ABSTRACT

Polycyclic aromatic hydrocarbons (PAHs) have been observed in absorption along lines of sight toward embedded protostars. In such cold space environments PAHs should condense into H2O-rich ice mantles at low temperature and be exposed to ionizing radiation. In this paper we present the first infrared spectra of PAH cations in solid H2O generated under conditions that resemble dense molecular clouds. After exposing PAHs in solid H2O at 15 K to low doses of UV radiation, we have observed both the vibrational absorptions of PAH cations in the mid-IR and electronic transitions in the near-IR. The PAHs observed as ions in solid H2O were naphthalene, anthracene, phenanthrene, benzo[k]fluoranthenes, and benzo[g]juliperylene. Peak positions, strengths, and temperature dependence are reported for the detected ion bands, and their astrophysical significance is discussed. These laboratory measurements suggest that absorption bands of PAH cations in H2O ice may be observable by astronomers in the near- and mid-infrared.


1. INTRODUCTION

Polycyclic aromatic hydrocarbons (PAHs) and related aromatic materials are proposed to be present in virtually all phases of the interstellar medium (ISM; Allamandola et al. 1999; Cox & Kessler 1999), including lines of sight toward protostellar objects. Infrared (IR) absorptions attributed to PAHs have been observed in spectra of objects embedded in dense clouds including features near 3030 cm\(^{-1}\) (3.3 \(\mu\)m; Smith et al. 1989; Sellgren et al. 1995; Brooke et al. 1999), 1600 cm\(^{-1}\) (6.2 \(\mu\)m; Chiar et al. 2000), and 890 cm\(^{-1}\) (11.2 \(\mu\)m; Bregman et al. 2000). H2O is by far the most abundant solid component present along these lines of sight, suggesting the possibility that some of the PAHs responsible for these absorption bands may be condensed into H2O-rich ices. We have previously presented laboratory IR spectra of neutral PAHs in H2O (Sandford et al. 2004; Bernstein et al. 2005) for comparison to the interstellar absorption features, but to our knowledge no IR spectra of PAH cations in H2O ices exist in the literature.

In our first paper on the ultraviolet (UV) photochemistry of PAHs in H2O (Bernstein et al. 1999), we hypothesized, and then others calculated (Rica & Bauschlicher 2000), that PAH cations are possible intermediates in the reaction(s) that formed oxidized PAHs. However, even if this were true, it does not necessarily follow that PAH ion intermediates would be abundant enough in the ices to be observed. However, the electronic transitions of PAH cations in solid H2O were observed by Gudipati (2004) and Gudipati & Allamandola (2006), suggesting that they can be present at abundances high enough that absorptions of PAH cations in solid H2O might be observable in the mid-IR. This prompted us to generate PAH cations in solid H2O, and identify their IR absorptions by comparison with previously measured spectra of PAH cations isolated in argon matrices (e.g., Hudgins & Allamandola 1995a, 1995b; Hudgins et al. 2000).

It is to be hoped that this report of IR absorptions of PAH cations in H2O ice will facilitate the detection of PAH cations in dense molecular clouds where neutral PAHs have already been reported in absorption (Smith et al. 1989; Sellgren et al. 1995; Brooke et al. 1999; Chiar et al. 2000; Bregman et al. 2000). In addition, detections of aromatics on icy Saturnian satellites (Clark et al. 2005; Cruikshank et al. 2005, 2007) via the Visual IR Mapping Spectrometer (VIMS) aboard the Cassini spacecraft suggest that PAH cations in H2O could also be present in our own solar system.

Finally, we have shown that low-temperature PAH photochemistry produces oxidized aromatic compounds (Bernstein et al. 1999, 2003) much like those in meteorites (Cronin & Chang 1993; Cody & Alexander 2005) and interplanetary dust particles (IDPs; Flynn et al. 2004). Studies of the organic molecules in primitive meteorites and IDPs show these objects contain deuterium enrichments that are best explained by an interstellar dense cloud chemical heritage (Sandford et al. 2000; Sandford 2002), consistent with reactions involving PAH cations in H2O ice at low temperature. Thus, this work also pertains to those branches of astrobiology and meteoritics that examine the organic matter delivered to the Earth by meteorites and IDPs, a majority of which are aromatic.

