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Subtle hydrogen bond preference and dual Franck–Condon activity – the interesting pairing of 2-naphthol with anisole[†]

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The hydrogen-bonded complexes between 2-naphthol (or β -naphthol) and anisole are explored by detecting their IR absorption in the OH stretching range as well as their UV absorption by means of laser-induced fluorescence and resonance-enhanced two-photon UV ionisation. For the more stable *cis* and the metastable *trans* conformations of the OH group in 2-naphthol, hydrogen bonding to the oxygen atom of anisole is consistently detected in different supersonic jet expansions. Alternative hydrogen bonding to the aromatic ring of anisole remains elusive, although the majority of state-of-the-art hybrid DFT functionals with London dispersion correction and – less surprisingly – MP2 wavefunction theory predict it to be slightly more stable at zero-point level, unless three-body dispersion correction is added to the B3LYP-D3(BJ) approach. This changes at the CCSD(T) level, which forecasts an energy advantage of 1–3 kJ mol^{−1} for the classical hydrogen bond arrangement even after including (DFT) zero-point energy contributions. The UV and IR spectra of the *cis* complex exhibit clear evidence for intensity redistribution of the primary OH stretch oscillator to combination states with the same low-frequency intermolecular bending mode by Franck–Condon-type vertical excitation mechanisms. This rare case of dual (vibronic and vibrational) Franck–Condon activity of a low-frequency mode invites future studies of homologues where aromatic ring docking of the OH group may be further stabilised, e.g. through anisole ring methylation.

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1 Introduction

The spectroscopic study of hydrogen-bonded complexes in the gas phase is a mature field, where different spectral regions and different detection techniques lead to a fruitful synergy and powerful conclusions.^{1–13} When two multifunctional molecules form a weakly bound complex, there are two interesting structural aspects which lend themselves to closer experimental investigation, because they can provide quantum chemical methods with suitable reference points. One is the ground state preference for one over other structural arrangements of the complex in the competition of intermolecular forces. It is addressed by the concept of intermolecular energy balances,^{14–16} where systems are

picked for being ambivalent in terms of two different arrangements of their constituents.^{17–24} The other aspect is the influence of molecular excitation of the components on this structural arrangement of the complex, due to the coupling of intramolecular degrees of freedom with intermolecular ones. This leads to the phenomenon of Franck–Condon effects in electronic and – less frequently observed in neutral complexes – vibrational spectroscopy of molecular aggregates.^{25–30}

In this work, we argue that the complex between 2-naphthol and anisole (Fig. 1) is a particularly interesting and challenging case of an intermolecular energy balance and of a dual Franck–Condon effect. It is a rare case where even rather high-level electronic structure calculations supplemented by (harmonic)

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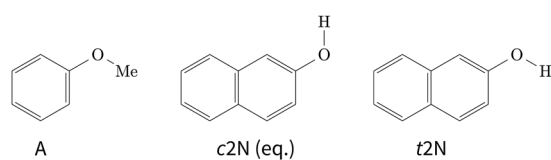


Fig. 1 Schematic Lewis structures of anisole (A) and conformational isomers of 2-naphthol (2N). The *cis* isomer is the global minimum conformation of 2-naphthol.



vibrational zero-point energy correction remain ambiguous about the global minimum structure. Only at the highest level, one can confidently conclude that the detectable structure is the energetically winning structure in a very subtle competition. The 2-naphthol-anisole complex is also a rare case where apparently the same slow intermolecular vibrational mode is demonstrated to be excited together with electronic or together with a suitable high-frequency vibrational excitation. This again reflects the sensitivity of the molecular complex structure to internal excitation, and thus the strongly competing character of different intermolecular forces in determining this structure.

2 Experimental and computational methods

2.1 FTIR jet spectroscopy

To obtain the FTIR jet spectra, two different versions of the heatable “popcorn-jet” setup were used (hereafter referred to as “old”¹⁹ and “new”³¹). In both cases, a stream of helium (Linde, 99.996%) was first seeded with anisole (Fluka, 99%) molecules in a saturator tempered to 0–20 °C (273–293 K) and then led into a 69 L reservoir at a stagnation pressure of 1.5 bar. Afterwards, the gas mixture was directed through a heatable chamber which contained the less volatile 2-naphthol (Merck, 99%), deposited on molecular sieve and heated to 115–120 °C (388–393 K), and was enclosed by two individually heatable poppet valves. The gas mixture was subsequently expanded in pulses through a slit nozzle. Further details on the measurement conditions can be found in the ESI.† In the old setup (spectra (i), (ii), (a), and (b) in Fig. 3), a straight double 0.5 × 10 mm² slit nozzle was used with a pulse length of 316 ms and an opening pressure of the poppet valves of 690 mbar. The new setup (spectra (c) and (d) in Fig. 3) features an individually heatable single 162° angled 0.2 × 60 mm² slit nozzle with a shortened pulse length of 200 ms and a reduced opening pressure of 350 mbar. In both versions of the setup, a 3.6 m³ buffer volume that is permanently evacuated by three consecutive vacuum pumps with a total pumping speed of 500 m³ h⁻¹ ensured a sufficiently low background pressure during expansions. The FTIR spectra (in both versions one double-sided interferogram scan per pulse) were recorded perpendicular to the expansion direction by a Bruker IFS 66 v/S spectrometer containing a ceramic glower as light source, a KBr beamsplitter and CaF₂ optics. A resolution of 2 cm⁻¹ was achieved.

In ref. 31, the main advantages of the new setup and differences to the previous setup have already been discussed in detail. The comparison of spectra recorded with the old (a and b) and new (c and d) setup in Fig. 3 nicely illustrates the overall signal enhancement achieved through the increased absorption path length in combination with the reduced opening pressure of the poppet valves. In essence, the new setup can produce more useful spectra despite the shortened pulse length and the partially lower temperature and thus lower substance concentration.

2.2 UV and IR-UV jet spectroscopy

Anisole (Sigma-Aldrich, 99.7%) and 2-naphthol (Sigma-Aldrich, 99%) were purchased and used without further purification. The supersonic expansion was obtained by expanding helium at a pressure of 4 bar through a 200 μm pulsed nozzle (General Valve). The carrier gas was seeded with 2-naphthol heated at 120 °C (393 K) and room-temperature anisole. Mass-resolved S₁ ← S₀ spectra were obtained by one-colour resonance-enhanced multiphoton ionisation (1c-REMPI). Molecules were excited by a tunable frequency-doubled dye laser (Sirah 0.02 cm⁻¹ resolution) that was pumped by a Nd:YAG laser (Quanta Ray, Spectra Physics). The ions were detected by a microchannel plate detector (RM Jordan, 25 mm) mounted on top of a 1 m linear time-of-flight mass spectrometer (RM Jordan, Wiley-McLaren type). Fluorescence excitation spectra were obtained by collecting the fluorescence by a two-mirror system composed of a toroidal mirror and a rotating planar mirror. The emitted light was focused on the entrance slit of a 25 cm monochromator (Huet M25) used under broadband conditions (~20 nm resolution) and was detected by a photomultiplier tube (PMT, Hamamatsu R2059).¹⁰ The ion or the PMT signal was averaged by an oscilloscope (Lecroy wavesurfer) and processed through a personal computer.

