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1. Introduction

The field of N-heterocyclic carbene (NHC) complexes of 3d metals has grown significantly in recent years since the first synthesis of an isolable imidazol-2-ylidene by Arduengo in 1991.¹ Consequently, NHCs have found numerous applications across fields such as catalysis, materials sciences and metallopharmaceuticals. Comprehensive and specialised reviews have described the synthesis, practical applications and recent

Department of Chemistry, National University of Singapore, 4 Science Drive 2, Singapore, 117544, Republic of Singapore. E-mail: chmkmj@nus.edu.sg developments (not limited to nickel) of NHCs including NHCenabled organocatalysis.²⁻⁹

The structure of NHCs can have a significant impact on their behaviour as ligands. Four key factors that influence this behaviour are the presence and location of heteroatoms (such as nitrogen), substituents at proximal positions, the backbone structure and the size of the ring.⁴ The adjacent electronegative nitrogen atoms can inductively withdraw electron density from the non-bonding lone pair at the carbenic carbon (HOMO). At the same time, nitrogen lone pairs can overlap with the empty p-orbital of the carbenic carbon as a π -electron donor. Steric



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N-Heterocyclic carbenes as privileged ligands for

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Alkene functionalisation is a powerful strategy that has enabled access to a wide array of compounds including valuable pharmaceuticals and agrochemicals. The reactivity of the alkene π -bond has allowed incorporation of a diverse range of atoms and functional groups through a wide variety of reaction pathways. N-Heterocyclic carbenes (NHCs) are a class of persistent carbenes that are widely employed as ancillary ligands due to their ability to act as strong σ -donors compared to widely-applied conventional phosphine-based ligands. NHCs are also unique as their molecular bulk provides steric influence for regio- and stereo-control in many alkene functionalisation reactions, illustrated by the examples covered in this review. A combination of the unique reactivity of NHC ligands and nickel's characteristics has facilitated the design of reaction pathways that show distinct selectivity and reactivity,

including the activation of bonds previously considered "inert", such as C-H bonds, the C-O bond of ethers and esters, and the C-N bonds of amides. This review summarises the advancements in Ni(NHC)

catalysed alkene functionalisation up to 2022, covering the following major reaction classes: Heck-type

nickel-catalysed alkene functionalisation

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reactions, hydrofunctionalisation and dicarbofunctionalisation.

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Chart 1 Characteristic and classes of common NHCs. Ad: adamantyl, Me: methyl, Mes: mesityl, Cy: cyclohexyl, ^tBu: *tert*-butyl, and iPr: iso-propyl.

bulk on the *N*-substituents serves to protect the reactive centre from kinetic dimerisation and decomposition whilst altering the electronic properties of the ring system. Unsaturation in the cyclic backbone provides enhanced thermodynamic stability due to the increased aromaticity of the ring.⁵ Different ring sizes in the backbone also have a marked influence on the overall stereoelectronic properties of the NHC ligand (Chart 1).

Combining the abovementioned factors, NHCs are generally regarded to have a highly nucleophilic carbenic carbon atom, making them strong σ -donors that bind strongly to metal or non-metal centres. This characteristic has led to their wide-spread use as ligands in numerous catalytic applications. NHCs are also considered to be more electron-donating than

phosphine ligands, as reflected in their lower calculated Tolman electronic parameter¹⁰ values. This results in the formation of thermodynamically stronger metal-ligand bonds in NHC-metal complexes, as reflected in the greater bond dissociation energies and shorter metal-ligand bond lengths observed compared to those of their phosphine-ligated complexes.⁵ This in turn confers greater thermal and oxidative stability to the Ni-NHC complex.¹¹ In the domain of steric factors, the measurement of percentage buried volume ($%V_{bur}$) can be used to quantitatively compare the steric bulk of NHC ligands with that of phosphine ligands.¹² Modelling studies on the $%V_{\text{bur}}$ value of most NHC ligands have shown that they are as bulky or even more so than most regular phosphine ligands, particularly IAd (1a) and I^tBu (1e). This ultimately has significant implications on the reactivity and selectivity of reactions, which we will cover in detail.

