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Gas-phase spectroscopy of photostable PAH ions from the mid- to far-infrared

Sandra D. Wiersma^(D),^{1,2,3} Alessandra Candian^(D),¹ Joost M. Bakker^(D) and Annemieke Petrignani^(D)

¹Van 't Hoff Institute for Molecular Sciences, University of Amsterdam, PO Box 94157, NL-1090 GD Amsterdam, the Netherlands ²Radboud University, Institute for Molecules and Materials, FELIX Laboratory, Toernooiveld 7, NL-6525 ED Nijmegen, the Netherlands ³Institut de Recherche en Astrophysique et Planétologie (IRAP), CNRS, Université de Toulouse (UPS), F-31028 Toulouse, France

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ABSTRACT

We present gas-phase InfraRed Multiple Photon Dissociation (IRMPD) spectroscopy of cationic phenanthrene, pyrene, and perylene over the 100–1700 cm⁻¹ (6–95 μ m) spectral range. This range covers both local vibrational modes involving C–C and C–H bonds in the mid-IR, and large-amplitude skeletal modes in the far-IR. The experiments were done using the 7T Fourier-Transform Ion Cyclotron Resonance (FTICR) mass spectrometer integrated in the Free-Electron Laser for Intra-Cavity Experiments (FELICE), and findings were complemented with Density Functional Theory (DFT) calculated harmonic and anharmonic spectra, matching the experimental spectra well. The experimental configuration that enables this sensitive spectroscopy of the strongly bound, photoresistant Polycyclic Aromatic Hydrocarbons (PAHs) over a wide range can provide such high photon densities that even combination modes with calculated intensities as low as 0.01 km mol⁻¹ near 400 cm⁻¹ (25 μ m) can be detected. Experimental frequencies from this work and all currently available IRMPD spectra for PAH cations were compared to theoretical frequencies from the NASA Ames PAH IR Spectroscopic Database to verify predicted trends for far-IR vibrational modes depending on PAH shape and size, and only a relatively small redshift (6–11 cm⁻¹) was found between experiment and theory. The absence of spectral congestion and the drastic reduction in bandwidth with respect to the mid-IR make the far-IR fingerprints viable candidates for theoretical benchmarking, which can aid in the search for individual large PAHs in the interstellar medium.

Key words: astrochemistry – methods: laboratory – techniques: spectroscopic – planets and satellites: atmospheres – ISM: molecules – infrared: ISM.

1 INTRODUCTION

Polycyclic aromatic hydrocarbons (PAHs) are one of the most stable families of organic molecules known. They are found in outer space, where they are formed through combustion-like chemistry in the outflows of carbon-rich stars (Allamandola, Tielens & Barker 1989; Latter 1991). The presence of PAHs has been established in many astronomical environments, and it has been estimated that PAHs comprise between 10 and 18 per cent of the elemental carbon budget (Joblin, Leger & Martin 1992; Tielens 2013). Since the interstellar PAH hypothesis was presented in 1984 (Leger & Puget 1984; Sellgren 1984), many astronomical observations, theoretical studies, and laboratory experiments have been conducted that have led to the conclusion that the dominant Aromatic Infrared Bands (AIBs) observed in the mid-infrared (MIR) at 3.3, 6.2, 7.7, and 11.3 µm can be attributed to emissions of UV-excited PAHs in the ISM (Allamandola et al. 1989; Hudgins & Allamandola 1995; van Diedenhoven et al. 2004; van Dishoeck 2004; Tielens 2008; Peeters 2011; Montillaud, Joblin & Toublanc 2013; Candian, Sarre &

Tielens 2014; Andrews, Candian & Tielens 2016; Croiset et al. 2016; Bauschlicher et al. 2018). These and many other studies have helped put limits on the expected size of interstellar PAHs and elucidated details of their life cycles; it is now well-established that a large fraction of the PAH population in the ISM is present in cationic form (see Tielens 2008 and references therein).

Recently, the radio wave fingerprints of both pure and nitrogencontaining PAHs with two rings have been detected in the dark cloud TMC-1 (Burkhardt et al. 2021; Cernicharo et al. 2021; McGuire et al. 2021). Fingerprints of closely related species have also been detected, leading to the discovery of benzene and benzonitrile (Cernicharo et al. 2001; Kraemer et al. 2006; McGuire et al. 2018), as well as the fullerenes C_{60} , C_{60} ⁺, and C_{70} (Cami et al. 2010; Sellgren et al. 2010; Berné, Mulas & Joblin 2013; Campbell et al. 2015; Cordiner et al. 2019). The presence of fullerenes has been confirmed by both their visible (Campbell et al. 2015; Cordiner et al. 2019) and MIR (Cami et al. 2010; Sellgren et al. 2010; Berné et al. 2013) fingerprints. Benzene, C₆₀, and C₇₀ possess characteristic IR emission lines that allowed interstellar MIR features to be assigned to them. However, an infrared match with the spectrum of a single PAH is still elusive, as PAHs in the MIR exhibit highly similar spectra, dominated by local C–C and C–H modes. This is especially true for a larger ones (>50 $\,$

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^{*} E-mail: sandra.wiersma@irap.omp.eu (SDW); a.petrignani@uva.nl (AP)



Figure 1. Skeletal vibrations of cationic pyrene, visualized with displacement vectors. The scaled frequencies are taken from the harmonic DFT calculations which are treated in Section 2.2 and listed from the drumhead mode on in Table 3.

C atoms), which are suspected to be the most abundant (Andrews et al. 2016).

