**Mechanisms and Kinetics for Sorption of CO₂ on Bicontinuous Mesoporous Silica Modified with \( n \)-Propylamine**

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5 Supporting Information

**ABSTRACT:** We studied equilibrium adsorption and uptake kinetics and identified molecular species that formed during sorption of carbon dioxide on amine-modified silica. Bicontinuous silicas (AMS-6 and MCM-48) were postsynthetically modified with (3-aminopropyl)triethoxysilane or (3-aminopropyl)methyldiethoxysilane, and amine-modified AMS-6 adsorbed more CO₂ than did amine-modified MCM-48. By in situ FTIR spectroscopy, we showed that the amine groups reacted with CO₂ and formed ammonium carbamate ion pairs as well as carboxylic acids under both dry and moist conditions. The carboxylic acid was stabilized by hydrogen bonds, and ammonium carbamate ion pairs formed preferably on sorbents with high densities of amine groups. Under dry conditions, silylpropylcarbamate formed, slowly, by condensing carboxylic acid and silanol groups. The ratio of ammonium carbamate ion pairs to silylpropylcarbamate was higher for samples with high amine contents than samples with low amine contents. Bicarbonates or carbonates did not form under dry or moist conditions. The uptake of CO₂ was enhanced in the presence of water, which was rationalized by the observed release of additional amine groups under these conditions and related formation of ammonium carbamate ion pairs. Distinct evidence for a fourth and irreversibly formed moiety was observed under sorption of CO₂ under dry conditions. Significant amounts of physisorbed, linear CO₂ were detected at relatively high partial pressures of CO₂, such that they could adsorb only after the reactive amine groups were consumed.

**INTRODUCTION**

Recent studies indicate that certain sorbents could reduce the cost of carbon capture very significantly.\(^1\)\(^–\)\(^2\) Choi et al.\(^3\) and Hedin et al.\(^4\) recently reviewed various aspects of such adsorbents. Sorbents modified with amines have high CO₂-over-N₂ selectivity and high capacity for adsorption of CO₂ at low partial pressures, due to preferential reactions with CO₂.\(^3\)\(^–\)\(^9\) A unique feature of sorbents modified with amines is that their capacity and selectivity for CO₂ are often enhanced by water vapor.\(^7\)\(^,\)\(^10\)\(^–\)\(^13\)

The main aspects of CO₂–amine chemistry in the liquid phase are known.\(^14\)\(^–\)\(^16\) On reaction of CO₂ with amines, ammonium carboxylates (1), ammonium bicarbonates (2), and carboxylic acids (3) form (Scheme 1).\(^17\)\(^–\)\(^22\) Carboxylic acids are often stabilized as dimers or oligomers.\(^23\) Masuda et al.\(^21\) observed that carboxylic acids formed when CO₂ reacted with naphthylalkylamines in protophilic solvents; however, in protophobic solvents ammonium carboxylate ion pairs formed, and in propanol/water HCO₃⁻ formed.

For amines supported, or tethered, on solids, the detailed CO₂–amine chemistry is under investigation.\(^3\)\(^,\)\(^24\)\(^–\)\(^27\) Pinto et al.\(^27\) concluded that both carboxylic acid and ammonium carbamate ion pairs formed on amines tethered on silica by using solid-state $^{13}$C NMR spectroscopic data and their related temperature dependencies; however, Fourier transform infrared (FTIR) spectroscopy has so far been the main tool to study this chemistry. Table S1 (Supporting Information) compiles such FTIR-related findings on this chemistry.\(^3\)\(^,\)\(^24\)\(^–\)\(^26\) Nevertheless, the heterogeneous nature of these amine-modified sorbents and their complex FTIR spectra have resulted in conflicting structural assignments.\(^3\)\(^,\)\(^24\)\(^–\)\(^26\) For example, IR spectra of chemisorbed CO₂ on SBA-15 modified with (3-aminopropyl)triethoxysilane (APTES) have been understood very differently.\(^12\)\(^,\)\(^26\)\(^–\)\(^29\) Hiyoshi et al.\(^12\) assigned bands in the IR spectra to propylammonium propylcarbamate (for simplicity we use “ammonium carbamate” hereafter) ion pairs, which formed with CO₂ under both dry and moist conditions, and showed that these ion pairs became dominant when the density of amine groups was high enough.\(^12\) For related samples, Chuang and co-workers\(^29\)\(^,\)\(^30\) assigned bands in similar IR spectra to carbonates and bicarbonates on adsorption of...
CO₂ under both moist and dry conditions, a conclusion that they might have inferred from the presence of bicarbonates from reactions known to occur in aqueous amine solutions. The presence of bicarbonates was consistent with the enhanced uptake of moist CO₂ on amine-modified silica observed by several groups. Still, the hypothesized doubling of the uptake of CO₂ under moist conditions has never been reached (CO₂:NH₂ = 1 for bicarbonates and CO₂:NH₂ = 0.5 for ammonium carbamate ion pairs). Danon et al. showed very recently in a systematic study that ammonium carbamate ion pairs and silylpropylcarbamates (so-called surface-bound carbamates) formed on sorption of CO₂ on SBA-15 modified with APTES and that their relative fractions depended on both the surface coverage of amine groups and the availability of silanol groups. They could firmly exclude any formation of bicarbonates or carbonates.

