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Rethinking the molecular structures of $W^{VI}O_x$ sites dispersed on titania: distinct mono-oxo configurations at 430 °C and temperature-dependent transformations†

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The structural properties of the $(WO_x)_n$ phase dispersed on TiO₂ (P25, anatase) at surface densities of 0.5-4.5 W nm⁻² (i.e. up to approximately a monolayer) were explored by using in situ Raman and FTIR spectroscopy, in situ Raman/18O exchange and Raman spectroscopy in static equilibrium at temperatures of 175-430 °C. Deciphering the temperature and coverage dependence of the spectral features under oxidative dehydration conditions showed that (i) the $(WO_x)_n$ dispersed phase is heterogeneous at 430 °C consisting of two distinct mono-oxo species: Species-I with C_{3v} -like O=W(-O-) $_3$ configuration (W=O mode at $1009-1014 \text{ cm}^{-1}$) and Species-II with C_{4v} -like $O=W(-O-)_4$ configuration (W=O mode at 1003–1009 cm⁻¹); (ii) the $O=W(-O)_3$ site is formed with first order of priority and its formation ceases after the complete consumption of the most basic hydroxyls that are titrated first, hence is abundant at low coverage (<1.5 W nm⁻²); (iii) the O= $W(-O-)_4$ site prevails over the O= $W(-O-)_3$ site at medium to high coverage (≥2 W nm⁻²) and partially occurs in associated (polymerized) coverages above 2 W nm⁻²; (iv) lowering the temperature in the 430 \rightarrow 250 \rightarrow 175 °C sequence does not affect the structural and vibrational properties of $O=W(-O-)_3$ but leads to the gradual transformation of the $O=W(-O-)_4$ site to a di-oxo $(O=)_2W(-O-)_3$ site (with a symmetric stretching mode at ~985 cm⁻¹) and the partial association of adjacent O=W(-O-)4 units. All temperature-dependent structural/configurational transformations are fully reversible in the 430-175 °C range and are interpreted at the molecular level by a mechanism involving water molecules retained at the surface that act in a reversible temperature-dependent mediative manner resulting in hydroxylation (upon cooling, e.g. to 250 °C) and dehydroxylation (upon heating, e.g. to 430 °C). The Raman spectra obtained for the hydroxyl region confirm the successive hydroxylation/ dehydroxylation steps during temperature cycles (e.g. $430 \rightarrow 250 \rightarrow 430$ °C). One can tune the speciation of the dispersed $(WO_x)_n$ phase under dehydrated conditions by appropriate control of temperature and coverage.

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1. Introduction

A wide range of reactions in industrial and environmental catalysis benefit from the formidable mix of strong acid and modest redox properties of titania-supported tungsta catalysts.

Applications worth mentioning are—although not restricted to —photocatalytic processes, *e.g.* the photooxidation of methane, ¹ the photodecompositions of 1,4 dichlorobenzene ² and 4-nitrophenol, ³ olefin processing reactions ⁴ and environmental processes such as the NH₃/SCR of NO_x, ⁵⁻⁷ the abatement of chlorinated VOCs and CFC-12^{8,9} and sorbitol transformation into biofuels. ¹⁰ A very recent review summarises the applications of tungsten oxide–based catalysts in environmental catalysis. ¹¹

The behavior of tungsta/titania as a catalyst is directly connected to the configuration and molecular structure of the oxo- W^{VI} phase, $(WO_x)_n$, dispersed on the titania support surface. The literature on catalysis science abounds with studies of the molecular structure of dispersed acidic oxometallic $(MO_x)_n$ sites (M = transition metal, e.g. V, Mo, W, and Re)

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[†]Electronic supplementary information (ESI) available: Fig. S1–S4 showing loading dependence of *in situ* Raman, *in situ* FTIR and *static equilibrium* Raman spectra obtained for $WO_x/TiO_2(P25)$ at 250 °C (Fig. S1) and $WO_x/TiO_2(a)$ at 430, 250 and 175 °C (Fig. S2–S4). See DOI: https://doi.org/10.1039/d2dt00595f

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and many pertinent reviews are available. 12-24 Despite the extent of the literature, the reports on key aspects such as the termination configuration (i.e. discriminating between the mono-oxo, di-oxo and/or trioxo termination configurations) of dehydrated MOx sites and the eventual heterogeneity of the dispersed oxometallic phase (i.e. the occurrence of more than one prevailing species within the deposited phase) are not in consensus. Diverging views and statements based on fragmented evidence and/or arbitrarily oversimplified generalizations have created a real puzzle. For example, an old, "established" view has been claiming that thermodynamic constraints drive the formation of the prevailing configurations and structures of transition metal oxide species dispersed on oxide supports and that the configurations and structures obtained by the deposited oxometallic sites are reported as independent of the catalyst preparation routes. 20,21,24,25 However, the preparation method affects the molecular structure of the deposited e.g. $(WO_x)_n$ phase, particularly when advanced synthesis routes based on molecular approaches are followed, thereby in some cases enabling catalyst design. 15,26,27 In addition, it has been reported, without sufficient evidence, that the prevalent configurations obtained after calcination by the deposited species do not revert/change with temperature under dehydrating conditions, 16-25 despite the sound evidence to the contrary documented in scarce reports.²⁸ Recently, however, we have provided strong evidence that controverts the soundness of the above statement and showed that a temperature-dependent transformation takes place under dehydrated conditions between the sites of distinct configurations in rhenia supported on TiO2 and low-loaded tungsta and molybdena supported on TiO2.29-31

Literature reports pertaining to the molecular structure of the (WO_x)_n/TiO₂ system are far from being in consensus. An early in situ Raman work, with the spectra recorded after cooling to 80 °C, on 1-10 wt% WO₃ on TiO₂(P25) samples made by incipient wetness impregnation³² reports the existence of one single band at 1010 cm⁻¹ in the W=O stretching region at all weight loadings, which is ascribed to the existence of one single species. A second in situ Raman report³³ based on the same samples and the same spectra as ref. 32 enriches the interpretation supporting the mono-oxo configuration, based on the correspondence of the Raman and IR W=O stretching wavenumbers. The claim for the existence of one single band appears to be abandoned later when a poorly resolved band in the W=O stretching region of in situ Raman spectra (recorded however at 25 °C, after cooling) is reported at 1007 cm⁻¹ for 2.7 W nm⁻² and 1011 cm⁻¹ for 4 W nm⁻² WO₃/ TiO₂(P25) samples made by incipient wetness impregnation.³⁴ An in situ structural analysis by XANES/EXAFS showed that under dehydrated conditions, wolfram is in the hexavalent state in the form of WVIO5 and WVIO4 units dispersed on TiO₂(P25) in mono-oxo W=O configuration with the W-O-W interlinking bridges and WO5 units prevailing over their WO4 counterparts with increasing loading.³⁵ The use of a one-pot co-precipitation synthesis route led, based on X-ray absorption and vibrational (infrared and in situ Raman combined with

¹⁸O/¹⁶O exchange) spectroscopy, to the first report on the heterogeneity of dispersed $(WO_x)_n$ on TiO_2 with two distinctly observed tungstate species (bands at 978 and 1016 cm⁻¹) forming an overlayer on TiO2 36 even at surface densities exceeding the reported³⁴ monolayer coverage of \sim 4.5 W nm⁻². Onfroy et al. and Lebarbier et al. reported the amorphous nature of the deposited oxo-WVI phase on titania up to 4.4 W nm⁻² and a correlation between the abundance of Brønsted acid sites, which reportedly prevail above 1.3 W nm⁻², and the catalytic activity towards isopropanol dehydration and *n*-hexane isomerization.^{37,38} An elegant spectroscopic study combining IR and Raman work examined the effect of surface density in the range of 0.6-3.4 W nm⁻² for WO_x/TiO₂(P25) catalysts and proposed the occurrence of three distinct mono-oxo species (bands at 1004, 1012 and 1017 cm⁻¹), albeit the spectroscopic measurements were undertaken after cooling, under controlled atmosphere. 39 The formation of two distinct surface mono-oxo WO5 species has also been confirmed recently by adopting a co-precipitation method in the catalyst synthesis route,40 although providing an alternative interpretation viewing angle, to be discussed below.

Hence, despite the extensive extant literature, the molecular structures, the site configurations, coordination and aspects related to the occurrence of multiple sites with distinct configurations are not fully resolved and understood. A plethora of published spectroscopic work suffers from one or more deficiencies including e.g. relying on only one or two samples with loading below monolayer coverage, undertaking "in situ" spectroscopic measurements at one single temperature or after cooling the samples (in which case the state of sample dehydration is questioned), performing fast spectroscopic measurements without being concerned with steady-state attainment etc.

The scope of the present work is to deploy the tools of in situ vibrational (Raman and FTIR) spectroscopy combined with 18O/16O isotope exchange for studying the molecular structures and termination configurations of WOx sites dispersed on two variations of titania (P25 and anatase) under dehydrated conditions in the temperature range of 175-470 °C and in the entire submonolayer surface density range of 0.5-4.5 W nm⁻². The study is complemented by neoteric³⁰ Raman measurements in sealed quartz cells under static equilibrium. The results point towards rethinking the molecular structure of the dispersed $W^{VI}O_x$ phase with respect to its heterogeneity and temperature dependence of its molecular structure.

