Revisiting the Use of 2,6-Dimethylpyridine Adsorption as a **Probe for the Acidic Properties of Metal Oxides**

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Received May 14, 2001. In Final Form: July 9, 2001

To check if the adsorption/desorption of 2,6-dimethylpyridine (2,6-DMP) is a suitable probe for the surface acidity of oxides, and to assign a band of controversial interpretation often observed upon 2,6-DMP adsorption, the ambient temperature uptake of 2,6-DMP on several oxidic systems was carried out. SiO₂ was observed to yield a plain H-bonding interaction, characterized by an 8a band of adsorbed 2,6-DMP that is typical of silanols. Several (spinel) transition aluminas and the corundum α -alumina phase were examined and allowed to test the suitability of 2,6-DMP in distinguishing different types of surface Lewis acidity. Sulfated alumina and a β -zeolite in its H-form were used to check the sensitivity of 2,6-DMP uptake toward either induced or intrinsic Brønsted acidity and the possibility of observing (weak) Lewis-coordinated 2,6-DMP also in the presence of (strong) Brønsted-bound 2,6-DMP.

Introduction

IR spectroscopy of the adsorption/desorption of basic molecules, like for instance ammonia or pyridine and its derivatives, is an often used method to characterize the surface acidity of solids, in that this method is considered to be able to titrate concentration and strength of surface acid sites.

The adsorption of pyridine (py) as a tool to test the acidity of oxidic systems has been proposed long ago.1 This molecule can distinguish between Lewis and Brønsted acidity, giving rise to specific IR absorption bands.^{1,2} Among the various bands, one located at \sim 1450 cm⁻¹ (due to Lewis-coordinated py) and one located at $\sim 1540~\rm cm^{-1}$ (due to Brønsted-bound py) are both associated with the 19b ring mode of py and are usually considered as analytical of the interaction of py with coordinatively unsaturated (cus) Me^{n+} cations and with protonic sites,³ respectively.

If py contains substituent groups in the 2- and 6-position, as in the case of lutidine (i.e., 2,6-dimethylpyridine, hereafter referred to as 2,6-DMP), the formation of coordinative chemisorption bonds is expected to be sterically hindered.

In the literature, many works are present dealing with the adsorption of sterically hindered pyridines onto oxidic systems. ^{4–13} Some authors ^{4–6} have suggested, on the basis of 2,6-DMP adsorption/desorption measurements on dif-

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ferent oxidic systems, that this base plays the role of specific proton probe, also in the presence of surface Lewis acid sites. In fact, they report that this base interacts first with Brønsted acid sites and subsequently and only under high adsorptive pressures with Lewis sites. Moreover, some authors claim that when 2,6-DMP is adsorbed, they cannot distinguish among Lewis acid sites of different acidic strengths (e.g., see refs 9 and 13), and most authors report that the Lewis coordinative interaction of 2,6-DMP is so weak (due to steric hindrance) that is never realized for adsorption/desorption at $T \ge 423$ K.

Sometimes, the adsorption of 2,6-DMP on oxidic systems has been reported to give rise to a band located at \sim 1618 cm^{-1} , 7,8 and this absorption seems to be never present in zeolitic systems. $^{6.9}$ The assignment of this peak is still a controversial matter, $^{7-12}$ as some authors assigned it to a very strong Lewis-coordinated species, whereas other authors ascribed it to a very weak Brønsted (protonic) interaction.

The aim of this work is 2-fold: (i) to prove that 2,6-DMP is a valid probe for Lewis surface acidity, as well as other basic molecules, 11,12 even if steric hindrance yields weaker Lewis complexes than in the case of other basic adsorbates; (ii) to assign the controversial band, located at \sim 1618 cm⁻¹, on the basis of 2,6-DMP adsorption/desorption experiments carried out onto different oxidic systems and in different hydration/dehydration conditions.

Experimental Section

Materials. As a SiO2 sample, we used Aerosil 200, which is a commercial product supplied by Degussa (Frankfurt aM, Germany). The material is amorphous and nonporous, and its BET surface area is 200 m² g⁻¹.

A sample of pure γ -Al₂O₃ was prepared by calcination at 773 K of pure bohemite ($\dot{\gamma}$ -AlOOH), prepared and supplied by Centro Ricerche Fiat (Orbassano, Italy). Starting from this γ -alumina, a subsequent calcination at either 1273 or 1473 K allowed us to obtain either the δ , θ -Al₂O₃ modification or the α -Al₂O₃ modification, respectively. The phase composition of the three alumina samples was checked by the XRD technique, and the diffractograms were found to be consistent with those reported in the literature (e.g., see ref 14). The mesoporous nature of the two transition aluminas and the nonporous nature of the corundum alumina phase were all consistent with those first described by Lippens et al. 14 The BET surface area was found to be 200 m 2 $g^{-1}(\gamma-Al_2O_3)$, 120 m² g⁻¹ ($\delta,\theta-Al_2O_3$), and 4 m² g⁻¹ ($\alpha-Al_2O_3$).

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A sample of sulfated alumina, hereafter referred to as γ -AS, was prepared by sulfating with 0.1 N H₂SO₄ the above-mentioned γ-Al₂O₃ preparation, so as to obtain a surface sulfates concentration of \sim 3 SO₄²⁻/nm². Both crystal phase and BET surface area of the γ -Al₂O₃ sample were found not to be significantly influenced by the sulfation process.

A sample of β -zeolite in its H-form (hereafter referred to as H- β) was kindly supplied by Dr. S. Bordiga (University of Turin, Turin, Italy). The material, usually employed for the alkylation of aromatics, was of industrial provenance 15 and was characterized by a Si/Al ratio of 14.5.

Techniques. Fourier transform infrared (FTIR) spectra were obtained at 2 cm⁻¹ resolution with a Bruker IFS113v spectrophotometer connected to a glass vacuum line (residual pressure <10⁻⁵ Torr) that allows to perform strictly in situ adsorption/ desorption runs. The samples were examined in the form of either self-supporting pellets ($\sim 15~\text{mg cm}^{-2}$) or thin layer depositions $(\sim 10 \text{ mg cm}^{-2})$ on Si wafers.

