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Formation and exploration of polymorphism in WO_3 nanostructures for improved catalytic and chemical sensing performance†

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Nowadays, the development of functional sensing materials to detect gaseous and volatile organic compounds is of great importance for health and the environment. Doping and functionalization of semiconductor materials, as well as the synthesis of composites, are the most considered methods to improve their sensing performance. In this respect, polymorphic transitions in semiconductor metal oxides and subsequent structural changes offer alternative and efficient strategies to tune the kinetics of redox reactions at their surface. Thus, the polymorphism in transition oxide nanostructures can lead to a significant improvement in their catalytic properties. Here, monoclinic/orthorhombic polymorphic WO₃ nanomaterials are synthesized by the precipitation method. Experimental studies and theoretical modeling of the prepared structures are carried out. The results indicate that the composition of the solvent is a crucial factor in the formation and shape of WO₃ nanoparticles. The use of polyethylene glycol as a surfactant increases the degree of structural disorder, suppresses the formation of hydrates, and affects the oxygen content in WO₃. These effects, in combination with the formation of monoclinic/ orthorhombic n-n junctions in the WO₃ polymorph, dramatically increase its sensitivity to acetone molecules. In particular, the response of the synthesized WO₃ polymorph is up to 8 times greater compared to the single-phase monoclinic structure. Hence, appropriate modification of the WO_₹ crystal structure and formation of monoclinic/orthorhombic junctions in the nanomaterial can significantly improve its sensing performance to acetone without using dopants, mixture materials, or catalytic layers. The performed experimental studies and theoretical modeling to describe the influence of monoclinic/ orthorhombic polymorphism on the sensing performance of WO₃ are important and may provide new insights into its applications in monitoring systems and heterogeneous catalysis.

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Introduction

Semiconductor nanomaterials, with their large surface area and tunable electronic properties, are among the most studied

structures for the development of heterogeneous catalytic systems. These materials generally exhibit promising properties leading to their application in chemical gas sensing, light harvesting, energy storage and conversion.1-5 Health and safety issues related to air pollution and the ever-growing industrial activities are increasing the need for chemical gas sensors based on semiconductor materials widely used for air quality monitoring and medical diagnosis. 6-9 The adsorption and desorption reactions of oxidizing and reducing gaseous compounds on the surface of semiconductor nanostructures lead to a change in their electrical conductivity. This characteristic feature enables the fabrication of small-size and easy-to-use sensing devices. 10-12 Various approaches, such as functionalization by catalytic layers and the synthesis of composite structures, have been proposed to improve the redox reactions at the surface of semiconductors.13-16 Furthermore, the synthesis of heterojunctions with improved charge separation and transport properties dramatically impacts the catalytic activity of gas sensors.2,8,17-20 In this context, polymorphism

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semiconductors is of great interest, and it is currently considered one of the most effective strategies for developing heterogeneous catalytic systems. The formation of phase junctions in polymorphic materials can lead to an improvement in their physicochemical properties.²¹⁻²³ Recently, it has been demonstrated that differences in band energy between phases stimulate the separation of photogenerated charge carriers in the oxide material and significantly enhance its photocatalytic activity. Here, the polymorphic phase transformation in TiO2 and the formation of the anatase-rutile phase junction affected the concentration of surface vacancies, changed the bandgap, and enhanced the separation and mobility of the charge carriers.22 This led to a drastic increase in the photocurrent of TiO2 and to the improvement of its photocatalytic activity for converting CO₂ to CO without adding impurities to the material. Thus, the proper formation of different crystalline phases in the same material makes it possible to avoid the introduction of additives and expensive catalytic materials to improve the electronic properties and reactivity.

WO₃ is a versatile material used in sensors and photocatalytic devices.²⁴⁻²⁷ It can be crystallized in different phases, such as triclinic, monoclinic, orthorhombic, hexagonal, and tetragonal, depending on the synthesis and annealing conditions.^{28,29} The experimental and theoretical findings show that the tunable crystalline structure of the material is a crucial factor for its catalytic applications.^{27,30,31} Therefore, WO₃ with different crystalline phases is widely used in the oxidation and decomposition of gaseous compounds.^{32,33} Research studies on the addition of suitable elements to the structure of WO₃ nanomaterials to improve their functionality have already achieved considerable success.^{25,34,35} Furthermore, coupling WO₃ with other materials and forming heterostructures with efficient charge separation leads to improved catalytic properties of the composite.³⁶⁻³⁹

Recent studies suggest that deeper insights into the structural polymorphism of transition metal oxide nanomaterials offer new opportunities to tune their electronic band structure and the kinetics of redox reactions at their surface. 22,40 Thus, the preparation of polymorphs could be an alternative to optimize the gas sensing response of pure WO₃. Theoretical and experimental studies show that the monoclinic WO3, with its acidity, is favorable for the interaction with acetone molecules and their decomposition.41-44 However, the activation energy for the chemisorption of acetone on the material and its decomposition is very high. Therefore, the optimum operating temperature of gas sensors based on pure WO₃ is around 400 °C. 42,45-47 Furthermore, a transition from monoclinic to orthorhombic phases in WO₃ is possible at these temperatures.²⁸ Although there is a great interest in WO₃ nanostructures, the chemical gas sensing properties of the monoclinic/orthorhombic WO₃, considering the role of phase transition, are not yet known. Therefore, the challenges of gas sensing response of the WO₃ polymorph still need to be addressed.

The use of surfactants for synthesizing WO₃ nanomaterials can play a crucial role in their crystalline properties. 48,49 Such

studies can make an important contribution to the development of structures with enhanced sensitivity based on the use of WO₃ polymorphism.

The aim of this work is to synthesize and investigate the gas sensing characteristics of monoclinic/orthorhombic polymorphic WO₃ nanomaterials. Thus, we evaluate the properties of monoclinic/orthorhombic junctions in WO₃ polymorph for application in chemical gas sensing. The structures were prepared by the precipitation method. The influence of synthesis parameters on the formation and shape of the material was analyzed. The gas sensing performance of monoclinic/ orthorhombic WO3 polymorph nanostructure has been comprehensively discussed considering its compositional, morphological, and structural properties. For the first time, the influence of oxygen and hydrate content, as well as the formation of monoclinic/orthorhombic n-n junctions on the charge transfer and chemical sensing characteristics of the material, is discussed in detail based on experimental studies and theoretical modeling. The obtained results are an important contribution to the development of chemical gas sensing materials with high sensitivity and selectivity, as well as to the production of highly efficient catalytic systems.