The vibrational spectra were obtained resorting to the IR-UV double resonance technique,² using either ion or fluorescence detection.^{32–35} To this end, two counter-propagating lasers were focused by a 500 mm focal length lens on the cold region of the supersonic expansion. The UV laser wavelength was fixed on a transition of a given conformer, while the IR laser was scanned in the region of the O–H and C–H stretch modes. An active baseline subtraction procedure was used to monitor the IR absorption by measuring the signal difference produced by successive UV laser pulses (one without and one with the IR laser present). The tunable IR source was a tabletop IR Optical Parametric Oscillator/Amplifier (OPO/OPA, LaserVision) with 3 cm⁻¹ resolution. The synchronisation between the lasers was performed by a computer-controlled home-made gate generator.

The IR-UV spectra were calibrated in the wavenumber domain against an atmospheric water reference using a linear calibration function $\tilde{\nu}_{\text{cal}} = 0.98677\tilde{\nu} + 55.37389 \text{ cm}^{-1}$. To convert from air to vacuum, the calibrated wavenumbers were subsequently divided by $n = 1.000273$ which is valid between 2.5 and 3.0 μm (from <https://refractiveindex.info/>, based on ref. 36). To convert the UV spectra into the wavenumber domain ($\tilde{\nu}^{-1} = n\lambda$), we used $n = 1 + 0.05792105 \cdot (238.0185 - [\lambda/\mu\text{m}]^{-2})^{-1} + 0.00167917 \cdot (57.362 - [\lambda/\mu\text{m}]^{-2})^{-1}$ (from <https://refractiveindex.info/>, based on ref. 37).

2.3 Conformer searches

Informed by the structures of related anisole complexes with aliphatic or aromatic hydroxy compounds (ref. 20, 21 and unpublished data), chemical intuition was used to generate different start structures and explore the conformational landscape of the 1:1 complex between *cis/trans*-2-naphthol and anisole. Initial geometries were constructed with GaussView



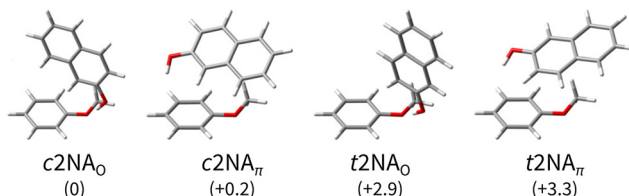


Fig. 2 Computed minimum-energy structures of 1:1 complexes between 2-naphthol (2N) and anisole (A). The 2-naphthol either forms a directional hydrogen bond with a lone pair of the anisole oxygen (O) or a distorted hydrogen bond with the electron-rich benzene ring, in favour of increased dispersion interaction between 2N and A (π). The complexes are sorted by their harmonically zero-point-corrected relative energies (in kJ mol^{-1}), as predicted at the B3LYP-D3_{abc}(BJ)/def2-QZVP level (Table 1).

and pre-optimised with Gaussian 09 Rev. D.01³⁸ at the B97-D/TZVP level.^{39,40} Unique configurations that remained after pre-optimisation were subsequently optimised at the B3LYP-D3_{abc}(BJ)/def2-TZVP level with TURBOMOLE 7.0.^{41–43} The absence of imaginary harmonic frequencies was taken as an indication for a (local) minimum on the multi-dimensional potential energy surface.

To independently check that no relevant structures were missed, we utilised the CREST program (Conformer-Rotamer Ensemble Sampling Tool).^{44,45} Using the recommended “-nci” option for non-covalently bound complexes, the four geometries from Fig. 2 were provided as initial start structures and the resulting configurations (within $E - E_{\min} \leq 25.1 \text{ kJ mol}^{-1}$) were again optimised at the B3LYP-D3_{abc}(BJ)/def2-TZVP level, this time using ORCA 5.0.3.^{46–48} Rewardingly, this second search did not reveal any new relevant structures. For consistency, all final DFT geometries utilised in this work were re-optimised with ORCA 5.0.3.

2.4 Employed density functionals

Density functional theory (DFT) results presented throughout this work, such as geometries, energies, and vibrational properties, were computed with two hybrid functionals, B3LYP^{49,50} and PBE0,⁵¹ using the so-called Karlsruhe basis sets (def2-XVP; X = S, TZ, QZ).⁵² In all cases, Grimme's two-body dispersion correction (D3; optionally with Axilrod-Teller-Muto three-body dispersion D3_{abc}) and Becke-Johnson damping (BJ) were included.^{53,54} Throughout all DFT geometry optimisation (“VeryTightOpt”) and analytic harmonic frequency calculations (“Freq Mass2016”), very strict thresholds were used (“DefGrid3 VeryTightSCF NoRI NoCosX”).^{46–48}

2.5 DLPNO and PNO single-point energy calculations

To obtain more accurate relative energy predictions beyond dispersion-corrected DFT, we used DFT-optimised reference geometries to compute frozen-core single-point electronic energies at the second-order Møller-Plesset (MP2 and its spin-component-scaled variant SCS-MP2⁵⁵) and the coupled-cluster singles, doubles, perturbative triples (CCSD(T)) level.

To expedite the calculations and achieve near-convergence with respect to the one-particle basis, their local (pair natural

orbitals, PNO) and explicitly correlated (F12) variants were used,^{56,57} as implemented in MOLPRO 2021.3^{58,59} and ORCA 5.0.3.^{47,48} The PNO implementation in ORCA is referred to as DLPNO which stands for “domain based local pair natural orbital”.

Strict thresholds ensured minimal errors on the relative energies due to the localisation approximations (“DomOpt = Tight PairOpt = Tight” for Molpro; “TightSCF TightPNO” for ORCA). In these correlated calculations, Dunning's augmented correlation-consistent basis sets (aug-cc-pVXZ; aVXZ) and their non-augmented F12 (cc-pVXZ-F12; VXZ-F12) variants by Peterson *et al.* were utilised from aVQZ up to VQZ-F12.^{60,61}

3 Survey of the conformational landscape

There are two stable conformers of 2-naphthol (Fig. 1), *t*2N and *c*2N, which are separated by a barrier of $\sim 15 \text{ kJ mol}^{-1}$ (from the lower-energy *c*2N well) with a relative energy difference of $\sim 2 \text{ kJ mol}^{-1}$. These values are computational predictions at the B3LYP-D3(BJ)/def2-TZVP level, including harmonic zero-point vibrational corrections, that were recently published by Hazrah *et al.*⁶² The relative energy between *c*2N and *t*2N has been estimated to be $\sim 1.7 \text{ kJ mol}^{-1}$,⁶³ although this value must be taken with caution. (Prior, Hollas and bin Hussein estimated a similar value,⁶⁴ providing error bars of $\pm 0.6 \text{ kJ mol}^{-1}$.) To the best of our knowledge, no experimental value of the barrier has been published so far.