Research in this area has also expanded to include the use of non-classical NHCs (see Chart 1), most of which possess an even stronger electron-donating ability compared to their classical counterparts and have been proven to stabilise low coordination number metal (NHC) complexes, as well as reactive transient metal species. Numerous published reviews and book chapters¹³ have already covered the current state of research on well-defined NHC complexes of copper,¹⁴ nickel,¹⁵ cobalt,¹⁶ and iron.¹⁷ These articles extensively examine the development of these systems in terms of their structural, catalytic, and stoichiometric reactivities, as well as the aspects of their synthesis, bioinorganic chemistry, material chemistry, and photophysical properties.

Nickel, located just above palladium in the periodic table, is a group 10 metal and can perform many similar elementary reactions to palladium or platinum. Due to its smaller atomic radius, lower electronegativity, and lower redox potential, nickel exhibits faster rates of oxidative addition and slower rates of β -hydride elimination when compared to palladium in the same ligand framework.¹⁸ Thus, nickel is often used as a cost-effective alternative to its noble metal counterparts and





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can also offer a more facile access to multiple oxidation states $(Ni^0, Ni^I, Ni^{II}, Ni^{III} and even Ni^{IV})$, rendering radical pathways more accessible.

There are several reported methods for the synthesis of Ni-NHC complexes. These include the interaction of isolated or in situ generated NHCs with the common Ni⁰ and Ni^{II} precursors, Ag transmetalation protocols, and C-X (X = H, halide) bond activation methodologies.¹⁶ Rare oxidation states of nickel (*i.e.*, Ni^I), proposed as reactive catalytic intermediates, can also be stabilised with various NHCs, and are usually synthesised from the disproportionation of Ni⁰ and Ni^{III} precursors or single-electron reduction protocols of Ni^{II}, as described in a recent review.¹⁹ Synthesis and characterisation of various Ni(NHC) complexes of different nuclearity and oxidation states (Ni⁰, Ni^{di} and Ni^{II}) are unfortunately outside the scope of this review. Nickel, being more electropositive than palladium, can under certain conditions activate bonds (C-O of ethers, C-N of amides) that are relatively inert to oxidative addition with palladium.²⁰ The strong electron-donating properties of NHCs further facilitate this oxidative addition process, allowing Ni(NHC) catalysts to activate even more challenging C-H bonds.

Ni(NHC) complexes have a long history of use in homogenous catalysis and have been thoroughly reviewed for their use in C-C cross-couplings, C-H bond functionalisation, C-X (X = O, N, S, B) bond formation, oligo/polymerisation, and oxidation-reduction reactions.21,22 This review will not cover these previously mentioned classes of reactions. In this review, recent developments in Ni(NHC) catalysed alkene functionalisation will be discussed, mainly categorised by the three main reaction classes of alkene functionalisation: Heck-type reactions, hydrofunctionalisations and difunctionalisations (Scheme 1). The contents for this review will not include reactions with strongly activated enones, alkynes²³⁻²⁷ or cycloaddition-type reactions.²⁸⁻³¹ The literature studies presented in this review are up to date until 2022. The origins of regioselectivity, diastereoselectivity and enantioselectivity in the discussed reactions will be thoroughly examined, including comprehensive mechanistic details, if deemed necessary.

