

Fe Promoted NO_x Storage Materials: Structural Properties and NO_x Uptake

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Fe promoted NO_x storage materials were synthesized in the form of FeO_x/BaO/Al₂O₃ ternary oxides with varying BaO (8 and 20 wt %) and Fe (5 and 10 wt %) contents. Synthesized NO_x storage materials were investigated via TEM, EELS, BET, FTIR, TPD, XRD, XPS, and Raman spectroscopy, and the results were compared with the conventional BaO/Al₂O₃ NO_x storage system. Our results suggest that the introduction of Fe in the BaO/Al₂O₃ system leads to the formation of additional NO_x storage sites which store NO_x mostly in the form of bidentate nitrates. NO₂ adsorption experiments at 323 K via FTIR indicate that, particularly in the early stages of the NO_x uptake, the NO_x storage mechanism is significantly altered in the presence of Fe sites where a set of new surface nitrosyl and nitrite groups were detected on the Fe sites and the surface oxidation of nitrites to nitrates is significantly hindered with respect to the BaO/Al₂O₃ system. Evidence for the existence of both Fe³⁺ as well as reduced Fe^{2+/(3-x)+} sites on the freshly pretreated materials was detected via EELS, FTIR, Raman, and XRD experiments. The influence of the Fe sites on the structural properties of the synthesized materials was also studied by performing ex situ annealing protocols within 323–1273 K followed by XRD and Raman experiments where the temperature dependent changes in the morphology and the composition of the surface domains were analyzed in detail. On the basis of the TPD data, it was found that the relative stability of the stored NO_x species is influenced by the morphology of the Ba and Fe containing NO_x-storage domains. The relative stabilities of the investigated NO_x species were found to increase in the following order: N₂O₃/NO⁺ < nitrates on γ-Al₂O₃ < surface nitrates on BaO < bidentate nitrates on FeO_x sites < bulk nitrates on BaO.

1. Introduction

Over the past several years, the global environmental protection efforts resulted in a stringent emission legislation regarding NO_x, CO, and hydrocarbon emissions originating from stationary as well as mobile sources.¹ In recent years, the transportation sector and in particular the diesel-equipped vehicles have become one of the primary sources for NO_x emissions. Therefore, regulations limiting NO_x emissions in industrialized countries have become very stringent. According to the European Union regulations (EURO VI),² by the end of 2014, diesel passenger cars will be subject to a 3-fold decrease in NO_x emissions from 0.25 to 0.08 g km⁻¹.

In the early 1990s, NO_x storage-reduction (NSR) technology, was developed by the Toyota Motor Corporation^{3,4} for NO_x removal from lean-burn gasoline engines. The operational principle is based on the alternating cycles of lean (abundant in oxygen) and rich (abundant in reductants such as unburned hydrocarbons, H₂ and CO) regimes.^{5,6} Numerous detailed experimental and theoretical studies^{4,7–10} investigated some of the fundamental chemical aspects related to the NO_x storage-reduction mechanism occurring on the surface of Pt/BaO/γ-Al₂O₃ NSR catalysts. Recently, detailed surface science studies on the NO_x storage capacity and the thermal aging of model NO_x storage materials in the form of BaO/θ-Al₂O₃/NiAl(100)^{11–13}

have also been reported. The spectroscopic characterization of the alumina support¹⁴ and its interactions with H₂O,¹⁴ NO₂,¹⁵ and H₂O + NO₂¹⁶ in the absence of the active Ba-containing phase were studied as well. More recently, a similar surface science approach was also extended to the BaO/θ-Al₂O₃/NiAl(110) model system.^{17,18}

Numerous studies^{19–21} were directed toward the design of NSR catalysts with long-term durability, high activity, and improved sulfur tolerance by incorporation of additional oxides such as TiO₂ or Fe₂O₃ to the catalytic Pt/BaO/γ-Al₂O₃ system. For instance, in one of our recent reports,²² the nature of the Ti–Ba interactions on the Ba/Ti/Al NO_x storage systems has been thoroughly investigated where it was found that the presence of TiO₂ has a considerable effect on the surface distribution of the NO_x storage domains and the thermal stability of the stored nitrates on the surface. These results²² indicated that TiO₂ domains present a significant affinity toward BaO and/or Ba(NO₃)₂, resulting in a strong Ti–Ba interaction and the formation of overlapping surface domains.

Several studies^{23–25} were performed in the literature in order to understand the influence of iron or iron oxides on the sulfur tolerance of NSR catalysts. It was reported that the addition of Fe improves the long-term stability of the catalysts by assisting the formation of sulfur resistant nitrate species in the Ba/Fe/γ-Al₂O₃ system. In separate studies,^{26,27} it was pointed out that the Fe component on the aged Pt/Ba/Fe/γ-Al₂O₃ catalyst played two important roles: (i) suppressing the growth of sulfate deposit on the catalyst and, thereby, (ii) lowering the desorption temperature of the sulfate domains by inhibiting the BaSO₄

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particle size growth. However, in these studies, minor attention has been devoted to the questions regarding the influence of Fe domains on the NO_x storage behavior, structure, and morphology of the NO_x adsorption sites on the Fe/BaO/ γ -Al₂O₃ systems.

Thus, a fundamental insight regarding the NO_x storage properties can be obtained by systematically investigating the nature of the NO_x species formed on the surface of the Fe/BaO/ γ -Al₂O₃ mixed oxide system as a function of the Fe and Ba loadings. In order to achieve this goal, in the current study, we provide a detailed spectroscopic and microscopic account of the structure of the Fe/BaO/ γ -Al₂O₃ system and focus on the NO_x uptake mechanism at low temperature (323 K) in comparison with the conventional BaO/ γ -Al₂O₃ system. Besides, the thermal behavior of the adsorbed NO_x species on the Fe/BaO/ γ -Al₂O₃ system along with the surface morphologies of the Fe and Ba surface domains were also investigated via the temperature programmed desorption (TPD) technique.

2. Experimental Section

2.1. Sample Preparation. A series of BaO/ γ -Al₂O₃ and BaO/FeO_x/ γ -Al₂O₃ samples with different Ba (8 and 20 wt % BaO) and Fe (5 and 10 wt % Fe) loadings were synthesized by conventional incipient wetness impregnation of γ -Al₂O₃ (PURAL-OX, 206 m²/g, SASOL GmbH, Germany). In this synthetic protocol, the support material was impregnated with aqueous solutions of barium nitrate (ACS Reagent, $\geq 99\%$, Riedel - de Han, Germany) (for BaO/ γ -Al₂O₃ samples). The mixed FeO_x/BaO/ γ -Al₂O₃ (8 or 20 wt % BaO and 5 or 10 wt % Fe) samples were prepared by simultaneous impregnation of γ -Al₂O₃ with the aqueous solutions of barium and ferric nitrates (Fe(NO₃)₃·9H₂O, $\geq 98\%$, Sigma Aldrich). Then, the synthesized materials were dried at 353 K and calcined in Ar atmosphere for 2 h at various temperatures, ranging from 423 to 1273 K. Hence, after each temperature treatment, representative samples from the treated batches were collected for ex situ analysis. For the FTIR and TPD studies, samples synthesized via impregnation were directly annealed in Ar at 873 K for 2 h before their introduction to the catalytic reactor for the NO₂(g) adsorption experiments.

Compositions of the NO_x storage materials and their acronyms used in the current text are as follows: 8Ba/Al and 20Ba/Al samples contained 8 and 20 wt % BaO, respectively (with balancing amounts of Al₂O₃). FeO_x/BaO/Al₂O₃ samples were synthesized in such a manner that they have BaO to Al₂O₃ mass ratios that are identical to the BaO/Al₂O₃ samples. Thus, 5Fe/8(20)Ba/Al samples contained 5 wt % Fe, 7.6 wt % (19 wt %) BaO, and balancing amounts of Al₂O₃, while 10Fe/8(20)Ba/Al samples contained 10 wt % Fe, 7.2 wt % (18 wt %) BaO, and balancing amounts of Al₂O₃. Fe/Al samples were also prepared for supporting experiments, in which γ -Al₂O₃ was impregnated with Fe(NO₃)₃·9H₂O where the Fe content was 5 and 10 wt % for the 5Fe/Al and 10Fe/Al samples, respectively.

2.2. Experimental Methods. BET specific surface area measurements were performed using a Micromeritics Tristar 3000 surface area and pore size analyzer by low-temperature isothermal adsorption–desorption of N₂. Before the surface area measurements, the samples were outgassed at 623 K for 2 h in a vacuum. The transmission electron microscopy (TEM) specimens were prepared by suspending the finely ground sample particles in ethanol and dispersing this suspension on a lacey-type carbon film coated copper TEM grid. TEM measurements were carried out in bright field mode using an FEI Tecnai G2 F30 microscope with a specified point resolution of 0.17 nm, a Gatan slow-scan CCD camera, and a Gatan serial energy-

loss spectrometer. The operating voltage of the microscope was 300 keV during the TEM imaging. Elemental analysis of the imaged samples was performed by acquiring the electron energy loss spectra (EELS) with an incident electron energy of 100 kV. The typical duration of the EELS spectral acquisition was 15–20 s.

The powder X-ray diffraction (XRD) patterns were recorded using a Rigaku diffractometer, equipped with a Miniflex goniometer and an X-ray source with Cu K α radiation, at $\lambda = 1.5418$ Å, 30 kV, and 15 mA. The powder samples were scanned in the 10–80° 2 θ range with a scan rate of 0.01° s⁻¹. Diffraction patterns were assigned using Joint Committee on Powder Diffraction Standards (JCPDS) cards supplied by the International Centre for Diffraction Database (ICDD). X-ray photoelectron spectroscopy (XPS) was acquired using a Perkin-Elmer PHI560 ESCA equipped with a Mg K α anode operated at 250 W and 15 kV. Binding energies were charge referenced at C 1s = 284.7 eV to denote adventitious carbon.

Raman spectra were recorded on a HORIBA Jobin Yvon LabRam HR 800 instrument, equipped with a confocal Raman BX41 microscope, a spectrograph with an 800 mm focal length, and a CCD detector. The Raman spectrometer was equipped with a Nd:YAG laser ($\lambda = 532.1$ nm). During the Raman experiments, the laser power was tuned to 20 mW, in order to minimize the sample heating effects. Before the Raman measurements, the powder samples were mechanically dispersed onto a single-crystal Si holder. The spectrometer was regularly calibrated by adjusting the zero-order position of the grating and comparing the measured Si Raman band frequency with the typical reference value of 520.7 cm⁻¹. All Raman spectra were acquired within 100–4000 cm⁻¹ with an acquisition time of 213 s and a spectral resolution of 4 cm⁻¹. Bulk Ba(NO₃)₂, Fe(NO₃)₃·9H₂O, and Fe₂O₃ nanopowder (Sigma-Aldrich, part no. 544884-5G) samples were used for the reference Raman spectra.

