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Probing the surfaces of interstellar dust grains: the adsorption of CO at bare grain surfaces

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ABSTRACT

A solid-state feature was detected at around 2175 cm⁻¹ towards 30 embedded young stellar objects in spectra obtained using the Infrared Spectrometer and Array Camera at the European Southern Observatory Very Large Telescope. We present results from laboratory studies of CO adsorbed at the surface of zeolite wafers, where absorption bands were detected at 2177 and 2168 cm^{-1} (corresponding to CO chemisorbed at the zeolite surface) and 2130 cm^{-1} (corresponding to CO physisorbed at the zeolite surface), providing an excellent match to the observational data. We propose that the main carrier of the 2175-band is CO chemisorbed at bare surfaces of dust grains in the interstellar medium. This result provides the first direct evidence that gas-surface interactions do not have to result in the formation of ice mantles on interstellar dust. The strength of the 2175-band is estimated to be $\sim 4 \times 10^{-19}$ cm molecule⁻¹. The abundance of CO adsorbed at bare grain surfaces ranges from 0.06 to 0.16 relative to H_2O ice, which is, at most, half of the abundance (relative to H_2O ice) of CO residing in H₂O-dominated ice environments. These findings imply that interstellar grains have a large (catalytically active) surface area, providing a refuge for interstellar species. Consequently, the potential exists for heterogeneous chemistry to occur involving CO molecules in unique surface chemistry pathways not currently considered in gas grain models of the interstellar medium.

Key words: astrochemistry – line: identification – methods: laboratory – dust, extinction – ISM: lines and bands – infrared: ISM.

1 INTRODUCTION

Interstellar dust grains are thought to consist of an amorphous silicate or carbonaceous core surrounded by a molecular ice layer (Draine 2003; Gibb et al. 2004). Surface reactions, on and in the icy mantle, or on the bare grains, are key routes to forming many molecules observed in star-forming regions (van Dishoeck & Blake 1998; Herbst 2000). Presently, little is known about the morphology or chemistry of these grain surfaces, or the porosity of the grains themselves. From observational evidence of star-forming regions and cometary dust, interstellar silicates are most regularly associated with amorphous olivine or pyroxene (Henning 2003), i.e. silicates incorporating Mg and Fe in their crystal structure. Studies of pre-solar nebula inclusions in meteoritic samples suggest that these grains may contain embedded metals or sulphides, even in their interstellar state (Bradley et al. 1999).

Spectral features in the 2000-2200 cm⁻¹ region are characteristically associated with stretching vibrations of doubly and triply bonded molecules. Pontoppidan et al. (2003) have performed an extensive survey in this spectral region with the Infrared Spectrometer and Array Camera (ISAAC) at the European Southern Observatory (ESO) Very Large Telescope (VLT), of around 40 low-mass (<50 L_{\odot}) and intermediate-mass (>50 L_{\odot}) embedded young stellar objects (YSOs). The spectra are dominated by a broad absorption feature, corresponding to condensed CO (CO ice) in pure CO-dominated (2139.9 and 2143.7 cm⁻¹) and H₂O-dominated (2136.5 cm⁻¹) environments (Tielens et al. 1991), with rovibrational transitions of gas-phase CO superimposed, either in absorption or emission (Boogert et al. 2002; Pontoppidan et al. 2002). Analysis of these data revealed a second, weak, interstellar feature in 30 of the objects, as illustrated (for six of these objects) in Fig. 1. The band's peak position ranged from 2165 to 2194 cm⁻¹, with an

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Figure 1. Detailed fits to six spectra of YSOs where the unidentified 2175-cm^{-1} band was observed with the ESO VLT ISAAC (Pontoppidan et al. 2003): (a) Reipurth 50, (b) TPSC78, (c) IRS63, (d) HH100 IR, (e) EC 90A, (f) Elias 32. For clarity, the scale is set to truncate the intense pure CO ice feature at 2139 cm^{-1} and this band is excluded from the fit. The overall fit (grey solid line) comprises the laboratory data shown in Fig. 2 (light-grey dashed line), and a Gaussian component, fixed at 2165 cm^{-1} , representing the 'XCN' feature, attributed to the presence of OCN⁻ in interstellar ice (light-grey dotted line). Observational data are shown in black; all observations show a combination of broad solid state and sharp narrow gas-phase CO features. Fits to Elias 32 and HH 100 IR appear to fit poorly in the 2152-cm^{-1} region due to the presence of the bright Pf β hydrogen recombination line at 4.687 µm.

average peak-centre of $2173 \pm 4 \text{ cm}^{-1}$, except in the 'higher'-mass stars in the sample (e.g. Reipurth 50) and those low-mass stars located near to the Trapezium O-stars in Orion (e.g. TPSC 78) whose average peak-centre position was $2167 \pm 2 \text{ cm}^{-1}$ (see table 4 in Pontoppidan et al. 2003 for full details). Its full width at half-maximum (FWHM) ranged from 9 to 36 cm⁻¹. Henceforth in this paper, this feature is referred to as the 2175-band.