The experimental techniques used for this work are summarized in § 2, and the data obtained from the various PAH cations in H2O are shown in § 3. Some interpretations of the laboratory results and astrophysical implications are presented in § 4.

2. EXPERIMENTAL TECHNIQUES

2.1. Sample Preparation

The H2O-PAH samples studied in this paper were vapor deposited as submicron-thick films on a CsI window. The window was held at \(\sim 15 \text{ K}\) and was suspended in an evacuated sample chamber at a pressure of \(\sim 10^{-8} \text{ torr}\). Detailed descriptions of the deposition process and apparatus can be found elsewhere (Hudgins & Allamandola 1995a; Mattioda et al. 2005).\(^1\)

All the PAHs examined in this study are solids at room temperature and, except for naphthalene (C10H8), all have a fairly low vapor pressure. As a result, only the H2O-naphthalene samples were deposited from a bulb premixed in the gas phase at room temperature before deposition, as was done in our previous study on the IR spectra of neutral naphthalene in H2O (Sandford et al. 2004). For all the other compounds, H2O and the PAH under study were codeposited onto the CsI substrate through separate inlets. In both cases the H2O was deposited from a gas bulb containing H2O.

\(^{1}\) See also http://www.astrochemistry.org.

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vapor prepared from a liquid sample on a glass manifold with a background pressure of \(1 \times 10^{-5}\) mbar. Since the pressures in the \(\text{H}_2\text{O}\) sample bulbs were 10–20 mbar, contaminant levels associated with the \(\text{H}_2\text{O}\) were negligible. The flow rate of \(\text{H}_2\text{O}\) onto the cold window was controlled by a microflow meter and by the pressure of \(\text{H}_2\text{O}\) in the bulb.

The PAHs were sublimed onto the substrate from a Pyrex tube wrapped in resistive heating wire (see Hudgins & Allamandola 1995a). The temperature to which the tube was heated depended on the sublimation point of the PAH in question and the approximate \(\text{H}_2\text{O}/\text{PAH}\) ratio desired. The temperatures used for deposition of the different PAHs were: anthracene (60°C), phenanthrene (130°C), benzo[k]fluoranthene (130°C), and benzo[ghi]perylen (150°C).

The PAH sample tube takes minutes to come up to temperature, and thus, it takes minutes for the PAHs to reach a steady rate of sublimation. To prepare samples with a uniform PAH concentration, we bring the PAH tube up to temperature with the substrate in a position shielded from the sample inlets. In all cases the samples were not permitted to deposit onto the CsI substrate until after the temperature of the PAH sample tube and the \(\text{H}_2\text{O}\) flow rate had stabilized. Similarly, the sample deposition was halted by rotating the sample to the shielded position as well as shutting off the flow of sample.

The \(\text{H}_2\text{O}-\text{PAH}\) sample gases were typically codeposited onto the 15 K CsI window at a rate that produced an 0.1 \(\mu\)m thick ice layer after a few minutes of deposition based on the observed strengths of the \(\text{H}_2\text{O}\) absorption features in the sample spectra (see §2.2). This deposition technique produces an ice composed of an intimate mixture of the PAH molecules in \(\text{H}_2\text{O}\). We know that pure \(\text{H}_2\text{O}\) deposited under these conditions is in its high-density amorphous form (Jenniskens & Blake 1994; Jenniskens et al. 1995). Our IR spectra of neutral PAHs in \(\text{H}_2\text{O}\) ices are consistent with the \(\text{H}_2\text{O}\) being amorphous (Sandford et al. 2004; Bernstein et al. 2005), the dominant form of \(\text{H}_2\text{O}\) ice present in interstellar molecular clouds (Allamandola & Sandford 1988; Jenniskens & Blake 1994; Jenniskens et al. 1995).