Fourier transform infrared (FTIR) spectroscopic measurements were carried out in transmission mode in a batch-type catalytic reactor coupled to an FTIR spectrometer (Bruker Tensor 27) and a quadrupole mass spectrometer (QMS) (Stanford Research Systems, RGA 200) for temperature programmed desorption (TPD) and residual gas analysis (RGA). FTIR spectra were recorded using a Hg–Cd–Te (MCT) detector, where each spectrum was acquired by averaging 128 scans with a spectral resolution of 4 cm⁻¹. The samples were mounted into the IR cell²⁸ consisting of a five-way stainless steel chamber equipped with optically polished KBr windows. This IR cell was connected to a gas manifold (including a dual-stage rotary vane pump and two turbomolecular pumps) so that the pressure in the cell could be varied within 1000–10⁻⁶ Torr. About 20 mg of finely ground powder sample was pressed onto a high-transmittance, lithographically etched fine-tungsten grid which was mounted on a copper sample holder assembly, attached to a ceramic vacuum feedthrough. A K-type thermocouple was spot-welded to the surface of a thin tantalum plate attached on the W-grid to monitor the sample temperature. The sample temperature was controlled within 298–1100 K via a computer-controlled DC resistive heating system using the voltage feedback from the thermocouple. After having mounted the sample in the IR cell, the sample was gradually heated to 373 K in vacuum and kept at that temperature for at least 12 h before the experiments in order to ensure the removal of water from the surface. In an attempt to minimize the effect of background water (i.e., readsorption of water on the sample from the reactor walls and the gas manifold), the system was frequently baked.

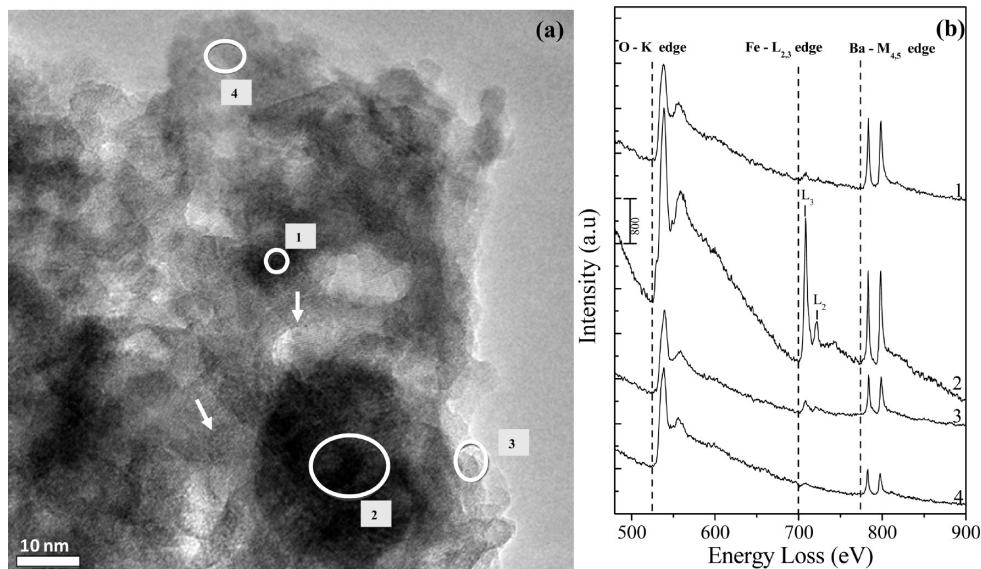


Figure 1. Representative TEM image (a) of a 10Fe/20Ba/Al sample that is initially thermally treated at 873 K for 2 h in Ar(g). EELS data (b) for a set of randomly selected points on the image are also given. Arrows emphasize the relatively ordered domains on the surface.

TABLE 1: BET Specific Surface Area Measurement Results for the Investigated Materials

sample ^{a,b}	γ -Al ₂ O ₃	8Ba/Al	20Ba/Al	5Fe/8Ba/Al	10Fe/8Ba/Al	5Fe/20Ba/Al	10Fe/20Ba/Al
	205	185	126	163	160	125	109

^a All of the samples were initially annealed at 873 K in Ar(g) for 2 h before the BET measurements. ^b Surface area values are given in m²/g.

Prior to each NO_x adsorption experiment, the walls of the vacuum system were passivated by flushing 2 Torr of NO₂(g) for an extended period of time (20 min) followed by evacuation at the same temperature. Next, in order to obtain a surface that is free of adsorbed NO_x and other adsorbates, the sample was annealed in vacuum by increasing the temperature to 1023 K in a linear fashion at a constant rate of 12 K min⁻¹. After this annealing protocol, the sample was cooled down to 323 K. Before acquisition of each spectral series, a background spectrum of the clean, adsorbate-free sample was obtained in a vacuum at 323 K (with a residual reactor pressure of $\sim 1 \times 10^{-5}$ Torr).

NO₂(g) (prepared by reacting NO(g), Air Products, Purity >99.9% and O₂(g) Ersoy Gas, Purity >99.9%) used in the adsorption experiments was further purified *via* several freeze–thaw–pump cycles before the experiments. NO₂(g) adsorption experiments were performed *via* two subsequent NO_x uptake protocols. In the first protocol, precisely controlled doses of 0.6 Torr of NO₂(g) (measured by a 1000 Torr MKS Baratron absolute pressure transducer) were introduced in a stepwise fashion onto the NO_x-free sample surface at 323 K. After having the acquired sample spectrum of each dose at 323 K, the system was evacuated to $\sim 1 \times 10^{-3}$ Torr before the next dosage. This protocol allowed us to follow the FTIR spectral changes during the gradual NO₂(g) adsorption on the samples at 323 K. In the second NO_x uptake protocol, subsequent to the gradual (stepwise) NO₂ doses, the sample was exposed to 8 Torr of NO₂(g) for 20 min at 323 K in order to ensure the complete saturation of the sample surface with NO_x and the FTIR spectra were acquired after having evacuated the reactor to $\sim 1 \times 10^{-3}$ Torr at 323 K.

TPD experiments were performed by using a QMS that is directly connected to the vacuum chamber through a pneumatic gate valve. Before the TPD experiments, the sample powder (mass = ca. 20 mg) pressed onto a tungsten grid was mounted in the IR/TPD cell and pretreated using the identical procedure

described above. After cooling to 323 K, the sample was exposed to 8 Torr of NO₂ for 20 min until the equilibrium was reached. Next, the system was outgassed to $\sim 1 \times 10^{-6}$ Torr in order to remove weakly physisorbed molecules. Subsequently, TPD experiments were carried out. Evolved NO_x species during the desorption process were monitored by recording the QMS signals with mass to charge ratios (*m/z*) equal to 18, 28, 30, 32, 44, and 46 in pressure vs time mode. A linear temperature ramp within 323–1023 K was used during the TPD experiments where the ramp rate was 12 K/min. In order to ensure the reproducibility of the results, TPD experiments were performed multiple times for each sample.

3. Results and Discussion

3.1. BET Surface Area Measurements. BET surface area measurements of the synthesized materials are given in Table 1. All of these samples were initially annealed in Ar(g) for 2 h at 873 K before the measurements. BET results suggest that the specific surface area values monotonically decrease with the increasing Ba and Fe loadings. This expected decrease in surface area values with respect to the bare γ -Al₂O₃ support material is due to the blocking of the porous alumina surface with low-surface-area Fe and/or Ba oxide domains as well as due to the solid state reactions described in sections 3.3 and 3.4.

3.2. TEM and EELS Measurements. Parts a and b of Figure 1 present TEM and EELS data for a 10Fe/20Ba/Al sample that is initially annealed in Ar(g) for 2 h at 873 K, respectively. The morphology of the 10Fe/20Ba/Al system presents complex and poorly defined features that are similar to the γ -Al₂O₃ (data not shown) and BaO/ γ -Al₂O₃ systems.²⁹ In addition to these, more ordered domains with well-defined orientations (indicated by the arrows in Figure 1a) are also clearly discernible which can be attributed to the presence of Fe and/or Ba-containing oxide domains exhibiting larger crystallites with a more ordered character than that of γ -Al₂O₃.

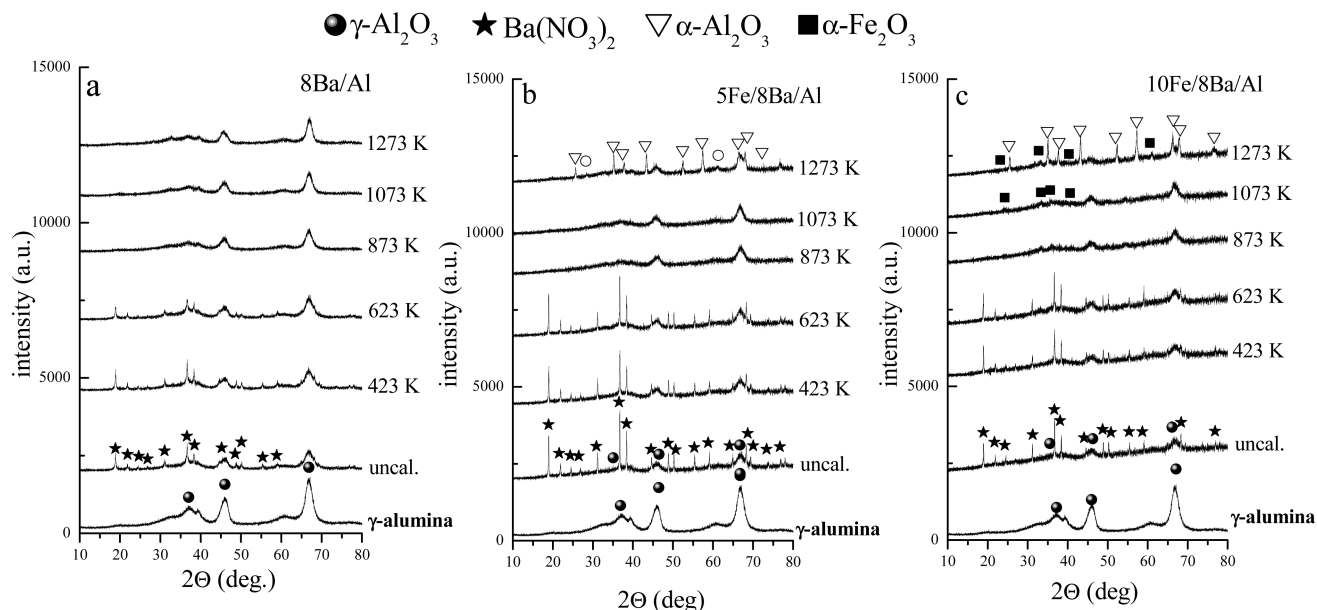


Figure 2. XRD profiles corresponding to the as-prepared and thermally treated (423–1273 K) (a) 8Ba/Al, (b) 5Fe/8Ba/Al, and (c) 10Fe/8Ba/Al samples (see text for details).