In this paper, we assign the 2175-band to CO adsorbed directly on the surfaces of bare interstellar grains. In Section 2 we address the possible carriers of the 2175-band, drawing on previous literature describing spectral features in this region. We also explain why OCN⁻, dangling Si–H and C–D stretching vibrations were eliminated as potential carriers of the band. Laboratory experiments were conducted under high-vacuum (HV) conditions, where zeolite wafers were exposed to CO gas, at surface temperatures from 300 to 100 K, to illustrate that when CO is chemisorbed on specific binding sites at the zeolite surfaces, characteristic spectral features are observed between 2180 and 2165 cm⁻¹ (see Sections 3 and 4). An excellent fit was obtained between these laboratory spectra and the 2175-cm⁻¹ feature in the observations of Pontoppidan et al. (2003) (see Section 4). The astrophysical implications of these results are addressed in Section 5.

2 POTENTIAL CARRIERS OF THE 2175-BAND

The 2175-band is unique. The vibrational modes of CO molecules trapped in low-temperature ices invariably range from 2135 to 2155 cm⁻¹ (Fraser et al. 2004). In several studies of COcontaining interstellar ice analogues, including for example CH₄, CH₃OH, HCOOH, CO₂ or H₂O (Sandford et al. 1988; Schmitt, Greenberg & Grim 1989; Gerakines et al. 1995; Ehrenfreund et al. 1996, 1997a,b; Palumbo 1997; Collings et al. 2003a,b; Fraser et al. 2004), no CO ice band has ever been observed as far into the blue as 2175 cm⁻¹. In spectra of pure CO ices, a combination band (arising from the fundamental vibrational and translational modes within the solid) appears as a weak, broad feature centred at 2200 cm^{-1} , with an FWHM of 80–100 cm⁻¹ (Ewing & Pimentel 1961), significantly broader than the interstellar feature. To observe this band in the laboratory with transmission infrared spectroscopy requires very thick samples of CO (>500 monolayers) in which the majority of the CO ice is crystalline; attempts to fit such laboratory features to the observations were unsuccessful, ruling out the combination band as a feasible carrier. The optical depth of the 2175-band does not scale with that of the CO or H₂O ice bands in these objects (van Broekhuizen et al. 2004a), although a tenuous correlation was found between the optical depths of the 2175-band and the 2136.5-cm⁻¹ component of the CO ice band (CO trapped in a H₂O-dominated environment; Pontoppidan et al. 2003), suggesting that both features may share the same, or a closely related, chemical species.

In spectra of high-mass star-forming regions, a band has sometimes been detected at around 2165 cm⁻¹ associated with the CN-stretching vibrations of the OCN- ion (e.g. Pendleton et al. 1999; Hudson, Moore & Gerakines 2001; Gibb et al. 2004; van Broekhuizen, Keane & Schutte 2004b). However, the peak of this 2175-band is clearly shifted from 2165 cm⁻¹. Although more difficult to ascertain because of spectral confusion, the optical depth of the 2175-band does not correlate with the optical depth of any 'XCN'-type component of the band (van Broekhuizen et al. 2004a). Even the most blueshifted laboratory spectra of solid OCN- reported in the literature peak at only 2170 cm⁻¹ (Raunier et al. 2003), as do CN-stretching vibrations in nitriles and isonitriles (Pendleton et al. 1999); alone, all are inadequate for fitting the 2175-cm⁻¹ peak position and band profile (van Broekhuizen et al. 2004a). For these reasons, solid OCN- was eliminated from this study as the primary carrier of the 2175-band.

These deductions, coupled with the detection of the 2175-band in cold, low-mass protostellar sources where ices are not thought to be energetically processed (Langer et al. 2000), suggest that the 2175-band should be attributed to a solid-state feature that is distinct from the ice mantles accreted in dense clouds. One carrier candidate could be the dangling Si–H bonds formed at silicate surfaces when they condense in the presence of H₂ (Blanco, Fonti & Orofino 1999). Dangling Si–H bonds give rise to very strong, broad, absorption features between 2200 and 2100 cm⁻¹, depending on the silicate, which have significantly broader FWHM (\approx 150 cm⁻¹) than the profile of this 2175-band. Similarly, infrared absorption spectra of hydrogenated amorphous carbon (HAC), particularly deuterated HAC, also exhibit strong, broad features between 2000 and 2250 cm⁻¹, associated with C–D and C–H stretching bands (Grishko & Duley 2003). However, once again such features are significantly broader (FWHM $\approx 200 \text{ cm}^{-1}$) than the 2175-band observed here, and would have to be accompanied by additional, even stronger and distinctive features between 1300 and 800 cm⁻¹, which have never been detected in the sources mentioned here for which complementary observations exist (Alexander et al. 2003). For these reasons C–D/C–H and Si–H bonds were also eliminated from this study as potential carriers of the 2175-band.

