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Multiply charged naphthalene and its $C_{10}H_8$ isomers: bonding, spectroscopy, and implications in AGN environments

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ABSTRACT

Naphthalene ($C_{10}H_8$) is the simplest polycyclic aromatic hydrocarbon (PAH) and an important component in a series of astrochemical reactions involving hydrocarbons. Its molecular charge state affects the stability of its isomeric structures, which is specially relevant in ionized astrophysical environments. We thus perform an extensive computational search for low-energy molecular structures of neutral, singly, and multiply charged naphthalene and its isomers with charge states +q = 0-4 and investigate their geometric properties and bonding situations. We find that isomerization reactions should be frequent for higher charged states and that open chains dominate their low-energy structures. We compute both the scaled-harmonic and anharmonic infrared (IR) spectra of selected low-energy species and provide the calculated scaling factors for the naphthalene neutral, cation, and dication global minima. All simulated spectra reproduce satisfactorily the experimental data and, thus, are adequate for aiding observations. Moreover, the potential presence of these species in the emission spectra of the circumnuclear regions of active galactic nuclei (AGNs), with high energetic X-ray photon fluxes, is explored using the experimental value of the naphthalene photodissociation cross-section, σ_{ph-d} , to determine its half-life, $t_{1/2}$, at a photon energy of 2.5 keV in a set of relevant sources. Finally, we show that the computed IR bands of the triply and quadruply charged species are able to reproduce some features of the selected AGN sources.

Key words: astrochemistry – molecular data – methods: numerical – infrared: galaxies – infrared: general – X-rays: galaxies.

1 INTRODUCTION

The recent detections of highly polar aromatic molecules, namely cyanocyclopentadiene (c-C5H5CN; McCarthy et al. 2021; Kelvin Lee et al. 2021), benzonitrile (c-C₆H₅CN; McGuire et al. 2018), and cyanonaphthalene (CNN; McGuire et al. 2021; McCarthy et al. 2021), in Taurus Molecular Cloud 1 (TMC-1) have been encouraging the search for other aromatic derivatives in interstellar and circumstellar environments. Indeed, in addition to these nitrogen-bearing species, pure hydrocarbon cycles, such as cyclopentadiene (c-C₅H₆; Cernicharo et al. 2021a), indene (c-C₉H₈; Burkhardt et al. 2021b; Cernicharo et al. 2021a), and *ortho*-benzyne (o-C₆H₄; Cernicharo et al. 2021b), were also recently and unambiguously detected in TMC-1. Benzyne and benzonitrile are proxies for benzene (Lee, McGuire & McCarthy 2019; Cooke et al. 2020) and can serve as molecular probes. Analogously, cyanonaphthalene can act as a proxy for naphthalene ($C_{10}H_8$), a small aromatic molecule featuring fused bicyclic six-membered rings and the simplest polycyclic aromatic hydrocarbon (PAH). The search for naphthalene and its derivatives is not restricted to TMC-1 because it is possible to find aromatic

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molecules in different astrophysical environments. A very illustrative example is benzonitrile, which was also detected in Serpens 1A, Serpens 1B, Serpens 2, and the dense core MC27/L1521F (Burkhardt et al. 2021a).

Due to the relevance of naphthalene in fields such as astrochemistry, combustion science, and environmental chemistry, several studies have focused on its spectroscopic properties and formation pathways at different charge states. This vast body of knowledge has provided clear evidence that naphthalene is an important intermediate of PAH growth routes (Alliati et al. 2019). Also, through ionization events, naphthalene can trigger the formation of diacetylene (C_4H_2) and the benzene radical cation (Solano & Mayer 2015), thus demonstrating that the charge state influences the formation of other species, including cyclic molecules. Furthermore, even for highly valenceexcited states of the naphthalene cation, relaxation pathways to the ground state due to vibronic coupling and conical intersections are present (Marciniak et al. 2015; Reitsma et al. 2019). This trend, which is also observed for larger PAHs (Wenzel et al. 2020), corroborates with the hypothesis that multiply charged PAHs (Tielens 2008; Zhen et al. 2017, 2018; Banhatti et al. 2021), together with their nitrogendoped, phosphorus-doped, and protonated counterparts (Mattioda et al. 2008; Ricks, Douberly & Duncan 2009; Álvaro Galué & Díaz Leines 2017; Meyer et al. 2021; Oliveira et al. 2021), are also viable candidates as carriers of the unidentified infrared (UIR) bands.

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In addition to fragmentation processes, the ionization of small cycles, including aromatic rings, can induce molecular isomerization (Jašík, Gerlich & Roithová 2015; Monfredini et al. 2016, 2019; Fantuzzi et al. 2018, 2019; Quitián-Lara et al. 2018; Wolff et al. 2020), which is usually accompanied by a severe loss of molecular integrity. Several examples were demonstrated by screening the potential energy surface (PES) of a given molecular structure for each charge state under consideration (Jašík et al. 2015; Fantuzzi et al. 2018; Cerqueira et al. 2020; Gutsev et al. 2020; Hendrix et al. 2020). For benzene, the basic unit of PAHs, the six-membered ring structure is the global minimum for the neutral and singly charged states, but the dicationic singlet ground state has an unusual pentagonal-pyramidal geometry (Krogh-Jespersen 1991; Jašík, Gerlich & Roithová 2014; Jašík et al. 2015; Fantuzzi, de Sousa & Chaer Nascimento 2017; Pozo-Guerron et al. 2020; Wolff et al. 2020). Additionally, while the dicationic global minimum of toluene (C7H8) and aniline (C6H7N) are characterized by the presence of a six-membered ring (Monfredini et al. 2016; Forgy, Schlimgen & Mazziotti 2018), the dicationic chlorobenzene (C_6H_5Cl) is composed of an unusual structure featuring a cyclopropenyl ring and a terminal C=Cl double bond in a formyl-like CHCl moiety (Fantuzzi et al. 2018). Finally, the cationic and dicationic global minima of biphenyl (C₁₀H₁₂), another important building block of larger PAHs, are composed of acenaphthene-like structures, where a C₂ motif bridges a naphthalene backbone (Quitián-Lara et al. 2020).

Given the prevalence of PAHs at differente charge states in astrophysical environments, and taking into account that the molecular charge state can dramatically influence the relative stability of isomeric structures, herein we investigate geometrical and spectroscopic properties of neutral and multiply charged cations of naphthalene $(C_{10}H_8)$ and its isomers at different +q charge states (q = 0-4). Initially, we perform an extensive search for molecular structures in order to reveal low-energy isomers and geometric patterns of $C_{10}H_8$ species for all charge states. Furthermore, we compute scaled-harmonic and anharmonic infrared (IR) spectra of the most representative species, which are discussed in the context of UIR bands. Finally, our findings are also extrapolated to the chemistry of environments with hard radiation fields, where PAHs with a high degree of ionization have been observed (Witt et al. 2006; Peeters et al. 2012), focusing in circumnuclear regions of dusty active galaxy nuclei (AGNs).

The paper is organized as follows: the methodology of geometry and spectroscopic calculations is shown in Section 2. The results about the multiply charged isomers' stability and their main transitions are described in Section 3. In Section 4, we contextualize our results using mid-IR spectra of active galaxies. Lastly, we sum up our findings in Section 5.