2. Experimental section

Synthesis and textural analyses of WO_x/TiO₂ materials

Two variations of titanium dioxide were used as supports, Degussa P25 (surface area of 49 m² g⁻¹; consisting of 80 wt% anatase and 20% rutile with a surface composition of 90% anatase and 10% rutile) and anatase (Alfa Aesar, surface area: 127 m² g⁻¹). The tungsta/titania samples were synthesised by

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Table 1 WO_x/TiO₂ catalysts' characteristics (surface density, wt% W loading, BET surface area, initial concentration of precursor solution, and support type). Calcination: 480 °C, 4 h

$n_{\rm s}$ (W nm ⁻²)	Loading (wt% W)	$S_{\rm BET}$ (m ² g ⁻¹)	$C_{\mathrm{W(VI)}}\left(\mathbf{M}\right)$	Sealed cells #	Support
0.51	0.81	52	9×10^{-4}		P25
1.2	1.67	47	1.9×10^{-3}	3, 8	P25
2.1	3.32	51	3.6×10^{-3}	11	P25
2.5	3.90	52	4.3×10^{-3}		P25
3.4	5.35	52	5.9×10^{-3}	4,7	P25
4.2	6.33	50	7.0×10^{-3}	,	P25
0.54	1.56	94	1.7×10^{-3}		Anatase
1.1	3.17	94	3.4×10^{-3}	5	Anatase
2.2	6.34	93	6.9×10^{-3}	12	Anatase
2.6	7.14	91	7.8×10^{-3}		Anatase
3.6	9.52	86	1.04×10^{-2}	6	Anatase
4.6	11.90	85	1.29×10^{-2}		Anatase
	0.51 1.2 2.1 2.5 3.4 4.2 0.54 1.1 2.2 2.6 3.6	0.51 0.81 1.2 1.67 2.1 3.32 2.5 3.90 3.4 5.35 4.2 6.33 0.54 1.56 1.1 3.17 2.2 6.34 2.6 7.14 3.6 9.52	0.51 0.81 52 1.2 1.67 47 2.1 3.32 51 2.5 3.90 52 3.4 5.35 52 4.2 6.33 50 0.54 1.56 94 1.1 3.17 94 2.2 6.34 93 2.6 7.14 91 3.6 9.52 86	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

wet impregnation using (NH₄)₁₀W₁₂O₄₁·5H₂O (Alfa Aesar, 99.999% metal basis) as the precursor and previously calcined (480 °C, 4 h) supports. The concentration of the aqueous precursor solutions was adjusted to correspond to the desired W loading (see Table 1). During the stages of precursor dissolution and support impregnation (both procedures lasted for 1 h under agitation), the pH of the solutions and suspensions was continuously measured and controlled to 7.5. In the applied concentration region of $C_{W(VI)} = 0.9 \times 10^{-3} - 1.3 \times 10^{-2}$ M and for pH = 7.5, the oxo-W^{VI} species in the electrolyte solutions as well as in the titania/electrolyte interface are exclusively in the mononuclear monomeric tetrahedral form. 41,42 The support/precursor solution suspensions were then heated under mixing at 45 °C for 1 h and subjected to rotary evaporation under reduced pressure at 45 °C for removing the solvent. The final samples were obtained after overnight drying at 120 °C and calcination at 480 °C for 4 h in a muffle furnace under static air.

The BET specific surface area (S_{BET}) of the calcined samples was determined by nitrogen adsorption-desorption, as earlier described. 42 The samples' coverage, n_s , expressed in the form of W surface density (W nm⁻²) was determined based on the wt% W loading and the S_{BET} . The pertinent results are compiled in Table 1 together with other characteristics. Each calcined catalyst sample is denoted as $xWTiO_2(P25)$ or $xWTiO_2(a)$ where x is the W surface density (W nm⁻²). The following formula is used for calculating the W surface density (W nm⁻²):

$$n_{\rm s} = \frac{\left[\frac{g_{\rm W}}{g_{\rm cat}}/A_{\rm W}\right] \times 6.023 \times 10^{23}}{\left[S_{\rm BET}/\left({\rm m^2\,g_{\rm cat}}^{-1} \times 10^{18}\,{\rm nm^2\,m^{-2}}\right)\right]}$$

where A_{W} is the atomic mass of W in amu.

2.2 Raman spectroscopy and protocols of measurements

2.2.1 In situ Raman spectra. A homemade Raman optical cell was used for recording the in situ Raman spectra of the WO_x/titania catalyst samples under oxidative dehydrating conditions in the temperature range of 175-470 °C. A detailed description of the in situ cell can be found elsewhere. 13,43

90–150 mg of gently powdered sample was pressed in the form of a disc pellet that was mounted on the optical cell sample holder. The cyan 491.5 nm line of a Cobolt Calypso DPSS laser was used as the irradiation source. The incoming laser beam had a ca. 10 mW power on the sample and was slightly defocused to further reduce sample irradiance. The scattered radiation was collected at right-angles in a horizontal scattering plane and was analyzed using a 0.85 m 1403 Spex double monochromator and detected with an RCA photomultiplier tube (cooled at −20 °C) that was coupled with the Labspec software.

Each sample was first heated under flowing (30 cm³ min⁻¹) 20% O₂/He gas mixture at 430 °C for 1 h and the in situ Raman spectra were recorded under dehydrated conditions at 430 °C. The gases used were He (99.999%) and O2 (99.995%) from L'Air Liquide and the incoming feed gas stream was passed through molecular sieve traps to eliminate any traces of moisture. The temperature was then lowered to 250 °C and 175 °C and in situ Raman spectra were recorded after subjecting the sample for 1 h and 45 min at each temperature under flowing 20% O2/He gas. Subsequently, the sample was reheated to 430 °C and the reinstatement of the initial structural condition could be verified by reproducing the in situ Raman spectrum at 430 °C. Alternative sequences of temperatures have been tested (always starting at 430 °C, i.e. 430 °C \rightarrow 175 °C \rightarrow 250 °C, 430 °C \rightarrow 175 °C \rightarrow 120 °C \rightarrow 250 °C etc.) and the in situ Raman spectra could always be reproduced. To achieve a satisfactory signal-to-noise ratio, a slow scanning protocol of 1.2 s of photon counting per point in 0.25 cm⁻¹ increments was applied, thereby giving rise to a 1.5 h duration per recording. To account for the "path length" effect resulting from the variations in sample coverage (0.5-4.5 W nm⁻²), the Raman spectra were normalized, as described earlier. 30,44

2.2.2 Raman/isotope exchange measurements. In situ Raman - 18O/16O isotope exchange measurements were performed by performing successive reduction/oxidation cycles. First, the sample was reduced under flowing 5% H₂/He at 470 °C for 20 min and subsequently reoxidized under flowing 2% ¹⁸O₂/He for 10 min and the *in situ* Raman spectrum was recorded. The gases used were H₂ (Linde, 99.999%) and 2% **Paper Dalton Transactions**

¹⁸O₂/He (Linde). Typically, up to 21 reduction/oxidation isotope exchange cycles were run for each sample studied. The temperature of 470 °C was chosen to achieve an adequate extent of reduction and a commensurate 18O/16O exchange without exposing the catalysts above the calcination temperature of 480 °C. The TPR measurements showed that the onset of the WO_x/TiO₂ catalyst reduction was at (or above) 550 °C. The protocol of the isotope exchange experiments was established by simulating the reduction/oxidation using a 2% ¹⁶O₂/ He reoxidant gas, as described previously. 45,46

2.2.3 Raman spectra in sealed quartz cells under static equilibrium dehydrated conditions. A special set of experiments were designed for studying the WOx/titania catalysts under static equilibrium dehydrated conditions. The concept of the cell design has recently been described³⁰ and is depicted in Fig. 1. The cell is made of quartz tubing and it has a 20 mm o.d. ~3 cm long main compartment that possesses a 6 mm o.d. ~2 cm long appendix in the bottom (for containing the sample in the powder form) and a 6 mm o.d. stem at the

In a typical experiment, 100 mg of each studied WO_x/ TiO₂(P25 or anatase) catalyst was filled into the cell that was then attached to the vacuum line, which is suitable also for gas additions. The cell was evacuated $(p = \sim 1.3 \times 10^{-4} \text{ atm})$ and a cylindrical furnace was subsequently connected around the cell. The sample containing cell was then heated at 200 °C and subjected to evacuation for 1 h, while maintaining the vacuum line trap immersed in liquid nitrogen (77 K). The main valve connecting the vacuum pump was then closed, the furnace surrounding the cell was removed, oxygen gas (99.999%) was injected into the evacuated line at a pressure of 0.19 atm (i.e. the vapor pressure of oxygen at 77 K) and finally the cell stem was sealed off by means of a propane/oxygen torch. Several such "static" cells were made and WO_x/TiO₂ catalysts with representative coverages were sampled as listed in

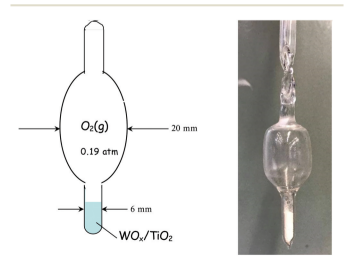


Fig. 1 Diagram and actual picture of a quartz cell used for measuring Raman spectra under static equilibrium. The cell is sealed under $O_2(g)$, $p_{O_2} = 0.19$ atm.

Table 1. The purpose of designing the static equilibrium experiments was twofold: first, we wanted to demonstrate that the temperature-dependent structural evolution reported is not due to humidity incoming with the gas feed, and second, we wanted to check the effect of limiting the amount of water retained directly by the surface and examine if this could inhibit the temperature-dependent changes observed.

A description of the optical furnace used for recording Raman spectra under static equilibrium has been given earlier. 47,48 The Raman spectra under static equilibrium were obtained using the linearly polarized 532.0 nm green line of a Spectra-Physics Excelsior DPSS laser operated at a power level of ~20 mW on the sample (defocused in order to reduce sample irradiance). The scattered light was passed through a notch filter to reject Rayleigh scattering. The Raman spectra were recorded in a 90° geometry using a collecting lens system (90 and 150 mm FL) coupled to a monochromator (IHR-320 JY, ISA-Horiba group) and equipped with a CCD detector thermoelectrically cooled to -70 °C. The resolution of the instrument was set at 2 cm⁻¹. Each cell was first heated to 430 °C and the Raman spectra under static equilibrium were recorded at a sequence of decreasing temperatures. The Raman setup used for the measurements under static equilibrium has been described earlier.49

2.3 In situ FTIR spectra

In situ FTIR spectra at temperatures ranging from 175-430 °C under flowing (30 cm³ min⁻¹) 2% O₂/He were recorded in order to complement the in situ Raman spectra. The FTIR spectrometer used was a Nicolet 6700 equipped with a DRIFT cell from Spectra Tech, an MCTB detector and a KBr beam splitter. The sample treatment protocol was identical to the one followed for recording the in situ Raman spectra. Before recording each spectrum, each studied sample was subjected to the applied temperature and gas atmosphere conditions for 1 h and 45 min. In situ FTIR spectra were also recorded under flowing 20%O₂/He and were similar (though of lower signal-tonoise ratio) to the ones reported. Due to the strong absorption in the wavenumber range of fundamental W=O stretching, the corresponding overtone region of the in situ FTIR spectra is explored.

