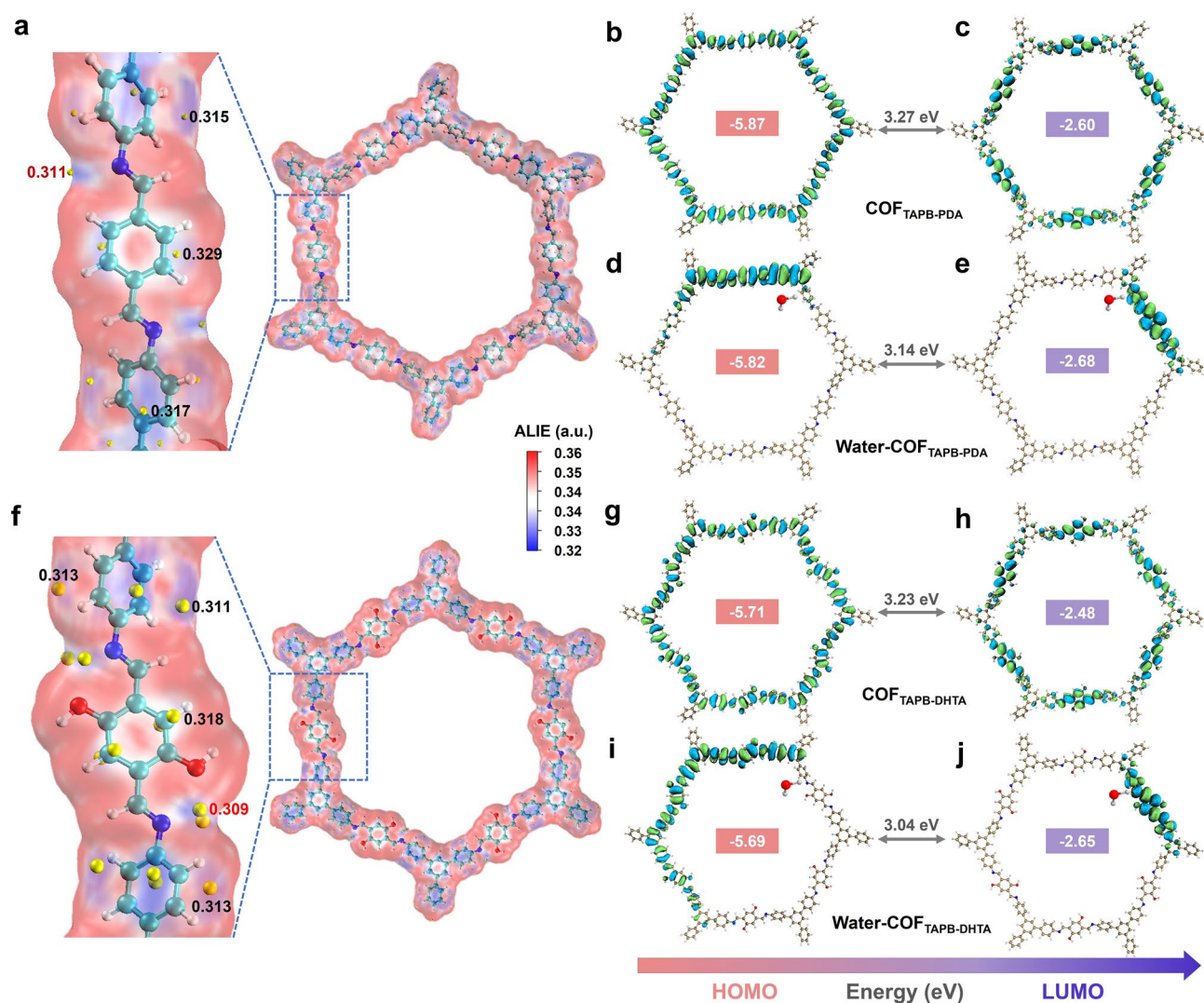


Fig. 6 **a** ALIE distribution on the surface of COF_{TAPB-PDA} film. **b** HOMO and **c** LUMO of COF_{TAPB-PDA} film. **d** W-HOMO and **e** W-LUMO of Water-COF_{TAPB-PDA} is formed by absorbing water molecules of COF_{TAPB-PDA} film. **f** ALIE distribution on the surface of COF_{TAPB-DHTA} film. **g** HOMO and **h** LUMO of COF_{TAPB-DHTA} film. **i** W-HOMO and **j** W-LUMO of Water-COF_{TAPB-DHTA} formed by absorbing a water molecule of COF_{TAPB-DHTA} film

in the light response test (Fig. S26). Analogous results are shown in the UV–Vis and UPS test spectra, as shown in Figs. S33–S35.

Subsequently, we carried out in situ Raman under different RH conditions (including dry, atmosphere, and wet vapor are 0, 45%, and 90% RH, respectively) with a 532 nm laser (Figs. 7 and S36). As shown in Fig. 7a and Table S3, the Raman shifts of the COF_{X-DHTA} skeleton are assigned to typical vibrations of different groups. All of the COF films and powders have distinct characteristic peaks around 1600 cm⁻¹, corresponding to the vibrational modes of (–C=N–) bonds, indicating that the skeleton structure of

COF films may be maintained after humidity sensing. The aldehyde group (C=O) bond of DHTA stretching vibration is around 1665 cm⁻¹ with a weak signal, which is caused by the photoluminescence phenomenon [50]. As the aforementioned humidity sensing test findings show, there is no shift in the Raman peaks of COF_{TAPB-BPDA} and COF_{TAPB-PDA} powders and films for the comparison samples (Fig. S36). The characteristic peak positions of COF_{X-DHTA} powders scarcely shift under any conditions (Fig. 7e–g).

The tautomerism generated by water molecules on the outermost surface alone is insufficient to produce a significant deflection peak of the COF_{X-DHTA} powders [51].

