High-resolution two-dimensional electronic spectroscopy reveals the homogeneous line profile of chromophores solvated in nanoclusters

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Doped clusters in the gas phase provide nanoconfined model systems for the study of system-bath interactions. To gain insight into interaction mechanisms between chromophores and their environment, the ensemble inhomogeneity has to be lifted and the homogeneous line profile must be accessed. However, such measurements are very challenging at the low particle densities and low signal levels in cluster beam experiments. Here, we dope cryogenic rare-gas clusters with phthalocyanine molecules and apply action-detected two-dimensional electronic spectroscopy to gain insight into the local molecule-cluster environment for solid and superfluid cluster species. The high-resolution homogeneous linewidth analysis provides a benchmark for the theoretical modelling of binding configurations and shows a promising route for high-resolution molecular two-dimensional spectroscopy.
The control and spectroscopic study of nanoconfined systems is of high relevance in many fields of research, ranging from quantum technologies to light harvesting applications, catalysis and spectroscopy. To understand the structure–function relationship on the nanoscale, individual structural configurations have to be resolved with high resolution, while, ideally, parasitic environmental perturbations are minimized. These challenges are best met in studies of isolated nano-objects in the gas phase.

In cluster-isolation spectroscopy rare-gas clusters serve as nanoconfined matrix to isolate individual spectroscopic probes in the gas phase. Helium (He) clusters form superfluid nanodroplets which efficiently cool dopant species to sub-Kelvin internal temperatures inside a homogeneous solvent, providing ideal conditions for high-resolution spectroscopy, e.g. of isolated molecules, radicals, aggregates, and complexes. More recently, solid rare-gas clusters were employed as a nanoconfined cryogenic substrate facilitating the formation of molecular networks with tunable interaction strength on the cluster surface. This approach has revealed cooperative molecular mechanisms such as superradiance and singlet fission with spectral resolution not achievable in thin film and bulk experiments. While the cluster-isolation technique was so far predominantly used for high-resolution spectroscopy of embedded species, the heterogeneous cluster systems may also serve as models for the study of structural configurations in isolated nanoconfined systems.

Insight into system–bath interactions, e.g. the coupling to the bath modes, the co-existence and configuration of different binding sites and their dynamic rearrangement are generally gained from the absorption line profiles. Thereby, the key challenge is the extraction of the homogeneous line profile from the inhomogeneously broadened ensemble response. However, most methods for homogeneous linewidth retrieval are not compatible with the low particle densities of gas-phase cluster samples. Hole burning, as an exception, has been applied to doped He nanodroplets, however did not resolve the homogeneous linewidth. This technique faces the difficulty of adapting the laser parameters to the time and frequency scales of the target system, reducing the approach mainly to photochemical hole burning where photoreactive chromophores are probed on quasi-infinite time scales. These challenges may explain why in heterogeneous cluster samples the homogeneous linewidth has not been determined, so far.

Recently, two-dimensional electronic spectroscopy (2DES) has been established, which is an ultrafast nonlinear spectroscopy technique enabling the disentanglement of homogeneous and inhomogeneous broadenings while automatically adapting to the time–frequency scale of the probed ensemble. 2DES and 2D infrared spectroscopy have proven as very useful in condensed-phase systems to extract line shape information where other methods are not applicable or do not provide the required time–frequency resolution. Due to technical challenges, the method’s potential in gas phase experiments is hardly explored.

Here, we apply the method to a cluster-isolated chromophore in the gas phase, which enables us to resolve the homogeneous absorption profile of the system. As chromophore we chose free-base phthalocyanine (H$_2$Pc), which belongs to a class of aromatic molecules of high relevance in optoelectronics, nonlinear optical materials and photobiology. Our results provide insight into the molecule-surface binding configurations with a resolution far beyond the accuracy and resolution of current density functional theory approaches. This offers a perspective for resolving the role of local configurations in photo-chemical reactions and opens a route for ultrafast multidimensional spectroscopy studies of isolated molecular systems with high spectral resolution.