In view of the high interconversion barrier between *t*2N and *c*2N and the relatively small *cis-trans* energy difference that leads to significant amounts of the higher-energy *t*2N conformer in the stagnation region, one may argue that there is no major complexation-assisted *trans-cis* relaxation during expansion (*vide infra*), thus leading to the formation of *c*2NA and *t*2NA complexes according to the monomeric free energy difference at the stagnation temperature. We, therefore, define the dissociation energy D' as

$$D'[x2NA] = E[A] + E[x2N] - E[x2NA] \quad \forall x \in \{c,t\} \quad (1)$$

which removes energy differences that are associated with the orientation of the hydroxy group in monomeric 2N.

As expected,^{20,21} there are two main bonding motifs predicted by the calculations (Fig. 2): one hinge-like structure where a rather directional, ‘classical’ O–H \cdots O hydrogen bond between 2N and a lone pair of the anisole oxygen atom is formed (2NA_O) and one where a distorted O–H \cdots π hydrogen bond to the benzene ring of anisole is realised along with increased dispersion interactions between the molecular planes (2NA _{π}). Other 2NA conformers found in the conformer search that are within a few kJ mol^{-1} above the minimum exhibit these two bonding motifs where only the relative 2N and A orientations slightly differ. Other than in the anisole or 1-naphthol homodimers,^{11,65} the π – π stacked motif is predicted to be unfavourable in the 1:1 2NA complex; the conformer searches did not yield a stable conformation. For the moment, we note that the hydrogen bond preference, π or

Table 1 Relative DFT energies (ΔE , in kJ mol^{-1}), dissociation energies (D' , eqn (1), in kJ mol^{-1}) and double harmonic OH stretching properties (signed shifts $\Delta\omega$ relative to the predicted c2N band centre 3820, in cm^{-1} ; IR band strengths S_{IR} , in km mol^{-1}) of 2N and 2NA complexes at the B3LYP-D3_{abc} (BJ)/def2-QZVP level. The subscript "e" indicates purely electronic and "0,h" harmonically zero-point-corrected properties

	Energies			OH stretch		
	ΔE_e	$\Delta E_{0,h}$	D'_e	$D'_{0,h}$	$\Delta\omega$	S_{IR}
c2N	0	0	—	—	0	67
t2N	2.1	1.7	—	—	+8	85
c2NA _O	0	0	35.2	30.7	-188	636
c2NA _π	1.2	0.2	34.0	30.5	-76	196
t2NA _O	3.3	2.9	34.0	29.6	-205	1091
t2NA _π	4.7	3.3	32.5	29.1	-39	138

oxygen lone pair, sensitively depends on the employed density functional, basis set, and inclusion of three-body dispersion (*abc*). These effects are quantified in more detail later (Section 6).

At the B3LYP-D3_{abc}(BJ)/def2-QZVP level, both bonding motifs are predicted to be very close in energy, well within 1 kJ mol^{-1} . We would, therefore, anticipate to observe at least four signals in the OH stretching spectra that correspond to c2NA_O, c2NA_π, t2NA_O, and t2NA_π. Their key electronic and (double harmonic) vibrational OH stretching properties at this level are summarised in Table 1. Based on the computed harmonic complexation shifts of the OH stretch, we anticipate to be able to easily distinguish between all four complexes *via* their spectroscopic signature in the OH stretching range (Table 1).

4 Analysis of the jet-cooled OH stretching spectra

4.1 FTIR spectra

4.1.1 The 2-naphthol spectrum. The FTIR spectra of jet-cooled 2-naphthol with varying water concentrations are shown in Fig. 3 (i and ii, grey) where overall six bands are observed which we label A–F. The 2-naphthol concentration and water impurity both increase from (i) to (ii). Based on relative intensities and previous spectroscopic studies,⁶⁶ A and B are easily identified as both conformers of monomeric 2-naphthol and assigned to t2N and c2N, respectively.

E and F clearly correspond to 1:1 2-naphthol hydrates as they are absent in spectrum (i) which has the lowest water concentration – the extent of water contamination can be assessed by the ro-vibrational lines of the antisymmetric water stretching vibration, labelled "W".⁶⁷ The vibrational signatures of 2-naphthol hydrates were previously characterised by Schütz *et al.*⁶⁸ and Matsumoto *et al.*⁶⁶ and the new band centres agree within the experimental resolution with previously reported values (*cf.* Table 2).

Lastly, C and D can be ascribed to clusters of 2-naphthol; on the basis of their scaling behaviour with increased 2-naphthol concentration (see ESI,† Fig. S1(iii)), we ascribe C to a homodimer whereas D corresponds to a larger, presumably trimeric, cluster. To the best of our knowledge, vacuum-isolated clusters

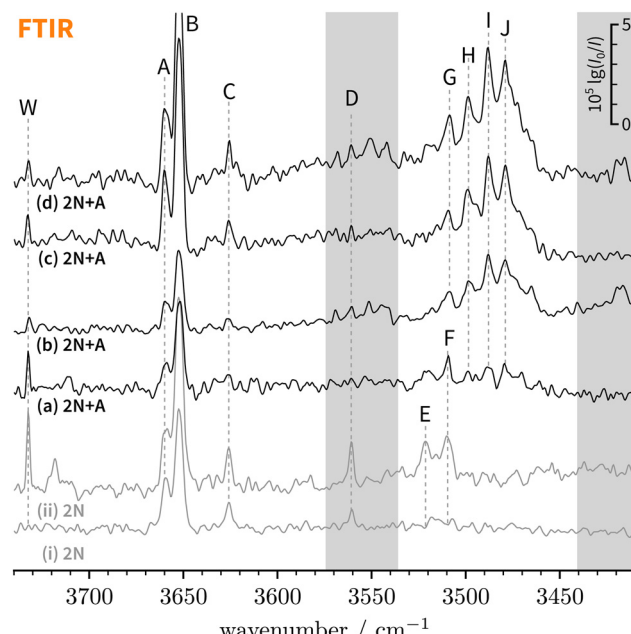


Fig. 3 FTIR decadic absorbance spectra ($\lg(l_0/l)$) of bare 2-naphthol expansions (i and ii, grey) and 2-naphthol and anisole co-expansions (a–d, black) with varying concentrations and water content. An extended version covering a broader spectral range can be found in the ESI† (Fig. S1). The overall signal enhancement from traces (a)–(d) illustrates the extended absorption path length in the new setup.³¹ Note that bands F and G partially overlap (see Section 4.1.2).

of 2-naphthol have so far not been spectroscopically investigated. The comparably small red-shift of C indicates that no classical hydrogen bond is formed. Explorative quantum chemical searches of the conformational landscape (not shown) further indicate dominant $\pi-\pi$ contributions to the binding energy, quite similar to the experimentally-characterised homodimers of 1-naphthol^{11,69} and 2-naphthalenethiol,⁷⁰ a heavier homologue of 2N. We refrain from assigning C to a particular dimeric 2N conformer which is not possible based on vibrational frequencies alone.¹¹

Table 2 Observed OH stretching vacuum band centres ($\tilde{\nu}_0$, in cm^{-1}) of 2-naphthol monomers and clusters (2N), 2-naphthol hydrates (2NW), and 2-naphthol-anisole complexes (2NA). Note that bands F and G partially overlap (see Section 4.1.2)

FTIR	$\tilde{\nu}_0$	IR-UV		Assignment
		$\tilde{\nu}_0$	$\tilde{\nu}_0$ (ref. 66)	
A	3659	3660	3661	t2N
B	3652	3653	3654	c2N
C	3626			(2N) ₂
D	3560			(2N) _{≥3}
E	3521		3523	t2NW
F	3510		3512	c2NW
G	3509	3509		c2NA _O
H	3498	3499		c2NA _O
I	3488	3489		c2NA _O
		3486		t2NA _O
J	3479	3479		t2NA _O



It is worth mentioning that under similar conditions, only traces of the higher-energy rotamer are observed in the jet-cooled FTIR OH stretching spectrum of 1-naphthol.²³ This can be ascribed to the nearly two-fold smaller *cis-trans* energy difference in 2-naphthol⁶² which leads to a higher initial population of the higher-energy rotamer than in 1-naphthol.