Prior to adsorption/desorption experiments, all samples were activated in situ at the selected temperatures in homemade quartz IR cell (equipped with KBr windows) connected to the vacuum

2,6-DMP adsorption/desorption tests were carried out as follows: (i) A relatively large amount of base (~4 Torr) was first allowed on the samples and left in contact at beam temperature (BT; \sim 333–343 K) for 2 min. (ii) The excess 2,6-DMP was then evacuated at BT for increasing times in the 1-15 min range. (iii) A desorption of the strongly bonded base fraction was eventually carried out at 423 K.

Results and Discussion

2,6-DMP Adsorption onto SiO₂. Silica was activated in vacuo at two different temperatures (473 and 1073 K), to obtain different surface dehydration levels. After vacuum activation at 473 K, no undissociated adsorbed water is present anymore, but the surface of Aerosil 200 is still largely hydrated, as the dehydroxylation process of nonporous silicas is completed only for much higher activation temperatures. In fact, the 3900-2500 cm⁻¹ spectral range (the v_{OH} spectral range; see curve 1 in Figure 1a) still contains at least two contributions: (i) at high frequency (\sim 3745 cm⁻¹), the well-known $\nu_{\rm OH}$ stretching mode of terminal silanols free from H-bonding interactions; (ii) at lower frequency (3650-3100 cm⁻¹), a broad and unresolved envelope due to the v_{OH} stretching modes of OH groups still interacting by H-bonding. When the activation temperature is as high as \sim 1073 K, the broad tailing at low ν totally disappears, and the high- ν spectral range (see curve 2 in Figure 1a) presents only a sharp peak located at ~3750 cm⁻¹, due to residual free OH groups.

After adsorption of 2,6-DMP onto silica activated at 423 K, in the ν_{OH} spectral range (3900–2500 cm⁻¹; see curve 1' in Figure 1a) several spectral changes are observed: (i) The $v_{\rm OH}$ stretching mode of free silanols (~3745 cm⁻¹) disappears. (ii) The broad envelope at $\sim 3650-3100$ cm⁻¹, ascribed to the H-bonding interactions among surface OH groups, shows a decreased intensity. (iii) A strong and broad band envelope forms at $\nu < 3200$ cm⁻¹. All these spectral features clearly indicate that 2,6-DMP uptake involved in H-bonding of medium strength ($\Delta \nu_{\rm OH} \approx -700$ cm⁻¹) all of the free surface OH species and an appreciable fraction of the surface OH species still involved in H-bonding.

Upon evacuation at BT (spectrum 1" of Figure 1a), the perturbation caused in the $\nu_{\rm OH}$ spectrum by 2,6-DMP

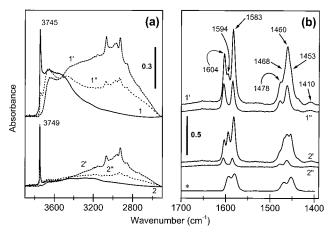
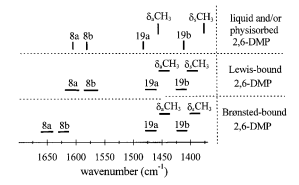


Figure 1. (a) OH spectral pattern (in the 3900-2500 cm⁻¹ range) of an Aerosil-200 sample activated in vacuo at 473 K (spectral set 1) and at 1073 K (spectral set 2). (b) IR spectra in the analytical range 1700-1400 cm⁻¹ (8a, 8b, 19a, 19b, and δ_{CH_3} modes) of 2,6-DMP adsorbed onto an Aerosil-200 sample activated in vacuo at 473 K (spectral set 1) and at 1073 K (spectral set 2). In both sections, the primed numbers refer to the adsorption of a large dose of 2,6-DMP (~4 Torr) and the double-primed numbers refer to the evacuation of 2,6-DMP at beam temperature (BT) for 15 min. The reference spectrum, marked with an asterisk, corresponds to liquid 2,6-DMP.

Chart 1. Spectral Location of Mid-IR Vibrational Modes of Free and Adsorbed 2,6-DMP



adsorption is only partly eliminated, whereas after evacuation at 423 K the original ν_{OH} spectrum is fully recovered.

As for the analytical mid-IR spectral range (1700–1400 cm⁻¹), 2,6-DMP species usually present two groups of bands. Figure 1b shows a series of spectra obtained after adsorption/desorption of 2,6-DMP onto the differently activated silica specimens. The spectral segments considered in the following comprise both the analytical range of ring stretching modes 19a, 19b and 8a, 8b and that of the two δ_{CH_3} deformation modes, as reported in Chart 1. (a) When the surface of SiO₂ is only partially dehydrated, in the presence of 2,6-DMP vapors many bands form: among them, the bands ascribable to a liquid-like (physisorbed) 2,6-DMP species are observed, lying at \sim 1594, \sim 1583, \sim 1468 (shoulder), \sim 1453 (shoulder), and \sim 1410 cm⁻¹ (very weak); the first four bands correspond to the only bands observable in the reference spectrum (marked with an asterisk in Figure 1b) relative to liquid 2,6-DMP. It is worthwhile noting that the peaks located at lower frequencies (~1400-1480 cm⁻¹) are little useful in analytical terms, because they undergo little spectral (upward) changes upon adsorption. On the other hand, the second group of bands (~1580-1605 cm⁻¹) gives more information: the strong band at 1583 cm⁻¹ (the 8b mode) is due to both physisorbed (liquid-like) species and H-bonding

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H-bonded 2.6-DMP Lewis-bound 2,6-DMP Brønsted-bound 2.6-DMP

interacting 2,6-DMP molecules (Chart 2). The \sim 1594 cm⁻¹ band is due to the 8a mode of physisorbed molecules, whereas the peak located at 1604 cm⁻¹ can be assigned to the 8a mode of a stronger form of adsorbed 2,6-DMP, i.e., to 2,6-DMP interacting by H-bonding with surface silanols.