3. Results

Vibrational (Raman and FTIR) spectra of WOx/TiO2 (P25 and anatase) catalysts under oxidative dehydration conditions -Temperature dependence and effect of coverage

Fig. 2-7 (panels (A) and (B)) show the temperature dependence of the in situ Raman and in situ FTIR spectra obtained for the WO_x/TiO₂(P25) and WO_x/TiO₂(anatase) catalysts under flowing O2/He gas in the temperature range of 430-175 °C at representative constant coverages of 1.2 W nm⁻² on P25 (Fig. 2), 1.1 W nm⁻² on anatase (Fig. 3), 2.1 W nm⁻² on P25 (Fig. 4), 2.2 W nm⁻² on anatase (Fig. 5), 3.4 W nm⁻² on P25 (Fig. 6) and 3.6 W nm⁻² on anatase (Fig. 7), i.e. at low, intermediate and high

1100

1050

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(A) (B) (C)WO/TiO₂(P25) WO/TiO(P25) flowing 20% O₂/He WO /TiO (P25) flowing 1.2 W/nm² 1.2 W/nm² 2%O₂/He 1.2 W/nm² P(O₂)= 0.19 atm (Static, Cell #8) 1010 T/°C T/°C T/°C 430 Normalized Raman Intensity Normalized Raman Intensity (II) Reflectance 250 1009 (I) 1997(II) 175 2011 1990 (II) 800

Fig. 2 WO_x/TiO₂(P25) with a surface density of 1.2 W nm⁻²: (A) Sequential in situ Raman spectra obtained under flowing 20%O₂/He at temperatures as indicated by each spectrum. Recording parameters: laser wavelength, $\lambda_0 = 491.3$ nm; laser power, w = 10 mW; time constant, $\tau = 1.2$ s; and spectral slit width, ssw = 8 cm^{-1} . (B) Sequential in situ FTIR spectra obtained under flowing $2\%O_2$ /He at temperatures as indicated by each spectrum. (C) Sequential Raman spectra at static equilibrium under $p_{O2} = 0.19$ atm at temperatures as indicated by each spectrum; spectral parameters: $\lambda_0 = 0.19$ 532.0 nm; laser power, w = 20 mW; resolution, 2 cm^{-1} .

2050

wavenumber, cm

2000

1950

800 2200

2150

2100

900

Raman Shift, cm

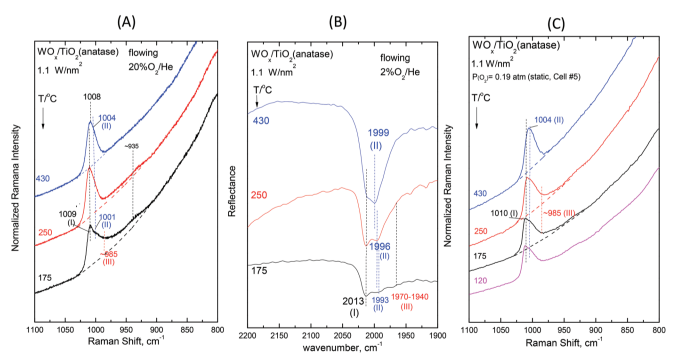


Fig. 3 WO_x/TiO₂(anatase) with a surface density of 1.1 W nm⁻²: (A)–(C): see Fig. 2 caption.

coverage for both studied systems. The left panels (i.e. Fig. 2 (A)-7(A)) pertain to in situ Raman spectra whilst the middle panels (i.e. Fig. 2(B)-7(B)) show the counterpart in situ FTIR spectra. Notably, the spectra are recorded at a sequence of decreasing temperatures. Before recording each in situ spectrum at 250 and/or 175 °C, each sample was treated at 430 °C

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1000

950

Raman Shift, cm⁻¹

900

850

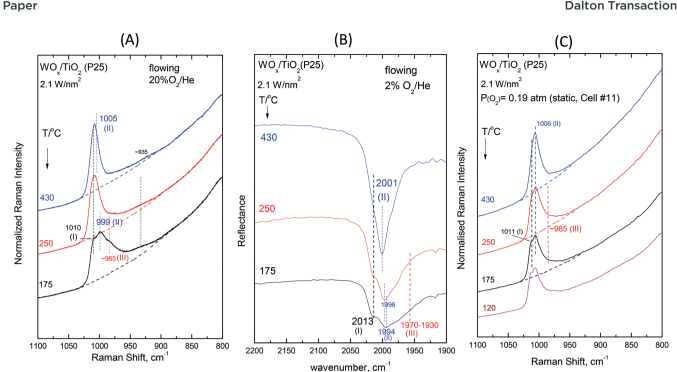


Fig. 4 WO_x/TiO₂(P25) with a surface density of 2.1 W nm⁻²: (A)–(C): see Fig. 2 caption.

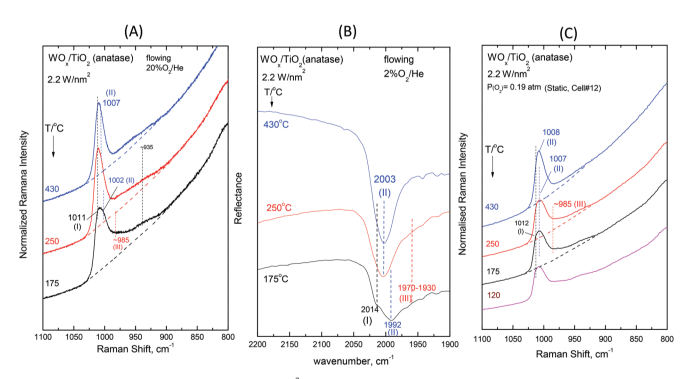


Fig. 5 WO_x/TiO_2 (anatase) with a surface density of 2.2 W nm⁻²: (A)–(C): see Fig. 2 caption.

for 0.5 h, the in situ spectrum was recorded at 430 °C and then temperature was lowered to 250 and/or 175 °C and allowed to attain the steady state. The observed changes are fully reversible, i.e. upon reheating each sample at 430 °C under flowing O₂/He gas, the respective spectral features are fully reinstated.

Fig. 2(C)-7(C) (i.e. the right panels) show the respective Raman spectra obtained under static equilibrium conditions in sealed quartz cells.

The effect of temperature shows trends that are mutatis mutandis common to both types of carriers (i.e. $WO_x/P25$ and

850

900

800

1100

1050

1000

950

Raman Shift, cm

900

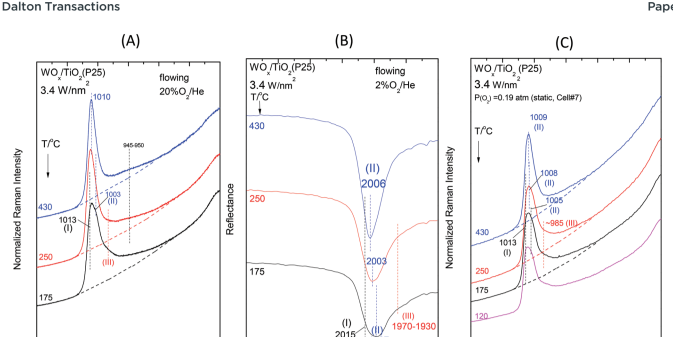


Fig. 6 WO_x/TiO₂(P25) with a surface density of 3.4 W nm⁻²: (A)–(C): see Fig. 2 caption.

800

2200

2150

2100

2050

wavenumber, cm⁻¹

850

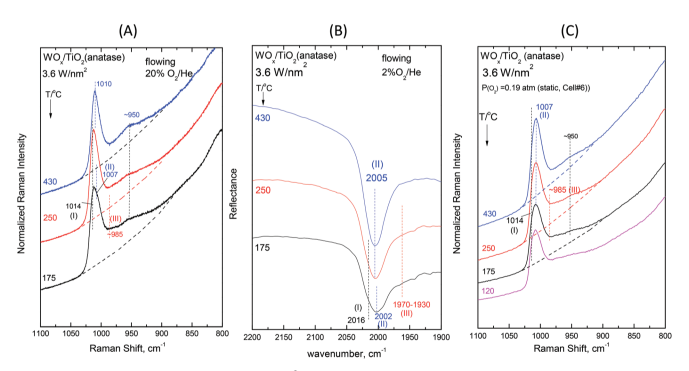


Fig. 7 WO_x/TiO₂(anatase) with a surface density of 3.6 W nm⁻²: (A)–(C): see Fig. 2 caption.

WO_x/anatase samples) and all coverages in the range of 1.1-3.6 W nm⁻² for the in situ Raman and in situ FTIR spectra displayed in Fig. 2-7 (panels (A) and (B)). As far as the extent of the observed effects is concerned, the latter varies slightly in coverage and type of carrier, as described below. A comparison between the in situ Raman spectra (Fig. 2(A)-7(A)) with the corresponding static Raman spectra (Fig. 2(C)-7(C)) shows (as further described and discussed below) that the structural changes taking place upon cooling are indeed partly inhibited after heating the samples in vacuo at 200 °C for 1 h and sealing their container cells under oxygen (p_{O_2} = 0.19 atm).

1100

1900

1050

1000

950

Raman Shift, cm⁻⁷

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3.1.1 Low-loaded WO_x/TiO₂ samples (1.1 and 1.2 W nm⁻²). For the low-loaded samples (i.e. 1.2 W nm⁻² on P25 and 1.1 W nm⁻² on anatase, Fig. 2 and 3), the in situ Raman spectra under oxidative dehydration conditions (left panels, i.e. Fig. 2 (A) and 3(A)) show at 430 °C a band feature (presumably hosting two components) above 1000 cm⁻¹, of which the tip becomes sharper on cooling to 250 and 175 °C, whilst concomitantly the low wavenumber side of the band is gradually weakened and becomes resolved as a well-defined shoulder that undergoes a gradual redshift to ~997 cm⁻¹ for $1.2 \text{WTiO}_2(\text{P25})$ and $\sim 1001 \text{ cm}^{-1}$ for $1.1 \text{WTiO}_2(a)$. The counterpart in situ FTIR spectra (middle panels, Fig. 2(B) and 3(B)) not only complement but also further enrich the observations. The double character of the band at 430 °C is ascertained because the distance of its component bands is doubled in the overtone region, thereby allowing the bands to become separated. Very clearly, upon cooling to 250 and 175 °C (Fig. 2(B) and 3 (B)), the low wavenumber component (1997 cm⁻¹ for 1.2WTiO₂(P25) and 1999 cm⁻¹ for 1.1WTiO₂(a)) undergoes a relative weakening, a broadening and a redshift, whilst the high wavenumber component (2011 and 2013 cm⁻¹ for 1.2WTiO₂(P25) and 1.1WTiO₂(a), respectively) becomes prevalent, without intensity change and remains at a fixed wavenumber position. The behavior is typical of temperature dependence between prevailing configurations, each one represented by one band near and above 1000 cm⁻¹. We denote the configuration represented by the high wavenumber band hereinafter band (I)—as Species-I, which is present in fixed amounts at all temperatures and is predominant at lower temperatures. The configuration represented by the lower wavenumber band-hereinafter band (II)-is denoted as Species-II, which prevails at high temperatures and gradually diminishes with decreasing temperature. The characteristic Species-II Raman band occurs at ~1003 and ~1004 cm⁻¹ at 430 °C and its IR overtone is observed at 1997 and 1999 cm⁻¹ for the low-loaded samples (Fig. 2 and 3). The characteristic Species-I Raman band is seen at 1008/1009 cm⁻¹ with IR overtones at 2011 and 2013 cm⁻¹ (Fig. 2 and 3). In each case the first overtones are observed at wavenumbers that are lower than the values obtained by doubling the respective Raman fundamentals, which is in agreement with the anharmonic oscillator model in the diatomic approximation. A final observation pertains to the possible existence of a peak mass at ~985 cm⁻¹ emerging at 175 °C in the in situ Raman spectra (Fig. 2(A) and 3(A)) with a corresponding overtone feature at

Additional subtle changes taking place upon cooling in the *in situ* Raman spectra (Fig. 2(A)–3(A)) pertain to the appearance and gradual growth of a weak and broad peak mass at \sim 935 cm⁻¹. This event proceeds parallel to the weakening and gradual shift of band (II).