In our view, IR spectra of CO₂ chemisorbed on silica modified by n-propylamine (mesocaged silica, SBA-15, MCM-48 (ref 36 and this study), AMS-6) depend more on the details in the n-propylamine coverage and less on the identity of the underlying silica substrate. In addition to the bands for ammonium carbamate ion pairs and silylpropylcarbamates (so-called surface-bound carbamates) formed on sorption of CO₂ on SBA-15 modified with APTES and that their relative fractions depended on both the surface coverage of amine groups and the availability of silanol groups. They could firmly exclude any formation of bicarbonates or carbonates.

By considering changes to the IR spectra recorded on sorption of CO₂ on silica with tethered amine groups, we will show aspects of the CO₂-sorption chemistry not described so far. We will show that ammonium carbamate ion pairs form on adsorption of CO₂ on silica with tethered amine groups under both dry and moist conditions and that bicarbonates or carbonates do not form. By time-resolved spectra we will show that significant amounts of hydrogen-bonded carbamic acid form rapidly on adsorption of CO₂, while silylpropylcarbamate (surface-bound carbamate) forms slowly and could be desorbed under moist conditions.

**EXPERIMENTAL SECTION**

**Substrates and Amine Modification.** We used substrates with interconnected structures and a cubic Ia3d symmetry. MCM-48 was synthesized with cetyltrimethylammonium bromide (0.88 g; Aldrich), water (9.981 g; Millipore), NaOH (1.15 g; Aldrich), and tetraethoxysilane (TEOS; Aldrich, 1.04 g). We used well-established procedures for synthesis, and the substrates were characterized by powder X-ray diffraction and N₂ adsorption. For the synthesis of AMS-6, N-lauroyl-1-alanine (C₁₂₃α; Nanolycia AB, Sweden) was used as the surfactant, APTES (Aldrich) as a co-structure-directing agent (CSDA), and TEOS as the silica source. In a typical synthesis, a homogeneous solution of C₁₂₃α (0.10 g) in distilled water (20 g) was kept at 80 °C for 24 h under static conditions. The surfactant solution was stirred for 10 min before APTES (0.10 g) addition, and TEOS (0.51 g) was added 3 min after APTES addition. The solution was stirred for another 15 min at 80 °C in a closed bottle. The synthesis gel was subsequently stored at room temperature under stirring for 24 h. The final synthesis mixture was kept sealed and without stirring at 100 °C for 3 days. The final pH of the synthesis mixture was 8.7. The solid product was filtered and dried at room temperature at atmospheric pressure. The molar composition of the reaction mixtures was C₁₂₃α:AMPS:TEOS:H₂O = 1:1.25: 6:7.309.1. The surfactant extraction method was as follows: 1 g of as-made AMS-6 was refluxed for 6 h in 80 g of ethanol and 20 g of HCl (37%), filtered, washed, and dried in air. The as-synthesized sample was extracted twice to remove all the surfactant. Both samples of AMS-6 and MCM-48 were calcined under static conditions in an oven isothermally at 550 °C under a stream of air for 6 h.

N-Propylamine groups were tethered and condensed to the internal surfaces of the silica. Reactions were carried out under dry nitrogen or argon in toluene obtained from a VAC solvent purifier system. Reagents were used as obtained from commercial suppliers without further purification. (3-Aminopropyl)methyldiethoxysilane (APMDES) was prepared by a method described by Sabourault et al. APTES was used as obtained from the supplier. An example for functionalization of AMS-6 with APMDES is given below (for MCM-48 and modifications with APTES see the Supporting Information).

AMS-6 was first dried in an oven for 20 h at 150 °C, 300 mg of it was suspended in toluene (18 mL), and the suspension was stirred for 30 min at 50 °C under argon in a closed tube. The tube was opened, and the internal surface of the silica substrate was hydrated by adding water (68 µL, 3.78 mmol) drop by drop. The tube was sealed, and the mixture was heated to 120 °C for 1.5 h. APMD (1.54 g, 8.03 mmol) was added, and the tube was closed and heated for 72 h at 120 °C. After being cooled to room temperature, the tube was opened, the solid was filtered and washed with toluene (50 mL), and free residual APMD was removed by Soxhlet extraction in EtOH. The solid was dried under reduced pressure overnight. Anal. C, 10.32; H, 2.38, N, 2.28.

**Elemental Analysis.** Elemental analyses of the adsorbents for C, H, and N were performed at the elemental analysis section at Santiago de Compostela University (Spain) using Fisons Instruments 1108.

**Brunauer – Emmett – Teller (BET) and Pore Analysis.** Nitrogen adsorption – desorption isotherms were recorded at –196 °C using a Micrometrics ASAP2020 volumetric adsorption analyzer. Samples were treated under dynamic vacuum conditions (<10⁻⁵ Torr) at a temperature of 130 °C for 5 h. Specific surface areas were calculated, according to the BET method, from the data recorded at P/P₀ = 0.05–0.15. The total pore volume was calculated at a relative pressure of P/P₀ of 0.988.