2 COMPUTATIONAL METHODS

All calculations performed in this work are based on density functional theory (DFT). Initially, we conduct an exhaustive exploration of the chemical space of $C_{10}H_8$ species at the DFT level for different charge (+q) states, i.e. neutral (q = 0), cation (q = 1), dication (q = 2), trication (q = 3), and tetracation (q = 4). To speed-up the calculations, we use the resolution-of-the-identity method (Weigend 2006) coupled with the chain-of-spheres (COSX) algorithm (Neese et al. 2009). For each charge state, we perform around 185 geometry optimizations and Hessian evaluations with the PBE0 functional (Adamo & Barone 1999; Ernzerhof & Scuseria 1999) combined with the def2-TZVP basis set (Weigend & Ahlrichs 2005). The initial structures were generated by the open molecule generator

(OMG) software (Peironcely et al. 2012). For all cases the lowest possible multiplicity was chosen (singlet for q = 0, 2, 4 and doublet for q = 1, 3). Further single-point calculations were conducted for selected species at higher multiplicities to compute singlet-triplet and doublet-quartet vertical energy gaps. All reported geometries are characterized as minimum energy structures at their corresponding potential energy surfaces by the vibrational frequency analysis as no imaginary frequency was found. All energy values discussed herein are enthalpies computed at the standard temperature of 298.15 K $(H^{298.15})$ and are reported as relative values with respect to the global minimum of each charge state q. Similar stability trends are observed for calculations considering only the electronic energy including zero-point energy corrections (see Table S1 in the Supporting Information). The isomers are labelled in the ascending order of $H^{298.15}$, with 1^{q+} being the corresponding global minimum of the charge state q.

In order to analyse the energetic trends of selected isomers as a function of the charge state q, we use the deviation plot (D-plot) analysis (Fernandez-Lima et al. 2006, 2007; Fantuzzi et al. 2013). This analysis consists of grouping species into distinct classes of compounds and calculating their deviation energies $D_{n,q}^{298,15}$ from the average energy ($\bar{H}_{a}^{298,15}$) for each distinct value of q:

$$D_{n,q}^{298.15} = H_{n,q}^{298.15} - \bar{H}_q^{298.15}, \tag{1}$$

where

$$\bar{H}_q^{298.15} = \sum_{n=1}^N \frac{H_{n,q}^{298.15}}{N}.$$
(2)

The electronic structure and bonding situation of selected species is analysed by inspection of the frontier Kohn–Sham molecular orbitals and additional calculations based on the intrinsic bond orbital (IBO; Knizia 2013) and natural bond orbital (NBO; Weinhold, Landis & Glendening 2016) methods. In order to investigate the aromaticity (or antiaromaticity) properties of the species, nucleus independent chemical shift (NICS; Schleyer et al. 1996; Chen et al. 2005) calculations are also performed.

Scaled-harmonic and anharmonic IR spectra of all structures with H^{298} values in-between 0–10 kcal mol⁻¹ from the global minima are simulated from further optimizations and frequency calculations at the B3LYP/def2-TZVPP level of theory (Vosko, Wilk & Nusair 1980; Lee, Yang & Parr 1988; Becke 1993; Stephens et al. 1994; Weigend & Ahlrichs 2005). The B3LYP functional was chosen due to its better performance at predicting the frequencies and intensities of the molecular IR bands in comparison to PBE0. The anharmonic spectra are calculated with the vibrational second-order perturbation (VPT2) method (Nielsen 1951; Barone, Biczysko & Bloino 2014). The NASA Ames PAH IR Spectroscopic Database (PAHdb; Bauschlicher et al. 2018) contains the largest library of both laboratory-measured and theoretically computed PAH spectra. Following its guidelines, we apply three scaling factors to the harmonic spectra, with ranges of $\nu > 2500 \text{ cm}^{-1}$ (1), $2500 > \nu > 1111.\overline{1} \text{ cm}^{-1}$ (2), and $\nu < 1111.\overline{1}$ cm^{-1} (3). The scaling factors were determined by minimizing the following expression:

$$\sum_{i=1}^{n} (s(i) * \omega_e(i) - \nu(i))^6,$$
(3)

where ω_e is the harmonic frequency, ν is the experimental fundamental counterpart, s(i) is the scaling factor, and n is the number of IR bands for which there are experimental data. The simulated spectra are convoluted with a Gaussian lineshape function with a full

(a) Most stable $C_{10}H_8^{q+}$ species (q = 0-4)



Figure 1. (a) Low-lying energy $C_{10}H_8^{q+}$ isomers at neutral, cation, dication, trication, and tetracation charge states obtained at the PBE0/def2-TZVP level of theory. Relative enthalpies (kcal mol⁻¹) are shown in parenthesis. (b) IBOs, relative energies, and NBO analysis of the neutral methyleneindene species **2** and **4**. (c) Structure and energetics of $C_{10}H_8^{q+}$ species (q = 2-4) featuring a pentagonal-pyramidal structure.

width at half-maximum of 15 cm^{-1} , in accordance with Boersma et al. (2014).

All geometry optimizations were conducted using ORCA 4.2 (Neese 2012). The IBO calculations were done with the IBOVIEW software (Knizia 2013). Scaled-harmonic and anharmonic IR spectra, as well as the NICS calculations, were performed using the GAUSSIAN 16, revision C.01 (Frisch et al. 2016).

3 RESULTS AND DISCUSSION

3.1 Bonding and relative stability

The seven most stable isomers of $C_{10}H_8$ for each charge state q = 0-4 at the PBE0/def2-TZVP level are shown in Fig. 1(a). Other

structures can be found in the Supporting Information. From q = 0 to q = +3, the vast majority of structures is composed of fused rings. Indeed, the global minima for q = 0 - 2 (1, 1⁺, and 1²⁺) are all naphthalenic structures, with fused six-membered rings. The situation changes dramatically for higher charge states. In the case of q = 3, four isomeric structures have enthalpies within 3.3 kcal mol⁻¹ from the computed global minimum 1³⁺, which contains two terminal C₃H₂ rings attached to an open-chain C₄H₄ motif. This structure is also attributed to the global minimum of the tetracationic C₁₀H₈ system, with the most stable structure containing fused rings being 46.0 kcal mol⁻¹ higher in energy than 1⁴⁺ (*vide infra* for more details).

The second most stable neutral isomer with $C_{10}H_8$ stoichiometry (2) is the 1-methyleneindene species, which lies 23.4 kcal mol⁻¹

above 1. This structure is characterized by fused six- and fivemembered rings, with a methylene (CH₂) group attached to the α -carbon of the five-membered ring. This structure is 21.4 kcal mol^{-1} more stable than the 2-methyleneindene isomer, where the CH₂ group is attached to the β -carbon atom of the five-membered ring. The higher stabilization of the 1-methyleneindene isomer 2 can be attributed to the formation of a Clar's aromatic sextet, as shown in Fig. 1(b). Indeed, calculations based on the NBO approach reveal that the difference between the Lewis energy (E_L) of the two isomers favours the neutral 2-methyleneindene 4 with respect to 2, as $E_I(2)$ – $E_L(4) = +22.3 \text{ kcal mol}^{-1}$. The E_L term considers only the Lewis-type NBOs and neglects electronic delocalization. The correct stability trend between 2 and 4 is only obtained if non-Lewis, delocalized contributions (E_{NL}) are also taken into account. In this case, $E_{NL}(2)$ – $E_{NL}(4) = -44.1 \text{ kcal mol}^{-1}$. The sum of the E_L and E_{NL} contributions leads to the full energy difference (E_{full}) between the two isomers, which is $E_{\text{full}}(2) - E_{\text{full}}(4) = -21.8 \text{ kcal mol}^{-1}$. Following ionization processes, one could expect that these effects would contribute less to the stability of the systems, decreasing the energy difference between the methyleneindene isomers and eventually inverting the energetic trend. In accordance to such expectations, the singly charged 1methyleneindene species 3^+ is merely 0.9 kcal mol⁻¹ more stable than the cationic 2-methyleneindene system 4^+ , while the latter structure is more stable than the former for charge states q = 2-4. The dicationic, tricationic, and tetracationic 2-methyleneindene isomers 2^{2+} , 2^{3+} , and 7^{4+} are, respectively, 12.9, 9.1, and 6.9 kcal mol^{-1} more stable than the doubly (5^{2+}), triply (5^{3+}), and quadruply charged (9⁴⁺, see the Supporting Information) 1-methyleneindene species.

