Conformer Selection by Matter-Wave Interference

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We establish that matter-wave diffraction at near-resonant ultraviolet optical gratings can be used to spatially separate individual conformers of complex molecules. Our calculations show that the conformational purity of the prepared beam can be close to 100% and that all molecules remain in their electronic ground state. The proposed technique is independent of the dipole moment and the spin of the molecule and thus paves the way for structure-sensitive experiments with hydrocarbons and biomolecules, such as neurotransmitters and hormones, which have evaded conformer-pure isolation so far.

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Introduction.—The conformation of a molecule can have a strong influence on its chemical reaction rates. This was demonstrated for a number of compounds [1–4], where the rate constants of the conformer-specific reactions varied by a factor of at least 2 even though the conformers differed only in the orientation of a single bond. To explore such conformer-specific traits, it is desirable to develop methods to separate conformers with high efficiency from an initially unsorted molecular ensemble.

Several conformer selection techniques have been developed in recent years. Charged molecules can be separated in collision cells using ion-mobility spectroscopy [5–7]. The separation of neutral molecules in high vacuum has been achieved using the Stark effect [8–10]. This method also allows the isolation of spin isomers [11] and clusters with polar [12] and nonpolar [13] particles in specific stoichiometries. However, separation due to the Stark effect requires that the relevant conformers differ substantially in their rotational spectrum [14] or their electric dipole moment [12,13]. Any method that can overcome these restrictions will be instrumental for subsequent slowing [15–20], trapping [21,22], or collision experiments [23–26], for high-resolution spectroscopy [27–29] and reaction studies [1].

Conformer-dependent reactions of hydrocarbons and aromatic radicals can strongly affect atmospheric and astrochemical processes, such as the oxidation of aromatics [30], the formation of smog [31,32], and the chemistry on Saturn’s moon Titan [33,34]. Beams of size- and conformer-selected water clusters may give new insights into astrochemical processes [35], atmospheric, and environmental reactions [36,37]. Finally, small biomolecules, such as neurotransmitters and amino acids, offer rich conformational spaces [38–40]. Experimental studies in the gas phase can shed new light on conformational preferences [41,42], isomerization barriers [43,44], and the influence of solvent molecules in hydrated beams [45,46].

Here, we propose to use matter-wave diffraction at a tunable standing light wave grating for separating different conformers with high purity and high efficiency. Our scheme exploits that most conformers exhibit spectroscopically well-separated electronic transitions in the ultraviolet, even if they are structurally similar and have comparable dipole moments. When the laser wavelength is resonant with the electronic transition of one specific conformer, the standing light wave will realize both an absorptive and a phase grating for the matter wave [47,48]. All other conformers will only be subjected to a pure phase grating [49,50] which can suppress certain diffraction orders depending on the laser wavelength and energy. By balancing these effects a specific diffraction order can be preferentially populated by a selected conformer. One may then select this conformer with high purity by spatially filtering the molecular interference patterns. This method can address a wide range of biomolecules, radicals, hydrocarbons, and their water clusters (Supplemental Material [51]).

 Proposed setup.—In our proposed setup (see Fig. 1) molecules are entrained in a pulsed supersonic expansion to prepare a rovibrationally cold ensemble with a fast but narrow velocity distribution. Several hundred rotational levels will still be occupied due to their small energy spacing. The molecules pass a slit skimmer $S_1$ and are diffracted at the source skimmer $S_2$ in accordance with Heisenberg’s uncertainty principle [76]. After the distance $L_1 = 1$ m the molecular wave packet of mass $M$ traveling with velocity $v_z$ has a transverse coherence width $4\pi\hbar L_1/Mv_z$, sufficiently large to illuminate several antinodes of a retro-reflected standing light wave. This optical grating with a period of $\lambda_L/2$ can be generated by a narrow-band pulsed UV laser of wavelength $\lambda_L$, tuned with a linewidth of better than 1 pm. State-of-the-art frequency-doubled dye lasers meet these requirements and deliver sufficient energy per pulse to diffract the molecules. At the grating the molecules are diffracted according to their
de Broglie wavelength \( \lambda_{\text{db}} = 2\pi \hbar / M v \). The required alignment of the grating mirror with respect to the molecular beam is determined by the pulse length of the laser and the collimation of the molecules. After traversing the grating, the molecular wave propagates the distance \( L_2 \) before it impinges onto a mask with two adjustable slits. These select, for instance, both first order diffraction peaks with a diffraction angle of \( 2\lambda_{\text{db}} / \lambda_L \).