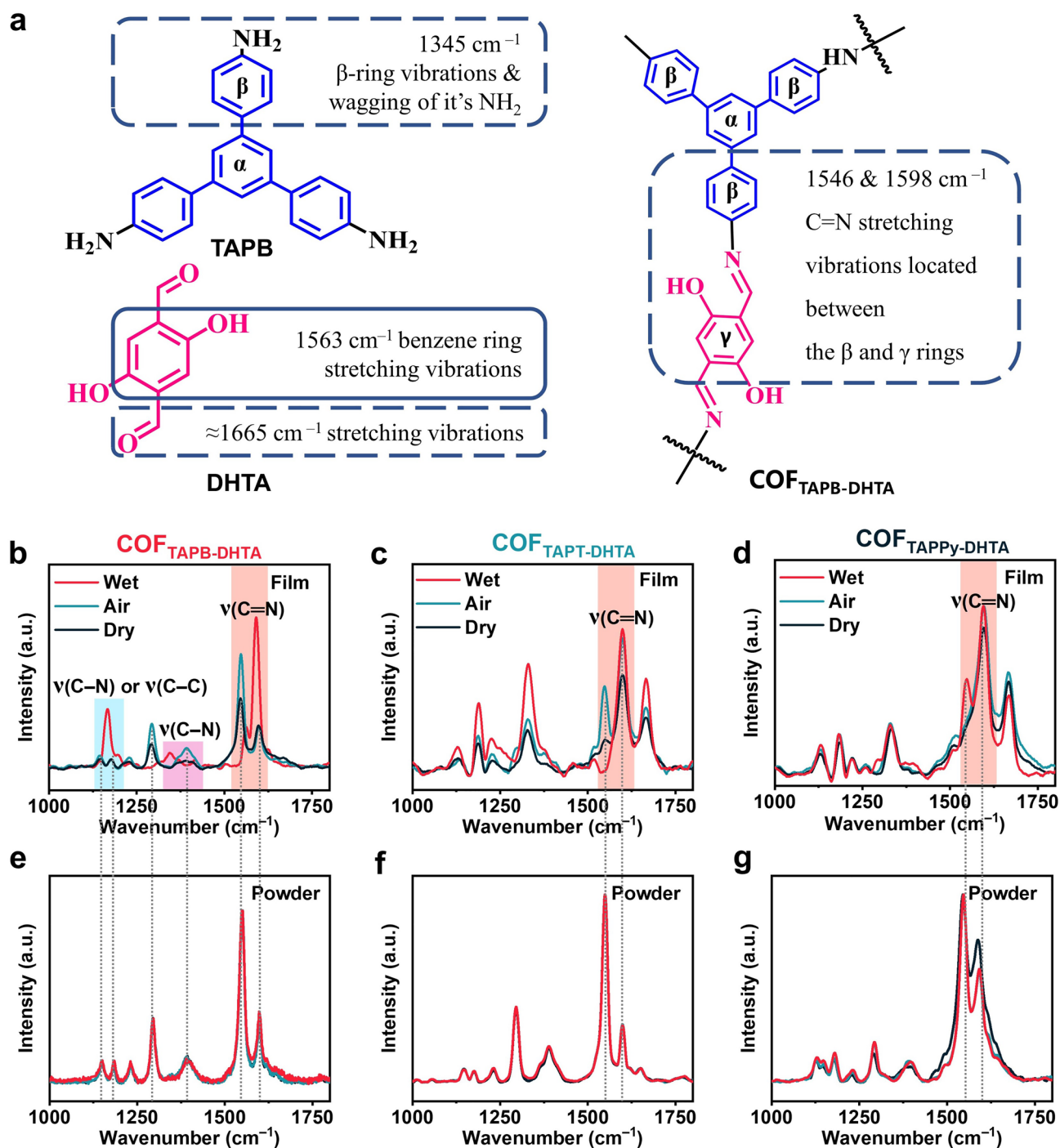


Fig. 7 **a** Raman shift assignment of $\text{COF}_{\text{TAPB-DHTA}}$. In situ Raman of **b** spectra $\text{COF}_{\text{TAPB-DHTA}}$, **c** $\text{COF}_{\text{TAPT-DHTA}}$, and **d** $\text{COF}_{\text{TAPPY-DHTA}}$ film. In situ Raman spectra of **e** $\text{COF}_{\text{TAPB-DHTA}}$, **f** $\text{COF}_{\text{TAPT-DHTA}}$, and **g** $\text{COF}_{\text{TAPPY-DHTA}}$ powder

Unlike the powder, COF film provides active sites and a conjugated conductive platform for water molecules through a large-scale and precise structural design, realizing outstanding humidity sensing performance. Under dry and air

atmosphere conditions belonging to the low humidity (black and blue curves in Fig. 7b, c, < 50% RH), the intensity difference of characteristic peaks can be observed. However, with increasing RH, the band positions shifted sharply with

the new torsional and stretching vibrations generated due to the tautomer induced by water molecules (red curves in Fig. 7b, c, 94.3% RH) [52]. The characteristic peaks of the COF_{X-DHTA} films are around 1550 cm⁻¹ (aromatic ν (CC) aromatic ring chain vibrations) and 1590 cm⁻¹ (ν (C=N) strong), representing imine bond stretching vibrations located between the β and γ rings drastically changed [53]. It implied that water molecules bind to the β -site stretching vibration close to the host framework of TAPB, proving the active site is indeed the N atoms [54]. The COF_{TAPT-DHTA} and COF_{TAPPY-DHTA} film have more prominent peaks at this position because they are increasingly electron-rich monomers than the TAPB of COF_{TAPB-DHTA} film and thus exhibit higher response values in humidity sensing tests (Fig. 7c, d). Especially, two bands at 1143.98 and 1178.54 cm⁻¹ in COF_{TAPB-DHTA} film can be assigned to ν (C–N) or (C–C) aromatic under the low RH while turning into a single peak after water vapor treatment at high RH, respectively (the light blue square in Fig. 7b) [55]. In addition, the band at 1392.07 cm⁻¹ (dry) shifts to 1344.93 cm⁻¹ (wet) assigns to the ν (C–N) of COF_{TAPB-DHTA} film vibration (pink square in Fig. 7b) [56]. It is also confirmed that COF_{TAPB-DHTA} film can generate two stretching vibration effects induced by *cis-Keto* imine tautomerism (Fig. 7b) [19]. It reasonably explains that the humidity sensor based on COF_{TAPB-DHTA} film has a more sensitive and quantitative detection ability than other sensors. In view of the above arguments, based on the above arguments, the imine groups in COF films acted as dual-active sites for humidity sensing, inducing an intrinsic enhanced mechanism of reversible protonated tautomerism via water molecule hydrogen bonding is summarized in Fig. S37.

