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“Switching Sides: Driving Reactions Uphill and Defining Design Rules for Photoswitchable Imines”

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Wykład w formie tradycyjnej

Sala Rady Wydziału 2.57, Wydział Chemii UAM

Abstract: Molecules and materials comprised of dynamic-covalent imine bonds display many desirable properties, including stimuli-responsiveness, recyclability, and effortless preparation, among others. However, the *E/Z* photochromism of imines has often been overlooked due to historically poor performance compared to other photoswitches, typically showing less than 50% conversion to the metastable state and thermal half-lives of under one minute.¹ Recently, we developed a strategy that significantly improves these photoswitching properties, achieving quantitative *E*-to-*Z* conversion with visible light and extending the thermal half-lives of the metastable *Z*-state to over one day.^{2,3}

With these improved properties, we explored the light-dependent dynamic-covalent chemistry of these photoswitches. To our surprise, we found that they can drive transamination reactions energetically uphill when irradiated

with light,⁴ operating via a mechanism akin to a light-driven information ratchet.⁵ Beyond pushing thermodynamic equilibria into non-equilibrium steady states, we found that the regiochemistry of the C=N bond plays a significant role in the switching behavior.^{3,6} Through a systematic investigation, a set of design rules for this novel class of imine photoswitch has now been reported, along with the first crystal structure of an arylimine in its metastable *Z*-conformation.⁶ This talk will provide an overview of our recent progress in this area.

Selected references:

- [1] L. Greb et al., in *Molecular Photoswitches*, Wiley, 2022, pp. 325–349.
- [2] J. Wu, J. L. Greenfield et al., *Chem. Sci.*, 2024, 15, 3872–3878.
- [3] J. Wu, J. L. Greenfield et al., *Chem. Commun.*, 2024, 60, 12365–12368.
- [4] J. Wu, J. L. Greenfield, *JACS*, 2024, 146, 20720–20727.
- [5] V. Serreli et al., *Nature*, 2007, 445, 523–527.
- [6] J. Wu, J. L. Greenfield et al., *Angew. Chem. Int. Ed.*, 2024, e202415464.