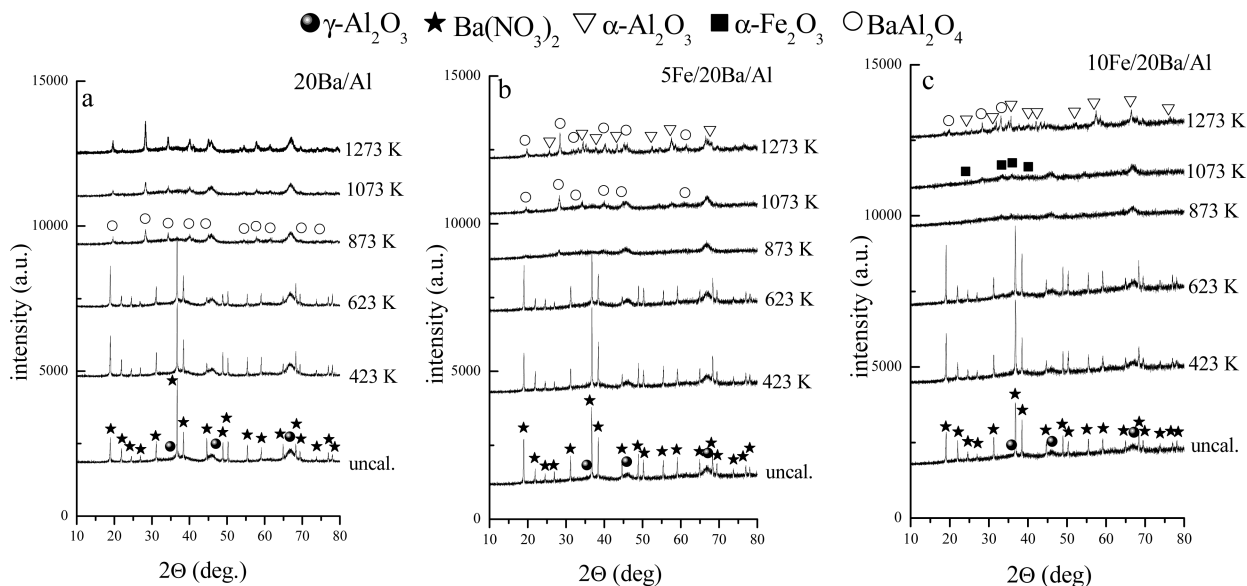


Figure 3. XRD profiles corresponding to the as-prepared and thermally treated (423–1273 K) (a) 20Ba/Al, (b) 5Fe/20Ba/Al, and (c) 10Fe/20Ba/Al samples (see text for details).

Elemental analysis of the imaged samples was also performed via EELS measurements. EELS spectra obtained from different locations of the image presented in Figure 1b reveal that the surface concentrations of the Fe and Ba containing domains are not homogeneous. Further information regarding the d-electron occupancy of the Fe sites can also be obtained by analyzing the relative integrated peak areas of the FeL_3 and FeL_2 white line peaks in the EELS data. It is well-known in the literature that the $\text{FeL}_3/\text{FeL}_2$ integrated area ratio can provide insight regarding the oxidation state of the Fe centers.³⁰ $\text{FeL}_3/\text{FeL}_2$ integrated area ratio values obtained from the EELS data presented in Figure 1b vary between 3.9 and 6.5 that is consistent with the presence of Fe_2O_3 , Fe_3O_4 , and possibly FeO_x domains. In other words, the oxidation states of the Fe centers on the surface of the 10Fe/20Ba/Al sample can exist in a variety of values such as Fe^{3+} , Fe^{2+} , and $\text{Fe}^{(3-x)+}$.

3.3. XRD Experiments. Ex situ X-ray diffraction studies were also performed after thermally treating the synthesized

materials at various temperatures ranging from 323 to 1273 K. Figure 2 presents XRD results corresponding to 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/Al samples with lower Ba content, whereas Figure 3 presents XRD data for the 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al samples with a higher Ba loading.

XRD patterns of all of the uncalined samples in Figure 2 present diffraction maxima that can be assigned to $\gamma\text{-Al}_2\text{O}_3$ (JCPDS 29-0063, marked with filled circles) and crystalline $\text{Ba}(\text{NO}_3)_2$ (JCPDS 24-0053, marked with stars). It should be pointed out that the $\text{Ba}(\text{NO}_3)_2$ -related reflections in the XRD profiles of the 5Fe/8Ba/Al and 10Fe/8Ba/Al samples are characterized with relatively sharper features with higher intensities in comparison with that of the 8Ba/Al sample. This observation is also consistent with our TEM results given in section 3.2, indicating the presence of crystallographically ordered domains on the surface. Heat treatment at 423 and 623 K does not result in any significant changes in the XRD profiles. A further increase in the annealing temperature to 873 K, leads

to the complete disappearance of the diffraction maxima due to the Ba(NO₃)₂ for all samples. It can be seen in Figure 2a that only very broad diffraction features associated with the γ -alumina support are visible in the XRD profiles of the 8Ba/Al sample calcined in the range 873–1273 K. However, at higher calcination temperatures such as 1273 K, both in the 5Fe/8Ba/Al and 10Fe/8Ba/Al samples, γ -Al₂O₃ support material starts to undergo a phase transformation toward the α -Al₂O₃ (corundum) phase (marked with open triangles in Figure 2b and c) (similarly, such a phase transition was observed for the 5Fe/Al and 10Fe/Al samples that do not contain any Ba domains, as shown in Figure 1 of the Supporting Information). It should be noted that such a phase transformation takes place at $T > 1273$ K on the pure alumina surface. Former studies in the literature suggest that the presence of small α -Fe₂O₃ (hematite) particles leads to a decrease in the γ -Al₂O₃ \rightarrow α -Al₂O₃ phase transformation temperature.³¹ It is also noteworthy to mention that diffraction features belonging to α -Fe₂O₃ (hematite) (JCPDS 72-0469) become visible at $T > 1073$ K for the 10Fe/8Ba/Al sample, indicating the formation of larger hematite crystallites with increasing Fe loading. In addition to these, very minor features associated with the BaAl₂O₄ phase (JCPDS 17-0306) (marked with open circles) are also visible in Figure 2b for the 5Fe/8Ba/Al sample at 1273 K. These results regarding the BaAl₂O₄ formation at elevated temperatures are also in good agreement with our XPS results that are provided in the Supporting Information (Figure 2).

In accordance with the higher Ba loading, XRD patterns given in Figure 3 have sharper and more intense Ba(NO₃)₂ features within 323–873 K where the Ba(NO₃)₂ decomposes into BaO at 873 K for all samples. Another common characteristic of the XRD patterns in Figure 3 is the more visible BaAl₂O₄ features appearing at temperatures as low as 873 K (e.g., 20Ba/Al case in Figure 3a) that are consistent with the higher Ba loading.^{7,10} In addition, comparison of Figure 3c to Figure 3a and b reveals that increasing Fe content in the Fe/Ba/Al system suppresses the BaAl₂O₄ formation at higher temperatures, suggesting a stronger interaction between the Fe domains and the underlying Al₂O₃ support material for the higher Fe loadings. These complex thermal changes occurring on the Ba/Al and Fe/Ba/Al systems particularly at $T \geq 873$ K leading to the formation of BaAl₂O₄, Fe₂O₃, and α -Al₂O₃ are consistent with the smaller specific surface areas of the Ba/Al and Fe/Ba/Al systems in comparison with the γ -Al₂O₃ support material (section 3.1).

3.4. Raman Spectroscopy Experiments. In order to obtain further structural information regarding the synthesized materials, Raman spectroscopic experiments were performed after thermally treating the samples at various temperatures within 323–1273 K. The spectra (i–iii) in Figure 4a also present reference Raman data for unsupported (pure) Ba(NO₃)₂, Fe(NO₃)₃·9H₂O, and α -Fe₂O₃ (hematite), respectively. Similarly, Figure 4b and c shows Raman spectroscopic results for the 5Fe/8Ba/Al and 10Fe/8Ba/Al samples, while Figure 5 shows corresponding results for the 5Fe/20Ba/Al and 10Fe/20Ba/Al samples with a higher Ba loading.

The Raman spectra given in Figures 4 and 5 reveal similar dominant features at $T \leq 623$ K. These dominant Raman features can be assigned to the relevant surface species with the help of the former reports in the literature^{32–34} as well as the supporting experiments given in Figures 3 and 4 of the Supporting Information. The spectra in Figures 4 and 5 at $T \leq 623$ K are dominated by the strong Raman bands associated with the symmetric (1052 cm⁻¹), antisymmetric stretching (~1357 and 1406 cm⁻¹), and bending modes (740 cm⁻¹) of the

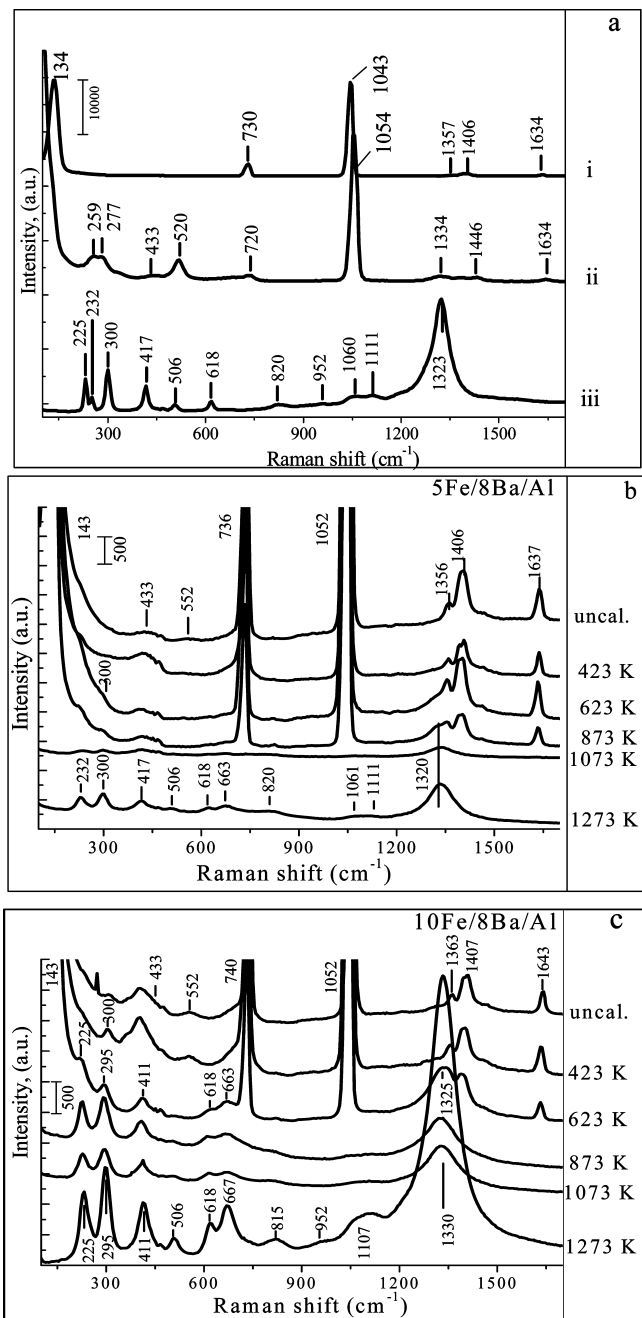


Figure 4. Ex situ Raman spectra corresponding to pure reference materials (a) i, Ba(NO₃)₂; ii, Fe(NO₃)₃·9H₂O; iii, α -Fe₂O₃ and to the as-prepared and thermally treated (423–1273 K) (b) 5Fe/8Ba/Al and (c) 10Fe/8Ba/Al samples.

