

Exploring light-driven molecular swing

October 18 2022



Calculated coherence and energy transfer ratios. Calculated vibrational molecular coherence in the symmetric stretching vibrational mode of the DMSO₂ molecule in solution displayed in a frame rotating with the vibrational eigenfrequency. The calculations are done with **a** the compressed pulse using the RWA, **b** the compressed pulse without the RWA, **c** the chirped pulse using the RWA, and **d** the chirped pulse without the RWA. The red curves include relaxation; the blue curves do not include relaxation. Each bump of the cycloid in **b** corresponds to one half-cycle of the electric field, showing that energy transfer from the excitation field to the molecular system is completed after 3 to 4 field cycles in the case of the FCE. In **d**, the time points that are maxima of CET(*t*)



caused by the symmetric stretching vibration are marked with black dots. **e** Reemitted (orange) and maximum absorbed (green) fraction of the impinging pulse energy versus concentration. The results were obtained from the impulsiveregime model (see Supplementary Information) with the FCE and ab initio Lorentz parameters. The solid lines include direct interactions with the surrounding water and the screening effect of the polarizable continuum, the dashed lines only the latter, and the dotted lines neither. For small concentrations, the maximum absorbed energy scales linearly with concentration. In contrast, the coherent re-emission, containing the spectroscopic information, scales quadratically with concentration. Credit: *Nature Communications* (2022). DOI: 10.1038/s41467-022-33477-5

When light impinges on molecules, it is absorbed and re-emitted. Advances in ultrafast laser technology have steadily improved the level of detail in studies of such light-matter interactions.

FRS, a laser spectroscopy method in which the electric field of laser pulses repeating millions of times per second is recorded with <u>time</u> resolution after passing through the sample, now provides even deeper insights: Scientists led by Prof. Dr. Regina de Vivie-Riedle (LMU/Department of Chemistry) and PD Dr. Ioachim Pupeza (LMU/Department of Physics, MPQ) show for the first time in theory and experiment how molecules gradually absorb the energy of the ultrashort light pulse in each individual optical cycle, and then release it again over a longer period of time, thereby converting it into spectroscopically meaningful light.

The study elucidates the mechanisms that fundamentally determine this <u>energy transfer</u>. It also develops and verifies a detailed quantum chemical model that can be used in the future to quantitatively predict even the smallest deviations from linear behavior.



A child on a swing sets it in motion with tilting movements of the body, which must be synchronized with the swing movement. This gradually adds energy to the swing, so that the deflection of the swing increases over time. Something similar happens when the alternating electromagnetic field of a short laser pulse interacts with a molecule, only about 100 trillion times faster: When the alternating field is synchronized with the vibrations between the atoms of the molecule, these vibration modes absorb more and more energy from the <u>light pulse</u>, and the vibration amplitude increases.

When the exciting field oscillations are over, the molecule continues to vibrate for a while—just like a swing after the person stops the tilting movements. Like an antenna, the slightly electrically charged atoms in motion then radiate a light field. Here, the frequency of the light field oscillation is determined by properties of the molecule such as atomic masses and bond strengths, which allows for an identification of the molecule.

Researchers from the attoworld team at MPQ and LMU, in collaboration with LMU researchers from the Department of Chemistry (Division of Theoretical Femtochemistry), have now distinguished these two constituent parts of the light field—on the one hand, the exciting light pulses, and on the other, the decaying light field oscillations—using timeresolved spectroscopy. In doing so, they investigated the behavior of organic molecules dissolved in water.

"While established laser spectroscopy methods usually only measure the spectrum and thus do not allow any information about the temporal distribution of the energy, our method can precisely track how the molecule absorbs a little more energy with each subsequent oscillation of the light field," says Ioachim Pupeza, head of the experiment.

That the measurement method allows this temporal distinction is best



illustrated by the fact that the scientists repeated the experiment, changing the duration of the exciting pulse but without changing its spectrum. This makes a big difference for the dynamic energy transfer between light and the vibrating molecule: Depending on the temporal structure of the laser pulse, the molecule can then absorb and release energy several times during the excitation.

In order to understand exactly which contributions are decisive for the energy transfer, the researchers have developed a supercomputer-based quantum chemical model. This can explain the results of the measurements without the aid of measured values. "This allows us to artificially switch off individual effects such as the collisions of the vibrating molecules with their environment, or even the dielectric properties of the environment, and thus elucidate their influence on the energy transfer," explains Martin Peschel, one of the first authors of the study.

In the end, the energy re-emitted during the decaying light field oscillations is decisive for how much information can be obtained from a spectroscopic measurement. The work thus makes a valuable contribution to better understanding the efficiency of optical spectroscopies, for example with regard to molecular compositions of fluids or gases, with the objective of improving it further and further.

The research is published in *Nature Communications*.

More information: Martin T. Peschel et al, Sub-optical-cycle lightmatter energy transfer in molecular vibrational spectroscopy, *Nature Communications* (2022). DOI: 10.1038/s41467-022-33477-5

Provided by Ludwig Maximilian University of Munich



Citation: Exploring light-driven molecular swing (2022, October 18) retrieved 23 July 2024 from <u>https://phys.org/news/2022-10-exploring-light-driven-molecular.html</u>

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