2. Experimental

2.1 Synthesis of materials

Tungsten(vi) chloride (CAS number: 13283-01-7, Sigma-Aldrich), benzyl alcohol (CAS number: 100-51-6, Sigma-Aldrich), and polyethylene glycol 200 (CAS number: 25322-68-3, Sigma-Aldrich) were used as reagents for the synthesis of the materials. Two solutions were used for preparing WO3 nanostructures: (i) 1 g of tungsten(vi) chloride (WCl₆) was dissolved in 50 ml of benzyl alcohol and stirred at room temperature for 30 minutes. (ii) The second solution was prepared similarly to the first. However, in this case, 0.2 g of polyethylene glycol 200 (PEG) was added dropwise to the prepared WCl₆ solution. Then both solutions were heated at 80 °C for 13 hours. Subsequently, the resulting powders were washed with ethanol (CAS number: 64-17-5, Sigma-Aldrich) by centrifugation and dried at room temperature. Finally, the precipitates were calcined at 450 °C for 2 hours. In this work, the powder prepared in benzyl alcohol solution with WCl₆ is referred to as WO3_P_1, and the one synthesized in benzyl alcohol solution with WCl6 and PEG 200 is referred to as $WO_3_P_2$.

2.2 Characterization

The crystalline structure of the fabricated materials was investigated using X-ray diffraction spectroscopy (XRD), electron diffraction, and Raman spectroscopy. The Raman spectra were recorded with a Jasco NRS-5100 confocal Raman microscope equipped with a diode laser at 785 nm and a grating of 600 l mm $^{-1}$, operating at a nominal power of 20 mW. An MPLFLN $100\times$ objective focuses the laser beam onto the sample with a 1

μm spot. The backscattered signal is collected with a cooled (-69 °C) CCD camera. The measurements were carried out in the range of 159–1890 nm with an effective power of 6.0 mW (accumulations: 50, exposure value: 7 s, resolution: 3 cm⁻¹). We performed four spectral measurements for each material and the spectra were processed (baseline correction and data smoothing) using the Spectra ManagerTM operating system connected to the Jasco Raman instrument. The second derivative was calculated to detect the spectrum maxima positions.

Attenuated total reflectance (ATR) Fourier-transform infrared (IR) spectra were performed using a FTIR Vertex70 interferometer (Bruker Optics, Ettlingen, Germany) equipped with a thermal source (Globar), an ATR crystal module with a single reflection diamond crystal at 45°, and an RT-DLaTGS wide range detector. The measurements were carried out under vacuum conditions. The IR spectra of WO₃ nanomaterials were acquired at an acquisition rate of 7 kHz in the spectral region of 50–6000 cm $^{-1}$, averaging 128 scans collected at a spectral resolution of 4 cm $^{-1}$. Second-derivative analysis was performed by applying a 9-point smoothing filter. 50 ATR-IR data pre- and postprocessing were performed using OPUSTM 8.2.

The morphological analysis of the materials was carried out by a field-emission scanning electron microscope (FESEM, MIRA3 FEG-SEM, TESCAN) and a transmission electron microscope (TEM, F-200, JEOL) equipped with a GATAN RIO16 CMOS camera and an energy dispersive X-ray (EDX) spectrometer for elemental analysis.

X-ray (XPS) and ultraviolet (UPS) photoemission spectra were acquired by using an Escalab 250Xi spectrometer (Thermo Fisher Scientific Ltd, East Grinstead, UK) with a monochromatic Al Ka (1486.6 eV) excitation source. The measurements were performed in an ultra-high vacuum at a base pressure of about 1×10^{-9} mbar, which increased to 1 \times 10⁻⁸ during the UPS measurements. The spectra were collected at pass energies of 40 and 5 eV for XPS and UPS, respectively, in the standard mode of the electromagnetic lens system, corresponding to an analysis area of approximately 1 mm in diameter. The binding energy (BE) scale was corrected for a low sample charging by positioning the main C 1s peak of aliphatic carbon at BE = 285.0 eV and verifying that the Fermi level corresponds to BE = 0 eV. The work function of the samples was calculated from the cutoff energy at zero intensity in the valence band spectra acquired by using He I source. A series of negative bias voltages in the range of 2-20 V was applied to shift these spectra from the spectrometer threshold. Spectroscopic data were processed by Avantage v.5 software (Thermo Fisher Scientific Ltd).

To perform gas sensing studies, the materials were deposited on alumina substrates (dimensions: 2 mm \times 2 mm \times 0.75 mm, Kyocera, Japan) using the drop-casting method. Initially, platinum (Pt) interdigitated electrical contacts and a Pt heater were deposited on the surface and the rear side of the substrates using DC magnetron sputtering. Next, the synthesized powdered materials were dispersed in distilled water (H₂O) and

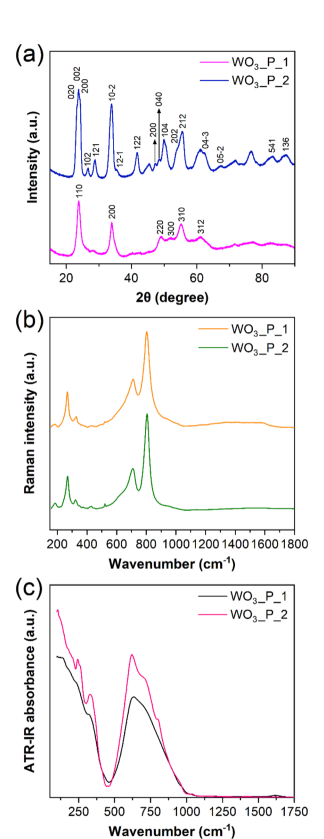


Fig. 1 (a) XRD patterns of WO $_3$ _P_1 and WO $_3$ _P_2 samples. (b) Raman spectra of WO $_3$ _P_1 and WO $_3$ _P_2. (c) ATR-IR spectra of WO $_3$ _P_1 and WO $_3$ _P_2.