The scenario is different for the far-infrared (FIR) fingerprints of PAHs. Here, global, skeletal deformation vibrations, with names such as drumhead, butterfly, or jumping jack modes (see Fig. 1) are much more molecule specific (Mulas et al. 2006a, b; Ricca et al. 2010, 2012). Although possible FIR PAH emission bands are often obscured by dust emissions in interstellar objects (Draine 2003), emission features in the MIR-to-FIR transition zone, at 15.8, 16.4, 17.4, 17.8, and 18.9 µm, have been attributed to PAHs (Boersma et al. 2010; Ricca et al. 2010). Further in the FIR, bands have been suggested to be even more characteristic for individual PAHs or PAH families (Mulas et al. 2006a, b; Boersma et al. 2011b, a; Joblin et al. 2011; Ricca et al. 2012). These studies are, however, entirely based on theoretical spectra from the NASA Ames PAH Database (Bauschlicher et al. 2018), for which the FIR benchmarking is primarily based on the spectrum of a single PAH, neutral naphthalene (Pirali et al. 2009).

Laboratory studies have revealed many characteristic IR spectral properties of PAHs, but have mostly been limited to wavelengths below 20 μ m (above 500 cm⁻¹), which do not involve the truly global modes. Matrix isolation spectroscopy studies have mostly focused on neutral PAH species, partially due to difficulties in discriminating signal originating from ions or neutral (Leger, D'Hendecourt & Defourneau 1989; Blanco, Fonti & Orofino 1990; Moutou, Leger & D'Hendecourt 1995; Mattioda et al. 2009; Cataldo et al. 2013; Zhang & Sander 2017). Other difficulties including impurities and backgrounds exist for gas-phase FT-IR or emission spectroscopy (Kurtz 1992; Zhang et al. 1996; Cané et al. 1997; Pirali et al. 2006, 2009, 2013; Goubet & Pirali 2014). For neutral PAHs, iondip spectroscopy allows for detailed high-resolution investigations of, e.g. anharmonic behaviour; however, its application to ionic species is not trivial (Maltseva et al. 2015; Lemmens et al. 2019). To record laboratory spectra for gas-phase ions over large spectral ranges

(covering mid- to far-IR), IR multiple-photon dissociation (IRMPD) spectroscopy is presently one of the most powerful methods (Oomens et al. 2000, 2006).

In this work, we used the high IR photon densities available in the Fourier Transform Ion Cyclotron Resonance (FT-ICR) mass spectrometer beamline of the FELICE intra-cavity free-electron laser to perform IRMPD spectroscopy of PAH cations in the FIR up to wavelengths of 100 μ m. We present IRMPD spectra for phenanthrene (C₁₄H₁₀⁺), pyrene (C₁₆H₁₀⁺), and perylene (C₂₀H₁₂⁺), in the range of 100–1700 cm⁻¹ with particular attention to the 100–600 cm⁻¹ FIR spectral range. We assign a large range of low-frequency bands using Density Functional Theory (DFT) calculations in both the harmonic and the anharmonic approximation. The narrow FIR modes in our experimental spectra are then considered as possible benchmarking data to improve the NASA Ames PAH Database in this spectral range (Bauschlicher et al. 2018). Finally, we discuss the implications of our findings for possible astronomical detection of molecule-specific signatures in the ISM.

2 METHODS

2.1 Experimental

IRMPD spectroscopy is a multiple-photon technique wherein absorption occurs repeatedly via the same transition. The absorbed energy is dissipated over a bath of dark vibrational states and once the fragmentation limit is overcome the species of interest can dissociate, signalling an IR active vibrational mode (Oomens et al. 2006). The multiple-photon character of IRMPD spectroscopy leads in general to slight spectral broadening and redshifts from the fundamental vibrational transition, as it is favourable to achieve fragmentation when higher transitions of the anharmonic vibrational ladder are also excited. Interestingly, the latter make the detected bands a closer representation for the emissions from hot, interstellar PAHs. IRMPD intensities tend to scale linearly with laser power and the single-photon absorption cross-section once a certain threshold is overcome (Oomens et al. 2006; Bakker et al. 2007; Berden et al. 2019) but because this threshold is band- and species-dependent, IRMPD spectra can differ from the linear absorption spectra.

The IRMPD spectra were recorded using the FT-ICR mass spectrometer beam line of the Free Electron Laser for Intra-Cavity Experiments (FELICE), at the FELIX Laboratory in Nijmegen. The FELICE FT-ICR apparatus, schematically shown in Fig. 2(a), has been described in detail before (Petrignani et al. 2016; Wensink et al. 2020). Here, additional experimental details are presented on features of the FELICE FT-ICR that were used for the first time in this work.

Phenanthrene, pyrene, and perylene (Sigma Aldrich/Merck) were brought into the gas phase using an effusive sublimation source, at temperatures of 50, 70, and 130°C, respectively. The gas-phase PAHs were then exposed to 20-eV electrons in an electron impact ionizer (Ardara Technologies), creating singly-charged radical cations. These cations were led through a series of ion optics, including a linear quadrupole mass spectrometer (QMS) operated in radiofrequency guiding mode and collected a sectioned, quadrupole ion trap with rectilinear rods (RIT), and collisionally cooled with argon gas ($\sim 10^{-2}$ mbar) (Ouyang et al. 2004). After accumulating and cooling over 50 ms in the RIT, an ion pulse was extracted towards an intra-cavity quadrupole deflector (IQD), which diverted the ion trajectory by 90° into the FELICE cavity. A home-built intra-cavity radiofrequency quadrupole ion guide (IQG) transported the ions towards one of the four FT-ICR storage cells. For each cell, the



Figure 2. (a) Schematic of the FELICE FT-ICR instrument. The path of the molecules through the apparatus from the sublimation source down to the ICR cells is depicted in blue. The fuchsia beam illustrates the FELICE waist through the cavity. Proportions in this figure are not to scale. (b) Cross section of the four storage cells (C1-4) and the intra-cavity quadrupole ion guide (IQG). The ions move from right to left and are stored in one of the storage cells (C#). The top and side plates are used as excitation (exc#) and detection (det#) plates, respectively. When using one cell, the electrodes of all other cells are grounded. The focus of the FELICE laser beam coincides with the centre of C1, and is characterised by a 82 mm Rayleigh length.

mass resolution exceeds $m/\Delta m = 10^5$ for mass-to-charge ratios of up to at least 1000 m/z.

