Every moment of ultrafast chemical bonding captured on film
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The bond formation reaction in a gold trimer complex is initiated by a laser pulse, and a three-dimensional structure after a certain time delay is detected by an x-ray scattering image. Credit: IBS

Targeted cancer drugs work by striking a tight bond between cancer cells and specific molecular targets that are involved in the growth and spread of cancer. Detailed images of such chemical bonding sites or pathways can provide key information necessary for maximizing the efficacy of oncogene treatments. However, atomic movements in a molecule have never been captured in the middle of the action, not even for an extremely simple molecule such as a triatomic molecule, made of only three atoms.

A research team led by Ihee Hyotcherl of the Institute for Basic Science (IBS, South Korea) (Professor, Department of Chemistry, KAIST), in collaboration with scientists at the Institute of Materials Structure Science of KEK (KEK IMSS, Japan), RIKEN (Japan) and Pohang Accelerator Laboratory (PAL, South Korea), reported the direct observation of the birthing moment of chemical bonds by tracking real-time atomic positions in the molecule.

"We finally succeeded in capturing the ongoing reaction process of the chemical bond formation in the gold trimer. The femtosecond-resolution images revealed that such molecular events took place in two separate stages, not simultaneously as previously assumed," says Associate Director Ihee Hyotcherl, the corresponding author of the study. "The atoms in the gold trimer complex remain in motion even after the chemical bonding is complete. The distance between the atoms increased and decreased periodically, exhibiting the molecular vibration. These visualized molecular vibrations allowed us to name the characteristic motion of each observed vibrational mode," adds Ihee.

Atoms move extremely fast at a scale of femtosecond (fs)—quadrillionths of a second. The movement is minute on the level of angstroms, equal to one ten-billionth of a meter. They are especially elusive during the transition state where reaction intermediates are transitioning from reactants to products in a flash. The research team made this experimentally challenging task possible by using femtosecond X-ray liquidography (solution scattering).

This experimental technique combines laser photolysis and X-ray scattering techniques. When a laser pulse strikes the sample, X-rays scatter and initiate the chemical bond formation reaction in the gold trimer complex. Femtosecond X-ray pulses obtained from a special light source called an X-ray free-electron laser (XFEL) were used to interrogate the bond-forming process. The experiments were performed at two XFEL facilities (4th generation linear accelerator), PAL-XFEL in South Korea and SACL in Japan, and this study was conducted in collaboration with researchers from KEK IMSS, Pohang Accelerator Laboratory (PAL), RIKEN, and the Japan Synchrotron Radiation Research Institute (JASRI).

Scattered waves from each atom interfere with
each other and thus their X-ray scattering images are characterized by specific travel directions. The IBS research team traced real-time positions of the three gold atoms over time by analyzing X-ray scattering images, which are determined by a three-dimensional structure of a molecule. Structural changes in the molecule complex resulted in multiple characteristic scattering images over time. When a molecule is excited by a laser pulse, multiple vibrational quantum states are simultaneously excited. The superposition of several excited vibrational quantum states is called a wave packet. The researchers tracked the wave packet in three-dimensional nuclear coordinates and found that the first half round of chemical bonding was formed within 35 fs after photoexcitation. The second half of the reaction followed within 360 fs to complete the entire reaction dynamics.

In this study, the IBS research team improved upon their 2015 study published by *Nature*. In the previous study in 2015, the speed of the X-ray camera (time resolution) was limited to 500 fs, and the molecular structure had already changed to be linear with two chemical bonds within 500 fs. (Figure 2, upper right) In this study, the progress of the bond formation and bent-to-linear structural transformation could be observed in real time, thanks to the improvement time resolution down to 100 fs. Thereby, the asynchronous bond formation mechanism in which two chemical bonds are formed in 35 fs and 360 fs, respectively, and the bent-to-linear transformation completed in 335 fs were visualized (Figure 2, lower right). In short, in addition to observing the beginning and end of chemical reactions, they reported every step of the intermediate, ongoing rearrangement of nuclear configurations with dramatically improved experimental and analytical methods.

The research team will push this method of 'real-time tracking of atomic positions in a molecule and molecular vibration using femtosecond X-ray scattering' to reveal the mechanisms of organic and inorganic catalytic reactions and reactions involving proteins in the human body. "By directly tracking the molecular vibrations and real-time positions of all atoms in a molecule in the middle of reaction, we will be able to uncover mechanisms of various unknown organic and inorganic catalytic reactions and biochemical reactions," says Dr. KIM Jong Goo, the first author of the study.

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The researchers also accurately illustrated molecular vibration motions in both temporally and spatially. This is quite a remarkable feat considering that such an ultrafast speed and a minute length of motion are very challenging conditions for acquiring precise experimental data.

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