Matter-wave diffraction at optical gratings has been realized in continuous and pulsed interferometers for atoms [77,78], electrons [79], and molecules [50,80,81]. The phase grating transfers an integer multiple of the grating momentum \( 4\pi \hbar / \lambda_L \) onto the traversing, rotating molecule. In addition, the molecule may absorb one or more photons depending on the laser wavelength, intensity, molecular absorption cross section, and interaction time. If only a single photon is absorbed, the coherent diffraction signal will be shifted in momentum by \( \pm 2\pi \hbar / \lambda_L \) and filtered out by the mask, provided that the diffraction peaks are sufficiently separated. Upon absorption of two or more photons, the molecule is ionized and removed from the beam.

**Molecule-laser interaction.**—The force acting on a polarizable molecule in the laser field is determined by the optical susceptibility, which depends on the laser wavelength \( \lambda_L \) and the rotational state \( r \). The real part of the susceptibility is the polarizability \( \alpha_r(\lambda_L) \) while the imaginary part \( \epsilon_0 \lambda_L \sigma_r(\lambda_L) / 2\pi \) depends on the total absorption cross section \( \sigma_r(\lambda_L) \).

The effect of the pulsed standing light wave on the transverse motional state of the molecule can be described by the phase shift \( \phi_r(\lambda_L, E_L) \) and the mean number of absorbed photons \( n_r(\lambda_L, E_L) \) at the antinodes as a function of \( \lambda_L \) and the pulse energy \( E_L \). The grating transit of a molecule in a rotational state \( r \) is then characterized by the state-dependent grating transformation [82,83]

\[
t_r(x) = \exp \left[ \left( i\phi_r - \frac{n_r}{2} \right) \cos \left( \frac{2\pi x}{\lambda_L} \right) \right].
\]

The phase and the mean photon number can be related to the real and imaginary part of the susceptibility,

\[
\phi_r(x, \lambda_L, E_L) = \frac{2\alpha_r(\lambda_L) E_L}{\hbar c \epsilon_0 \alpha_L},
\]

and

\[
n_r(x, \lambda_L, E_L) = \frac{2\epsilon_r(\lambda_L) E_L \alpha_r(\lambda_L)}{\pi \hbar c \epsilon_0 \alpha_L},
\]

with the spot size \( \alpha_L \).

**Interference pattern.**—The transverse wave function of a molecule’s center-of-mass motion, starting at position \( x_0 \) with velocity \( v_z \) in the rotational state \( r \) can be evaluated in the paraxial approximation as [83,84]

\[
\Psi_r(x, x_0) \propto \int_{-S_{1/2}}^{S_{1/2}} dx' t_r(x') \exp \left[ \frac{i M v_z}{2\hbar} \left( \frac{1}{L_1} + \frac{1}{L_2} \right) x'^2 \right] \\
\times \exp \left[ - \frac{i M v_z}{\hbar} \left( \frac{x_0}{L_1} + \frac{x}{L_2} \right) x' \right].
\]

Here, \( S_3 \) is the total width of the grating as determined by the third skimmer. Averaging over all source points and rotation states with population \( p_r \) then yields the molecular interference pattern

\[
S(x) \propto \int_{-S_{1/2}}^{S_{1/2}} dx_0 \sum_r p_r |\Psi_r(x, x_0)|^2.
\]

