Spatially separating the conformers of the dipeptide Ac-Phe-Cys-NH₂

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Atomic-resolution-imaging approaches for single molecules, such as coherent x-ray diffraction at free-electron lasers, require the delivery of high-density beams of identical molecules. However, even very cold beams of biomolecules typically have multiple conformational states populated. We demonstrate the production of very cold (Trot ∼ 2.3 K) molecular beams of intact dipeptide molecules, which we then spatially separate into the individual populated conformational states. This is achieved using the combination of supersonic expansion laser-desorption vaporisation with electrostatic deflection in strong inhomogeneous fields. This represents the first demonstration of a conformer-separated and rotationally-cold molecular beam of a peptide, and will enable future single biomolecule x-ray diffraction measurements.

To investigate biomolecules in the gas-phase requires their vapourisation without fragmentation or ionisation. Laser desorption (LD) has been demonstrated as a technique to vapourise such thermally labile molecules and the combination with supersonic expansion allows for rapid cooling of the desorbed molecules. However, even in such cold molecular beams different conformers, which differ by rotations about single bonds, can coexist. In order to produce a pure beam containing only a single conformer, we combine LD with electrostatic deflection. This allows the spatial separation of molecular species based on their distinct interaction with the applied electric field. This so-called Stark effect is dependent on the quantum-state-specific effective dipole moment and this technique has been demonstrated to spatially separate conformers of small aromatic molecules and for very small molecules it can even produce single-quantum-state samples. Furthermore, due to the rotational-state-dependence of the Stark effect deflection allows the creation of very cold (Trot < 100 mK) molecular ensembles. This can significantly improve the degrees of laser alignment and mixed-field orientation of molecules in space and thus enable ensemble-averaged single-molecule imaging.

Here, we present the first combination of laser desorbed biomolecules with electrostatic deflection and demonstrate the spatial separation of the two main conformers of the dipeptide Ac-Phe-Cys-NH₂, shown in Figure 1a. These two conformers differ in their hydrogen-bonding interactions and, hence, 3D structure. One conformer, indicated by red colour throughout the paper, forms a hydrogen bond from the SH group to the oxygen on the carboxamide group, while the other conformer, blue colour, forms a hydrogen bond from the SH to the delocalised π-system. These two "beautiful molecules" have been previously identified using vibrational and electronic spectroscopy. In a cold molecular beam these two conformers cannot interconvert, however, their significantly

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different dipole moments of 3.2 D and 8.1 D result in different Stark interactions, see Figure 1 b. This allows for their spatial separation with the electrostatic deflector if a sufficiently cold molecular ensemble can be created. This would, furthermore, also separate the sample of interest from unwanted fragments or contaminants present in the beam, such as carbon clusters from the LD process. Compared to the separation of molecular ions in ion mobility measurements, our method enables the separation of neutral species, avoiding space-charge density limitations that severely affect diffractive-imaging experiments. Furthermore, the low temperatures of the generated molecular ensembles allow for strongly fixing the molecules in space — two prerequisites for the recording of atomically resolved molecular movies.

RESULTS AND DISCUSSION

Our implementation of the combination of LD with electrostatic deflection is shown schematically in Figure 2. Details are given in Methods. Briefly, the laser-desorbed molecular beam enters a 15 cm long deflector sustaining electric field strengths on the order of 150 kV/cm. The different conformers experience a different vertical deflection within this field, which originates from the Stark-effect interaction between the molecules’ space-fixed dipole moment \( \mu_{\text{eff}} \) and the applied electric field \( \epsilon \). This leads to a force \( F = -\mu_{\text{eff}}(\epsilon) \cdot \nabla \epsilon \) acting on the molecules. Thus, the observed deflection depends on the effective-dipole-moment-to-mass ratio and the two conformers experience different forces, i.e., transverse accelerations, in the electric field, leading to their spatial separation. The molecular beam and the separation of conformers was characterised by recording spatial profiles of the beam. This was achieved by vertically translating the ionisation laser beam through the horizontal molecular beam, and recording the relative density as a function of laser height. The ionisation laser was tuned to specific resonances to selectively detect a single conformer.