drop-cast on the alumina substrates. A Gilson dispenser was used for drop casting. Gas sensing analyses were conducted using the flow-through technique in a thermostatic test chamber, where the humidity level was regulated with a humidity sensor. The gas flow rate was 0.3 l min⁻¹. The concentration of each analyte was obtained by mixing the synthetic air with the gas. The sensing response of fabricated structures to reducing gases was calculated according to the following equation:

$$S = rac{\left(G_{
m f} - G_0
ight)}{G_0} = rac{\Delta G}{G_0}$$

herein, G_0 is the conductance of material in air and G_f is the steady-state conductance value of the structure in the presence of gas. The response to oxidizing gases was defined as the relative variation of the electrical resistance of sensors (R_0 : the resistance of the sensor in air, R_f : the steady-state resistance in the presence of an analyte):

$$S=rac{\left(R_{
m f}-R_0
ight)}{R_0}=rac{\Delta R}{R_0}$$

2.3 DFT calculations

Theoretical modeling of the band structure was performed based on the *ab initio*-density functional calculations⁵³ with plane wave basis and norm-conserving pseudopotentials (CASTEP code).⁵⁴ The hybrid PBE0 exchange-correlation functional⁵⁵ was used, and the plane wave basis set was limited to a cut-off energy of 800 eV. To sample the Brillouin zone for the W_4O_{12} unit cell, a Monkhorst–Pack grid $2 \times 2 \times 2$ was applied,⁵⁶ resulting in two irreducible *k*-points.

Results and discussion

3.1 Structural and morphological analysis

Fig. 1(a) shows the XRD patterns of WO₃_P_1 and WO₃_P_2 materials. The XRD pattern of WO₃_P_1 reveals the following main diffraction peaks at 24.1°, 33.96°, 49.36°, 51.96°, 55.13°, and 61.27°, which can be assigned to the (110), (200), (220), (300), (310), and (312) planes of monoclinic phase (JCPDS no. 98-003-9121, space group: P1c1 and space group number: 7), respectively. In the XRD pattern of WO₃_P_2, the peaks at 23.228°, 23.801°, 24.495°, 26.777°, 28.858°, 50.006°, 80.297°, 87.111° are indexed as (020), (002), (200), (102), (121), (104), (541), (136) planes of orthorhombic phase (JCPDS no. 98-004-9836, space group: Pbcn and space group number: 60). The ones at 33.951°, 35.651°, 41.802°, 47.404°, 48.430°, 54.035°, 55.509°, 62.474° , and 67.365° correspond to the (10-2), (12-1), (122), (200), (040), (202), (212), (04-3), and (05-2) planes of monoclinic phase (JCPDS no. 98-007-1692, space group: P12₁/c1 and space group number: 14), respectively. The peak intensities of WO₃_P_2 are sharper and narrower than those of WO₃_P_1, which indicates that PEG 200 enhances the crystallinity of WO₃ nanomaterial. Hence, the application of PEG in the synthesis

Table 1 Raman peak positions and chemical group assignment of WO_3 _P_1 and WO_3 _P_2 nanomaterials

Frequency position (cm ⁻¹) Sample WO ₃ _P_1	Frequency position (cm ⁻¹) Sample WO ₃ _P_2		References
268	270	$\delta (ext{O-W-O})$	59, 62 and 64
328	324	O–W–O deformation mode ν (W–OH ₂)	62, 64 and 66 59
370	353		
	376 397	Orthorhombic	64
430	431		59
522	521	Si-O	
	616		
	633	Orthorhombic	64
713	713	ν(O–W–O) Monoclinic	42 and 64 59
804	806	Stretching (O-W-O)	59 and 64
940	940	Stretching (W=O)	63 and 64

procedure affects the formation of crystalline WO_3 nanostructure. 48,57,58

Raman spectra of WO₃_P_1 and WO₃_P_2 are shown in Fig. 1(b). They both exhibit main vibrational bands at 184, 324, 710, and 805 cm⁻¹, which are generally assigned to the monoclinic phase. The bands located at around 805 and 710 cm^{-1} can be ascribed to the stretching vibration of O-W-O,59-63 while the one at 324 cm⁻¹ corresponds to the O-W-O bending vibration. The peak at 184 cm⁻¹ is attributed to the lattice mode, and it is possible to observe a shift to higher frequencies for the $WO_3_P_2$ sample. Furthermore, unlike the $WO_3_P_1$ material, peaks and shoulders corresponding to the orthorhombic phase are identified in the spectrum of WO_{3_P_2} (Table 1). Here, three main vibrational bands are observed at 269, 376, and 632 cm⁻¹. The bands at 632 and 376 cm⁻¹ can be assigned to the stretching vibration of the bridging oxygen O-W-O,64 while the peak at 269 cm⁻¹ is associated with W-O-W deformation. 59,64 The highest frequency peak at 940 cm⁻¹ present in both samples, can be attributed the W=O stretching vibration of the terminal oxygen,63,65 which is usually not present in threedimensional structures, such as orthorhombic monoclinic.64

The ATR-IR spectra of WO₃_P_1 and WO₃_P_2 are reported in Fig. 1(c), while the absorption frequencies are listed in Table 2 with assignment to the corresponding vibrational modes. The spectrum of the sample WO₃_P_1 recorded in the far-IR region shows main characteristic bands at 348 cm⁻¹,

Paper

Table 2 Positions of IR absorption peaks and their assignment to vibrational modes of functional groups in $WO_3_P_1$ and $WO_3_P_2$ nanomaterials