 $1940-1970 \text{ cm}^{-1}$ in the *in situ* FTIR spectra (Fig. 2(B) and 3(B)),

probably representing a third species (Species-III, formed at

the expense of Species-II), the existence of which is further evi-

We now turn our attention to a comparison between the in situ Raman spectra (Fig. 2(A)-3(A)) and the corresponding

static equilibrium Raman spectra shown in Fig. 2(C)-3(C). A diversified behavior is observed when studying the spectral changes taking place upon cooling the sealed cells #8 and #5 containing the low-loaded samples (see Table 1). Upon cooling, band (II) initially weakens without undergoing any redshift and eventually its intensity remains stable. Concomitantly, i.e. with decreasing temperatures, band (I) becomes prevalent without gaining intensity. Moreover, in agreement with the in situ conditions (panels (A)), the emergence of Species-III is (albeit to a limited extent) evidenced in the low-temperature region under static conditions (panels(C)). However, the observed changes do not show further progress below 250 °C. The spectra obtained at 120 °C are also shown to strengthen the last statement.

3.1.2 WO_x/TiO₂ samples with intermediate coverage (2.1 and 2.2 W nm⁻²). The temperature-dependent trends of the in situ Raman and in situ FTIR spectra under flowing O2/He gas at intermediate coverages of 2.1 and 2.2 W nm⁻² for the P25 and anatase-supported samples (Fig. 4(A, B) and 5(A, B)) are mutatis mutandis similar to the ones described above pertaining to low coverage, though slightly affected by the coverage variation from low to intermediate. Hence, upon cooling (i) the Species-II band (predominant at 430 °C) undergoes a relative weakening and a redshift on going from 430 to 175 °C $(1005 \rightarrow 999 \text{ cm}^{-1} \text{ (R)} \text{ and } 2001 \rightarrow 1994 \text{ cm}^{-1} \text{ (IR) for}$ $2.1 \text{WTiO}_2(\text{P25})$ (Fig. 4) and $1007 \rightarrow 1002 \text{ cm}^{-1}$ (R) and $2003 \rightarrow$ 1992 cm⁻¹ (IR) for $2.2WTiO_2(a)$ (Fig. 5)); (ii) the Species-I band (minority at 430 °C) gradually emerges (due to weakening and redshift of the Species-II band) remaining in the fixed wavenumber position and becomes clearly resolved at 175 °C $(1010 \text{ cm}^{-1} \text{ (R) and } 2013 \text{ cm}^{-1} \text{ (IR) for } 2.1 \text{WTiO}_2(\text{P25}) \text{ (Fig. 4)},$ though somewhat less resolved for 2.2WTiO₂(a) (~1011 cm⁻¹ (R) and 2014 cm⁻¹ (IR) as seen in Fig. 5); (iii) the evidence for the presence of Species-III with characteristic peak mass at ~985 cm⁻¹ (R) and 1930–1970 cm⁻¹ (IR) is clear at 175 °C for 2.1WTiO₂(P25) (Fig. 4) and for the anatase-supported counterpart sample 2.2WTiO₂(a) in Fig. 5; (iv) a weak and broad peak progressively grows at ~935 cm⁻¹ in the in situ Raman spectra (Fig. 4(A) and 5(A)). Contrary to the case of samples with low coverage (Fig. 2(A) and 3(A)), the band feature at \sim 935 cm⁻¹ can be discerned already at 430 °C.

The non-single character of the observed Raman band at 430 °C is once again ascertained from the *in situ* FTIR spectra due to the doubling of the distance between the component bands in the overtone region. In addition, with decreasing temperature, the *in situ* FTIR spectra clearly confirm the emergence and clear appearance of the Species-I band and crosscheck the emergence of the Species-III peak mass at 175 °C. Significantly, the peak mass assigned to Species-III spreads to ~1930 cm⁻¹ in the IR overtone region, thereby implying the activity of a vibrational stretching mode that is less active in the Raman spectra, as further discussed below.

The spectral changes that take place upon cooling under *static equilibrium* conditions (Fig. 4(C) and 5(C), cells #11 and #12) follow similar trends to the ones described in the context of Fig. 2(C) and 3(C), *i.e.* band (I) undergoes neither an inten-

denced below.

sity change upon cooling to 250 °C and 175 °C nor any shift but band (II) relatively weakens with decreasing temperature. In addition, the weakening of band (II) goes along with the emergence of a peak mass ascribed to the formation of Species-III. Moreover, in this case (*i.e.* intermediate coverage), the observed changes appear to terminate upon cooling to

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250 °C.

3.1.3 High-loaded WO_x/TiO₂ samples (3.4 and 3.6 W nm⁻²). A much sharper band feature is observed in both the in situ Raman and in situ FTIR spectra (Fig. 6(A, B) and 7(A, B)) obtained at 430 °C for the high-loaded samples 3.4WTiO₂(P25) and 3.6WTiO₂(a), suggesting a higher prevalence of the "hightemperature" Species-II. The doubling of the bandwidth in the FTIR overtone region barely indicates a multiple band character at 430 °C (Fig. 6(B) and 7(B)). Upon cooling to 250 and 175 °C, the Species-II band consistently undergoes a relative weakening and redshift, while concomitantly a high wavenumber shoulder emerges, thus revealing the presence of Species-I. With decreasing temperature, the convoluted peak feature broadens in the in situ Raman and in situ FTIR spectra (Fig. 6 (A, B) and 7(A, B)) due to the gradual redshift of the Species-II band, as noted also for the low and intermediate coverages (vide ante). In addition, a peak mass suggestive of Species-III formation appears at 175 °C in both types of spectra that extends towards ~1930 cm⁻¹ in the FTIR spectra, thereby cross-checking that Species-III possesses an additional lower wavenumber mode, which is selectively more active in the infrared spectra. Moreover, a low resolution among the various bands is observed for the high loaded samples (Fig. 6 and 7), suggesting the occurrence of multiple closely related configurations resulting in the distribution of bond orders for the terminal W=O modes that give rise to closely packed overlapping bands. In addition, a broadband at ~945–950 cm⁻¹ extant already at 430 °C, progressively gains intensity upon cooling. Remarkably, the ~945–950 cm⁻¹ band is stronger for the anatase-supported sample compared to its P25-supported counterpart.

The changes in the Raman spectra upon cooling under *static equilibrium* conditions (Fig. 6(C) and 7(C), cells #7 and #6) are in agreement with the trends described for lower loaded samples (Fig. 2(C)–5(C)), *i.e.* the Species-II band gradually weakens and the Species-I band emerges. Therefore, the high wavenumber component of the convoluted band becomes more prominent. Moreover, clear evidence for the formation of Species-III is obtained, thereby confirming that Species-III is formed with decreasing temperature at the expense of Species-II.

3.1.4 Viewing angle of varying coverage at a constant temperature. One can organise the *in situ* Raman, *in situ* FTIR and *static equilibrium* Raman spectra shown in Fig. 2–7 by plotting the dependence on coverage at a constant temperature. Fig. 8 and 9 show representative pertinent examples for WO_x/TiO₂(P25) at 430 °C and 175 °C, respectively, *i.e.* at the highest and lowest temperatures as a function of coverage in the 1.2–3.4 W nm⁻² range. Fig. S1–S4† shows the corresponding dependencies for WO_x/TiO₂(P25) at 250 °C (Fig. S1†) and WO_x/

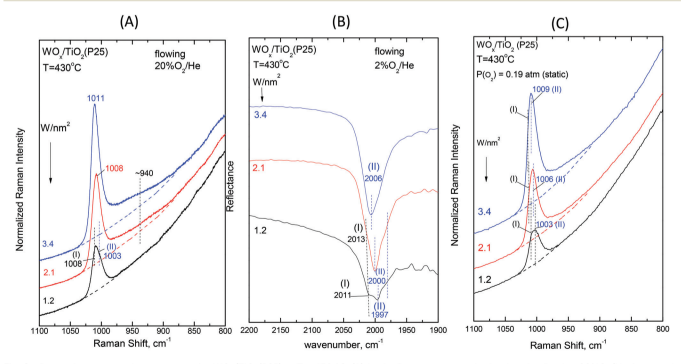
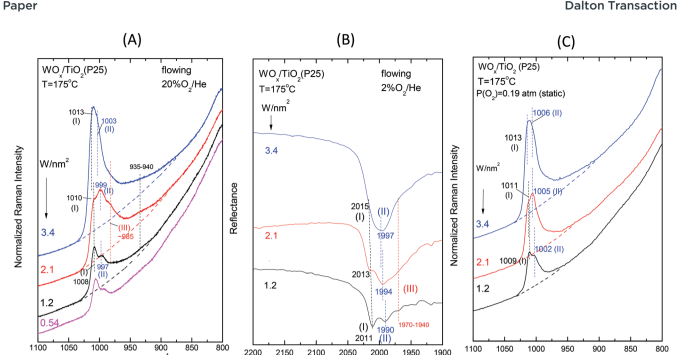


Fig. 8 Dependence on surface coverage for $WO_x/TiO_2(P25)$ at T=430 °C. (A) In situ Raman spectra obtained under flowing $20\%O_2/He$ for samples with W surface densities (W nm⁻²) as indicated by each spectrum; spectral conditions: see Fig. 2(A) caption. (B) In situ FTIR spectra obtained under flowing $2\%O_2/He$ for samples with W surface densities (W nm⁻²) as indicated by each spectrum. (C) Raman spectra at static equilibrium under $p_{O_2}=0.19$ atm for samples with W surface densities (W nm⁻²) as indicated by each spectrum; spectral parameters: see Fig. 2(C) caption.



wavenumber, cm

Fig. 9 Dependence on surface coverage for $WO_v/TiO_2(P25)$ at T = 175 °C. (A)–(C): see Fig. 8 caption.

