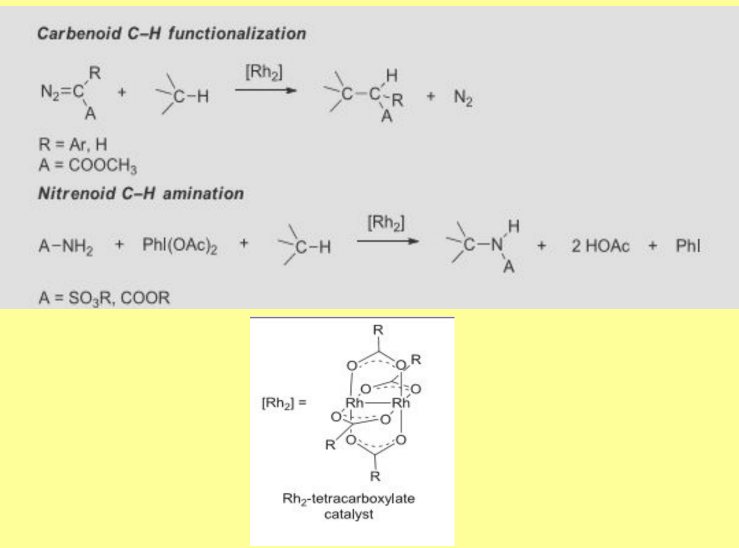
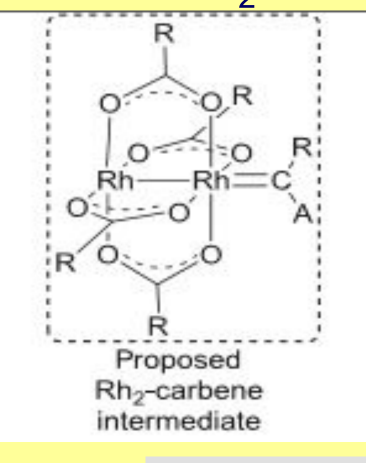
Metal–metal multiple bonded intermediates in catalysis

(for example,  $Rh_2$  and  $Ru_2$  complexes)

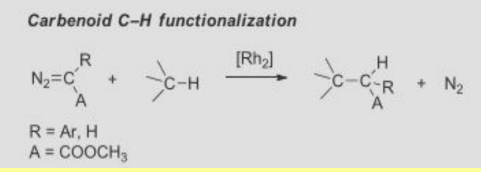
#### Overview of Rh<sub>2</sub>-catalysed C–H functionalization and C–H anination chemistries



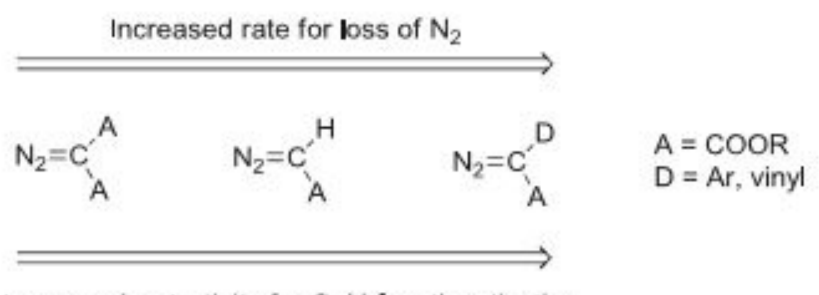
#### Rh<sub>2</sub> carbene chemistry



The key electronic feature of this intermediate is delocalized Rh–Rh–C three-centre bonding with appropriate three-centre orbitals of  $\sigma$  and  $\pi$ symmetry