After evacuation at BT (see curve 1" in Figure 1b), bands remain, though with a drastically decreased intensity, at \sim 1604, \sim 1478, \sim 1460, and \sim 1410 cm $^{-1}$ (very weak), together with a residual component at 1580 cm⁻¹, due to the 8b mode common to all adsorbed 2,6-DMP species. On the basis of the spectral behavior, these bands can be confirmed to be ascribable to the 8a, 8b, 19a, 19b, and δ_{CH_3} modes of 2,6-DMP interacting by H-bonding with surface OH groups still abundantly present on the surface. In particular, the large upward shift undergone by the 8a mode of H-bonded 2,6-DMP (~1604 cm⁻¹ is a frequency often observed for 2,6-DMP coordinated to Lewis acid sites) is not unexpected, as, also with other molecules (e.g., acetonitrile¹⁶), surface silanols were observed to yield, by H-bonding, large spectral shifts comparable to the shiffs normally observed with Lewis acid-base interactions of medium strength.

(b) When silica is activated at high temperatures (\sim 1073 K; see curve 2' in Figure 1b), in the 1650-1400 cm⁻¹ spectral range no new bands are added to the abovementioned ones. The only difference is that the overall 2,6-DMP uptake is somewhat decreased, and there is a largely enhanced ratio between physisorbed and H-bonded species, as a consequence of a far decreased surface concentration of OH species capable of H-bonding the adsorbate. In the ν_{OH} spectral range, the spectral changes $(\Delta \nu_{OH} \approx -700~\text{cm}^{-1})$ confirm the profile already evidenced for the sample activated at lower T (see curves 2 and 2' in Figure 1a). Upon evacuation at BT, a much larger fraction of H-bonded 2,6-DMP is desorbed (with respect to the sample activated at low temperature), and when 2,6-DMP is desorbed at 423 K, all the spectral features due to adsorbed species disappear and the background spectrum recovers the starting shape in all spectral ranges examined.

2,6-DMP Adsorption onto Aluminas (γ -, δ , θ -, and **α-Phases).** (a) γ -Alumina. γ -Alumina was chosen as a test adsorbent because of its large use in catalytic applications and for its strong Lewis acidity (see, for instance, ref 17 and references therein). Alumina was vacuum activated at three increasing temperatures (473, 773, and 1023 K, respectively), to achieve a progressive surface dehydration, and thus create at the surface larger and larger concentrations of cus cationic sites, acting as surface Lewis acid sites.

When the sample is activated at 473 K, it is freed from all undissociated (coordinated) water, but it is only very

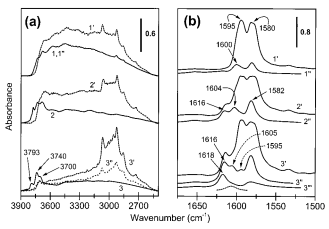


Figure 2. (a) OH spectral pattern (in the 3900-2500 cm⁻¹ range) of a γ -alumina sample activated in vacuo at 473, 773, and 1023 K (spectral sets 1–3, respectively). (b) Spectral features in the analytical range of the 8a, 8b modes (1700-1500 cm $^{-1}$) of 2,6-DMP adsorbed onto γ -alumina activated in vacuo at 473, 773, and 1023 K (spectral sets 1-3, respectively). In both sections, the curves marked with the prime symbol refer to 2,6-DMP adsorption at BT and under a relatively high pressure (~4 Torr), whereas the curves marked with the doubleand triple-primed symbols refer to 2,6-DMP desorption at BT and at 423 K, respectively.

slightly dehydroxylated (if at all). This is clearly observable in the v_{OH} stretching region (see Figure 2a, curve 1), where a broad and unresolved band envelope, due to H-bonded OH species, is still present at $\nu < 3700 \text{ cm}^{-1}$. Only two shallow peaks lying at high frequency are ascribable to OH groups already free from H-bonding. Activation at higher temperatures (see curves 2 and 3 in Figure 2a) produces a more and more dehydrated surface, and the ν_{OH} spectral pattern acquires a well-known five-band profile, 17-19 whose shape and overall intensity is largely determined by the activation temperature reached.

Figure 2b shows the IR spectra in the 8a, 8b spectral range of 2,6-DMP adsorbed onto γ -alumina activated at the three different temperatures mentioned above. Only the higher-v subrange, due to 8a, 8b modes, is here examined, because the presence on alumina of several 2.6-DMP adsorbed species renders the band's envelope at $\nu < 1550 \ {\rm cm^{-1}}$ broad and very poorly resolved, so that its analytical use is virtually null.

When the activation is carried out at the lowest temperature (see the spectral set 1 in Figure 2b), a twoband profile is observed centered at ~ 1595 and ~ 1580 cm⁻¹, mostly due to a liquid-like 2,6-DMP species. But the broad and asymmetric band at \sim 1595 cm⁻¹ is most likely made up of several contributions: (i) One is certainly due to a fast reversible weak physisorbed species (i.e., the mentioned liquid-like species). (ii) A second nonresolved component could be due to 2,6-DMP interacting by H-bonding with surface OH species, but a specific 8a mode could not be singled out. Still, the dramatic spectral changes observed in the ν_{OH} spectral region (see Figure 2a) indicate that surface OH species are indeed perturbed by H-bonding upon 2,6-DMP adsorption. (iii) Another component could be ascribable to 2,6-DMP molecules interacting with weak Lewis acid centers. In fact, when the excess 2,6-DMP is outgassed at BT, the global intensity of the 8a, 8b mode envelope decreases dramatically, while the v_{OH} profile is fully recovered, and curve 1" in Figure

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2b indicates that an 8a peak remains, centered at $\sim\!1600$ cm $^{-1}$, together with a residual 8b component at 1581 cm $^{-1}$. It is recalled that, upon activation at 473 K, alumina is freed from coordinated undissociated water, and this ligand removal leaves behind the first (weak) Lewis sites, as it has been evaluated by the adsorption of CO₂ (at room temperature) or of CO at low temperature. The band at $\sim\!1600$ cm $^{-1}$ is thus assignable to the 8a mode of (a first family of) weak Lewis acid sites.