CO is frequently used as an infrared probe to characterize the surfaces of oxides and zeolites (Hadjivanov & Vayssilov 2002), particularly to identify the active catalytic binding sites on such materials. Zeolites can be thought of as 'exceptional' silicates, with some Si replaced by Al; geologically they are formed by processing of silicates in aqueous environments. Binding sites, corresponding to CO chemisorbed at Brønsted acid-type sites (Al₂O₃, Al-OH), Si-OH-Al or at cation inclusions on the surfaces of these materials, have characteristic frequencies between 2160 and 2180 cm⁻¹ (Gruver, Panov & Fripiat 1996); physisorbed CO is observed at around 2136 cm⁻¹ (Hadjivanov & Vayssilov 2002). This does not mean to suggest that zeolites are present in interstellar space, although they have been identified in meteorites (Woltzka & Wark 1982), but that similar binding sites could exist on interstellar grains, where Al would be replaced by Mg or Fe. The binding sites associated specifically with absorptions at 2175 cm⁻¹ correspond to CO adsorbed at cation sites, e.g. Mg²⁺, Fe³⁺, and have also been found on cation-doped silicate surfaces (Hadjivanov & Vayssilov 2002). As such spectral features are weak and narrow, associated with secondary (physisorption) features at around 2130 cm^{-1} , and even require the same carrier species (CO) as the 2136.5-cm⁻¹ feature (itself associated with CO trapped in a H₂O-dominated matrix), these interactions were adopted as the most likely carrier of the interstellar 2175-band.

3 EXPERIMENTAL METHOD

To test the hypothesis that the 2175-cm⁻¹ band is related to CO adsorbed directly on 'bare' interstellar grain surfaces, a synthetically pure sample of a naturally occurring zeolite, caged clinoptilolite [(K₂ Ca₂ Na₂)O-Al₂O₃-10SiO₂-8H₂O] was exposed to CO gas in our laboratory, under pseudo-interstellar conditions. This material was used to 'prove a concept', being readily available, known to possess the binding sites of interest, with a caged structure (i.e. large surface area), and relatively easy to press into a 'pure' wafer of material. 200 µm thick, ≈1-cm diameter wafers of pure clinoptilolite were produced by high-pressure compression of the powdered material. These were then secured on to a specifically designed goldcoated, oxygen-free high-conductivity copper holder, mounted on an He-cooled cold finger, capable of reaching temperatures as low as 10 K. The whole system was positioned at the centre of a HV chamber, with base pressures of 1×10^{-7} mbar, described in detail elsewhere (Gerakines et al. 1995). The sample was baked overnight to over 400 °C to remove its water of crystallization, then allowed to cool to room temperature, and pumped for a further 24 h, to ensure the surface did not become rehydrolysed. The sample was then cooled from 300 to 100 K in a CO atmosphere of a few Torr, to ensure (i) a constant dynamic monolayer coverage of CO at the zeolite surface, (ii) maximum occupancy of the chemisorption sites, and (iii) that any vacuum contaminants (especially H₂O), which may preferentially occupy or freeze-out on the wafer surface under rare vacuum conditions, were displaced. 0.5-cm⁻¹ resolution Fourier transform infrared (FTIR) transmission spectra were recorded prior and during CO exposure, at surface temperatures from 300 to



Figure 2. (a) Laboratory spectrum of CO adsorbed on the internal and external surfaces of a clinoptilolite wafer (solid black line) at 100 K. Substructure and line centres of the absorption feature components were obtained from a derivative plot of the spectrum (not shown). A L–M non-linear least-squares regression fit to the spectrum is shown as a solid grey line. The main peak comprises three components at ~2177 cm⁻¹ (dash-dotted line; assigned to CO chemisorption at cation and Si–OH–Al sites), 2168 cm⁻¹ (dashed line; assigned to CO chemisorption at Al–OH/Al₂O₃ sites) and 2130 cm⁻¹ (dotted line; assigned to CO physisorption); weak features are also observed at 1870, 1887 and 1900 cm⁻¹. (b) The locations of various CO binding environments within and on the caged structure of clinoptilolite are indicated, and the chemical structure of each binding site shown schematically.

100 K. The surface temperature was never lowered beyond this value to avoid CO condensation to form CO ice, which occurs at surface temperatures of around 30 K under these experimental conditions (Fraser et al. 2004). Although the zeolite wafer is transparent in the 2200–2000 cm⁻¹ region, spectra of the wafer prior to CO exposure were subtracted from the experimental data to ensure that any spurious or temperature effects from the zeolite spectrum did not influence the final spectrum.

4 RESULTS

4.1 Laboratory spectra

Three absorption features, related to CO adsorbed at the surface of the zeolite wafer, were observed from 300 to 100 K. The 100-K spectrum is shown in Fig. 2(a). Substructure and line centres of the absorption feature components were obtained from a derivative plot of the spectrum. Gaussian components at 2177, 2168 and 2130 cm⁻¹ (plus three minor components at 1870, 1887 and 1900 cm⁻¹) were required to make a Levenberg–Marquardt (L–M) non-linear least-squares regression fit to the absorption spectrum (dark-grey trace, Fig. 2a). The strongest of these component features were assigned, by comparison with reported CO vibrational frequencies on zeo-lite surfaces (Datka et al. 1999), to physisorbed CO (2130 cm⁻¹), and chemisorbed CO, at either Brønsted acid-type binding sites (2168 cm⁻¹), or a combination of the Si–OH–Al and cation binding

Table 1. Distribution of the available binding sites in caged clinoptilolite versus their occupation in these experiments.