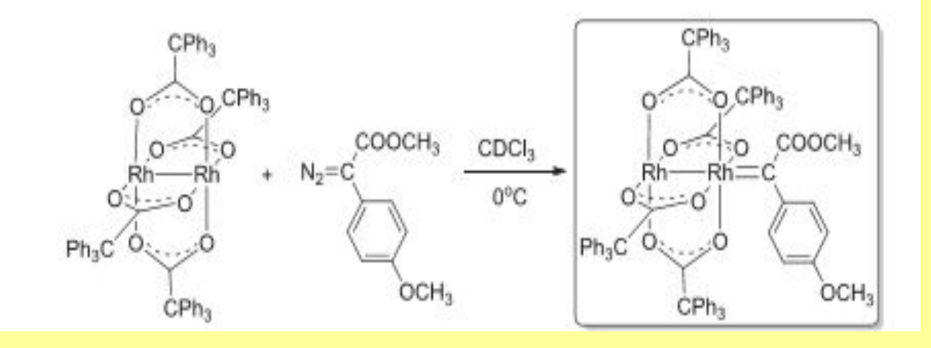


### Trends in reactivity for the different classes of organic diazo compounds

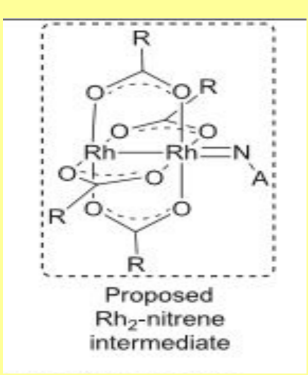


Increased selectivity for C-H functionalization

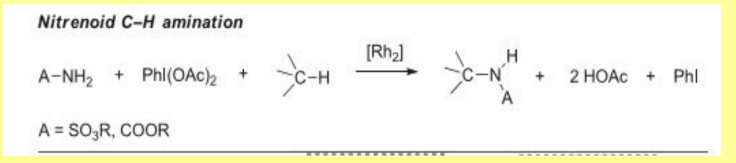
## Preparation of the first Rh<sub>2</sub> D/A carbene complex



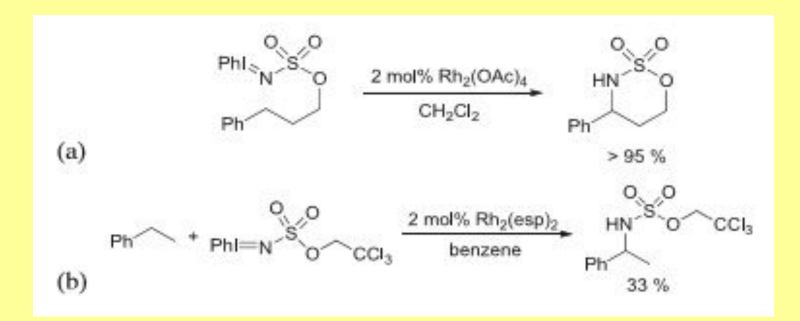
#### Rh<sub>2</sub> nitrene chemistry



Rh<sub>2</sub>-catalysed nitrenoid chemistry is mechanistically more complex than the corresponding carbenoid chemistry