NO₃⁻ ions which bear significant resemblance to the reference Ba nitrate and Fe nitrate spectra given in Figure 4a (spectra i and ii). Furthermore, the lattice phonon band at 143 cm⁻¹ which clearly indicates the presence of crystalline Ba(NO₃)₂ is also present in all of the samples in Figures 4 and 5 at $T \leq 623$ K. Furthermore, the weak band at ~1640 cm⁻¹ can be readily assigned to the longitudinal optical (LO) mode and the group of weak bands at about 1360 and 1410 cm⁻¹ to the transverse optical (TO) modes of the antisymmetric nitrate stretch, respectively.^{32–34} In addition, a low frequency band at 552 and a weak shoulder at ~433 cm⁻¹ corresponding to the symmetric A_{1g}(Fe–OH)₂ stretching mode of Fe(H₂O)₆³⁺ ion can also be observed³⁴ in Figures 4 and 5 which reveal the coordination of the Fe centers to the surface hydroxyls or water molecules after the impregnation procedure. Observations given above point to

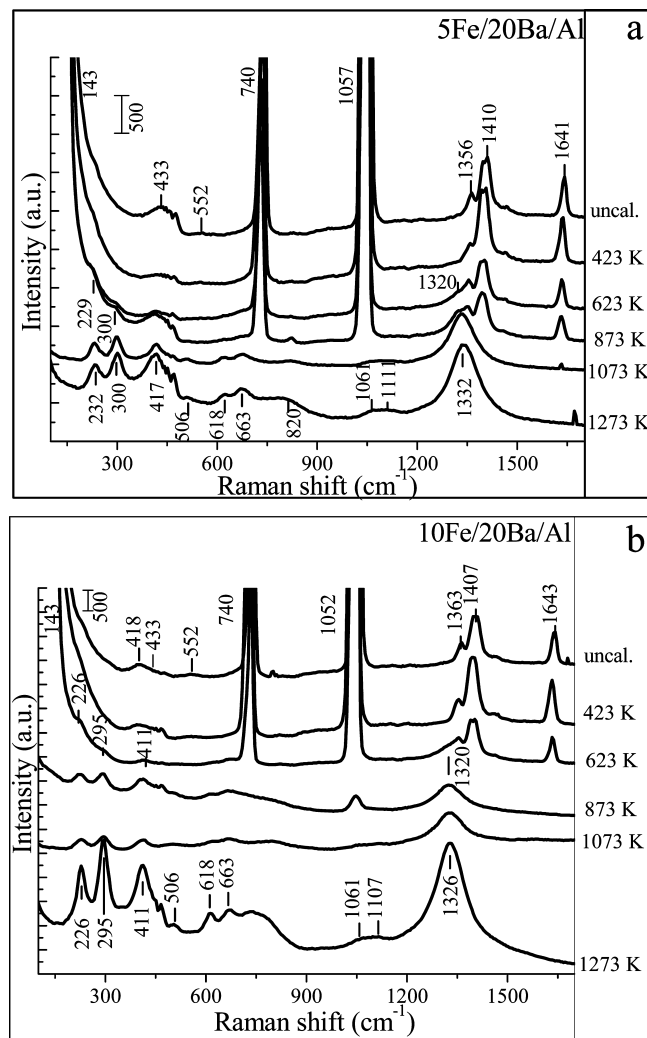


Figure 5. Ex situ Raman spectra corresponding to the as-prepared and thermally treated (423–1273 K) (a) 5Fe/20Ba/Al and (b) 10Fe/20Ba/Al samples.

the fact that $\text{Fe}(\text{NO}_3)_3$ and $\text{Ba}(\text{NO}_3)_2$ domains coexist on the Fe/Ba/Al samples at low temperatures which yield similar nitrate-related overlapping bands in the Raman spectra.

The samples analyzed after calcining at 623 K in Figures 4 and 5 are still dominated by the strong nitrate features; however, they also unveil an important aspect, the $\alpha\text{-Fe}_2\text{O}_3$ (hematite) formation. Hematite formation at 623 K is evident in the 5Fe/8Ba/Al and 5Fe/20Ba/Al samples (Figures 4b and 5a) as weak shoulders at ca. 300 and 1320 cm^{-1} . They are more clearly discernible in the samples with higher Fe content such as the 10Fe/8Ba/Al and 10Fe/20Ba/Al samples (Figures 4c and 5b), as a result of the stronger bands at ~ 225 , 295, 411, 618, and 1328 cm^{-1} . Thus, it is apparent that the decomposition of the deposited ferric nitrates on the surface results in the formation of $\alpha\text{-Fe}_2\text{O}_3$ in the Fe/Ba/Al system. $\text{Ba}(\text{NO}_3)_2$ decomposition also occurs in a similar thermal window (i.e., 623–873 K) where $\text{Ba}(\text{NO}_3)_2$ decomposes to form $(\text{BaO})_x$. Barium oxide crystallizes in the rock salt structure, and the first order Raman scattering is symmetry forbidden.^{32,33} However, broad and weak bands within 350–500 cm^{-1} due to the lattice modes of the defective BaO domains can still be visible via Raman spectroscopy (see for instance, Figure 3 in the Supporting Information). However, due to the stronger and complex Fe_2O_3 -related features in this spectral window, these weaker BaO lattice features are significantly obscured in Figures 4 and 5.

Raman spectra of the Fe/Ba/Al systems annealed at $T \geq 1073$ K in Figures 4 and 5 are almost exclusively dominated by the Fe_2O_3 features regardless of the relative composition of the Fe/Ba/Al structure. The crystalline $\alpha\text{-Fe}_2\text{O}_3$ (hematite) belongs to the D_{3d}^6 crystal space group, which exhibits the following Raman features:^{35,36} seven phonon lines (two A_{1g} modes at 232 and 506 cm^{-1} and five E_g modes at 252, 293, 299, 417, and 618 cm^{-1}) and a magnon scattering feature at ~ 1323 cm^{-1} . Furthermore, a number of additional minor bands at 820, 952, 1060, and 1111 cm^{-1} which are also characteristic of the hematite structure³⁶ can also be seen in the spectra. The increased band intensities corresponding to the hematite domains at higher temperatures may suggest enrichment in the surface concentration of $\alpha\text{-Fe}_2\text{O}_3$ species, ordering of the hematite crystal structure, as well as an increase in the $\alpha\text{-Fe}_2\text{O}_3$ particle size (i.e., sintering). Raman spectra of all Fe containing oxide materials corresponding to 1273 K also reveal a readily distinguishable band at 663 cm^{-1} . According to the former Raman spectroscopy studies in the literature on various iron oxides,^{35,36} the Raman spectrum of Fe_3O_4 (magnetite) is characterized by strongly expressed bands located at 663 cm^{-1} and weaker bands at 302, 514, and 534 cm^{-1} . Thus, it is apparent that, within 623–1273 K, the Raman spectra of the Fe/Ba/Al samples given in Figures 4 and 5 consist of combinations of bands related to the simultaneous presence of $\alpha\text{-Fe}_2\text{O}_3$ (hematite) and Fe_3O_4 (magnetite). It should also be noted that the presence of minority iron oxide domains with dissimilar stoichiometries to the ones discussed above cannot be ruled out, as these structures may be elusive to detect via ex situ Raman spectroscopy and XRD techniques.

3.5. Low-Temperature NO_x Uptake of the Fe/Ba/Al System: FTIR Spectroscopy Experiments. After having acquired a detailed understanding of the structural properties of the Fe/Ba/Al NO_x storage systems, the low-temperature NO_x storage behavior of these materials was analyzed via FTIR spectroscopy. NO_2 adsorption experiments were performed on the Fe/Ba/Al systems, as described in the Experimental Section. NO_x uptake results of the Fe/Ba/Al systems with a lower (8 wt %) Ba loading are presented in section 3.5.1, while the results for the samples with a higher Ba loading (20 wt % Ba) are given in section 3.5.2.

3.5.1. Stepwise $\text{NO}_2(\text{g})$ Adsorption on 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/Al at 323 K. Figure 6 presents FTIR data for $\text{NO}_2(\text{g})$ adsorption on 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/10Al samples. NO_2 adsorption on the BaO/ $\gamma\text{-Al}_2\text{O}_3$ system has been thoroughly studied in the literature via FTIR spectroscopy^{37–40} where the nature of the adsorbed NO_x species was mostly interpreted by considering the BaO adsorption sites that belong to different morphologies on the support ($\gamma\text{-Al}_2\text{O}_3$) surface. Along these lines, it was suggested that there exists two major types of BaO adsorption sites relevant to NO_2 adsorption, the so-called “surface” and “bulk” BaO sites. The surface BaO sites were suggested to exist in the form of a $(\text{BaO})_x$ structure where well-dispersed BaO monomers, dimers, or other small oligomers directly interact with the underlying alumina support sites in a very strong fashion.³⁷ It was further suggested that this well-dispersed surface $(\text{BaO})_x$ species prefers to anchor particularly on the (100) facets of the $\gamma\text{-Al}_2\text{O}_3$ surface which constitutes about 16% of the total surface area of the $\gamma\text{-Al}_2\text{O}_3$ structure.⁴² In addition, these (100) facets showing high affinity toward BaO monomers and dimers, were proposed to expose exclusively penta-coordinated Al^{3+} sites on the surface.^{37–42} The second major type of BaO adsorption sites relevant to the NO_2 adsorption is the so-called “bulk” BaO sites, which were