4.1.2 The 2-naphthol-anisole spectrum. The FTIR jet spectra of co-expanded 2-naphthol and anisole are shown in Fig. 3 (a-d, black). Traces (a) and (b) were measured with the old setup, (c) and (d) with the new one. The respective upper traces (b and d) have an increased 2-naphthol and anisole concentration relative to the respective lower traces (a and c). Details can be found in the ESI.[†] Even upon adding anisole to the expansion, both monomeric 2-naphthol signals persist in the spectra. Using calculated relative band strength ratios and experimental band integrals for *c*2N and *t*2N (see ref. 71), we obtain a conformational temperature¹⁵ of roughly 160–210 K when using the calculated energy difference of 2 kJ mol⁻¹ recently reported by Hazrah *et al.*⁶² Our own calculations (see ref. 72), including cautious estimates for thermal corrections⁷³ and other uncertainties, confirm an effective energy difference of (1.8 ± 0.4) kJ mol⁻¹, translating into a conformational temperature range of 120–220 K. This is only consistent with the nozzle temperature of about 390 K if there is significant relaxation over the ~ 15 kJ mol⁻¹ barrier in the jet,¹⁵ either in the initially expanding monomers or in hot re-dissociating complexes.^{74,75}

Turning now to mixed complexes, the question of relative abundance and visibility of *t* and *c* isomers arises. Assuming the same abundance ratio as for monomers, the *c* isomers will dominate. Anisole as a binding partner does not switch this energetic preference but rather seems to enhance it somewhat (Table 1). Based on spectral intensities in Table 1, O-bound isomers will generally be more visible than π -bound isomers. We thus do not expect to observe *t*2NA _{π} which is predicted somewhere between bands C and D. However, the OH stretch of *t*2NA_O is predicted to considerably gain in intensity relative to *t*2N. Furthermore, it is consistently predicted to be 1.6–2 times more visible than that of *c*2NA_O (all twelve combinations of B3LYP, PBE0, D3(BJ), D3_{abc}(BJ), def2-SVP, def2-TZVP, def2-QZVP). Thus, we would anticipate to also observe *t*2NA_O with a relative *t*2NA_O:*c*2NA_O intensity ratio between 2:3 and 1:3, if complexation-induced *trans-cis* relaxation is negligible. The latter assumption may not hold quantitatively. The energy freed upon complexation is in principle enough to induce thermal isomerisation even in a supersonic jet expansion.^{16,74,75} Such processes are still poorly understood, unlike corresponding laser-induced processes.^{76,77} Therefore, partial relaxation of *t*2NA_O to the further solvent-stabilised *c*2NA_O should not be strictly ruled out but is unlikely to be quantitative.

Overall, there are three main new absorption regions that correspond to mixed 2-naphthol-anisole clusters in Fig. 3(a-d). They are highlighted through two grey bars and the white space in-between. The main absorption feature in-between is observed even in the lowest-concentrated spectrum (a) whereas the features in the grey bars seem to be correlated. Because the signals on the right (near 3400 cm⁻¹) appearing in higher-

concentration spectra (b) and (d) are likely due to trimeric or larger aggregates, this could also be the case on the left (near 3550 cm⁻¹), close to D marking 2N homocluster traces, where *c*2NA _{π} would be expected at low intensity. The same applies to the even less abundant and visible *t*2NA _{π} , which is expected near 3600 cm⁻¹. In search of π -bound 2NA complexes, we tried to optimise the expansion conditions to maximise the new signals around D (left grey bar) while keeping the formation of trimers or higher clusters minimal (right grey bar), which proved to be difficult. All in all, one can at best derive upper bounds for π -bound 2NA complexes in the FTIR jet spectra and these bounds will not be very tight due to the smaller IR visibility of π -bound 1:1 complexes in the OH stretching region (Table 1). We return to the analysis in Section 6 and focus on the more conclusive O-bound absorption features in-between the grey bars for the remainder of this section.

Within the broad absorption around 3500 cm⁻¹, which roughly spans 60 cm⁻¹, there are several distinguishable peaks that we label G–J. Due to the presence of water impurities in the jet spectra, one might argue that G is not a new 2-naphthol-anisole band but actually a 2-naphthol hydrate signal (F, see Table 2). This is to a large extent the case for spectrum (a) but not (b) which displays only negligible amounts of water. By extension, this argument applies to spectra (c) and (d) where G, H, and I exhibit a similar intensity scaling ratio as in (b). Similarly, we can exclude anisole hydrates as possible origins.⁷⁸ While the overall broad absorption may well include contributions from trimeric or larger 2-naphthol-anisole clusters, we are cautiously optimistic that peaks H–J correspond to 1:1 complexes as they persist in spectrum (a). Therefore, this argument similarly applies to G. On the basis of the computed complexation shifts of different 2NA conformers (Table 1) and known shifts of similar O–H \cdots O hydroxy compound-anisole 1:1 complexes (phenol –174 and 1-naphthol –197 cm⁻¹, ref. 23), G–J are plausible candidates for *c*2NA_O and *t*2NA_O. But if this were the case, we would at most expect two signals in this spectral region and not four or more.

Left with three plausible explanations to account for the excessive number of persistent absorption features in Fig. 3(a-d) – conformational richness of 2NA 1:1 complexes, larger clusters, or spectral transitions beyond the standard double harmonic oscillator selection rules –, we recorded jet-cooled conformer- and mass-selective IR-UV double-resonance spectra to gain clarity. In view of the last possibility, it should be noted that the absorption features near D exhibit a similarly regular pattern of near-evenly spaced bands.

4.2 REMPI and IR-UV spectra

4.2.1 The 2-naphthol spectrum. The laser-induced fluorescence (LIF) spectrum of jet-cooled 2-naphthol is shown in Fig. 4(a). The vibrationless electronic band centres of the S₁ \leftarrow S₀ transitions of monomeric 2-naphthol are well-known and our band centres agree to within a few cm⁻¹ with previously reported values for *c*2N and *t*2N.^{63,64,66,68,79–81} Their IR-UV double-resonance OH stretching spectra are shown as insets (b) and (c) and the vibrational band centres agree to within 1 cm⁻¹ with



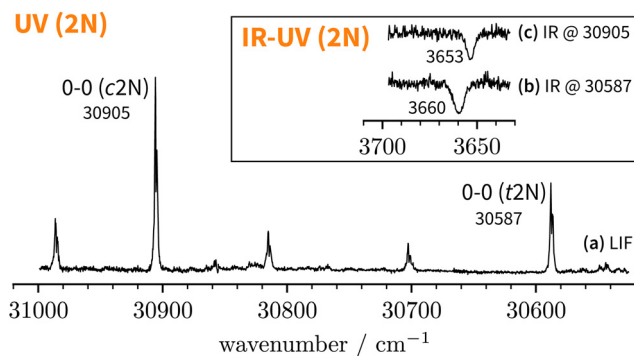


Fig. 4 LIF spectrum of jet-cooled 2-naphthol. IR-UV double-resonance spectra at selected $S_1 \leftarrow S_0$ transitions are shown as insets.

the FTIR and previously reported IR-UV values (Table 2). Having verified the calibration accuracy of the electronic and vibrational spectra of bare 2-naphthol, we can proceed to the discussion of the mixed spectra with anisole.