A more complex situation is observed when γ -alumina is vacuum activated at 773 and 1023 K (see, in Figure 2b, the spectral sets 2 and 3, respectively). The dehydration process has proceeded, and in both cases the adsorption of 2,6-DMP produces, in the 1700-1500 cm⁻¹ range, a complex band envelope made up of, at least, three broad components (centered at \sim 1575, \sim 1590, and \sim 1610 cm⁻¹, respectively). If the excess 2,6-DMP is pumped off, the labile (physisorbed and H-bonded) species are removed and in the analytical spectral region three peaks (activation at 773 K, curve 2" of Figure 2b) and four peaks (activation at 1023 K, curve 3" of Figure 2b) remain. The first one, located at ~ 1582 cm⁻¹, is assigned to the ovelapping 8b modes of all 2,6-DMP species, whose 8a modes lie at higher wavenumbers.6 As in the case of pyridine adsorption, also for 2,6-DMP the rule is that the higher is the 8a mode of the coordinated base, the stronger is the interaction between the coordinated base and the adsorption site. For this reason, the proposed assignment for the (up to) three 8a mode bands observed in Figure 2b is as follows:

(i) The band located at the highest ν (\sim 1616 cm $^{-1}$), and thus corresponding to 2,6-DMP complexes formed at sites characterized by the highest Lewis acidity, can be ascribed to 2,6-DMP molecules interacting with cus Al^{3+} cations possessing an incomplete tetrahedral coordination. It is known that these sites are the most important component of the strong Lewis acidity for which transition aluminas are appreciated for many catalytic applications. (It is recalled that, on the highly acidic low-temperature transition aluminas (γ - and η -Al₂O₃), py yields with cus Al^{IV} sites strong complexes whose 8a mode is observed at frequencies as high as \sim 1622 cm $^{-1}$.)

(ii) As for the 8a band located at $\sim\!1603-1606~cm^{-1},$ it can be ascribed to 2,6-DMP complexes formed at coordinative vacancies bridging between Al^{IV} and Al^{VI} surface cationic centers. (It is recalled that, on low-temperature transition aluminas ($\gamma\!$ - and $\gamma\!$ -Al $_2O_3$), py yields with these coordinative vacancies complexes whose 8a mode lies at $\sim\!1614~cm^{-1}$.)

(iii) Finally, the 8a band located at $\sim 1595~\text{cm}^1$ (some of which remains on γ-Al₂O₃ treated at 1073 K also after evacuation at BT) may have a double nature, as in this spectral position two different contributions are possible. One contribution can derive from the H-bonding interaction of 2,6-DMP with some surface OH species, whereas the other one can be due to the interaction of 2,6-DMP with Lewis acid sites weaker than the ones already mentioned, i.e. with coordinative vacancies on octahedrally coordinated (AlVI) sites. Both of these assignments imply some surprising aspects: it is surprising that some H-bonded 2,6-DMP species remain, after evacuation at BT, only on the sample activated at the highest temperature (on which all five different OH species are still present, but there should be the lowest overall amount of OH species), and it is surprising that (some of) the base Lewis coordinated to the weakest acid sites remains after evacuation at BT only on the sample activated at the highest temperature, on which the surface concentration of strong Lewis acid sites is maximum. In fact, on

insulating systems, most frequently the coordination to weak charge-withdrawing sites is observable only in the absence of complexes formed at stronger charge-withdrawing sites. Still, we favor the latter assignment, as on all other systems examined (but silica) the elimination of H-bonded 2,6-DMP is observed to be complete upon evacuation at BT.

After a vacuum desorption treatment at 423 K (see, for instance, the spectral set 3 of Figure 2b, relative to $\gamma\text{-}Al_2O_3$ activated at 1073 K), the overall bands intensity decreases but is not eliminated. The high- ν 8a mode (now centered at $\sim\!1618~\text{cm}^{-1}$) remains with almost unchanged intensity, whereas only some 15% of the 8a mode at $\sim\!1606~\text{cm}^{-1}$ remains, as indicated in the figure by the dotted-lineresolved component. (Obviously, also a contribution of the unresolved 8b mode remains at $\sim\!1584~\text{cm}^{-1}$.) This indicates that an adsorption/desorption temperature of $\sim\!423$ K is not sufficient to prevent/eliminate the coordination of 2,6-DMP onto/from strong Lewis sites of alumina, i.e., on coordinative vacancies involving Al^IV centers.

The data for 2,6-DMP adsorption/desorption on γ -alumina so indicate that, unlike what previously reported, ¹³ at temperatures as high as \sim 423 K this base still probes selectively the strongest Lewis acid sites and that, unlike what reported by other authors, ¹⁰ this base can distinguish Lewis acid sites of different strength, much as other bases do. Moreover, the present data indicate that, at least on γ -Al₂O₃, the controversial 8a band at \sim 1618 cm⁻¹ is ascribable to a strong Lewis interaction (actually, the strongest one) and not to a weak Brønsted interaction that, on pure alumina, is not present at all.

(b) δ, θ - and α -Aluminas. To confirm the above hypothesis and spectral assignments, we performed some adsorption/desorption experiments of 2,6-DMP onto δ , θ - and α-alumina specimens activated at 1073 K, i.e., in highly dehydrated conditions. The interest in these alumina modifications consists of the following facts: (i) δ , θ -Al₂O₃ (one of the so-called high-temperature transition aluminas) still possesses a defective spinel structure but should possess a lower surface concentration of highly defective Al³⁺ centers with an incomplete tetrahedral coordination.¹⁷ This system is thus suitable for confirming the ability of 2,6-DMP to differentiate among Lewis sites of different strength. (It is recalled that, using a δ , θ -Al₂O₃ modification, other authors¹³ observed only one Lewis coordinated species and, thus, concluded that 2,6-DMP is not a suitable probe for the Lewis acidity of metal oxides.) (ii) α -Al₂O₃ has a different crystal structure (corundum), and at least in the bulk (i.e., in regular conditions), should possess Al³⁺ ions only in the octahedral coordination. Also this system is thus suitable for confirming the ability of 2,6-DMP to differentiate among Lewis sites of different strength.