Binding site	$\nu(CO)_{ads}$ $(cm^{-1})^a$	Per cent of sites on wafer surface	Number of occupied surface sites in these experiments	
Gas ^b	2143	-	_	
Ice ^b	2139	-	_	
Al–OH, Al ₂ O ₃	2169	12.6	6.3×10^{16} (2168 cm ⁻¹ component)	
Si-OH-Al	2175	64.7	(2100 cm component)	
Na ⁺	2178		9.5×10^{16}	
K^+	2180	22.7	(2177 cm ⁻ component)	
Ca ²⁺	2177	JJ		
Mg ²⁺	2178 ^c	_	_	
Fe ²⁺	2180 ^c	_	-	

^aDatka et al. (1999).

^bIncluded for comparison only.

^cNot relevant to this zeolite but included for interstellar relevance.

sites (2177 cm^{-1}) ; see Table 1. Illustrations of these binding sites, and their locations within or on the zeolite structure are shown in Fig. 2(b). No obvious distinction could be made between the spectra of the chemisorbed bands recorded at different temperatures,

although the weak physisorbed CO feature shifted from 2120 to 2130 cm⁻¹ as the surface temperature was lowered. As expected, no 'pure' CO ice features (around 2139 cm⁻¹) were observed.

4.2 Estimating the band strength of the 2175-cm⁻¹ laboratory feature

In the absence of published absorption coefficients for CO adsorbed on zeolite, the results from Section 4.1 were used to estimate the occupancy of CO molecules at the binding sites on the wafer surface, assuming that a dynamic monolayer of CO was fully covering all the available (internal and external) surface area – a reasonable assumption under these experimental conditions. The total surface area of each wafer was estimated to be around 500 cm², given that the specific surface area of clinoptilolite is $20-30 \text{ m}^2 \text{ g}^{-1}$ (Yasyerli et al. 2002) and the mass of one wafer was approximately 2 mg. It is important to note that this total surface area is significantly greater than the geometric surface area of the wafer itself ($\approx 1 \text{ cm}^2$). Assuming a binding site density of 1×10^{15} binding sites cm⁻², this equates to a total of 5 $(\pm 1) \times 10^{17}$ binding sites over the whole wafer surface, which will be distributed according to the stoichiometry of the zeolite, i.e. 12.6 per cent Brønsted acid-type binding sites (Al-OH, Al₂O₃), 64.7 per cent silicate-OH binding sites (Si-OH-Al) and 22.7 per cent cation sites (see Table 1). Assuming only one CO molecule was adsorbed at each binding site, and that the absorption coefficient of CO at each of the chemisorption sites is approximately equivalent (again a reasonable assumption as there is very little difference in the vibrational frequencies of CO adsorbed at the different sites, indicating that there is almost no change in the strength of the C \equiv O bond), one would expect the integrated absorbencies of the fitted components in the absorption spectrum to track the relative abundances of the binding sites at the wafer surface, i.e. around seven times as many (cation + silicate) binding sites on the surface in comparison to Brønsted acid-type binding sites. However, as can be seen from the fitted components shown in Fig. 2(a), the integrated intensity of the 2168-cm⁻¹ fitted component is almost equivalent to that of the 2177-cm⁻¹ component. Furthermore, some CO is observed to be physisorbed to the surface, so it is likely that not all the chemisorption sites are populated. Therefore, assuming that the Al₂O₃ and Al-OH binding sites are fully occupied, but that the cation and Si-OH-Al sites are not (fully occupied), it was estimated that over the total surface area of the wafer \approx 1.6 \times 10¹⁷CO molecules were occupying surface chemisorption sites and contributing to the laboratory spectrum, as detailed in Table 1. This corresponds to a chemisorbed CO density of 3.2×10^{14} molecules cm⁻², with a band strength A of $\sim 4 \times 10^{-19}$ cm molecule⁻¹, given that the optical depth of the laboratory spectrum at 2175 cm^{-1} is \sim 0.06. This number, uncertain by a factor of about 3, is consistent with the concept that the band is weak, but accounts for a significant population of CO molecules on a geometrically small wafer, which due to its morphology has a very large total surface area.

As this experiment was conducted in a CO gas environment, the remainder of the binding sites at the surface (of the whole wafer) were occupied by a dynamic monolayer of physisorbed CO molecules evinced by the (weak) 2130-cm⁻¹ component absorption feature in Fig. 2, and possibly a few chemisorbed H₂O molecules that were not desorbed during the wafer preparation (see Section 3). At least as many, and probably more CO molecules would have been physisorbed ($\leq 3.4 \times 10^{17}$) than chemisorbed ($\approx 1.6 \times 10^{17}$) to the wafer surface, illustrating, as expected, that the absorption coefficient of physisorbed CO is significantly less than that of chemisorbed CO. Under these experimental conditions, the lifetime of physisorbed CO at the wafer surface will be less than 20 ns; consequently, if the CO gas environment is removed, the physisorption band will also disappear. In interstellar regions, at grain temperatures above the condensation temperature of CO, a CO molecule may occupy a physisorption site, but never for long enough that a significant physisorbed population would amass, so the 2120– 2130 cm⁻¹ band will never be observed. As is already well established in the literature, below the condensation temperature of CO, CO ice may form, with its characteristic vibration at around 2139 cm⁻¹. No further discussion of physisorbed CO is made in this paper.