**Adsorption Measurements.** Adsorption and desorption isotherms of pure CO₂ were measured using a Micromeritics ASAP2020 device. Volumetric uptake of gas under equilibrium conditions was measured at two different temperatures (0 and 21 °C) from ~0.1 Torr.
Table 1. BET Specific Surface Areas, Pore Volumes, and Amine (N) Contents of the Materials

| Material          | BET specific surface area (m²/g) | Pore volume (cm³/g) | NH₂ group content (mmol/g) | NH₂ group content per nm² |
|-------------------|---------------------------------|--------------------|---------------------------|--------------------------|
| MCM-48            | 1236–1320°                      | 0.921–1.067        | 2.822                     | 1.37                     |
| MCM-48/APTES      | 310                             | 0.306              | 1.609                     | 0.73                     |
| MCM-48/APMDES     | 712                             | 0.474              | 1.628                     | 1.06                     |
| AMS-6             | 921                             | 1.046              |                           |                          |
| AMS-6/APTES       | 288                             | 0.218              | 3.467                     | 2.26                     |
| AMS-6/APMDES      | 599                             | 0.603              | 1.628                     | 1.06                     |

a Determined by elemental analysis. b Assuming a homogeneous coverage. c Two batches.

to atmospheric pressure. A low temperature was set by mixing water and ice in a slurry in the 5 dm³ Dewar. For room temperature experiments, pre-temperature-equilibrated water was used (temperature 21 ± 0.1 °C). A 100–200 mg portion of sample was used. Samples were treated under dynamic vacuum conditions at 130 °C for at least 3 h.

FTIR Spectroscopy. A Varian 670-IR spectrometer equipped with a homemade transmission cell and vacuum system25 and a mercury cadmium telluride (MCT) detector was used. A 40 mg portion of the adsorbents was pressed to self-supporting pellets with a 20 ton pressing tool (1 ton/cm², 1 min, die with a diameter of 16 mm). To investigate possible loss of porosity during pressing, pressed pellets of AMS-6 were investigated by nitrogen sorption. The data are presented in the Supporting Information. Pellets were treated under dynamic vacuum conditions (<10⁻⁶ Torr) at a temperature of 140 °C for 6 h in the cell, and IR background spectra were measured after such treatments (unless stated otherwise) with 256 scans accumulated at a spectral resolution of 4 cm⁻¹. Probe gases (pure CO₂ (>99.9%) or 20 vol % CO₂ in N₂, Linde Gas Co. (AGA), dry or saturated with 2.1% water vapor) were introduced into the cell and equilibrated at room temperature (cell temperature ∼25 °C when recording spectra). For certain experiments CO₂ was humidified to its saturation level by bubbling gas through liquid water at 23 °C. The water content of this mixture was 2.1%, which meant that the excess of water in the system increased with increasing pressure. Equilibrium IR spectra were recorded in situ when the integrals in the spectra remained constant. Along the adsorption isotherm, spectra were recorded up to pressures of 760 Torr, and spectra along the desorption branch were recorded by successively decreasing the pressure. Contributions from gaseous CO₂ to the IR spectra were compensated for by subtracting corresponding gaseous CO₂ spectra recorded in the cell (without the sample) at different pressures. Time-resolved in situ IR spectra were recorded using the same IR setup and sample preparation method. A 0.3 dm³ portion of dry or humid CO₂ was rapidly let into the cell. (The system had a buffer volume of 0.5 dm³, and the total pressure was 200 Torr.) Single-scan in situ IR spectra were recorded at a frequency of time resolution of 4 Hz with a spectral resolution of 4 cm⁻¹, corresponding to ∼14 400 spectra/h. To ensure equilibrium, additional spectra were collected every 5 min for several hours. The formation of species formed was monitored by recording the absorbance values of the corresponding bands.

Results and Discussion

The specific surface area was reduced more for MCM-48 than for AMS-6 on modification with n-propylamine, even though the degree of modification was lower. The highest degree of APTES functionalization was achieved for AMS-6. The degree of functionalization was higher for APTES than for APMDES, which is consistent with the findings of Rosenholm et al.46 Specific surface areas (BET) and pore volumes, as determined from the adsorption at P/P₀ = 0.98, were determined and are presented in Table 1. (Pore volume distributions, in the regime 2–7.5 nm domain, are presented in Figure S1 in the Supporting Information.) MCM-48 was synthesized at higher pH conditions than AMS-6, and in the absence of a costructuring agent, which could affect the degree of cross-linking in amorphous silica.

In situ IR spectra were recorded under high-vacuum conditions for MCM-48, MCM-48/APMDES, and MCM-48/APTES and are presented in Figure 1. In Figure 1a, the strongest band in the spectra, at 3745 cm⁻¹, corresponded to OH stretching of free silanol groups of MCM-48, and the broad band at ∼3600 cm⁻¹ mainly reflected internal or hydrogen-bonded silanol groups. No free silanol groups were detectable in the spectra for MCM-48/APTES in Figure 1c, but a small band remained in the spectrum of MCM-48/APMDES in Figure 1b. The broad band at ∼3500 cm⁻¹ in Figure 1b, arose from hydrogen-bonded silanol groups, and hence, silanol groups from non fully condensed APTES and APMDES appeared to be hydrogen-bonded to the tethered amine groups. This latter band was more intense for MCM-48/APMDES than for MCM-48/APTES and corresponded to more hydrogen-bonded OH groups. Bands from small amounts of adsorbed water were observed at ∼3500 and 1645 cm⁻¹ in Figure 1a for unmodified MCM-48. Hydrogen bonding affected NH₂ stretchings and shifted their frequencies to lower wavenumbers as compared to those of free amine groups,12,47–48 reflected in bands at 3373 and 3309 cm⁻¹ (stretchings), 3170 cm⁻¹ (first overtone of the bending), and 1593 cm⁻¹ (bending) for MCM-48/APMDES. These stretching frequencies were slightly lower in the spectra of MCM-48/APTES than of MCM-48/APMDES because of the higher surface coverage of amine groups in the former. Ammonium groups (propyl–NH₂⁺ · · · O–Si) were observed by the bands at 1640 and 1545 cm⁻¹ and were more intense for low surface coverage of amines. Methylene fragments were observed by bands at 2930 and 2869 cm⁻¹ (stretchings), saturated for MCM-48/APTES
and at 1470, 1445, and 1389 cm\(^{-1}\) (scissoring; deformation). Methyl groups were detected by bands at 2966 cm\(^{-1}\) (stretching) for MCM-48/APMDES. Si–CH\(_2\) moieties were detected by the band at 1412 cm\(^{-1}\). Broad bands for the overtones of the Si–O modes were observed at 2000 and 1860 cm\(^{-1}\). Note that the silica materials absorb all IR radiation below \(\sim 1250\) cm\(^{-1}\) because of the strong absorbance of Si–O vibrations. Corresponding IR spectra of AMS-6, AMS-6/APMDES, and AMS-6/APTES were practically identical to those in Figure 1 (see Figure S2 in the Supporting Information).