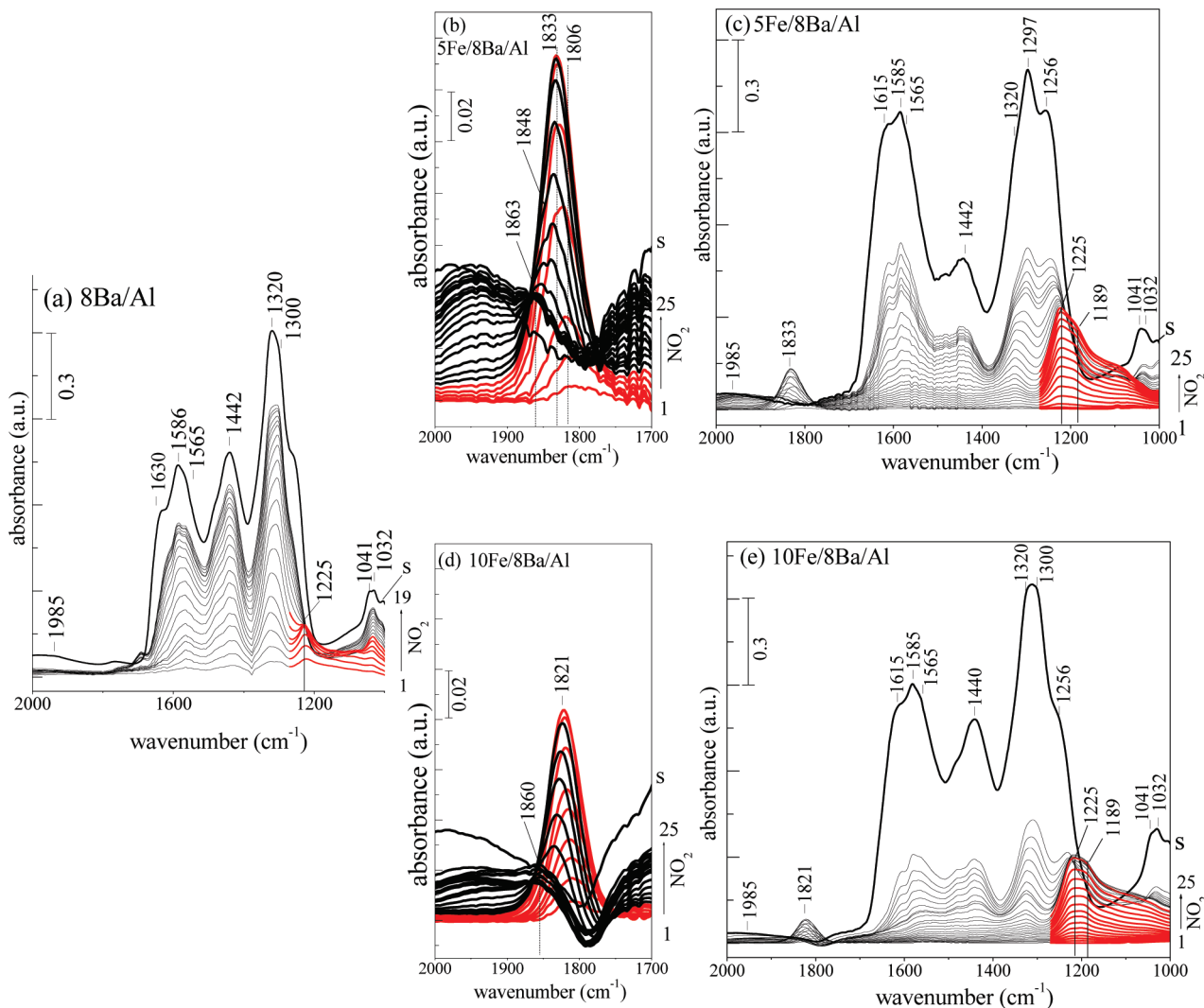


Figure 6. FTIR spectra corresponding to the stepwise NO₂ adsorption at 323 K on the (a) 8Ba/Al, (b, c) 5Fe/8Ba/Al, and (d, e) 10Fe/8Ba/Al samples. The spectra corresponding to the fully saturated sample surfaces are marked with an “S” (see text for details). Red lines emphasize the early stages of the NO_x uptake.

suggested to be in the form of three-dimensional BaO clusters on the alumina surface which are capable of not only adsorbing NO_x on the BaO cluster surface but also capable of storing NO_x in the bulk of the BaO cluster.³⁷

Along these lines, the FTIR data corresponding to the NO₂ adsorption on the 8Ba/Al sample in Figure 6a can be readily analyzed in the light of the detailed former studies in the literature.^{37–42} During the initial NO₂ doses (the spectra marked with red in Figure 6a), the FTIR data is mostly dominated by a feature at 1225 cm⁻¹ which can be attributed to nitrites on the BaO sites. Concomitant with the formation of nitrites, further NO₂ doses lead to the oxidation of the surface nitrite species into nitrates. This is evident by the appearance of additional vibrational bands at 1585, 1565, and ~1300 cm⁻¹ that can be attributed to bidentate (surface) nitrates. Furthermore, the shoulder at 1320 cm⁻¹ and the doublet with a characteristic splitting of the asymmetric stretching vibration bands at 1442 cm⁻¹ can be assigned to the bulk nitrates. Symmetric vibrational modes (ν_{sym}) of the nitrate groups mentioned above also reveal themselves as weak features at 1030 and 1041 cm⁻¹. In addition to these, a broad vibrational feature at ~1985 cm⁻¹ is also visible which can be attributed³⁹ to the weakly adsorbed N₂O₃ or NO⁺ on the surface. Figure 6a shows that, upon increasing NO₂ doses at 323 K, most of the NO_x species on the 8Ba/Al surface are

oxidized to form surface and bulk nitrates while some of the nitrate species are adsorbed on the exposed γ -Al₂O₃ support material in the form of bidentate and/or bridging nitrates, indicated by the observed shoulders at 1630 and 1256 cm⁻¹. It is well-known in the literature^{37,38} that, at relatively low loadings of Ba such as 8 wt %, the NO_x storage component is not able to cover the support surface completely, leaving bare support sites available for NO_x adsorption.

Using the 8Ba/Al system as a benchmark, NO₂ adsorption experiments on the 5Fe/8Ba/Al and 10Fe/8Ba/Al systems presented in Figure 6b–e can be elucidated in detail. It is apparent that the initial NO₂ uptake of the 5Fe/8Ba/Al and 10Fe/8Ba/Al systems at 323 K shows significant discrepancies with that of the 8Ba/Al system. First of all, although surface oxidation of nitrite species to form nitrates on the 8Ba/Al system readily occurs after the first few initial NO₂ doses, on the 5Fe/8Ba/Al and 10Fe/8Ba/Al surfaces, nitrate formation is not visible in these initial doses and becomes visible only after dosing a large number of NO₂ pulses (note that the pressures and exposure durations used in the NO₂ pulses were identical for all doses and for all surfaces). This suggests that the presence of Fe sites significantly alters the oxidation of NO_x species on the surface by hindering the transformation of nitrites into nitrates. The structures of the nitrite bands are also relatively more complex

in the Fe-containing systems (Figure 6c and e), as an additional nitrite band exists at 1189 cm^{-1} , signifying the existence of at least one more type of nitrite species that does not exist in the case of the 8Ba/Al (Figure 6a).

Another important difference in the initial NO_x uptake behavior of the 5Fe/8Ba/Al and 10Fe/8Ba/Al systems reveals itself in the spectral region located within $1700\text{--}2000\text{ cm}^{-1}$. It is visible in Figure 6b and d that NO_2 adsorption on the Fe-containing systems leads to a new feature around $1800\text{--}1860\text{ cm}^{-1}$. This additional feature is visible even after the very first NO_2 pulse and increases its intensity for a certain period, with a slight shift to higher frequencies. After reaching a maximum intensity at $1821\text{--}1833\text{ cm}^{-1}$, the intensity of this new band starts to attenuate with a further blue shift in frequency. Concomitant with this intensity attenuation, a significant growth in the intensity of the nitrate bands ($1250\text{--}1620\text{ cm}^{-1}$) as well as the broad $\text{N}_2\text{O}_3/\text{NO}^+$ band (1950 cm^{-1}) is observed. A vibrational feature within $1800\text{--}1860\text{ cm}^{-1}$ was also observed in a former study where NO adsorption was performed on both oxidized and reduced hematite particles supported on SiO_2 .⁴³ On the basis of these former results, the new feature given in Figure 6b and d can be readily assigned to various types of nitrosyls on reduced Fe centers (i.e., Fe^{2+} and/or $\text{Fe}^{(3-x)+}$). It should be noted that the presence of reduced Fe centers as a result of the initial thermal activation and the cleaning protocol (described in the Experimental Section) before the introduction of $\text{NO}_2(\text{g})$ into the FTIR cell is also supported by the ex situ characterization results given in sections 3.1–3.4. Thus, it is apparent that, immediately after the very first NO_2 dose, NO_2 species preferentially adsorb on the highly reactive coordinatively unsaturated (and reduced) $\text{Fe}^{2+}/\text{Fe}^{(3-x)+}$ centers,⁴³ and decompose on these centers, leaving an oxygen atom which can further oxidize the neighboring reduced iron sites. Consequently, during the first few NO_2 doses at 323 K, it is likely that most of the NO_x species are converted into nitrosyls on the Fe centers while the remaining NO_x species exist exclusively in the form of nitrites that are located on both BaO and FeO_x domains. The presence of an additional nitrite feature at 1189 cm^{-1} (Figure 6c and e) in the Fe/Ba/Al systems is also consistent with the fact that this additional nitrite species is likely located on the Fe centers.

Therefore, the NO_x coverage-dependent changes within the $1800\text{--}1860\text{ cm}^{-1}$ region can be described by considering the formation of dinitrosyl species (1810 cm^{-1}) on the reduced $\text{Fe}^{2+}/\text{Fe}^{(3-x)+}$ centers with a lower (4-fold) coordination⁴³ followed by a transformation into mononitrosyl (1830 cm^{-1}) species located on the high-coordination (6-fold) $\text{Fe}^{2+}/\text{Fe}^{(3-x)+}$ centers at higher NO_x coverages. After the complete oxidation of the reduced Fe sites and the healing of the oxygen defects on the surface, oxidation of the surface NO_x species starts to occur with the further $\text{NO}_2(\text{g})$ pulses. This leads to the formation of the stable nitrates ($1250\text{--}1620\text{ cm}^{-1}$) as well as the weakly bound $\text{N}_2\text{O}_3/\text{NO}^+$ species (1950 cm^{-1}). It should be mentioned that the nitrosyl groups on Fe^{3+} sites are relatively unstable at 323 K which is in very good agreement⁴³ with the attenuation of the nitrosyl intensity after the completion of the oxidation of the reduced Fe centers on the surface. It is also worth mentioning that similar low-temperature NO_2 adsorption experiments performed on the 5Fe/Al and 10Fe/Al systems (data not shown) which do not possess any Ba domains also reveal a similar initial NO_x uptake behavior to the 5Fe/8Ba/Al and 10Fe/8Ba/Al systems, while such a hindrance regarding the nitrite oxidation or the generation of the surface nitrosyls was not detected during the NO_2 adsorption on the pure $\gamma\text{-Al}_2\text{O}_3$ support

material (data not shown). These control experiments also support the fact that the dissimilar initial NO_x uptake of the Fe/Ba/Al systems is directly linked to the Fe-containing domains.