Alkene functionalisation offers chemists a platform to rapidly assemble molecular complexity and stereogenicity from simple starting materials such as alkenes, due to their versatile reactivity and abundance as a feedstock chemical.³²

There have been numerous developments in the use of homogenous nickel catalysis (not limited to NHC ligands) for the functionalisation of alkenes in recent years, which demonstrate its effectiveness in performing such valuable transformations. This has also been highlighted in several reviews.^{32,33} Nickel's high effectiveness as a catalyst for alkene functionalisation can be attributed to the following factors: nickel readily donates d-electrons to π -acceptors, resulting in strong olefin bonding. The less electronegative nickel undergoes more facile oxidative addition with electrophiles. β -Hydride elimination tends to be slower with nickel as compared to palladium, due to the significantly higher energy barrier for Ni–C bond rotation prior to β -hydride elimination in the organonickel species.³⁴



Scheme 1 Different classes of alkene functionalisation covered in this review.

As such, Ni(NHC) catalytic systems have also enabled the hydrofunctionalisation and difunctionalisation of alkenes beyond Heck-type reactions.

2. Heck-type reaction

The Mizoroki–Heck reaction has been established as one of the most important methods for the formation of valuable C–C bonds to afford substituted alkenes, and has been utilised significantly in organic chemistry, including the synthesis of various natural products.³⁵ It falls under the class of "palladium-catalysed cross-coupling reactions" for which the 2010 Nobel Prize in Chemistry was awarded to Richard F. Heck, Ei-ichi Negishi, and Akira Suzuki for their pioneering efforts.³⁶

Nickel has also been found to be effective in catalysing Hecktype reactions,³⁷ which follow a mechanism similar to that of the well-explored palladium-catalysed reaction.³⁸ This is illustrated in a four-step catalytic cycle as shown in Scheme 2. First, the metal complex undergoes oxidative addition of the alkyl, aryl or alkenyl-(pseudo)halide bond, facilitated by the electrondonating ligand(s). Then, the olefin associates with the metal to form a π -complex, followed by the migratory insertion of the olefin into the carbon-metal bond. The regioselectivity of the Heck reaction, if any, is governed by this olefin insertion step, which can produce either 1,1-disubstituted or 1,2-substituted alkenes as the possible regioisomers. In the third step, β -hydride elimination occurs, furnishing the coupled product and a metal-hydride intermediate. Finally, the active metal catalyst is regenerated after the base-assisted removal of HX, completing the catalytic cycle. Variations in the Heck mechanism have also surfaced, suggesting the involvement of cationic Ni(II) intermediates.39



Scheme 2 Typical Heck catalytic cycle (where M = nickel or palladium).

Although numerous examples of the Heck reaction enabled with various phosphine-ligated Ni complexes have been reported,³⁷ the use of NHC ligands to effect such transformations has been largely elusive and unexplored. The use of nickel–NHC complexes in the Heck reaction was first described by Sakamoto *et al.*⁴⁰ in 2005. Their Ni(NHC) catalyst system was found to be highly efficient for the Heck reaction between aryl bromides/iodides with acrylates. The desired cinnamate-type products were obtained in high yields and with excellent *trans*-selectivity (Scheme 3). The reaction system was also evaluated for functional group tolerance and was found to work efficiently with nitriles (**8**), esters (**10**), aldehydes (**9**) and thiophenes (7).

In 2007, Schleicher and Jamison developed a Ni(0)/IPr catalysed formal intermolecular Heck coupling between α -olefins and isocyanates, affording N-alkylacrylamides in good yields and with excellent regioselectivity⁴¹ (Scheme 4). The observed regioselectivity for the C-C bond formation (prior to β -hydride elimination) was attributed to the sterically demanding NHC ligand, where the R^1 substituent of the α -olefin prefers to be oriented away from the bulky isopropyl group of the IPr 1f ligand (Scheme 4), providing 1,1-disubstituted N-alkylated acrylamide as the major product. This is opposite in selectivity to previous reports under the Ni/PR3 catalytic system,42 where the major product was a *trans*-1,2-disubstituted α , β -unsaturated amide, instead. Aliphatic olefins with branching at the allylic position were highly effective substrates (11 and 13), furnishing the desired product in excellent yields and with regioselectivity. Compared to linear alkyl chains (n-hexyl in 14), the presence of coordinating groups (ketone in 12) in the olefins also resulted in enhanced selectivity of the products. However, the reaction appears to be only selective for monosubstituted terminal olefins and is also limited to bulky and electron-rich (Cy or ^tBusubstituted) alkyl isocyanates.