The FT-ICR storage cells are labelled C1–C4, and are depicted in Fig. 2(b). These open-ended, cylindrical cells (5 cm diameter) are located within a movable 7-Tesla superconducting magnet (Cryomagnetics Inc.). This magnet can be positioned in such a way that the volume of homogeneous magnetic field (<19 ppm) overlaps with the centre of either of the four cells. The multicell configuration was introduced to enable IR spectroscopy with varying IR fluences inside the laser cavity. C1 is positioned in the laser focus, and ions stored here experience the largest laser fluence. With a spacing between the cells of 10 cm, and the laser Rayleigh range of 82 mm, ions in each next cell are submitted to a fluence reduced by a factor of

2.3. For compounds with a relatively strong absorption efficiency, spectroscopy is preferably done in C4, as the overlap between the ion cloud and the laser beam is larger, and the reduced fluence will lead to less spectral broadening. C1 is used for the most photostable species or weakest absorptions. For pyrene and perylene, spectra were recorded in C4 in the $600-1700 \text{ cm}^{-1}$ spectral range, and spectra at lower frequencies in C1. For phenanthrene, the 235–650 cm⁻¹ spectral range was recorded using C4, and the $105-300 \text{ cm}^{-1}$ range in C1.

After transfer to the selected cell, the cations of interest were massisolated via a stored waveform inverse Fourier Transform (SWIFT) pulse (Marshall, Hendrickson & Jackson 1998). The ions were then exposed to one or several (max. 20) FELICE macropulses, depending on the spectral range and studied molecule. At 1000 cm^{-1} (10 µm), fragmentation for a PAH cation requires more than 6 eV or 50 photons to break a C–H bond, translating to more than 600 photons at (100 cm⁻¹) 100 µm (Ricca et al. 2012). For each spectrum, the IR frequency was scanned in steps of 1, 2.5, or 5 cm⁻¹, depending on the spectral range. At each frequency step, 5–20 time transients were averaged and Fourier transformed. The whole experimental sequence is controlled by home-built software developed by Mize et al. (2004).

The IRMPD spectrum was obtained by monitoring the fragmentation yield Y(v) at wavenumber v, defined as

$$Y(\nu) = \ln\left(\frac{-N_{\text{par}}(\nu)}{N_{\text{par}}(\nu) + N_{\text{frag}}(\nu)}\right)$$
(1)

with $N_{\rm frag}$ and $N_{\rm par}$ as the total fragment and parent mass counts, respectively. Y is then divided by P(v), the macropulse energy, which is inferred from coupling a fraction of the light out through a 1 mmdiameter hole of the cavity end mirror, and directing it on to a power meter. The IR wavelength was calibrated using a grating spectrometer (Princeton Instruments SpectraPro). The 100-1700 cm⁻¹ spectral range was recorded with three separate FEL settings. In the 600-1800 cm⁻¹ range, the FELICE macropulse energy peaked between 1500 and 1600 cm^{-1} at 1.6 J. In the 200–600 cm^{-1} range, it was slightly lower with its maximum at 1.3 J around 475 cm^{-1} . For the 100-300 cm⁻¹ range, the maximum was reached around 275 cm⁻¹ at 0.4 J. In spite of attempts to moderate the macropulse energy, saturation of the strongest bands in the 1000–1800 cm⁻¹ wavenumber range occurred. For these bands, all ions overlapped by the laser fragmented, leading to a maximum in the fragmentation yield. Saturated peak heights in the spectrum are still modulated by the power correction. After power correction of the three individual curves, they are joined by normalizing to the overlapping resonances, or in the case there is no overlapping resonance, by correcting for the drop in fluence from one cell to the next. The composite spectrum is then normalized to the resulting highest intensity, giving the final normalized IRMPD yield. For the presented experiments, 4-10 µs long macropulses were used, at a FELICE repetition rate of 5 or 10 Hz, which in turn consist of ps-long micropulses at 1 GHz. The spectral bandwidth (FWHM) was ~ 0.6 per cent of the central wavelength (in µm).

2.2 Computational

DFT calculations were employed to predict the IR spectra and compare to experiment, using the Gaussian16 Suite (Frisch et al. 2016). For all three molecules, geometry optimizations, and harmonic vibrational calculations were performed using the B3LYP functional (Lee, Yang & Parr 1988; Becke 1993) and 6-311++G(2d,p) (Frisch, Pople & Binkley 1984) basis set. The harmonic calculations were scaled to match the experimental frequencies by a factor of 0.96. For pyrene and perylene, we present both the scaled harmonic calculations and unscaled anharmonic calculations at the B3LYP/N07D level of theory (Barone, Cimino & Stendardo 2008; Mackie et al. 2018). The individual modes are visualized and identified using the computational chemistry software suite GABEDIT (Allouche 2011). We adopted the FIR mode naming conventions introduced by Ricca et al. (2010).