Such spatial molecular beam profiles for the individual conformers in the absence of an electric field, i.e., with the deflector at 0 kV, are shown in Figure 3 a, to which all beam-profile intensities have been normalised. These show that both conformers are centered around \( y = 0 \) mm and exhibit the same spatial distribution. The measured width of the molecular beam is predominately defined by the apertures of skimmers and the electrostatic deflector placed in the molecular beam, see Methods. The relative population of the two conformers in the beam was assessed by placing the ionisation laser focus at the center of the profile, as indicated by the black arrow in Figure 3 a, and scanning the ionisation wavelength across the electronic-origin transitions of the two conformers around 37325 cm\(^{-1}\) and 37450 cm\(^{-1}\), respectively. The resulting resonance-enhanced multiphoton-ionisation (REMPI) spectrum is shown in Figure 3 d and yielded an intensity ratio of \( \sim 2 : 1 \) for the SH–O and SH–π bound conformers, respectively. Assuming identical ionisation probabilities for the REMPI process, this ratio can be taken as a measure of the relative conformer populations in the molecular beam.

Charging of the electrostatic deflector lead to deflection of the molecular beam in the positive, upward direction, as shown in Figure 3 b, c. Application of 4 kV to the deflector, Figure 3 d, lead to a clear shift of both spatial profiles, with the more polar SH–π-bound conformer shifting significantly more. This created an area, between \( \sim 1.7–2.5 \) mm, were a highly enriched sample of this conformer was obtained, as confirmed by the REMPI
FIG. 3. Spatial molecular beam profiles (a–c) and corresponding REMPI spectra (d–f) for the two main conformers of Ac-Phe-Cys-NH₂. These are collected at deflector voltages of 0 kV (a,d), 4 kV (b,e) and 10 kV (c,f). Solid lines in the deflection profile plots (a–c) are taken from quantum-state resolved trajectory simulations with a 2.3 K thermal state weighting. REMPI spectra are taken at the spatial position indicated by the black arrow in the spatial profiles.

A REMPI spectrum collected at position \( y = 1.75 \text{ mm} \) and shown in Figure 3c. To separate and create a pure sample of the SH–O-bound conformer, a voltage of 10 kV was applied, leading to depletion of the SH–\( \pi \)-bound system from the interaction region, as shown in Figure 3c. This is due to the large deflection experienced by this more polar conformer, such that these molecules collided with the deflector or following apertures and no clear beam was observable anymore. Instead, a position-independent small background signal was present. A REMPI spectrum recorded in the deflected beam is shown in Figure 3f, confirming the highly-enriched sample of the SH–O-bound conformer created under these conditions.

Using a calibrated ion detector, we estimated the number of ions produced per laser shot to be \( \sim 1 \) for REMPI ionisation. By using more efficient strong-field ionisation (SFI) we extracted a lower limit for the absolute number density of \( 10^7 \text{ cm}^{-3} \), see appendix A for details. Derivation of this density assumes an ionisation efficiency of 1 for SFI and only takes into account the major assigned fragmentation channels for Ac-Phe-Cys-NH₂ and thus strictly represent a lower limit of the density.

Further to the deflection of the molecular beam, we observed a significant broadening of the spatial profiles. This is due to the dispersion of the different rotational states in the electric field, arising from the rotational-state-dependence of the Stark effect. This is shown in Figure 1b for \( J = 0 \ldots 3 \) states, indicating the larger effective dipole moment of lower-lying rotational states, leading to these states being deflected more, and hence the creation of a rotationally colder sample in the deflected beam. To extract approximate rotational temperatures and quantum-state distributions in the deflected beam, we have simulated particle trajectories through our setup for the different populated rotational states details are given in appendix B. Resulting simulated deflection profiles are shown as solid lines in Figure 3a–c which were obtained by applying a thermal-distribution weighting to the individual-state simulations, corresponding to the rotational temperature distribution from our LD-molecular-beam source. We extract an approximate rotational temperature of \( 2.3 \pm 0.5 \text{ K} \) for the laser-desorbed molecular beam.