The uptake of CO\(_2\) on MCM-48/APTES and AMS-6/APTES was \(>2\) mmol/g at 0 \(^\circ\)C and 760 Torr, see Figure 2, which is among the highest values reported for silica modified with APTES.\(^{3,4}\) In particular, silica modified with APTES, but also to some extent those modified with APMDES, showed high uptake of CO\(_2\) at low pressures (see Figure 2). The high uptake values and weak temperature dependencies suggested chemisorption of CO\(_2\) at low pressures. Despite the fact that the amine coverage of AMS-6/APTES was approximately double that of AMS-6/APMDES (Table 1), AMS-6/APTES adsorbed \(\sim 6\) times the amount of CO\(_2\) (~0.2 vs \(\sim 1.2\) mmol/g) in the low-pressure domain of the isotherms, Figure 2c,d. This nonlinear response for the uptake of CO\(_2\) with respect to the density of amine groups is consistent with the formation of ammonium carbamate ion pairs, which requires proximity among amine groups to form.\(^{12,49,50}\) At high CO\(_2\) pressures, the isotherms were basically linear and suggestive of physisorption on both modified and nonmodified silica. Nominal amine efficiencies (adsorbed CO\(_2\) (mmol) vs number of amine groups (mmol)) for AMS-6/APTES and MCM-48/APTES at 21 \(^\circ\)C and 760 Torr were around 55% and 60%, respectively. If we propose that approximately 50% of NH\(_2\) groups transform to NH groups in the case of ammonium carbamate ion pair formation,\(^{25}\) the difference in the amine efficiencies may be explained by the physisorbed CO\(_2\).

Mechanisms of Reactions and Interactions of CO\(_2\) with Propylamines and Silica. As the details of the reactions of CO\(_2\) with propylamines are very rich, we present a proposed mechanism with a short summary. Evidence for this proposed mechanism is shown in the following sections. Ammonium carbamate ion pairs and hydrogen-bonded carbamic acids formed rapidly on contact with dry CO\(_2\). More ion pairs formed on samples with high amine densities (silica/APTES) than on those with low (silica/APMDES) amine densities. When contacted with CO\(_2\) for a significant time under very dry conditions, an additional

\[\text{Scheme 2. Proposed Mechanisms for Chemisorption of CO}_2\text{ on Silica Modified with n-Propylamine}^{\text{a}}\]

\[\begin{align*}
\text{NH}_2 &\quad \text{NH}_2 &\quad \text{NH}_2 \\
\text{+CO}_2\text{(slow)} &\quad \text{+CO}_2\text{(fast)} &\quad \text{+CO}_2\text{(fast)} \\
\text{1715 cm}^{-1} &\quad 1564,1433 \text{ cm}^{-1} &\quad \text{H}_2\text{O} \\
\text{Silylpropylcarbamate forms slowly on single amine groups where a neighboring amine group is not available. If water is present, silylpropylcarbamate does not form. IR frequencies for C=O and COO groups are given below the species. (I) and (II) are two possibilities for hydrogen bonds.}\end{align*}\]
moiety formed. We assigned it as silylpropylcarbamate, which is consistent with the findings and assignments by Danon et al.\(^{26}\) This ester did not form when water was present during the uptake of CO\(_2\). (See Scheme 2 for a representation of the species formed.) These assignments agree well with those of Danon et al.\(^{26}\) However, they did not investigate the effect of moisture on silylpropylcarbamates and the detailed dependencies for the carbonyl band in the spectral region of 1700 − 1800 cm\(^{-1}\) (depending on the CO\(_2\) coverage and density of the species on the surface) was detected. The 1715 − 1710 cm\(^{-1}\) band was discussed in some extent by Bacsik et al.\(^{25}\) elsewhere and is here assigned to silylpropylcarbamate.

The relevant infrared bands\(^{51−53}\) for chemisorption of CO\(_2\) are tabulated in Table 2. Knöfel et al.\(^{24}\) assigned similar frequencies (with different intensities of the bands) to ammonium carbamate ion pairs and carbamic acid. For hydrogen-bonded carbamic acid an intense C=O stretching at \(\sim 1700 − 1680\) cm\(^{-1}\) (depending on the CO\(_2\) coverage and density of the species on the surface) was detected. The 1715 − 1710 cm\(^{-1}\) band was discussed in some extent by Bacsik et al.\(^{25}\) elsewhere and is here assigned to silylpropylcarbamate.