In the same year, Ho and Jamison developed a Ni-catalysed three-component Heck-type coupling reaction between alkenes,



 $\mbox{Scheme 3}$ First Ni/NHC-catalysed Heck reaction of aryl halides with acrylates. 40

aldehydes and silvl triflates to deliver the 1,1-disubstituted allylic alcohol product with excellent regioselectivity⁴³ (Scheme 5) Their methodology utilises the synergistic relationship between a strong σ -donor ligand (NHC) and a strong π -acceptor phosphite ligand $(P(OPh)_3)$. Only when both are used in tandem will the reaction proceed catalytically with high selectivity and efficiency. From their preliminary optimisation and mechanistic studies, in the absence of the IPr 1f ligand, no product was observed. In the absence of P(OPh)₃, there was no catalytic turnover, resulting in much lower conversion and yield, but without diminishing its selectivity. Their postulated mechanism (Scheme 5) involves the initial formation of the oxa-nickelacycle 20 from the Ni-NHC complex, alkene, and aldehyde, assisted by the Lewis Acid; silyl triflate. This step determines the regioselectivity of the oxanickelacycle assembly 20, which ultimately leads to the 1,1disubstituted allylic alcohol product. The observed regioselectivity could be attributed to the bulky IPr 1f ligand which disfavours the other regioisomer (oxanickelacycle 20'). The role of the phosphite co-ligand was to accelerate the reductive elimination of the [Ni(IPr)H]OTf species 21 (via 22 and 23), which in its absence, accumulates in the reaction mixture, leading to side reactions and hence, diminished yields. It was postulated that the formal reductive elimination of 21 to 19 to





close the catalytic cycle was too slow (hence no catalytic behaviour in the absence of phosphite) due to the presence of the strong σ -donating IPr ligand.⁴³ Anisaldehydes proved to be excellent substrates (**15** and **17**) while the presence of chlorine (**16**) and heteroaryl substituted aldehydes was less efficient (**18**).

In addition to the commonly employed $[Ni(COD)_2]/IPr$ system, which can be used as a precursor for the *in situ* generation of [(NHC)NiH]OTf, Jamison and co-workers have developed an alternative, novel bench-stable nickel^{II}-NHC precatalyst (24), which can be efficiently reduced to the NHC–Ni⁰ active species in the presence of silyl triflate (Scheme 6).⁴⁴ The architecture of the nickel–NHC precatalyst was inspired by phosphine–nickel complexes, featuring a bidentate aryl ligand containing a piperidine moiety that occupies the coordination sites around the nickel, thereby stabilising the complex. Additionally, a pendant alkene linkage is present, which serves to reduce the Ni^{II} to Ni⁰ (in its active form) through an intramolecular Heck reaction.

This bright yellow nickel^{II}–NHC precatalyst (**24**) also exhibits remarkable stability in air and can be easily isolated using column chromatography with neutral alumina. The precatalyst was found to be very effective in promoting the carbonyl-ene type coupling of alkenes and aldehyde, exhibiting even greater



 $\label{eq:scheme 5} \begin{array}{l} \mbox{Regioselective formal Heck coupling of alkenes and aldehydes} \\ \mbox{with } [Ni(NHC)P(OPh_3)].^{43} \end{array}$

catalytic activity than the previously developed $Ni(COD)_2/IPr$ system (Scheme 5).