3 RESULTS

The fragments observed in the IRMPD spectroscopy of the three respective PAHs are listed in Table 1. The smallest PAH ion, phenan-

threne, exhibits both H and hydrocarbon loss. For pyrene, H-loss is dominant, and the hydrocarbon loss is very weak. For perylene, only H-loss is observed. This follows previously established experimental trends for cationic PAHs using IRMPD (Bouwman et al. 2019), collision-induced dissociation (West, Lesniak & Mayer 2019), and UV-photodissociation (Ekern et al. 1998). Theoretical studies on the reaction paths for [2C, 2H] loss processes showed that pericondensed PAHs lack the structural flexibility to form the four- and five-ringed intermediates necessary for [2C, 2H] loss, and that structural rigidity increases for larger, more aromatically stabilized PAHs (Holm et al. 2011; West et al. 2019). Furthermore, the ease with which -2H/-H2is lost from highly-excited large PAHs is also reflected in other work (Zhen et al. 2014, 2017, 2018; Castellanos et al. 2018; Rodriguez Castillo, Simon & Joblin 2018).

The IR spectra of the respective PAH ions were constructed according to equation (1) using the fragments listed in Table 1 followed by power correction and normalization, providing the normalized IRMPD yield. Minor loss channels (italicised) were omitted due to their low S/N ratios.

3.1 Phenanthrene

The IR spectrum of cationic phenanthrene ($C_{14}H_{10}^{+}$, m/z = 178) is shown in the 105–650 cm⁻¹ spectral range in Fig. 3 (black, top). The spectrum is a composite of two scans, which overlap between 240 and 290 cm⁻¹. Four well-resolved resonances are found for which frequencies and intensities are listed in Table 2. The bandwidths observed are roughly 20 cm⁻¹ full-width at half-maximum (FWHM), except for the 199 cm⁻¹ band, which has a width of ~ 5 cm⁻¹. We attribute this smaller bandwidth to a reduction of the power broadening at longer wavelengths, to the reduced FEL spectral bandwidth, and a possibly diminished anharmonicity of lower frequency modes (Joblin et al. 1995; Oomens et al. 2006; Lemmens et al. 2020; Chakraborty et al. 2021). The 390 cm^{-1} band exhibits a highfrequency shoulder, and the 571 cm^{-1} band is asymmetric, both likely caused by underlying low-intensity resonances. The shown spectrum partially overlaps the IR spectra reported by Piest et al. (2001). Piest et al. (2001) presents the IRMPD spectrum of the phenanthrene cation and an IR photodissociation spectrum of the cold phenanthrene-Ar cation down to approximately 400 cm^{-1} . For the untagged phenanthrene cation, a band was observed at 578 cm^{-1} , while the tagged spectrum reported bands at 482 and 581 cm⁻¹. The same publication also reported multiple Franck-Condon active vibrational states detected via mass-analysed threshold ionization (MATI) further into the FIR, of which only the band at 402 cm^{-1} matches with this work; the other reported bands are IR-inactive. The results from Piest et al. are consistent with this work, with a maximum frequency deviation of 10 cm^{-1} .

Our DFT-calculated spectrum of phenanthrene is also shown in Fig. 3 (bottom panel, red) with the calculated frequencies scaled by a factor of 0.96 and listed in Table 2. To facilitate comparison with the experiment, the stick spectrum is convolved with a Gaussian line shape function with a 30 cm^{-1} FWHM. One can observe that there is a good agreement between calculated and experimental band positions. The calculated relative intensities compare reasonably well, with the largest mismatch for the 479 cm⁻¹ experimental band, which is predicted far weaker than observed. The high-frequency shoulder of the 390 cm⁻¹ band is rationalised by a (weak) predicted band at 420 cm⁻¹, and the raised baseline between the 479 and 571 cm⁻¹ bands is attributed to a calculated band at 532 cm⁻¹. A detailed comparison between the experiment and the DFT calculation can be found in Table 2, along with descriptions of the vibrational modes.

Table 1. Cationic fragment masses detected in the IRMPD spectroscopy of phenanthrene ($C_{14}H_{10}^+$, m/z = 178), pyrene ($C_{16}H_{10}^+$, m/z = 202), and perylene ($C_{20}H_{12}^+$, m/z = 252). The dominant channel/fragment is denoted in bold. Detected fragment channels that were too weak to be included in the IRMPD spectra are listed in italic.

$C_{14}H_{10}^{+}$ m/z = 178		$C_{16}H$ $m/z =$	10 ⁺ 202	$C_{20}H_{12}^{+}$ m/z = 252	
fragment m/z	loss of	fragment m/z	loss of	fragment m/z	loss of
177	-H	201	-H	251	-H
176	$-2 \mathrm{H}$	200	$-2 \mathrm{H}$	250	- 2 H
152	-[2C, 2H]	199	-3 H	249	-3 H
151	-[2C, 3H]	198	$-4\mathrm{H}$	248	$-4 \mathrm{H}$
150	-[2C, 4H]	176 174	-[2 C, 2 H] -[2 C, 4 H]	246	-6 H



Figure 3. The experimental IRMPD spectrum (top) and theoretical IR spectrum (bottom) of cationic phenanthrene. The harmonic theoretical stick spectrum is scaled with a factor 0.96, and convolved with a 30 cm⁻¹ FWHM Gaussian line shape function.

Table 2. Phenanthrene cation observed band frequencies (cm^{-1}) and relative intensities, together with calculated frequencies (cm^{-1}) and IR intensities $(km \text{ mol}^{-1})$, and descriptions of the theoretical modes. The listed harmonic frequencies are scaled by a factor 0.96. *oop: out-of-plane; ip: in-plane; E/C: elongation/compression*.

Experiment		Harmonic calc.		Mode description	
Freq. (cm ⁻¹)	Norm. Y	Freq. (cm ⁻¹)	IR int. (km mol ⁻¹)		
199	0.22	204	6.3	long-axis butterfly	
390	0.18	395	9.5	quarto HCCH _{oop} twist, duo CH _{oop} wag	
		420	0.9	armchair ring rock	
479	0.48	475	3.7	ip rock central ring, E/C outer phenyls	
		532	0.7	E/C middle ring	
571	1.00	576	48.9	E/C outer phenyls	

These mode descriptions show that as we move further into the FIR, the skeletal motions become increasingly out-of-plane.