4.3 Comparison with observations

Fig. 1 shows detailed comparisons between the laboratory COzeolite spectra (grey dashed line) and six of the YSOs observed by Pontoppidan et al. (2003) (black line). In certain objects, such as IRS63, HH100 IR and Elias 32 (Figs 1c, d and f, respectively), only the CO-zeolite spectrum makes a significant contribution to the 2175-band, and the laboratory spectrum can be scaled to provide an excellent match to the observational profile. However, in many cases it seems a second carrier contributes to the 2175-cm⁻¹ profile, so each spectrum was subsequently fitted (grey solid line) by a combination of the laboratory CO-zeolite spectrum (from Fig. 2), plus a Gaussian fixed at a centre position of 2165 cm⁻¹ and with FWHM of 25 cm⁻¹ (grey dotted line), representative of laboratory OCN⁻ spectra (Hudson et al. 2001; van Broekhuizen et al. 2004b). For convenience, the intense CO ice band was excluded from the fit. Details of the fitting of this band and a combined fit to the 2175-cm⁻¹ and CO ice features can be found elsewhere (Pontoppidan et al. 2003; van Broekhuizen et al. 2004a). As can be seen from Fig. 1, no OCN- contribution was required to fit the 2175-band profile in the low-mass sources, e.g. IRS63, HH100 IR or Elias 32. Along lines of sight that are heated or possibly subject to strong ultraviolet radiation from nearby objects, e.g. Reipurth 50 (Fig. 1a) and TPSC 78 (Fig. 1b), the OCN- feature contributes more significantly to the overall profile, and in these cases the weak interstellar feature peaks closer to 2165 cm⁻¹. In Fig. 1(e), EC90a is an example of an object where the CO gas grain-surface interaction and OCNfeature contribute almost equally to the overall band profile.

5 ASTROPHYSICAL IMPLICATIONS

5.1 Adsorbed CO abundance

The column density associated with CO adsorbed directly at grain surfaces along any one observational line of sight (henceforth called the CO gas grain or CO_{gg}), was estimated (within a factor of about 3) from the optical depth at 2175 cm⁻¹ of the scaled laboratory CO_{gg} spectrum required to fit the observational data, assuming a band strength (+ estimated error) equivalent to that derived in Section 4.2. These data, shown in Table 2, range from 1.1 to 3.2×10^{17} cm⁻², suggesting that about twice as many CO molecules are found in water-rich ice environments compared to those found on bare grain surfaces. In Table 2, the abundances (relative to H₂O ice) of pure CO ice, CO in a H₂O-dominated ice matrix and CO_{gg} are also compared. The relative abundance of CO_{gg} is reasonably constant between all the YSOs. No relationship is apparent between the relative abundances of CO_{gg} and pure CO or H₂O ice. This lack of scaling with ice column density is addressed further in Section 5.2.

CO may adsorb at bare grain surfaces arising, for example, (i) where ices have not yet accreted or formed (see Fig. 3a), or (ii)

Source	COgg		Pure CO ice		CO in H ₂ O-rich ice	
	Column density ×10 ¹⁷ cm ⁻²	Relative abundance $(to H_2O)^a$	Column density ×10 ¹⁷ cm ⁻²	Relative abundance $(to H_2O)^a$	Column density $\times 10^{17} \text{ cm}^{-2}$	Relative abundance (to H ₂ O) ^a
Reipurth 50	3.2	0.06	3.4	0.07	7.3	0.14
TPSC 78	2.1	0.08	0.8	0.03	3.7	0.14
IRS 63	2.6	0.16	11.2	0.68	4.9	0.29
HH100IR	2.9	0.12	8.8	0.37	4.4	0.18
EC 90A	1.1	0.08	13.9	1.07	4.4	0.34
Elias 32	24	0.15	11.7	0.75	6.0	0.44

Table 2. Comparison of CO column densities in ice versus bare grain environments.

^aH₂O column densities taken from van Broekhuizen et al. (2004a), using $N = 300 \times \tau (3.07 \text{ }\mu\text{m})/A_{\text{H}_2\text{O}}$ where $A_{\rm H_2O} = 2.0 \times 10^{-16}$ cm molecule⁻¹ (d'Hendecourt & Allamandola 1986).

where CO molecules migrate through the ice layer and adsorb to the underlying grain surface (see Fig. 3b). Depending on the nature of the binding sites, CO desorption enthalpies on zeolite surfaces are reported to range from \approx 2400 to \approx 12 990 K (Hadjivanov & Vayssilov 2002), with the majority of CO desorbing from sites with desorption enthalpies above ≈ 6000 K. The reported binding energies for CO adsorbed at Mg2+ and Si-OH-Al-type binding sites on doped zeolite surfaces are ≈4930-10850 K and ≈5530-7500 K, respectively (He et al. 1992; Hadjivanov & Vayssilov 2002). These values are significantly higher than the CO-CO binding energy in CO ices (860 K) or the CO-H₂O binding energy in binary ice systems (1180 K; Collings et al. 2003a,b), and (at the lowest limits) comparable with the desorption enthalpy of H₂O ice: 5600 K (amorphous) to 5570 K (crystalline; Fraser et al. 2001).