Another important C_{10} backbone is that of the azulene species (3), the most stable neutral $C_{10}H_8$ system with a seven-membered ring. This structure lies 34.3 kcal mol⁻¹ above 1 and is composed of fused seven- and five-membered rings. The energy difference between the naphthalene skeleton and that of azulene decreases as the systems become more ionized. For q = 1, the singly charged azulene species (2⁺) is 18.5 kcal mol⁻¹ higher than 1⁺, an energy drop of more than 50 per cent. This value decreases to 9.2 and 6.1 kcal mol⁻¹ for q =2 and q = 3, respectively. For q = 4, both structures are more than 50 kcal mol⁻¹ above the tetracationic global minimum 1⁴⁺, with the azulene-like structure lying above the naphthalene backbone by around 6 kcal mol⁻¹.

During our investigation, we became interested if structures bearing a pentagonal-pyramidal geometry, such as that of the global minimum of the benene dication, could also be found amongst low-lying $C_{10}H_8$ isomers. All obtained structures are high-energy isomers (see Fig. 1c), indicating that, when the benzene dication global minimum motif is present, the energy always increases.

Amongst the most stable $C_{10}H_8$ structures, we also found species featuring four-membered rings. Some examples are the neutral dimethylenebenzocyclobutene isomer **6**, which lies 56.0 kcal mol⁻¹ above **1**, and the tetracationic structure **5**⁴⁺, which is composed of non-planar four- and six-membered rings connected by a carbon– carbon bond of 1.461 Å. Molecular structures featuring the aromatic, three-membered cyclopropenyl moiety are significantly more common, particularly for dicationic, tricationic, and tetracationic structures. For q = 2, the most stable structure featuring the cyclopropenyl ring is **4**²⁺, which lies 13.9 kcal mol⁻¹ above **1**²⁺ and presents such motif connected to a planar C_7H_6 seven-membered ring with overall C_{2v} symmetry. For q = 3 and q = 4, cyclopropenyl rings are found for species **1**³⁺, **3**³⁺ (2.7 kcal mol⁻¹ above **1**³⁺), **1**⁴⁺, **2**⁴⁺, **3**⁴⁺, and **4**⁴⁺ (3.1, 25.0, and 31.0 kcal mol⁻¹ above **1**⁴⁺, respectively).



Figure 2. Selected optimized dicationic structures featuring large rings. Relative enthalpies (kcal mol^{-1}) are shown in parenthesis.

We note that singlet structures featuring eight- to 10-membered rings are not found amongst the low-energy isomers. These include the monocyclic dicationic structures proposed by Leach and coworkers (Leach, Eland & Price 1989a,b) as intermediate species for twobody charge separation of doubly charged naphthalene and azulene. Selected optimized dicationic structures featuring large rings and their corresponding energies are shown in Fig. 2. These structures lie from 50 to 80 kcal mol⁻¹ above 1^{2+} . For other charge states, similar structures are also found, but they have even higher energies. Isomer 14^{2+} is characterized by a quasi-planar structure featuring fused eight- and four-membered rings. Another eight-membered ring structure characterized herein is 19^{2+} , where two cyclopropene rings are attached to a planar C_8H_4 motif forming a C_{2h} -symmetry geometry. In turn, 15^{2+} is composed of an asymmetrical ninemembered ring fused to a cyclopropenyl ring. The most stable 10membered ring amongst those found in our investigation is 16^{2+} , where the two hydrogen-free carbon atoms are connected through a formal triple bond of 1.240 Å. This structure lies 77.0 kcal mol⁻¹ above 1^{2+} and is nearly degenerate to 13^{2+} , where the hydrogenfree carbon atoms are in para-position. Small energy differences are also observed between these structures and 18^{2+} , which contains the hydrogen-free carbon atoms in meta-position. These results indicate that the dicationic 10-membered ring species with C₁₀H₈ stoichiometry are very fluxional. Despite several attempts, no singlet 10-membered ring formally produced from disruption of the central carbon-carbon bond of the naphthalene dication 1^{2+} has been found. Its triplet counterpart $({}^{3}20^{2}+)$, however, is indeed a minimum energy structure, lying 105.7 kcal mol⁻¹ above 1^{2+} .

After discussing structural properties of the global minima, lowand high-lying energy structures of neutral, singly, and multiply charged $C_{10}H_8$ isomers, we now focus on particular aspects of their energies and bonding situation. Fig. 3(a) shows the structures distribution plot (SDP; Sánchez et al. 2016) of the species investigated herein in the energy window of 0–2.0 eV from the corresponding $C_{10}H_8^{q+}$ (q = 0–4) global minima (1 eV = 23.06 kcal mol⁻¹). These energies are significantly smaller than those of the first ($S_0 \rightarrow S_1$) electronic transition (4.0 eV) and the ionization potential (8.1 eV) of



Figure 3. (a) Structures distribution plot (SDP) of $C_{10}H_8^{q+}$ isomers (q = 0-4) in the energy window of 0-2.0 eV from the corresponding global minima. (b) D-plot analysis of selected $C_{10}H_8^{q+}$ isomers (q = 0-4). (c) Canonical Kohn–Sham molecular orbitals of $C_{10}H_8^{q+}$ isomers (q = 0-4) featuring naphthalenic structures. Their corresponding HL-gap, ST-gap, NICS(0), and NICS_{zz}(0) are also shown. (d) Bonding situation of the tetracationic 1^{4+} species from the IBO perspective.

naphthalene (Cockett et al. 1993). In the SDP plot, each vertical line represents the relative energy position of a stable isomer, with the energies referenced to the global minimum energy structure of each charge state. Amongst the distinct charge states, the population of isomers within the 2.0 eV energy range is the smallest for q = 0. The neutral charge state also shows the largest energy difference between the global minimum (1) and its first low-lying energy isomer (2). These features highlight the strong stabilizing effects at play in the neutral isomers featuring aromatic rings, which are particularly large for naphthalene.

The population of isomers within the 2.0 eV energy window increases for ionized states, reaching its maximum for q = 3. Isomerization reactions at the triply charged potential energy surface should therefore occur more easily, and the minimum energy structures are expected to have more fluxional character.

Increasing the charge state to q = 4 leads to the disappearance of most of the lines and only one low-lying isomer (2^{4+}) is found within the 1.0 eV energy range from the tetracationic global minimum 1^{4+} . Inspection of Fig. 3(b) evidences that the structures featuring fused rings with five, six, and seven members, which accounts for the global minimum and low-lying isomers for q = 0-3, are particularly

unstable for q = 4 as a consequence of charge repulsion. In fact, even the tetracationic naphthalene 8^{4+} (see Fig. 3c) being a Hückel aromatic species, as evidenced by its larger HOMO-LUMO (HLgap, 2.91 eV) and singlet-triplet (ST-gap, 26.2 kcal mol⁻¹) gaps in comparison to those of the antiaromatic 1^{2+} (2.38 eV and 16.8 kcal mol⁻¹, respectively), as well as by its negative NICS(0) (-5.7 ppm) and NICS_{zz}(0) (-11.8 ppm) values, it lies energetically above 1^{4+} by more than 50 kcal mol⁻¹. On the other hand, structures featuring aromatic cyclopropenyl rings and open-chain CH motifs, such as those of 1^{4+} (see Fig. 3d for the corresponding IBOs) and 2^{4+} , are able to distribute the charge more effectively through the carbon backbone, which explains their particular relative stability for higher values of q.

