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Origin of the 6.85 μ m band near young stellar objects: The ammonium ion (NH⁺₄) revisited*

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Abstract. We have investigated whether the v_4 feature of NH₄⁺ is a viable candidate for the 6.85 μ m absorption band seen towards embedded young stellar objects. To produce NH₄⁺ astrophysical ice analogs consisting of H₂O, CO₂, NH₃ and O₂ were UV photolysed. The IR spectra reveal peaks that are identified with the NH₄⁺, NO₂⁻, NO₃⁻ and HCO₃⁻ ions. It is shown that the NH₄⁺ matches two absorption features that are observed towards embedded young stellar objects, i.e., the strong 6.85 μ m feature and the 3.26 μ m feature. The characteristic redshift with temperature of the interstellar 6.85 μ m feature is well reproduced. The abundance of NH₄⁺ in interstellar ices would be typically 10% relative to H₂O. The experiments show that the counterions produce little distinct spectral signature but rather a pseudo-continuum if a variety of them is present in a H₂O dominated environment. The anions could therefore go undetected in IR spectra of interstellar ice. In the ISM, where additional mechanisms such as surface chemistry and additional elements such as sulfur are available many acids and an even wider variety of anions could be produced. These components may be detectable once the ices sublime, e.g., in hot cores.

Key words. methods: laboratory – stars: individual: W33A – stars: individual: MonR2:IRS3 – ISM: abundances – ISM: molecules – infrared: ISM – ISM: lines and bands

1. Introduction

The nature of the 6.85 μ m absorption feature towards embedded Young Stellar Objects (YSO's) has remained an enigma since its discovery 25 years ago (Russell et al. 1977). While the first high resolution observations of this band were recently obtained by the Short Wavelength Spectrometer (SWS) on board the Infrared Space Observatory (ISO; Schutte et al. 1996; Dartois et al. 1999a; Keane et al. 2001), this did not yet help to clarify its origin. A number of candidates, such as carbonates, and the CH deformation modes in organic molecules like methanol, could be excluded based on the feature's spectral properties and the absence of strong additional bands. One possibility, however, the v_4 mode of the ammonium ion (NH₄⁺), first proposed by Grim et al. (1989b), could not be excluded. NH_4^+ is produced in astrophysical ice analogs by acid-base reactions. This is achieved either by deposition and warmup of NH₃ together with acids such as HNCO or HCOOH

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(Novozamsky et al. 2001; Schutte et al. 1999), or by photolysis of ice mixtures containing NH₃ (e.g., NH₃/O₂; Grim et al. 1989a) where the acids are produced by the photochemistry. Recently, the good spectral correspondence between the v_4 feature of NH₄⁺ and the interstellar 6.85 μ m absorption as seen by ISO/SWS was again demonstrated (Demyk et al. 1998). Nevertheless, this assignment faces a fundamental problem, since it requires a large abundance of NH₄⁺ of ~10% in the ices near YSO's. An equal amount of negative charge would need to be present. Although the ions OCN⁻ (also referred to as NCOin the chemistry literature) and (probably) HCOO⁻ have been identified, their abundance falls far short of what is needed for the balance (Schutte et al. 1996a; Gibb et al. 2000; Keane et al. 2001, 2002). No sign of other negative solid state species has shown up in the ISO data.

This paper revisits the possibility that NH_4^+ is responsible for the interstellar 6.85 μ m band. We investigate in the laboratory whether a *variety* of anions can supply the required countercharge, and that such a combination of species is either lacking in outstanding IR features, or fall below current detection limits. Taking the cosmic abundances of the elements into consideration, the counterions would have to consist of H, O, C, N and S in order to make a significant contribution. If we furthermore constrain the acid precursors to

^{*} Based on observations with ISO, an ESA project with instruments funded by ESA Member States (especially the PI countries: France, Germany, The Netherlands and the UK) and with the participation of ISAS and NASA.

species with at most 1 carbon atom, possibilities would still include a variety of thermodynamically stable species, i.e., OH⁻, CN⁻, OCN⁻, HCOO⁻, HCO⁻₃, CO²₃⁻, NO⁻₂, NO⁻₃, SO²₄⁻, HSO_3^- , and SO_3^{2-} . In principle an astrophysical ice analog containing NH_4^+ and a mixture of such negative ions could be prepared by depositing NH₃ together with a variety of acids. However, experimental limitations compel us to produce the ions in situ with UV photolysis. To this end, we processed mixtures of H₂O/CO₂/ NH₃/O₂ to produce acids such as H₂CO₃ (carbonic acid), HNO₃ (nitric acid) and HNO₂ (nitrous acid). These acids react with NH_3 forming NH_4^+ and a mixture of counterions, as desired. Earlier work on similar mixtures indeed showed the formation of such species (Moore & Khanna 1991; Gerakines et al. 2000; Grim et al. 1989b). Of course, this experimental method of producing the ions may be different from what happens in the ISM. For example, the high abundance of O₂ in our samples, necessary to stimulate the photochemical production of oxygen rich acids, may considerably exceed the abundance in interstellar ices (Vandenbussche et al. 1999). In space acids are possibly produced by other mechanisms such as surface chemistry (e.g., Keane & Tielens 2002, in preparation).

To verify whether NH_4^+ is a plausible candidate for the identification of the 6.85 μ m band a number of crucial issues need to be adressed. First of all, detailed spectroscopy of the v_4 mode is needed to investigate its behaviour as a function of ice composition and temperature. Next, the issue whether other infrared features of NH_4^+ could be present in interstellar ice spectra must be adressed. Subsequently, the counterions that are produced in the experiments should be identified to assess their astrophysical relevance. Finally, the spectral signature of the counterions under various conditions is studied, where the prime question is whether conditions exist at which their infrared signatures become inconspicuous.

The paper is organized as follows. In Sect. 2 we review the experimental techniques. Section 3 summarizes the results. NH₄⁺ is created by UV photolysis of astrophysical ice analogs. We describe the spectral properties of NH₄⁺ under such conditions. Furthermore we establish which counterions are formed in the experiments and study their spectroscopic properties as well. In Sect. 4 the obtained NH₄⁺ spectra are compared with observations of YSO's. Besides matching the 6.85 μ m band, it is furthermore shown that the interstellar 3.26 μ m feature closely corresponds to one of the NH⁺₄ features. Also, using the experimental results, it is investigated whether the absence of features due to counterions in the observations can be reconciled with an assignment of the 6.85 μ m band to NH₄⁺. In Sect. 5, the astrophysical implications of the NH_4^+ identification are discussed. Section 6, finally, summarizes the conclusions of this paper.

2. Experimental

Detailed descriptions of the general procedure for creation and photolysis of ice samples and the measurement of their infrared spectra have been published earlier (Gerakines et al. 1995, 1996). In summary, the set-up consists of a high vacuum chamber (10^{-7} mbar) , with an IR transparent CsI substrate mounted

on a cold finger which is cooled to ~12 K. Samples were slowly deposited ($\sim 3 \times 10^{15}$ molec. cm⁻² s⁻¹/4 μ m hr⁻¹) through a narrow tube controlled by a regulation valve. Photolysis by vacuum UV was subsequently performed by a hydrogen flow discharge lamp (~5 × 10¹⁴ photons cm⁻² s⁻¹; $E_{\text{photon}} \ge 6 \text{ eV}$). For thorough photolysis, the thickness of the sample should be ~<0.2 μ m. To overcome this limit, in one case deposition and photolysis were performed simultaneously. In this way a photolysed sample can be produced of several microns thickness, which greatly enhances the amount of photoproducts and thus the S/N ratio of the IR spectrum. Afterwards the sample was warmed in steps. The evolution of the sample throughout the photolysis and warm-up sequence was monitored by infrared transmission spectroscopy. In addition to the photochemical experiments, we did band strength measurements for NH⁺₄ by deposition and warm-up of ices containing NH₃ and HCOOH.

The reagents used in these experiments were H_2O (purified by three freeze-thaw cycles), CO_2 (Praxair, 99.996% purity), NH_3 (Praxair, 99.99% purity), O_2 (Praxair, 99.999% purity) and HCOOH (Baker, 98%). When ice samples containing both NH_3 and CO_2 or NH_3 and HCOOH were prepared, the CO_2 or HCOOH was deposited through a separate tube, to prevent reactions with NH_3 prior to deposition. This method is described in Gerakines et al. (1995).