Similar behaviour to (ii) has been observed in haloform-iceplatinum systems (Greca et al. 2004). The haloform molecules diffuse 'through' pores and cracks in H₂O ice, and on reaching the interface between the ice and the supporting substrate chemisorb to the substrate rather than remaining bonded to the ice surface, as the haloform-substrate bond is an energetically more favourable state. From laboratory work (Collings et al. 2003a,b) and observational data (Pontoppidan et al. 2003), it is clear that CO ice adsorbed above H₂O ice can subsequently diffuse and become trapped within pores in the H₂O ice matrix. In the laboratory the process is initiated by heating, analogous to thermal processing in protostellar regions. However, as the energy barriers to 'diffusion of CO into H₂O ice pores' and 'CO desorption from CO dominated ice layers' are almost identical (Collings et al. 2003b), it is also possible, given the time-scales available in interstellar regions, that CO could equally migrate through the porous structure of interstellar H₂O ices and reach the ice-grain interface at grain temperatures as low as 15 K. This scenario may additionally help to explain why Pontoppidan et al. (2003) observed a weak correlation between the 2175- and 2136.5-cm⁻¹ features in the analyses of their observational spectra, because both features rely on the migration of CO molecules within the porous structure of H₂O ice.

A schematic representation of the different (postulated) ways a COgg population could accumulate at interstellar grain surfaces is shown in Fig. 3. Initially, a COgg population is accumulated in regions where ices are just starting to (or have not yet) formed, and surface formation of H₂ and the gas-phase formation of CO (from C^+ + OH) are key molecular processes (see Fig. 3a, i). From simple abundance arguments, COgg accretion must compete with other 'bare grain' processes, e.g. H_2 formation and $\mathrm{H}_2\mathrm{O}$ formation, so it is unlikely that the whole grain surface would ever become saturated by a monolayer of only CO_{gg} (see Fig. 3a, ii). As a consequence of some CO molecules being chemisorbed at the grain surface, formation of CO_2 (CO + O) and HCO (H + CO) may also be feasible (see Section 5.3). Eventually, ice mantles, dominated by H₂O, will accumulate on the grain surface. COgg acquired prior to ice formation will be more tightly bound to the grain surface than the ice layer itself (see Section 5.3) and therefore become buried under the ice layer as it grows (see Fig. 3a, iii and iv). CO may still be accreted from the gas phase, but from energetics arguments would still be more likely to seek a chemisorption binding site at the remaining regions of bare grain surface than on the ice surface itself. As the ice becomes 'thicker' the whole grain will appear to be covered by a mantle, although due to the porous nature of amorphous H₂O ice, the molecular solid would only be tethered to the grain surface at a few points, leaving much of the surface 'bare' even though incoming molecules accreting at the ice surface can no longer 'see' the grain (see Fig. 3a, iv).

Fig. 3(b) illustrates the coldest densest interstellar regions, where H₂O-dominated ices have already formed and CO is frozen out, forming CO ice; some COgg may already be buried under the ice layer, accumulated in direct gas grain interactions (see Fig. 3b, v). Over the time-scales available in interstellar regions or in regions where the grains are even mildly heated (15 < grain 'temperature')< 40 K), some of the CO will migrate along the surfaces of pores in the H2O ice. Because the pore structure is three-dimensional and interconnected, some pores will inevitably lead from the H₂O-CO ice interface to the H₂O ice-grain interface. CO cannot migrate through the H₂O ice bulk. As some of the grain surface is still 'bare', at least some of the migrating CO molecules could preferentially adsorb at the grain surface rather than the ice surface (see Fig. 3b, vi). This process continues until all the binding sites at the grain surface are occupied, or no more CO molecules can reach the bare grain surface (see Fig. 3b, vii). The COgg population becomes saturated. In this instance, the optical depth of the CO_{gg} band would also become saturated, as first observed by Pontoppidan et al. (2003). Such suggestions are entirely consistent with the fact that a COgg component is also observed on 'warmer' lines of sight to high-mass YSOs (van Broekhuizen et al. 2004a), where the COgg feature exists, but is overwhelmed by the nearby, stronger OCNband, peaking at 2165 cm⁻¹, arising from chemical processing of the ice mantle. Finally, as the binding energy of COgg is much greater than the desorption enthalpy of H_2O ice (see Section 5.3), CO_{gg} molecules remain bound to the grain surface during, and even after, ice desorption (for example, in hot cores and close to the star in protostellar regions). Consequently, CO is available to react with radicals, ions, and larger molecules that may have been formed in the ice, and may even provide a carbon repository to react with transient species from the gas phase in subsequent stages of stellar evolution.



Figure 3. A schematic representation of the processes leading to the accumulation of CO molecules adsorbed directly at interstellar grain surfaces. (a) CO_{gg} accreted directly from the gas phase, in regions where ices are just starting to (or have not yet) formed. (b) CO_{gg} formed indirectly, in regions where CO and H₂O ice mantles have amassed and CO migrates to the grain surface. See text for a more detailed discussion.