4.3 The 2-naphthol-anisole spectrum

Since the UV spectrum of the 2-naphthol-anisole complex had not been measured before, we first explored the UV region using mass detection (REMPI), then switching to fluorescence detection (LIF). The one-colour REMPI spectrum obtained by monitoring the 2NA mass channel is shown in Fig. 5.

Starting above $30\,500\text{ cm}^{-1}$, the REMPI spectrum exhibits several UV band systems. All the main electronic band systems exhibit a vibronic Franck-Condon progression built on a 10 cm^{-1} mode. By probing these UV band systems – blue arrows (I and II) and red arrows in Fig. 5 – with IR-UV double-resonance spectroscopy, it was revealed that all observed band systems correspond to only one conformer. The IR-UV spectra of this species are shown in Fig. 6. (II) was measured using mass detection and (I) using fluorescence detection to improve the signal-to-noise ratio. These spectra clearly exhibit three IR bands, at 3489 , 3499 , and 3509 cm^{-1} , which are again separated by 10 cm^{-1} . The three band centres match the FTIR band positions of I, H, and G, respectively (cf. Table 2).

The conformer-selectivity of IR-UV double-resonance spectroscopy only leaves two possible explanations for the two surplus bands. Either the measured ion intensity corresponds to ionic $1:1$ 2-naphthol-anisole complexes that fragmented from larger clusters or the neutral $1:1$ complex gives rise to more than one IR signal in the OH stretching range due to anharmonic effects. The former can be excluded for two main reasons. First, the preceding analysis of the FTIR spectra makes a mixed trimeric, let alone tetrameric, assignment for these three IR bands highly unlikely. Second, if more than one 2-naphthol were in the complex, we would either expect at least one band near(er) to the OH stretching band of bare 2-naphthol due to weaker $\text{O}-\text{H}\cdots\pi$ hydrogen bonds (or dangling OH groups) or at least one much more shifted IR band due to cooperativity effects since all OH groups would form $\text{O}-\text{H}\cdots\text{O}$ hydrogen bonds.

With the main UV band systems in Fig. 5 assigned to 2NA, we can determine the relative orientation of the 2-naphthol hydroxy group (*cis* or *trans*) and the hydrogen bond motif ($\text{O}-\text{H}\cdots\text{O}$ or $\text{O}-\text{H}\cdots\pi$) based on the UV and IR-UV spectra alone. This facilitates the discrimination of the four predicted 2NA complexes purely experimentally (Fig. 2). The magnitude of the OH stretch shift upon complexation is indicative of an $\text{O}-\text{H}\cdots\text{O}$ hydrogen bond and quantitatively matches previously measured complexation shifts of phenol-anisole and 1-naphthol-anisole.²³ Based on the complexation-induced UV shift, we can discriminate between $c2\text{NA}_\text{O}$ and $t2\text{NA}_\text{O}$ as the electronic $S_1 \leftarrow S_0$ transition is a much more sensitive probe to the 2-naphthol conformation ($30\,905$ and $30\,587\text{ cm}^{-1}$) than the OH stretch fundamental in the S_0 state (3653 and 3660 cm^{-1}). The several vibronic Franck-Condon progressions in Fig. 5 complicate a precise assignment of the $0-0$ band origin which we approximately locate at $30\,550\text{ cm}^{-1}$. If we assign it to $c2\text{NA}_\text{O}$, the resulting UV shift relative to $c2\text{N}$ is approximately -350 cm^{-1} . If referenced to $t2\text{N}$ instead, the shift is about -50 cm^{-1} . The suspected assignment to $c2\text{NA}_\text{O}$ can be confirmed by comparing to the UV shifts of $1:1$ complexes with a similar connectivity and bonding motif. The UV shift of anisole-phenol⁸² amounts to -352 cm^{-1} and $1:1$ hydrates⁶⁶ of *cis*- and *trans*-2-naphthol exhibit similar shifts of -371 and -332 cm^{-1} . Even

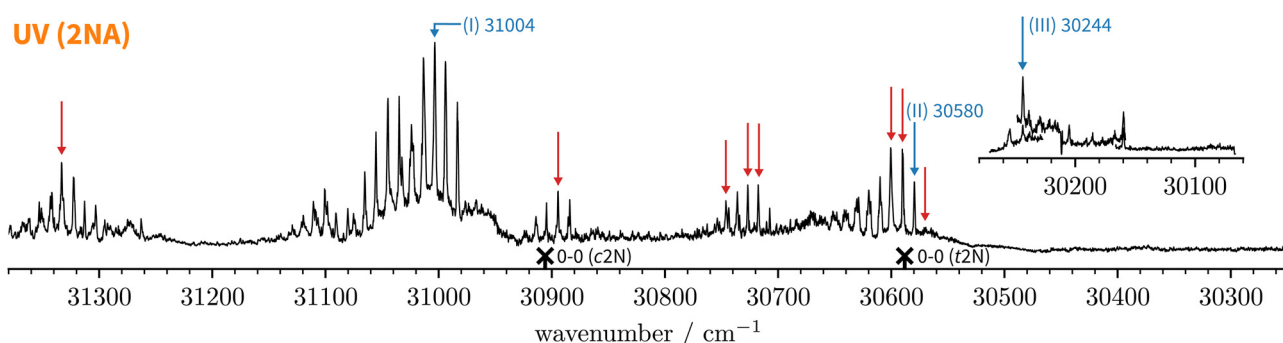


Fig. 5 Normalised one-colour REMPI spectrum of the jet-cooled co-expansion of 2-naphthol and anisole monitored at the 2NA mass channel (252 amu). The $0-0$ band positions of the $S_1 \leftarrow S_0$ transitions of $c2\text{N}$ and $t2\text{N}$ are shown for reference (crosses). The normalised LIF spectrum below $30\,280\text{ cm}^{-1}$ is shown as an inset. Blue arrows mark selected $S_1 \leftarrow S_0$ transitions where IR-UV double-resonance spectra were recorded (see Fig. 6). Red arrows mark all $S_1 \leftarrow S_0$ bands that were checked to exhibit the same IR spectrum as bands (I) and (II).



the $\pi-\pi$ stacked 1-naphthol⁶⁹ and anisole⁸³ homodimers show similarly large red shifts. This purely experimental assignment to $c2\text{NA}_\text{O}$ is in agreement with quantum chemical calculations that predict this conformer to be the global minimum structure (Table 1). It is worth noting that the observed vibrational spacing of 10 cm^{-1} in the electronic ground state (Fig. 6) effectively matches the UV fine structure of several observed Franck-Condon bands in the first excited electronic state (Fig. 5), indicating that the potential of the involved intermolecular vibrational mode is similar in the S_0 and S_1 states and rather harmonic.