TiO₂(a) at 430 °C, 250 °C and 175 °C (Fig. S2-S4†). Fig. 8 and 9 and Fig. S1-S4† do not show sequential transformations but portray collections of snapshots obtained for different samples at the same temperature. The following main observations are verified: (i) Species-I (bands at 1008 cm⁻¹(R) and 2011 cm⁻¹(IR)) is prevalent at low W surface densities and low temperatures; (ii) the band representing Species-II undergoes a blue shift with increasing surface density, best observed in the overtone region of the in situ FTIR spectra (Fig. 8(B) and 9(B)) where the wavenumber shifts are nearly doubled in the overtone region; (iii) Species-III is favoured at intermediate W surface densities (i.e. 2.1 and 2.2 W nm⁻²) and lower temperatures as seen in Fig. 9(A) and Fig. S4(A);† and (iv) the ~940 cm⁻¹ band gains intensity with increasing coverage and is entirely absent at 430 °C and low coverages of 1.2 and 1.1 W nm^{-2} (Fig. 8(A) and Fig. S2†).

Raman Shift, cm

3.2 Surface hydroxyl groups—temperature and coverage effects

To gain insight into the mechanisms governing the configurational changes that affect the vibrational properties of the dispersed WO_x phase, it is of importance to also address the spectral region of the surface hydroxyls. The vibrational properties of the surface hydroxyls on titanium dioxides have been extensively studied under dehydrated conditions, primarily by means of infrared spectroscopy. Worth mentioning are the often-referred pioneer studies of Primet et al.50 and Busca et al.51 In contrast, the Raman spectra of titania surface hydroxyls under dehydrated conditions are scarce.⁵² Most commonly, the OH stretching region of 3500-3750 cm⁻¹ is

explored. 50-53 The Ti-O(H) Raman stretching vibrations are masked by strong bands due to bulk titania; however, meticulous studies addressing the difference in Raman spectra were able to discern the Ti-O(H) stretching bands due to the isolated (\sim 700 cm⁻¹) and geminal (\sim 800 cm⁻¹) hydroxyls.⁵⁴

Raman Shift, cm⁻¹

Fig. 10 and 11 portray the temperature dependence of the Raman spectra obtained under static equilibrium in the wavenumber region pertaining to the OH groups. Fig. 10 pertains to the WO_x/TiO₂(P25) samples with coverages of 1.2 W nm⁻² (panel Fig. 10(A)), 2.1 W nm⁻² (panel Fig. 10(B)) and 3.4 W nm $^{-2}$ (panel Fig. 10(C)), respectively. Correspondingly, Fig. 11 pertains to the WO_x/TiO₂(anatase) samples with coverages of 1.1 W nm⁻² (panel Fig. 11(A)), 2.2 W nm⁻² (panel Fig. 11(B)) and 3.6 W nm $^{-2}$ (panel Fig. 11(C)), respectively. The Raman spectrum obtained for the pure support materials under static equilibrium at 430 °C (i.e. in cells containing pure $TiO_2(P25)$ and $TiO_2(anatase)$ sealed under 0.19 atm of $O_2(g)$ is included in each panel for reference. Four bands due to hydroxyl species can be discerned in the Raman spectra of each support type under O2(g) at 430 °C, albeit with different relative intensities. The bands at 3710, 3675, 3640 and 3540 cm⁻¹ are assigned to surface hydroxyls on the P25 carrier (spectra (a) in panels 10(A), 10(B) and 10(C)), whilst the bands at ~ 3705 , 3665, 3640 and 3540 cm⁻¹ due to hydroxyl species are observed for the counterpart anatase support (spectra (a) in panels 11(A), 11(B) and 11(C)).

Fig. 12 (panels (A) and (B)) shows the effect of coverage on the Raman spectra under static equilibrium at a constant temperature of 430 °C and sheds light on the progressive consumption of surface hydroxyls with increasing WO_x coverage. The **Dalton Transactions** Paper

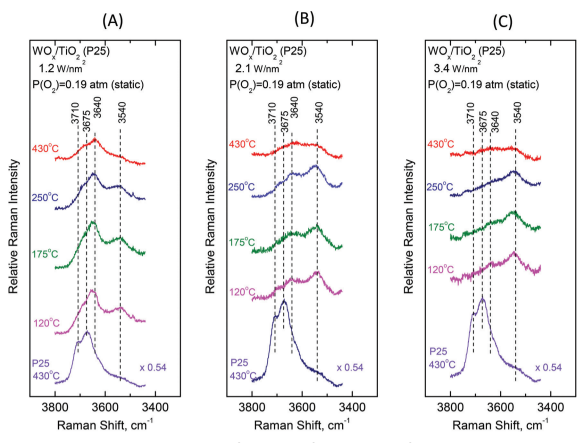


Fig. 10 WO_x/TiO₂(P25) with surface density of (A) 1.2 W nm⁻²; (B) 2.1 W nm⁻²; and (C) 3.4 W nm⁻². Sequential Raman spectra at static equilibrium under $p_{O_2} = 0.19$ atm at temperatures as indicated by each spectrum. Each panel includes the Raman spectrum of pure TiO₂(P25) at 430 °C as reference. Spectral parameters: $\lambda_0 = 532.0$ nm; laser power, w = 20 mW; and resolution, 2 cm⁻¹.

titration of surface hydroxyls and deposition of WO_x species results in the diminishing of hydroxyl groups and condensation of water molecules that remain on the surface by hydrogen bonds.27,30 Hence, with increasing coverage a gradual weakening of the hydroxyl bands is observed as shown in Fig. 12. Significantly, the OH band intensities are not attenuated simultaneously, but in a specific order of priority as follows. In the case of P25-supported materials (Fig. 12, panel (A)), the OH group represented by the 3710 cm⁻¹ band is selectively titrated first and it nearly disappeared for coverage of 1.2 W nm⁻², followed by the OH group represented by the 3675 cm⁻¹ band and finally the 3640 cm⁻¹ band is also eventually diminished, whilst the 3540 cm⁻¹ OH band is not at all affected during the deposition. Likewise, a corresponding order of priority is evidenced in the attenuation of the OH bands with increasing WO_x coverage for the anatase-supported catalysts (Fig. 12, panel(B)), namely the ~3705 cm⁻¹ band is attenuated first, followed by the 3665 cm⁻¹ band and then by the 3640 cm⁻¹ band whereas the 3540 cm⁻¹ band does not undergo any weakening with increasing coverage. Notably, a similar order of priority of OH band attenuation as a function of coverage has been reported in in situ FTIR studies of vanadia/titania catalysts.^{28,55} Finally, it is worth pointing out that lower basicity for surface hydroxyls is evidenced for the anatase-supported samples compared to their P25-supported counterparts, thereby accounting the lower respective band wavenumber⁵² for the respective hydroxyls (3705 vs. 3710 cm⁻¹ and 3665 vs. 3675 cm⁻¹).

Significantly, the order of priority for the hydroxyl bands' intensities' diminishing with increasing WOx coverage at 430 °C is in perfect agreement with the corresponding order of WO_x species appearance and evolution of species' prevalence as a function of coverage. Hence, the diminishing of the 3710/ 3705 cm⁻¹ hydroxyl band (due to the type-I terminal and the most basic OH group⁵² that is titrated first, vide infra) at low coverage is accompanied by the appearance and significant presence of Species-I, whilst the successive diminishing of the 3675/3665 and 3640 cm⁻¹ bands (due to the OH groups of lower basicity and the type-II hydroxyls, 52 vide infra) comes along with the appearance and progressive prevalence of Species-II with increasing WO_x coverage (see Fig. 8 and 9 and Fig. S1-S4†). Notably, the formation of Species-I ceases upon the elimination of the 3710/3705 cm⁻¹ OH band, which is further discussed below.

¹⁸O/¹⁶O isotope exchange and *in situ* Raman spectra

Examining the vibrational isotope effects and the band splitting pattern caused by the ¹⁸O/¹⁶O isotope exchange can shed

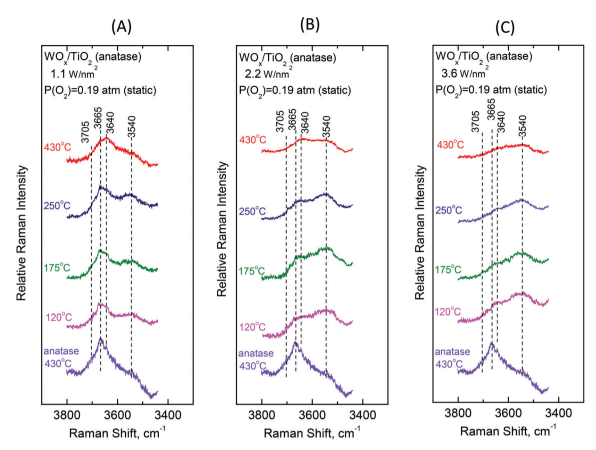


Fig. 11 WO_x/TiO_2 (anatase) with surface density of (A) 1.1 W nm⁻²; (B) 2.2 W nm⁻²; and (C) 3.6 W nm⁻². Sequential Raman spectra at static equilibrium under $p_{O_2} = 0.19$ atm at temperatures as indicated by each spectrum. Each panel includes the Raman spectrum of pure TiO_2 (anatase) at 430 °C as reference. Spectral parameters: $\lambda_0 = 532.0$ nm; laser power, w = 20 mW; resolution, 2 cm⁻¹.

light on the termination configuration of the dispersed oxometallic sites. 13,23 The present isotope exchange study is limited to the temperature of 470 °C that is required in order to attain a satisfactory reduction of the dispersed WVIOx species under flowing 5% H₂/He for 20 min which would enable ¹⁸O/¹⁶O exchange by reoxidation under flowing 2% ¹⁸O₂/He for 10 min (i.e. the two steps comprise one isotope exchange redox cycle). As described above, at high temperatures (i.e. at 430 °C and a fortiori at 470 °C) Species-I is prevalent at low coverage and Species-II prevails at high coverage. Hence, the oxygen-isotopeexchange Raman measurements primarily contribute to determine the termination configuration of the high-temperature prevailing species in each case. Therefore, representative samples of low and high surface densities (i.e. ~0.5 W nm⁻² and $\sim 3.5 \text{ W nm}^{-2}$) were chosen for the isotope exchange study and Fig. 13 and 14 show the pertinent results.