Frequency position (cm ⁻¹)	Frequency position (cm ⁻¹)		
Sample WO ₃ _P_1	Sample WO ₃ _P_2	Assignment	
57	59	Lattice modes	
72	75	Lattice modes	
87	_	Lattice modes	
_	92	Lattice modes	
105	105	Lattice modes	
120	120	Lattice modes	
139	137	Lattice modes	
145	148	Lattice modes	
_	178	Lattice modes	
182	_	Lattice modes	
194	194	Lattice modes	
_	212	Lattice modes	
234	_	ν (W–O–W)	
	243	ν (W–O–W)	
326	323	$\nu(W-OH_2)$	
348	343	δ(W-O)	
369	370	$\nu(W-OH_2)$	
_	417	Water libration	
430	_		
_	453		
	554		
602	601		
618	617	(0)	
633	633	ν(W-O)	
715	717	$\nu(W-O)$	
730	_	ν(W-O)	
756	753	ν (O–W–O)	
787	_		
016	806	-(O W O)	
816		ν(O-W-O)	
	869	ν (O–W–O)	
	926	"(W—O)	
070	961	$\nu(W=0)$	
970	1012	$\nu(W=O)$	
1020	1012 1038		
1039			
	1093		
	1130		
	1184	OH in plane bend	
	1272 1343	OH in plane bend	
1377	1949	OH-bend	
1413	 1408	Deformation OH	
1413	1436	Deformation Off	
	1481	C=C-C aromatic ring	
	1401	stretching due to PEG	
1540		saccoming due to I Ed	
1600		Deformation OH	
1621	1621	Deformation OH	
1805	1804	Aromatic combination bands	
1828	1828	Aromatic combination bands	
1947	1946	Aromatic combination bands	
	2660	moment combination bands	
2848	2000	ν(C-H)	
	2863	ν (C–H) due to PEG	
2871	_000	ν(C-H) ddc to 1 EG ν(C-H)	
2918		ν(C-H)	
	2928	ν(C-H)	

Table 2 (Contd.)

Frequency position (cm ⁻¹)	Frequency position (cm^{-1})	
Sample WO ₃ _P_1	Sample WO ₃ _P_2	Assignment
2955	2958	ν(C-H)
3234	3234	ν(O-H)
3343	3343	ν(O-H)
3545	3545	ν (O–H) due to WO ₃ ·2H ₂ O
3560		ν(O-H)

326 cm⁻¹, and 234 cm⁻¹, which can be assigned to δ (W-O), $\nu(W-OH_2)$ and $\nu(W-O-W)$ vibrations, respectively. ^{59,67-69} The band at 369 cm⁻¹ is attributed to the vibrational modes of water molecules. 67,69 The IR bands, located in the very low-frequency region (below 200 cm⁻¹) can be ascribed to lattice modes of WO₃ and its hydrates.⁶⁷ In particular, more intense ones are located at 182, 145, 120, and 105 cm⁻¹. The dominant broad band at 633 cm⁻¹, as well as the shoulders at 714 and 730 cm⁻¹, can be ascribed to the $\nu(W-O)$ vibrational modes.^{67,69} The two shoulders identified by 2nd derivative analysis 50,70 are located around 816 and 970 cm⁻¹, and indexed as ν (O-W-O) and ν (W=O), respectively.⁶⁸ Furthermore, the ATR-IR spectrum of the sample WO₃_P_1 shows two IR bands at 1621 and 1600 cm⁻¹ corresponding to the bending mode of H₂O molecules, which can be attributed to the water content in the sample.71,72

The ATR-IR spectrum of the WO₃_P_2 sample in the far-IR region shows main characteristic bands at 343 cm⁻¹ (δ (W–O)), 323 cm⁻¹ (ν (W-OH₂)), and 243 cm⁻¹ (ν (W-O-W)).^{59,67-69} The band at 370 cm⁻¹ corresponds to the vibration modes of water molecules. 67,69 In the low-frequency region, main absorption bands due to WO3 lattice modes and its hydrates are located at 178, 148, 120, and 105 cm⁻¹.67 The intense broad band at 633 cm⁻¹ and the evident shoulder at 718 cm⁻¹ correspond to the ν (W–O) vibrational modes.^{67,69} Two shoulders located around 753 and 869 cm⁻¹ are indexed as ν (O-W-O) vibrations and can be ascribed to the bridging vibration of the cornersharing WO₆ octahedron in the WO₃ crystal.^{68,73} The weak bands in the spectral regions of 1450-1510 cm⁻¹, 1310-1410 cm⁻¹, and 1260-1350 cm⁻¹ are associated with C=C-C aromatic ring stretching, O-H bend, and O-H in plane bend (observed in phenol), respectively. In particular, the bands located at 1481, 1343, and 1272 cm⁻¹ can be attributed to the PEG 200.74,75 The ATR-IR spectrum of WO₃_P_2 structure also shows an IR band at 1621 cm⁻¹ due to the water content in the sample.

Considering Fig. 1(c), slight differences between $WO_3_P_1$ and $WO_3_P_2$ spectra can be recognized in the 600–1000 cm⁻¹ spectral range. It is well established that all WO_3 polymorphs have vibrational spectra that differ from each other. However, most of the works in the literature do not describe the low-frequency range of 200–500 cm⁻¹.69,72,76 Meanwhile, Fig. 1(c) clearly shows that the most significant differences between

WO₃_P_1 and WO₃_P_2 materials are observed in the low-frequency range (50–500 cm⁻¹). Here, both samples show absorption bands at 57, 72, 105, 120, 138, 145, 194, 323, and 370 cm⁻¹. Moreover, some significant differences in peak positions are observed between the WO₃_P_1 and WO₃_P_2 samples. The band located at 87 cm⁻¹ (lattice mode) in the spectrum of WO₃_P_1 shifts to a higher frequency (92 cm⁻¹) in the spectrum of WO₃_P_2, while the peak at 182 cm⁻¹ shifts to a lower frequency (178 cm⁻¹) in the spectrum of WO₃_P_2. The absorption band at 234 cm⁻¹ in the spectrum of WO₃_P_1 (due to ν (W–O–W) vibration) shifts to a higher frequency (243 cm⁻¹) in that of WO₃_P_2. Finally, the absorption due to the W–O bending located at 348 cm⁻¹ in the spectrum of WO₃_P_1 sample shifts to 343 cm⁻¹ in that of WO₃_P_2. Furthermore,

between the spectra of the materials, strong differences are evident in the mid-IR region, which can be attributed to the application of PEG 200 as a surfactant.^{74,75}

The H_2O bending mode signal located at 1621 and 1600 cm⁻¹ is more intense in the spectrum of $WO_3_P_1$ compared to that of $WO_3_P_2$, revealing a major water content in the sample prepared without using PEG. Besides, both samples exhibit vibrational bands above 3000 cm⁻¹ due to the CH stretching of the benzyl alcohol solvent.