4 Conclusions

In this work, we presented humidity-responsive COF-based films for monitoring human respiration. The COF_{X-DHTA} films exhibited excellent crystallinity and reversible tautomerism, resulting in intrinsic resistance variations while maintaining high anti-interference and long-term sensitivity. The structure–property relationship between imine reversible isomerization and ultra-sensitive sensing of COF_{TAPB-DHTA} film has been established by analyzing the functional group

species, bonding types, minimum polarity point, and N content of COF films with the support of in situ Raman spectroscopy measurement and DFT calculations. A respiration monitoring experiment was performed to generate a respiration frequency floating signal with a response/recovery time of 0.4/1 s. Additionally, this COF_{X-DHTA} film can be directly used as a sensor for humidity sensing, without any additional processing steps. This work utilizes a facile synthesis method to fabricate uniform and stable COF films, which have the potential for applications of flexible electronics and wearable devices in quantitative human respiratory monitoring. These findings are crucial for the development of new sensing materials and devices in the field of respiratory monitoring.

Acknowledgements This work was supported by the National Key Research and Development Program of China (2022YFB3205500, and 2022YFC3104700), the National Natural Science Foundation of China (62101329 and 61971284), the Shanghai Sailing Program (21YF1421400), the Natural Science Foundation of Shanghai (23ZR1430100), the Oceanic Interdisciplinary Program of Shanghai Jiao Tong University (SL2020ZD203, SL2021MS006 and SL2020MS031), Scientific Research Fund of Second Institute of Oceanography, Ministry of Natural Resources of P. R. China (SL2003), and Startup Fund for Youngman Research at Shanghai Jiao Tong University. We also acknowledge analysis support from the Instrumental Analysis Center of Shanghai Jiao Tong University and the Center for Advanced Electronic Materials and Devices of Shanghai Jiao Tong University. The computations in this paper were run on the π 2.0 cluster supported by the Center for High Performance Computing at Shanghai Jiao Tong University.

Funding Open access funding provided by Shanghai Jiao Tong University.

Conflict of Interest The authors declare no interest conflict. They have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s40820-023-01107-4>.