Single isotopic splitting is observed for all the samples subjected to oxygen isotopic substitution studies, *i.e.* 0.51 WO_x/TiO₂(P25), 0.54 WO_x/TiO₂(a), 3.4 WO_x/TiO₂(P25) and 3.6 WO_x/TiO₂(a). One single symmetric band due to W= 18 O stretching emerges for all the samples at a fixed wavenumber in each case. Progress in the extent of the gradual 18 O/ 16 O isotope exchange is completed under the applied conditions after 13–15 isotope exchange redox cycles. Moreover, worth men-

tioning is a gradual slight redshift ($ca. 2 \text{ cm}^{-1}$) of the W= 16 O mode due to the remaining (i.e. non-substituted) W= 16 O sites that can be discerned with the help of the corresponding vertical inspection lines in Fig. 13 and 14.

Discussion

The heterogeneity of the WO_x amorphous phase dispersed on titania supports (P25 and anatase) has preliminarily been addressed in previous reports. The unprecedented effect of tunable speciation of the dispersed heterogeneous oxometallic phases has already been comprehensively explored for the ${\rm Re^{VII}O_x/TiO_2(P25)}$ catalysts by means of *in situ* Raman and FTIR spectroscopy and *in situ* Raman/¹⁸O isotope exchange. The role of surface non-titrated hydroxyls and water molecules retained by the surface²⁷ (formed during the condensation steps involved in the impregnation/deposition steps) has been given documented prominence. The surface of the work of of the wo

Arguments attributing the occurrence of water molecules as if they were part of the incoming gas feed during the *in situ* spectroscopic measurements were precluded by means of neoteric (as far as catalysis science is concerned) experiments, namely temperature-dependent *static equilibrium* Raman

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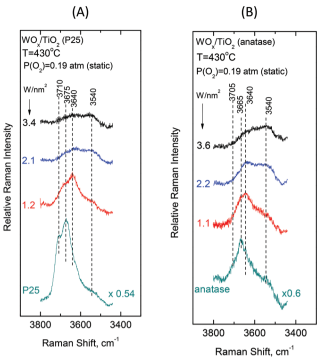


Fig. 12 Dependence on surface coverage at T = 430 °C for (A) WO_x/ TiO₂(P25) and (B) WO_x/TiO₂(anatase). Raman spectra at static equilibrium under p_{O_2} = 0.19 atm for samples with W surface densities (W nm⁻²) as indicated by each spectrum. Each panel includes the Raman spectrum of the corresponding pure TiO2 support at 430 °C as

studies of samples sealed under an $O_2(g)$ atmosphere after evacuation at an elevated temperature (200 °C).³⁰

In the present work, we are concerned with the challenge of elucidating the molecular structures and the termination configurations (i.e. mono-oxo vs. di-oxo etc.) of the species present within the oxo-wolfram(VI) sites dispersed on titania and proposing consistent molecular-level explanations for the reversible temperature-dependent transformations taking place. The above described in situ Raman, in situ FTIR, in situ Raman/¹⁸O isotope exchange and static equilibrium Raman results (the latter extending also in the hydroxyl spectral region) provided adequate evidence for the occurrence of three species in the temperature range of 430-175 °C (Species-I, Species-II and Species-III) depending on temperature and surface density (W nm⁻²). Below, we propose our full interpretation of the results and discuss adequately its full consistency with all the observed effects.

4.1 Interpretation of spectroscopic (in situ Raman, in situ FTIR, in situ Raman/¹⁸O isotope exchange and static Raman) results

Three distinct WO_x configurations (I, II and III) occur within the oxo-wolfram phase dispersed on TiO₂(P25) and TiO₂(anatase). On grounds of mononuclear species representation, Species-I has a mono-oxo termination configuration and fourfold coordination for W in an O=W(-O-Ti)₃ distorted C_{3v} like arrangement; likewise, Species-II has a mono-oxo termin-

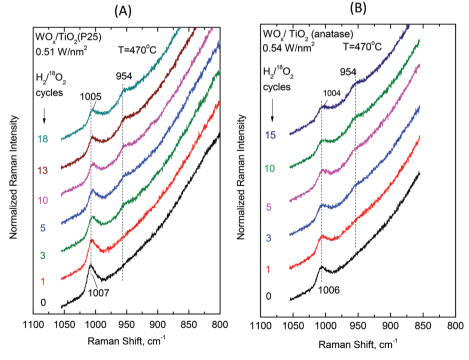


Fig. 13 (A) WO_x/TiO₂(P25) with surface density of 0.51 W nm⁻² and (B) WO_x/TiO₂(anatase) with a surface density of 0.54 W nm⁻². Sequential in situ Raman spectra obtained under flowing (10 cm³ min⁻¹) 2% ¹⁸O₂/He after subsequent H₂/¹⁸O₂ reduction/oxidation isotope exchange cycles as indicated by each spectrum. Recording parameters: see Fig. 2(A) caption.

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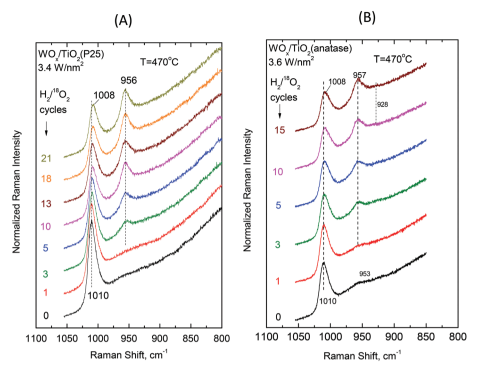


Fig. 14 (A) WO_x/TiO₂(P25) with a surface density of 3.4 W nm⁻² and (B) WO_x/TiO₂(anatase) with a surface density of 3.6 W nm⁻². Sequential in situ Raman spectra obtained under flowing (10 cm³ min⁻¹) 2% ¹⁸O₂/He after subsequent H₂/¹⁸O₂ reduction/oxidation isotope exchange cycles as indicated by each spectrum. Recording parameters: see Fig. 2(A) caption.

ation configuration albeit with a fivefold coordination for W in an $O=W(-O-Ti)_4$ distorted C_{4v} -like arrangement and Species-III has a di-oxo termination configuration with a fivefold W coordination in an (O=W)2(-O-Ti)3 arrangement. Species-II and Species-III occur also in the associated (W-O-W linked) oligomeric form with increasing surface coverage. Fig. 15 depicts the molecular models for Species-I, Species-II and Species-III in mononuclear representations.

4.1.1 Species-I and Species-II with mono-oxo termination configuration-vibrational properties (Raman and IR) and vibrational isotope effects. The temperature-dependent changes in the relative intensities of bands (I) and (II) observed in Fig. 2-7 show undoubtedly that they represent the W=O stretching modes of two distinct species. The striking similarity of the *in situ* Raman spectra in the fundamental W=O stretching region with its counterpart in situ FTIR spectra in the respective overtone region provides sound evidence that both species have a mono-oxo termination configuration. Considering the anharmonicity (accountable for observing the overtone modes) for W=O in diatomic approximation, the observed fundamental and observed first overtones are given by 13,56,57

$$\nu_{W=0,1\leftarrow 0} = \omega_{W=0}(1-2\chi_{W=0})$$

$$\nu_{\mathrm{W=O,2\leftarrow0}} = \omega_{\mathrm{W=O}} (1\text{--}3\chi_{\mathrm{W=O}})$$

where $\omega_{W=0}$ is the wavenumber corrected for anharmonicity and $\chi_{W=0}$ is the anharmonicity constant. Hence, the FTIR overtones would be expected at wavenumbers slightly lower than the

respective doubled Raman fundamentals because the vibrational energy levels are not equidistant, but their separation decreases with increasing vibrational quantum number.⁵⁶ This is exactly the case for bands marked as "I" and "II" in all spectra displayed in Fig. 2(A, B)-7(A, B). Utmost care is needed in assigning bands as due to overtones. Assignments must comply with the requirement for the first overtone wavenumber to be lower than the doubled wavenumber of the respective fundamental. Sometimes, wrong pertinent interpretations are reported, e.g. previously, a band at ~2000 cm⁻¹ has been erroneously assigned to the first overtone of a 985 cm⁻¹ W=O fundamental.⁴⁰

It is shown below that Species-II occurs partly in the associated $(WO_x)_n$ form at coverages above half monolayer at high temperatures (i.e. 430 °C) and also undergoes partial association with oligomeric units possessing W-O-W bridges when exposed to gradually decreasing temperatures (430 \rightarrow 250 \rightarrow 175 °C) under oxidative dehydrated conditions. Note that with increasing coverage, Species-II gradually prevails over Species-I (Fig. 8 and 9, and Fig. S1-S4†). Hence, in agreement with the very well documented study of Hilbrig et al.,35 which shows the existence of the WO5 and WO4 units with the WO5/WO4 ratio increasing with increasing coverage, Species-I is assigned to tetra-coordinated O=W(-O-)3 units and Species-II to penta-coordinated $O=W(-O-)_4$ units, in conformity with the relative band wavenumbers for their respective W=O terminal stretching modes (i.e. $\nu_{W=O, Species-II} > \nu_{W=O, Species-II}$).

Notably, there exists a vital difference between the temperature dependence of bands (I) and (II). With decreasing temp**Dalton Transactions** Paper

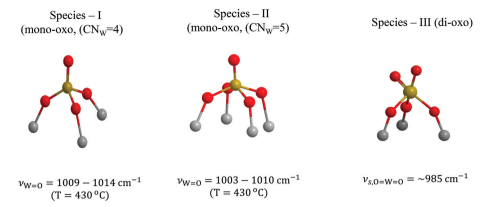


Fig. 15 Structural models and W coordination/termination characteristics for Species-I, Species-II and Species-III in mononuclear representations.

erature (panels (A) and (B), Fig. 2-7), band (II) loses intensity and undergoes a progressive redshift, whilst band (I) remains unaffected both in terms of its position as well as its intensity. As will be explained below, the structural transformation accountable for the weakening and redshift of band (II) is mediated by surface water molecules formed by condensation of surface hydroxyls during the deposition titration steps and retained at the surface by H-bonds. The stability exhibited by Species-I with decreasing temperature in the sequence 430 °C \rightarrow 250 °C \rightarrow 175 °C precludes the provenance of the water molecules as being in the incoming feed gas (20% O₂/He). If water molecules were incoming as part of the feed gas, then there would be no selectivity of their "action" on Species-I and Species-II.