Table 1 provides a log of our experiments. Column densities and abundances of H_2O , CO_2 and NH_3 were directly obtained from the IR spectrum using band strengths from the literature (Gerakines et al. 1995; Kerkhof et al. 1999).

3. Results

3.1. Photolysis of $H_2O/CO_2/NH_3/O_2$ mixtures

We photolysed a number of mixtures of H2O/CO2/NH3/O2 to produce NH_4^+ and a variety of negative ions (Table 1). Figure 1 shows the evolution of spectrum of photolysed $H_2O/CO_2/NH_3/O_2 = 10/2/1.1/1$ (Expt. 4; Table 1) during warm-up from 12 K to 220 K. First of all we want to verify whether NH_4^+ is formed. Therefore, Fig. 1 also shows the spectrum of the binary mixture $HNCO/NH_3 = 1/1.2$, after warm-up to 120 K (no photolysis). In this sample NH_4^+ is readily formed by proton transfer between the isocyanic acid and the ammonia (Novozamsky et al. 2001). The NH_4^+ features fall at 3000 cm^{-1} (broad; v_1), 3200 cm^{-1} ($2v_2$), 3060 cm^{-1} (v_1+v_5), near 2860 cm⁻¹ (2 v_4) and near 1450 cm⁻¹ (v_4), where v_5 is a lattice mode (assignments from Nakamoto 1972). The first four features blend together in a broad structure which extends from \sim 3500–2400 cm⁻¹. Directly after the photolysis of the $H_2O/CO_2/NH_3/O_2$ mixture, the v₄ feature is already clearly present at ~1500 cm⁻¹. The H₂O ice absorption in the 3400- 2800 cm^{-1} region obscures the other absorptions due to NH₄⁺. Upon warming to 220 K, causing the ice to sublime, the other NH_4^+ features can all be seen. Clearly, the ammonium ion is produced by the photolysis.

As discussed in Sect. 1, we want to investigate the spectral properties of the v_4 NH₄⁺ feature. To study the temperature dependence, Fig. 2 shows the 1900–1000 cm⁻¹ spectra of photolysed H₂O/CO₂/NH₃/O₂ = 10/2/1.1/1 (expt. 4). Spectra

Table 1. Log of experiments.

	Mixture				N	d^a	dose	dose
	H_2O	CO_2	NH ₃	O_2	mol. cm^{-2}	μ m	photons cm ⁻²	eV mol. ^{-1d}
1	10	10	0	10	7.2(17)	0.37	9(17)	<11 ^c
2	10	5	7	10	4.0(17)	0.17	9(17)	$<20^{c}$
3^b	10	2.2	1.2	1	2.5(19)	9.5	2.4(19)	8.6
4	10	2	1.1	1	8.5(17)	0.32	6(17)	<6.4 ^c
5	10	1.2	0.9	0.9	4.8(17)	0.17	9(16)	<1.7 ^c
6	0	1.2	1	1	4.4(17)	0.23	9(16)	<1.8 ^c
7	0	0	1	1	4.5(17)	0.18	9(17)	<18 ^c

^{*a*} Thickness obtained from the column density using a nominal density of 1 g cm^{-3} .

^b Obtained by simultaneous deposition and photolysis.

^{*c*} Sample not optically thick, therefore dose is an upper limit.

^d Using an average photon energy of 9 eV (Jenniskens et al. 1993).



Fig. 1. Spectra of the photolysed sample $H_2O/CO_2/NH_3/O_2 = 10/2/1.1/1$ (expt. 4): **a**) before warm-up (12 K); **b**) at 120 K; **c**) at 220 K. Curve **c**) was scaled up by a factor 3. Curve **d**) shows the spectrum of HNCO/NH₃ = 1/1.2 at 120 K (after the transformation of most of the original species into OCN⁻ and NH₄⁺; arrows identify features of NH₄⁺). The spectra have been offset for clarity.

are shown directly after photolysis at 12 K, and at a variety of temperatures up to 240 K. Some important spectral properties can be gleaned from the figure. Upon warm-up to 120 K the depth of the v_4 feature near 1460 cm⁻¹ increases by a factor \sim 2. While this increase could result from further acid-base reactions during the warm-up, the absence of any feature attributable to acids such as H₂CO₃ or HNO₃ after the photolysis (Fig. 2, H_2CO_3 has bands at 1727, 1480 and 1275 cm⁻¹; Gerakines et al. 2000; HNO₃ has a strong feature at 1300 cm^{-1} ; McGraw et al. 1965) argues against this possibility. Thus, the growth of the NH⁺₄ feature during warm-up is likely caused by an increase of its intrinsic strength. Furthermore, the v_4 feature shifts strongly redward with temperature. The enhancement and the shift reflect the strong interaction of the ion with its environment. Figure 3 plots the position of the v_4 band vs. temperature for a number of samples (expts. 2, 4 and 6 of Table 1).



Fig. 2. Thermal evolution of the photolysed ice mixture $H_2O/CO_2/NH_3/O_2 = 10/2/1.1/1$ (expt. 4), blow-up of the 1900–1000 cm⁻¹ region: **a**) before warm-up (12 K); **b**) after warm-up to 120 K; **c**) to 150 K; **d**) to 180 K, **e**) to 220 K; **f**) to 240 K. Apart from the multiplication factors indicated in the figure, the vertical scale is identical for all curves. However, the spectra have been offset for clarity.

It can be seen that, while there is a steady redshift throughout the warm-up, the shift is particularly pronounced during H_2O sublimation between 160 and 180 K.

As discussed in Sect. 1, it is essential to identify the negative ions produced in the experiments to assess their astrophysical significance. To this end, we study the IR spectrum of the residue which remains after the ices have fully evaporated. As a basis for the interpretation of more complex samples, Fig. 4 shows the warm-up behaviour of the residue of photolysed $NH_3/O_2 = 1/1$ (expt. 7; Table 1). The evolution is best understood by a "hot to cold" interpretation, i.e., starting with the most refractory component at the final stages of the warm-up, and then working backwards in the sublimation sequence to see which spectral elements are sequentially lost. After warm-up to 220 K, a component remains which is stable up to ~280 K. Its spectrum resembles that of ammonium nitrate (Fig. 4; NH_4NO_3 ; throughout the paper we will use the



Fig. 3. Redshift of the v_4 NH⁴₄ feature as a function of temperature: H₂O/CO₂/NH₃/O₂ = 10/2/1.1/1 (expt. 4; open circles); 1/0.5/0.7/1 (expt. 2; open squares); 0/1.2/1/1 (expt. 6; filled circles).



Fig. 4. The thermal evolution of the residue of photolysed $NH_3/O_2 = 1/1$ (Expt. 7). From central to top: the residue at 180 K; at 220 K; subtraction of the 220 K from the 180 K spectrum. The vertical scale is identical for all curves, however, the spectra have been offset for clarity. For comparison, ammonium nitrate (NH_4NO_3) and sodium nitrite ($NaNO_2$) are shown at the bottom (Miller & Wilkins 1952).

standard chemical notation for salts. It must be noted however that the constituents of these compounds are ions, in this case, NH_4^+ and NO_3^-). The NO_3^- ion was previously identified in photolysed NH₃/O₂ ices (Grim et al. 1989b). However, while there is a general correspondence with the literature spectrum of ammonium nitrate there are some differences in the relative intensities and positions of the bands. The literature spectrum was produced from fine crystalline powder in nujol mull, while our samples consist of a mixture of salts at low temperature. Spectra of ions are generally quite sensitive to factors like temperature, matrix and degree of annealing, due to the strong interaction of the ions with the matrix (cf. OCN⁻ in various salt matrices; Maki & Decius 1959). This effect possibly causes the differences. During warm-up from 180 to 220 K two strong features disappear at 1270 and 1220 cm^{-1} , together with a fraction of the 1450 cm⁻¹, and 2400–3500 cm⁻¹ complex due to NH₄⁺.



Fig. 5. The thermal evolution of the residue of photolysed $H_2O/CO_2/NH_3/O_2 = 10/2.2/1.2/1$ (Expt. 3). From bottom to top: The spectrum of the sample at 200 K; at 240 K; at 280 K; subtraction of the 240 K from the 200 K spectrum; subtraction of the 280 K from the 240 K spectrum. The vertical scale is identical for all curves, however, the spectra have been offset for clarity. For comparison, ammonium nitrate (NH₄NO₃; Miller & Wilkins 1952) and ammonium bicarbonate (NH₄HCO₃; Khanna & Moore 1999) are shown.