Furthermore, we extract the quantum-state distribution within the deflected beam in Figure 3d. These are shown in Figure 4 and indicate that the deflector creates a significantly colder ensemble. While this has a non-thermal rotational state distribution, the highest rotational states populated are approximately corresponding to a 1.5 K distribution. Even colder ensembles can be probed by moving the interaction region further into the deflected beam, this is indicated by the magenta and cyan distributions in Figure 4 evaluated at position 2.2 mm in the deflected beam, which are comparable to a 1.0 K average.

These results highlight the quantum-state-sensitivity of the electrostatic-deflection technique, allowing us to control conformer populations and rotational state distri-
CONCLUSION

We demonstrated the combination of laser desorption for the vaporisation of labile biological molecules with the electric deflector for the spatial separation of conformational states and the creation of pure and rotationally-cold samples of individual conformers. Using the prototypical (d)peptide Ac-Phe-Cys-NH$_2$ as a model system, we showed that its two conformers, in the gas-phase, can be spatially separated and samples of either conformer can be obtained. The measured deflection was quantitatively understood using trajectory calculations, which furthermore allowed us to assign a rotational temperature of $2.3 \pm 0.5$ K for the beam from our laser desorption source. The generally good agreement between experiment and simulation also confirms the calculated dipole moments and that Stark effect calculations based on the rigid-rotor approximation are sufficient even for these large systems.

The created molecular samples will enable novel x-ray diffractive imaging experiments: they are conformer-pure beams that are well-separated from carrier gas and rotationally cold enough for strong laser alignment and orientation. The achieved densities of around $10^7$ cm$^{-3}$ are sufficient for high-resolution diffraction experiments at free-electron laser sources such as the European XFEL, which will deliver up to 26,000 pulses per second, allowing fast collection of data. This enables the collection of a diffraction image within 1 h and simulated aligned-molecule diffraction patterns for the two conformers, showing marked differences, are shown in appendix C. Our laser desorption source, with its low overall repetition rate, but reasonably long gas pulses of 100s of µs is well-suited to the pulse-train structure of superconducting-LINAC-based XFELs. The produced rotationally cold samples are well suited to strong-field alignment, which can be achieved using the available in-house laser systems available at FELs.

Our developed technique will more generally enable experiments on conformer-selected biological molecules with inherently non-species-specific experimental techniques, such as (sub-)femtosecond dynamics, reactive collision studies or diffractive imaging. This will open new pathways to study the intrinsic structure-function relationship of these basic molecular building blocks of the complex biochemical machinery.

METHODS

A laser desorption source, described in detail elsewhere, is used to vapourise the dipeptide Ac-Phe-Cys-NH$_2$ (APCN, 95% purity, antibodies-online GmbH), which is used without further purification. The resulting cold supersonic molecular beam is skimmed twice before entering the strong inhomogeneous field of the electrostatic deflector: once by a 2 mm skimmer (Beam Dynamics Inc. Model 50.8) 75 mm downstream of the expansion, and again by a 1 mm skimmer (Beam Dynamics Inc. Model 2) 409 mm downstream of the expansion. Within the strong inhomogeneous electric field of the deflector, molecules are dispersed according to their effective dipole moment-to-mass-ratio. The molecular beam is skimmed once more with a 1.5 mm skimmer (Beam Dynamics Inc. Model 2) prior to entering the interaction region. This skimmer can be translated.
in height to ensure no part of the molecular beam is cut off. During measurements, data is collected for two skimmer positions and subsequently combined by keeping the highest intensity measured. The relative density of the conformers is probed via resonance-enhanced multi-photon ionisation (REMPI). The ultraviolet probe light is produced by frequency doubling the output of a dye laser (Radiant Dyes NarrowScan, using Coumarin 153 dye in methanol), pumped by the third harmonic of a Nd:YAG laser (Innolas, SpitLight 600). Typical laser-pulse energies were around 19 µJ loosely focused to a 100 µm spot in the interaction region.