3.2 Infrared spectra

In Fig. 4(a), we present the scaled-harmonic, anharmonic, and experimental spectra of neutral naphthalene (1), the latter taken from the NIST data base (Stein 2018). The calculated peak positions and relative intensities are listed in Table 1. The scaling factors derived for this species are 0.956 for the first group (i.e. $\nu > 2500 \text{ cm}^{-1}$),



Figure 4. (a) Comparison of the scaled-harmonic (blue) and anharmonic (red) IR spectra of neutral naphthalene (1). (b), (c), and (d) are the same as (a), but for (1^+) , (1^2^+) , and (2^{2^+}) , respectively. The experimental spectrum of 1 from NIST is also shown with the black dashed line in panel (a). It was obtained in the gas phase at 300 K.

0.978 for the second group (i.e. $2500 > v > 1111.\bar{1} \text{ cm}^{-1}$), and 0.973 for the third group (i.e. $v < 1111.\bar{1} \text{ cm}^{-1}$). Overall, the fundamental bands of the harmonic spectrum after scaling are fairly close to the experiment, yielding average errors of only 5.8 cm⁻¹ with a maximum value of 18 cm⁻¹ associated with one of the C–H stretching bands. The harmonic spectrum before scaling shows an error trend that leads to more blueshifted lines as a function of their frequency. However, in relative terms, the blueshift is higher for the bands in the first group, second highest for the bands in the third group, and the least intense for the bands in the second group. This discrepancy pattern is satisfactorily mitigated by the scaling factors. None the less, relatively higher errors associated with the C–H stretching bands persist. Considering only the frequency range of groups 2 and 3, the average and maximum errors are reduced to 2.3 and 4.3 cm⁻¹, respectively.

Likewise, the relative intensities of the fundamental bands are also in good agreement with laboratory measurements. The most intense vibrational mode in the scaled spectrum is the C–H wagging mode at ~782 cm⁻¹ (12.8 μ m), followed by the C–H stretching modes at ~3055 cm⁻¹ (3.3 μ m). The third most intense fundamental band corresponds to a C–H bending mode, at ~476 cm⁻¹ (21.0 μ m). In order to quantify the agreement between relative intensities, we normalized the theoretical and experimental intensities to their strongest band. The harmonic spectra yielded an excellent average relative intensity ratio of I_{harmonic}/I_{experimental} ~ 0.96. The relative intensities of the fundamental bands in the anharmonic spectrum are also in very good agreement with the experiment, with $I_{anharmonic}/I_{experimental} \sim 0.92$. Its frequencies, however, are not as accurate as the scaled-harmonic counterpart. Considering the whole spectrum, the average and maximum errors were of 9.96 and 45.9 cm⁻¹, respectively, which are only marginally adequate to aid astronomical observations. The fundamental modes with b_{3u} symmetry were the least accurately predicted by the anharmonic spectrum, with an average absolute error of ~ 24 cm⁻¹ – mainly due to the out-of-plane bending mode at ~ 1003 cm⁻¹ (10.0 µm), which disagrees with the experimental frequency by almost 46 cm⁻¹. Comparatively, the anharmonic b_{1u} and b_{2u} modes are significantly closer to the experiments, with an average error of ~ 6 cm⁻¹.

The scaled-harmonic and anharmonic spectra of the naphthalene cation (1^+) are presented in Fig. 4(b). In Figs 4(c) and (d), we show the spectra of the two least energetic structures of the naphthalene dication $(1^{2+} \text{ and } 2^{2+})$. Given the lack of experimental C–H stretching bands for the ions, we were only able to derive their scale factors for groups 2 and 3. For the naphthalene cation, the calculated scaling factors are 0.985 (2) and 0.973 (3), whereas for the dication they are both 0.976. The harmonic bands pertaining to the first group were scaled using the scaling factor derived for the neutral species (i.e. 0.956).

The calculated peak positions and relative intensities of the naphthalene cation (1^+) are listed in Table 1. Differently from the neutral species, the IR spectra of the cation is absent of intense C-H stretching bands, in accordance with other experimental and

Table 1. Calculated band positions v_{vib} (cm⁻¹) and relative intensities I_{rel} of the fundamental modes of naphthalene (1), (1⁺) and (1²⁺). We compare the calculations to the experimental measurements and give the respective discrepancies δv_{vib} (cm⁻¹).

Experimental	Harmonic calculations			Anharmonic calculations			Assignment / symmetry	
-	$v_{\rm vib}$	$\delta v_{\rm vib}$	I _{rel}	$v_{\rm vib}$	$\delta v_{\rm vib}$	I _{rel}		
				1 (neutral, $q =$	0)			
474 ^a	476.543	2.543	0.199	488.462	14.462	0.188	$H-C-C$ o-o-p bend i-ph / b_{3u}	
620 ^a	620.123	0.123	0.029	631.343	11.343	0.030	C-C str. + H-C-C i-p bend / b_{2u}	
782 ^{<i>a</i>}	782.321	0.321	1.000	793.512	11.512	1.000	H-C-C o-o-p bend i-ph / b_{3u}	
957 ^b	958.852	1.852	0.028	1002.883	45.883	0.059	H-C-C o-o-p bend o-o-ph / b_{3u}	
1011 ^b	1009.115	1.885	0.070	1019.943	8.943	0.066	C-C str. + H-C-C i-p bend / b_{2u}	
1129 ^b	1128.541	0.459	0.041	1138.311	9.311	0.028	H–C–C i-p bend / b_{1u}	
1267 ^b	1262.735	4.265	0.057	1265.503	1.497	0.057	H–C–C i-p bend / b_{1u}	
1389 ^b	1393.221	4.221	0.038	1398.787	9.787	0.032	$H-C-C$ i-p bend / b_{1u}	
1514 ^b	1517.102	3.102	0.076	1515.407	1.407	0.069	C-C str. + H-C-C i-p bend / b_{2u}	
1603 ^b	1605.984	2.984	0.032	1605.447	2.447	0.024	C-C str. + H-C-C i-p bend / b_{1u}	
3006 ^b	3023.277	17.277	0.049	3013.342	7.342	0.022	Aromatic C–H str. / b_{1u}	
3032 ^b	3025.488	6.512	0.007	3031.971	0.029	0.057	Aromatic C–H str. / b_{2u}	
3058 ^b	3040.001	17.999	0.467	3053.25	4.750	0.582	Aromatic C–H str. / b_{1u}	
3068 ^b	3050.965	17.035	0.347	3058.197	9.803	0.289	Aromatic C–H str. / b_{2u}	
				1^+ (cation, $q =$	1)			
410	416.754	6.754	0.210	419.938	9.938	0.229	$H-C-C$ o-o-p bend i-ph / b_{3u}	
589	594.893	5.893	0.091	604.196	15.196	0.109	C-C str. + H-C-C i-p bend / b_{2u}	
759	763.377	4.377	1.000	763.826	4.826	1.000	H-C-C o-o-p bend i-ph / b_{3u}	
1019	1012.563	6.437	0.135	1027.797	8.797	0.179	C-C str. + H-C-C i-p bend / b_{2u}	
1121	1109.893	11.107	0.048	1109.852	11.148	0.047	H-C-C i-p bend / b_{1u}	
1168	1176.030	8.030	0.152	1178.361	10.361	0.183	H-C-C i-p bend / b_{2u}	
1215	1220.391	5.391	2.228	1215.891	0.891	2.181	C-C str. + H-C-C i-p bend / b_{2u}	
1284	1289.241	5.241	0.096	1286.409	2.409	0.101	$H-C-C$ i-p bend / b_{1u}	
1523	1530.011	7.011	0.869	1524.492	1.492	0.109	C-C str. + H-C-C i-p bend / b_{1u}	
1539	1548.630	9.630	0.271	1535.244	3.756	0.170	$C-C$ str. + $H-C-C$ i-p bend / b_{2u}	
				1^{2+} (dication, q	= 2)			
726	740.135	14.135	0.348	731.427	5.427	0.938	$H-C-C$ o-o-p bend i-ph / b_{3u}	
992	979.396	12.604	0.132	988.371	3.629	0.485	$C-C$ str. + $H-C-C$ i-p bend / b_{2u}	
1060	1053.848	6.152	0.256	1062.148	2.148	0.673	H-C-C i-p bend / b_{1u}	
1184	1183.872	0.128	0.075	1192.328	8.328	0.462	H-C-C i-p bend / b_{2u}	
1229	1227.573	1.427	0.899	1234.300	5.300	2.200	H-C-C i-p bend / b_{2u}	
1292	1287.015	4.985	0.037	1291.553	0.447	0.082	H–C–C i-p bend / b_{1u}	
1409	1407.175	1.825	0.391	1417.240	8.240	0.784	H–C–C i-p bend / b_{1u}	
1453	1451.843	1.157	0.509	1449.829	3.171	1.514	C-C str. + H-C-C i-p bend / b_{1u}	
1461	1463.351	2.351	1.000	1461.650	0.650	1.000	C-C str. + H-C-C i-p bend / b_{2u}	
1567	1571.056	4.056	0.195	1569.174	2.174	0.217	C-C str. + H-C-C i-p bend / b_{2u}	