3.2 Pyrene

The IR spectrum of cationic pyrene ($C_{14}H_{10}^+$, m/z = 202 amu) is shown in Fig. 4(a) (black curve, top panel). It spans the range of 110–1590 cm⁻¹ and consists of a composite of three individual curves,

which overlap between 240-300 cm⁻¹ and 560-600 cm⁻¹. We observe twelve distinct bands, for which frequencies and intensities are listed in Table 3. The bands at 1320 and 1381 cm⁻¹ have almost merged, but two distinct shapes can still be discerned. The observed bandwidths vary between approximately 40 and 60 cm⁻¹ in the range between 450 and 1590 cm⁻¹, decrease to 30 cm⁻¹ for the features at 335 and 434 cm^{-1} , and drop sharply to 3 cm^{-1} for the feature at 188 cm⁻¹. The spectrum between 1000–1600 cm⁻¹ is possibly affected by saturation due to the high power and strong absorptions; as a consequence, laser power normalization leads to an underestimation of the relative intensity. Although the high power may thus lead to saturation, it also enables the detection of very weak modes, especially in the FIR. A vertical close-up (black, dotted curve) in the FIR section clearly reveals modes at 335 and 434 cm⁻¹ with a high signal-to-noise (S/N) ratio, and predicted intensities of only 1.2 and 0.5 km mol⁻¹, respectively. These are around two orders of magnitude weaker than the strongest band in the MIR experimental spectrum.

The 600–1590 cm⁻¹ region of our spectrum agrees well with a previously reported spectrum of Oomens et al. (2000) in terms of peak positions and band shapes. We detect an additional band at 1100 cm⁻¹ and confirm the band at 990 cm⁻¹, which was detected at a very low S/N ratio in the previous work. Panchagnula et al. (2020) also published an IRMPD spectrum of pyrene in their Supplementary Material, in the 600-1700 cm⁻¹ range, which matches even better with our spectrum. They reproduced both our experimentally observed 990 and 1100 cm⁻¹ bands with a similarly high S/N, and the shape and peak positions line up nicely. In their main paper, they presented a high-resolution IR photodissociation spectrum of cold, neon-tagged pyrene in the same range. Many of the narrow bands in that spectrum fall under one, broadened feature in their IRMPD spectrum. Notably, the left shoulders on the 840, 990, and 1230 cm⁻¹ bands in our spectrum can be explained by blending of the higher-intensity transitions with lower-intensity features. Furthermore, Panchagnula et al. report a band at 768 cm⁻¹ where we do not observe any fragmentation. The agreement is good, but this work exhibits a consistent redshift in the region of 1200-1600 cm⁻¹ compared to the Ne-tagged spectrum, to a maximum of 23 cm⁻¹, which we attribute to the single-photon character of tagging spectroscopy.

Fig. 4(a) also displays the scaled harmonic (red curve, middle panel) and the anharmonic (blue curve, bottom panel) DFT spectra. The relative intensities for both theoretical spectra match the experiments reasonably well, although there are notable deviations in intensity for the observed 671, 840, and 1515 cm^{-1} bands. Most band frequencies in the scaled harmonic spectrum match within 10 cm⁻¹,



Figure 4. (a) The experimental IR spectrum of cationic pyrene (top, black), accompanied by DFT calculated spectra in harmonic (middle/red, frequencies scaled by a factor 0.96) and anharmonic (bottom/blue, unscaled) approximations; both calculated spectra are convolved with a 30 cm⁻¹ FWHM Gaussian line shape function. The dotted curves are vertical enlargements of the solid curves, slightly raised from the baseline for clarity. (b) Close-up of 400–550 cm⁻¹ spectral range of panel (a), with a convolution of 15 cm⁻¹ for both calculated spectra.

Table 3. Pyrene cation observed band frequencies (cm^{-1}) and relative intensities, together with calculated frequencies (cm^{-1}) and IR intensities (km mol⁻¹), and descriptions of the theoretical modes. The listed harmonic frequencies are scaled by a factor 0.96. *oop: out-of-plane; ip: in-plane; E/C: elongation/compression.*

Experiment		Harmonic calculations		Anharmonic calculations			
Freq. (cm ⁻¹)	Norm. Y	Freq. (cm ⁻¹)	IR int. (km mol ⁻¹)	Freq. (cm ⁻¹)	IR int. (km mol ⁻¹)	Mode description	
188	0.390	191	12.4	203	11.7	drumhead	
335	0.036	340	1.2	352	1.2	short-axis jumping jack	
434	0.003	451	0.5	459	0.3	long-axis HCCH _{oop} twist	
488	0.130	485	7.3	500	8.3	long-axis jumping jack	
	_	518	0.5	533	0.4	short-axis E/C	
671	1.000	668	9.0	687	8.4	long-axis E/C	
	_	678	57.2	691	45.3	in-phase duo and trio CH _{oop} wag	
840	0.759	845	88.7	858	95.7	in-phase duo and trio CH _{oop} wag	
990	0.298	953	11.1	981	10.6	short-axis E/C	
	_	968	15.4	1000	11.5	long-axis E/C	
1100	0.096	1077	13.4	1108	11.1	antisymm. CH _{ip} trio scissor	
	_	1184	17.3	1217	16.3	_	
1230	0.533	1218	74.9	1246	52.4	in-phase CH _{ip} duo rock	
1320	0.528	1311	138.3	1354	95.2	short-axis ipCC stretch	
	_	_	_	1360	10.2	_	
	_	_	_	1389	6.3	_	
1381	0.276	1391	23.2	1428	11.1	CH _{ip} scissor, short-axis CC _{ip} stretch	
	_	1411	28.1	1437	19.3	CH _{ip} scissor, long-axis CC _{ip} stretch	
	_	_	_	1445	20.3	_	
1515	0.341	1517	171.2	1550	108.2	CH _{ip} scissor, short-axis CC _{ip} stretch	
	-	1523	24.9	1577	57.6	- **	