To summarise, the overwhelming majority of observed UV signals in the jet-cooled REMPI spectrum (Fig. 5) correspond to $c2\text{NA}_\text{O}$. By employing size- and conformer-selective IR-UV spectroscopy, we could confirm that the FTIR bands G, H, and I all correspond to $c2\text{NA}_\text{O}$. We did not find evidence for the $c2\text{NA}_\pi$ complex, which is predicted by harmonically zero-point-corrected DFT to be near-isoenergetic, leaving the origins of the FTIR absorption features around D (left grey panel in Fig. 3) unclear. The absence of π -bound 2NA conformers in the UV spectra is not a conclusive proof of their absence in the jet expansion but could be attributed to rapid isomerisation to $c2\text{NA}_\text{O}$, rapid dissociation, or – rather unlikely – very small UV absorption cross sections.

Lastly, we must address the assignment of the FTIR band J at 3479 cm^{-1} which, based on its high intensity and scaling behaviour, most likely corresponds to a $1:1$ complex. $t2\text{NA}_\text{O}$ is the obvious choice. Based on the complexation-induced UV shift of $c2\text{NA}_\text{O}$, the 0-0 band of $t2\text{NA}_\text{O}$ is expected around 30250 cm^{-1} . The REMPI spectrum in Fig. 5, however, does not provide evidence for a $t2\text{NA}$ complex. This is probably due to the fact that the photon energy is too low for ionising the complex below $\sim 30500\text{ cm}^{-1}$. Indeed, the ionisation energy of bare 2-naphthol is 63 670 for $c2\text{N}$ and 63189 cm^{-1} for $t2\text{N}$.⁶³ Even if one considers the decrease of the ionisation energy upon complexation, it is possible that at 30300 cm^{-1} – the low-energy limit of the UV spectrum obtained by REMPI –, the photon energy is not sufficient for ionising the system. Therefore, we recorded the $S_1 \leftarrow S_0$ spectrum in the same region using LIF. Normalised LIF spectra that cover the relevant spectral range are shown as an inset in Fig. 5. We measured the IR-UV double-resonance spectrum by setting the probe on the UV band at

30244 cm^{-1} which is shown in Fig. 6(III). The OH stretching band exhibits a broad dip that appears to consist of several overlapping bands with a ‘maximum’ at 3479 cm^{-1} . The band centre is in very good agreement with the FTIR band centre of J. Moreover, it appears that the perturbations in $c2\text{NA}_\text{O}$ that give rise to additional IR bands are similarly present in $t2\text{NA}_\text{O}$, albeit with a decreased splitting. The third member in the progression is not well resolved in Fig. 6(III) but appears to be roughly centred at 3493 cm^{-1} . Indeed, there is an additional peak between the FTIR bands H and I at 3495 cm^{-1} close to the noise level which can best be seen in Fig. 3(a) and (c), providing further evidence that also $t2\text{NA}_\text{O}$ gives rise to (at least) three IR bands in the OH stretching range.

5 The split OH stretching band of $c2\text{NA}_\text{O}$

The experimental analysis in the preceding section unequivocally showed that the OH stretch fundamental of $c2\text{NA}_\text{O}$ in the electronic ground state is split into three (possibly more but less visible) evenly spaced (*i.e.* within the resolution uncertainty) bands at 3488 , 3498 , and 3509 cm^{-1} with decreasing intensity. The similarity of the IR-UV and FTIR intensity pattern rules out hot band interpretations. In increasing wavenumber, we assign them to ν_{OH} – the pure OH stretch fundamental –, $\nu_{\text{OH}}+\nu_{\text{inter}}$, and $\nu_{\text{OH}}+2\nu_{\text{inter}}$. ν_{inter} is an intermolecular vibrational mode with a fundamental wavenumber of approximately 10 cm^{-1} which is promoted by OH stretch excitation in the same way in which it is promoted by electronic excitation (Fig. 5), namely structural change along the mode in higher-energy states of the complex.

Based on the experimental separation of 10 cm^{-1} , the harmonic wavenumber of a suitable vibrational coupling partner should ideally be slightly larger than 10 cm^{-1} to leave room for some (diagonal) anharmonicity. To get a feeling for the numerical sensitivity of the low-frequency vibrations, we compared several different DFT calculations (all eight combinations of B3LYP, PBE0, D3(BJ), D3_{abc}(BJ), def2-TZVP, def2-QZVP). Very strict SCF and geometry optimisation thresholds and large DFT grids ensured numerical stability (see Section 2.4). In all these calculations, two intermolecular normal modes are consistently predicted below 20 cm^{-1} , which we label ν_{101} and ν_{102} following the nomenclature by Herzberg.⁸⁴ Fig. 7 shows that the second-lowest vibration (ν_{101} , $11\text{--}14\text{ cm}^{-1}$) is a plausible candidate as it modulates the O-H \cdots O hydrogen bond angle in a seesaw-like movement of the anisole.⁸⁵ This hydrogen bond angle, in turn, will likely be stiffened when the OH stretch is excited and gains in dipole moment. As a result, OH stretch intensity is redistributed to the intermolecular combination bands in a vibrational Franck-Condon fashion.^{25–30} In first approximation, the total intensity of this progression should be compared to the harmonic intensity predicted for the OH stretch. In the adiabatic picture, the intensity pattern can be rationalised through the squared wavefunction overlap of different n_{inter} states – the ‘slow’ mode – between the adiabatic $n_{\text{OH}} = 0$ and

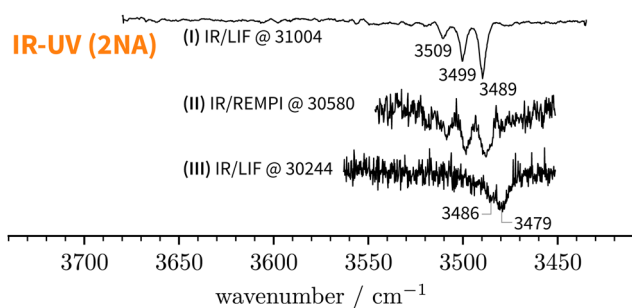


Fig. 6 IR-UV double-resonance spectra at selected $S_1 \leftarrow S_0$ transitions of 2NA (see Fig. 5).



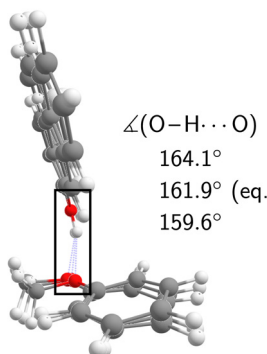


Fig. 7 Equilibrium geometry of $c2NA_O$ at the B3LYP-D3_{abc}(BJ)/def2-QZVP level superimposed with normal mode displacements in opposing directions along ν_{101} which modulates the O-H···O angle.