Fig. 2(a) and (b) show the surface morphologies of WO₃_P_1 and WO₃_P_2 materials obtained by FESEM. As can be seen, the structures consist of nanoparticles. It is worth noting that metal chloride, such as WCl₆, exhibits a strong polarization effect, and its high valence metal ions can be easily attacked by

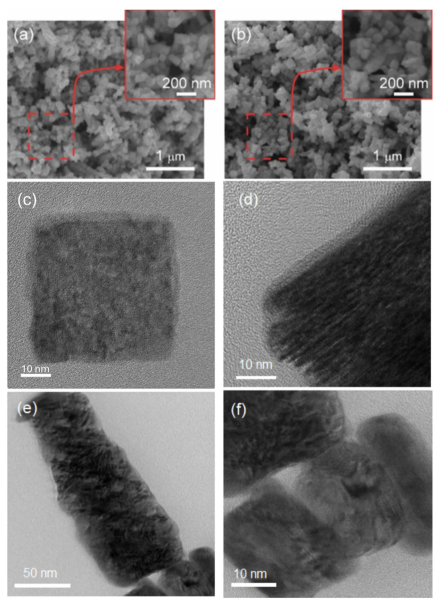


Fig. 2 (a) and (b) FESEM images of WO_3 nanostructures: (a) the morphology of WO_3 –P_1 material and (b) the morphology of WO_3 –P_2 material. (c) and (d) HRTEM images of a square nanoparticle WO_3 –P_1: (c) WO_3 nanoparticle showing a very thin layer of amorphous material, and (d) another HRTEM image showing that the entire nanoparticle structure appears to be composed of a large number of very small nanorods. (e) and (f) HRTEM images of WO_3 –P_2 nanoparticles.

a nucleophilic agent, resulting in high reactivity. Therefore, controlling the precipitation of WO₃ in a protic solvent is difficult.77,78 The obtained morphological observations indicate that benzyl alcohol is a very suitable solvent to control the reaction rate of WCl₆ and the formation of particles. Benzyl alcohol can act as a solvation medium, oxygen supplier, and capping agent.⁷⁹⁻⁸¹ During our experimental process, the reaction of WCl₆ with benzyl alcohol leads to a color change from yellow to orange, light blue, and dark blue, which may indicate the formation of different charge transfer complexes by ligand exchange on tungsten atoms. 82,83 The color change to dark blue illustrates the partial reduction of WVI.82 Furthermore, the formation of WO3 particles without agglomeration can be attributed to the stability of the carbocation intermediate formed during the reaction due to the direct bonding of the benzene ring with the α -carbon.⁷⁸

Fig. 2(c) and (d) report high-resolution TEM (HRTEM) images of WO₃_P_1. In this case, the WO₃ nanoparticles are mainly square/rectangular and are surrounded by a very thin layer of amorphous material (about 3 nm). According to the results reported below (X-ray distribution of the elements, Fig. 3), this layer may be due to some residual ethanol used to cast the material on the TEM grid. In addition, as shown in

Fig. 2(d), the entire nanoparticle consists of a large number of very thin nanorods (thickness: 1-2 nanometers), approximately aligned along one axis. Instead, the presence of PEG (sample WO₃ P₂), as shown in Fig. 2(e) and (f), increases the degree of disorder of the overall structure. This type of nanoparticle generally appears larger, thicker, and more irregular in shape, while the small nanorods described in Fig. 2(c) and (d) seem to disappear. The particle size distribution of WO₃, P₁ is shown in Fig. S1(a),† where it can be seen that most of the distribution is concentrated around 50 and 100 nm, with an average value of 93.86 \pm 57.89 nm. From the size distribution shown in Fig. S1(b),† it is clear that the presence of PEG (WO₃_P_2 sample) leads to an increase in the size of WO₃ nanoaggregates, the average value of which is 120.45 \pm 75.17 nm. The Gaussian particle size distribution for both materials is reported in Fig. S2.†

Fig. 3 shows the distribution of W, C, and O elements by EDX mapping for a WO₃_P_1 nanoparticle. The distribution of elements is quite homogeneous, and based on observations of intensity changes, the ratio of W and O is stable, while the presence of C is mainly due to the influence of the carbon-based membrane of the grid, although the presence of a slight

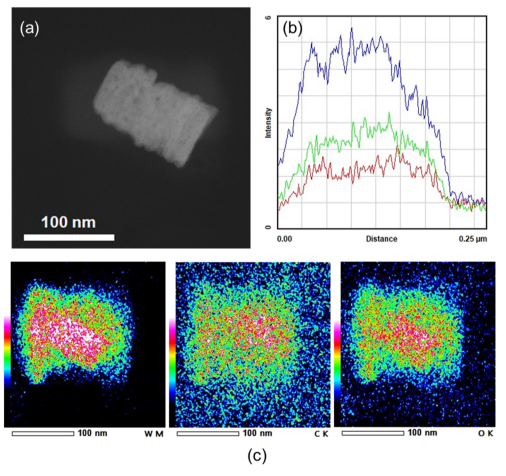


Fig. 3 EDX mapping of W, C, and O elements for a WO_3 – P_1 nanoparticle: (a) STEM image of a WO_3 nanoparticle, (b) X-ray intensity profile of a single WO₃ nanoparticle (W - blue line, O - green line, C - red line), and (c) X-ray distribution map of the nanoparticle reported in (a) for C, O, and W.

increase in its intensity can be associated with some ethanol residues.

The X-ray intensity line profile in Fig. 4(b) indicates that the carbon and oxygen contents of WO₃_P_2 nanoparticles are higher than those of WO₃_P_1. An increased amount of carbon is also shown on the distribution map (Fig. 4(c)), where the presence of this chemical element is in the same area of the nanoparticle but also just outside the single particle, probably due to an excess of PEG.

Fig. S3(a)† shows the electron diffraction pattern of nanostructured WO₃_P_1, which shows a polycrystalline structure in which the different domains of the crystal have the same orientation on the main axis, but also small variations that transform the single diffraction spot into a short line. The SAED interpretation is consistent with the XRD signature corresponding to a monoclinic WO₃ structure oriented along the [100] zone axis, although the resulting *d*-spacings are higher than the expected values for monoclinic WO₃.⁸⁴ Table S1† shows the *d*-spacing of spots with higher intensity. In addition, the plane (002) can also be observed in the HRTEM image (Fig. S3(b)†).