except for the two experimental features at 990 and 1100 cm⁻¹. Here, the scaled harmonic positions at 953/968 cm⁻¹ and 1077 cm⁻¹ deviate 30–40 cm⁻¹ from experiment (see Table 3). These two bands are predicted better in the anharmonic calculation, which only deviate by 8 and 9 cm⁻¹, respectively. The peak positions of the 0-K anharmonic spectrum fall under the broad IRMPD features. This may reflect how the IRMPD spectra are measured at room

temperature and are effectively redshifted from the 0-K band position.

Due to the sparse nature that the experimental and theoretical spectra exhibit in the FIR, unambiguous assignments are possible. Similar to the phenanthrene spectrum, the predicted FIR bands show activity in the entire skeletal frame, for which the exact descriptions are given in Table 3. Above 671 cm^{-1} , the modes become



Figure 5. (a) The experimental IR spectrum of cationic perylene (top, black), accompanied by DFT calculated spectra in the harmonic (middle/red, frequencies scaled by a factor 0.96) and anharmonic (bottom/blue, unscaled) approximations; both calculated spectra are convolved with a 30 cm⁻¹ FWHM Gaussian line shape function. The dotted curves are vertical enlargements of the solid curves. (b) Close-up of 270–410 cm⁻¹ spectral range of panel (a) Note that the values on the y-axis of the two bottom panels have been multiplied by 100 for increased readability, and that the convolution has been decreased to 10 cm⁻¹.

increasingly local, and the experimental features are attributed to several close-lying local modes. The MIR assignments in Table 3 are therefore only given for the highest-intensity mode under the convolved peak.

Arguably, the weakest band is found at 434 cm⁻¹, a band that does not seem to be predicted in either of the DFT spectra. However, a close-up of this wavelength region in Fig. 4(b) shows that the strong, predicted band at 485/500 cm⁻¹ (harmonic/anharmonic) is accompanied by two much weaker satellites. The low-frequency satellite could very well account for the experimental 434 cm⁻¹ band, while the high-frequency satellite could account for the asymmetric lineshape of the experimental 488 cm⁻¹ band. Including anharmonicity does not provide a better match with the experiment. The calculated intensity of the harmonic 451 cm⁻¹ band has an intensity of only 0.5 km mol⁻¹. Our ability to detect this feature again demonstrates the high experimental sensitivity.

3.3 Perylene

For perylene ($C_{20}H_{12}^+$, m/z = 252 amu), the IR spectrum in the 105– 1760 cm⁻¹ range is shown in Fig. 5(a) (black curve, top panel). The spectrum comprises three individual curves, with overlap between 215–300 cm⁻¹ and 620–675 cm⁻¹. In total, we observe sixteen bands (see Table 4) of which twelve can be clearly distinguished. The dotted curve shows a vertical enlargement for the weaker bands observed in the FIR, and clearly reveals high S/N bands at 234, 305, 388, 444, 522, and 578 cm⁻¹. The weakest 388 cm⁻¹ feature has a relative intensity three orders of magnitude lower than that for the highestintensity observed band at 737 cm⁻¹. Bandwidths decrease for lower frequencies, down to 5 cm⁻¹ for the 188 cm⁻¹ band. The 188 cm⁻¹ band has a line width of 5 cm⁻¹. A previously reported spectrum of perylene for the MIR region (Bouwman et al. 2019), agrees well with our spectrum in that range. Band frequencies observed here are slightly redshifted (below 10 cm⁻¹) compared to Bouwman et al. (2019). Additionally, our work reveals two weak bands at 923 and 1059 cm⁻¹.

Scaled harmonic (red curve, middle panel) and anharmonic (blue/bottom) spectra are also shown in Fig. 5(a). The harmonic prediction agrees well with the experimental curve. Most peak positions are reproduced within 10 cm⁻¹, although in the 1000-1550 cm⁻¹ range, deviations of up to 18 cm⁻¹ occur, potentially due to saturation effects. As in pyrene, the peak positions of the 0-K anharmonic spectrum fall under the broad IRMPD features, showing what is likely a temperature-induced redshift. Following the same procedure as for phenanthrene and pyrene, the modes of most bands in the spectrum were assigned using the scaled harmonic spectrum, as listed in Table 4. The sparse FIR spectrum allowed for unambiguous assignment, whereas the more densely populated MIR shows several calculated transitions present under one merged band. In the MIR, the assignment corresponds with the highest-intensity mode(s) under such a peak. Similar to pyrene, the modes in the $600-1600 \text{ cm}^{-1}$ range have an increasing in-plane component for increasing frequency. In the FIR (100-600 cm⁻¹) region, we find skeletal out-of-plane and elongation/compression (E/C) modes.