After deposition onto the CsI window, IR spectra of the samples were measured at a resolution of 1 \(\text{cm}^{-1}\) with a Biorad FTS 3000 spectrometer with LN\(_2\)-cooled mercury telluride (MCT) detector and salt beamsplitter for the mid-IR and an ambient temperature deuterated triglycine sulfate (DTGS) detector and quartz beamsplitter for the near IR. Analyses of the areas of the resulting IR absorption bands of both the \(\text{H}_2\text{O}\) and PAHs verified that our sample ice layers had thicknesses of a few tenths of a micron and had \(\text{H}_2\text{O}/\text{PAH}\) ratios falling between 30 and 100 (see §2.2). In some cases, additional spectra were subsequently measured after the samples had been warmed to temperatures of 50 and 100 K. Samples were warmed at a rate of \(2 \text{ K min}^{-1}\) between the temperature steps and were allowed to equilibrate for more than 5 minutes at each temperature before spectra were taken.

The \(\text{H}_2\text{O}\) used in these experiments was purified via a Millipore Milli-Q water system to 18.2 M\(\Omega\) and freeze/pump/thawed three times in vacuum to remove dissolved gases prior to use. The PAHs (purchased from the Aldrich Chemical Company unless otherwise noted) were used without any further purification, and were reported to have the following purities: naphthalene (99%), anthracene (99%), phenanthrene (99.5%+), benzo[k]fluoranthene (Janssen Chimica; 99%+), and benzo[ghi]perylen (98%).

UV photolysis was performed using a microwave-excited hydrogen flow lamp with a “McCarrol” cavity from Optohs, as described in detail elsewhere (Warneck 1962). This lamp produces \(10^{15}\) photons \(\text{cm}^{-2} \text{s}^{-1}\) evenly divided between Ly\(\alpha\) photons at 1216 Å and a several hundred nm wide broad emission band.
centered at 1600 Å. Typical samples were photolyzed for only a few minutes, corresponding to a dose of $3 \times 10^{10}$ photons cm$^{-2}$, equivalent to roughly $10^7$ yr in a dense cloud assuming the most conservative UV flux based solely on the cosmic-ray–induced UV photons at a rate of $1 \times 10^3$ cm$^{-2}$ s$^{-1}$ (Prasad & Tarafdar 1983).

2.2. H$_2$O/PAH Sample Ratios

With H$_2$O-naphthalene mixtures it was possible to carefully control the H$_2$O/C$_{16}$H$_8$ ratios by premixing samples in the gas phase. The other PAHs considered in this work have vapor pressures that are too low for gas-phase mixing, making it difficult to precisely control the ultimate H$_2$O/PAH ratio of the ice samples. We estimated the H$_2$O/PAH ratio of each sample after deposition by integrating H$_2$O and PAH peaks and ratioing the numbers of molecules calculated from the band strengths. We determined the H$_2$O column density by measuring the areas of the O–H stretching mode (3300 cm$^{-1}$; 3.0 µm) and H–O–H bending mode (1660 cm$^{-1}$; 6.0 µm) bands of H$_2$O ice and assuming inherent strengths of 1.7 x $10^{-16}$ and 1.0 x $10^{-17}$ cm molecule$^{-1}$, respectively (Hudgins et al. 1993). The column density of the given PAH in a sample was estimated by measuring the areas of multiple bands of the neutral PAH and using their calculated gas-phase band strengths (Langhoff 1996, Hudgins & Sandford 1998a, 1998b, 1998c). This method is expected to provide a good estimate of PAH column density, since our previous work on the H$_2$O-naphthalene and other H$_2$O-PAH systems has demonstrated that the presence of the H$_2$O produces modest changes in the relative band strengths of most PAH bands (Sandford et al. 2004; Bernstein et al. 2005). For all the PAHs but naphthalene, this method gave concentrations of PAHs relative to H$_2$O in the range of 1%–3%, so the PAHs in all our samples were in an H$_2$O-dominated environment (see, for example, Fig. 1). For naphthalene we were able to control the concentration by premixing the H$_2$O and naphthalene in the gas phase. This allowed us to examine the effect of concentration on ionization efficiency for this PAH. We performed photolysis on samples with naphthalene concentrations of 1% and 10% relative to H$_2$O (see, for example, Fig. 10).