Note. str. = stretch; o-o-p = out of plane; i-p = in plane; i-ph = in phase; o-o-ph = out of phase.

^a(Mackie et al. 2015).

^b(Chakraborty, Banik & Das 2016).

theoretical works (Pauzat et al. 1992; Szczepanski et al. 1992; Hudgins, Sandford & Allamandola 1994; Langhoff 1996; Piest, von Helden & Meijer 1999). In the scaled harmonic approximation, its most intense bands correspond to, in decreasing order, a skeletal bending mode at ${\sim}1220~\text{cm}^{-1}$ (8.2 $\mu\text{m}),$ a C–H bending mode at \sim 763 cm⁻¹ (13.1 µm), and a convolution of two skeletal bendings at ~1540 cm⁻¹ (6.5 μ m). This spectral profile agrees well with jet-cooled and matrix-isolation experiments (Piest et al. 1999). The scaled frequencies were also considerably close to the experimental spectrum, yielding an average absolute error of $\sim 7 \text{ cm}^{-1}$ with a maximum of $\sim 11 \text{ cm}^{-1}$. Comparatively, the anharmonic frequency predictions of the fundamental bands were only slightly worse than the scaling approach, yielding average and maximum discrepancies of 6.88 cm⁻¹ and 15.2 cm⁻¹, respectively. All three IR-active symmetries $(b_{1u}, b_{2u}, and b_{3u})$ yield similar errors, i.e. all symmetries are predicted by the anharmonic calculations to a comparable level of accuracy. The resulting spectral profile was very similar to the harmonic approximation, with the two most intense bands at ${\sim}1215~{\rm cm}^{-1}~(8.2~{\mu}{\rm m})$ and ${\sim}763~{\rm cm}^{-1}~(13.1~{\mu}{\rm m})$ corresponding to the aforementioned skeletal distortion and C–H bending modes. Moreover, this spectrum also shows an intense band at ${\sim}1500~{\rm cm}^{-1}~(6.7~{\mu}{\rm m})$. However, in contrast with the strong fundamental band that gives rise to this peak in the harmonic approximation, in the anharmonic spectrum this band is mainly composed of two partially resolved combination bands of relatively high intensity – characterized by the $\nu_{37} + \nu_{35}$ and $\nu_{39} + \nu_{32}$ modes. Moreover, both $C_{10}H_8^+$ spectra are overall significantly more intense than those of the neutral species, due to the latter's lack of a permanent electric dipole moment.

The dicationic naphthalene (1^{2+}) IR spectra have a slightly more complicated profile than the singly charged counterparts. Its calculated peak positions and relative intensities are listed in Table 1. Both the scaled-harmonic and the anharmonic spectra show an intense double band at ~1500 cm⁻¹ (6.7 µm) due to out-of-



Figure 5. Scaled-harmonic (blue) and anharmonic (red) spectra of all the remaining calculated naphthalene ions.

phase H–C–C bending and C–C stretching modes, as well as another intense band at ~1250 cm⁻¹ (8.0 μ m) related to in-phase H–C–C bending. They also show moderately intense bands at ~730 cm⁻¹ (13.7 μ m), ~1400 cm⁻¹ (7.1 μ m), and ~3100 cm⁻¹ (3.2 μ m), respectively related to wagging, rocking, and C–H stretching modes. Comparatively, the naphthalene cation rocking bands are significantly less intense. Moreover, combination bands of moderate intensity in the ~1400–1600 cm⁻¹ (6.3–7.1 μ m) range considerably change the profile of the spectrum in comparison to the harmonic approximation. In fact, anharmonic corrections seem to play an important role in the IR spectra of doubly charged naphthalene (1²⁺) (Banhatti et al. 2021): the average and maximum frequency errors of the anharmonic spectra were, respectively, 3.9 and 8.3 cm⁻¹ – ~20 per cent and ~41 per cent less than the ones of the scaled-harmonic spectrum. These errors are consistent throughout the entire frequency range of groups 2 and 3 (2500 cm⁻¹ > ν), and are independent of the mode symmetry. Both scaled-harmonic and anharmonic spectra are reasonably to significantly close to the experimental spectrum of Banhatti et al. (2021) at the measured frequency range.

The scaled-harmonic and anharmonic spectra of the remaining calculated structures with $H^{298} < 10$ kcal mol⁻¹ are shown in Fig. 5. Due to the lack of experimental data, the harmonic bands of all ionic species with charges above q = +2 were scaled using the factors derived for the naphthalene dication 1^{2+} . The profiles of the spectra change appreciably with the charge state, even for

similar naphthalene-like geometries. Different structures of the same charge states also result in very distinct spectra. Combination bands of intermediate to high intensity were derived in the majority of the anharmonic spectra, but the resulting spectral profiles remained

the intensity of fundamental bands that fall in their vicinity. As observed for the dication global minimum (1^{2+}) , the IR spectra of the remaining multiply charged species are considerably more complex than those of the neutral and singly charged species, which hampers the characterization of potential fingerprint regions. With the exception of the naphthalene cation, all charged species show C-H stretching features of moderate to high strength. In fact, for most of the trication structures and all of the tetracation structures, these are the dominant bands of the spectra. The tetracationic, secondlowest-energy $C_{10}H_8$ structure 2^{4+} is unique in that it has an aliphatic extremity that results in C-H stretching bands with frequencies ranging from 2800 to 3000 cm^{-1} (3.3–3.6 μm) additionally to the higher energy features typical of unsaturated species. The 3^{3+} (thirdlowest-energy structure) of the trication species also has aliphatic C-H bonds, but the resulting IR features have higher frequencies at around 3090 cm⁻¹ (3.2 μ m). This shift is likely due to the close proximity to the aromatic ring.