Interferometric conformer selection can be applied to all molecules with spectroscopically separated transitions in the visible or ultraviolet. This can be seen by considering the approximate intensity \( S_n \) of the \( n \)th diffraction order in the far field, i.e., for \( M v_z (L_1 + L_2)^2 S_{3/2}^2 / 2 \hbar L_1 L_2 \ll 1 \),

\[
S_n(\lambda_L, E_L) \propto \sum_r p_r e^{-n_r/2} J_n \left( \frac{\phi_r}{2} + i \frac{n_r}{4} \right)^2,
\]

where \( J_n(\cdots) \) are Bessel functions. Spectroscopic separability of the conformers implies that the wavelength \( \lambda_L \) can be tuned in such a way that the polarizability of any chosen conformer exceeds all others substantially. By adjusting the pulse energy sufficiently low so that the Bessel function in Eq. (6) vanishes for all but the selected
species, the first diffraction order will be populated predominantly by this conformer. This demonstrates the general applicability of the proposed method, even though there can be regions of $\lambda_L$ and $E_L$ for which interferometric conformer selection works with even higher efficiency.

**Conformer-selective interference of PEA.**—To illustrate the method, we study the neurotransmitter 2-phenylethylamine (PEA), see Fig. 2. In jet experiments it exhibits four conformers, which are spectroscopically well separated [85,86]. They differ in the conformation of the C–N bond and the lone pair of the NH$_2$ group. While the former is either in anti or gauche position to the C$_6$–C$_7$ bond, the latter points up or out with respect to the chromophore. Since PEA has only a single polar group, the dipole moments of all four observed conformers are predicted to be virtually identical (1.25 ± 0.05 D) [87,88]. Stark separation is thus practically impossible [13]. The susceptibility of the four different conformers is calculated from data compiled in the Supplemental Material [51]. The complex optical susceptibility of each conformer can be described by a Lorentz oscillator model [89,90] where the transition frequencies of the rotational states are calculated from the conformer-dependent rotational constants [86,91]. The mean static polarizability $\alpha_0/4\pi\varepsilon_0 = 14.6$ Å$^3$ was calculated using density functional theory and agrees well with the value obtained from the refractive index in solution [92]. The maximum deviation between different conformers is at 0.2 Å$^3$ [51] and negligible compared to the optical polarizability which changes by tens of Å$^3$ over the considered wavelength region. The rotational substructure of the electronic transition consists of several hundred individual peaks even at a rotational temperature of 3 K. To account for this, we divide the rotational spectrum into 20 intervals and calculate for each bin its mean spectroscopic weight $p_r$.

**Conformer-selection efficiency.**—In Fig. 3(a) we show the interference patterns of the four conformers diffracted at a grating with $\lambda_L = 265.8$ nm, $E_L = 0.67$ mJ, and $A_L = 1$ mm$^2$. The grating wavelength lies between the electronic transitions of the anti(up) and the gauche(out) conformers. Hence, the respective values of $\alpha_r(\lambda_L)$ deviate strongly from the static polarizability, while they are close to that value for the gauche(up) and anti(out) conformers. At the chosen parameters the intensity in the first diffraction order is maximal for the gauche(out) conformer comprising 35% of the total population of this conformer.

From the interference patterns we calculated the **conformer-selection efficiency** $\eta$ in the first diffraction order, that is the fraction of one conformer compared to the sum of all four. For the patterns shown in Fig. 3(a) $\eta$ reaches a value of >93%, illustrating that high-purity conformer-selection is possible with this method. To prove the feasibility of a clear separation for all molecular conformers, we have extended the simulations to the wavelength region between 265 and 267 nm and to a surface energy density $E_L/A_L$ up to 1.1 mJ/mm$^2$. For each simulated pattern we calculated $\eta$ and compiled the results in Fig. 3(b) for the four conformers. Here we color the plot only when an efficiency of 50% is exceeded; areas of lower selectivity are left blank. It shows a rich pattern due to the strong wavelength-dependence of $\sigma_r(\lambda_L)$ and $\alpha_r(\lambda_L)$ near the resonances. The size of the parameter space which leads to selectivity depends strongly on the relative population of the conformer in the beam. For the gauche(out) conformer a selectivity greater than 50% is observed in large parts of the parameter space. However, even for the weakly populated anti(out) conformer $\eta$ exceeds 80% over a range of 0.03 nm which is experimentally easily accessible. It demonstrates that every single conformer of PEA can be selected with high conformer selectivity and that specific conformers can be addressed by tuning the laser wavelength and power. Since the details of the selection efficiency shown in Fig. 3(b) are highly sensitive on the oscillator strength $f$, measuring the conformer-selection efficiency $\eta$ offers a new way to probe it for all conformers.