The proportions of PAH ionized by photolysis were estimated by measuring the areas of the new ion bands and using literature values for the absorption strengths of the PAH cations studied in argon matrices (Hudgins & Allamandola 1995a, 1995b). PAH ion abundances determined in this manner provided ion yields in general agreement with complementary estimates based on measuring the simultaneous loss of the original PAH neutrals. The conversion of neutral PAHs to PAH ions is the dominant radiation process at low temperature and low concentrations (Gudipati & Allamandola 2006), as opposed to oxidation or reduction reactions, for example (Bernstein et al. 1999). Our estimates of the fraction of the neutral PAH ionized on exposure of our H$_2$O/PAH mixtures to UV photolysis is consistently higher than that.
observed for PAHs isolated in argon (e.g., Hudgins & Allamandola 1995a, 1995b), but less than that observed by Gudipati & Allamandola (2006) (see § 4.1).

3. RESULTS

The mid-IR spectra of H$_2$O-PAH mixtures with astrophysically relevant concentrations of PAHs are dominated by the strong vibrational-mode bands of H$_2$O, as is exemplified in the 4000–5000 cm$^{-1}$ (2.5–20 µm) mid-IR spectrum of an H$_2$O/anthracene $\approx 70$ sample shown in Figure 1. The prominent absorptions in the spectrum in Figure 1 near 3300, 1650, and 750 cm$^{-1}$ (3.0, 6.1, and 13.3 µm) are those of the O–H stretch, H–O–H bend, and librational modes of the H$_2$O, respectively.

The presence of 1% PAH (anthracene in the case of Fig. 1) produces additional, weaker, sharper peaks superimposed on the broad H$_2$O bands. This is typical for the IR spectra of neutral PAHs at these concentrations in H$_2$O (Sandford et al. 2004; Bernstein et al. 2005). The peaks of neutral anthracene seen in Figure 1 are least obscured in the region between the 1650 and 750 cm$^{-1}$ H$_2$O features, and the 1440–1120 cm$^{-1}$ (6.94–8.93 µm) region appears inset, expanded, baseline-corrected, and magnified 100 times in transmittance. It is in spectra of this sort that new cation bands must be identified after such ices are UV photolyzed.

3.1. IR Spectra of PAH Cations in H$_2$O

3.1.1. Anthracene (C$_{14}$H$_{10}$)

In previous work by Szczechanski et al. (1993) and Hudgins & Allamandola (1995b) it had been shown that when anthracene isolated in argon was exposed to UV photolysis, some of the neutral starting anthracene was converted to cations. The positions and strengths of the absorptions of the anthracene cation in argon were reported in their papers. We exposed the H$_2$O/anthracene $\approx 70$ mixture (shown above in Fig. 1) to exactly the same lamp and dose of UV photolysis that Hudgins & Allamandola (1995b) used to produce the anthracene cation in argon and obtained completely analogous results.

In Figure 2 we return to the 1440–1120 cm$^{-1}$ (6.94–8.93 µm) region of the IR spectrum seen inset in Figure 1, but now focusing on changes that occur with UV photolysis. The top two traces are spectra of anthracene isolated in solid argon at 15 K before (Fig. 2, line “A”) and after (line “B”) exposure to UV photolysis, and the bottom two are spectra of the above H$_2$O/anthracene $\approx 70$ mixture at 15 K before (line “C”) and after (line “D”) exposure to UV photolysis. The peaks of the anthracene cation in argon previously assigned by Hudgins & Allamandola (1995b) are marked with dashed lines connecting them to the new peaks produced by UV photolysis of the H$_2$O-anthracene sample.

There is a remarkable correspondence, both in position and relative intensity, between the peaks of the anthracene cation in argon and the new bands in the spectrum of H$_2$O/anthracene $\approx 70$ after exposure to UV photolysis. An exception in this and some subsequent experiments is a new peak near 8.06 µm (1240 cm$^{-1}$), which we routinely see in pure H$_2$O photolysis control experiments, suggesting this is simply an H$_2$O photolysis product. The relative areas of the peaks of neutral anthracene diminish by 30% on average after photolysis, suggesting that approximately one-third of the starting anthracene has been converted to another species, presumably the cation. The positions and relative intensities for these and all subsequent mid-IR cation peaks can be found in Table 1.