In the case of the WO₃_P_2 material synthesized using PEG, the electron diffraction results at zero tilt are quite chaotic. They

show weak signals indicating the presence of a polycrystal, as well as some intense spots that can be interpreted as (002) family lines (3.99 Å) for both orthorhombic and monoclinic structures, although they deviate slightly from the expected value (3.87 Å) (Fig. S4(a)†), probably due to the presence of PEG in the main crystalline structure. This interpretation is supported by the HRTEM image in Fig. S4(b),† where the calculated d-spacing is 4.02 Å, corresponding to the (002) line. These distances are suitable for both orthorhombic and monoclinic structures.

The XPS analyses indicate that the surface of both oxide films is composed of almost stoichiometric WO_3 (atomic ratio of about 2.8–3.0) with relatively low surface contamination by adventitious carbon and hydroxyl groups (Tables S2 and S3†). Therefore, for the compositional comparison of both samples, the film surface was cleaned by low-energy (1 keV) Ar^+ ion sputtering for a short time (60 s), removing about 0.6 nm of the film. After this cleaning, the surface contamination was removed, and the films were composed of three W species: WO_3 , WO_2 , and metallic W(0). The spectra of W 4f acquired for WO_3 _P_1 and WO_3 _P_2 samples after surface cleaning are presented in Fig. 5(a) and (b). After the application of the peak

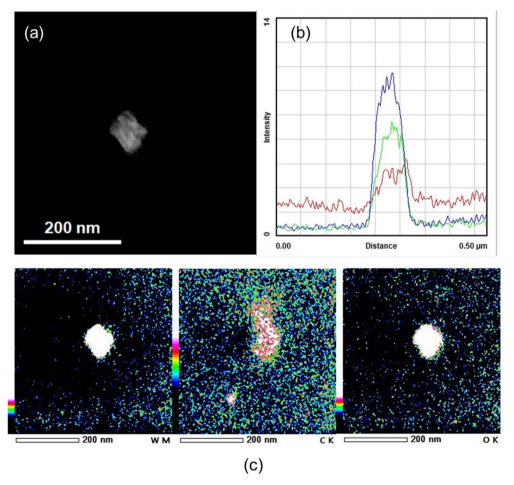


Fig. 4 EDX mapping of W, C, and O elements for a WO_3 _P_2 nanoparticle: (a) STEM image of a WO_3 nanoparticle, (b) X-ray intensity profile of a single WO_3 nanoparticle (W – blue line, O – green line, C – red line), and (c) X-ray distribution map of the nanoparticle reported in (a) for C, O, and W.

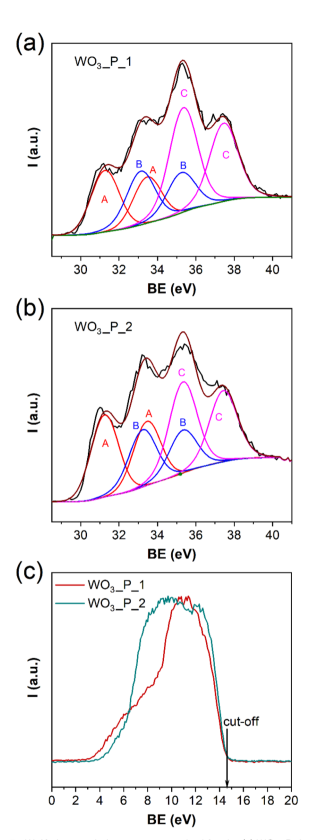


Fig. 5 W 4f photoemission spectra acquired for the (a) WO₃_P_1 and (b) WO₃_P_2 materials after surface cleaning by ion sputtering. Three spin-orbit doublets of W species revealed by peak fitting are marked as A, B, and C for W⁰, W²⁺, and W³⁺, respectively. (c) He I valence band spectra acquired for the WO₃_P_1 and WO₃_P_2 materials.

fitting routine was revealed the presence of three W species with W $4f_{7/2}$ peaks at BE = 31.6 eV (A), 33.6 eV (B), and 35.7 eV (C), attributed to the species of W⁰, W²⁺, and W³⁺, respectively. As follows from this comparison, the content of less coordinated W species (W⁰ and W²⁺) is noticeably higher in the WO₃ P₂ sample, where it is about 0.9 and 0.6 of stoichiometric W³⁺, while in the WO₃ P₁ sample, it was only about 0.6 and 0.5, respectively. The UPS spectra of the valence band acquired for the WO₃_P_1 and WO₃_P_2 using a He I (21.2 eV) source are shown in Fig. 5(c). The obtained results indicate that there is no noticeable change in the valence band maxima (VBM).

Gas sensing results

Fig. 6(a) shows the response of WO₃ P 1 and WO₃ P 2 sensors to 10 ppm of acetone as a function of their operating temperature at 40% relative humidity (RH) in the test chamber. The response values of materials are listed in Table S4.† As can be seen, WO_{3_P_1} is almost not sensitive to acetone (response value: 0.01) at 200 °C. The structure exhibits a good response (response value: 3) at 250 °C, which slightly enhances with an increase in the operating temperature to 300 °C (response value: 3.1). From 300 to 400 °C, the response curve has a larger slope and the response of the material reaches a maximum value (8.7) at 400 °C. A similar sensing behavior is observed for the WO_{3_}P_2 structure when the operating temperature is increased from 200 to 300 °C and then from 300 to 400 °C. However, in the temperature range of 300-400 °C, a significantly better enhancement of the response of WO3_P_2 is observed (the response value increases from 9 to 62) compared to WO₃_P_1. Thus, the response of WO₃_P_2 nanomaterial to acetone is remarkably higher than that of WO₃ P₁.