**Experimental feasibility.**—The separation between neighboring resonances of the polarizability spectrum is on the order of 0.1 nm, and therefore well resolved by a dye laser with a laser linewidth of 1 pm, even in the presence of rotational broadening of the spectrum. For medium-sized molecules such as PEA, the rotational energy spread at 3 K is roughly 0.02 nm [86]—much smaller than the separation between the electronic resonances of the neighboring conformers. In a realistic experimental situation a certain fraction of the molecular beam will not interact with the pulsed laser beam and pass on to the detector in the blocked zeroth diffraction order. Furthermore, as the carrier gas is much lighter than the molecules, it experiences larger diffraction angles than the analyte molecule. Hence, the diffraction orders do not overlap and the proposed method is virtually background-free. This is comparable to Stark deflection where polar conformers are deflected out of the initial beam [9].
The selectivity of our proposed method depends on the power stability of the laser system. When operated in saturation the fluctuation of a dye laser depends solely on the performance of the diode-pumped solid state laser, which can reach <3%. After frequency doubling this leads to ±6% stability in power. However, even for power variations of ±15% the simulations predict values of $\eta$ between 70% for the anti(out) and 95% for the gauche(out) conformer [51].

The flux for a specific conformer behind the selection slits can be estimated using the vapor pressure of PEA at 410 K [93], the characteristics of a pulsed Even-Lavie valve [94,95], a 100 Hz laser illuminating an area of 1 mm², and the diffraction pattern of the gauche(out) conformer shown in Fig. 3(a). Assuming that 78% of the molecules are in their vibrational ground state and a mean number of absorbed photons of $<10^{-4}$, we expect a mean flux of $1.3 \times 10^8$ cm⁻² s⁻¹ [51]. The peak density reaches $1.3 \times 10^8$ cm⁻³ which is comparable to densities used in x-ray diffraction at free electron lasers [96] and in crossed beam studies [26]. The mean flux can be increased by a factor of 100 using a 10 kHz laser illuminating the same area.

Since the presented matter-wave assisted separation technique requires the conformers only to have sharp and separated transitions in the ultraviolet, it can be used for a wide range of species and applications. This includes large families of molecular systems such as hydrocarbons, small biomolecules, and aromatic radicals [39,97,98] (Supplemental Material [51]). It can provide conformational samples for x-ray diffraction [99], Coulomb explosion studies [100,101], merged beam experiments [26,102], and collisions with trapped neutral particles [23] or cold ions [1,24,25]. The selected structure can be varied during the experiment by changing the wavelength and the pulse energy of the grating laser. This way several conformers can be compared within one experimental run. The combination with molecular cooling schemes might also prepare high-resolution spectroscopy [27–29].

Conclusion.—We have presented a robust scheme to select molecular conformers in the vibrational ground state with purities of up to 95%. The method requires an intense and tightly collimated molecular beam giving rise to matter-wave diffraction at a tunable laser grating. It is applicable to large families of conformers that can be distinguished by individual and sharp electronic transitions in the UV. Being independent of internal dipole moments or the spin state, it can be applied to nonpolar molecules, radicals, and their clusters alike. Furthermore, this technique eliminates most of the vibrationally excited molecules from the region of interest, selecting colder molecules in the beam.

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\[ \lambda_L = 265.8 \text{ nm} \]

\[ E_L/A_L = 0.67 \text{ mJ/mm}^2 \]

\[ \eta > 93\% \]

\[ \text{Performance of the diode-pumped solid state laser, saturation the fluctuation of a dye laser depends solely on power stability of the laser system.} \]

\[ \text{The mean flux can be increased by a factor of 100 using a 10 kHz laser illuminating the same area.} \]

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