The band at 1220 cm⁻¹ may correspond to the main feature of NO_2^- . As compared with the literature spectrum of $NaNO_2$ (in nujol mull; Fig. 4), the band is shifted by 35 cm⁻¹, possibly due to matrix interactions. The assignment of this component with ammonium nitrite (NH₄NO₂) is supported by the identification of the NO_2^- ion in the photolysed ice (Grim et al. 1989b). The 1270 cm⁻¹ feature corresponds to an unidentified photoproduct.

Figure 5 shows the spectral evolution of the residue of photolysed $H_2O/CO_2/NH_3/O_2 = 10/2.2/1.2/1$ (expt. 3). To probe the nature of the anions, we will again analyse this diagram "hot to cold". As with the NH₃/O₂ sample, the most refractory component, remaining at 280 K after all other material has sublimed, resembles ammonium nitrate (Fig. 5). During warmup from 240 to 280 K a component sublimes which is characterized by two strong bands at 1600 and 1300 cm⁻¹, together with four weaker features at 1375, 1010, 830, and 690 cm^{-1} . The disappearance of these bands is accompanied by a decrease of the NH_4^+ features. All these features have counterparts in the spectrum of ammonium bicarbonate in a KBr pellet (NH₄HCO₃; Fig. 5). Still, some differences are apparent in width and relative intensity, while furthermore some extra bands are present in the literature spectrum of which only minor indications are seen in the 240 to 280 K component. Again, we note that the spectra of ions are generally quite sensitive to such factors as temperature, matrix and degree of annealing, and it seems likely that the differences could derive from such factors.

To verify the presence of HCO_3^- in the 240–280 K fraction, we photolysed the ice mixture $H_2O/CO_2/O_2 = 1/1/1$ (Expt. 1, Table 1) to see whether its precursor, carbonic acid (H_2CO_3) is formed when NH₃ is omitted from the mixture. The photolysis



Fig. 6. Spectra of photolysed ice samples, after warm-up to 160 K, enlargement of the 1900–1000 cm⁻¹ region: **a**) NH₃/O₂ = 1/1 (expt. 7); **b**) H₂O/CO₂/NH₃/O₂ = 1/0.5/0.7/1 (expt. 2); **c**) 10/2/1/1.1 (expt. 4); **d**) and 10/1.2/0.9/0.9 (expt. 5). The spectra have been offset for clarity.

gave rise to features at 3040, 2840, 2627, 1727, 1480, and 1275. These bands can all be ascribed to carbonic acid (Moore & Khanna 1991; Gerakines et al. 2000).

The residue fraction subliming between 200-240 K is characterized by a rather complex spectrum. Features at 1605, 1375, 1300, and 1010 cm⁻¹ show that some HCO₃⁻ sublimes in this temperature interval. However, the weakness of these bands shows that the contribution of bicarbonate to the total negative charge in the 200-240 K fraction is minor. The band at 1670 cm⁻¹ and the broad structure between 1000 and 600 cm⁻¹ likely correspond to a small quantity of H₂O which is embedded in the more refractory material (Schutte & Buys 1961; Ryskin 1974). Likewise, the 2340 cm⁻¹ feature can be ascribed to a small amount of embedded CO₂. Like for the photolysed NH_3/O_2 , the 1220 cm⁻¹ may be due to the NO_2^- ion (Fig. 4). However, the intensity of this band shows that $NO_2^$ accounts for only ~half of the countercharge of the NH_4^+ which sublimes in the 200-240 K interval. The 2170 cm⁻¹ feature indicates the production of some OCN⁻ (Hudson et al. 2001), however, it is too weak to make a substantial contribution to the countercharge. We conclude that an important fraction of this relatively volatile residue component must consist of unknown anions. No sign of such species, or of the acids from which they originate, was found in experiments with less than four initial ice components (i.e., H₂O/CO₂/O₂, NH₃/O₂; Sect. 3.1). It therefore seems probable that they are relatively complex ions whose formation pathway involves all four original components.

As outlined in Sect. 1, a prime issue for the viability of the NH₄⁺ assignment of the 6.85 μ m band concerns whether it is possible to produce a clear v_4 spectral signature while features of counterions are weak or absent as for the interstellar spectra. From Fig. 2 it can be seen that intense broad bands near 1320 and 1220 cm⁻¹ are present once the ices have sublimed at ~180 K. As discussed above, these bands are caused by negative ions, probably HCO₃⁻, NO₃⁻, and NO₂⁻. Before ice sublimation, these bands are however quite inconspicuous, resulting in a smooth sloping spectrum in the 1400–1200 cm⁻¹ range with only minor substructure due to the 1300 and 1270 cm⁻¹ features of unidentified photoproducts.

To further investigate the influence of the H₂O matrix on the appearance of the anion bands, Fig. 6 compares the 1900– 1000 cm⁻¹ spectra of the ice samples H₂O/CO₂/NH₃/O₂ = 0/0/1/1, 1/0.5/0.7/1, 10/2/1/1.1 and 10/1.2/0.9/0.9 (Expts. 2, 7, 4 and 5) All spectra were taken after warm-up to 160 K and sublimation of NH₃, which enables a better view of the features of the photoproducts. Samples 2 and 7 show strong bands due to negative ions at 1385 and 1335 cm⁻¹ (NO₃) and at 1230 cm⁻¹ (NO₂). There is hardly a sign of these or other features of anions in the H₂O dominated samples (expts. 4 and 5), even though the intensity of the v_4 NH₄⁺ band at ~1475 cm⁻¹ is similar. This again shows that in the H₂O dominated samples features due to anions are inconspicuous.

3.2. Summary

The photolysis of $H_2O/CO_2/NH_3/O_2$ ice mixtures efficiently produces NH_4^+ . These experiments revealed the following four important spectral properties of NH_4^+ and the negative counterions which pose important constraints for an identification of NH_4^+ in interstellar ices:

1. During warm-up from 12 to 280 K, the NH₄⁺ v_4 feature shows a pronounced shift from ~1480 cm⁻¹ to ~1420 cm⁻¹.

2. Between 12 to 120 K the $NH_4^+ \nu_4$ feature grows smoothly by a factor 2.

3. In H₂O ice three relatively weak features of NH₄⁺ are only evident after the ice sublimation at ~180 K. These are the $2v_2$ band near 3200 cm⁻¹, the v_1+v_5 band near 3060 cm⁻¹, and the $2v_4$ band near 2860 cm⁻¹.

4. In H_2O dominated ice, the spectral features of the *counterions* are only apparent after the sublimation of the ice, but are unconspicuous in the H_2O dominated ice matrix.

Concerning this last point, analysis of the spectrum of the residue shows that HCO_3^- , NO_3^- , NO_2^- are probably present. The total abundance of these species is somewhat unsufficient for balancing the NH_4^+ , indicating the production of additional, yet unidentified, anions.

The identifications of anions from the IR spectra are in some cases tentative, while in other cases no identification was possible at all. A positive identification of the anions and corresponding acid neutrals may help to constrain the composition of interstellar ices (see Sect. 5 below). In future, the nature of the negative species could be studied by analysis of the molecules that evaporate during the warm-up, for example, by mass spectroscopy.

3.3. Strength of the v_4 NH⁺₄ band

To obtain column densities, it is necessary to measure the intrinsic strength of the $NH_4^+ v_4$ feature. This can be derived from experiments in which NH_4^+ is formed by simple warmup (no photolysis) of an ice containing NH_3 and an acid. In this case the quantity of NH_4^+ formed during warm-up is

Table 2. Band strength of the v_4 NH⁺₄ band.

mixture	<i>T</i> (K)	$A (\text{cm ion}^{-1})$
H ₂ O/NH ₃ /HCOOH		
100/3.6/3.6	120 K	4.4 (-17)
$NH_3/HCOOH = 4/10$	80 K	4.0 (-17)
H ₂ O/CO ₂ /NH ₃ /O ₂		
10/1.2/0.9/0.9, + hv	150 K	≥2.7 (−17)
10/1.2/0.9/0.9, + hv	150 K	≥2.7 (−17)

equal to the amount of NH₃ which is lost. Since the band strength of the NH₃ umbrella mode in various matrices is known (Kerkhof et al. 1999), this allows a precise determination of the NH_4^+ band strength. We performed two experiments involving NH3 with formic acid (HCOOH; see Schutte et al. 1999 for experimental details). The results are listed in Table 2. Since the strength may depend on the composition of the matrix and nature of the counterion, the band strength of NH_4^+ in the photolysis experiments may differ from the HCOOH/NH₃ mixtures. No direct measurement in the photolysis experiments is possible, because, besides acid-base reactions, other processes may contribute to the NH₃ destruction during the photolysis. Therefore the amount of NH₃ destroyed exceeds the formation of NH₄⁺ by an unknown factor and only a lower limit to the NH_4^+ band strength can be obtained. The most constraining lower limits are derived for the experiments with H₂O-dominated ices (see Table 2). This is not surprising, since in these experiments features of N-containing photoproducts other than NH_4^+ are small (Sect. 3.1).