Fig. 6(b) shows the dynamic response of the two materials to different concentrations of acetone at their optimum operating temperature. The baseline conductance of the sensors stabilizes in the air when the environmental oxygen is adsorbed on their surface. The capture of electrons by chemisorbed oxygen species is accompanied by the formation of an upward band bending at the surface of the semiconductor.11 Then, the conductance values of structures enhance when exposed to acetone. This behavior is typical for n-type semiconductor materials due to the adsorption of reducing gaseous or volatile organic compounds on their surface.85-88 In this case, the acetone molecules interact with adsorbed oxygen species on the surface of the structure, which leads to the trapped charge carriers being returned to the conduction band (eqn (1)).42

$$CH_3COCH_3(gas) + 8O^-(chemisorbed) \rightarrow 3CO_2 + H_2O + 8e^-$$
 (1)

Afterward, the acetone flow is turned off, and the gas chamber is purified with a synthetic air flow. Herein, the recovery of the conductance of each sensor to its initial value in the air confirms the reversible sensing features of the materials. Furthermore, the response curves of both sensors as a function of acetone concentration (Fig. S5(a)†) confirm the superior functionality of WO₃_P_2 compared to WO₃_P_1 even at relatively low operating temperatures. Moreover, the WO₃_P_2

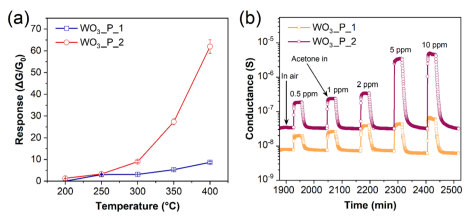


Fig. 6 (a) Response vs. temperature dependence of WO_3 – P_1 and WO_3 – P_2 sensors toward 10 ppm of acetone (RH, 40%). (b) Dynamic response of WO_3 – P_1 and WO_3 – P_2 toward different concentrations of acetone at a RH of 40%.

material exhibits an excellent response to very low concentrations (200 ppb) of acetone (Fig. S5(a) and Table S5†).

As we can see in Fig. 6(b) and S5(c),† the conductance of WO₃ P₂ increases when PEG is used as a surfactant. TEM studies of the sample prepared using PEG as a surfactant (WO₃_P_2) indicate that the nanoparticles are surrounded by a very thin layer of amorphous material. Besides, ATR-IR analyses demonstrate that several bands (e.g., located at 1481, 1343, and 1272 cm⁻¹) can be attributed to residual PEG in the WO₃ matrix. Since PEG is a conductive material, the higher conductance of WO₃_P_2 compared to WO₃_P_1 is consistent with the results obtained from TEM and ATR-IR studies. Furthermore, the second slope of the Arrhenius plot for the WO₃ P₂ sample (0.19 eV, Fig. S5(b)†) correlates with the slope inherent in the pure PEG material (0.2 eV).89 Based on the abovementioned facts, it cannot be excluded that PEG can influence the sorption processes, thereby increasing the sensitivity of the WO₃_P_2 material.

However, what is much more important is that the PEG surfactant displaces the water content of the material. Chacón *et al.* reported that when synthesizing WO₃ with low water concentration in the feed solution, the orthorhombic crystal structure is favored, while at higher water content, the monoclinic phase predominates. This conclusion is fully consistent with our observations. Indeed, the XRD and Raman studies (Fig. 1(a) and (b)) indicate the presence of mixed orthorhombic and monoclinic phases in the WO₃_P_2 sample, while in the WO₃_P_1, only the monoclinic phase is detected.

Thus, suppressing hydrates and forming monoclinic/orthorhombic n-n junction in the WO₃ polymorph leads to a drastic increase in its sensitivity to acetone. At relatively low working temperatures (\leq 250 °C), acetone molecules are adsorbed on the WO₃ surface through the oxygen atoms of the carbonyl group attached to under-coordinated W cations on the surface. Inhibition of hydrate formation by using PEG as a surfactant should increase the density of active adsorption sites on the material surface. Indeed, XPS analysis shows that the concentration of uncoordinated W atoms increased in WO₃-P₋2 compared to WO₃-P₋1 (Tables S2 and S3†). This could

be the reason for the improved sensitivity of the WO₃_P_2 sample in the relatively low operating temperature range.

The formation of the monoclinic/orthorhombic junctions and their effect on the sensitivity of the material require more careful analysis. Our DFT calculations (Fig. 7) show that the monoclinic structure has a slightly larger indirect bandgap than the orthorhombic phase (2.57 eV vs. 2.51 eV, respectively). Although our theoretical simulations carried out with PBE0 functional slightly underestimate the experimental values determined for the monoclinic (2.62–2.7 eV)^{92,93} and orthorhombic (2.58 eV)⁹⁴ phases of the WO₃ polymorph, the predicted trend remains the same.

Based on the Mott–Schottky plot, Kang *et al.* found that the conduction band minimum (CBM) of hexagonal h-WO₃ is located at about 0.1 eV higher than the CBM of the monoclinic γ -WO₃.⁹³ On the other hand, Y. Li *et al.* determined a downshift of 0.13 eV between the CBM of hexagonal h-WO₃ and orthorhombic β -WO₃ phases.⁹⁵ Our UPS investigations show that the VBM of nanomaterials WO₃_P_2 and WO₃_P_1 are located approximately at the same level (Fig. 5(c)).

Given our theoretical and experimental results in this work, we can deduce that the CBM of orthorhombic β -WO $_3$ is slightly lower (0.03–0.06 eV) than the CBM of monoclinic γ -WO $_3$ structure. Therefore, the electron transfer from the γ -WO $_3$ to the β -WO $_3$ counterpart leads to a further increase in the surface acidity of WO $_3$ -P $_2$, where the monoclinic phase still dominates. The increased acidity of the WO $_3$ -P $_2$ sample significantly facilitates the adsorption of acetone, which has weakly basic properties.

TEM characterization of the samples (Fig. 4(b)) demonstrates that the WO₃_P_2 sensor has a much higher oxygen content than the WO₃_P_1. According to eqn (1), the presence of oxygen is a necessary condition for the catalytic oxidation of acetone at elevated temperatures. Eventually, the increased oxygen content in the disordered WO₃_P_2 sensor can explain its significant sensitivity to acetone in the high-temperature operating range.