Physisorbed CO\(_2\) on pure silicas was detected by various bands, shown for AMS-6 in Figure 3. The intense band for asymmetric stretching of linear CO\(_2\) \((\nu_3)\) at 2344 cm\(^{-1}\) was accompanied by that of \(^{13}\)CO\(_2\) \((\nu_3)\) at 2280 cm\(^{-1}\). A very weak band at 1382 cm\(^{-1}\) was detected for the normally IR inactive symmetric stretching mode \((\nu_1)\) of linear CO\(_2\). Combination bands \((\nu_3 + \nu_1, \nu_3 + 2\nu_2)\) were observed at 3705 and 3602 cm\(^{-1}\). At high partial pressures of CO\(_2\) a band at 2365 cm\(^{-1}\) was observed, and recently, Roque-Malherbe et al.\(^{54}\) observed such a band at 2363 cm\(^{-1}\), which they assigned to a weakly bonded adduct between silanol groups and CO\(_2\): \(-\text{OH} \cdots \text{O} = \text{C} = \text{O}\) for CO\(_2\) adsorbed on silicas. When silica was contacted with humid CO\(_2\), only bands for physisorbed water and CO\(_2\) were detected; bands for carbonates or bicarbonates were not observed as there were no

**Figure 3.** In situ FTIR spectrum for AMS-6 contacted with dry CO\(_2\) gas at 100 Torr after being treated at \(<10^{-6}\) Torr and 140 °C. Contributions from gaseous CO\(_2\) and the substrate were compensated for.

Table 2. Major IR Bands for Amine-Modified Silica Contacted with CO\(_2\)

| frequency (cm\(^{-1}\)) | assignation | group | ref |
|-------------------------|-------------|-------|-----|
| 3440                    | NH str      | ammonium carbamate ion pairs, carbamic acid (hydrogen-bonded) | 12, 24−26 |
| 3375                    | NH₂ asym str | primary amine (negative band) | 12, 31, 52 |
| 3300                    | NH₂ sym str | primary amine (negative band) | 12, 31, 52 |
| 3300−1800               | NH str, hydrogen-bonded | NH₃⁺ | 52 |
| 2920                    | CH asym str | CH₂ in propyl groups (negative band) | 51−53 |
| 2850                    | CH sym str | CH₂ in propyl groups (negative band) | 51−53 |
| 2338 (and 2365)         | CO₂ asym str | physisorbed linear CO₂ | 25, 26 |
| 2160                    | NH₃⁺ combination | silylpropylcarbamate (hydrogen-bonded) | 26 |
| 1715−1710               | C=O str (amide I) | carboxylic acid (hydrogen-bonded) | 24 |
| 1700−1680               | C=O str (amide I) | NH₃⁺ | 12, 24−26, 36, 37, 51 |
| 1633                    | NH₃⁺ asym def | NH₃⁺ | 51 |
| 1580⁺                   | NH₃⁺ asym def | carbamate (ionic form) | 12, 24−26, 36, 37, 51 |
| 1564                    | COO⁻ asym str | silylpropylcarbamate carbamic acid (hydrogen-bonded; not resolved) | 26, 52 |
| 1520                    | NH def + C−N str (amide II) | carbamate (ionic form) | 51 |
| 1484                    | NH₃⁺ sym def | carbamate (ionic form) | 26, 51−53 |
| 1480                    | NH def      | carbamate (ionic form) | 26, 51−53 |
| 1433                    | COO⁻ sym str | carbamate (ionic form) | 26, 51−53 |
| 1381                    | COO⁻ sym str (?) | carbamate (ionic form) (?) | 26, 51−53 |

\(^{a}\) High-frequency shoulder on the band at 1564 cm\(^{-1}\).
bands detectable in the region of 1700–1300 cm$^{-1}$. (Figure S4 in the Supporting Information presents an FTIR spectrum for pure silica AMS-6 contacted with moist CO$_2$ gas.)

Much more CO$_2$ physisorbed on pure silica than on amine-modified silica, but the frequencies were lower for physisorbed CO$_2$ ($\nu_3$) on modified silica, with frequencies of 2344, 2340, and 2338 cm$^{-1}$ for MCM-48, MCM-48/APMDES, and MCM-48/APTES at 100 Torr of CO$_2$. The shift in frequency could suggest stronger adsorption for physisorption of CO$_2$ on modified MCM-48 than on pure MCM-48, possibly relating to a higher degree of hydrogen bonding. By measuring the relative intensities of $\nu_3$ bands —using the Si$^-$O overtone bands as internal references and the CO$_2$ uptake of the pure silica—the amounts of physisorbed CO$_2$ on MCM-48/APMDES and MCM-48/APTES were estimated to be $\sim2\%$ and $\sim0.2\%$ of that on MCM-48 at 100 Torr of CO$_2$.