In addition to the new mid-IR features (seen in Fig. 2) exposure of the H$_2$O/anthracene $\approx 70$ mixture to UV photolysis also produces a strong new absorption in the near-IR that is consistent with the anthracene cation. Figure 3 shows the 14,800–13,250 cm$^{-1}$ (0.676–0.755 µm) near-IR spectrum of the same H$_2$O/anthracene $\approx 70$ mixture after exposure to UV photolysis at 15 K (bottom) compared to that of the anthracene cation (C$_{14}$H$_{10}^+$) in the gas phase (top) from Sukhorukov et al. (2004). A dashed vertical line at 13,832 cm$^{-1}$ (0.72296 µm) indicates the position of the anthracene cation isolated in an argon matrix (from Table II of Hudgins & Allamandola 1995b). We assign the broad band centered near 14,000 cm$^{-1}$ (0.714 µm) to the anthracene cation in solid H$_2$O.

3.1.2. Benzo[k]fluoranthene (C$_{20}$H$_{12}$)

In a manner analogous to that seen above for anthracene, new absorptions in mid- and near-IR spectra of benzo[k]fluoranthene in H$_2$O are seen after UV photolysis that are consistent with the benzo[k]fluoranthene cation in H$_2$O. Figure 4 depicts the 1400–1250 cm$^{-1}$ (7.14–8.0 µm) IR spectra of benzo[k]fluoranthene in solid H$_2$O at 15 K (Fig. 4, line “A”) before and (line “B”) after exposure to UV photolysis. The four prominent new peaks (marked by an asterisk) that appear after photolysis agree with those previously assigned to the benzo[k]fluoranthene cation in argon by Hudgins et al. (2000). The new ion absorption band near 1340 cm$^{-1}$ (7.46 µm) falls on top of one of the preexisting peaks of neutral benzo[k]fluoranthene, and blends with it because peak widths are greater in H$_2$O than in argon. However, given that the spectra are displayed on the same scale, the increased...
intensity at this position makes it clear there is new absorption occurring where a new ion band would be predicted based on the argon data of Hudgins et al. (2000), see Table 1.

Based on the unobscured neutral benzo[k]fluoranthene peak at 1192 cm$^{-1}$ (8.34 $\mu$m), which was seen to have diminished to 60% of its original area, we estimate that 40% of the starting PAH was converted to ion by photolysis. However, it is possible that some of the loss of starting material was a result of reaction with water to produce oxidized or hydrogenated products (Bernstein et al. 1999, 2003).

Similarly, a new strong band that appears in the near-IR spectrum of benzo[k]fluoranthene in solid H$_2$O after photolysis at 15 K (Fig. 5) is consistent with the benzo[k]fluoranthene cation. The dashed line in Figure 5 at 9520 cm$^{-1}$ (1.05 $\mu$m) marking the location of a strong absorption of the benzo[k]fluoranthene cation in solid argon (Mattioda et al. 2005) agrees well with the broad new band near 9700 cm$^{-1}$ (1.03 $\mu$m) seen in the spectrum of benzo[k]fluoranthene after photolysis in H$_2$O. Again, the strength of this feature suggests it is an electronic transition. Moreover, we note that this apparent cation peak is diminished after warming to 100 K at 2 K minute$^{-1}$, consistent with it being an unstable species such as a cation.

3.1.3. Phenanthrene (C$_{14}$H$_{10}$)

In a manner akin to that described above, IR spectra of the PAH phenanthrene (C$_{14}$H$_{10}$) in solid H$_2$O and exposed to UV photolysis at 15 K display new peaks that are consistent with previous measurements of the phenanthrene cation in argon. Figure 6 shows the 1350–1100 cm$^{-1}$ (7.41–9.09 $\mu$m) IR spectra of the PAH phenanthrene in solid H$_2$O at 15 K before and after exposure to UV photolysis. Four new peaks (marked by an asterisk) agree with those previously assigned to the phenanthrene cation in argon by Hudgins & Allamandola (1995a; see also our Table 1). The new ion absorptions near 1140 cm$^{-1}$ (8.77 $\mu$m) fall on top of a preexisting peak of neutral phenanthrene. By comparing the relative areas of the peaks of neutral phenanthrene in Figure 6 before and after photolysis, i.e., those at 1246, 1203, and 1166 cm$^{-1}$ (8.026, 8.313, and 8.576 $\mu$m), we find that 40% of the starting phenanthrene has been converted to another species, presumably the cation.