**Results**

**Experimental scheme.** Figure 1 summarizes the experimental scheme. A supersonic beam of rare-gas clusters is generated in a molecular beam apparatus and is doped with H$_2$Pc molecules (details in the “Methods” section). In our study, we compare the molecular response of a single chromophore dissolved inside a superfluid He droplet with the response of 2–3 chromophores attached to the surface of a solid Ne cluster (Fig. 1a). At the low equilibrium temperatures of the nano-systems (He droplet: 0.37 K, Ne cluster: 10 K), only the lowest vibrational state of the dopant molecule is thermally occupied. Nearest-neighbor interactions between the chromophores are well-suppressed in both cluster-isolation experiments and no spectral signatures (line shifts/splittings) of inter-molecular couplings are observed despite the high spectral resolution of the experiment.

So far, 2DES is mainly performed in the condensed phase and the desired nonlinear signals are separated from the background by non-collinear four-wave mixing geometries (coherent-detected 2DES). Conversely, for the dilute cluster beam samples a collinear beam geometry is needed combined with the detection of an optical action (action-detected 2DES). In the latter, the sample is excited with a sequence of four femtosecond laser pulses (Fig. 1b) and the fourth-order light–matter response is deduced from the detected fluorescence. The pulse delays τ, t are interferometrically scanned to track the free polarization decay of the sample induced by pulses 1 and 3, respectively. Accordingly, a Fourier transform with respect to τ, t yields 2D frequency-spectra, which directly correlate the excitation (x-axis) and detection (y-axis) frequencies, while the time delay T determines the time evolution of the correlation spectra (Fig. 1c, d).
The ultralow optical density of doped cluster beam samples\textsuperscript{27}, requires a highly sensitive 2DES apparatus\textsuperscript{29}, which adapts the phase modulation technique developed by Marcus and co-workers\textsuperscript{38}. This method implements carrier-envelope-phase modulation of the optical pulses on a shot-to-shot basis at a high laser repetition rate (200 kHz). The phase modulation leads to characteristic beat notes in the fluorescence yield based on which the linear and nonlinear signal contributions are efficiently separated and amplified using lock-in detection. Thereby, the lock-in detection suppresses phase noise in the interferometric measurement and greatly enhances the detection sensitivity (details in the “Methods” section). These properties enabled several studies of highly dilute samples\textsuperscript{27,39,40} and quantum interference measurements at extremely short wavelengths\textsuperscript{41,42}.

The relevant information content of the 2D spectra is schematically shown in Fig. 1c, d. Peaks on the diagonal position of the 2D spectra closely resemble the linear excitation spectrum of the system, whereas off-diagonal peaks directly disclose nonlinear couplings, which are normally obscured in linear absorption measurements (Fig. 1c)\textsuperscript{13}. In addition, 2D lineshapes provide insight into ensemble inhomogeneities and their dynamic evolution (Fig. 1d). With the coherent multipulse excitation scheme, nonlinear rephasing and non-rephasing signals are recorded which lead to a separation of the inhomogeneous (homogeneous) lineshape along the diagonal (anti-diagonal) projection of the 2D spectra, accordingly. At time $T=0$ fs this effect leads to an elongated peakshape indicating a strong correlation between the absorption and detection frequency. For $T>0$ statistical fluctuations of the local environment lead to a loss of the correlation resulting in increasingly symmetric peak shapes (termed spectral diffusion). This concept is known from 2D photon echo spectroscopy as performed in coherence-detected 2DES\textsuperscript{20,43}. The equivalence between coherence and action-detected 2DES in terms of lineshape information was shown in ref. 44. We note that coherence and action-detected 2DES may differ in the detection of excited state absorption and multiple quantum coherence signals\textsuperscript{45,46} which are, however, not relevant in the current study.