References

1. X. Peng, K. Dong, C. Ning, R. Cheng, J. Yi et al., All-nanofiber self-powered skin-interfaced real-time respiratory monitoring system for obstructive sleep apnea-hypopnea syndrome diagnosing. *Adv. Funct. Mater.* **31**(34), 2103559 (2021). <https://doi.org/10.1002/adfm.202103559>
2. P. Zhu, S. Li, X. Jiang, Q. Wang, F. Fan et al., Noninvasive and wearable respiration sensor based on organic semiconductor film with strong electron affinity. *Anal. Chem.* **91**(15), 10320–10327 (2019). <https://doi.org/10.1021/acs.analchem.9b02811>
3. J. Liu, J. Zhou, J. Yao, X. Zhang, L. Li et al., Impact of meteorological factors on the COVID-19 transmission: a multi-city study in China. *Sci. Total Environ.* **726**, 138513 (2020). <https://doi.org/10.1016/j.scitotenv.2020.138513>
4. L.B. Baker, J.B. Model, K.A. Barnes, M.L. Anderson, S.P. Lee et al., Skin-interfaced microfluidic system with personalized sweating rate and sweat chloride analytics for sports science applications. *Sci. Adv.* **6**(50), eabe3929 (2020). <https://doi.org/10.1126/sciadv.abe3929>
5. R. Xie, Q. Du, B. Zou, Y. Chen, K. Zhang et al., Wearable leather-based electronics for respiration monitoring. *ACS Appl. Bio Mater.* **2**(4), 1427–1431 (2019). <https://doi.org/10.1021/acsabm.9b00082>
6. C. Wang, K. Xia, H. Wang, X. Liang, Z. Yin et al., Advanced carbon for flexible and wearable electronics. *Adv. Mater.* **31**(9), e1801072 (2019). <https://doi.org/10.1002/adma.201801072>
7. X. Xiao, C. Li, S. Fan, Y. Liu, Y. Liu, Optical-thermally actuated graphene mechanical resonator for humidity sensing. *Sens. Actuators B Chem.* **374**, 132851 (2023). <https://doi.org/10.1016/j.snb.2022.132851>
8. B. Adhikari, S. Majumdar, Polymers in sensor applications. *Prog. Polym. Sci.* **29**(7), 699–766 (2004). <https://doi.org/10.1016/j.progpolymsci.2004.03.002>
9. R. Freund, O. Zaremba, G. Arnauts, R. Ameloot, G. Skorupskii et al., The current status of MOF and COF applications. *Angew. Chem. Int. Ed.* **60**(45), 23975–24001 (2021). <https://doi.org/10.1002/anie.202106259>
10. X. Chen, T. Wang, J. Shi, W. Lv, Y. Han et al., A novel artificial neuron-like gas sensor constructed from CuS quantum dots/Bi₂S₃ nanosheets. *Nano-Micro Lett.* **14**(1), 8 (2021). <https://doi.org/10.1007/s40820-021-00740-1>
11. P. Li, N. Su, Z. Wang, J. Qiu, A Ti₃C₂T_x Mxene-based energy-harvesting soft actuator with self-powered humidity sensing and real-time motion tracking capability. *ACS Nano* **15**(10), 16811–16818 (2021). <https://doi.org/10.1021/acsnano.1c07186>
12. H. Xing, X. Li, Y. Lu, Y. Wu, Y. He et al., Mxene/MWCNT electronic fabric with enhanced mechanical robustness on humidity sensing for real-time respiration monitoring. *Sens. Actuators B Chem.* **361**, 131704 (2022). <https://doi.org/10.1016/j.snb.2022.131704>