Figure 7 shows the position of a new near-IR band at 11,100 cm$^{-1}$ (0.901 $\mu$m) that is formed on UV photolysis of phenanthrene in H$_2$O. This band coincides remarkably well with that of the electronic transition of the phenanthrene cation in argon, indicated by a dashed vertical line (Hudgins & Allamandola 1995a; Mattioda et al. 2005). The absorptions in H$_2$O at 15 K differ by less than 10 cm$^{-1}$ (0.001 $\mu$m). In contrast to the benzo[k]fluoranthene cation (Fig. 5), which continued to persist even after warming to 100 K, the behavior of the band in Figure 7 with warming at 2 K minute$^{-1}$ suggests that the phenanthrene cation is not as stable under these conditions; it diminishes considerably by 50 K, and is no longer detectable by 100 K.

3.1.4. Benzo[ghi]perylene (C$_{22}$H$_{12}$)

As with the previous PAHs, IR spectra of benzo[ghi]perylene (C$_{22}$H$_{12}$) in solid H$_2$O show, on exposure to UV photolysis at 15 K, the appearance of new absorptions consistent with the
The cation of naphthalene was more difficult to detect than those of the previous PAHs studied. We were able to detect only the single strongest of the mid-IR absorptions of the naphthalene cation in H₂O after photolysis (see below). In addition, we were not able to observe the lowest lying electronic transition of the naphthalene cation in solid H₂O, because it falls near 14,500 cm⁻¹ (0.690 µm) in H₂O ice (Gudipati 2004), slightly longward of its position in an argon matrix (Salama & Allamandola 1991), but still beyond the reach of our near-IR detector.

However, an advantage of naphthalene is that it is volatile enough that one can easily premix it with H₂O vapor and control its concentration (§ 2.2). Figure 10 compares the 1230–1200 cm⁻¹ (8.13–8.33 µm) portion of the mid-IR spectra of photolyzed H₂O-naphthalene mixtures before and after photolysis at two different concentrations (H₂O/naphthalene = 10 and 70). In the H₂O/naphthalene = 70 case the strength of the putative cation peak at 1217 cm⁻¹ (8.217 µm) is greater despite a lower starting neutral naphthalene concentration. Apparently, lower concentrations of naphthalene in H₂O permits a higher proportion to be converted to ion, presumably because of fewer naphthalene-naphthalene interactions or less shielding than occurs at higher naphthalene concentrations. The matter is somewhat complicated by the overlap between the cation peak and an absorption of the neutral naphthalene starting material at 1212 cm⁻¹ (8.251 µm). As in previous cases, the position of the corresponding band of the cation isolated in argon (Hudgins et al. 1994) is indicated by a dashed vertical line.

4. SUMMARY, DISCUSSION, AND IMPLICATIONS

4.1. Laboratory Measurements

Submicron-thick films of five different PAHs (naphthalene, anthracene, phenanthrene, benzo[k]fluoranthene, and

![Fig. 6](image)

**Fig. 6.** The 1350–1100 cm⁻¹ (7.41–9.09 µm) IR transmission spectra of the PAH phenanthrene (C₁₄H₁₀) in solid H₂O at 15 K (line “A”) before and (line “B”) after exposure to UV photolysis. The new peaks marked by an asterisk (*) agree with those previously assigned to the phenanthrene cation in argon by Hudgins & Allamandola (1995a). We therefore attribute them to the phenanthrene cation in solid H₂O. The new absorptions near 1140 cm⁻¹ (8.77 µm) fall on top of a preexisting peak of neutral phenanthrene. The two spectra are on the same scale, but the spectra are offset for clarity. The peaks of neutral C₁₄H₁₀ in the lower spectrum have diminished relative to those above, corresponding to conversion of 25%–30% of the original neutrals into cations.