### 1D coherence scans

So far, the majority of 2DES studies have been performed in the liquid and solid phases\textsuperscript{29}. To relate our approach to experiments in the condensed phase, we perform 1D coherence scans (using pulses 1 and 2 only) of H$_2$Pc attached to Ne clusters (denoted H$_2$Pc–Ne$_N$) and He nanodroplets (denoted H$_2$Pc–He$_N$) and compare the Fourier spectrum to the linear absorption spectrum of H$_2$Pc in an organic solvent (Fig. 2). Figure 2a shows the 1D coherence scan of the S$_1 \leftrightarrow$ S$_0$ transition in H$_2$Pc–He$_N$, featuring a clean, long-lived oscillation of the electronic coherence in the time domain. While in the condensed phase, electronic coherences are strongly perturbed and decay typically within <100 fs\textsuperscript{47}, they persist for more than 100 ps in H$_2$Pc–He$_N$ (Fig. 2a) and ~3 ps in H$_2$Pc–Ne$_N$ (not shown), implying a weak coupling of the cluster environments to the molecule’s electronic degrees of freedom. We note, that the full coherence decay in H$_2$Pc–He$_N$ extends beyond the experimental observation window. The observation of a single oscillating frequency in Fig. 2a suggests, that the molecule is initially prepared in a single state and the laser field drives a single vibronic transition. This offers ideal conditions for coherent control applications as well as optical trapping and cooling of molecules\textsuperscript{48,49}.

A Fourier transform of the 1D coherence scans yield the linear absorption spectra of both systems (Fig. 2b). For reference we show the H$_2$Pc absorption spectrum in a 1-chloronaphthalene solution. Comparing the molecular response in the three different environments, a clear trend can be observed. An increasing line broadening and red shift of the absorption band occurs when going from H$_2$Pc–He$_N$ to H$_2$Pc–Ne$_N$ and H$_2$Pc in 1-chloronaphthalene. This can be explained by the reduced perturbation and lower temperature in the cluster-isolated molecular samples (cf. Fig. 2c). A zoom on the Q$_x$ absorption band of the cluster-isolated molecules is shown in Supplementary Fig. 1 along with a comparison to high-resolution steady-state laser excitation spectra.

In the H$_2$Pc–Ne$_N$ spectrum several sharp peaks at much lower amplitude are observable clearly separated from the Q$_x$ absorption. These lines are assigned to complexes of H$_2$Pc and clusters of H$_2$O, N$_2$, or O$_2$ as also discussed in refs. 50,51. While the superfluid, homogeneous He solvent favors the formation of heterostructures, complex formation on the surface of solid Ne clusters is less likely, explaining the absence of respective spectral signatures in this system.

### 2D spectra

Having compared the linear spectra to studies in solution, we turn now to the nonlinear 2DES experiments. Figure 3a shows the 2D spectrum of H$_2$Pc–He$_N$ featuring several sharp, highly resolved diagonal peaks and no discernible off-diagonal resonances. Note, that the color scale is saturated by a >3p s in H$_2$Pc–He$_N$ (Fig. 2a) and ~3 ps in H$_2$Pc–Ne$_N$ (not shown), implying a weak coupling of the cluster environments to the molecule’s electronic degrees of freedom. We note, that the full coherence decay in H$_2$Pc–He$_N$ extends beyond the experimental observation window. The observation of a single oscillating frequency in Fig. 2a suggests, that the molecule is initially prepared in a single state and the laser field drives a single vibronic transition. This offers ideal conditions for coherent control applications as well as optical trapping and cooling of molecules\textsuperscript{48,49}.

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For H$_2$Pc–Ne$_N$ the 1D absorption spectrum (Fig. 2b) reveals a much broader line profile reminiscent of systems with multiple vibronic transitions. Figure 3c shows the respective 2D spectrum of H$_2$Pc–Ne$_N$. Here, we observe a single, strongly elongated diagonal peak. At the cryogenic temperatures of the Ne cluster environment, only the lowest vibrational state of H$_2$Pc is populated. The absence of cross-peaks thus implies the coupling of the ground state to a single vibronic transition in analogy to the H$_2$Pc–He$_N$.