The harmonic spectrum shows no intensity in the range between $300 \text{ and } 400 \text{ cm}^{-1}$, where two features are observed experimentally at $305 \text{ and } 388 \text{ cm}^{-1}$. Fig. 5(b) displays a deeper close-up of this range. The scaled harmonic curve only shows activity at the very edges. The anharmonic prediction shows minor activity, resulting from combination bands – not exceeding intensities above 0.02 km mol⁻¹. The 305 cm^{-1} band is assigned to two combination bands calculated at 295 cm^{-1} (the $133 \text{ cm}^{-1} \text{ H-C} - \text{C} - \text{H}$ twist/scissor vibration and the 163 cm^{-1} drumhead mode) and at 299 cm^{-1} (two ring twisting modes at $26 \text{ and } 274 \text{ cm}^{-1}$), whereas that at 388 cm^{-1} is assigned to a combination band calculated at 291 cm^{-1} (the 26 cm^{-1} ring twisting mode and a 364 cm^{-1} diagonal E/C mode). The experimental 388 cm^{-1} feature is assigned to a combination band

Table 4.	Perylene cation observe	ed band frequencies (cm-	1) and relative	intensities, to	ogether with cal	lculated frequenci	es (cm^{-1})	and IR
intensities	$(km mol^{-1})$, and description	iptions of the theoretical m	nodes. The list	ed harmonic fi	requencies are s	scaled by a factor	0.96. <i>oop:</i>	out-of-
plane; ip:	in-plane; E/C: elongati	on/compression.						

Experiment Harmonic calculations Anharmonic calcula	ations
Norm. Y Freq. IR int. Freq. IR int. (cm^{-1}) $(km mol^{-1})$ (cm^{-1}) $(km mol^{-1})$	nt. Mode description ol^{-1})
0.495 162 8.4 163 8.0	0 drumhead
0.044 242 1.3 247 1.3	3 short-axis jumping jack
295 0.0	007 combination
0.024 – – 299 0.0	006 combination
0.001 – – 391 0.0	024 combination
0.134 444 1.5 450 0.8	8 long-axis jumping jack
- 447 0.8 462 0.8	8 –
- 509 1.5 525 1.1	1 –
0.104 517 8.2 526 8.2	2 zigzag CH _{oop} wag
0.087 564 7.5 576 6.1	long-axis E/C
1.000 737 108.8 741 51.5	5 trio CH _{oop} wag
- 746 6.5 762 6.1	1 –
- 773 1.2 778 0.6	
- 793 2.1 793 1.5	5 –
0.232 805 52.1 807 110.2	2 armchair CH _{oop} wag
0.035 917 3.6 931 2.5	5 antiphase zigzag CH_{oop} , bay CH_{oop}
- 1013 13.8 1037 9.1	1 short-axis E/C
0.085 1077 1.8 1055 5.9	9 trio CH _{ip} scissor
0.157 1104 20.9 1131 13.2	2 zigzag CH _{ip} rock
- 1135 5.6 1149 3.9	9 –
- 1174 41.0 1205 50.7	7 antiphase armchair CH _{ip} scissor
0.270 1197 27.5 1220 14.5	5 in-phase zigzag CH _{ip} rock
- 1267 128.3 1305 93.9	9 antiphase bay HCCCCH _{ip} scissor
1324 19.8	8 -
0.462 1305 187.4 1347 158.4	4 CC _{ip} stretch
0.189 1365 39.4 1393 8.7	7 armchair CH _{ip} rock
- 1421 8.7 1398 12.5	5 –
1402 18.8	8 –
1523 158.8 1553 42.6	5 zigzag CC _{in} stretch, zigzag HCCCH _{in} scissor
0.341 1531 87.4 1564.6 78.4	4 antiphase armchair CC _{ip} stretch
- 1544 9.5 1565.0 80.4	4 –
1584 55.7	7 –
1603 12.5	5 –

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predicted at 391 cm⁻¹ with an intensity of 0.024 km mol⁻¹, which is a combination of an IR-inactive ring twisting mode at 26 cm⁻¹ and an IR-inactive diagonal E/C mode, at 364 cm⁻¹. Calculation predicts an opposite intensity ratio between the two features observed in experiment. It may very well be that the closeness of the 295 and 299 cm⁻¹ bands leads to cooperative absorption, resulting in one stronger resonance.

4 DISCUSSION

The MIR-to-FIR spectra presented in this work are the first spectra of gas-phase, cationic PAHs to go down to 100 cm^{-1} , with bands below 200 cm^{-1} having bandwidths of $\leq 5 \text{ cm}^{-1}$. These spectra represent an important venture into the FIR, adding to very limited experimental data (Bakker et al. 2010). There is no theoretical limit to how large a PAH can become before it becomes impossible to measure its FIR IRMPD spectrum. Although reduced IR cross sections, laser fluence, and photon energy will inevitably form a practical barrier, this has not been reached yet, and spectra of larger, more astronomically representative PAHs will likely be within the measurable range. As anharmonic effects at the basis of redshifts and line broadening are not well-understood in the FIR, especially narrow experimental bands might be more suitable than calculated bands in

comparison to interstellar features. Beyond this measurable range, noble-gas tagging IRMPD spectroscopy becomes a potent method.

Astronomical FIR spectroscopy has been proposed in various studies as a method for molecule-specific PAH identification in the ISM (Mulas et al. 2006a, b; Boersma et al. 2011b, a; Ricca et al. 2010, 2012). Such studies ended up largely fruitless due to the lack of astronomical sensitivity at the time. With the *Origins Space Telescope* (*OST*) mission on the horizon, we are hopeful that efforts to find FIR signatures will be reinvigorated. With its high-resolution OSS and HERO FIR instruments, the *OST* mission will provide a resolution and sensitivity that are orders of magnitude better than Herschel's PACS and AKARI's FIS instruments were able to offer (Battersby et al. 2018). This mission thus offers a realistic (and exciting) prospect to discern PAH FIR modes from the dust background.

The improvements on the observation that *OST* will bring require an increased accuracy for the theoretical predictions. All FIR calculations have so far been benchmarked one experimental value of neutral naphthalene alone (166.4 cm⁻¹; Pirali et al. 2009) as experimental data in the FIR is lacking. Here, we discuss the comparison with this work (pyrene and perylene) and a previous study by one of the authors of this work (anthracene and coronene; Bakker et al. 2010), following the approach of Ricca et al. (2010, 2012) and Boersma et al. (2011b, a).