13. A.P. Côté, A.I. Benin, N.W. Ockwig, M. O’Keeffe, A.J. Matzger et al., Porous, crystalline, covalent organic frameworks. *Science* **310**(5751), 1166–1170 (2005). <https://doi.org/10.1126/science.1120411>
14. K. Geng, T. He, R. Liu, S. Dalapati, K. Tan et al., Covalent organic frameworks: design, synthesis, and functions. *Chem. Rev.* **120**(16), 8814–8933 (2020). <https://doi.org/10.1021/acs.chemrev.9b00550>
15. R.-R. Liang, S.-Y. Jiang, R.-H. A. X. Zhao, Two-dimensional covalent organic frameworks with hierarchical porosity. *Chem. Soc. Rev.* **49**(12), 3920–3951 (2020). <https://doi.org/10.1039/d0cs00049c>
16. L. Guo, L. Yang, M. Li, L. Kuang, Y. Song et al., Covalent organic frameworks for fluorescent sensing: recent developments and future challenges. *Coord. Chem. Rev.* **440**, 213957 (2021). <https://doi.org/10.1016/j.ccr.2021.213957>
17. D.D. Medina, V. Werner, F. Auras, R. Tautz, M. Dogru et al., Oriented thin films of a benzodithiophene covalent organic framework. *ACS Nano* **8**(4), 4042–4052 (2014). <https://doi.org/10.1021/nn5000223>
18. A.M. Evans, M.J. Strauss, A.R. Corcos, Z. Hirani, W. Ji et al., Two-dimensional polymers and polymerizations. *Chem. Rev.* **122**(1), 442–564 (2022). <https://doi.org/10.1021/acs.chemrev.0c01184>
19. S. Jhulki, A.M. Evans, X. Hao, M.W. Cooper, C.H. Feriante et al., Humidity sensing through reversible isomerization of a covalent organic framework. *J. Am. Chem. Soc.* **142**(2), 783–791 (2020). <https://doi.org/10.1021/jacs.9b08628>
20. T.Q. Trung, N.E. Lee, Flexible and stretchable physical sensor integrated platforms for wearable human-activity monitoring and personal healthcare. *Adv. Mater.* **28**(22), 4338–4372 (2016). <https://doi.org/10.1002/adma.201504244>
21. H. Yuan, N. Li, J. Linghu, J. Dong, Y. Wang et al., Chip-level integration of covalent organic frameworks for trace benzene sensing. *ACS Sens.* **5**(5), 1474–1481 (2020). <https://doi.org/10.1021/acssensors.0c00495>
22. S. Harpreet, T. Vijay, J. Nityasaga, B. Indu, S. Nidhi et al., A porous, crystalline truxene-based covalent organic framework and its application in humidity sensing. *J. Mater. Chem. A* **5**(41), 21820–21827 (2017). <https://doi.org/10.1039/c7ta05043g>
23. Y. Liang, Q. Ding, H. Wang, Z. Wu, J. Li et al., Humidity sensing of stretchable and transparent hydrogel films for wireless respiration monitoring. *Nano-Micro Lett.* **14**(1), 183 (2022). <https://doi.org/10.1007/s40820-022-00934-1>
24. W. Wang, W. Zhao, H. Xu, S. Liu, W. Huang et al., Fabrication of ultra-thin 2D covalent organic framework nanosheets and their application in functional electronic devices. *Coord. Chem. Rev.* **429**, 213616 (2021). <https://doi.org/10.1016/j.ccr.2020.213616>

