## Nonacethrene as a magnetic photoswitch: can one methyl group change the game?

<u>P. Pfister</u><sup>1</sup>, D. Čavlović<sup>1</sup>, C. Moreno Cruz<sup>2</sup>, M. Juríček<sup>1</sup>\*

<sup>1</sup>University of Zurich, Department of Chemistry, Winterthurerstrasse 190, 8057 Zurich, <sup>2</sup>University of Granada, Department of Chemistry, Avenida de la Fuente Nueva S/N 18071 Granada

In this project we aim to design a new all organic chiral magnetic photoswitch. The target compound is called nonacethrene and is the bigger homolog of cethrene, a chiroptical diradicaloid photoswitch. Pristine cethrene is too reactive but functionalization with two methyl groups in the fjord region leads to dimethylcethrene, which can be switched between an open and a closed form via light but does not possess an electron paramagnetic resonance (EPR) signal at room temperature. By expanding the  $\pi$ -backbone and therefore lowering the singlet-triplet gap, nonacethrene is EPR active. Nonacethrene undergoes an unwanted cascade reaction and dimethylnonacethrene is not reactive enough to act as a photoswitch. The adjustment of the steric bulk in the fjord region with only one methyl group represents an opportunity for further optimization to achieve bistability and is a viable strategy to realize a magnetic photoswitch operating at ambient temperature.