(i) The adsorption/desorption of 2,6-DMP onto a highly dehydrated δ,θ -Al₂O₃ system (see spectral set 1 in Figure 3) brings about the formation of the same 8a, 8b bands described above for the γ -alumina specimen vacuum activated at the same high temperature, but the intensity ratios are here quite different. In particular, if we compare the spectra obtained on γ -Al₂O₃ and on δ,θ -Al₂O₃ after 2,6-DMP evacuation at BT (curve 3" in Figure 2b and curve 1" in Figure 3, respectively), it is quite evident that the amount of irreversible adsorbate is small in both cases (as witnessed by the weak 8b mode remaining at \sim 1580 cm $^{-1}$ after evacuation), but on the δ,θ -Al₂O₃ system there is an appreciably decreased amount of the strongest species (\sim 1616 cm $^{-1}$, assigned to cus tetrahedral Al sites) and an increased relative amount of the other two species (\sim 1604 cm $^{-1}$, assigned to coordinative vacancies bridging between

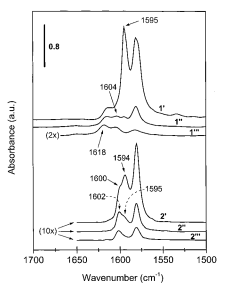


Figure 3. Analytical spectral range of the 8a, 8b modes (1700–1500 cm $^{-1}$) of 2,6-DMP adsorbed onto different alumina samples activated in vacuo at 1023 K (set 1, δ , θ -alumina; set 2, α -alumina). The curves marked with the prime symbol refer to 2,6-DMP adsorption at BT and under a relatively high pressure (\sim 4 Torr), whereas the curves marked with the double- and triple-primed symbols refer to 2,6-DMP desorption at BT and at 423 K, respectively. For the sake of clarity, some of the spectra were ordinate-magnified, as reported on the curves.

tetrahedral and octahedral Al^{3+} ions, and $\sim 1594~\text{cm}^{-1}$, (tentatively) assigned to cus Al3+ ions with octahedral coordination). Also after 2,6-DMP evacuation at 423 K (curve 3" in Figure 2b and curve 1" in Figure 3), the residual 8a, 8b bands confirm that, at this temperature, the strongest Lewis acid sites can still adsorb the base, and the relative intensity of the two 8a modes (now at \sim 1618 and \sim 1604 cm $^{-1}$, respectively) confirms that, on δ , θ -Al₂O₃, the surface concentration of strongly defective Al^{IV} cus sites is lower than in the case of γ -Al₂O₃. The fact that, on a similar δ , θ -Al₂O₃ system, other authors¹³ did not observe the multiplicity of species here observed is (at least partly) due to the fact that, in that case, the activation was carried out at a much lower temperature, so that most of the more energetic Lewis acid sites were still missing (i.e., the surface of the transition alumina was still in a highly hydrated form).

(ii) The adsorption/desorption of 2,6-DMP onto an α-Al₂O₃ system activated at 1073 K (at this temperature, α -Al₂O₃ is virtually fully dehydrated) leads to the formation of a lower number of spectral components (see set 2 in Figure 3). This is better seen in the spectral pattern 2", relative to the BT evacuation of the excess 2,6-DMP and removal of all weakly adsorbed species. Besides the band located at $\sim 1580~\text{cm}^{-1}$ (the 8b mode of all 2,6-DMP adsorbed species), the 8a spectral range presents two components of odd intensity: one (stronger) component is centered at ~ 1602 cm⁻¹, and a second one is present as an evident shoulder at ~ 1595 cm⁻¹. On the basis of the spectral position, the former band can be ascribed to the interaction of 2,6-DMP with quasi-tetrahedral Al³⁺ sites, sharing a coordination vacancy with an octahedral Al3+ ion, and the latter band to the interaction of 2,6-DMP with purely octahedral Al^{3+} cus sites. (In the same spectral position there might be also a contribution from H-bonded 2,6-DMP species, but this possibility is low in that, as reported above, α-Al₂O₃ activated at 1023 K is virtually fully dehydroxylated.) The band due to 2,6-DMP complexes formed at cus octahedral sites should, on α -Al₂O₃, be the sole coordinated species formed, as the corundum structure

implies for cationic species the sole octahedral coordination. As a matter of fact, the 8a band at $\sim\!1595~cm^{-1}$ was definitely much stronger under a high 2,6-DMP pressure (see curve 2') and was then drastically reduced during the BT evacuation, but it was not the only Lewis-coordinated species present. The presence at the surface of the corundum $\alpha\text{-Al}_2O_3$ phase of defective quasi-tetrahedral Al^{3+} sites is not unexpected and was already postulated some years ago by the use of other adsorbates. But is certainly interesting to note that also this peculiar aspect of Lewis acidity at the surface of a nominally monophasic oxidic system can be easily evidenced by the adsorption/desorption of 2,6-DMP.

The high acidic strength of the sites responsible for the 8a band at ${\sim}1602~cm^{-1}$ on ${\alpha}\text{-}Al_2O_3$ is further confirmed by the persistence of most of the band (now absorbing at ${\sim}1605~cm^{-1})$ after 2,6-DMP evacuation at 423 K (see curve 2″ in Figure 3).