We also investigated the effect of RH on the electrical and sensing properties of synthesized materials by gradually increasing its concentration in the test chamber to 95%

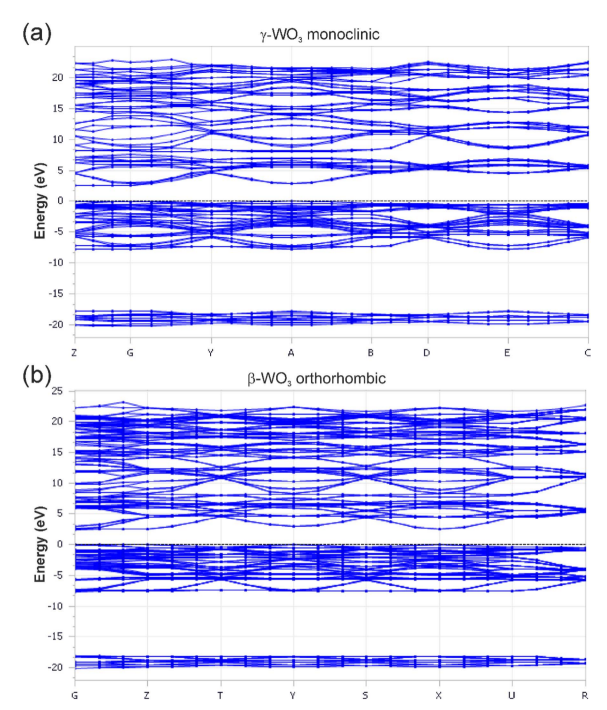


Fig. 7 Calculations of the electronic band structure of the monoclinic γ -WO $_3$ (a) and orthorhombic phases of β -WO $_3$ (b). Zero energy corresponds to the top of the valence band.

(Fig. S5(c)† and 8(a)). The obtained results indicate that there are no significant changes in the conductance and response of the WO₃_P_1 and WO₃_P_2 sensors due to the increase in the RH level in the chamber, even at a low concentration of acetone (200 ppb). Fig. 8(b) shows the response of WO₃_P_1 and WO₃_P_2 toward other gaseous compounds. Here, the high selectivity of both materials is evident. In particular, WO₃_P_2 exhibits excellent sensing performance to acetone. Furthermore, we measured the dynamic response of the polymorphic WO3_P_2 sensing material to different concentrations of

acetone at 400 °C over two consecutive exposure cycles. The structure conductance returns to its baseline value in air after the injection of each concentration of acetone into the test chamber and its subsequent purification during both test cycles (Fig. S6†), demonstrating the reliability of the obtained results. We also compared its acetone sensing parameters with those of other semiconductor gas sensors based on pure and composite structures (Table S6†). As can be seen, the nanostructuration of WO₃ and appropriate modification of its crystalline structure have a crucial effect on its functionality.

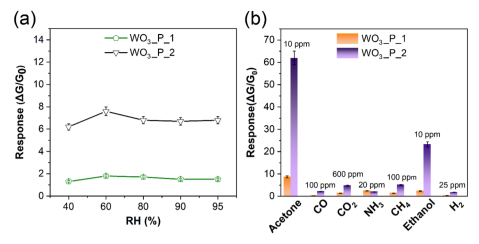


Fig. 8 (a) Response values of $WO_3_P_1$ and $WO_3_P_2$ sensors to 200 ppm of acetone at different concentrations of RH (40–95%) in the test chamber. The operating temperature of the sensors is 400 °C. (b) Sensing response of $WO_3_P_1$ and $WO_3_P_2$ toward different gaseous and volatile organic compounds at their optimum operating temperature (400 °C).

4. Conclusions

In summary, we synthesized and studied the monoclinic/orthorhombic WO_3 polymorph nanomaterials for their application in chemical gas sensing devices. The precipitation method is an effective approach to obtain nanostructured WO_3 , where benzyl alcohol is crucial for controlling the reaction rate of the WCl_6 precursor and the precipitation of material. Thus, it can be used to prepare WO_3 nanoparticles. The addition of PEG in the synthesis process increases structural disorder and suppresses hydrate formation in the resulting WO_3 nanomaterial. Furthermore, PEG surfactant affects the oxygen content in WO_3 and promotes partial crystallization of the material into the orthorhombic structure.

Without the addition of PEG, the water content of WO₃ annealed at 450 °C is higher, and the material tends to crystallize into the monoclinic phase. In contrast, the material synthesized using PEG and annealed at 450 °C still predominantly exhibits the monoclinic phase, but a contribution from orthorhombic phases can also be observed. Our experimental and theoretical results demonstrate that hydrate reduction and the formation of monoclinic/orthorhombic n-n junctions in the WO₃ polymorph are critical factors in enhancing its interaction with acetone. The reduction of hydrates increases the density of active adsorption sites on the surface of the material, which is favorable for the adsorption of acetone. Moreover, the electronic band structure in the monoclinic/orthorhombic polymorph results in electron transfer from the γ -WO₃ to the β -WO₃, increasing the acidity of WO3 and enhancing its interaction with acetone. Furthermore, the higher oxygen content of the material prepared using PEG facilitates the catalytic oxidation of acetone at elevated temperatures and thus improves its sensing response at relatively high operating temperatures.

The high sensing response and selectivity of the WO₃ sample prepared with the addition of PEG, as well as its stable functionality at different humidity levels, indicate that the polymorphism in the WO₃ nanomaterial, considering the appropriate modification of crystalline phases in the structure,

may lead to a wider use of WO₃ in chemical gas sensing and photocatalytic processes.

Data availability

All data supporting the findings of this study are available within the article and its ESI. \dagger

Author contributions

Hakimeh Pakdel: conceptualization, formal analysis, investigation, methodology, validation, visualization, writing - original draft; Vardan Galstyan: conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, supervision, validation, visualization, writing - original draft, writing - review & editing; Viacheslav Golovanov: data curation, formal analysis, funding acquisition, investigation, methodology, software, validation, visualization, writing – original draft, writing – review & editing; Annalisa D'Arco: formal analysis, investigation, methodology, validation, visualization, writing - original draft, writing review & editing; Tiziana Mancini: formal analysis, investigation, methodology, validation, visualization, writing - original draft; Francesco Mura: formal analysis, investigation, methodology, validation, writing - original draft; Marco Rossi: funding acquisition, methodology, resources, writing - review & editing; Stefano Lupi: funding acquisition, methodology, resources, writing - review & editing; Alessio Mezzi: formal analysis, investigation, methodology, validation, visualization, writing original draft; Saulius Kaciulis: funding acquisition, investigation, methodology, resources, writing - original draft, writing review & editing; Elisabetta Comini: funding acquisition, methodology, resources, project administration, writing review & editing.

Conflicts of interest

There are no conflicts to declare.

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