![Fig. 7](image)

**Fig. 7.** The 12,200–10,000 cm⁻¹ (0.82–1.0 µm) near-IR transmission spectra of the PAH phenanthrene in solid H₂O at 15 K after UV photolysis (bottom) and after warming to 50 and 100 K. The new broad band centered near 11,100 cm⁻¹ (0.901 µm) corresponds remarkably well to that at 11,110 cm⁻¹ (0.900 µm) assigned to the phenanthrene cation in argon by Hudgins & Allamandola (1995a) and Mattioda et al. (2005). We note that the feature diminishes on warming and is essentially gone by 100 K. These spectra are displayed on the same scale, but offset for clarity.

benzo[ghi]perylene cation. In Figure 8, new peaks in the 1600–1100 cm⁻¹ (6.25–9.09 µm) region of the mid-IR spectrum (marked with an asterisk) correspond well to peaks previously assigned to the benzo[ghi]perylene cation in argon by Hudgins & Allamandola (1995a; see also our Table 1). The relative areas of the peaks of neutral before and after photolysis suggests that only 15%–20% of the starting benzo[ghi]perylene has been converted to another species, presumably the cation.

The near-IR spectrum of the UV photolyzed H₂O-benzo[ghi]perylene mixture presented in Figure 9 contains two strong new absorptions near 13,150 cm⁻¹ (0.7605 µm) and 12,100 cm⁻¹ (0.826 µm), the latter being twice as broad as the former. The centers of the three near-IR peaks of the benzo[ghi]perylene cation in argon identified by Hudgins & Allamandola (1995a) and Mattioda et al. (2005) at 13,050, 12,480, and 12,190 cm⁻¹ (0.766, 0.8013, and 0.8203 µm, respectively) are indicated by dashed vertical lines. The two near-IR peaks we observed for the photolyzed H₂O-benzo[ghi]perylene mixture correspond quite well with the shortest and longest wavelength peaks of these electronic transitions of the benzo[ghi]perylene cation in argon. The middle, 12,480 cm⁻¹ (0.8013 µm), peak of the benzo[ghi]perylene cation in argon is presumably blended with the longer wavelength peak we observe at 12,190 cm⁻¹ (0.8203 µm), and this is why it is broader than the other.

3.1.5. Naphthalene (C₁₀H₈)

The cation of naphthalene was more difficult to detect than those of the previous PAHs studied. We were able to detect only
benzo[ghi]perylenes) frozen individually at the 1%–3% level in H₂O at 15 K and then exposed to UV photolysis show mid-IR absorptions indicative of the formation of PAH cations (see Table 1 for a summary). The positions and relative proportions of these peaks in the mid-IR are consistent with previously measured IR spectra of these cations isolated in argon (Hudgins et al. 1994, 2000; Hudgins & Allamandola 1995a, 1995b) and the greater peak widths are consistent with previously measured mid-IR spectra of PAH neutrals in H₂O (Sandford et al. 2004; Bernstein et al. 2005).

In addition, for all but naphthalene, new strong bands were also observed in the near-IR, and these are consistent with previous measurements of electronic transitions of the corresponding PAH cations isolated in argon (Hudgins & Allamandola 1995a, 1995b; Mattioda et al. 2005) and/or in the gas phase (Sukhorukov et al. 2004). Whereas the mid-IR peak positions of the PAH cations in H₂O at 15 K were always within the peak width (a few cm⁻¹) of the corresponding absorptions of the PAH cation isolated in argon, the stronger electronic transitions in the near-IR were seen to differ by as much as 180 cm⁻¹ (0.02 µm) from those in argon. This behavior is consistent with that observed for other PAHs (Gudipati 2004, and references therein). The greater variance of the position of the near-IR absorptions (compared to the mid-IR) is consistent with the fact that electronic transitions are well-known to be more sensitive to matrix interactions than are vibrational transitions.

The peaks attributed to the PAH cations in H₂O diminish on warming, but not all at the same temperature; the benzo[k]fluoranthene cation in solid H₂O is still present at 100 K, but the phenanthrene cation is not (compare Figs. 5 and 7).