25. N. Keller, T. Bein, Optoelectronic processes in covalent organic frameworks. *Chem. Soc. Rev.* **50**(3), 1813–1845 (2021). <https://doi.org/10.1039/d0cs00793e>
26. C. Fan, H. Wu, J. Guan, X. You, C. Yang et al., Scalable fabrication of crystalline COF membranes from amorphous polymeric membranes. *Angew. Chem. Int. Ed.* **60**(33), 18051–18058 (2021). <https://doi.org/10.1002/anie.202102965>
27. Q. Hao, C. Zhao, B. Sun, C. Lu, J. Liu et al., Confined synthesis of two-dimensional covalent organic framework thin films within superspreading water layer. *J. Am. Chem. Soc.* **140**(38), 12152–12158 (2018). <https://doi.org/10.1021/jacs.8b07120>
28. H. Jiang, Z. Guo, H. Wang, X. Liu, Y. Ren et al., Solvent-processable 0D covalent organic framework quantum dot engineered composite membranes for biogas upgrading. *J. Membr. Sci.* **640**, 119803 (2021). <https://doi.org/10.1016/j.memsci.2021.119803>
29. Z. Pang, T. Zhou, R. Liang, Q. Qi, X. Zhao, Regulating the topology of 2D covalent organic frameworks by the rational introduction of substituents. *Chem. Sci.* **8**(5), 3866–3870 (2017). <https://doi.org/10.1039/C6SC05673C>
30. K. Dey, M. Pal, K.C. Rout, S. Kunjattu, A. Das et al., Selective molecular separation by interfacially crystallized covalent organic framework thin films. *J. Am. Chem. Soc.* **139**(37), 13083–13091 (2017). <https://doi.org/10.1021/jacs.7b06640>
31. D. Zhou, X. Tan, H. Wu, L. Tian, M. Li, Synthesis of C–C bonded two-dimensional conjugated covalent organic framework films by Suzuki polymerization on a liquid–liquid interface. *Angew. Chem. Int. Ed.* **58**(5), 1376–1381 (2019). <https://doi.org/10.1002/anie.201811399>
32. S. Marco, C. Alessio, M. Manuela, Z. Massimo, C. Gabriella et al., Self-assembly of functionalized oligothiophene into hygroscopic fibers: fabrication of highly sensitive and fast humidity sensors. *Adv. Electron. Mater.* **4**(10), 1700382 (2018). <https://doi.org/10.1002/aelm.201700382>
33. B. Zhang, L. Chen, Z. Zhang, Q. Li, P. Khangale et al., Modulating the band structure of metal coordinated salen COFs and an in situ constructed charge transfer heterostructure for electrocatalysis hydrogen evolution. *Adv. Sci.* **9**(22), e2105912 (2022). <https://doi.org/10.1002/advs.202105912>
34. Q. Xu, Y. Tang, X. Zhang, Y. Oshima, Q. Chen et al., Template conversion of covalent organic frameworks into 2D conducting nanocarbons for catalyzing oxygen reduction reaction. *Adv. Mater.* **30**(15), e1706330 (2018). <https://doi.org/10.1002/adma.201706330>
35. Y. Li, W. Cui, Q. Jiang, R. Liang, X. Li et al., Arousing electrochemiluminescence out of non-electroluminescent monomers within covalent organic frameworks. *ACS Appl. Mater. Interfaces* **13**(40), 47921–47931 (2021). <https://doi.org/10.1021/acsaami.1c12958>
36. J. Zhao, J. Ren, G. Zhang, Z. Zhao, S. Liu et al., Donor-acceptor type covalent organic frameworks. *Chem. Eur. J.* **27**(42), 10781–10797 (2021). <https://doi.org/10.1002/chem.202101135>
37. S. Kano, K. Kim, M. Fujii, Fast-response and flexible nanocrystal-based humidity sensor for monitoring human respiration and water evaporation on skin. *ACS Sens.* **2**(6), 828–833 (2017). <https://doi.org/10.1021/acssensors.7b00199>
