

Physicists edge closer to controlling chemical reactions

December 11 2018



Control valve. Credit: MIPT

A team of researchers from the Moscow Institute of Physics and Technology, and Aarhus University in Denmark has developed an algorithm for predicting the effect of an external electromagnetic field on the state of complex molecules. The algorithm, which is based on a theory developed earlier by the same team, predicts tunneling ionization rates of molecules. This refers to the probability that an electron will bypass the potential barrier and escape from its parent molecule. The new algorithm, presented in a paper in the *Journal of Chemical Physics*,



enables researchers to look inside large polyatomic molecules, observe and potentially control electron motion therein.

Physicists use powerful lasers to reveal the electron structure of <u>molecules</u>. To do this, they illuminate a molecule and analyze its reemission spectra and the products of the interaction between the molecule and the electromagnetic field of the laser pulse. These products are the photons, electrons, and ions produced when the molecule is ionized or dissociates (breaks up).

Previous research involving MIPT's theoretical attosecond physics group led by Oleg Tolstikhin showed that besides elucidating the electronic structure of a molecule, the same approach may enable physicists to control the electron motions in the molecule with attosecond precision. An attosecond, or a billionth of a billionth of a second, is the time it takes <u>laser light</u> to travel the distance comparable to the size of a small molecule.

"If you place a molecule in a field of powerful laser radiation, <u>ionization</u> occurs: An electron escapes the molecule," explains Andrey Dnestryan, a member of the theoretical attosecond physics group at MIPT. "The motion of the electron is then affected by the variable laser field. At some point, it may return to the parent molecular ion. The possible outcomes of their interaction are rescattering, recombination, and dissociation of the molecule. By observing these processes, we can reconstruct the motions of electrons and nuclei in molecules, which is of profound interest to modern physics."





The orientation of the naphthalene molecule relative to the external electric field can be described by angles β and γ in the following manner: Electric field F is directed along axis z?, while β denotes the angle between z? and the molecular axis z, and γ is the rotation angle around axis z. The latter angle specifies an arbitrary orientation of the molecule relative to the field F. The two angles β and γ are known as Euler angles. The figure also shows two outer orbitals (a and b) of the naphthalene molecule -- that is, the areas where the two outer electrons are localized in this molecule. The outer electrons are the first to undergo ionization in the presence of an electric field. Credit: Moscow Institute of Physics and Technology

The interest in <u>tunneling</u> ionization stems from its role in experiments observing electronic and nuclear motion in molecules with attosecond time resolution. For example, tunneling ionization may enable



researchers to track the motions of electrons and holes—positively charged empty spots resulting from the absence of electrons—along the molecule. This opens up prospects for controlling their motion, which would help control the outcomes of chemical reactions in medicine, molecular biology, and other areas of science and technology. Precise calculations of tunneling ionization rates are vital to these experiments.

The tunneling ionization rate could be interpreted as the probability of an electron escaping the molecule in a particular direction. This probability depends on how the molecule is oriented relative to the external magnetic field.

Currently used theories tie tunneling ionization rates to electron behavior far away from atomic nuclei. However, the available software for quantum mechanical calculations and computational chemistry fail to predict the state of electrons in those regions. The researchers found a way around this.

"We recently managed to reformulate the asymptotic theory of tunneling ionization so that the ionization rate would be determined by electron behavior near nuclei, which can be calculated rather precisely using the methods available now," Dnestryan said.







Dependence of calculated structure factors for the two highest occupied molecular orbitals -- HOMO and HOMO-1 -- of the naphthalene molecule on electric field orientation, that is, on the Euler angles β and γ from figure 1. The absolute values of the structure factor are color-coded, with red indicating the minimum values, and yellow and purple indicating the maximum values. The squared absolute value of the structure factor determines the rate of tunneling ionization from a given orbital in the direction opposing that of the field, since the electron is negatively charged. Credit: Moscow Institute of Physics and Technology

"Until now, researchers could only calculate tunneling ionization rates for small molecules made of a few atoms. It is now possible for significantly larger molecules. In our paper, we demonstrate this by running the calculations for benzene and naphthalene," the physicist added.

The authors of the paper calculated tunneling ionization rates for several molecules as a function of their orientation relative to the external field. To perform the calculations, the team developed software, which it plans to make openly available. This will enable experimenter to rapidly determine the structure of large molecules with attosecond precision based on observed spectra of the molecules.

"This work turns the asymptotic theory of tunneling ionization, which we developed in 2011, into a powerful tool for calculating ionization rates for arbitrary polyatomic molecules. This is essential for solving a wide range of problems in strong-field laser physics and attosecond physics," Tolstikhinsaid.

More information: Andrey I. Dnestryan et al, Structure factors for tunneling ionization rates of molecules: General grid-based methodology



and convergence studies, *The Journal of Chemical Physics* (2018). DOI: 10.1063/1.5046902

Provided by Moscow Institute of Physics and Technology

Citation: Physicists edge closer to controlling chemical reactions (2018, December 11) retrieved 1 May 2024 from <u>https://phys.org/news/2018-12-physicists-edge-closer-chemical-reactions.html</u>

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