Later stages of the NO_x uptake of the 5Fe/8Ba/Al and 10Fe/8Ba/Al systems (Figure 6c and e) show also significant dissimilarities with respect to the 8Ba/Al system (Figure 6a). After the saturation of the 5Fe/8Ba/Al and 10Fe/8Ba/Al surfaces with NO_2 (spectra labeled with the letter “S” in Figure 6c and e), the relative intensities of the nitrate bands present a different profile than the 8Ba/Al system. Overall, the presence of Fe leads to an increase in the bidentate nitrate bands that are located at 1300 , 1585 , and 1565 cm^{-1} . Detailed analysis of the thermal behavior of these species followed by the TPD technique described in section 3.7 suggests that these bidentate nitrate species are associated with the Fe^{3+} sites on the Fe_2O_3 (hematite), Fe_3O_4 (magnetite), and FeO_x domains in addition to the regular surface nitrates on the Ba domains. Therefore, it is apparent that the promotion of the conventional Ba/Al system with Fe oxides leads to the creation of new NO_x storage sites that can store NO_x in the form of bidentate nitrates. The influence of the Fe loading on the Fe/Ba/Al systems with relatively low Ba content (i.e., 8 wt % BaO) during the low-temperature NO_x uptake can also be investigated by comparing parts b and c of Figure 6 with parts d and e. Such a comparison clearly indicates that increasing the Fe loading to 10 wt % decreases the relative intensities of the Fe-nitrosyls ($1800\text{--}1860\text{ cm}^{-1}$) during the initial uptake. In a similar fashion, the intensities of the bidentate nitrate species formed after the saturation of the surface with $\text{NO}_2(\text{g})$ are also lower in Figure 6d and e with respect to Figure 6b and c. These observations point to the fact that the NO_x uptake capability of the Fe domains diminishes with increasing Fe loading. This can be attributed to the slightly lower specific surface area of the 10Fe/8Ba/Al sample (section 3.1), as well as the increase in the crystallite size of the Fe domains with increasing Fe content (sections 3.3–3.4) and the Fe-induced structural changes in the BaO units and the $\gamma\text{-Al}_2\text{O}_3$ support material (section 3.3).

3.5.2. Stepwise $\text{NO}_2(\text{g})$ Adsorption on 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al at 323 K. Figure 7 shows the stepwise NO_2 adsorption on 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al samples at 323 K. Comparison of the low-temperature $\text{NO}_2(\text{g})$ uptake of the 20Ba/Al sample with the 8Ba/Al sample reveals that the IR intensity ratio of the bulk nitrates to surface nitrates is greater for the 20Ba/Al case, suggesting that Ba domains in the 20Ba/Al case contain larger 3D clusters that prefer to store NO_2 as bulk nitrates.^{37–42} The presence of more significantly expressed bulk nitrates is also apparent in the 5Fe/20Ba/Al and 10Fe/20Ba/Al systems (Figure 7c and e) especially when compared to the 5Fe/8Ba/Al and 10Fe/8Ba/Al cases (Figure 6c and e). Initial NO_x uptake characteristics of the 5Fe/20Ba/Al system show significant resemblance to the 5Fe/8Ba/Al and 10Fe/8Ba/Al cases, suggesting that the initial oxidation of the reduced Fe centers and the nitrite formation are followed by the oxidation of the surface NO_x species to nitrates. The Fe-induced hindrance of the nitrite \rightarrow nitrate transformation is slightly less visible in the 5Fe/20Ba/Al case with respect to the 5Fe/8Ba/Al case, which may be attributed to the partial covering of the Fe sites by Ba in the 5Fe/20Ba/Al system.

A significant deviation in the NO_x uptake behavior as well as in the nature of the adsorbed nitrate species is observed particularly in the case of the 10Fe/20Ba/Al system. It is clearly seen in Figure 7d and e that, during the initial NO_2 doses on the 10Fe/20Ba/Al surface, oxidation of the nitrites to form

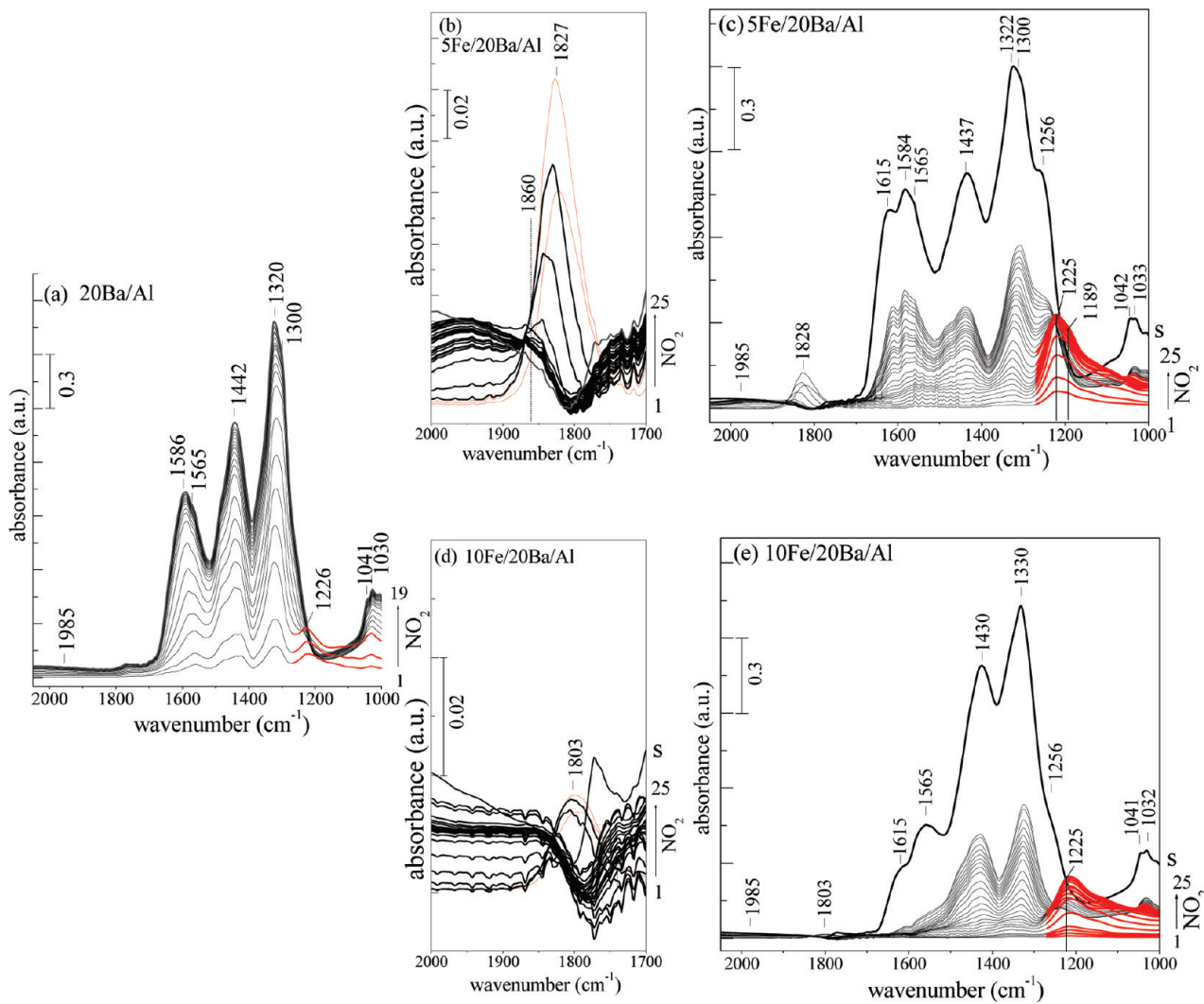


Figure 7. FTIR spectra corresponding to the stepwise NO₂ adsorption at 323 K on the (a) 20Ba/Al, (b, c) 5Fe/20Ba/Al, and (d, e) 10Fe/20Ba/Al samples. The spectra corresponding to the fully saturated sample surfaces are marked with an “S” (see text for details). Red lines emphasize the early stages of the NO_x uptake.

nitrate starts almost simultaneously with the formation of the nitrite species. Furthermore, nitrosyl, nitrite, and bidentate nitrate formation on the Fe domains are significantly suppressed on the 10Fe/20Ba/Al surface, whereas the formation of bulk nitrates on the Ba domains is greatly enhanced. In a parallel fashion, increasing abundances of Fe and Ba in the composition of the Fe/Ba/Al systems also decrease the fraction of the exposed γ -Al₂O₃ support material evident by the attenuation of the shoulders at 1630 and 1256 cm⁻¹ in Figure 7e. All of these indicators point to the fact that the morphology of the NO_x storage domains is profoundly altered in the case of the 10Fe/20Ba/Al system where Fe domains agglomerate to form large crystallites with a lower surface area that are less accessible to NO₂ adsorption (consistent with our BET measurements for this surface), possibly accompanied by covering of the Fe domains with the bulk Ba domains. A more comprehensive account of the morphological changes occurring on the different NO_x storage units will be discussed in sections 3.7 and 3.8 below.

3.6. Regeneration of the Reduced Fe Sites in the Fe/Ba/Al NO_x Storage System. An important aspect that was also investigated in our studies regarding the presence of the additional NO_x storage sites that are associated with the reduced Fe sites was whether these reduced sites could be regenerated in a reducing environment, after an oxidative treatment. The

significance of this aspect can be realized particularly when the alternating oxidizing (i.e., lean)⁷ and reducing (i.e., rich)⁷ operational regimes in the NSR catalytic process are considered. Therefore, a series of simple (proof of principle) experiments were performed whose results are presented in Figure 8. In these experiments, a freshly activated 5Fe/8Ba/Al sample (preannealed in a vacuum at 1023 K) was exposed to 10 Torr of O₂ for 20 min at 623 K. Then, the NO_x uptake of the preoxidized surface was followed during the NO₂ adsorption at 323 K via FTIR (Figure 8a). It is clearly seen in Figure 8a that the nitrosyl features that are associated with the presence of the reduced Fe sites are not visible for the preoxidized surface (Figure 8a). Furthermore, as expected from the oxidized surface, the nitrate region of the spectra reveals a simultaneous increase in the nitrite and nitrate bands without a visible hindrance in the surface oxidation of the NO_x species during the initial stages of the NO_x uptake. The preoxidized surface, which was saturated with NO_x, was subsequently exposed to a reducing environment. This was achieved by annealing the sample in a vacuum first at 1023 K (for 2 min) and then at 923 K (for 30 min). After this reduction protocol, NO₂ adsorption on this surface was performed at 323 K (Figure 8b). Figure 8b reveals the reappearance of the nitrosyl features during the initial stages of the NO_x uptake (which are converted into other NO_x species during the later