More recently, Koh and co-workers reported a novel Ni(i)/NHC-catalysed regio- and stereo-selective coupling reaction of monosubstituted alkenes with a variety of electrophilic



 $\label{eq:scheme 6} \begin{array}{l} \text{Bench-stable Ni}^{II}/\text{NHC precatalyst developed by Jamison and coworkers.} \end{array}$

reagents to access tri- and tetra-substituted alkenes in high yields (up to 92%) and selectivity (>98% E:Z).⁴⁵ Mechanistic and computational studies support the idea of a novel tandem pathway involving a site-selective Heck-type reaction followed by a stereoselective C=C bond isomerisation, under a Ni^I/Ni^{III} catalytic pathway (Scheme 7). Based on their proposed mechanism, initial dissociation of the dinuclear IPr-derived Ni(1) dimeric complex (25), and ligand exchange from sodium tertpentoxide (NaOt-Am) forms the catalytically active Ni(I)alkoxide 30. In the presence of an aryl triflate and a monosubstituted alkene, the Mizoroki-Heck reaction occurs, through an initial oxidative addition of 30 with aryl triflate, followed by a regioselective arylnickelation and β -hydride elimination. This produces an intermediate π -complex of Ni^{III}-hydride and the 1,1-disubstituted alkene 33. Base-promoted reductive elimination of 33 then generates another olefin-coordinated Ni^Ialkoxide 34. Mechanistic studies and computational modelling of the transition states (of trans-TS-34 to 35 versus cis-TS-34 to 35) suggest a probable stereoselective non-radical allyl isomerisation sequence of 34. 34 undergoes an uphill (hence requiring higher reaction temperatures) trans-selective allylic C-H insertion to give the η^1 -allylnickel(m) intermediate 35, followed by a subsequent allyl migration to form 36 and a final irreversible reductive elimination to furnish the desired trisubstituted alkene product, regenerating the catalytically active Ni^Ialkoxide 30. Despite the uphill olefin migration step, the overall process was computed to be thermodynamically favourable, with an energy change of $-58.1 \text{ kcal mol}^{-1.45}$

A plethora of functionalised monosubstituted alkenes and aryl triflates were found to be compatible under the optimised protocol. Electrophilic ketone-containing polycyclic triflates (26), epoxide (28) and silane (29) were effective substrates. With slight modifications, this reaction can also be extended to the synthesis of tri- or tetra-substituted alkenyl boronates (27),



Koh 2021

Scheme 7 Ni()/IPr-catalysed stereoselective synthesis of alkenes *via* formal Heck reaction/alkene isomerisation.⁴⁵

which are valuable intermediates in the preparation of many medicinally relevant pharmaceuticals.^{46,47} Analogous to the above proposed mechanism (Scheme 7), the alkenyl boronate product is likely formed from an initial boryl-Heck type reaction followed by an olefin isomerisation.

An extension to their previous work (Scheme 7), Koh and co-workers reported a Ni(i)/IPr catalysed branched-selective Heck-type benzylation-stereoselective olefin isomerisation methodology to access tri-substituted alkenes.⁴⁸ The same dinuclear IPr-derived Ni(i) dimeric complex (25) was used as the precatalyst, which efficiently enabled the reaction of styrenes/vinylboronates with various functionalised benzyl chlorides under basic conditions, to afford trisubstituted alkenes with high *trans*-selectivity. The excellent branched selectivity (where the benzyl group is attached to the more substituted olefinic carbon) of the initial Heck-type benzylation is attributed to the steric control provided by the bulky NHC ligand, which effectively overrides the olefin's strong electronic bias for linear selectivity.

The proposed mechanism for the tandem Heck-type benzylation–isomerisation reaction (Scheme 8) is similar the nonradical Ni^I/Ni^{III} pathway as illustrated previously (*cf.* Scheme 7). Reaction of the catalytically active monomeric Ni^I–X (X = Cl or Br) complex **40** with benzyl chloride forms the putative π -allyl nickel intermediate **41**. Subsequent ligand exchange with TMSOTf forms a cationic Ni^{III} intermediate **42** with the weakly bound non-nucleophilic triflate anion, accompanied with the TMS-halide by-product. The incoming styrene/vinylboronate substrate can form an initial π -complex with electrophilic nickel **42**, followed by the regioselective migratory insertion of the benzyl group to yield a primary alkylnickel species **43**.