A critical inspection of Fig. 12 sheds light on the order of priority of Species-I and Species-II formation. The most basic surface hydroxyls (bands at ~3710/3705 cm⁻¹)⁵² are titrated first i.e. at low coverages of e.g. 1.2 W nm⁻² and 1.1 W nm⁻² (spectra 12Ab and 12Bb); hence, Species-I that prevails at low coverage (Fig. 8 and 9, and Fig. S1-S4†) is formed preferentially on those particular receptor sites where the most basic hydroxyls have first been titrated. Eventually, such sites are completely consumed and do not exist above ca. 1.5 W nm⁻², when only hydroxyls with lower basicity remain for being titrated leading to the formation of Species-II (CN_W = 5), which thereby gradually prevails over Species-I (CN_W = 4) with increasing coverage.

Hence, Species-I and Species-II are distinct, they both have a mono-oxo termination configuration and Species-II prevails over Species-I with increasing coverage. Therefore, in alignment with the previously documented prevalence of WO₅ sites over WO₄ sites with increasing coverage, 35 the C_{3v} -like and C_{4v} like configurations are the most plausible ones, as depicted in Fig. 15. The $CN_W = 4$ for Species-I justifies the higher wavenumber for $\nu_{W=O, Species-II}$ compared to $\nu_{W=O, Species-II}$.

The Raman spectra obtained under the static equilibrium also show: (i) the gradual prevalence of Species-II over Species-I with increasing coverage (panels (C) shown in Fig. 8 and 9 and Fig. S1-S4†) and (ii) the progressive weakening of band (II) with decreasing temperature (panels (C), Fig. 2–7). In contrast, the

gradual redshift of band (II) observed with decreasing temperature in the in situ Raman and FTIR spectra (Panels (A) and (B), Fig. 2-7) is hardly observed under static conditions. This is due to limiting the effect of surface water molecules ascribed to their partial removal during the 1 h evacuation of the samples at 200 °C before sealing off the cells for the static equilibrium Raman experiments (see the Experimental section).

Under in situ conditions, a temperature decrease of the order 430 °C \rightarrow 175 °C results in the activation of water molecules retained by the surface (vide ante)27,30 for hydrolyzing the anchoring W-O-Ti bonds, thereby resulting in the formation of Species-III (to be further discussed below) and the partial association of WO5 Species-II sites by means of W-O-W linkages. The latter is justified by observing the broad peak mass evolved with decreasing temperature in the ~935-950 cm⁻¹ range (Fig. 2(A)–7(A)), of which the relative intensity increases with increasing coverage at a constant temperature (Fig. 8(A)-9 (A) and $S1(A)-S4(A)^{\dagger}$). Hence the broadband at 935–950 cm⁻¹ is attributed to the W-O-W bridging modes. Previously, in the in situ Raman studies pertaining to the WO_x/ZrO₂ catalysts, 58-60 broadbands at 905-910 cm⁻¹ have been observed with relative intensities that increased with increasing loading and assignment to W-O-W has been suggested to account for the observations.⁵⁸ Significantly, such a peak mass at 935-950 cm⁻¹ already exists for coverages above the half monolayer (Panels (A), Fig. 6 and 7) and can clearly be discerned for coverages above 2 W nm⁻² already from 430 °C as shown in Fig. 8(A) and Fig. S2(A)†, that show the dependence on coverage at 430 °C. Hence, although the monomer-to-associated species transformation is completely reversed upon heating for low coverages of e.g. \sim 1.2 W nm⁻², this is not the case for high coverages where associated species existed already at 430 °C (see e.g. Fig. 6(A) and 7(A)). The formation of W-O-W bridges, which are inherently stronger than their counterpart W-O-Ti anchors, results in the redistribution of charge and valency around the W atom, thereby weakening of the Species-II W=O terminal bond and hence accounting for the redshift undergone by band (II) with decreasing temperature.

The W=O terminal stretching mode wavenumbers for Species-I and Species-II undergo a blue shift with increasing

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coverage (Fig. 8, 9 and S1-S4†) due to the varying basicity of the corresponding "receptor" surface OH sites. It is evident that the most basic hydroxyls are titrated first, i.e. at low coverage preferably the most basic hydroxyls are titrated. With increasing coverage, hydroxyls with lower basicity are also titrated. The lower basicity of a titrated hydroxyl accounts for the lower electron donating ability of O towards W along with Ti-O-W and by the cascade effect to the strengthening of the terminal W=O bond with increasing coverage. Actually, with increasing coverage, the $\nu_{W=0}$ for Species-II (band (II)) undergoes a pronounced blue shift (e.g. in Fig. 8, pertaining to WO_x/ $TiO_2(P25)$ at 430 °C: 1003 \rightarrow 1010 cm⁻¹ (R), 1997 \rightarrow 2006 cm⁻¹ (IR overtone) and $1003 \rightarrow 1009 \text{ cm}^{-1}$ (R, static)), which is justified by the progressively lower basicity of the titrated hydroxyls, thereby—by the cascade effect—resulting in the strengthening of the terminal W=O bond. Concomitantly, whereas the formation of Species-I ceases at low-to-medium coverage due to the full consumption of the most basic hydroxyls, increasing amounts of Species-II are formed at medium-to-high coverage and band (II), on account of its intensity increase and blue shift, obscures the unshifted band (I) at high temperature (430 °C) and high coverage (e.g. Fig. 6 (panels (A),(B)) and Fig. 7 (panels (A),(B))).

Therefore, Species-II is the dominant species at high coverage and high temperature, whilst Species-I is prevailing at low coverage. Hence, the in situ Raman/18O isotope labeling measurements performed for samples with coverages of ~0.5 W nm⁻² (Fig. 13) and \sim 3.5 W nm⁻² (Fig. 14) at the temperature of 470 °C address exclusively the confirmation of the termination configuration for Species-I and Species-II, respectively. In all the cases, a single isotopic splitting is observed, thereby cross-checking the proposal for mono-oxo termination configuration for both Species-I and Species-II. Notably, multiple band splitting would be expected in the case of ¹⁸O isotope labeling Raman measurements pertaining to poly-oxo (e.g., dioxo) termination configuration for the examined dispersed $(WO_x)_n$ sites. ^{13,31,61} Such multiple isotopic band splitting has indeed been observed in recent in situ Raman/18O isotope exchange studies for di-oxo (O=)2ReOx sites dispersed on TiO₂(P25)³¹ and CeO₂.⁶²

One can exploit the diatomic model in the harmonic approximation 13,23,31,61 for calculating the corresponding theoretically predicted wavenumber for the W=18O, in each case. This is done on the grounds of equal W-O distances for the W=16O and substituted W=18O sites. Hence, by applying the isotopic ratio of 1.0554 (ref. 61) the calculated wavenumbers for the W=18O modes in each case coincide with the values indicated in Fig. 13 and 14, thereby further strengthening the validity of the proposed mono-oxo configuration for the distinct Species-I and Species-II.

A point worth discussing in the in situ Raman/18O exchange measurements is the slight gradual redshift (e.g. 1010 > 1008 cm⁻¹ in Fig. 14) of the remaining non-substituted W=16O sites. This is ascribed to the previously reported nextnearest-neighbor vibrational isotope effect. 13,45 Very briefly, the DFT theoretical calculations⁶³ predicted that the hydrogen

most favorably adsorbs dissociatively on titania surface unsaturated Ti-O-Ti and Ti-O sites (deprotonated former hydroxyl sites), hence preferably reducing titania rather than a W=16O site, by leading the initial steps of ¹⁸O/¹⁶O exchange to take place on lattice oxygens and by surface diffusion to anchoring oxygens of W-O-Ti linkages (W- 16 O-Ti \rightarrow W- 18 O-Ti). The lower nuclear charge density of ¹⁸O (compared to ¹⁶O) causes —by the cascade effect—a weakening of the remaining (yet non substituted) W=16O bonds.13,45

Another comment worth mentioning on the vibrational properties of the W=O terminal stretching modes of Species-I and Species-II pertains to their respective band intensities in the Raman and FTIR spectra. A comparison of the in situ Raman spectra with the in situ FTIR spectra, e.g. at low coverages of 1.2 and 1.1 W nm⁻² (i.e. Fig. 2(A, B) and 3(A, B)), shows that the intensity ratio of band (II) to band (I) is somewhat lower in the Raman spectra compared to the FTIR spectra, i.e.

$$\left(\frac{I_{W=O(II)}}{I_{W=O(I)}}\right)_{R} < \left(\frac{I_{W=O(II)}}{I_{W=O(I)}}\right)_{IR}$$

This is ascribed to the different selection rules pertaining to the Raman vs. IR band intensities, namely change in polarizability vs. change in dipole moment.

4.1.2 Species-III with di-oxo termination configuration vibrational properties (Raman and IR). The spectral changes upon cooling the WO_x/TiO₂(P25 and anatase) samples under dehydrated conditions invariably show the gradual appearance and progressive growth of band (III), which is a peak mass at ~985 cm⁻¹, i.e. located in the low wavenumber wing of band (II) (see Fig. 2-7, panels (A) and (C)). The corresponding observation in the overtone region of the in situ FTIR reveals upon cooling to 175 °C, the appearance of a broad peak mass extending in the 1970-1930 cm⁻¹ range (see Fig. 2-7, panels (B)). Band (III) becomes clearly discernible at 175 °C and its width and wavenumber position suggest that the band consists of two components, namely the symmetric and antisymmetric stretching (ν_s/ν_{as}) of a di-oxo O=W=O core possessed by a dispersed WO_x site formed at the expense of Species-II, of which the representative band (II) undergoes a corresponding weakening (vide ante). A mechanistic interpretation at the molecular level (presented below) suggests a five-fold coordination for W (CN_W = 5) for the di-oxo (O=)₂W(-O-)₃ site formed at the expense of Species-II following the hydrolysis of an anchoring bond by surface-retained water, activated and favored by lowering the temperature to and below 250 °C.