The broad linear absorption spectrum can be thus purely attributed to ensemble inhomogeneities. A quantitative 2D lineshape analysis yields the inhomogeneous and homogeneous line profiles of the S$_1$ ← S$_0$ transition in H$_2$Pc–Ne$_N$. To this end, we performed a 2D peak fit by adapting the model from ref. 55 (details in Supplementary Note 2). The 1D projections of the fit result are shown in Fig. 3d, e. We find excellent agreement between the fit model and the experimental data, except for an outlier at 15,040 cm$^{-1}$. This outlier is not observed in our other data (Fig. 2) and is attributed to an experimental artifact.

We deduce for the inhomogeneous broadening a value of 23 cm$^{-1}$ and a remarkably narrow homogeneous broadening of 0.42 ± 0.01 cm$^{-1}$ (both full-width at half maximum values). These values are not limited by the resolution of the experimental apparatus. To the best of our knowledge, this is the first experimental determination of the homogeneous broadening in such hetero nanosystems. We experimentally determined the fluorescence lifetime to be >10 ns (details in the “Methods” section), in good agreement with fluorescence lifetimes in bulk rare gas matrices. This implies a negligible lifetime contribution to the homogeneous linewidth. The homogeneous broadening thus reflects in good approximation the pure dephasing rate of the system, which, in analogy to studies in bulk matrices, is attributed to an elastic scattering with phonon modes of the rare gas cluster. A sample-temperature-dependent study of the homogeneous linewidth would provide further insight into the phonon scattering as commonly done in the condensed phase. However, the infrared inactivity and rapid evaporative cooling of the nanoclusters prevent any means for heat injection.

The high-resolution data separating the homogeneous and inhomogeneous linewidth allows us to estimate a lower limit for the number of binding configurations between the H$_2$Pc molecule and the cluster surface. To this end, we fitted the inhomogeneous lineshape with a simplistic model assuming an equidistant spread of homogeneous line profiles. Fit parameters were the density of homogeneous absorption lines $n_l$ and their amplitudes $A_l$. We optimized the fit by gradually increasing the line density and considered as convergence criterion the point where the spectral modulations in the fit function are on the level of the noise of the experimental data (Fig. 4). We find that a minimum line density of $n_l \geq 4.0$ lines/cm$^{-1}$ (corresponds to 360 configurations) is necessary to fit the data. Taking the finite spectral resolution of the experiment into account, this constrains the number of binding configurations to a lower limit of ≥216 (details in the Supplementary Note 3), which corresponds to a mean energetic separation of the binding configurations of only 0.42 cm$^{-1}$.

This information about binding configurations is clearly beyond the resolution and accuracy of current theory and may help to gauge new models of molecule-surface binding configurations, as they play an important role in, e.g. surface chemistry. Here, we attribute the large number of binding configurations to the different geometric orientations of the molecule with the cluster lattice and to cluster surface defects (icosahedral cluster structure), while only a minor effect is expected from the statistical cluster size variation in the probed ensemble. The latter is rationalized by the fact, that the current study employed fairly
small clusters (=400 Ne atoms per cluster)\textsuperscript{58}, while basically the same inhomogeneous line profile was obtained for much larger cluster sizes (see nanosecond laser excitation experiment in Supplementary Fig. 1). In contrast, experiments in bulk Ne matrices showed an almost one order of magnitude broader inhomogeneous linewidth of \( \approx 200 \text{ cm}^{-1} \)\textsuperscript{156}. The same behavior is observed for 3, 4, 9, 10-perylenetetracarboxylic dianhydride (PTCDA)\textsuperscript{59}. The much broader inhomogeneous linewidth in the bulk environment may be explained by the formation of a solvation shell when the molecule is fully embedded in the bulk matrix compared to when bound to the surface of a cluster. At the same time, the homogeneous broadening of H$_2$Pc–Ne$_{50}$ seems in reasonable agreement with the values obtained in bulk matrices of heavier rare-gases where a factor of \( \approx 5 \) smaller homogeneous broadening is observed\textsuperscript{60}. The reduced phonon broadening may be explained by the lower matrix temperature (4 K) as compared to the cluster samples with temperatures of \( \approx 10 \text{ K} \).