Spectra of other molecules adsorbed on zeolites are well documented in the literature (Bordiga et al. 2000), as are the catalytic properties of such surfaces (Weitkamp 2000). In the interstellar case, H_2O is one other obvious candidate that could also chemisorb at these catalytic binding sites, besides CO. In an extensive search of the literature, it was not possible to find any measurements of H_2O binding energies at zeolite surfaces; possibly because surface analysis of such materials is conducted using CO, and requires the water of crystallization to be removed. However, to 'dry' zeolite crystals, temperatures in excess of 650 K are required, indicating that the majority of H₂O molecules must desorb at energies in excess of $\approx 20\,000$ K. In interstellar regions, the difficulty in detecting H₂O adsorbates lies in distinguishing the surface hydroxyl (Si–OH) and water–hydroxyl (H–OH) stretching vibrations from each other, and from any H₂O ice bands in the same line of sight, because all three sets of vibrations lie between 3500 and 3000 cm⁻¹. The only potential band that might be observed is a sharp, narrow, intense feature at around 3620 cm⁻¹, related to the stretching vibration of a subset

of the surface hydroxyl groups, which increases in intensity and broadens if a strong hydrogen bond forms between the surface and the adsorbate, with the adsorbate acting as the acceptor molecule (Hadjivanov & Vayssilov 2002).

5.2 Surface area of interstellar dust

Assuming that all grains are chemically equivalent in these different objects, one may at first suspect that the $\rm CO_{gg}$ column density should scale with dust column density (or H2 column density) and CO gas column density. H₂ column densities are not available for the objects discussed here, but are typically a factor of 10^4 to 10^5 greater than that of the H_2O ice column density in YSOs (e.g. Whittet et al. 2001; Pontoppidan, van Dishoeck & Dartois 2004). However, no clear scaling pattern emerges, as illustrated in Table 2. The average abundance of COgg adsorbed directly on the grain surface binding sites is $\approx 2.4 \times 10^{-6}$ per H nucleus. Assuming only one CO molecule occupies each chemisorption site and that the occupied surface area per binding site is approximately 30 Å², the total surface area occupied by CO_{gg} on interstellar grains is at least 7 (±3) × 10^{-21} cm² per H nucleus, and possibly much larger if only a fraction of the catalytically active surface sites are occupied. The uncertainty in this figure is derived from a combination of the factor of 3 uncertainty in band strength derived from the estimated occupancy of the surface binding sites in the experiment, and an additional maximum estimated error of 20 per cent in the COgg abundance per H nucleus, which arises from assuming that $n(H_2O)/n(H_2)$ is $\approx 9 \times 10^{-4}$ on all the lines of sight presented here. For comparison, the total surface area of interstellar silicates has previously been estimated as $1-2 \times$ 10^{-21} cm² per H nucleus, assuming that the grains are solid spheres (e.g. Mathis, Rumpl & Nordsieck 1977; Walmsley et al. 2004).

Many authors characterize interstellar grains using the concept of porosity, defined as the ratio between the solid volume of the grain to the total volume of the grain. Indeed, Mathis (1998) has suggested that the observed strength of the interstellar silicate band is only reproducible if interstellar silicate grains are formed from amorphous silicates, containing almost 100 per cent of the Si, Fe and Mg abundance, with >25 per cent of their total volume being 'vacuum' (Mathis & Whiffen 1989). Porous grains have both internal and external surface areas, which would be accessible if CO molecules were first adsorbed on the external grain surface, then migrated into the pores. This could account for the difference between the two estimates of total surface area (of interstellar grains) per H nucleus given above. However, porosity is only a measure of the volume-filling factor of a grain and says little of its morphology: the collision cross-section of a porous grain hardly differs in comparison to a solid grain containing the same mass of material, so the probability of CO molecules colliding with the grains would be much less than if the same mass were contained in fractal grains, whose open and filamentary structures exhibit increases in both collision cross-section and surface area when compared to solid grains of the same mass (Fogel & Leung 1998). The large interstellar grain surface area derived above from the COgg interaction, coupled with the COgg column density, seem compelling evidence that interstellar grains deviate significantly from 'solid spheres' and are most likely porous, irregularly shaped aggregates with very large surface areas.

Consequently, the CO_{gg} abundance traces the total grain surface area, i.e. the morphology of interstellar grains. Assuming interstellar grains are porous and fractal, the CO_{gg} abundance will not just depend on CO gas-interstellar grain collision rates, but also the number of available binding sites on the grains in the line of sight, which may depend on diffusion rates and geometric or chemical ac-

cessibility to the binding sites, factors related to the dust morphology rather than its abundance. As grains aggregate and grow larger, the total grain surface area will reduce, and the COgg abundance will go down. Where gas phase CO is adsorbed on bare grains, CO gas phase abundances $(2-3 \times 10^{-4}$ relative to H₂; Lacy et al. 1994) could be depleted by around 5 per cent, even in regions where no CO ice is detected. At an extinction $A_I = 1$ mag (roughly corresponding to $A_{\nu} \approx 3-4$ mag, depending on the extinction law applied), this depletion corresponds to an optical depth of the CO_{gg} band of ≈ 0.013 , assuming $N_{\rm H} = 5.6 \times 10^{21} \,{\rm cm}^{-2} \,{\rm mag}^{-1} \,A_J$ (Vuong et al. 2003). It is therefore possible that COgg bands could be detected above the noise limit (i.e. at optical depths ≈ 0.02) from lines of sight where $A_I \ge 1.5$ (roughly corresponding to $A_v \approx 4.5-6$ mag). This is above the critical extinction threshold for H₂O ice detection in Taurus (Whittet et al. 2001), but below the A_J at which CO ice could be detected. It will be an exciting prospect to find a line of sight where the weak 2175-cm⁻¹ feature is observed but CO ice is not.

