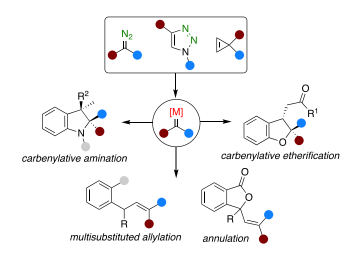
Functionalization of Metallocarbenes Derived from Diazo and Non-Diazo Surrogates

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Abstract:

Metallocarbenes show versatile reactivity in organic synthesis and offers access to diverse complex frameworks in single step. Most often these metallocarbenes are generated from α -diazocarbonyl compounds in the presence of suitable transition metal.¹ Recently, *N*-sulfonyl-1,2,3-triazoles and cyclopropenes have emerged as unique surrogates and offers structurally different metallocarbenes, which possess distinct reactivity.² The unique reactivity of these metallocarbene precursor have been efficiently utilized in our lab for the carbenylative functionalization of *o*-vinylaniline/phenol derivatives, multisubstituted allylation of arenes and annulation reactions.³ In addition, we are also studying their asymmetric approach with suitable chiral catalyst. In this presentation, our efforts on the catalytic functionalization of metallocarbenes and their asymmetric version will be discussed.



References and Notes:

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