2DES also offers time-resolved information about binding configurations (cf. Fig. 1d). The principle is well established in the condensed phase, where thermal fluctuations in the surrounding bath lead to a rearrangement of the binding configurations over time\textsuperscript{23,61}. The spectral diffusion of 2D lineshapes thus mainly reflects the solvent dynamics. In contrast, in the cryogenic cluster environment thermal fluctuations are absent and changes in the binding configurations may be directly attributed to the dynamics of the dopant molecules. In this context the question arises, if an electronic excitation of the dopant can change the binding configuration with the environment. Such mechanisms are common for species dissolved in He droplets, leading, e.g. to a migration towards the droplet surface followed by the ejection of the dopant on a pico to femtosecond time scale\textsuperscript{62,63}.

In the 2D spectra of H$_2$Pc–Ne$_{50}$, the onset of any configuration change would be observable by a broadening of the anti-diagonal linewidth or a reduction of the correlation between excitation and detection frequencies with increasing time \( T \). We studied the time evolution of the hetero-nanosystem for an evolution time of 0–500 ps at a reduced spectral resolution (2.3 cm\(^{-1}\) for anti-diagonal linewidth) to limit the measurement time (Fig. 5). No significant temporal changes can be observed (Fig. 5c), implying that fluctuations in the binding configurations are below the experimental resolution of 2.3 cm\(^{-1}\) (0.3 meV). This accounts for molecules populated in the electronic ground and excited state, since the response of both is reflected in the 2D spectra. We conclude that the molecules are mainly immobile on the cluster surface on the time scale of 500 ps, which was previously only indirectly confirmed for ground state molecules\textsuperscript{59}. We note, that the rotation of the cluster in the laboratory frame adds in principle another contribution to the inhomogeneous broadening due to the cluster size distribution in the probed ensemble. This broadening is however expected to be negligible due to the large moment of inertia of the clusters.

On the contrary, dopant species are highly mobile on the surface of superfluid He droplets as reflected in the formation of alkali complexes on the droplet surface\textsuperscript{64}. Likewise, species fully immersed in the quantum fluid clusters exhibit high mobility. Their rotational motion is well studied and served as a probe of the internal temperature of He clusters\textsuperscript{35}. The solvation mechanisms of organic chromophores in superfluid He droplets have been previously studied with high-resolution frequency-domain spectroscopy\textsuperscript{65}. It was rationalized that solutes are embedded in a non-superfluid He solvation layer. The different configuration of the solvation complexes lead to spectral finestructures\textsuperscript{19,66} while the dispersive solute–cluster interaction causes asymmetric lineshapes due to the statistical cluster size distribution\textsuperscript{30}. The homogeneous linewidth for the ZPL of solutes in superfluid He has not been determined to date. High-resolution 2DES may solve this problem and provide new insight into the solvation mechanisms in superfluid media. For Pc–He$_{50}$, an inhomogeneous linewidth of \( \geq 0.1 \text{ cm}^{-1} \) was found and an estimate for the homogeneous broadening of 0.02 cm\(^{-1}\) was indirectly deduced\textsuperscript{30}. Both are beyond our current experimental resolution. This could be solved by deploying advanced sampling strategies\textsuperscript{67–69} or implementing frequency-comb technology\textsuperscript{53} in order to increase the resolution to the required regime, thus, opening a promising perspective for the study of solvation mechanisms in superfluid environments.