38. J. Liang, X. Zhang, J. Ji, Hygroscopic phase change composite material—a review. *J. Energy Storage* **36**, 102395 (2021). <https://doi.org/10.1016/j.est.2021.102395>
39. H. Li, X. Li, X. Li, C. Chong, J. Jin et al., Multifunctional smart mask: enabling self-dehumidification and self-powered wearables via transpiration-driven electrokinetic power generation from human breath. *Chem. Eng. J.* **461**, 142083 (2023). <https://doi.org/10.1016/j.cej.2023.142083>
40. Z. Zhen, Z. Li, X. Zhao, Y. Zhong, L. Zhang et al., Formation of uniform water microdroplets on wrinkled graphene for ultrafast humidity sensing. *Small* **14**(15), e1703848 (2018). <https://doi.org/10.1002/sml.201703848>
41. R. Wei, D. Breite, C. Song, D. Gräsing, T. Ploss et al., Biocatalytic degradation efficiency of postconsumer polyethylene terephthalate packaging determined by their polymer microstructures. *Adv. Sci.* **6**(14), 1900491 (2019). <https://doi.org/10.1002/advs.201900491>
42. Y. Zhang, J. Guo, G. Han, Y. Bai, Q. Ge et al., Molecularly soldered covalent organic frameworks for ultrafast precision sieving. *Sci. Adv.* **7**(13), eabe8706 (2021). <https://doi.org/10.1126/sciadv.abe8706>
43. W. Humphrey, A. Dalke, K. Schulten, VMD: visual molecular dynamics. *J. Mol. Gr.* **14**, 33–38 (1996). [https://doi.org/10.1016/0263-7855\(96\)00018-5](https://doi.org/10.1016/0263-7855(96)00018-5)
44. F. Neese, The ORCA program system. *WIREs Comput. Mol. Sci.* **2**(1), 73–78 (2011). <https://doi.org/10.1002/wcms.81>
45. T. Lu, F. Chen, Multiwfn: a multifunctional wavefunction analyzer. *J. Comput. Chem.* **33**(5), 580–592 (2012). <https://doi.org/10.1002/jcc.22885>
46. H. Zhang, L. Zhang, S. Dong, X. Duan, D. Zhu et al., Regulating energy band structures of triazine covalent organic frameworks with electron-donating/withdrawing substituents for visible-light-responsive photocatalytic tetracycline degradation and Cr(VI) reduction. *J. Hazard. Mater.* **446**, 130756 (2023). <https://doi.org/10.1016/j.jhazmat.2023.130756>
47. S. Liu, X. Zhou, W. Han, J. Li, X. Sun et al., Theoretical and experimental insights into the OH-mediated mineralization mechanism of flutriafol. *Electrochim. Acta* **235**, 223–232 (2017). <https://doi.org/10.1016/j.electacta.2017.03.062>
48. Q. Wei, K. Paul, K. Yevhen, X. Qiu, K. Hartmut et al., Conjugation-induced thermally activated delayed fluorescence (TADF): from conventional non-TADF units to TADF-active polymers. *Adv. Funct. Mater.* **27**(7), 1605051 (2017). <https://doi.org/10.1002/adfm.201605051>
49. J. Xu, S. An, X. Song, Y. Cao, N. Wang et al., Towards high performance Li–S batteries via sulfonate-rich COF-modified separator. *Adv. Mater.* **33**(49), e2105178 (2021). <https://doi.org/10.1002/adma.202105178>
50. X. Xu, R. Xiong, Z. Zhang, X. Zhang, C. Gu et al., Space-partitioning and metal coordination in free-standing covalent organic framework nano-films: over 230 mWh/cm³ energy density for flexible in-plane micro-supercapacitors. *Chem.*