Figure 6. Double logarithmic plot of the harmonic DFT calculated (0,1) drumhead modes (circles) and short-axis jumping jack (triangles) for twenty cationic PAHs present in the NASA Ames PAH Database (PAHdb; Bauschlicher et al. 2018), together with experimentally observed band positions of four of these molecules as reported previously and in this work (open symbols) (anthracene and coronene from Bakker et al. (2010), and pyrene and perylene from this study). Linear fits of the predicted bands as function of PAH surface area for the different molecular families are given by the solid lines. A full list of included species is given in the Appendix. Cationic perylene was not present in the PAHdb, and we therefore used our own harmonic calculation, scaled with a factor of 0.96 to match in both the MIR and the FIR.

Fig. 6 shows the calculated FIR frequency trends for cationic PAHs as a function of PAH shape (family) and size (surface area *A*) together with the experimental data. Three PAH families (coronenes, pyrenes, and acenes) are displayed as well as an assorted set of PAHs that are capable of jumping-jack and/or drumhead vibrations (see Appendix for the complete list with further details). Also shown are the fitted trend lines for each predicted vibrational mode and family separately. The calculated values are taken from the NASA Ames PAH Database (PAHdb; Bauschlicher et al. 2018). Bond lengths are based on those of benzene (both C–C \cong 1.39 Å and C–H \cong 1.09 Å), after which we calculated the surface areas including H atoms in Å², for each PAH individually, using the geometries defined in Ricca et al. (2012). For clarity, only the drumhead modes with a single node (the (0,1) modes, circles), and short-axis jumping jack modes (triangles) are plotted.

There is an overall good agreement between experiment (open symbols) and PAHdb prediction (solid symbols). However, there is a systematic redshift of $6-11 \text{ cm}^{-1}$ of the experimental value with respect to the theoretical prediction. Two potential causes are the finite temperatures at which the experiments are conducted and the intrinsic red-shifting of IRMPD bands compared to the calculated fundamental $\nu = 1 \leftarrow \nu = 0$ transitions. This work could possibly contribute to new benchmarking efforts to improve predictions of FIR modes for hot, interstellar molecules.

As reported in Boersma et al. (2010), the drumhead modes (circles) are sensitive to PAH size, whereas the jumping-jack modes (triangles) are both strongly shape and size dependent. Indeed, the jumping-jack trend lines for each family are well separated, differing in offset and slope. Surprisingly, despite the strong shape dependence, the bent, single-row PAHs from the displayed assorted set follow the same jumping-jack trend as the linear acenes. This trend line seems to not necessarily be limited to the acene family. Interestingly, there is also a weak shape dependence of the drumhead modes visible, with the

trend lines of the coronene and pyrene family similar, but separated. Benchmarking theory with accurate experimental data, especially for larger PAHs, can provide the means to fully disentangle shape from size dependence and identify individual PAHs. Given the current demonstration of IRMPD spectroscopy extended toward 100 μ m, we foresee that such data will become available in the coming years (Boersma et al. 2011b).

Again, two mixed-shape PAHs, perylene and ovalene, follow the trend observed, potentially because both they and pyrenes in general have D_{2h} symmetry. Because the drumhead trends for the coronenes and pyrenes show a weak, but significant difference, drumhead modes are slightly, but significantly shape dependent, contrary to what was previously postulated. Triphenylene and benzo[e]pyrene, two other mixed-shape molecules with a drumhead mode, do not coincide with the derived trend lines.

5 SUMMARY AND CONCLUSION

We present gas-phase spectra of cationic PAHs in the MIR to FIR region, up (down) to 100 μ m (100 cm⁻¹). These spectra match our scaled, harmonic spectra very well. We also present anharmonic DFT calculations for cationic PAHs in the FIR, which we used to assign pure combination bands with a predicted intensity down to 0.006 km mol⁻¹. The lowest-frequency bands also match calculations from the PAHdb (Bauschlicher et al. 2018) fairly well, although a small, systematic redshift is found. The bandwidths below 200 cm⁻¹ are ≤ 5 cm⁻¹, making the IRMPD data in this wavelength range suitable for benchmarking efforts. In future studies, narrow FIR bands of astronomically sized PAHs – measured with IRMPD – would make good candidates for identification in astronomical surveys.

In this context, we revisited the hypothesis that FIR bands can be used to search for the IR emissions of single PAH species (Mulas et al. 2006a, b; Boersma et al. 2011b, a; Ricca et al. 2012). We report a larger dependence on molecular shape than was previously found, but emphasize the importance of reliable benchmarking before the deduced trends are used to search for astronomical features. Future experimental efforts should be focused on the measurement of FIR spectra of large, symmetric PAHs, both neutral and cationic. Using up to 20 FELICE macropulses, we were able to measure these FIR spectra. Although an upper limit for the size of PAH that we could measure is hard to determine without insight in the density of states, recent MIR studies using an extra-cavity FELIX setup on C48H18 + and $C_{48}H_{20}$ + give us confidence that at least this lower limit for astronomical PAHs is attainable Zhen et al. (2017, 2018). The future OST will be able to provide the sensitive observational data that is necessary to find these elusive FIR bands (Joblin et al. 2011; Battersby et al. 2018).

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DATA AVAILABILITY

The data underlying this article is available at www.doi.org/10.219 42/uva.17023880. Theoretical data from Fig. 6 is available at https: //www.astrochemistry.org/pahdb/theoretical/3.20/default/view, and several experimental data points are derived from Bakker et al. (2010).

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SUPPORTING INFORMATION

Supplementary data are available at *MNRAS* online.

Table S1: List of PAH cations plotted in fig. 6 in the main paper, with their size in number of rings *R* and their area *A* in $Å^2$.

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