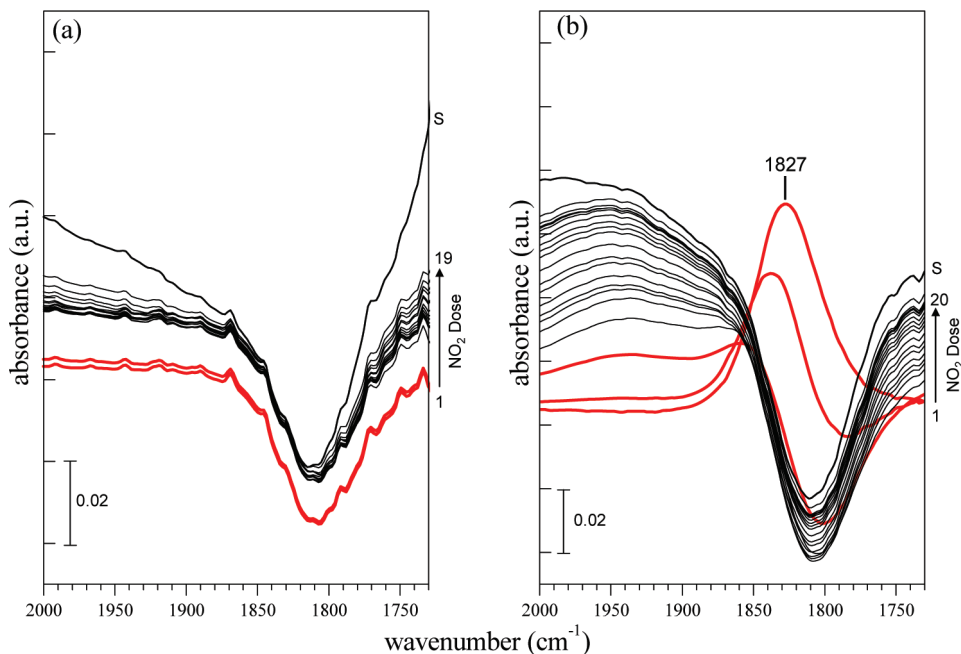


Figure 8. FTIR spectra corresponding to the stepwise NO_2 adsorption at 323 K on the preoxidized (a) 5Fe/8Ba/Al sample surface with 10 Torr of O_2 for 20 min at 623 K and on the subsequently reduced (b) sample by annealing in a vacuum first at 1023 K (for 2 min) and then at 923 K (for 30 min). The spectra corresponding to the fully saturated sample surfaces are marked with an “S” (see text for details). Red lines emphasize the early stages of the NO_x uptake.

stages of the NO_x adsorption, as described in the previous sections), indicating the regeneration of the reduced Fe centers after the reduction protocol. It should be noted that the intensities of the nitrosyl features in Figure 8b are slightly less than that of the freshly activated 5Fe/8Ba/Al surface (Figure 6b), implying a partial regeneration of the reduced Fe sites. It is worth mentioning that, under the realistic operational conditions of an NSR catalyst, powerful gas phase reducers (such as hydrocarbons, CO, and hydrogen) are present in the rich regime. Furthermore, these reducing agents can be readily activated on a real NSR catalyst particularly on the surface of the precious metal (e.g., Pt) particles serving as redox centers. Thus, although the actual operational temperature of the NSR process (ca. 523–623 K) is lower than the reduction temperature used above, it is plausible that the regeneration of the Fe centers under real operational conditions could also occur during the rich cycle due to the presence of strong gas phase reducing agents as well as the Pt redox centers on the NSR catalyst surface.

3.7. Thermal Behavior of the Adsorbed NO_x Species on the 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/Al Systems via TPD.

In order to elucidate the thermal behavior and the relative surface stabilities of the different types of NO_x species that are stored on the Fe/Ba/Al systems, FTIR (Supporting Information, Figures 5 and 6) and TPD experiments were performed.

Figure 9 shows TPD data obtained from the $\gamma\text{-Al}_2\text{O}_3$, 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/Al surfaces that are initially pretreated and subsequently saturated with $\text{NO}_2(\text{g})$, as described in the Experimental Section. The bottom-most TPD profiles given in Figure 9 correspond to the $\gamma\text{-Al}_2\text{O}_3$ support material. TPD experiments upon NO_2 adsorption on the $\gamma\text{-Al}_2\text{O}_3$ system have been thoroughly discussed in various former reports.^{40,44} In the light of these studies, it can be readily argued that TPD profiles for the $\gamma\text{-Al}_2\text{O}_3$ support material contain a low temperature feature at 389 K corresponding to the desorption of weakly bound N_2O_3 and NO^+ species, that desorb in the form of NO_2 and $\text{NO} + \text{O}_2$. The main desorption feature of the $\gamma\text{-Al}_2\text{O}_3$ support material appears at 625 K which corresponds

to desorption/decomposition of bridging and bidentate nitrates yielding predominantly a NO_2 desorption signal.^{40,44}

TPD profiles for the 8Ba/Al sample in Figure 9 indicated similar thermal decomposition of the weakly adsorbed species and the nitrates formed on the Al^{3+} NO_x adsorption centers.^{38,40} The low-temperature desorption signal located at 389 K can be attributed to the N_2O_3 and NO^+ species as in the case of the $\gamma\text{-Al}_2\text{O}_3$ surface, although the intensity of this feature is relatively smaller for the 8Ba/Al case due to the partial blocking of the $\gamma\text{-Al}_2\text{O}_3$ surface by the BaO domains. The shoulder at ca. 625 K can also be readily associated with desorption and decomposition of the nitrates that reside on the exposed sites of the $\gamma\text{-Al}_2\text{O}_3$ support material. The major desorption feature for the 8Ba/Al system at 685 K can be assigned to the decomposition of the surface (bidentate) nitrates associated with the BaO monomers/dimers strongly interacting with the support material. As discussed above, surface nitrates are relatively less stable than the bulk nitrates, which appear as a shoulder within 750–900 K.^{38,40} It is also worth mentioning that Figure 9 shows that surface nitrate decomposition yields mostly NO_2 species while bulk nitrate decomposition leads to mostly $\text{NO} + \text{O}_2$.

Using the TPD data for the $\gamma\text{-Al}_2\text{O}_3$ and the 8Ba/Al given in Figure 9 as a basis, TPD results of the 5Fe/8Ba/Al system can be analyzed in detail. TPD data for the 5Fe/8Ba/Al surface in Figure 9 is dominated by a new distinct feature that is located at 760 K. It should be noted that the thermal characteristics of this dominant feature cannot be explained by solely using the TPD results for the $\gamma\text{-Al}_2\text{O}_3$ and/or 8Ba/Al systems. This observation suggests that the new desorption feature at 760 K is associated with a new species that is present on the 5Fe/8Ba/Al surface and absent on the $\gamma\text{-Al}_2\text{O}_3$ and 8Ba/Al surfaces. Considering the FTIR results for the 5Fe/8Ba/Al surface given in Figure 6b indicating a significant accentuation of the bidentate nitrate species, it seems plausible to assign the desorption feature at 760 K to the desorption/decomposition of the bidentate nitrates associated with the Fe-containing domains. It is worth mentioning that this feature has similar desorption characteristics

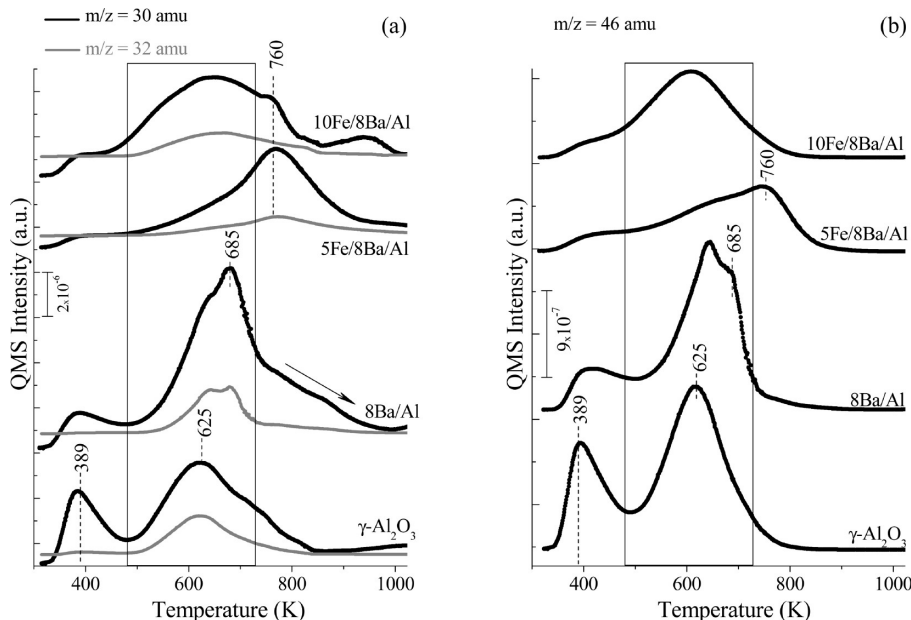


Figure 9. TPD profiles obtained from pure γ -Al₂O₃, 8Ba/Al, 5Fe/8Ba/Al, and 10Fe/8Ba/Al samples which were initially saturated with NO₂(g) at 323 K: (a) 30 and 32 amu signals; (b) 46 amu signal.

(i.e., desorbing mostly in the form of NO₂) to the surface nitrates on the well-dispersed BaO units that are strongly interacting with the alumina support surface. In addition to this major signal at 760 K, desorption features associated with the weakly bound N₂O₃ and NO⁺ species (389 K), nitrates on the γ -Al₂O₃ support surface (625 K), and surface (bidentate) Ba nitrates (685 K) are also visible as convoluted features on the low-temperature tail of the desorption trace of the 5Fe/8Ba/Al system, while the minor feature associated with the bulk Ba nitrate desorption (750–900 K) mostly appears in a heavily convoluted form, on the asymmetric high-temperature tail of the TPD trace. When these TPD results are assessed in conjunction with the FTIR results presented in Figure 6c, it can be argued that the activated 5Fe/8Ba/Al surface contains well-dispersed BaO and Fe₂O₃/Fe₃O₄/FeO_x domains which are readily accessible for NO_x adsorption. This is also consistent with the significant suppression of the major TPD feature that belongs to the nitrates on the exposed alumina surface (625 K). Therefore, it is apparent that nitrates located on the Fe and Ba domains have different thermal stabilities where the nitrates stored on the additional Fe domains are more stable than the surface nitrates on the well-dispersed BaO units by ca. 80 K.

Along these lines, the TPD profile for the 10Fe/8Ba/Al surface can also be analyzed in detail. The FTIR results for the NO₂ adsorption on the 10Fe/8Ba/Al surface given above (Figure 6e) as well as the structural characterization data point to the fact that the Fe-containing domains on the 10Fe/8Ba/Al surface exist as relatively large crystallites which have a relatively limited access to NO₂(g). This observation is in very good agreement with the corresponding TPD results given in the topmost desorption profiles in Figure 9. In Figure 9a, it is clearly seen that the desorption feature at 760 K corresponding to the bidentate nitrates on the Fe domains is significantly suppressed while the intense desorption features associated with the surface nitrates on the BaO units (685 K) as well as the signal related to the nitrates on the bare alumina support surface (625 K) are noticeably visible. Thus, an overall interpretation of these observations implies that, on the 10Fe/8Ba/Al sample, Fe domains agglomerate on the surface to form large crystallites with a low surface to volume ratio. These crystallites are

relatively inaccessible for NO₂ adsorption and leave a significant number of available adsorption sites on the surface Ba nitrate sites and on the bare Al₂O₃ support surface. The presence of a large number of available adsorption sites on the Al₂O₃ support surface in the 10Fe/8Ba/Al system is also consistent with the large TPD signal at 625 K in the topmost TPD profile in Figure 9a, in addition to the comparable surface areas of the 10Fe/8Ba/Al and 5Fe/8Ba/Al systems.