mostly unchanged - or only slightly different - due to decreases in

For the neutral species, the out-of-plane C-H modes that fall in the frequency range of \sim 666–1000 cm⁻¹ (10–15 µm window) are the dominant bands in the spectrum. Comparatively, the in-plane C-H bending and C-C stretching modes that fall in the frequency range of 1000–2000 cm⁻¹ (5–10 µm window) are not as important in terms of intensity. However, this profile changes for the cation and dication species: the latter modes become the dominant ones in the spectra. Based on this well-known phenomenon, the intensity ratios of the bands in the 5-10 and 10-15 µm regions have been widely used as both qualitative and quantitative proxies to track the charge balance of PAH populations in astronomical sources (Draine & Li 2001; Boersma, Bregman & Allamandola 2015, 2016; Maragkoudakis, Peeters & Ricca 2020). Indeed, this trend has been observed for the most part of the cationic species of Figs 4 and 5. In spite of that, the IR spectra of the trication naphthalene-like structure (4^{3+}) (see Fig. 5d) do not reproduce this behaviour. In fact, its out-of-plane C-H modes are more intense than the in-plane C-H bending and C-C stretching modes, which better resembles the spectral profile of the neutral species.

4 ASTROPHYSICAL IMPLICATIONS

In this section, we discuss the present results in the context of the chemistry of circumnuclear regions of AGNs. From the observed X-ray luminosities, we estimate the half-lives of naphthalene in the circumnuclear regions of the selected AGN sources, whose results are shown in Fig. 6(a). As discussed by Monfredini et al. (2019), multiply charged naphthalene molecules are indeed present in the mass spectrum of the system after interaction with 2.5 keV photons, revealing that such species could contribute to the emission spectrum of objects featuring high energetic fluxes, such as AGNs. The mid-IR emission spectra of the selected AGN sources, namely NGC 3227, NGC 7469 and NGC 7582, are shown in Fig. 6(b). They were compiled by the ATLAS project (Hernán-Caballero & Hatziminaoglou 2011) and are also available through IRSA (NASA/IPAC Infrared Science Archive; https://irsa.ipac.caltech.edu/). The main spectral features are compared with the anharmonic vibrational transitions of the most representative $C_{10}H_8^{q+}$ species calculated herein, whose computed data in the 6-15 µm range is shown in Fig. 6(c).

UIR bands – and therefore their carriers, which include PAHs – are widely observed in the Universe, even in highly dissociative regions such as the circumnuclear regions of AGNs (e.g. Alonso-Herrero et al. 2020). To explain the emission of PAH around AGNs, Voit (1992) proposed that these molecules should be protected by a dense torus surrounding the highly ionizing central source. Knowing that the gas entering a central AGN can form a circumnuclear disc, one could expect that this structure would play an important role in obscuring the AGN (Netzer 2015, and references therein). There is in fact evidence of PAH detection in the proximities of type-1 Seyfert and intermediate-type Seyfert galaxies (Sales, Pastoriza & Riffel 2010; Jensen et al. 2017), suggesting that PAHs can on the one hand survive in these regions and, on the other, that PAH excitation is not necessarily limited to star-forming regions.

ISM dust grains follow a size distribution in the range of 0.4 nm to larger than 10 nm (Tielens 2008), with PAHs at the smallest end being characterized in terms of the number of carbon atoms contained (Galliano et al. 2008). Despite occupying the small end of dust size distribution, PAHs are large molecules and their vibrational modes result in broad emission features in the mid-IR region associated with the different stretching and bending modes of C-H and C-C bonds, mainly at the \sim 3–13 µm region (Allamandola, Tielens & Barker 1989). As shown by e.g. Draine & Li (2001) and Li & Draine (2001), some of these features can be used to trace the structure, size, and charge of the molecules. It has been shown that neutral and singly ionized PAHs are not sufficient to model the features and multiply ionized PAHs must be accounted for in the ISM to provide good matches between laboratory and astrophysical data (Bakes, Tielens & Bauschlicher 2001a). Based on Bakes et al. (2001b), the features of multiply charged PAHs are more pronounced in the C-C modes in comparison to the C-H modes. The possibility of identifying ionized PAHs in the ISM has been proposed and discussed for several decades. In this regard, Leach (1986) showed the importance of studying the photostabilities of PAH monocations and dications and their astrophysical implications. Subsequently, Bakes et al. (2001a) proposed that the probability of finding PAHs in a state of multiple ionization increases with their molecular size. In regions with highenergy fluxes, PAHs therefore might be doubly, triply, or even more highly charged. From the experimental point of view, only some of the simplest ionized species have been extensively studied in the astrophysical context (Szczepanski & Vala 1993; Ruiterkamp et al. 2002; Tielens 2008; Zhen et al. 2015; Banhatti et al. 2021). PAHs with higher charge states are significantly more challenging to study both from the experimental and computational points of view, and their unambiguous detection in astrophysical environments is still unachievable (Zhen et al. 2017).

4.1 Half-lives of naphthalene in PAH-containing AGN sources

In order to compare our spectroscopic data with astrophysical objects, we selected three distinct PAH-containing AGNs, namely NGC 3227, NGC 7469, and NGC 7582, taken from Jensen et al. (2017). These AGNs are classified as intermediate Seyfert-type galaxies, with PAH bands identified in mid-IR and predominance of small, neutral, and pure PAH species (Martins-Franco & Menéndez-Delmestre 2021). The X-ray luminosities in the range of 2–10 keV of these selected AGNs are shown in Table 2.

NGC 3227 is a spiral-type galaxy interacting with a neighbouring dwarf elliptical galaxy (Delgado & Perez 1997). This object presents a large concentration of molecular gas in the central part, equivalent to 80 per cent of the total CO emission that is detected in a radius of \sim 1.2 kpc (Delgado & Perez 1997). NGC 3227 is located at a



Figure 6. (a) Half-life values of the PAH molecules studied herein after interaction with the X-ray radiation field of selected AGNs. Each data point refers to a specific distance between 10 and 500 pc from the Seyfert nucleus. (b) Mid-IR IRS/Spitzer spectra of the Seyfert galaxies that were analysed in this work, where the shaded areas represent the wavelength regions of the main PAH features. (c) Spectra of singly, doubly, and thirdly charged naphthalene scaled and compared to selected AGNs.

distance of ~22 Mpc, at a luminosity of log $L_X = 42.14 \pm 0.21$ erg s⁻¹ (Jensen et al. 2017). The galaxy also exhibits a short-term hard X-ray variability likely related to variation in the intrinsic continuum emission (Ramos Almeida et al. 2009).

In turn, NGC 7469 is an active core spiral galaxy with a largescale stellar bar (several kpc) identified in the near-IR (Knapen, Shlosman & Peletier 2000). Its circumnuclear region is surrounded by a 1.5–2.5 arcsec starburst ring (Soifer et al. 2003). In the mid-IR, the most central region of NGC 7469 (~1 arcsec of the nucleus) is dominated by emissions in the *K* band (2.2 µm) and the 3.3 µm band (Izumi et al. 2015), the latter indicating a strong presence of PAHs in a region very close to the nucleus. NGC 7469 is at ~67.9 Mpc and has a luminosity of log $L_X = 43.19 \pm 0.07$ erg s⁻¹ (Jensen et al. 2017).