2,6-DMP Adsorption onto Sulfated γ -Al₂O₃. The presence of sulfate groups at the surface of oxidic systems is known to modify their intrinsic Lewis acidic properties (e.g., see ref 21). In general terms, the overall Lewis acidity should decrease (as part of the surface is occupied by thermally stable sulfate groups and, thus, a lower amount of cus surface cations can be produced upon activation), and the residual Lewis acidity has been sometimes observed to increase in strength (due to inductive effects from the charge-withdrawing sulfate groups). Moreover, surface sulfates most often induce Brønsted (protonic) acidity, as indicated by the adsorption of strong bases like, for instance, ammonia or pyridine, which can form surface ammonium or pyridinium ions, respectively. 22,23

Surface sulfate species on oxidic systems are characterized by typical vibrational modes located at $\nu < 1500$ cm⁻¹.²³ The spectral features of these modes depend on the hydration/dehydration degree achieved by the sample, and in this respect, γ -AS makes no exception. On passage from a highly hydrated system (curve 1 in Figure 4a) to a more dehydrated one (curve 2 in Figure 4a), the spectral profile of surface sulfates evolve from a typical narrow three-band spectrum (ionic sulfates, similar to mono- and bidentate sulfato complexes²⁴) to a well-separated twoband spectrum (typical of highly covalent multibridged sulfates²⁵), with a high- ν component at \sim 1380 cm⁻¹ (the S=O stretching) and a complex low- ν envelope at ν < 1200 cm^{-1} (the S-O stretching range). The dotted-line traces in Figure 4a represent the spectral profile of the corresponding pure γ -alumina, vacuum activated in the same conditions.

If we carry out 2,6-DMP adsorption onto γ -AS activated in vacuo at 573 K (see Figure 4b, curve 2'), several bands forms in the 1700–1500 cm⁻¹ spectral range, as detailed in the following. There is a broad three-band profile at $\nu < 1610 \text{ cm}^{-1}$, with peaks evident at ~ 1602 , ~ 1595 , and $\sim 1580 \text{ cm}^{-1}$, and an additional two-band envelope located at $\nu > 1620 \text{ cm}^{-1}$, with broad peaks evident at $\sim 1650 \text{ and} \sim 1630 \text{ cm}^{-1}$. On the basis of existing literature data on the adsorption of 2,6-DMP, the latter bands can be ascribed to the 8a, 8b vibrational modes ($\sim 1650 \text{ and} \sim 1630 \text{ cm}^{-1}$,

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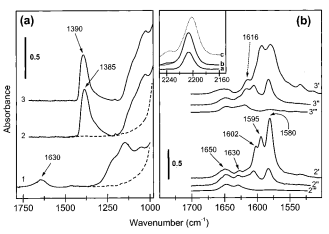


Figure 4. (a) IR spectra in the 1800–900 cm⁻¹ region ($\nu_{S=0}$ and v_{S-O} modes of surface sulfates) of a γ -AS sample activated in vacuo for 1 h at BT (curve 1), 573 K (curve 2), and 773 K (curve 3). The dashed traces represent the spectral profile of the corresponding unsulfated $\hat{\gamma}$ -Al₂O₃ samples. (b) Spectral features in the analytical range of the 8a, 8b modes (1700-1500 cm $^{-1}$) of 2,6-DMP adsorbed onto γ -AS activated in vacuo at 573 K (spectral set 2) and at 773 (spectral set 3). The curves marked with the prime symbol refer to 2,6-DMP adsorption at BT and under a relatively high pressure (~4 Torr), whereas the curves marked with the double- and triple-primed symbols refer to 2,6-DMP desorption at BT and at 423 K, respectively. The inset presents, for comparison, the spectra of 100 Torr CO adsorbed at BT onto γ -AS activated in vacuo at 573 K (curve a) and 773 K (curve b) and onto a plain γ -Al₂O₃ sample activated in vacuo at 773 K (curve c).

respectively) of protonated 2,6-DMP species (2,6-DMPH⁺). Protonated species are typically formed in systems carrying a Brønsted acidity, and their presence indicates that, as in many other oxidic systems (e.g., see ref 26), also on $\gamma\text{-Al}_2O_3$ the presence of strongly acidic surface anions induced the formation of a protonic acidity, which is promptly revealed by the adsorption of 2,6-DMP.

After evacuation of the excess 2,6-DMP at BT (curve 2" of Figure 4b), the spectral profile remains unchanged at $\nu > 1620 \, \text{cm}^{-1}$ (meaning that the Brønsted-bound species are strongly adsorbed), whereas at $\nu < 1620 \text{ cm}^{-1}$ only one 8a band remains at ~ 1604 cm⁻¹, together with a residual fraction of the 8b band at 1580 cm⁻¹. This suggests that the process of sulfation may have brought about the selective screening (by sulfate groups) of the cationic surface terminations that, upon dehydration, lead to the formation of the strongest Lewis acid sites (whose complexes with 2,6-DMP yield the 8a band at ~1616-1618 cm⁻¹). But certainly not all of the Al^{IV} surface cationic terminations were screened by sulfates, as the mediumstrength Lewis acid sites, ascribed to bridging coordinative vacancies, still remain and yield the 8a band at ~1604 cm^{-1} .

If the desorption of 2,6-DMP is carried out at 423 K (see curve 2''' in Figure 4b), only the two bands at $\nu > 1620$ cm $^{-1}$ remain with virtually unchanged intensity (the lutidinium surface species are so deduced to be very strongly held), whereas no spectral components remain at lower frequency, meaning that all Lewis-coordinated 2,6-DMP species are desorbed during this mild thermal treatment.

To check if the sulfation process eliminated entirely the cationic sites capable of yielding the strongest Lewis acid sites, the sample γ -AS was activated at a higher temperature (773 K). The spectral profile 3 in Figure 4a

shows that, at this temperature, the concentration and nature of surface sulfates did not change (in fact, surface sulfates are stable up to ~ 900 K), while the surface dehydration proceeded (not shown). The spectral set 3 in Figure 4b indicates that, further to activation at high temperature, an appreciable amount of the Lewis-bound 2,6-DMP species absorbing at \sim 1616 cm⁻¹ formed, although with an intensity definitely lower than that exhibited in the case of a plain γ -Al₂O₃ sample treated in similar conditions (dealt with in Figure 2). This trend is confirmed by the adsorption of CO at BT and presented for comparison in the inset to Figure 4b: the strongest Lewis complex of CO with cus Al^{IV} sites, which absorbs on γ - Al_2O_3 at \sim 2240 cm⁻¹, 17 does not form on γ -AS activated at 573 K (spectrum a) and forms weakly on γ -AS activated at 773 K (spectrum b), but the intensity is far lower than that produced on γ -Al₂O₃ activated at the same temperature (broken-line spectrum c).