3.8. Thermal Behavior of the Adsorbed NO_x Species on the 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al Systems via TPD. Figure 10 summarizes the TPD data obtained after the saturation of the γ -alumina, 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al surfaces with NO₂(g) at 323 K. The TPD profile corresponding to the 20Ba/Al sample is dominated by an intense, high-temperature desorption feature at 907 K, which can be attributed to the desorption of the bulk nitrates from the Ba domains. Furthermore, this bulk nitrate feature desorbs mostly in the form of NO + O₂.^{38,40} In addition to this dominant feature, the asymmetric low-temperature tail can be attributed to a convolution of the signals originating from the surface nitrates on Ba domains (650–780 K), nitrates on the alumina support (600–650 K), and weakly bound N₂O₃/NO⁺ species (ca. 389 K).

The TPD profiles given in Figure 10 regarding the 5Fe/20Ba/Al surface present noteworthy differences when compared to the 20Ba/Al case. It is apparent that the addition of 5 wt % Fe to the 20Ba/Al system results in a significant thermal destabilization of the adsorbed NO_x species. Considering the lack of a very significant amount of accessible Fe domains available for NO₂ adsorption in Figure 7c, it can be argued that the morphology of the 5Fe/20Ba/Al surface is such that most of the bulk Ba domains are located on the top of the well-dispersed Fe domains where the access of the Fe domains to NO₂(g) is partially blocked by the Ba sites. Thermal destabilization of the bulk nitrates on the Ba domains can also be attributed to the loss of the strong interaction between the bulk Ba nitrates with the underlying Al₂O₃ support material which is replaced by a weaker interaction between the Fe domains that are located below the bulk Ba clusters. Therefore, the desorption feature at 760 K for the 5Fe/20Ba/Al TPD profile can be mostly assigned

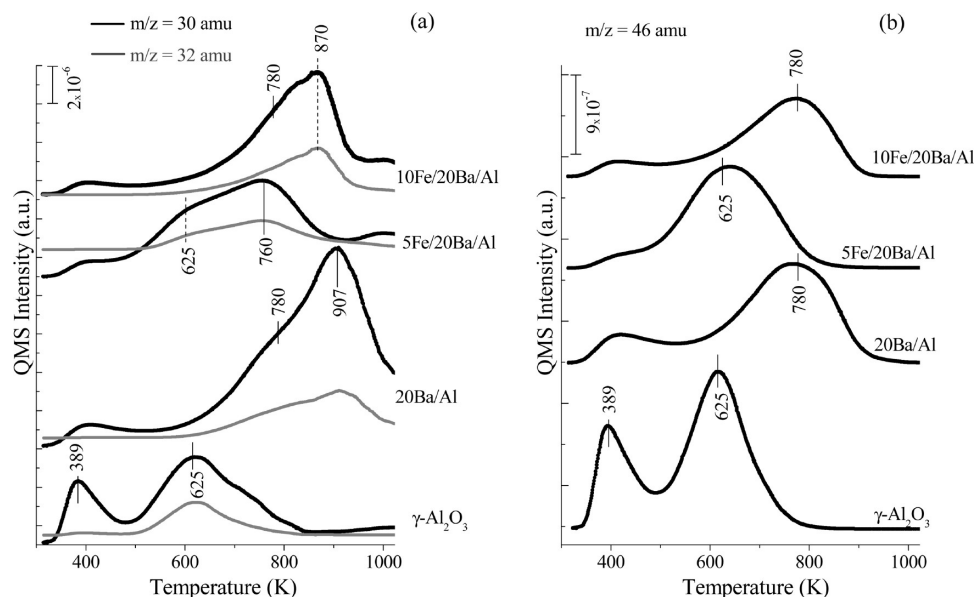
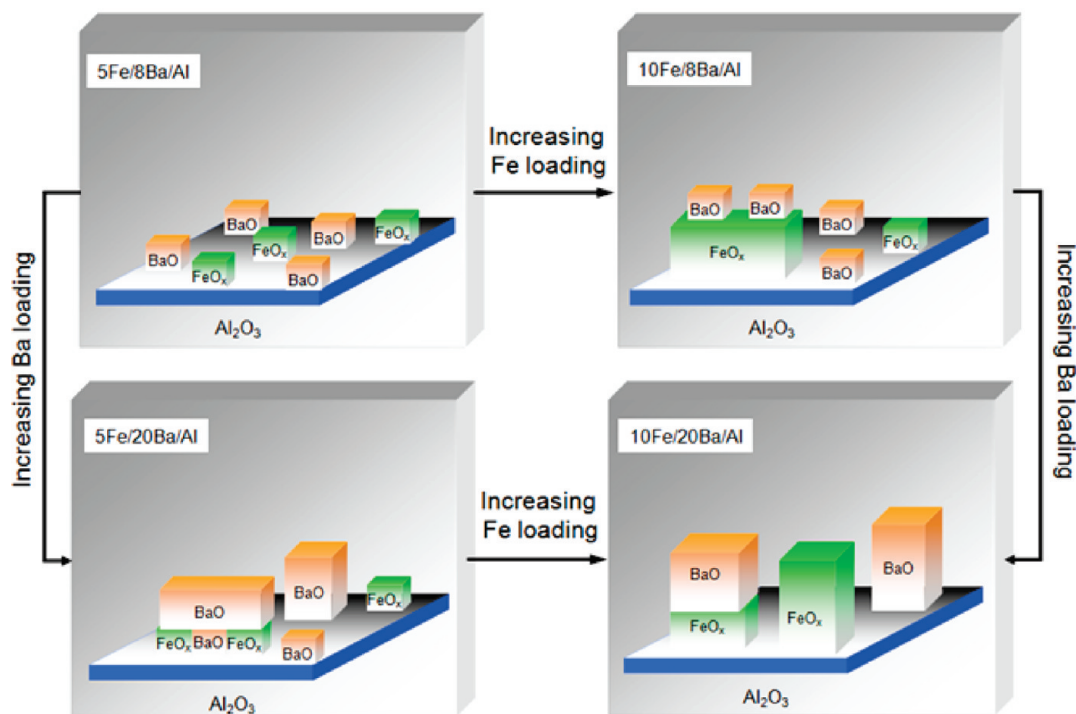


Figure 10. TPD profiles obtained from pure γ - Al_2O_3 , 20Ba/Al, 5Fe/20Ba/Al, and 10Fe/20Ba/Al samples which were initially saturated with $\text{NO}_2(\text{g})$ at 323 K: (a) 30 and 32 amu signals; (b) 46 amu signal.

SCHEME 1

A general view of the surface morphology of different NO_x Storage domains on the Fe/Ba/Al System:



to destabilized bulk Ba units that are on the top of the Fe domains (with a minor contribution from NO_x desorption due to Fe domains). This assignment is also consistent with the observation that the 760 K feature desorbs predominantly in the form $\text{NO} + \text{O}_2$, as expected from the bulk Ba nitrates. Along these lines, the lower-temperature features within 600–700 K can be associated with the surface nitrates on BaO monomers/dimers as well as the nitrates located on the bare alumina surface. These latter assignments are in harmony with the desorption characteristics of the surface Ba nitrates and the nitrates located on the bare alumina surface which yield mostly NO_2 during their thermal desorption/decomposition.

The morphology-related arguments mentioned above can also be extended to explain the TPD results for the 10Fe/20Ba/Al

system, given in the topmost spectra of Figure 10. The TPD trace for the 10Fe/20Ba/Al system shows a strong resemblance to the 20Ba/Al surface particularly in terms of its line shape. However, in the case of the 10Fe/20Ba/Al system, the major desorption feature is located at a lower temperature (870 K) with respect to that of the 20Ba/Al system (907 K). Considering the FTIR results for the 10Fe/20Ba/Al system, it can be argued that almost none of the Fe sites are available for NO_2 adsorption during neither early nor later stages of the NO_x uptake. These results can be explained by envisioning a surface morphology which is composed of mostly large and isolated Ba or Fe clusters. This explanation is also in good agreement with the results in sections 3.4 and 3.5.2 given above. Thus, the low surface to volume ratio of the large Fe clusters may render the Fe sites to be less accessible for NO_2 adsorption.

In addition, the slight thermal destabilization of the bulk Ba nitrates may also be due to the fact that some of the bulk Ba clusters are located on the large Fe₂O₃/Fe₃O₄/FeO_x crystallites, effectively blocking the access of the Fe domains to NO₂(g). Finally, it is worth mentioning that the desorption characteristics of the TPD features located at 870 and 780 K are also in good accordance with the suggested assignments, as the 870 K feature mostly desorbs in the form of NO + O₂ (consistent with the bulk Ba nitrates), whereas the shoulder within the 600–780 K feature desorbs mostly in the form of NO₂ that is in line with the surface Ba nitrates (with minor contributions from Fe nitrates and nitrates on Al₂O₃).

3.9. A General View of the Surface Morphology of Different NO_x Storage Domains on the Fe/Ba/Al System. The composition-dependent changes in the morphology of the various NO_x storage components present on the Fe/Ba/Al system that are investigated in the current text are summarized in Scheme 1.

4. Conclusions

The structure and low-temperature NO_x uptake properties of FeO_x/BaO/Al₂O₃ materials having different Fe and Ba loadings were investigated via a variety of spectroscopic, imaging, and diffraction techniques. Our results can be summarized as follows:

(a) The presence of Fe centers significantly alters the NO_x uptake mechanism at 323 K. Fe domains provide additional NO_x adsorption sites in the Fe/Ba/Al system where the adsorbed NO_x species are ultimately stored as bidentate nitrates.

(b) Freshly prepared and activated Fe/Ba/Al samples exhibit Fe^{2+(3-x)+/3+} centers whose relative abundances vary with the Fe and Ba loadings. In addition, reduced Fe centers hinder the oxidation of the adsorbed nitrites into nitrates until the complete oxidation of the reduced Fe centers into Fe³⁺. Besides, these reduced Fe sites were found to be partially regenerating under alternating oxidation/reduction treatments.

(c) Fe domains on the Fe/Ba/Al systems were found to exist in the form of Fe₂O₃, Fe₃O₄, and FeO_x whose surface morphology and the accessibility to gas phase NO_x are significantly dictated by the composition of the Fe/Ba/Al system.

(d) The decrease in the NO_x storage capacity of the Fe domains was attributed to the agglomeration of the Fe domains leading to a relatively poor Fe dispersion and/or blocking of the Fe sites by Ba domains, hindering the accessibility of the Fe domains toward gas phase NO_x adsorption.

(e) On the basis of the TPD data, the relative stabilities of the stored NO_x species can be ranked in terms of increasing stability as N₂O₃/NO⁺ < nitrates on γ-Al₂O₃ < surface nitrates on BaO < bidentate nitrates on FeO_x sites < bulk nitrates on BaO.

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Supporting Information Available: Figures showing XRD profiles, XPS measurements, ex situ Raman spectra, and temperature-dependent FTIR spectra. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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