$n_{OH} = 1$ potentials which take the roles of the ground and excited electronic Born–Oppenheimer potentials in vibronic Franck–Condon theory. Dropping the adiabatic approximation in combination with realistic electric dipole surfaces may introduce further effects.⁸⁶ Studies along the lines of ref. 30, where only the relevant vibrational degrees of freedom are retained, could provide instructive insights into the vibrational couplings, complementing the adiabatic picture.

6 To π or not to π ?

The preceding analysis of the FTIR and IR-UV spectra showed that the O-H···O-bound 2NA complex is formed in the jet expansions and in both conformers, $c2NA_O$ and $t2NA_O$, the S₀ OH stretch fundamental is found to exhibit a Franck–Condon-like progression.

It remains to be addressed if the π -bound 2NA complex is formed in the FTIR jet expansion but perhaps concealed by other spectral contributions. For this purpose, we will investigate experimental uncertainty and computational error sources in the following.

6.1 Assessment of experimental uncertainty sources

There are essentially three uncertainties, preventing the accurate FTIR estimation of a reliable $c2NA_\pi : c2NA_O$ concentration ratio.

Firstly, the three OH stretch bands of $c2NA_O$ are superimposed by other absorption features, most notably the J band, preventing the determination of reliable experimental band integrals.

Secondly, and most importantly, it is unclear where $c2NA_\pi$ is spectrally located. Moreover, in view of the previously confirmed Franck–Condon-like progression in $c2NA_O$ and $t2NA_O$, this begs the question if similarly the already weakly IR-active OH stretch of $c2NA_\pi$ is affected. Indeed, we noted earlier that the band system around D in Fig. 3(b) could plausibly correspond to $c2NA_\pi$. By closely comparing the scaling behaviour of bands C and D from spectrum (a) through to (c), we can infer that the absorption feature at 3561 cm^{-1} in spectrum (b) only partially corresponds to a 2-naphthol homodimer (band D). Its spacing to the two neighbouring bands at 3570 and 3552 cm^{-1}

is about 9 cm^{-1} . A similar vibrational Franck–Condon progression in $c2NA_\pi$ thus appears feasible.

Thirdly, even if reliable band integrals of $c2NA_\pi$ and $c2NA_O$ could be obtained, there is a sizeable variation in their predicted band strength ratio from 0.21 to 0.47 (all eight combinations of B3LYP, PBE0, D3(BJ), D3_{abc}(BJ), def2-TZVP, def2-QZVP), emphasising the low and erratic visibility of the π conformer. It is particularly noteworthy that with B3LYP-D3, the band strength in the π conformer decreases by roughly 30% once subtle three-body terms are added to the two-body dispersion. Calculations with and without three-body dispersion on the respective other optimised geometry show that these are entirely geometric effects.

Depending on the computational method for band strengths and the assumed spectral spread of the $c2NA_\pi$ contribution, its non-observation could still be consistent with a slight excess over $c2NA_O$ in the FTIR-probed expansions. However, a trace role of $c2NA_\pi$ in relation to $c2NA_O$ is at least as likely, considering its non-observation by all explored spectral techniques. Because the driving force of hydrogen bond isomerisation is weak and the barrier rather broad, it is difficult to say whether a subtle energy penalty will result in substantial depopulation by simple carrier gas collisions or more elaborate mechanisms.^{16,74,75}

6.2 Assessment of computational error sources

The key property we wish to obtain from quantum chemistry is the vibrational zero-point-corrected relative energy $\Delta E_0^{\pi-O}$ which consists of a purely electronic ($\Delta E_e^{\pi-O}$) and zero-point vibrational contribution ($\Delta ZPVE^{\pi-O}$). For positive relative energies, the O-H···O hydrogen bonding motif is energetically preferred.

Starting with the relative harmonic ZPVE, the data in the ESI† (Table S1) show that this quantity quickly converges with respect to the basis set size. The impact of three-body dispersion and the density functional are of the order of a few 100 J mol^{-1} . Harmonic zero-point vibrational effects persistently predict the relative stabilisation of the π over the oxygen bonding site in the 2NA 1:1 complex. Until now, we have completely omitted vibrational anharmonicity, by assuming $\Delta E_0 \approx \Delta E_{0,h}$. To lowest-order, anharmonic contributions to the ZPVE can be approximated using second-order vibrational perturbation theory.^{87–89} The results (not shown) are not reliable due to a multitude of large-amplitude vibrations leading to imaginary anharmonic fundamentals. We do not expect the anharmonic ZPVE corrections to dominate the harmonic ones. Based on the harmonic DFT predictions in Table S1 (ESI†), we estimate a generous error bar of $\pm 0.5\text{ kJ mol}^{-1}$ to account for the missing vibrational anharmonicity, yielding a best estimate of $\Delta ZPVE^{\pi-O} = -(1.0 \pm 0.5)\text{ kJ mol}^{-1}$.

Remaining in the realm of harmonic DFT, we can study the impact of the basis size, the choice of exchange-correlation functional, and inclusion of three-body dispersion on $\Delta E_{0,h}^{\pi-O}$. The main results are visualised in Fig. 8, a more complete list of data can be found in the ESI† (Table S1). Starting from our initial method in the centre, *i.e.* B3LYP-D3_{abc}(BJ)/def2-QZVP, the vertical deviation reflects the basis set error, the deviation to the left the impact of three-body dispersion, and the deviation to the right the influence of the exchange–correlation



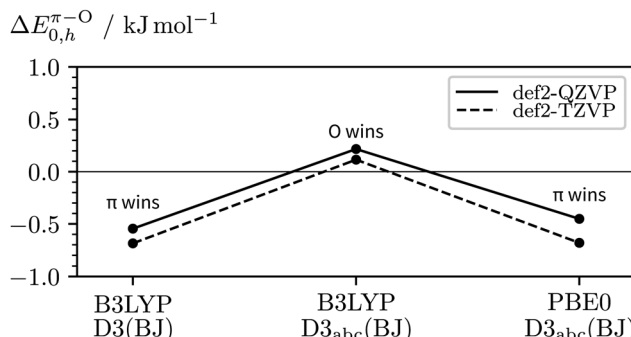


Fig. 8 Impact of the basis size, exchange–correlation functional, and inclusion of three-body dispersion on the relative stabilities of $c2NA_O$ and $c2NA_\pi$ ($\Delta E_{0,h}^{\pi-O}$).

functional when using PBE0 instead. Both, the change from PBE0 to B3LYP and inclusion of three-body dispersion lead to a relative destabilisation of the π hydrogen bond acceptor site which agrees with the primary experimental interpretation. While the trend of the DFT calculations weakly points towards $c2NA_O$ being more stable, even at the B3LYP-D3_{abc}(BJ)/def2-QZVP level, only a 0.2 kJ mol⁻¹ harmonically zero-point-corrected preference of O over π is predicted.