We will adopt a standard band strength $A(v_4, \text{NH}_4^+) = 4.4 \times 10^{-17}$ cm mol.⁻¹. This is close to the value of $(2.5-3.5) \times 10^{-17}$ cm mol.⁻¹ found in aqueous solution (Lowenthal & Khanna, in preparation). As discussed in Sect. 3.1, the band strength varies by a factor 2 with temperature in our photolysis experiments. It is unclear whether the standard band strength corresponds to the highest or lowest bandstrength in the photolysis experiments. We will therefore adopt a standard uncertainty of a factor 2 in this value.

4. Comparison to observations of Young Stellar Objects

In this section we investigate a number of criteria for the identification of the interstellar 6.85 μ m absorption with the v_4 feature of the ammonium (NH₄⁺) ion. First of all, the feature should provide a good match to the observed band. Second, the characteristic redshift of the feature with temperature (Sect. 3.1; Fig. 3) should show up in the observational data. Third, additional bands of NH₄⁺ are sought, specifically the $2v_2$, $v_1 + v_5$, and $2v_4$ features near 3200, 3060, and 2860 cm⁻¹ (3.12, 3.27, and 3.50 μ m). Fourth, there should be no spectral structure due to counter-ions that is inconsistent with the interstellar data. It will be shown that the available observational information indicates that all these criteria are satisfied. For further reference, Table 3 lists the relevant observational data for all lines of sight where high quality spectra of the 6.85 μ m feature (i.e., by ISO-SWS) are available. Data on the interstellar 3.26



Fig. 7. ISO/SWS 1 (thin line) and SWS 6 spectra (1900–1200 cm⁻¹ thick line) of W33A (from Gibb et al. 2000 and Keane et al. 2001, respectively). The smooth dashed curve is a third-order polynomial continuum fit.

and 3.48 μ m features are included because they are possibly associated with the NH⁺₄ $\nu_1 + \nu_5$ and $2\nu_4$ bands (see below). The depth of the 6.85 μ m band was measured from the original data. These were obtained from the ISO-SWS database (for NGC 7538:IRS1) and from Keane et al. 2001 (all other objects). The depth was obtained by subtracting a linear baseline in the log(*F*) vs. μ m plane. The baseline was drawn through the 5.5 and 7.5 μ m points.

4.1. Comparison with the interstellar 6.85 μ m feature

4.1.1. W33A

Figure 7 shows the 4000–1200 cm⁻¹ (2.5–8.3 μ m) spectra of the high mass YSO W33A obtained by ISO/SWS in observation modes 1 and 6 (de Graauw et al. 1996). Spectra were adopted from Gibb et al. (2000) and Keane et al. (2001), respectively. While the SWS6 spectrum gives the highest S/N, the wide range of the SWS1 spectrum allows an accurate baseline determination. We defined the baseline over the entire 4000-1200 cm⁻¹ range by drawing a smooth third order polynomial through the continuum regions $4100-3700 \text{ cm}^{-1}$, 2010-1980 cm⁻¹, and 1840–1810 cm⁻¹ (Fig. 7). This procedure differs in some respects from the baseline definition in Keane et al. and Gibb et al. Keane et al. defined the baseline only over a limited data range (1900-1300 cm⁻¹), and included the region around 1300 cm⁻¹, just before the onset of the silicate feature, in the baseline fit. It is however clear from Fig. 7 that, at least for W33A, there is considerable absorption in this region. Gibb et al., while also fitting the baseline over a similar broad range, included the 2500-2440 cm⁻¹ region in the fitting procedure. However, it is probable that both the long-wavelength shoulder of the H₂O band (Willner et al. 1982; Smith et al. 1989, see also Fig. 7) as well as the broad H_2O combination feature near 2220 cm⁻¹ (Gibb et al. 2000) extend into this

Table 3. Properties of the 3.26, 3.48, and 6.85 μ m features towards YSO's.

Object	$N(H_2O)^a$	$\frac{F(45)}{F(100)}b$	$3.26 \mu\mathrm{m}^c$		$3.48 \mu \mathrm{m}^c$	6.85 μm		$\frac{\tau(3.26)}{\tau(6.85)}^h$	$\frac{N(\mathrm{NH}_{4}^{+})}{N(\mathrm{H}_{2}\mathrm{O})^{\mathrm{i}}}$
	10^{18} cm^{-2}	. ,	$v (cm^{-1})$	τ	au	ν (cm ⁻¹)	$ au^d$		%
W33A	12	1.27		< 0.4	>0.29	1471	1.1	< 0.33	11
NGC 7538:IRS9	7	2.07			0.13 (0.02)	1473	0.29		5
GL2136	5.0	2.75		< 0.14	0.14 (0.02)	1459	0.23	< 0.6	6
S140:IRS1	2.8	3.24	3072	0.036 (0.007)	0.027 (0.007)	1451	0.16	0.23 (0.06)	7
			3085^{g}	0.050^{g}	0.050^{g}			0.31	
W3:IRS5	5.8	4.12			0.13	1459	0.25		5
MonR2:IRS3	1.6^{f}	4.32	3071	0.049 (0.007)	0.036	1437	0.23	0.21 (0.03)	17
AFGL 7009S	12					1462	1.1		10
GL 989	3		3098 ^g	0.069^{g}	0.069^{g}	1471	0.11	0.63	4
ho Oph Elias 29	3			< 0.03	0.089 (0.007)	1470	0.07	< 0.4	3
			3080 ^g	0.085^{g}	0.065^{g}			1.2	
HH100	2.4		3087	0.032 (0.010)	0.041 (0.005)	1470	0.15 (0.06)	0.21 (0.09)	8 (3)
NGC 7538:IRS1	3		3065	0.078 (0.013)	0.052 (0.014)	1455	0.32^{e}	0.24 (0.04)	13

^{*a*} Water column density obtained from the integrated depth of the 3 μ m band: NGC 7538:IRS9/Allamandola et al. (1992); AFGL 7009S/Dartois et al. (1999a) (from 6 μ m feature); W33A/Gibb et al. (2000) (obtained by fitting the 3 μ m feature with laboratory analog spectra); AFGL 989/Smith et al. (1989); AFGL 2136/Schutte et al. (1996a); Elias 29/Boogert et al. (2000); S140:IRS1 (or GL 2884)/Willner et al. (1982); W3:IRS5/Allamandola et al. (1992); MonR2:IRS3/Smith et al. (1989); HH100 (or RCra:IRS1)/Whittet et al. (1996); NGC 7538:IRS1/Willner et al. (1982).

^b Ratio of flux at 45 and 100 μ m; from Keane et al. (2001).

^c Ground-based observations (Sellgren et al. 1995; Brooke et al. 1996; Brooke et al. 1999), unless otherwise noted. Error noted in parentheses.

^d Keane et al. (2001); error typically 20%, unless otherwise noted.

^e From the ISO-SWS database.

f It is unclear whether the 3 μ m feature of MonR2:IRS3 is fully caused by H₂O ice, due to the large width of this band (Smith et al. 1989). Therefore this number should be considered an upper limit.

^{*g*} Observed by ISO-SWS (Bregman et al. 2001).

^h Boldface entries give the 3.26 μm depth derived from ISO-SWS data (Bregman et al. 2001). Errors are given in parentheses.

^{*i*} The uncertainty, due to the error in the bandstrength, is a factor ~ 2 .

region. Indeed, by excluding it we obtain a smoother baseline than Gibb et al. (i.e., third, rather than fourth order polynomial).

