

## Seminar

## Rationally Designed Carbon-Carbon and Carbon-Nitrogen Diradicaloids and Boron-Boron Diradicals

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Open-shell compounds with second-row p-block elements boron, carbon, nitrogen, and oxygen are crucial for various applications. The synthesis of carbon-carbon-centred diradicaloids typically involves redox processes of their corresponding cationic/anionic forms or the dehydrogenation of judicious polycyclic aromatic hydrocarbons. Nitrogen-nitrogen-centred dicationic diradicaloids are also known by the oxidation of corresponding neutral form. However, the synthesis of heteronuclear carbon-nitrogen-centred diradicaloids is challenging due to the simultaneous presence of open-shell centres at similar electrochemical potentials at both carbon and nitrogen centres, often resulting in short lifetime. Boron-boron-centred diradicals are only known with considering borane radical-anion motif.

In my talk, I will showcase our finding on the synthesis of i) carbon-carbon-centred diradicaloids under two-electron oxidation of  $\pi$ -conjugated bis-N-heterocyclic olefins, ii) carbon-nitrogencentred tricationic diradicaloids by judicious choice of alkene and amine under two-electron oxidation, and iii) boron-boron-centred neutral diradicals considering NHC-stabilised boryl radical motif boron-centred spin carrier.

*Thursday, Dec 26<sup>th</sup> 2024* 16:00 Hrs (Tea / Coffee 15:45 Hrs) Auditorium, TIFRH