The peak areas of the starting PAHs diminished by up to 40% when exposed to UV photolysis (as seen in Figs. 2, 4, 6, and 8). Presuming that the entire loss of starting material corresponds to formation of the cation, this is a higher ionization efficiency than was observed previously for these same PAHs in argon (Hudgins & Allamandola 1995a, 1995b), but lower than that inferred for other PAHs in H₂O by Gudipati & Allamandola (2006). It is reasonable to suppose that there should be a higher proportion ionized in H₂O because of its greater capacity to stabilize both the cations and the ejected electrons, and this probably explains why we report higher fractions of the PAHs ionized than did Hudgins & Allamandola (1995a, 1995b) in argon. Furthermore, Figure 10 strongly suggests a concentration effect, with more dilute mixtures of PAHs ionizing to a much greater extent. This may be the explanation for the lower proportion of ions produced in this work where the concentrations were 1%–3%, compared to that reported by Gudipati & Allamandola (2006) where the concentrations were 0.2%–0.5%.

4.2. Astrophysical Implications

IR astronomy suggests aromatics are ubiquitous in the interstellar medium (Allamandola et al. 1999; Cox & Kessler 1999) and the presence of deuterium-enriched and oxidized aromatics in carbonaceous meteorites (Cronin & Chang 1993; Cody & Alexander 2005) and IDPs (Flynn et al. 2004) suggests that
those aromatics may have experienced radiation at low temperature in H$_2$O-rich ices (Sandford 2002). The laboratory results reported here confirm that PAHs are ionized by UV in solid H$_2$O, and show that the resulting PAH cations can be directly observed in the IR. The IR absorptions of neutral aromatics in the ISM have already been observed (Smith et al. 1989; Sellgren et al. 1995; Brooke et al. 1999; Chiar et al. 2000; Bregman et al. 2000).

Given the prevalence of H$_2$O ice, both in the dense ISM and in the outer solar system, the modest radiation required to produce ions, and their apparent stability at low temperature (at least in pure H$_2$O), these PAH cations should also be observable in astrophysical ices at near- and mid-IR wavelengths. Interestingly, the strongest mid-IR PAH cation bands fall between 6 and 8 μm, the region in which there are several absorption features in dense clouds that are of uncertain origin (Lacy et al. 1991; Boogert et al. 1998; Keane et al. 2001). For example, the strong absorptions of the benzol[\i]fluoranthene cation could contribute to interstellar absorption seen near 1300 cm$^{-1}$ (7.6 μm) (Boogert et al. 1998). The positions of mid-IR absorptions of PAH cations in H$_2$O matrices are very similar to those of the corresponding cation in argon, so our mid-IR database of laboratory and modeled spectra (Hudgins et al. 1994, 2000; Hudgins & Allamandola 1995a, 1995b; Langhoff 1996$^*$) should provide an adequate guide to PAH cation band positions and strengths for IR astronomers.

In the near-IR the situation is potentially more problematic; the peak positions of PAH cations in H$_2$O can shift by as much as 180 cm$^{-1}$ relative to gas-phase/ inert matrix positions because of the sensitivity of electronic transitions to matrix interactions. However, the electronic transitions in the near-IR are $10^2$--$10^3$ times stronger than the vibrational absorptions in the mid-IR, so they could be easier to detect spectroscopically. As noted by Mattioda et al. (2005), these bands could impose detectable weak broadband structure(s) on the interstellar extinction curve, especially near 10,000 cm$^{-1}$ (1 μm). These absorptions are also relevant for H$_2$O-rich surfaces in the outer solar system where near-IR spectra are most commonly measured and aromatics have been detected, such as in Cassini VIMS spectra of Phoebe (Clark et al. 2005) and Iapetus (Cruikshank et al. 2005).

PAH cations in H$_2$O ices in solar system or protostellar environments where the temperature is above 50 K may not be indefinitely stable. When warmed to 100 K, some cations survive, but some do not (compare Figs. 5 and 7; see also Gudipati & Allamandola 2006). Clearly, temperature stability of a PAH cation will depend on its structure, size, and probably on its concentration in the ice as well.

Combined, these results suggest that it should be possible to use the current database of spectra from Ar matrix-isolated PAH cations to tentatively identify PAHs cations in dense interstellar clouds, around protostars, and on icy satellites in the outer solar system.

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2 See also http://www.astrochem.org.

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