Although there is a slight offset between the SWS1 and SWS6 data, their overall spectral shape is quite consistent. Therefore the SWS6 data were converted to an optical depth scale by subtraction of the baseline of Fig. 7. The result is shown in Fig. 8. The small offset between the SWS1 and SWS6 data was compensated by normalizing to 0 in the 1900–1800 cm⁻¹ region.

The CH₃ deformation feature of methanol falls near 6.85 μ m and will contribute a fraction of the interstellar band. The methanol column density towards W33A, as determined from the v_3 and combination bands near 3.54 and 3.9 μ m, respectively, equals 1.85×10^{18} cm⁻² (Dartois et al. 1999b). Figure 8 compares the optical depth spectrum of W33A with the v_3 band of methanol scaled to this column density. This feature was measured in a mixture H₂O/CH₃OH/CO₂ = 1/1/1 (Ehrenfreund et al. 1999). This matrix should be representative of the ices in which methanol in circumstellar objects is embedded (Gerakines et al. 1999; Dartois et al. 1999a). It is clear from Fig. 8 that methanol only gives a small contribution to the observed 6.85 μ m band of ~20% (see also Grim et al. 1991; Schutte et al. 1996). To enable a more suitable comparison with the laboratory NH₄⁺ spectra, we have subtracted this small

contribution from the W33A spectrum. The resulting curve is shown in Fig. 8. Since the correction is small, using spectra of methanol in other matrices, such as pure methanol, or a waterdominated ice, gives a very similar result.

Figure 9 compares the 6.85 μ m band of W33A (after subtraction of the methanol feature) with the v_4 band of NH⁺₄ as produced in one of our water-dominated samples after photolysis and warm-up (expt. 4). To take into account the variety of dust temperatures probed by the line of sight, the laboratory spectrum is an average of the 12, 120, and 180 K (×2) spectra. We note that good overall matches can be obtained with all photolysed water-dominated ices (expts. 3, 4 and 5), since they produce little substructure in the 1400–1250 cm⁻¹ (7.1–8.0 μ m) range. On the other hand, the photolysed ice mixtures with less water show strong peaks in this region due to NO⁻₃ and NO⁻₂ (Sect. 3.1; Fig. 6), which have no interstellar counterparts.

Due to the superposition of ices at different temperatures along the line of sight, components of different volatility will have different average temperature. This is illustrated by the temperature averaged laboratory match of the W33A spectrum (Fig. 9). The NH₄⁺ band arises primarily in the 120 and 180 K components, since at 12 K the intensity of this feature is small (Fig. 2). On the other hand, the H₂O ice features are produced in the 12 and 120 K spectra, since at $T \ge 170$ K H₂O sublimes. 1056



Fig. 8. Correction of the 6.85 μ m band of W33A for the contribution of CH₃OH. **a**) H₂O/CH₃OH/CO₂ = 1/1/1, at 112 K (from Ehrenfreund et al. 1999; band strength has been scaled to correspond to the CH₃OH column density, see text); **b**) optical depth spectrum of W33A obtained by subtraction of the continuum from the SWS 6 data (See Fig. 7); **c**) corrected optical depth spectrum of W33A obtained by subtraction of curve b. The vertical scale is identical for all curves, however, curve c has been offset for clarity.



Fig. 9. The optical depth spectrum of W33A after correction for CH₃OH (Fig. 8), compared to: **a**) photolysed $H_2O/CO_2/NH_3/O_2 = 10/2/1.1/1$ (expt. 4), average of 12 K, 120 K, 180 K (×2) spectra. Curve a has been offset for clarity.

Thus, the average temperature of the solid H₂O in the laboratory spectrum is 66 K, while the average temperature of NH₄⁺ is ~150 K. This clearly shows that temperatures derived by fitting the 6.85 μ m band should not be taken as representative of the temperature of more volatile components located in cooler regions along the line of sight.

4.1.2. MonR2:IRS3

MonR2:IRS3 is an extreme object in a number of ways. First, the solid CO₂ abundance is exceptionally low (\sim 1%;



Fig. 10. The optical depth spectrum of MonR2:IRS3 (From ISO/SWS; Keane et al. 2001) compared with some photolysed laboratory samples: **a**) $H_2O/CO_2/NH_3/O_2 = 1/0.5/0.7/1$ at 160 K (expt. 2); **b**) same, at 170 K; **c**) 10/2/1.1/1 at 180 K (expt. 4); **d**) 10/1.2/0.9/0.9 at 180 K (expt. 5). The laboratory spectra are offset for clarity.

Keane et al. 2001). Second, its 6.85 μ m feature is the reddest of all YSO's (Table 3). Third, the strength of the 6.85 μ m feature relative to the column density of H₂O is very high (Table 3). This all may indicate that the ice along the line of sight is exceptionally warm, as is also indicated by the high flux ratio $F(45 \ \mu\text{m})/F(100 \ \mu\text{m})$ (Keane et al. 2001; Table 3). Indeed, the large relative intensity of the 6.85 μ m band, assuming that the carrier of this band is a relatively refractory species, suggests that H₂O ice has begun to evaporate along most of the line-of-sight. This is confirmed by the large column density of water vapor (Boonman et al. 2000). Ice sublimation should greatly enhance the visibility of features of more refractory components, making MonR2:IRS3 an excellent testing ground for the presence of NH⁴₄ and associated counterions.

Figure 10 compares the 6.85 μ m band of MonR2:IRS3 with the v_4 NH⁺₄ features obtained in several of our experiments. This set of experimental data will further down be used to evaluate the presence of other interstellar NH_4^+ bands. No correction was made for the contribution of methanol to the 6.85 μ m band, since no CH₃OH has been observed towards this source (abundance $\leq 4.5\%$; Dartois et al. 1999b). The best matches are obtained for the photolysed water-rich ices (expts. 4 and 5), since the water-poor ice (expt. 2) gives strong features at 1385 and 1335 cm⁻¹ due to NO₃⁻¹ which are not observed. In all cases, the position of the laboratory band is somewhat blue-shifted from the MonR2:IRS3 feature. This situation could be mended by selecting laboratory spectra at higher temperatures. However, for such high temperatures a broad band near 1320 cm⁻¹, which is caused by a superposition of features of various negative ions (Fig. 2, Sect. 3.1) shows up strongly, resulting in a decrease of the overall quality of the match. Even at the temperatures selected for Fig. 10 too much absorption appears to be present in the 1350 cm⁻¹ region. However, this difference could be caused by the original choice of baseline,



Fig. 11. Position of the 6.85 μ m feature vs. *F*(45 μ m)/*F*(100 μ m) for the YSO's listed in Table 3.

which made the a priori assumption of little absorption in the 1350 cm⁻¹ region (Keane et al. 2001). As for W33A, considerable absorption may be present in this region. Thus, the position of the 6.85 μ m feature of MonR2:IRS3 is not fully reproduced by the experimental spectra, although the difference is small (~3–10 cm⁻¹) for the three H₂O-dominated samples. In view of the strong matrix dependence of the position of this band (Fig. 3), it seems plausible that the offset may be caused by a shift associated with the composition of the interstellar matrix.

Besides the 6.85 μ m band, the 6 μ m feature of MonR2:IRS3 also has a counterpart in the laboratory spectra. Especially expt. 4 at 180 K gives a band which is quite similar (Fig. 10). At this temperature most H₂O has sublimed, and the laboratory features near 6 μ m are dominated by a superposition of bands from various negative ions, i.e., HCO₃⁻ and one or more unidentified species (Fig. 4; Sect. 3.1). This suggests that most of the 6 μ m feature of MonR2:IRS3 is caused by non-volatile components rather than H₂O ice. Such a conclusion is consistent with the observations, which show that the 6 μ m feature of MonR2:IRS3 is 3.5 times stronger than expected at the H₂O column density derived from the 3 μ m band (Keane et al. 2001).

4.2. The redshift with temperature

It was shown in Sect. 3.1 that the ammonium band shifts strongly towards the red with increasing temperature. Even though the position of the NH_4^+ band depends considerably on the starting matrix, this general trend held for all our experiments. Thus, if the ammonium ion is the carrier of the interstellar 6.85 μ m feature, a correlation should be found between the position of the 6.85 μ m feature and indicators of the dust temperature along the line of sight.