Previously, reference model compounds possessing di-oxo WO2 cores with W in environments consisting exclusively of oxo-ligands with W in six-fold coordination, namely $W^{VI}O_2(SO_4)_2^{\ 2-}$ and $W^{VI}O_2(SO_4)_3^{\ 4-}$ molten complexes, ⁶⁴ exhibited the (ν_s/ν_{as}) pairs at 972/937 cm⁻¹ and 933/909 cm⁻¹, respectively. The five-fold coordination evidenced for W in the di-oxo Species-III justifies the slight wavenumber upshift. Hence, ν_s is located at ~985 cm⁻¹ (Fig. 2–7) and its ν_{as} counterpart, expected with much lower intensity in the Raman and

some 15–40 cm⁻¹ below $\nu_{\rm s}^{57}$ is obscured by the decaying envelope of band (II) and the ascent of the background towards the low wavenumber side of the spectra. Ideally, in the triatomic "gas phase" approximation for WO₂, the symmetric stretching $\nu_{\rm s}$ mode should be IR-silent⁵⁷ but distortions within the amorphous dispersed (WO_x)_n phase relax the symmetry selection rules and both $\nu_{\rm s}$ and $\nu_{\rm as}$ become active in both types of vibrational spectra.

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4.1.3 Effect of support type. Certain comparative observations between the two sets of catalysts (i.e. supported on P25 and anatase) are worth to be discussed. The comparison of the vibrational spectra obtained for the two series of catalysts shows that there are no significant consistent variations in the distributions of Species-I and Species-II on the two supports. Specifically, at low coverage of 1.2 and 1.1 W nm⁻² on P25 and anatase (Fig. 2 and 3) where bands (I) and (II) are clearly separated due to the prevalence of monomeric sites, thereby allowing for a direct comparison, the distribution is mutatis mutandis similar on the two supports. Variations observed by visual comparison at constant coverage and temperature are primarily ascribed to the differences in band wavenumbers. Hence, there is a consistent tendency for the band (I) and band (II) wavenumbers to be slightly higher for anatase-supported samples compared to their P25-supported counterparts of similar coverage. Such a difference is justified in view of the respective lower basicity of the receptor surface hydroxyls, e.g. for Species-II on anatase compared to P25 (OH bands at 3665 vs. 3675 cm^{-1} respectively, see section 3.2).

The FTIR spectra obtained for high loadings are dominated by band (II) both due to Species-II prevalence and due to the intrinsically higher IR activity of band (II) (see section 4.1.1). Moreover, as seen in Fig. 2–7, the redshift incurred by band (II) upon cooling is higher for P25-supported samples. Hence, when comparing the band (II) redshifts upon cooling in the *in situ* FTIR spectra for the highest W coverage (Fig. 6(B) ν s. 7(B)) where Species-II is predominant, a larger shift is evidenced for the P25-supported sample (2006 \rightarrow 1997 cm⁻¹) compared to its anatase-supported counterpart (2005 \rightarrow 2002 cm⁻¹).

4.2 Structural models and molecular-level mechanism for temperature-dependent structural transformations

The structural transformations taking place under dehydrated conditions upon cooling the WO_x/TiO₂(P25, anatase) catalysts were interpreted above, based on the combined *in situ* Raman, *in situ* FTIR and *static equilibrium* Raman experiments by means of two processes at the molecular level: (i) Species-II (e.g. O=W(-O-)₄ in mononuclear representation) transforms to Species-III (e.g. (O=)₂W(-O-)₃ in mononuclear representation) and (ii) Species-II undergoes oligomerization. Notably, Species-I (O=W(-O-Ti)₃) was found to be stable and not subjected to temperature-dependent structural/transformational changes in the sequence 430 °C \rightarrow 175 °C \rightarrow 430 °C. The structural transformations undergone by Species-II are ascribed to the action of surface water molecules retained by the support²⁷

that become activated and hydrolyze the anchoring sites of Species-II upon lowering the temperature.

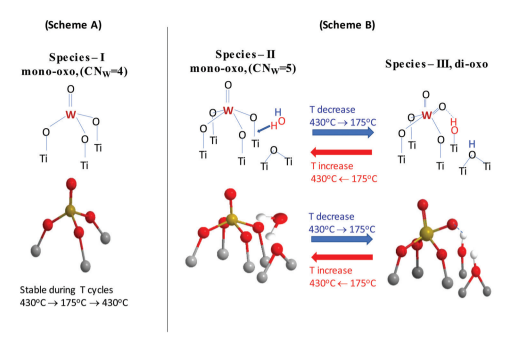
Fig. 16 portrays a consistent molecular level mechanism accounting for the spectral observations and the proposed transformations. Scheme (A) highlights the stability of Species-I that as evidenced by its vibrational properties does not undergo changes when cycling the temperature in the 430-175 °C range. Scheme (B) shows schematically the hydrolyzing action of a surface H₂O molecule (activated upon lowering the temperature) on an anchoring W-O-Ti bond of Species-II nearby a deprotonated Ti-O-Ti site (it could also be a Ti-O site) that had been formed by the titration of surface hydroxyls during the deposition process steps. For simplicity, a mononuclear representation for Species-II is shown. As a result, on account of the H2O molecule hydrolyzing action, the O atom of the W-O-Ti chain is detached from the Ti atom, thereby Species-II [(O=W(-O-Ti)₄] is transformed to Species-III [(O=)2W(-O-Ti)3] and two hydroxyls are formed on the surface. One hydroxyl group is formed by the hydroxylation of the liberated Ti atom and a second hydroxyl group is formed by the protonation of the Ti-O-Ti site. Scheme (B) also includes a perspective molecular model view of the abovedescribed step.

Scheme (C) shows the parallel oligomerization undergone by two adjacent Species-II moieties, mediated by two surface water molecules. The neighboring dehydroxylated (Ti–) and deprotonated (Ti–O–Ti) sites are also shown. The activation of the two surface H_2O molecules results in an oxygen-bridged associated dinuclear Species-II unit (*i.e.* comprised of two oxygen-bridged mono-oxo W=O sites with $CN_W = 5$) and four hydroxyls in a stoichiometrically consistent step. Scheme (C) includes also the corresponding structural model perspective view. The process shown in Fig. 16(C) does not take place under *static equilibrium* conditions due to the partial removal of the retained water molecules caused by the samples' evacuation for 1 h at 200 °C, thereby partly inhibiting the temperature-dependent effects.

4.3 Implications on WO_x/TiO₂-based catalysis

The results of the present work cannot be used directly to assign a particular constituent of the $(WO_x)_n$ dispersed phase as being of relevance for catalytic activity in WO_x/TiO₂-based catalysis. However, sound evidence is provided for ascertaining the heterogeneity of the dispersed phase and it remains to differentiate the spectator species and the catalytically active sites for each particular catalytic process without excluding the possibility of the active phase itself being heterogeneous, i.e. consisting of more than one active site. In contrast, it is clearly shown that a controlled tuning of the prevailing sites/configurations is feasible by appropriate control of temperature and surface coverage. Most importantly, it turns out that existing results and interpretations pertaining to temperature-dependent and coverage-dependent reactivity/selectivity for WOx/ TiO₂-based catalytic reactions probably need to be reevaluated by taking into account the corresponding dependence of configurations/site prevalence.

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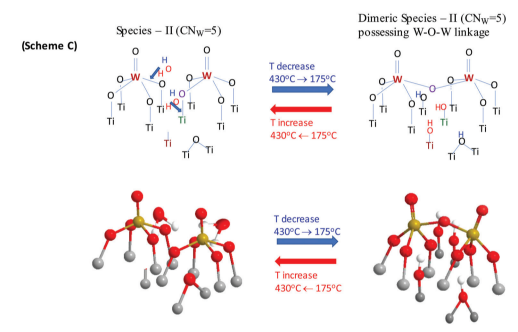


Fig. 16 Temperature dependent structural transformations. Scheme (A) highlights the stability of Species-I in the 430 °C \rightarrow 175 °C \rightarrow 430 °C cycle. Scheme (B): molecular level mechanism of Species-III T-dependent transformation. Scheme (C): molecular level mechanism of Species-II association via W-O-W bridging linkages.

5. Conclusions

- (a) The dispersed $(WO_x)_n$ phase supported on TiO_2 is heterogeneous. At an elevated temperature of 430 °C it consists of two distinct species with the mono-oxo termination configuration exhibiting C_{3v} -like (Species-I, mono-oxo, $CN_W = 4$) and C_{4v} -like (Species-II, mono-oxo, $CN_W = 5$) arrangements.
- (b) Species-I (WO₄) is formed by titration of the most basic support hydroxyls and is abundant at low coverage; its for-

mation ceases upon consumption of the most basic surface hydroxyls that are exclusive receptor sites for Species-I formation. The $\nu_{\rm W=O}$ mode for Species-I is at 1009–1014 cm⁻¹ and remains stable in terms of intensity and position in the temperature sequence of 430 \rightarrow 250 \rightarrow 175 °C.

(c) Species-II (WO₅) is formed by titration of the less basic support hydroxyls, in amounts that increase with increasing coverage and prevails over WO₄ at high coverage and high temperature. The association of WO₅ units takes place at and

above half monolayer, evidenced by a band attributed to W–O–W links. The $\nu_{\rm W=O}$ mode for Species-II is at 1003–1010 cm⁻¹ (blue shifting and strengthened with increasing coverage) and undergoes a reversible weakening and redshift in the temperature sequence of 430 \rightarrow 250 \rightarrow 175 °C.

- (d) Two reversible temperature-dependent transformations of Species-II mediated by water molecules retained on the surface take place in the $430 \rightarrow 250 \rightarrow 175 \rightarrow 430$ °C sequence. First, Species-II by hydrolysis of one of its W–O–Ti anchors is transformed into Species-III (di-oxo, characteristic symmetric stretching at ~985 cm⁻¹) and second, Species-II undergoes partial association to oligomerized units. The latter effect is more pronounced at low-to-medium coverage where isolated O=W (–O–)₄ units occur in adjacent locations; at medium-to-high coverage, the polymeric units already exist after calcination.
- (e) Subjecting the catalysts to 1 h evacuation at 200 °C, thus limiting the amount of surface retained water, results in the partial inhibition of the observed effects thereby restricting the Species-II polymerisation upon lowering the temperature.
- (f) The results show the feasibility of tuning the speciation of the $(WO_x)_n$ phase dispersed on TiO_2 by appropriate control of temperature and coverage.

Conflicts of interest

There are no conflicts of interest to declare.

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