5.3 Heterogeneous chemistry

The potential for heterogeneous chemistry to occur at catalytically active surfaces on interstellar grains opens up new pathways towards molecular complexity that could be as important in the solid-state chemistry of star-forming regions as the chemistry occurring at the ice surface or in the ice film itself. Put simply, heterogeneous catalysis requires the surface binding site to facilitate the surface reaction, participating in the reactions without being incorporated in the product. Effectively, the presence of a catalyst always lowers the energy barrier to a reaction, either by acting as a donor or acceptor of electrons, weakening the molecular bond and forming a transition state complex, or sterically orientating the reagents, so specific products are formed. The more effective the catalyst, the more significant the effects. For example, Woon and co-workers have used quantum chemistry to show that in the presence of H₂O ice, the barriers to H-atom addition and abstraction reactions may be lowered by between \approx 700 and 1900 K, e.g. H + H₂CO \rightarrow CH₃O (Woon 2002), and that ionization barriers are reduced by up to 50 per cent when certain species are solvated in H₂O ice, e.g. H₂, CH₃OH (Woon 2004). Silicon and metal cations are very effective catalysts because they have a range of accessible unoccupied atomic orbitals, making electron transfer and transition-state complex formation particularly simple. Consequently, silicon-based and cation-doped surfaces will make better catalysts than H2O, although in the absence of any comparative reaction studies in the literature this is difficult to prove unequivocally.

In 'non-icy' environments, it is already known that H₂O and H₂ form on bare grains from their atomic constituents (O'Neil & Williams 1999; Herbst 2001). Similar reactions, involving COgg, could account for the high abundances of CH3OH and CO2 in interstellar ices, which may be difficult to explain by ice chemistry alone (Tielens & Hagen 1982; van der Tak, van Dishoeck & Caselli 2000; Watanabe, Shiraki & Kouchi 2003). On cation-doped silica surfaces, more representative of the interstellar case, CO binding energies have been reported to range from around \approx 4930 to 8900 K (Xu & Goodman 1993), dominated by CO desorption from the most strongly bound sites. Assuming the adsorption and desorption enthalpies of these systems are similar, these figures imply that CO could become chemisorbed at bare grain surfaces prior to ice formation. In the absence of grain heating or sufficient energy to break the grain-CO bond, CO would then be likely to participate in Langmuir-Hinshelwood or Eley Rideal-type surface reactions, most likely involving an adsorbed or 'incoming' O or H atom,

leading to HCO and CO₂ formation (Fig. 3a). New reaction pathways may also be opened up, for example, if the chemisorbed CO becomes trapped under the ice layer when it forms, the COgg may react with ice constituents, e.g. free radicals or atoms, or may recombine with larger molecules forming heavier organics as the ice layer subsequently desorbs (Fig. 3b). Alternatively, as recent experimental work has shown, photochemistry of the grain-metal-cationadsorbate-ice system could generate a plethora of new molecules in the solid state (Gleeson et al. 2003; Bergeld et al. 2004). Consequently, the 'high-temperature' desorption of chemisorbed CO from bare grain surfaces will have an important impact on the organic inventory of warm protostellar regions, retaining a C source in the solid state to much higher temperatures than previously assumed, and possibly even beyond the desorption threshold of the ice layers themselves. Finally, chemisorbed CO will participate in surface reactions at higher temperatures when compared to the same reactions occurring on or in ice mantles, which may well have influenced the organic inventory of the early solar system (Pearson et al. 2002).

The adsorption of other molecular species, such as N_2 , to the same surface binding sites may lead to molecular dissociation, which activates otherwise chemically inert molecules, providing a direct route towards the formation of bio-molecular precursors, some of which have been tentatively detected in interstellar regions, e.g. glycine (Kuan et al. 2003). Furthermore, reactions involving heavy isotopes are known to be enhanced at zeolite surfaces (Lee et al. 1999), the heavier isotopes desorbing at higher temperatures, leading to isotope enhancement effects, (e.g. 13 C and D containing molecules).

6 CONCLUDING REMARKS

Prior to this work, no direct evidence existed for gas-surface dynamics in the interstellar medium. It was assumed that CO gas and interstellar dust collide and interact, but only such that CO molecules might condense and form CO ice. The experimental data presented in this paper corroborate our initial assumption that the (weak) 2175-cm⁻¹ feature is likely to be associated with CO molecules chemisorbed to bare interstellar grain surfaces. This identification suggests that gas and grains can interact without forming ice, and that CO, as well as other atomic and molecular species, may be adsorbed directly at bare interstellar grain surfaces. This observation has immediate implications: first, as to how much CO gas might be depleted on to grain surfaces, apparently invisible in gas phase or ice spectra; secondly, impacting on our understanding of the chemical and physical morphology of interstellar grain surfaces; and finally in terms of the subsequent heterogeneous chemistry that may result from CO being chemisorbed, rather than frozen out, on grain surfaces. These deductions open up a new set of discussions on the synergy between gas and dust in star-forming regions, the nature of interstellar dust, and the chemistry occurring there.

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