Figure 11 plots the position of the 6.85 μ m band in the data set of Keane et al. (2001) against the ratio of the fluxes at 45 and 100 μ m as measured by ISO/LWS. This ratio is a general indicator of the dust temperature along the line of sight. It can be seen that there is a clear correlation between the position of the band and the flux ratio, in the sense that the interstellar band shows a strong redshift with increasing average dust temperature. A similar conclusion was already formulated by Keane et al., with the difference that they interpreted the change as due to a systematic variation with line of sight temperature in the contribution of two independent components of the $6.85 \,\mu$ m feature. It must be stressed that interpretation of the temperature dependence of the band as either a variation of independent components, or a shift of a single component, is solely a matter of interpretation and cannot be decided on the basis of the interstellar band profile, since the feature does not show substructure.

The systematic redshift of the 6.85 μ m band with line-ofsight dust temperature is in good agreement with the trend observed for the v_4 feature of the ammonium ion. Comparing Figs. 11 and 3 of Sect. 3.2.1, it can be seen that the spectral region in which the interstellar feature is found falls well within the range spanned by the position of the v_4 band. Thus, the systematic redshift of the interstellar 6.85 μ m band gives strong support to its identification with the v_4 mode of NH⁴₄.

The range of positions of the interstellar band corresponds to laboratory temperatures of ~120-240 K. It must be noted though that the position of the feature not only depends on temperature, but also on the ice matrix (Fig. 3). Therefore, this temperature range should be taken as a rough indication. Nevertheless, it seems clear that solid NH_4^+ is found at higher temperatures than more volatile ice components. Matching the profile of the $3 \mu m H_2 O$ feature typically yields 20–80 K (Smith et al. 1989), while matching the profile of the $15.2 \,\mu\text{m}$ bending mode of CO₂ gives 110-140 K (Gerakines et al. 1999). This difference in temperature is consistent with the refractory nature of NH_4^+ , which will survive in the warmer regions close to the embedded source where the more volatile ice components have sublimed. In addition, the band strength of NH⁺₄ increases considerably above ~100 K (Sect. 3.1; Fig. 2). This effect will minimize the contribution of the coldest regions to the 6.85 μ m band.

4.3. Additional NH₄⁺ features

Besides the v_4 feature near 6.85 μ m, the ammonium ion shows a number of additional features, i.e., $2v_2$, $v_1 + v_5$, and $2v_4$. These are centered near ~3200 cm⁻¹ (3.12 μ m), ~3060 cm⁻¹ (3.27 μ m), and ~2860 cm⁻¹ (3.50 μ m), respectively (Figs. 1 and 5, Sect. 3.2.1). The optical depth of each of these bands is typically 30% of that of the v_4 feature.

The $2v_2$ feature would be very difficult to distinguish, since it nearly coincides with the peak of the 3 μ m water band (Smith et al. 1989). The two other bands, while relatively weak, may be observable as substructure on the red wing of the water band. Indeed, features have been observed towards many protostellar sources near these positions (e.g., Sellgren et al. 1995; Brooke et al. 1996; Brooke et al. 1999; summary in Table 3). For example, Fig. 12 shows the mid-IR spectrum of MonR2:IRS3 obtained by ISO-SWS (Gibb et al. 2001) and by ground-based spectroscopy (between 3.16–3.65 μ m; Sellgren et al. 1995). As already mentioned above, this source appears



Fig. 12. Comparison between the 3.26, 3.48 and 6.85 μ m features of MonR2:IRS3 and the $v_2 + v_4$, $2v_4$ and v_4 features of NH₄⁺ in a number of laboratory samples. The various NH₄⁺ features are indicated by vertical dashed lines. The labelling of the laboratory spectra corresponds to the description given with Fig. 10. Dotted lines indicate the adopted baselines for the two features. The observational data comprise ground-based observations between $3.16-3.65 \mu$ m (Sellgren et al. 1995; bold solid line) and ISO/SWS data (2.6–8 μ m; Gibb et al. 2001; solid line). The MonR2:IRS3 data were converted to optical depth by subtraction of a 2nd order polynomial baseline defined by fitting the continuum regions around 3800 and 2000 cm⁻¹ in the original flux data. All spectra were smoothed to resolution 400. The spectra have been offset for clarity.

to be exceptionally warm and therefore should have enhanced abundances of refractory components like NH_4^+ .

To facilitate comparison with the laboratory data, a continuum has been subtracted from the MonR2:IRS3 original flux data (Ground-based as well as ISO-SWS). This continuum was obtained using the ISO-SWS data by drawing a second order polynomial through the zero absorption regions around 3800 and 2000 cm⁻¹. Before continuum subtraction, the groundbased data were multiplied by 0.88 to match them to the ISO-SWS spectrum.

A small excess is apparent in the ISO-SWS data relative to the ground based spectrum around 3.29 μ m. This excess is obviously caused by the presence of the well-known emission feature in the wide aperture ISO data (This is a common phenomenon for ISO observations towards YSO's; Gibb et al. 2001).

As already discussed by Sellgren et al., MonR2:IRS3 shows two features at 3.26 and 3.48 μ m. Figure 12 also shows the $v_2 + v_4$ and $2v_4$ NH⁺₄ bands as obtained in our experiments. To allow a detailed comparison, we subtracted baselines from both the laboratory and interstellar spectra, as indicated in Fig. 12. The resulting optical depth curves are shown in Fig. 13. Only the ground-based observations were used, due to the "contamination" of the ISO spectra with the 3.3 μ m emission feature.



Fig. 13. Comparison between the 3.26 and 3.48 μ m (3075 and 2875 cm⁻¹) features of MonR2:IRS3 and the $v_2 + v_4$ and $2v_4$ bands of NH₄⁺ in the laboratory samples of Fig. 12, after subtraction of the baselines indicated in this figure. The laboratory spectra have been offset for clarity.

The comparison in Figs. 12 and 13 indicates a good general agreement between the 3.26 and 3.48 μ m features of MonR2:IRS3 and the $v_2 + v_4$ and $2v_4$ NH⁺₄ bands. Nevertheless, no single spectrum is able to fit in detail both the position and relative intensity of the observed features. However, the variation among the laboratory spectra shows that the features are quite matrix and temperature dependent, and it therefore seems plausible that this moderate difference could derive from matrix effects. Besides the agreement in profile, the agreement between the intensity of the laboratory and the MonR2:IRS3 features lends support to an identification with NH⁺₄.

It is clear from Fig. 12 that if the 3.26 and 3.48 μ m features of MonR2:IRS3 are caused by NH₄⁺, the ion would also partially account for the overall broad absorption in the 3.1 μ m feature. Indeed, some 50% of the unidentified excess absorption between 3.25 and 3.8 μ m (3080–2630 μ m⁻¹), which cannot be accounted for by H₂O ice (Smith et al. 1989), could arise from the ion.

If the 3.26, 3.48, and 6.85 μ m features are all due to the same carrier, a tight correlation between these bands is expected. Ground based observations of the 3.26 μ m feature exist for four of the objects whose 6.85 μ m feature has been observed by the SWS (Table 3). For these objects the data give a stable ratio $\tau(3.26)/\tau(6.85) = 0.22 \pm 0.02$. The constancy of the ratio is even somewhat better than what could be expected on the basis of the rather large uncertainty in the depth of the 3.26 μ m band (Table 4). Besides from telluric interference, this error derives from the baseline definition, since this shallow feature is located in a complex spectral region, where various spectral features are located, such as the broad feature of ammonia hydrate around 3.5 μ m (2860 cm⁻¹) and the 3.36 μ m (2980 cm⁻¹) feature of CH₃OH, as well as possibly structure caused by scattering (Dartois & d'Hendecourt 2001;

Brooke et al. 1999). These features may contribute to the broad long wavelength shoulder of the 3 μ m band, on which the feature is superimposed.