The 8a mode of the strongest Lewis 2,6-DMP complex, absorbing at $1616-1618~cm^{-1}$, remains with a somewhat decreased intensity also after evacuation at 423 K (see curve 3'" of Figure 4b). In all stages of the adsorption/desorption process, the $\sim 1618~cm^{-1}$ band overlaps severely the low- ν mode (centered at $\sim 1630~cm^{-1}$) of the Brønstedbound 2,6-DMP species. Still, the evolution of the spectral profile in set 3 of Figure 4b leaves no doubts on the fact that the $\sim 1618~cm^{-1}$ band is due to a strong Lewiscoordinated 2,6-DMP species and not to a weak Brønstedbound 2,6-DMP species. Moreover, these data confirm that, also in the presence of strong Brønsted acid centers, 2,6-DMP adsorption/desorption at temperatures up to 423 K can sample at least the strongest fraction of Lewis acidic sites.

2,6-DMP Adsorption onto β **-Zeolite.** The last system to be examined by the adsorption/desorption of 2,6-DMP is the sample termed H- β , i.e. a sample of β -zeolite in its H-form (all cations have been exchanged for H ions). This system is of interest in the present study because of the following: (i) It possesses an abundant natural protonic acidity, deriving from the presence of abundant bridging OH species (typical of zeolitic systems). (ii) It is characterized by large channels (5.7 \times 7.5 Å), so that 2,6-DMP molecules should not be prevented from coming into contact with sites located in all possible positions of the zeolitic structure. (iii) No cations are left within the zeolitic structure. (iv) When treated in vacuo at medium-high temperatures, some of the framework Al atoms (that in zeolites possess the tetrahedral coordination) may dehydroxylate and, thus, originate coordinative vacancies acting as Lewis acid centers. The nature of the latter can now be suitably investigated by the adsorption of 2,6-DMP, in the light of what already observed with the various alumina systems dealt with in previous sections.

Section a of Figure 5 reports the 3900-2600 cm⁻¹ spectral range of H- β activated at 873 K. Before adsorption, the complex OH spectral pattern (curve 1) contains several components, among which are clearly evident a strong sharp peak at 3747 cm⁻¹, due to abundant terminal silanols, and a broad peak at \sim 3610 cm⁻¹, due to bridging OH species. A minor component at ~3670 cm⁻¹ is also present and is normally ascribed to OH groups bound to quasi-extraframework Al³⁺ ions. ¹⁵ Upon 2,6-DMP uptake (curve 1'), the two major OH components become perturbed by H-bonding ($\Delta v_{\rm OH} \approx -700 \text{ to } -800 \text{ cm}^{-1}$), and a very complex pattern appears in the 3300-2650 cm⁻¹ region, mainly due to the ν_{C-H} vibrations of the various 2,6-DMP adsorbed species. (Note that this pattern is far more complex than that observed in the case of 2,6-DMP adsorbed on SiO₂ (Figure 1), as in the latter case only

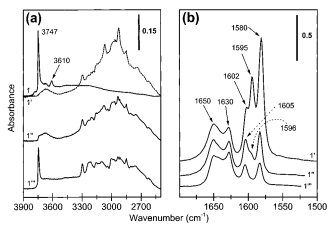


Figure 5. (a) 3900–2500 cm⁻¹ spectral range (analytical of $\nu_{\rm OH}$ and $\nu_{\rm CH}$ modes) of a β -H zeolite activated in vacuo at 873 K (curve 1), after the adsorption of a high pressure of 2,6-DMP (\sim 4 Torr; curve 1') and after 2,6-DMP evacuation at BT (curve 1") and at 423 K (curve 1""). (b) Spectral features in the analytical range of the 8a, 8b modes (1700–1500 cm $^{-1}$) of 2,6-DMP adsorbed onto β -H activated in vacuo at 873 K. The curve marked with the prime symbol refers to 2,6-DMP adsorption at BT and under a relatively high pressure (~4 Torr), whereas the curves marked with the double- and triple-primed symbol refer to 2,6-DMP desorption at BT and at 423 K, respectively.

liquid-like and H-bonded 2,6-DMP species were formed, whereas here at least one additional Brønsted-bound species is expected to be present.) Upon 2,6-DMP evacuation at BT (curve 1"), the starting OH pattern is only very marginally restored, and this is not so unexpected as also with pure SiO₂ it was observed (see Figure 1) that the H-bonding interaction between 2,6-DMP and silanols is only partially reversible at beam temperature. 2,6-DMP evacuation at 423 K (curve 1" of Figure 5a) gives back only \sim 2/3 of the silanol band at 3747 cm⁻¹ (meaning that other forms of silanol perturbation must be present besides the plain $(N \cdot \cdot \cdot \cdot H - O)$ H-bonding) and none of the bridging OH band at \sim 3610 cm⁻¹.

Section b of Figure 5 reports the effects of 2,6-DMP adsorption/desorption in the spectral range of the 8a, 8b modes. There is a very strong doublet of bands at \sim 1650 and $\sim 1630~\text{cm}^{-1}$, due to the abundant formation of lutidinium species. The intensity of these bands does not change further to 2,6-DMP evacuation at up to 423 K and so explains why the band of bridging OH groups at \sim 3610 cm⁻¹ (responsible for Brønsted acidity) is not restored at all upon evacuation.

