

New synthesis method for producing fluorinated piperidines

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Pressure vessel (autoclave) for the hydrogenation of fluorinated pyridines. The reactions are carried out at a hydrogen pressure of 50 bar (normal atmospheric pressure is 1 bar). Credit: Frank Glorius

Synthetic molecules are essential for many products, including medicines, crop protection agents and special materials such as Teflon.



These molecules have several components, which can be combined in a variety of ways, resulting in different properties. Both so-called piperidines and fluorinated groups are particularly important.

Piperidines are small, ring-shaped <u>chemical compounds</u>. As a result of their particular properties, fluorine atoms bring about dramatic changes in the properties of certain products. Thus, they are often integrated in pharmaceuticals. Around 20 percent of all medicines sold worldwide contain fluorine. Up to now, however, combining fluorine atoms and piperidines has always been an extremely laborious process. Now, for the first time, chemists at the University of Münster have developed a simple synthesis method for producing such fluorine-bearing piperidines. The study, written by Dr. Zackaria Nairoukh, Marco Wollenburg, Dr. Christoph Schlepphorst, Dr. Klaus Bergander and Prof. Frank Glorius, has just been published in the online edition of the *Nature Chemistry* journal.

In 2017, the Münster chemists had already published a study in *Science* in which they presented a method by which cyclic, fluorinated molecules could be produced simply and quickly. The chemists' new study follows on from this breakthrough.

"Up to now, it had been very difficult to combine piperidines and fluorine molecules," explains Prof. Frank Glorius from the Institute of Organic Chemistry at Münster University, "and this was despite the fact that together, they have outstanding chemical properties relevant to the production of active ingredients."

Methodology

The method developed by Frank Glorius and his team runs in two consecutive steps, but in the same vessel: The starting <u>molecules</u> are easily accessible, fluorinated pyridines, so-called "aromatic" compounds.



These compounds are flat and have a particularly high stability, which makes them inert for many chemical processes. In the method now published, the first step involves removing the aromaticity in a process called, unsurprisingly, dearomatization. This makes the second step possible, in which hydrogen atoms are transferred specifically to one side of the ring system (hydrogenation). The resulting fluorinated piperidines are now no longer flat, in contrast to the aromatic starting substances, and this can be helpful for the formation of complex threedimensional structures. For both reactions, the chemists used a catalyst.

The results can now be used to produce new types of valuable components previously unavailable for research in the fields of pharmaceuticals and agrochemicals. "We hope that these components can soon be produced in large quantities," says Frank Glorius. The Münster University chemists want to use part of the grant of 2.5 million euros recently awarded by the European Research Council to work on a "renaissance" of the hydrogenation of aromatics, leading to the efficient production of such new types of molecule. "We're going flat out in our efforts to try to understand more about the mechanism of catalysis, so that better catalysts can be developed and the potential of these reactions can be exploited to the full," says Glorius.

More information: Zackaria Nairoukh et al. The formation of allcis-(multi)fluorinated piperidines by a dearomatization–hydrogenation process, *Nature Chemistry* (2019). DOI: 10.1038/s41557-018-0197-2

Provided by Münster University

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