For some of these objects ISO-SWS observations on the $3.26 \,\mu\text{m}$ feature have also been reported (Bregman et al. 2001). Using these data, there is little or no correlation between the 3.26 and 6.85 μ m bands (Table 3). Indeed, the ground based and ISO data give conflicting results on the depth of the 3.26 μ m band, most notably in the case of ρ Oph Elias 29, where the discrepancy is at least a factor of 3. It not quite clear what causes these differences and which dataset is most reliable. For Elias 29, Bregman et al. used a speed 3 scan in AOT SWS1. However, a higher quality SWS6 scan in this region does not show the 3.26 μ m feature, consistent with the ground-based observations (Boogert et al. 2000; Boogert, private communication). A general problem with the ISO data could be their wide aperture. For this reason, the 3.3 μ m emission feature often shows up in the SWS spectra of embedded YSO's (Gibb et al. 2001; see also Fig. 12). Indeed, some 3.3 μ m emission seems consistent with the ISO spectra of S140 and GL 989 (Gibb et al. 2001). The presence of even a small amount of this feature would severely complicate a correct baseline definition and therefore introduce a large error in the depth of the shallow 3.26 μ m band. On the other hand, in ground-based observations the center of the 3.26 μ m band has an enhanced noise level due to telluric methane absorption. However, the ISO and ground-based observations also deviate for the 3.48 μ m band (Table 4), even though it falls in a relatively undisturbed region. This deviation therefore reinforces the concern that errors in the baseline definition caused by the $3.3 \,\mu\text{m}$ emission feature is a major pitfall when using ISO-SWS data to study the 3.26 and 3.48 μ m absorption bands.

Good candidates to search the 3.26 μ m feature are NGC 7538:IRS9 and W3:IRS5, for which we predict, from the strength of the 6.85 μ m band, $\tau(3.26) \approx 0.064$ and 0.055, respectively. A detection of these bands would be an important test of the NH⁴₄ assignment.

No tight correlation exists between the 3.26 μ m and 3.48 μ m features. Relative intensities vary between 0.13 (MonR2:IRS3) and 1.3 (Elias 29; see Table 3). While the relative intensity of the feature of MonR2:IRS3 is consistent with an assignment of the 3.48 μ m band to NH⁴₄ (Fig. 13), it is clear that in general other species must also contribute to this feature. This is not too surprising, since additional structure is present in this region. In particular, the long wavelength shoulder or wing of the 3 μ m H₂O band, which extends from ~3.1–3.8 μ m (3200–2600 cm⁻¹; Smith et al. 1989; their Fig. 6) produces structure in this region (Smith et al. 1989; their Fig. 4). This absorption will blend in with any additional absorbers, e.g., the CH stretching mode of aliphatic hydrocarbons and the ~3.5 μ m feature of ammonia hydrate (e.g., Dartois & d'Hendecourt 2001).

4.4. The 3.26 μ m feature and PAHs

Previously, an assignment of the 3.26 μ m feature to the CH stretching modes of polycyclic aromatic hydrocarbons has

been proposed (PAHs; Sellgren et al. 1995; Bregman et al. 2000). Such an assignment would of course limit the contribution of the $v_2 + v_4$ NH⁺₄ feature to this band (Sect. 4.3). Here we quantitatively assess the contribution of the PAH CH stretching mode to the 3.26 μ m feature of MonR2:IRS3.

In support of, at least, part of the 3.26 μ m feature deriving from the PAH CH tretching mode is the observation of a weak absorption band at $11.2 \,\mu m$ towards MonR2:IRS3, which is ascribed to the out-of-plane bending mode of isolated CH groups on the aromatic skeleton (Bregman et al. 2000). The intensity of this band can be used to constrain the PAH contribution to the 3.26 μ m feature. The integrated optical depth are $\tau_{int}(11.2) = 0.81 \text{ cm}^{-1}$ and $\tau_{int}(3.26) = 5.6 \text{ cm}^{-1}$. The ratio of the band strength of the CH stretching mode to that of the CH bending mode for isolated CH groups equals 0.5 for gaseous, neutral PAHs. For ionized or solid state PAHs the ratio is at least 4 times smaller (Bregman et al. 2000 and references therein). Therefore the CH groups associated with the 11.2 μ m feature can, at most, cause an absorption of $\tau_{int}(3.26 \,\mu\text{m};\text{PAH}) = 0.4 \text{ cm}^{-1}$, i.e., only 7% of the observed intensity. Besides isolated CH groups, aromatic species will have CH groups on their periphery which have 1 or more adjacent CH groups. For these the CH bending mode falls at longer wavelength (11.6–13.6 μ m; Hony et al. 2001). Besides discrete features near 12.7 and 13.5 μ m, the bending modes of these groups give rise to a broad plateau between $11-13 \,\mu m$ caused by overlapping absorptions. In emission, the fraction of the total emission associated with the CH bending modes (i.e., 11.2, 12.7, 13.5 μ m bands and the plateau) which is emitted by the 11.2 μ m band varies between 0.3–0.7 (Hony et al. 2001). Since the band strength of the CH bending mode for adjacent aromatic CH groups is two times weaker than for isolated CH groups, this implies that these groups could add $\tau_{\text{int}} = 0.4 - 1.9 \text{ cm}^{-1}$ to the 3.26 μ m feature. This implies that 14–40% of the 3.26 μ m band may be due to the PAH CH stretch. This fraction could actually be much lower if a large fraction of the PAHs would be frozen on the grains, causing the band strength of the CH stretching mode to drop by a factor ~4 (Joblin et al. 1994). Indeed, it seems probable that in a high density environment like the circumstellar regions of a young stellar object a large fraction of highly refractory molecules such as PAHs would accrete on the dust grains. In conclusion, it seems probable that the contribution of PAHs to the 3.26 μ m band is rather small. This agrees with our result in Sect. 4.3, namely, that, based on the assignment of the 6.85 μ m band to NH₄⁺, most of the 3.26 μ m feature should be caused by the NH₄⁺ $\nu_2 + \nu_4$ band.

Contrary to our result it was concluded by Bregman et al. (2000) that PAHs may account for the entire $3.26 \,\mu\text{m}$ absorption band. Some important differences between his calculations and ours must be pointed out. First, the integrated intensity of the $3.26 \,\mu\text{m}$ band was approximated by depth \times *FWHM* in this earlier paper, while we use exact numerical integration, arriving at a 1.7 times higher intensity. Second, Bregman et al. estimated that typically only 17–30% of the entire emission in the CH bending region originates from the 11.2 μ m band. However, a thorough recent survey of this region in a wide variety of

objects (Hony et al. 2001) showed that this fraction is considerably higher, i.e., 30–70%.

4.5. The abundance of NH_4^+ and the counterions

Applying the intrinsic strength of the v_4 NH⁺₄ band, the NH_4^+ column density can be derived from the observations. With $A(NH_4^+) = 4.4 \times 10^{-17} \text{ cm ion}^{-1}$ (Sect. 3.3; uncertainty factor 2), we find for W33A, with an integrated optical depth of $\tau_{int}(6.85 \ \mu m) = 57 \ cm^{-1}$ (after correction for the CH₃OH contribution), $N(\text{NH}_{4}^{+}) = 1.3 \times 10^{18} \text{ cm}^{-2}$. For W33A we adopt $N(H_2O) = 1.2 \times 10^{19} \text{ cm}^{-2}$. This number was derived by Gibb et al. 2000 by fitting the blue wing of the 3 μ m band with laboratory ice bands. It must be noted that those authors adopted a slightly lower column density as their end result, based, in part, on matches of the blue wing of the 3 μ m band of W33A with other interstellar 3 μ m features. However, the correspondence of the W33A 3 μ m band with the other interstellar features was considerably less than the correspondence with the selected laboratory ice bands (presumably caused by source to source variations in ice composition and temperature). Therefore, we prefer to adopt here the value which is solely based on the laboratory comparison. This gives an NH_4^+ abundance of 11% for W33A, with an uncertainty of a factor 2 due to the bandstrength error.

From the data summarized in Table 3, it can be derived that the NH₄⁺ abundance varies from 3% to 17% relative to H₂O. Here we use a constant correction for the contribution of methanol to the 6.85 μ m band of 20%, as for W33A. This is perhaps a bit large, since in general the methanol abundance in YSO's is lower than for W33A (Dartois et al. 1999b). However, the correction is small anyway, and no attempt was made for a detailed source to source analysis. One exception is GL7009S, which has an exceptionally high abundance of solid methanol, for which we adopted a 30% correction of the 6.85 μ m feature (Dartois et al. 1999a).