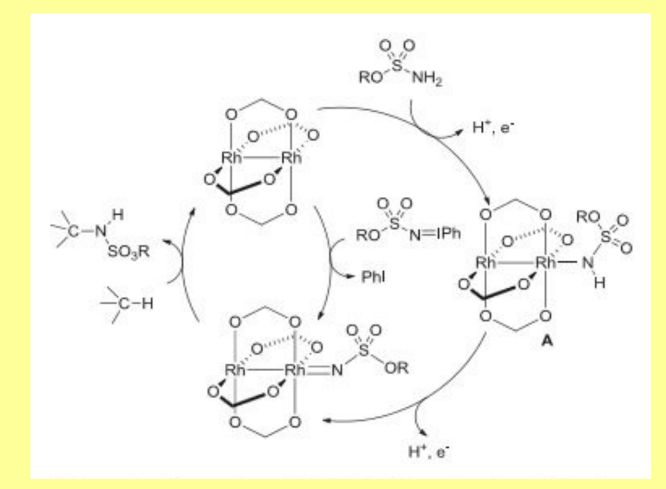


## Reactions using pre-formed iminoiodinane compounds



(a) – intramolecular cyclization(b) – intermolecular reaction

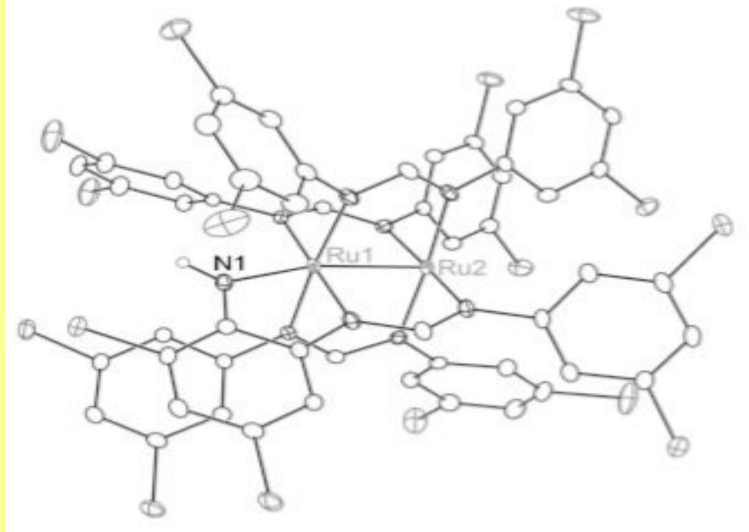
# Proposed mechanism for intermolecular C–H amination



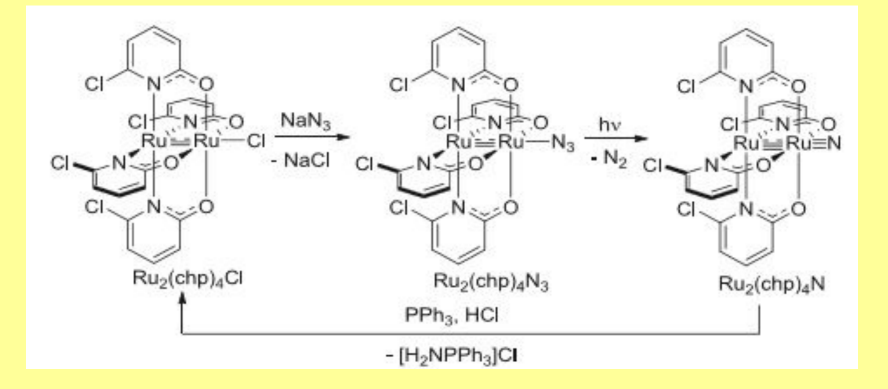
Organic groups on the catalyst are removed for clarity

Ru<sub>2</sub> nitrido chemistry  $Rh-Rh=E \square M-M=E \square Ru-Ru=N$ structures structures structure  $(E = CR_2/NR)$ The first Ru<sub>2</sub> nitrido compound  $-Ru_2(DPhF)_AN$ (DPhF = N,N'-diphenylformamidinate) – was found to be thermally unstable In an effort to understand the nature of this instability, the related Ru<sub>2</sub>(D(3,5-Cl<sub>2</sub>)PhF)<sub>4</sub>N<sub>3</sub> azide complex was investigated

# $Ru_{2}[(D(3,5-Cl_{2})PhF)_{3}(D(3,5-Cl_{2}-2-N)H)PhF)]$



# Synthetic cycle for N-atom transfer using the Ru<sub>2</sub>(chp)<sub>4</sub> core



### Summary

Efforts to identify reactive metal-metal bonded complexes having a linear M–M=E structure have led to the observation of important intermediates in Rh<sub>2</sub>-catalysed carbenoid and nitrenoid transformations. Inspired by the structures of these intermediates, chemists have been able to explore novel reactivity of the Ru–Ru≡N core including intramolecular C-H amination as well as intermolecular N atom transfer.

#### Source

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