Table 2. Some properties of the AGN sources studied herein. L_X (eV s⁻¹) stands for the X-ray luminosities integrated from 2–10 keV. $N_{\rm H}$ (cm⁻²) is the column density of atomic hydrogen. τ_x and \bar{F}_X (photons cm⁻² s⁻¹) are the estimated X-ray optical depths and the average X-ray photon fluxes at 2.5 keV within distances of 10–500 pc from the Seyfert nucleus of the corresponding sources.

Source	Туре	L_X^a (eV s ⁻¹)	$\frac{N_H^b}{(\mathrm{cm}^{-2})}$	τ_x	$\frac{\bar{F}_X}{(\mathrm{cm}^{-2}\mathrm{s}^{-1})}$
NGC 3227	\$1.5	8.62×10^{53}	1.86×10^{22}	0.52	5.57×10^{8}
NGC 7469 NGC 7582	S1.5 S1i	9.67×10^{54} 2.11×10^{54}	1.15×10^{21} 4.90×10^{23}	0.03 13.64	1.02×10^{10} 2.73×10^{3}

*Notes.^a*Taken from Jensen et al. (2017).

^bTaken from Jaffarian & Gaskell (2020).

Lastly, NGC 7582 is a nearby spiral galaxy, located at a distance of ~23.0 Mpc and a luminosity of log $L_{\rm X} = 42.53 \pm 0.38$ erg s⁻¹ (Jensen et al. 2017). NGC 7582 presents a circumnuclear region of active star formation and an AGN-powered outflow, making it an ideal site for studies focusing on the feeding and feedback processes in which AGN lead to an increase or decrease in star formation (Riffel et al. 2009).

In order to estimate the survival of naphthalene in the proximity of the circumnuclear regions of the aforementioned AGN sources, we used the experimental data taken from Monfredini et al. (2019) at photon energy $E = h\nu = 2.5$ keV to obtain the photodissociation cross-section, σ_{ph-d} (cm²), of the molecule. For that, the average X-ray photon flux, \bar{F}_X (photons cm⁻² s⁻¹) of each source, in a fixed range of distances from the circumnuclear region (10–500 pc), was calculated using the following equation:

$$\bar{F}_X = \frac{L_X}{4\pi r^2 h\nu} e^{-\tau_x},\tag{4}$$

where L_X is the X-ray luminosity (eV s⁻¹) integrated from 2 to 10 keV, *r* is the distance from the nucleus to a position between 10 and 500 pc, and τ_x is the X-ray optical depth given by

$$\tau_x = \sigma_{\rm H}(E) N_{\rm H},\tag{5}$$

where $\sigma_H(E)$ and N_H are the X-ray photoabsorption cross-section and the column density of the hydrogen atom, respectively. The values of $\sigma_H(E)$ can be obtained by (Gorti & Hollenbach 2004)

$$\sigma_{\rm H}(E) = 1.2 \times 10^{-22} \left(\frac{E}{1 \rm keV}\right)^{-2.594}.$$
(6)

Therefore, at a 2.5 keV photon energy, $\sigma_{\rm H} = 1.11 \times 10^{-23} \text{ cm}^2$.

The calculated \bar{F}_X values are shown in Table 2. Now, taking the X-rays photon flux values and using the photodissociation cross-section values from Monfredini et al. (2019), we estimate the half-life of naphthalene using the following equation (for more details, see Monfredini et al. 2019):

$$t_{1/2} = \frac{\ln 2}{k_{ph-d}},$$
(7)

where k_{ph-d} , the photodissociation rate (s⁻¹), is equal to the product of σ_{ph-d} and \bar{F}_X at the same photon energy. The average half-lives of naphthalene for each source vary from 7.4 × 10⁹ yr for NGC 7582 to 2.0 × 10³ yr for NGC 7469 (see Fig. 6a). In the case of NGC 7582, the half-life times obtained show that naphthalene can present comparable times, or even higher, than that necessary for the injection of PAHs in the interstellar medium (ISM; 2.5 × 10⁹ yr; Jones et al. 1994). The conditions of NGC 7582 are optimal for the formation of these molecular species and this is due to the fact that, although its X-ray luminosity of 2.11×10^{54} eV s⁻¹ is considerably large, its high $N_{\rm H}$ value of 4.90×10^{23} cm⁻² makes it the most obscured AGN source amongst those investigated herein. Therefore, we can speculate that the thick dust surrounding the circumnuclear region is an effective blocker, which exerts a high degree of protection for PAHs, including smaller PAHs such as naphthalene.

On the other hand, for the other two AGNs, the half-lives obtained are smaller than the PAH injection time. For NGC 7469, the ratio between the PAH injection time and the average half-life of naphthalene is as large as ~10⁶, even though the X-ray luminosity of this source is higher than that of NGC 7582 by a factor of ~4.6. This is due to the fact that the column density $N_{\rm H}$ in NGC 7469 is particularly thin ($\tau_x = 0.03$, see Table 2), around 450 times smaller than that of NGC 7582 ($\tau_x = 13.64$). In order to account for the presence of PAHs, particularly those of smaller sizes, in such objects that combine highly dissociative fluxes and optically thin column densities, one or more PAH protection mechanisms (see Quitián-Lara et al. 2018; Monfredini et al. 2019) should be at play in order to reduce the effectiveness of dissociation induced by photon–molecule interaction.

4.2 Broad-band emission features in the mid-IR spectra of AGNs

In Fig. 6(c), we display naphthalene ionization states addressed in this work with their respective structures and spectra. The first feature (a), appearing at 6.2 μ m, is associated with the C–C stretching band of PAHs and is common to the main $C_{10}H_8^{q+}$ structures with q =0-4. However, as expected due to their higher dipole moments, the intensity contributions coming from the ionized species are larger than that of the neutral counterparts. The second feature (b) coincides with the 7.7 µm emission band. This feature is broadly considered to be the combination of two emission bands, at 7.6 and 7.8 µm, the relative intensities of which are employed to classify different sources (Canelo et al. 2021). All AGNs investigated herein are characterized by the larger intensity of the 7.6 µm component, thus pertaining to the spectral class A' (Peeters et al. 2002). This component is indeed consistent with PAH emission, which in this wavelength is originated mostly from in-plane C–H bending modes. For the $C_{10}H_8^{q+}$ species, vibrations within this band are particularly intense for q = 3, which presents strong features at 7.7 and 8.6 µm. The latter is also assigned to C-H in-plane modes and is seen in feature (c). The features (d) and (e), 11.3 µm and 12.7 µm, respectively, are associated with C-H out-of-plane bending modes and, as predicted, are virtually unseen for ionized $C_{10}H_8^{q+}$ species, except for q = 4. In opposition to the other ionized $C_{10}H_8^{q+}$ species, the tetracation presents a magnified flux at higher wavelengths – and especially at feature (d). Finally, the C-H out-of-plane mode at 12.7 µm is the most intense vibrational contribution of neutral naphthalene, and coincides with the feature (e) present in all AGN sources investigated herein. Regarding the ionized naphathalenic structures, this band becomes increasingly redshifted as the charge state increases (see also Table 1). Consequently, the peak position of band 12.7 µm could potentially serve as a tracer of the medium's ionization state. Finally, we compare the NGC 7582 mid-IR spectrum with a composed theoretical curve obtained by a linear combination of the computed anharmonic spectra of Fig. 6(c). These results are summarized in Fig. 7. As expected, the 12.7 µm band is mainly reproduced by the neutral naphthalene. Interestingly, we also obtained that the combination of highly ionized $C_{10}H_8$ species is able to reproduce some features of the 6.2 and 7.7 µm bands, with trications contributing mainly to the higher wavelength



Figure 7. Comparison of the NGC 7582 mid-IR spectrum with a theoretical curve obtained by a linear combination of the computed anharmonic spectra of distinct $C_{10}H_8^{q+}$ species. For a better visualization, the computed spectra were scaled.

portion of the latter and tetracations to the lower wavelength portion of the former. These findings hint at the importance of considering not only the charge state of PAHs to the composition of AGN IR bands, but also the distinct low-lying isomeric structures that might be produced following multiple ionization events. However, the small size of our prototypical model is not large enough to provide definite answers. Therefore, further investigations of this phenomenon with larger multiply charged PAH species are warranted.