- Eng. J. **447**, 137447 (2022). <https://doi.org/10.1016/j.cej.2022.137447>
51. W. Xu, Y. Sun, X. Dong, S. Li, H. Wang et al., Local order and vibrational coupling of the C=O stretching mode of gamma-caprolactone in liquid binary mixtures. *Sci. Rep.* **7**(1), 12182 (2017). <https://doi.org/10.1038/s41598-017-12030-1>
52. C.R. DeBlase, K.H. Burgos, K.E. Silberstein, G.G. Rodríguez, R.P. Bisbey et al., Rapid and efficient redox processes within 2D covalent organic framework thin films. *ACS Nano* **9**(3), 3178–3183 (2015). <https://doi.org/10.1021/acs.nano.5b00184>
53. S.A. Ahmed, Q. Shen, Q. Liao, J. Zhou, S. Hanif et al., Mass transfer modulation and gas mapping based on covalent organic frameworks-covered theta micropipette. *Anal. Chem.* **92**(10), 7343–7348 (2020). <https://doi.org/10.1021/acs.analchem.0c01152>
54. L. Zhang, Y. Shi, Y. Wang, N.R. Shiju, Nanocarbon catalysts: recent understanding regarding the active sites. *Adv. Sci.* **7**(5), 1902126 (2020). <https://doi.org/10.1002/adv.201902126>
55. L. Zhang, T. Ritter, A perspective on late-stage aromatic C–H bond functionalization. *J. Am. Chem. Soc.* **144**(6), 2399–2414 (2022). <https://doi.org/10.1021/jacs.1c10783>
56. D. Zhang, P. Ren, W. Liu, Y. Li, S. Salli et al., Photocatalytic abstraction of hydrogen atoms from water using hydroxylated graphitic carbon nitride for hydrogenative coupling reactions. *Angew. Chem. Int. Ed.* **61**(24), e202204256 (2022). <https://doi.org/10.1002/anie.202204256>