Full-scale FTIR spectra are first presented for a general understanding and quantitative comparison of the chemisorbed species on MCM-48/APMDES and MCM-48/APTES, after which we concentrate on the details in the 1800–1260 cm$^{-1}$ region. Spectra for MCM-48/APMDES and MCM-48/APTES contacted with 100 Torr of CO$_2$ are shown in Figure 4. The areas for the chemisorbed bands were much larger for MCM-48/APTES than for MCM-48/APMDES and corresponded very well with the significant differences in the uptake level of CO$_2$ in the low-pressure regime, compared with the isotherms in Figure 2. Apart from the band at 1715 cm$^{-1}$, the spectra in Figure 4 are very similar to those reported by others$^{12,29,36}$ and practically agree with spectra measured by Danon et al.$^{26}$ on silica with high coverage of amines. Here, bands were assigned to ammonium carbamate ion pairs with the corresponding frequencies presented in Table 2. Besides these ion pairs, frequencies were assigned to hydrogen-bonded carboxylic acid and silylpropylcarbamate (1715–1680 cm$^{-1}$). It was not possible to identify these latter species in single spectra due to overlapping bands, but time-resolved measurements and the absence/presence of water vapor helped us to resolve the spectra (see the details in the forthcoming discussion). Negative bands appeared in Figure 4 with signatures of CH$_2$ and NH$_2$. The negative signals for primary amines corresponded to the formation of ammonium carbamates and carboxylic acid. At this point we are unable to explain the negative bands for CH$_2$ fragments, although some suggestions have been given by Chang et al.$^{29}$ Similar spectra for AMS-6/APMDES and AMS-6/APTES are presented in the Supporting Information (Figure S5).

**Time-Dependent Study of CO$_2$ Sorption on Amine-Modified Silica.** To clarify the mechanism of CO$_2$ adsorption on propylamine-modified silica, time-dependent measurements were performed on AMS-6/APMDES. Figure 5 shows the recorded IR spectra and buildup curves for ammonium carbamate ion pairs, hydrogen-bonded carboxylic acid, and silylpropylcarbamate (dashed lines, humid; solid lines, dry CO$_2$).
sudden increase of the pressure of dry and humid CO$_2$ in N$_2$ for AMS-6/APMDES. Ammonium carbamate ion pairs formed rapidly, and within 1 min, 83% of the final amount had formed, and within 10 min, 90% had formed, both with dry and humid CO$_2$. Then adsorption slowed considerably, and the equilibrium level for these ion pairs was reached only after ~7 h. In the time-resolved measurements, the amounts of ion pairs formed were close to identical for adsorption of dry and humid CO$_2$, although this could be explained by the small amounts of water in these particular experiments. The spectra for adsorption of dry and humid CO$_2$ were indeed initially very similar and showed a very rapid formation of hydrogen-bonded carbamic acid (at 1700 cm$^{-1}$) within the first 25 s (Figure 5a,b). After this initial stage, for dry CO$_2$, the carbonyl band continued to increase slowly, while it “shifted” to higher frequencies. This shift could not be explained by H-bonding of the carbonyl groups, which would have decreased its frequency. A possible explanation for the shift is a formation of a new band on the high-frequency side of the band at 1700 cm$^{-1}$. We considered several facts to assign this new band: (i) The band formed slowly. (ii) Esters always have a higher frequency for the carbonyl group than the corresponding acids. (Stapleton et al.$^{57}$ observed a related shift for esterified acetic acid on fumed silica. The C=O frequency for that ester, silylacetate, was in the region of 1760–1740 cm$^{-1}$, while it was 1724–1717 cm$^{-1}$ for hydrogen-bonded acetic acid.) (iii) This compound was very stable; it could not be removed by evacuation or even by heat treatment. (iv) This species was observed only under dry conditions (i.e., it did not form if water was present). When treated with water, it transformed into hydrogen-bonded carbamic acid (cf. hydrolysis of the ethoxy groups).$^{58}$ Considering these facts, we assigned the band at 1715 cm$^{-1}$ to a silylpropylcarbamate (see Scheme 2) (or surface-bound carbamate using the terminology of Danon et al.$^{26}$). These observed frequencies (1715–1680 cm$^{-1}$) are rather low for both an acid and an ester, but can be explained by the effects of neighboring N atoms and the hydrogen bonds, which decrease the frequency further. No intermediate species were detected in the spectra during the full adsorption process (the time resolution was 245 ms). It strongly appeared as ammonium carboxamates did not transform to hydrogen-bonded carbamic acid or silylpropylcarbamate. The bands corresponding to ammonium carboxamates did not decrease during the slow buildup of carbamic acid. For dry CO$_2$, the rate of formation of ammonium carbamate ion pairs slowed after 0.5 min, but silylpropylcarbamate continued to form until the experiment was aborted; see Figure 5c.

Kinetically resolved difference spectra for dry CO$_2$ adsorbed on AMS-6/APMDES are presented in Figure 6. IR spectra recorded after 2 and 60 min are presented in Figure 6a. Their difference spectrum in Figure 6b related, mainly, to slowly forming silylpropylcarbamates. The difference spectrum in Figure 6c is the difference between the solid-line spectrum of Figure 6a and the difference spectrum in Figure 6b. In Figure 5c, a carbonyl band remained at 1687 cm$^{-1}$, which indicated that there were two species with carbonyl bands in this region. The one with a band at 1687 cm$^{-1}$ formed relatively rapidly, could be removed by evacuation easily, and formed with moist CO$_2$ as well (as will be shown in the next paragraphs). We assigned this band to hydrogen-bonded carbamic acid. (In the liquid state, functional group frequencies for naphthyl alkylcarbamic acid at room temperature$^{21}$ are 1700, 1545, and 1250 cm$^{-1}$; for methylvcarbamic acid at low temperature$^{20}$ they are 1680, 1533, and 1263 cm$^{-1}$. Tseng et al.$^{59}$ assigned a carbonyl band at 1677 cm$^{-1}$ to similar structures on ethanolamine-modified TiO$_2$ reacted with CO$_2$.) The presence of the hydrogen bonding was confirmed by the negative OH bands at around 3675 cm$^{-1}$ (the corresponding positive, lower frequency and broader H-bonded OH band cannot be observed clearly because of the overlap with NH stretchings), which can be observed due to the formation of both carbamic acid and silylpropylcarbamate (see Figure S6, Supporting Information). For the latter compound an intense NH stretching band was observed at 3445 cm$^{-1}$. Danon et al.$^{26}$ also observed a similar shift (from 1715 to 1701 cm$^{-1}$) of the silylpropylcarbamate when ammonium carbamate was present on materials with high amine density. They explained this shift with the underlying broad band for NH$_3$ at 1625 cm$^{-1}$. In our opinion, that band appeared to be too far from the carbonyl frequencies to affect it significantly. (Full-scale spectra are given in the Supporting Information, Figure S6.)