The mid-IR broad-band features have been explored in the literature as a diagnostic tool for the ISM's radiation field and the ionization state of PAHs (e.g. Lambrides et al. 2019; Martins-Franco & Menéndez-Delmestre 2021). Draine & Li (2001) demonstrate that the 11.3-to-7.7 µm band flux ratios can tells us something about the overall characteristics of the PAH population in the source: the former is associated with the ionization fraction, whereas the latter is a strong function of the PAH size. This correlation, however, is not observed for the $C_{10}H_8^{q+}$ species explored here (see Fig. S6 in the Supporting Information). In fact, we detect no trend for the IR band intensities with respect to the molecule's charge state nor structure. That is rather unsurprising, considering that the model of Draine & Li (2001) was constructed for what the authors called a 'generic PAH' with C > 16 and that is either neutral or singly ionized. Moreover, their model considers environments that are not completely compatible with AGNs, such as the cold neutral and warm ionized ISM, as well as photodissociation regions. In this sense, further studies that focus on the IR spectra of multiply ionized PAHs of different sizes are warranted to derive adequate proxies for higher ionization states.

5 CONCLUSIONS

We investigated the geometric properties and patterns of neutral, singly, and multiply charged naphthalene and its $C_{10}H_8$ isomers by searching extensively for molecular structures with charge states q = 0-4. The most stable $C_{10}H_8$ isomers with q = 0-2 are predominantly composed of fused rings. However, for higher charge states, this profile changes sensibly, with open chains dominating the low-energy structures of the tetracationic $C_{10}H_8$. The effect of electronic delocalization is shown to become gradually less important to the stabilization of the systems as the charge state increases, and the increase in charge state also results in a significantly smaller energy difference between naphthalenic and azulenic backbones. High-energy pentagonal-pyramidal structures fused to six- or fivemembered rings were found for the $C_{10}H_8$ systems with q = 2-4, whereas for the species with lower charge states they were not observed. Additionally, four- and three-membered rings were also repeatedly observed, with the latter being notably common for the tetracationic systems. However, regarding larger rings with 8– 10 members, we do not find singlet structures with those features amongst the low-energy isomers.

The neutral charge state is evidently highly stabilized by the presence of aromatic rings, as is conveyed by the isomer population and their energy difference in the SDP plot. For higher charge states, and especially for q = 3, the population of isomers increases significantly. Thus, isomerization reactions should be more frequent and consequently play a more important role in the characterization of these systems. For q = 4, charge repulsion leads to the destabilization of structures with five-, six-, and seven-membered fused rings, whereas aromatic cyclopropenyl rings and open-chain CH motifs increase the system's stability by allowing to distribute the charge more effectively.

We also assessed the spectroscopic properties of the most stable structures ($H^{298.15} < 10 \text{ kcal mol}^{-1}$) in the IR region, with the goal of aiding in the interpretation of astronomical observations. We simulate both their harmonic and the anharmonic IR spectra, and derive scaling factors in accordance to the PAHdb guidelines based on the experimentally measured bands of the neutral, singly, and doubly charged global minima. Overall, the scaled-harmonic spectra were reasonably close to experimental results (when available), with average errors well within 10 cm⁻¹ from the measured values. Comparatively, the anharmonic spectrum of neutral naphthalene was sensibly less accurate than the scaled-harmonic counterpart. For higher charge states, however, anharmonic corrections become increasingly more important, eventually yielding a more accurate spectrum than the scaled-harmonic approach for the dicationic species. None the less, all spectra obtained here were satisfactorily close to experiments and thus both approaches seem adequate to aid astronomical observations.

The computed spectra of the remaining low-energy $C_{10}H_8$ structures are also provided. Overall, the complexity of the IR emission spectra increases significantly for higher ionization states. Intense bands in the C–H stretching region are consistent for the vast majority of the species studied here, with the only exception of the naphthalene cation (1⁺). Furthermore, the species with higher charge states also typically present spectra dominated by the bands in the 5–10 µm region. The naphthalenic trication (4³⁺), however, does not follow this trend, resembling more closely the profile of the neutral species.

The hard radiation generated by supermassive black holes accreting material may lead to the formation of singly, doubly, and even multiply charged PAHs, including in the case of $C_{10}H_8$. Therefore, the species explored in this work are potentially part of the chemistry of circumnuclear regions of AGNs, and could thus be enriching their emission spectra. For this reason, we explore the survival of naphthalene in the proximity of a set of PAH-containing AGNs, using our previous experimental value of photodissociation crosssection, σ_{ph-d} , and its half-life, $t_{1/2}$, at photon energy of 2.5 keV. For NGC 7582, the naphthalene half-life is comparable to, or even higher than the time necessary for the injection of PAHs in the ISM, which is likely due to the shielding of the PAHs from the high X-ray flux by the dust surrounding the circumnuclear region. For the rest of the selected sources, however, the dust available is not as effective in shielding the PAH molecules; other protection mechanisms, such as PAH superhydrogenation (Reitsma et al. 2014;

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Quitián-Lara et al. 2018; Stockett et al. 2021) and PAH-to-dust adsorption and incorporation processes (Monfredini et al. 2019), are required.

We also compared the computed IR spectra of the most representative low-energy $C_{10}H_8$ structures with that of the selected AGNs. The objects' spectra are consistent with PAH emission, and are especially compatible with the IR bands of the q = 3 and q = 4 species. The C-H out-of-plane modes of the naphthalenic species present increasingly redshifted frequencies as a function of the charge state, pointing in the direction of a promising tracing capability of the 12.7 µm band regarding the ionization state. Finally, we assessed the capability of typical ionization-state proxies to predict the charge of the multiply ionized $C_{10}H_8$, but found no correlation. Thus, further works dedicated to exploring the IR band ratios of PAHs with multiple charge states are desirable.

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DATA AVAILABILITY

Data available on request.

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SUPPORTING INFORMATION

Supplementary data are available at MNRAS online.

- **Table S1.** Comparison between the standard enthalpy at 298.15 K (H298:15) and the electronic energy with zero-point energy correction (E+ZPE) of selected dicationic species.
- Figure S1. Structure and relative stability of selected neutral, $C_{10}H_8$ isomers.
- Figure S2. Structure and relative stability of selected $C_{10}H_8^+$ isomers.
- Figure S3. Structure and relative stability of selected $C_{10}H_8{}^2$ + isomers.

Figure S4. Structure and relative stability of selected neutral, $C_{10}H_8^{3+}$ isomers.

Figure S5. The 11.3 μ m/7.7 μ m intensity ratio of the computed and observed IR bands as a function of their corresponding 6.2 μ m/7.7 μ m intensity ratio.

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