The structures and dipole moments of Ac-Phe-Cys-NH$_2$ were calculated using the GAMESS software suite using the B3LYP functional with a 6-311(p) basis set and confirmed against published structures.

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Appendix A: Determination of a lower limit of the number density

For the density determination a time-of-flight mass spectrum using strong-field ionisation by femtosecond laser pulses (800 nm central wavelength, 40 fs duration, typical pulse energies of 100 µJ) is recorded in the time-of-flight mass spectrum all peaks originating from the Ac-Phe-Cys-NH$_2$ molecule are integrated and the total ion current on the detector determined. This is compared to the known calibrated current for a single ion hit, which leads to approximately 18 ions/shot in the $\omega_0 = 50$ µm focus of the laser. Assuming an ionisation efficiency of 1 for strong-field ionisation and a molecular-beam width of 1 mm, this yields a density of $9 \times 10^6$ cm$^3$.

Appendix B: Numerical simulations and temperature determination

Spatial molecular-beam profiles were simulated by first calculating the Stark energies for each conformer for all rotational states up to $J = 70$, including all states up to $J = 70$ in the calculation, using the freely available CMIstark software package. Rotational constants and dipole moment vectors were taken from the DFT calculations and are summarized in Table A1. Subsequently, for molecules in each quantum state we carried out classical trajectory simulations through the experimental setup, taking into account apertures and applying the appropriate forces when molecules are within the electrostatic deflector. Finally, histograms of the final particle-position densities were determined at the interaction point and the contributions from each quantum state weighted by a Maxwell-Boltzmann distribution for a given initial temperature. Simulated intensities for given conditions – species and deflector voltage – were scaled with a single amplitude-scaling factor to compare with experimental data to account for additional losses and detection efficiency in the setup. The temperature that best described the experimental observations was determined by comparing the combined residuals, that is the absolute deviation between simulation and data, from all deflected data sets, excluding the SH—π conformer at 10 kV where only a constant background was observed. The sum of these residuals (black line) yielded a rotational temperature of 2.3 K.

| Species     | SH—O isomer | SH—π isomer |
|-------------|-------------|-------------|
| A (MHz)     | 340.181593  | 345.067516  |
| B (MHz)     | 203.443113  | 215.965933  |
| C (MHz)     | 159.877010  | 175.580323  |
| $\mu_A$ (D) | 0.768       | 6.789       |
| $\mu_B$ (D) | 2.406       | -2.701      |
| $\mu_C$ (D) | 1.975       | 3.406       |

TABLE A1. Rotational constants and dipole moment vectors used for calculating the Stark effect.
constant low background was observed experimentally, for different rotational temperatures. These are shown in Figure A5 and from the combined residuals (black trace) a rotational temperature of 2.3 K for our molecular beam was extracted. Since the minima for individual deflection profiles deviate by ~0.5 K, conservative error bounds for the rotational temperature are ±0.5 K.

Appendix C: Simulated X-Ray diffraction patterns

Simulated x-ray diffraction patterns at 9.5 keV photon energy achievable at current XFEL sources such as LCLS and the European XFEL, for the two conformers of Ac-Phe-Cys-NH$_2$ are shown in Figure A6. The simulation assumes a detector distance of 80 mm and maximum scattering angle on the detector of 50.2°, corresponding to a resolution of d ≈ 154 pm at the edge. These calculations assume perfectly aligned molecules and no contribution from background gas in the interaction region, with the most-polarizable axis of the molecules aligned vertically and the second-most-polarizable axis aligned horizontally within the image plane, as it would be obtained in a typical aligned-molecule-diffraction experiment. A clear, detectable difference between the two patterns is visible. Utilizing the 10 Hz bunch-structure of the upcoming European XFEL will enable the recording of such patterns as well as the reaction path of conformer interconversion.

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