In conclusion, the current study explored the nonlinear response of isolated chromophores in different nanosystems with femtosecond time resolution. The unique combination of 2DES with cluster-isolation techniques uncovered the homogeneous lineprofile beyond the ensemble average, providing insight into
the molecule-surface binding configurations with high resolution. Other methods for homogeneous linewidth retrieval cannot provide the combined time–frequency resolution of 2DES. The presented high-resolution 2DES approach, thus, provides a promising perspective for the analysis of molecular dynamics as a function of the local environment with femtosecond time resolution. Conversely, the here studied nanosystems represent highly resolved, well-defined molecular two-level systems which offer ideal conditions to explore many-body mechanisms of high interest in photovoltaics, such as molecular excitation migration and annihilation processes, as well as in quantum information science which is, so far, predominately limited to atomic samples. The potential of studying molecular networks on nanoconfined cluster surfaces was recently demonstrated and the advantage of 2DES in uncovering inter-particle interactions and cooperative dynamics was proven in previous studies.

Methods

Sample preparation. The experimental apparatus including the cluster beam preparation is described in detail in a recent review article. Further information about the sample preparation is also given in refs. 6, 7. Briefly, a continuous jet of the rare gas (5 μm orifice). The nozzle temperature is 14 K for He (mean cluster size of N\text{He} = 15,000 atoms) and 70 K for Ne (N\text{Ne} = 400\text{,}000), respectively. The cluster beam is skimmed by a 400 μm conical skimmer positioned in a distance of 15 mm from the nozzle. The cluster beam is doped in the adjacent chamber, where it passes a 10 mm long, heated oven containing H2Pc powder (Sigma Aldrich, 29H13P1Hthalocyanine, 98%). The doping follows a statistical pick-up process described by Poi- ssonian statistics and is controlled by the cell temperature and, thus, the density of evaporated molecules. The cell temperature is optimized for maximum fluorescence signal, which corresponds to molecule densities of 5 × 10^12 cm^-2 (350 °C cell temperature) for He and 3.2 × 10^13 cm^-3 (380 °C) for Ne, respectively. The fluorescence lifetime is due to the effusively emitted hot molecules from the oven cell. The doped cluster beam passes an orifice of 5 mm into the next chamber to suppress the latter contribution. In this chamber (3.8 × 10^-17 mbar base pressure) the femtosecond laser pulses interact with the cluster beam at right angle and the sample fluorescence is imaged with a lens onto a photo multiplier tube mounted in perpendicular direction to the laser and cluster beam propagation. The detector arrangement collects ~17% of the total fluorescence. The detector response time is 0.57 ns. The fluorescence lifetime was determined by tracking the decay of the fluorescence yield with a fast digital-to-analog converter (bandwidth: 500 MHz, sampling rate 1 G/s).

Spectroscopy method. The high-repetition-rate femtosecond laser system and phase-modulated 2DES setup is described in detail elsewhere. Laser parameters were: center wavelength l = 670 nm, spectral width 25 nm (FWHM), pulse duration ~50 fs, pulse energies 25–35 nJ per pulse, laser focus diameter 200 μm (both at the target) and laser repetition rate 200 kHz. A typical pulse spectrum is shown in Fig. 2b. For the 2DES experiments, the cluster beam was excited by a collinear pulse train of 4 fs laser pulses and the fluorescence is detected as parametric function of the inter-pulse delay τ, t. A 2D Fourier transform with respect to τ, t yields the rephasing and non-rephasing 2D spectra of which the absorptive correlation functions decay to 5% of its amplitude along each dimension. Note, that the resolution along the diagonal (anti-diagonal) dimension is higher. In case of the H2Pc–Ne3 sample, data apodisation was not necessary and the inhomogeneous (homogeneous) linewidths of the sample were determined without spectral broadening introduced by the experimental apparatus.

Data availability

The data that support the findings of this study are available from the corresponding author upon request.

Code availability

The code used to process the data of this study is available from the corresponding author upon request.

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Competing interests
The authors declare no competing interests.

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