

Chemists find 'huge shortcut' for organic synthesis using C-H bonds

May 11 2016



A model of the new catalyst. "We've designed a catalyst that provides a huge shortcut for how chemists can turn a simple, abundant molecule into a much more complex, value-added molecule," says Emory chemist Huw Davies. Credit: Huw Davies lab/Emory University



Chemists have taken another major step in the quest to use carbonhydrogen bonds to create new molecules, a strategy that aims to revolutionize the field of organic synthesis.

The journal *Nature* is publishing the work by chemists at Emory University. They demonstrated the ability to selectively functionalize the unreactive carbon-hydrogen (C-H) bonds of an alkane without using a directing group, while also maintaining virtually full control of site selectivity and the three-dimensional shape of the <u>molecules</u> produced.

"The catalyst control we have found goes beyond what has been achieved before," says Huw Davies, an Emory professor of organic chemistry whose lab led the research. "We've designed a catalyst that provides a huge shortcut for how chemists can turn a simple, abundant molecule into a much more complex, value-added molecule. We hope this gives people a fundamentally new view of what can be achieved through C-H functionalization."

The first author of the *Nature* paper is Emory chemistry graduate student Kuangbiao Liao.

The streamlined process described in the paper holds tremendous potential for the synthesis of fine chemicals, such as those needed for the development of pharmaceuticals.

"Organic synthesis is all about simplicity," Davies says. "It may lead to a sophisticated outcome, but it has to be simple to carry out in order to have practical applications."

Davies is also director of the National Science Foundation's Center for Selective C-H Functionalization (CCHF), which is based at Emory and encompasses 15 major research universities from across the country, as well as industrial partners.



The CCHF is leading a paradigm shift in organic synthesis, which has traditionally focused on modifying reactive, or functional, groups in a molecule. C-H functionalization breaks this rule for how to make compounds: It bypasses the reactive groups and does synthesis at what would normally be considered inert carbon-hydrogen bonds, abundant in organic compounds.

The fact that multiple C-H bonds are commonly present in a single organic molecule, however, presents a significant challenge to this new type of chemistry. Chemists experimenting with C-H functionalization normally use a directing group - a chemical entity that combines to a catalyst and then directs the catalyst to a particular C-H bond.

"The directing group has to be introduced and then removed," Davies explains. "It works fine but the process is cumbersome."

The Davies lab bypassed the need for a directing group by developing a series of dirhodium catalysts encased within a three-dimensional scaffold. The scaffold acts like a lock and key to allow only one particular C-H bond in a compound to approach the catalyst and undergo the reaction.

"We had already demonstrated that we could do site selectivity of C-H bonds in molecules where C-H bonds are fairly activated," Davies says. "Here, we've gone for the ultimate challenge - the extremely unreactive C-H bonds of alkanes."

Alkanes are the simplest of organic molecules, consisting only of hydrogen and carbon atoms. In chemistry, the original name for alkane was paraffin, which is Latin for "lacking reactivity."

"Alkanes are cheap, plentiful raw materials that are considered nonfunctional, or unreactive, except in uncontrollable situations, such as



burning them," Davies says. "Our work, however, shows that it is indeed possible to efficiently functionalize an alkane in a controlled manner. We've actually changed the way that the description of alkanes in organic chemistry textbooks needs to be written."

In addition to controlling site selectivity, the scaffold of the dirhodium catalysts developed by the Davies lab controls the chirality of the product produced in the reaction.

Chirality, also known as "handedness," refers to a property of threedimensional symmetry. Just as the human hand is chiral, because the right hand is a <u>mirror image</u> of the left, molecules can be "right-handed" or "left-handed."

The handedness of a molecule is important in organic chemistry, since this 3D shape affects how it interacts with other handed molecules. When developing a new drug, for instance, it is vital to control the chirality of the drug molecules because biological molecules recognize the difference.

Thalidomide is the most notorious example of the handedness problem. During the late 1950s, thalidomide was sold as an over-the-counter drug for pregnant women suffering from morning sickness.

"It's difficult to make molecules with one mirror image, so many older drugs were a mixture of both," Davies explains. "It was thought that one mirror image would be biologically active, and the other wouldn't matter."

In the case of thalidomide, however, while one mirror image cured morning sickness, the other turned out to cause birth defects.

Today, pharmaceutical makers must either limit a drug's molecules to a



single chiral shape, or go through the extra time and expense of testing the safety of a mixture of left-handed and right-handed molecules.

The new C-H functionalization catalyst may pave the way to a whole new realm of materials for drug discovery research.

"The starting material we used, pentane, is as cheap as gasoline, but we are able to efficiently generate a sophisticated product in a single step and control which mirror image is formed," Davies says.

The process is also more sustainable than traditional <u>organic synthesis</u>, which typically involves the use of many reagents, and can produce toxic, inorganic byproducts.

"In contrast, our catalyst speeds up a reaction but is not used up in a reaction," Davies says. "Only a tiny amount of the catalyst is needed to produce a lot of product. And the only byproduct is nitrogen, which is innocuous."

While the *Nature* paper "opens the door" to a new method for C-H functionalization, more work is needed, Davies says. "We need to understand how to use it predictably and demonstrate its use in complex target applications."

The Center for Selective C-H Functionalization (CCHF) speeds up this exploration process by breaking down the walls of individual labs and research institutes to form collaborations and teaching networks that reach across the country and across continents. An exchange program lets students spend time in labs from any of the 15 U.S. research universities that form the core of the CCHF, as well as major research universities in Japan, South Korea and the United Kingdom.

"The center is much better known since we started in 2009," Davies



says. "We now have a world presence through virtual symposia on C-H functionalization that we give three times a year."

Anyone can join the symposia live, as they are broadcast on YouTube, or access them later via the CCHF web site. The most recent symposium drew a live audience of more than 1,300 chemists from 45 countries. "C-H functionalization is an important new way of thinking about synthesis, and we have leading speakers in the field giving talks on it, so there is a lot of interest," Davies says.

More information: Kuangbiao Liao et al, Site-selective and stereoselective functionalization of unactivated C–H bonds, *Nature* (2016). DOI: 10.1038/nature17651

Provided by Emory University

Citation: Chemists find 'huge shortcut' for organic synthesis using C-H bonds (2016, May 11) retrieved 16 August 2024 from <u>https://phys.org/news/2016-05-chemists-huge-shortcut-synthesis-c-h.html</u>

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