Clearly, an equal amount of negative charge should be present in the ices to balance NH_4^+ . Recently, Gibb et al. (2000) investigated the full inventory of ice species along the line-of-sight towards the high mass YSO W33A. Their analysis showed that the negative species that have been detected towards this source, i.e., OCN⁻ and HCOO⁻ (assuming that OCN- is the carrier of the interstellar XCN feature; Grim & Greenberg 1987; Schutte & Greenberg 1997; Demyk et al. 1998; Hudson & Moore 2000, 2001; Novozamsky et al. 2001) have abundances of 3.2% and 0.8%, respectively (these values are slightly lower than those of Gibb et al., due to the 10% higher H_2O column we adopt for W33A). Since the NH_4^+ abundance for this source is $\sim 11\%$, this implies that only $\sim 30\%$ of the countercharge is provided by the observed negative species. Our experiments suggest that the residual charge could be provided by a combination of ions such as HCO_3^- , NO_2^- , NO_3^- . As was shown, a mixture of these species embedded in an H₂Odominated ice does not produce significant spectral structure. Their total abundance would be $\sim 7\%$ of H₂O for W33A. For other sources, the fraction of the charge balance that can be provided by the observed negative species (i.e., OCN⁻) is even smaller (Gibb et al. 2000). Thus, assuming that they make up for the residual balance, the total abundance of "invisible" anions in the ices near YSO's would be in general \sim 70–100% of NH⁴₄, i.e., 3–15%.

It has been argued that the poor correlation between the 4.62 μ m feature of OCN⁻ and the 6.85 μ m band disagrees with an identification of this feature with NH₄⁺ (Keane et al. 2001; cf. Figs. 7 and 12). However, as pointed out above, OCN⁻ provides only a small fraction ($\leq 20\%$) of the countercharge. In such a case, a clear correlation between the OCN⁻ and NH₄⁺ abundances is only expected if the make-up of the anion mixture is stable between YSO's. The lack of correlation shows that the fraction of OCN⁻ in the total ensemble of anions strongly varies.

5. Astrophysical implications

When the NH_4^+ assignment was originally made (Grim et al. 1989a), the identity of the counterions was uncertain. This was widely considered a prime concern (Schutte et al. 1996a; Keane et al. 2001; Demyk et al. 1998; Tielens & Whittet 1997). Our new experiments demonstrate that in H₂O dominated ices the charge balance can be achieved without introducing spectral structure of negative ions that is inconsistent with the observations. Thus it appears that the objections to the NH_4^+ assignment have been met. All things considered, NH_4^+ now stands as the best assignment for the interstellar 6.85 μ m feature (apart form the generally small fraction caused by CH₃OH; Sect. 4), based on the band's position, shape, and width. Moreover, the ammonium ion matches both the 6.85 and 3.26 μ m features, and is able to reproduce the temperature-dependent redshift of the 6.85 μ m band. Chemical considerations also support the assignment as NH_4^+ is easily produced by either photolysis or ion irradiation of known, or suspected, interstellar molecules, or by mild warm-up of ammonia and acids which could be present in the ice (see below). A straightforward observational test would be to probe the tight correlation between the 3.26 and the 6.85 μ m features which is expected if both are due to NH⁺₄.

It is unclear whether interstellar NH_4^+ could arise from irradiation or photolysis as in our experiments. The weakening of the 3.4 μ m CH stretching mode of aliphatic grain material in going from diffuse to dense regions indicates that energetic processing could play an important role in dense regions (Muñoz Caro et al. 2001; Mennella et al. 2001). However, the processing of our samples gives rise to some weak features which have not yet been observed, most notably the O₃ band at 1038 cm⁻¹, and the unidentified 1270 cm⁻¹ feature (Fig. 2). Neither of these bands are visible in the ISO-SWS observations of YSO's (Fig. 9; Gibb et al. 2001). Alternatively, the formation of NH_4^+ may not be associated with energetic processing at all. Perhaps NH₃ and acids like H₂CO₃, HNO₃, HNO₂, HNCO or even H₂SO₃ and H₂SO₄ can be formed by surface chemistry. Whether such relatively large species are formed by surface reactions depends on the extent at which the exothermicity of the reactions can be channeled into translational energy of the products (Brown 1990; Hasegawa et al. 1992). Acid-base reaction and ion formation could also be part of the surface reaction

sequence, or be promoted by warm-up or a small amount of radiation (Novozamsky et al. 2001; van Broekhuizen et al. 2003).

While in this view the anions would be undetectable in the solid state, possibly the corresponding acids could be found in the gas phase in the hot cores of star forming regions where the ice mantles evaporate. A search for molecules like H_2CO_3 , HNO_3 , HNO_2 or H_2SO_3 in such regions seems therefore warranted.

From the limited sample of objects listed in Table 3 it is unclear whether NH_4^+ has different abundances for low and highmass objects. The low-mass protostar Elias 29 in the ρ Oph cloud has an NH_4^+ abundance of 3%, substantially less than for the high mass sources. However, the other low mass object in our sample, HH100, gives a relatively high abundance of 8% (albeit with a large errorbar). More observations of the 6.85 μ m feature of low mass protostars and background field stars are desirable to clarify which dense environments favor the formation of NH_4^+ .

In hot core regions where elevated grain temperatures cause sublimation of the icy mantles, large gas phase abundances of ammonia have been found $((1-10) \times 10^{-6}$ relative to hydrogen; Blake et al. 1987; Heaton et al. 1989; Cesaroni et al. 1994). This considerably exceeds the abundance of NH₃ in cold dense regions $(N(NH_3)/N(H_2) \approx 10^{-7}$; Federman et al. 1990), indicating that ice sublimation is an important source of NH₃. However, recent observations failed to detect the 2.21 μ m feature of NH₃ towards W33A, indicating that the NH₃ abundance in interstellar ice may be low (<5%; Taban et al. 2003). A high abundance of solid NH₄⁺ in interstellar ices could alleviate this dilemma, since sublimation of this compound takes place during warm-up upon a reverse acid-base reaction and the reformation of NH₃.

 NH_4^+ can only be formed from NH₃. The presence of solid NH_4^+ thus indicates that solid NH₃ was efficiently formed during the ice accretion, presumably by hydrogenation of atomic nitrogen on the grain surface (Hiraoka et al. 1995). Whether in dense regions gaseous nitrogen will reside in its molecular or its atomic form depends on the depletion of oxygen, since reactions of atomic nitrogen with OH is an essential step towards N₂ formation (Charnley & Rodgers 2002). Thus the quantity of NH_4^+ that is present in the ices gives important information on the gas phase conditions at the epoch of condensation. Thus probing the strength of the 6.85 μ m band could become an invaluable tool for studying the chemical conditions associated with the different regions in dense clouds, i.e., quiescent, as well as near embedded high mass and low mass YSO's.

6. Conclusions

Experiments involving UV photolysis of H₂O/CO₂/NH₃/O₂ ice mixtures efficiently produce ammonium (NH₄⁺) together with probably HCO₃⁻, NO₃⁻, and NO₂⁻. These ions are produced by acid-base reactions between the NH₃ and photo-chemically produced acids. The ν_4 mode of NH₄⁺ gives a strong feature at 6.85 μ m. Other NH₄⁺ features fall at 3.27 and 3.50 μ m. Due to strong interaction of the ion with the matrix, the ν_4 NH₄⁺ band shifts strongly redward with increasing

temperature. The infrared signature of the counterions is very weak for H₂O-dominated ices.

Comparison to the 6.85 μ m feature towards embedded young stellar objects shows that the ν_4 NH₄⁴ feature obtained in our experiments provides a good match. The observed red shift of the interstellar band with increasing line-of-sight dust temperature is also consistent with this assignment. Furthermore, features are observed at 3.26 and 3.48 μ m towards these objects which could be (partly) due to the ammonium ion. The implied abundance of NH₄⁴ relative to H₂O is 3–17%.

The two anions observed in interstellar ices, OCN^- and (possibly) HCOO⁻, are insufficiently abundant to balance the positive charge from the NH_4^+ present. Therefore heavier negative ions of the kind formed in our experiments appear to be required. It is yet unclear whether such ions are formed by energetic processing, as in the laboratory experiments, or by grain surface chemistry.

Future laboratory studies should include analysis by mass spectroscopy to exactly define the nature of the photochemically produced acids. Sulfur could be added to the mixture to extend the number of acids that are produced. On the observational side, the NH₄⁺ assignment could be tested by probing the correlation between the 3.26 and 6.85 μ m bands. In addition, it would be very interesting to search for rotational emission bands of evaporating acids like HNO₂, H₂CO₃ or H₂SO₃ in hot core regions.

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