In the spectral range below 1620 cm⁻¹, the adsorption of 2,6-DMP brings about the formation of a strong threeband envelope that, on the basis of what observed in the case of the 2,6-DMP/SiO₂ system, could be assigned as follows: The band at $\sim 1580 \, \text{cm}^{-1}$ is the 8b mode, common to all physisorbed and coordinated species, the band at \sim 1595 cm⁻¹ is the 8a mode of physisorbed (liquid-like) 2,6-DMP, and the band at \sim 1602 cm⁻¹ is the 8a mode of 2,6-DMP H-bonded to surface silanols. This assignment, though certainly correct, is not sufficient, and other adsorbed species must be present and contribute to the 8a, 8b envelope. In fact, the following can be stated: (i) After evacuation at BT (see curve 1" of Figure 5b), a weak but clearly visible component remains at $\sim 1596~{\rm cm}^{-1}$ (besides a dominant band at ${\sim}1605~\text{cm}^{-1}\text{)}$ that could be ascribed to a fraction of physisorbed 2,6-DMP resisting evacuation at \sim 340 K, as suggested by one of the reviewers of this article. On the basis of what is discussed in the sections dedicated to the 2,6-DMP/Alumina systems, it seems more logical to ascribe the residual 8a band at \sim 1596

cm⁻¹ in curve 1" (and thus part of the strong component at \sim 1595 cm⁻¹ in curve 1') to the coordination of 2,6-DMP to Al ions carrying an incomplete octahedral coordination (Al^{VI}_{CUS}). (ii) After evacuation at 423 K (see curve 1" of Figure 5b), an appreciable fraction of an 8a band at \sim 1605 cm⁻¹ remains that cannot be ascribed to a fraction of merely H-bonded 2,6-DMP resisting evacuation at T >BT. On the basis of what is discussed in the sections dedicated to the 2,6-DMP/Alumina systems, it seems logical to ascribe the residual 8a band at $\sim 1605~\text{cm}^{-1}$ in curve 1"' (and thus part of the band centered at \sim 1602 cm⁻¹ in curve 1') to 2,6-DMP coordinated to bridging coordinative vacancies (for instance, AlIV-AlVI pairs, as observed in all transition aluminas and at the surface of α-Al₂O₃). The persistence, after evacuation at 423 K, of appreciable amounts of Lewis-coordinated 2,6-DMP species justifies the incomplete recovery of the silanol band at \sim 3747 cm⁻¹, as it is known that terminal OH species at the surface of oxidic systems often interact by H-bonding with the π cloud of Lewis coordinated aromatic heterocyclic hases

The adsorption of 2,6-DMP on β -H never leads to the appearance of a band at ~1618 cm⁻¹, as also observed with other zeolitic systems. Since, on the basis of what is discussed above, there is no doubt that, when present, the 2,6-DMP band at \sim 1618 cm⁻¹ is due to a strong Lewis coordination, the absence of this band on all zeolitic systems means that, on zeolites, the thermal dehydroxylation of Al centers (either in-framework or extraframework ones) never leads to the formation of the highly uncoordinated Al^{IV}_{CUS} sites, typical of the highly acidic (spinel) transition aluminas.

Conclusions

With reference to the 2-fold aim of the present work proposed in the Introduction section, the following can be concluded:

(i) The assignment of a band sometimes observed at \sim 1618 cm $^{-1}$ upon adsorption of 2,6-DMP on oxidic systems is no longer controversial. The band is due to the 8a ring mode of 2,6-DMP Lewis coordinated to Al^{IV} centers carrying a coordinative vacancy. To the best of our knowledge, this is the highest frequency observed for the 8a mode of a Lewis coordinated 2,6-DMP species, and this fact is not unexpected, as Al^{IV}_{CUS} yields the highest upward spectral shift also with other Lewis bases (e.g., pyridine uptake at BT on Al^{IV}_{CUS} presents the 8a mode at frequencies as high as ~ 1625 cm⁻¹, CO adsorption at BT on Al^{IV}_{CUS} presents the ν_{CO} mode at frequencies as high as \sim 2242 cm⁻¹, and acetonitrile- d_3 adsorption at BT on Al^{IV}_{CUS} presents the ν_{CN} mode at frequencies as high as \sim 2330 cm⁻¹). The very strong Lewis acid sites yielding, on adsorption of 2,6-DMP, the 8a ring mode at \sim 1618 cm⁻¹ are typical of aluminas with a spinel structure (transition aluminas). All other Al-containing oxidic systems do not present an 8a mode of adsorbed 2,6-DMP at $\sim 1618~\text{cm}^{-1}$, even if they may possess coordinative vacancies involving either tetrahedral Al centers (zeolites) or quasi-tetrahedral Al centers (defective α -Al₂O₃).

(ii) The adsorption of 2,6-DMP is a very valuable probe for Brønsted (i.e., protonic) acidic sites. In fact, unlike pyridine (actually, the most commonly used probe) that yields, in its pyridinium form, a weak and broad 19b analytical mode at \sim 1550 cm $^{-1}$, 2,6-DMP adsorbed in its lutidinium form yields two strong and well-recognizable 8a, 8b bands centered at ~ 1650 and ~ 1630 cm⁻¹, respectively. Moreover, surface lutidinium species are very strongly held and remain adsorbed also when all, or most, of all other 2,6-DMP adsorbed species have been desorbed upon evacuation. But, unlike what is reported in some literature works, 2,6-DMP uptake turns out to be a suitable probe also for Lewis acid centers. If the adsorption is carried out at BT and, possibly, under a reasonably high pressure, all types of coordinated and/or H-bonded species can be differentiated, on the basis of the relevant 8a modes. If 2,6-DMP is evacuated (or the adsorption is carried out under very low pressure), the various coordinated species can be differentiated, on the basis of the persistence of the relevant 8a modes. In particular, if the evacuation is carried out at $\sim\!423$ K (or the adsorption is carried out at

 $\sim\!423$ K), only the strongest fraction of the strongest Lewis acid sites can still be probed. But in none of the various systems here examined did we observe (as sometimes reported in the literature) that, at temperatures as high as 423 K, steric hindrance prevents 2,6-DMP molecules from being adsorbed at Lewis acid centers, so that only Brønsted acidity (if present) can be probed.

Acknowledgment. This work was partly financed by the Italian MURST (Rome), Cofin 2000, Sect. 03.

LA010707E