Equilibrium Study of CO$_2$ Sorption on Amine-Modified Silica. Chemisorbed species formed over the full range of partial pressure of CO$_2$ for AMS-6/APMDES and are visible in the equilibrium FTIR spectra measured in situ during adsorption and desorption. Spectra in Figure 7a show bands for hydrogen-bonded carbamic acid, silylpropylcarbamates, and ammonium carbamate ion pairs (see Table 2 for assignments). Bands at 1715 and 1518 cm$^{-1}$ (the latter frequency could not be seen in Figure 7a, but had been assigned from the time-resolved measurements) belong to silylpropylcarbamate. The band at 1518 cm$^{-1}$ (amide II) was rather intense due to overlap with the ammonium carbamate ion pair bands. Under dry conditions only the carbonyl band of the silylpropylcarbamate “shifted” from 1715 to 1703 cm$^{-1}$ on increasing the pressure; see Figure 7a. We rationalized this apparent shift by the facts that

![Figure 6. IR spectra measured during the adsorption of dry CO$_2$ on AMS-6/APMDES material: (a) after 2 min (dotted line) and after 60 min (solid line), (b) 2 min spectrum subtracted from the 60 min spectrum representing rapidly forming ammonium carbamate ion pairs and hydrogen-bonded carbamic acid. Assignations are presented in Table 2.](image-url)
the relative fractions of hydrogen-bonded carbamic acid to silylpropylcarbamate changed and that these bands are overlapping. Figure 7b shows that ammonium carbamate ion pairs could be removed by evacuation. As the pressure was reduced, the band for silylpropylcarbamate shifted back to 1715 cm$^{-1}$.

The spectra clearly showed that hydrogen-bonded carbamic acid was removed by evacuation, but the silylpropylcarbamate was not. In fact, the silylpropylcarbamate was stable to heat treatment under conditions of dynamic vacuum. Spectra in Figure 7c show a continuous formation of chemisorbed moieties on AMS-6/APMDES when raising the pressure of humid CO$_2$. The amount of carbamic acid was significantly reduced for adsorption under moist as compared with dry conditions. This difference was probably due to a water-mediated formation of ammonium carbamates from carbamic acids being hydrogen-bonded with $\text{NH}_2$ groups. Another major difference was that no signatures for silylpropylcarbamates were observed in spectra recorded under moist conditions. The band at 1715 cm$^{-1}$ was absent at
all but the very low pressures. Its presence at very low pressure could possibly have been related to mass transport limitation. The carbamate bands at 1560 and 1491 cm$^{-1}$ were well resolved for humid CO$_2$, because of the absence of the band of silylpropylcarbamate at 1518 cm$^{-1}$. At high pressures, spikelets from water vapor were detected on the spectra (1700–1400 cm$^{-1}$), indicating the excess of water vapor. The increased intensity of the band at 1640 cm$^{-1}$ related to an increased amount of physisorbed water, as compared to lower pressures. In the low-pressure domain of the desorption branch, a negative band remained at 1715 cm$^{-1}$; see the spectra in Figure 7d. This band related to hydrolysis of silylpolycarbamates that were present in AMS-6/APMDES from the very beginning of the experiment. The band at 1595 cm$^{-1}$ present during desorption, see Figure 7d, showed that under moist conditions additional primary amine groups were liberated. The amines were formed from hydrolysis of silylpolycarbamates, and possibly hydrogen-bonded ones. Liberated amine groups could explain the often observed higher uptake of moist CO$_2$ as compared with dry CO$_2$; additional available amine groups would allow more ammonium carbamate ion pairs to form. This amine band remained intense after further treating the sorbent with heat and dynamic vacuum.

Figure 8. Region of the IR spectra of MCM-48/APTES contacted with 20 vol % CO$_2$ in N$_2$ (a) at pressures of 2, 5, 10, 50, 100, 200, 400, 600, and 760 Torr (from bottom to top) in the adsorption branch and (b) at pressures of 760, 600, 400, 200, 100, 50, 10, and 2 Torr after 20 min of evacuation and after heat treatment (from top to bottom) in the desorption branch and MCM-48/APTES contacted with water-saturated 20 vol % CO$_2$ in N$_2$ (c) at pressures of 1, 5, 12, 50, 100, 200, 400, 600, and 760 Torr (from bottom to top) in the adsorption branch and (d) at pressures of 760, 600, 400, 200, 100, 50, 10, and 2 Torr after 20 min of evacuation and after heat treatment (from top to bottom) in the desorption branch.
(see the bottom trace in the collection of spectra in Figure 7d). Signatures of bicarbonates or carbonates were not seen in the spectra. Full-scale spectra for AMS-6/APMDES and spectra for MCM-48/APMDES are presented in the Supporting Information, Figures S7 and S8.