To see if there are significant electron correlation errors at the DFT-D3 level, we lastly consult PNO-coupled-cluster. As geometry optimisations are too expensive, the single-point energies are computed using the DFT equilibrium geometries. In the absence of microwave data (or other structural information) on the 2NA 1:1 complex, one can only speculate which DFT geometry is closest to the experimental one. One can also compare *a posteriori* which method gives geometries with the lowest PNO energy and should, therefore, be used in the PNO-LCCSD(T)-F12b/VDZ-F12 calculations (see Table S2, ESI[†]). Out of all optimised $c2NA_O$ geometries included in Fig. 8, B3LYP-D3_{abc}(BJ)/def2-TZVP is closest to the ‘true’ coupled-cluster minimum, closely followed by B3LYP-D3_{abc}(BJ)/def2-QZVP (+0.1 kJ mol⁻¹) whereas the PBE0-D3_{abc}(BJ)/def2-TZVP geometry is already 1.7 kJ mol⁻¹ higher in energy, providing further evidence that B3LYP is structurally more reliable than PBE0 for this system. Remarkably, when instead comparing the relative energies ($\Delta E_e^{\pi-O}$), PNO-LCCSD(T)-F12b/VDZ-F12 is found to be relatively insensitive to the reference geometry with deviations of the order of a few 100 J mol⁻¹, the main error source being the geometry of $c2NA_\pi$. In Fig. 9, the basis set convergence of $\Delta E_e^{\pi-O}$ is examined for Møller–Plesset and coupled-cluster predictions. Additional PNO data, including dissociation energies, are reported in the ESI[†] (Table S2). Fig. 9 shows two things. Firstly, PNO-LCCSD(T)-F12b/VDZ-F12 unambiguously favours $c2NA_O$ over $c2NA_\pi$. The F12 approximation errors are insignificant and relative electronic energies between 3.2 and 3.5 kJ mol⁻¹ are obtained with Molpro’s PNO-LCCSD(T)-F12a and -F12b and ORCA’s DLPNO-CCSD(T)-F12 and -F12D when using the VDZ-F12 basis. Secondly, PNO-LMP2 consistently overestimates the π binding site due to its well-known overestimation of dispersion effects. PNO-SCS-LMP2 significantly improves upon PNO-LMP2 but the

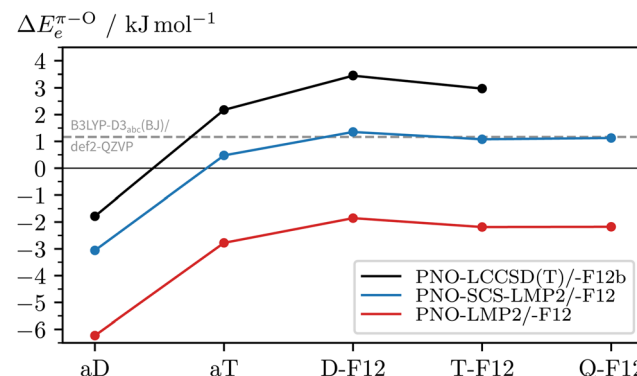


Fig. 9 Comparison of PNO (with $aX \equiv aVXZ$) and PNO-F12 (with $X-F12 \equiv VXZ-F12$) predictions of the relative electronic stabilities of $c2NA_O$ and $c2NA_\pi$ ($\Delta E_e^{\pi-O}$) for increased coverage of the one-particle basis space. The optimised B3LYP-D3_{abc}(BJ)/def2-QZVP geometries are used (reference values shown in grey).

converged relative energies are not an improvement over B3LYP-D3_{abc}(BJ)/def2-QZVP. Inspection of the absolute complex stabilities, *i.e.* the dissociation energies (Tables S1 and S2, ESI[†]), reveals that the main deviations between PNO-LCCSD(T) and the investigated DFT-D3 functionals arise from an overestimation of the binding energy of the π -bound complex by the latter, even when three-body dispersion is included. Informed by the parallel LMP2/SCS-LMP2 basis set progression in Fig. 9, we expect a stabilisation of the O binding site slightly below 3 kJ mol⁻¹ at the PNO-LCCSD(T) basis set limit.

Based on the estimates for the relative electronic and zero-point vibrational energies, we arrive at 1.0–3.0 kJ mol⁻¹ for the predicted total stabilisation of $c2NA_O$ over $c2NA_\pi$ at $T = 0$ K. It should be noted that we have entirely neglected calculated temperature effects, *e.g.* by comparing rigid-rotor-harmonic-oscillator Gibbs free energies $\Delta G_{RRHO}^{\pi-O}$,⁷³ which are inconclusive and highly erratic. Furthermore, the complexes are believed to form and still interconvert at relatively low temperatures, other than monomers with high interconversion barriers from the start of the expansion.

7 Conclusions

When 2-naphthol complexes anisole, the orientation of its OH group may either be *cis* or *trans* and it may either form a hydrogen bond to the ether group or to the π system of anisole. Because the barrier between *cis* and *trans* is high, it is not easily overcome in a supersonic jet expansion. Experimental evidence presented in this work by FTIR and IR-UV spectroscopies is consistent with a partially frozen distribution. The hydrogen bond preference of the complex is more interesting and has been explored in this work by a combination of experimental and theoretical methods. It results from a balance of hydrogen bonding and less directional London dispersion forces between the two aromatic ring systems. Experimentally, the question has to remain open because of the strong π hydrogen bond distortion and thus weak IR intensity and because of an



interesting dynamical feature in the vibrational spectrum of at least one of the isomers. We unambiguously showed that OH stretching excitation of *cis*-2-naphthol bound to the ether group leads to such a large change in the preferred complex geometry that it is accompanied in about 50% of all vertical transitions by the simultaneous excitation of a global scaffold vibration, which is three orders of magnitude slower. Such a Franck-Condon-type progression based on the adiabatic separation of fast and slow degrees of freedom is less frequently observed in vibrational spectroscopy than in electronic spectroscopy. However, it seems to be characteristic of the interaction between a phenol-containing molecule and an ether.^{27–29} For the present system, ring methylation of the anisole, as a means to stabilise the π binding site²¹ and to modulate the Franck-Condon progressions, appears particularly promising in view of some exploratory quantum chemical calculations.

The experimental elusiveness of the π -bonded structure has triggered theoretical investigations which are indeed diverse and thus educational in terms of the quality of electronic structure methods in describing hydrogen bonds to different sites in a molecule. Popular dispersion-corrected density functional calculations close to the basis set limit with important harmonic zero-point vibrational energy corrections predict an almost perfect energy balance between the competing isomers where even three-body dispersion corrections or different hybrid density functionals switch the sign of the energy sequence. Basis-set converged wavefunction methods also span the entire range from clear π -favouring (MP2) over undecided (SCS-MP2) to clearly ether-favouring (CCSD(T)) coordination. Direct detection of the π -bound conformation by a structural method^{11,12} may add further insight to this subtle and demanding intermolecular energy balance. It could turn this complex into a true benchmark for electronic structure theory, in addition to its here demonstrated role as a case of vibrational excitation-induced structural change which leads to a vibrational Franck-Condon-like progression detected by both direct absorption and size- as well as conformation-selective spectroscopies.

Author contributions

Conceptualisation: AN, AZ, MAS; formal analysis: AN; funding acquisition: AZ, MAS; investigation: AN, AFPM, ML, IA; supervision: AZ, MAS; visualisation: AN; writing – original draft: AN (lead), ML, AZ, MAS; writing – review & editing: AN, AZ, MAS.

Conflicts of interest

There are no conflicts to declare.

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