Mainly ammonium carbamate ion pairs formed on adsorption of dry or moist CO$_2$ on MCM-48/APTES. Ammonium carbamate ion pairs and hydrogen-bonded carbamic acid desorbed easily under dry conditions, and the adsorption/desorption processes were fully reversible for the ion pairs (Figure 8b). Silylpropylcarbamates also formed on MCM-48/APTES, and after 20 min of evacuation silylpropylcarbamate remained (see the remaining bands in the trace next to the bottom in Figure 8b). In contrast to the data shown in Figure 7b, the silylpropylcarbamate could be removed by heat treatment (a difference that we cannot explain at this point). After heat treatment, two bands belonging to a new and irreversibly formed species was observed at 1660 and 1550 cm$^{-1}$. Its formation was associated with a consumption of amine groups, which was reflected in the negative band at 1595 cm$^{-1}$ in Figure 8b, present in the spectra after heat treatment. Spectra recorded for CO$_2$ adsorption under moist conditions on MCM-48/APTES are presented in Figure 8c. The carbamate bands had slightly higher intensities for moist than for dry conditions. The characteristic band for silylpropylcarbamate was absent, and the band for hydrogen-bonded carbamic acid was smaller. The desorption spectra recorded with humid CO$_2$ were similar to those measured by others. Both hydrogen-bonded carbamic acid and ammonium carbamate ion pairs were easily removed, and bands corresponding to free amines appeared at 3370, 3300, and 1597 cm$^{-1}$ (full-scale spectra in the Supporting Information, Figure S10). After heat treatment, the bands at 1660 and 1550 cm$^{-1}$ remained, but with lower intensities than for the “dry” experiments. (Full-scale spectra recorded along the adsorption and desorption branches for CO$_2$ adsorption on AMS-6/APTES showed similar trends and are presented in the Supporting Information, Figure S9.)

**Regeneration of the Adsorbents.** Three repeated CO$_2$ uptake experiments on MCM-48/APMDES showed that the capacity to adsorb dry CO$_2$ was not reduced (1.150, 1.147, and 1.157 mmol/g). For each cycle, the adsorbent was regenerated by applying heat and dynamic vacuum. Sayari and co-workers and Tanthana and Chuang showed that several cycles were needed to show capacity loss for CO$_2$ sorption on related sorbents. In a previous paper, we discussed that a small amount of irreversibly formed products may build up during adsorption of CO$_2$. This buildup could lead to a significantly reduced capacity during repeated cycles. Sayari et al. recently showed that irreversibly formed species remained on silica modified with $n$-propylamines after repeated sorption of CO$_2$. They proposed that urea formed at high temperatures and that water could hinder urea formation. We observed such compounds in Figures 7 and 8 without repeated cycling; however, this was not very surprising as we allowed much longer equilibration times than Sayari et al. This compound was reflected in FTIR bands at 1660 and 1550 cm$^{-1}$ and a “half-band” at around the cutoff (1260 cm$^{-1}$) already after one cycle and a band at around 3400 cm$^{-1}$. The assignments of these bands are C=O stretching (amide I), NH deformation and CN stretching (amide II), CN stretching (amide III), and NH stretching, respectively. This species appears to have been an amide, possibly a urea. This compound could not be removed during regeneration with dynamic vacuum and heat treatment at 140 °C. Furthermore, the appearance of negative bands corresponding to primary amines indicated that amine groups were consumed. In addition, negative bands corresponding to OH groups were observed, perhaps due to further condensation of silanol groups (forming water and siloxane linkages). (The full-scale IR spectra after the first and several cycles are given in the Supporting Information, Figure S11.)

**4. CONCLUSIONS**

On silica modified with $n$-propylamines, three major chemisorbed moieties formed on adsorption of dry CO$_2$: propylammonium propylcarbamate ion pairs, hydrogen-bonded propylcarbamic acids, and silylpropylcarbamates. Silylpropylcarbamate formed by slow condensation of carbamic acid with silanol groups under very dry conditions, but did not form if water vapor was present. As expected, a large amount of ammonium carbamate ion pairs formed when the density of amine groups was high. Kinetic adsorption experiments showed that ammonium carbamate ion pairs and carbamic acid formed rapidly, which is important for eventual applications in CO$_2$ capture. The ammonium carbamate ion pairs and hydrogen-bonded propylcarbamic acid were weakly chemisorbed and could be removed by applying dynamic vacuum. No bicarbonates or carbonates formed under moist conditions, and the notation that the uptake of CO$_2$ can be doubled with water on this kind of amine-modified silica is very questionable. Instead, we detected additional free amine groups in the presence of water vapor, and more ammonium carbamate ion pairs formed, which can rationalize the often observed small increase of the uptake of humid CO$_2$ as compared with dry CO$_2$. A small amount of irreversibly chemisorbed species, plausibly ureas, formed as well.

**ASSOCIATED CONTENT**

* Supporting Information. Functionalization of the silica, table for FTIR detection of formed species on CO$_2$ adsorption, pore volume distributions, and in situ FTIR spectra of the formed species on CO$_2$ adsorption on amine-modified AMS-6 and MCM-48. This material is available free of charge via the Internet at http://pubs.acs.org.

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