Regiochemical outcome for this branched-selective benzylation (instead of linear-selective insertion) can be rationalised by the favoured transition state of lowest energy with minimised steric repulsions between the bulky IPr NHC ligand and the R¹ substituent of the olefin. β -H elimination then forms an olefin-associated Ni^I complex 44, which undergoes facile deprotonation with the weak Et₃N base to give 45. Lastly, with motivation from the results of previous DFT calculations,⁴⁵ *trans*-selective insertion (from 45 to 46) into the benzylic C–H bond takes place (to minimise steric interactions between the aryl group Ar, halide X and IPr ligand) followed by a non-radical allyl migration to form the final allylic η^1 -Ni complex 47, which reductively eliminates to deliver the final trisubstituted alkene in high *trans*-selectivity.

In the presence of TBAB additives, various benzyl chlorides bearing heteroaryl groups (37) were compatible with vinylboronate, to access highly substituted alkenylboronates in moderate yields. Highly biologically relevant carbohydrate-derived styrene substrates (39) were also effective. The utility of this regime was also demonstrated with the efficient synthesis of a retinoid precursor (38).

3. Hydrofunctionalisation of alkenes

In a broad sense, hydrofunctionalisation of alkenes refers to the installation of a hydrogen atom and a functional group across the C==C π bond of the alkene, thereby generating a new C(sp³) centre. Numerous reports have surfaced in recent years,^{49–51} documenting the use of Ni catalysts for the hydrofunctionalisation of alkenes, including the development of asymmetric variants which enables the synthesis of valuable, optically active molecules using simple alkene feedstock. In this review, we will not restrict our analysis to C-C bond forming reactions (*i.e.*, hydrocarbofunctionalisation), despite their importance. We will also be discussing various studies that demonstrate the versatility of hydrofunctionalisation using Ni/NHC catalysts in the hydrosilylation, hydroboration, and hydroamination of alkenes.

Not all hydrofunctionalisation reactions follow a universal reaction pathway. Herein, we illustrate certain subtle mechanistic nuances of hydrofunctionalisation reactions often disclosed in literature studies (Scheme 9). The C–H activation of the substrate as well as the migratory insertion into C=C can proceed *via* a single transition state,⁵² known as metalmediated ligand-to-ligand hydrogen transfer (LLHT), first termed by Perutz *et al.*⁵³ in their DFT-supported mechanistic study of Nakao's Ni-catalysed hydrofluoroarylation reaction.⁵⁴ This concerted pathway (Scheme 9a) proceeds without the involvement of nickel-hydride (Ni–H) species intermediates, which are commonly proposed in catalytic cycles of Ni-catalysed alkene transformation reactions.⁵⁵

A non-concerted stepwise mechanism of hydrofunctionalisation is also plausible, with Ni–H involved as intermediary species. An initial insertion of the alkene into the Ni–H bond generates an alkyl-Ni intermediate, which can react with the





2) non-oxidative in-situ generation of [Ni]-H



c) Stepwise insertion of Ni-O followed by reductive elimination of Ni-H



= C(sp³), C(sp²), C(sp), B, Si, N



desired electrophile (via insertion or oxidative addition). Subsequent β -hydride elimination or reductive elimination affords the final product (Scheme 9b). The formation of the Ni-H intermediate can occur through an oxidative route which involves the oxidative addition of C-H or Si-H bonds by low valent Ni complexes^{55,56} (Scheme 9b1). The non-oxidative route usually involves the in situ formation of Ni-H via reaction of a main group hydride (e.g., LiEt₃BH) and σ -bond metathesis with silanes (Si-H) or boranes (B-H).55 Other methods include the treatment of Ni⁰/IPr with silvltriflate, aldehyde and alkene, which effectively produces "[IPrNiH]OTf" species from an oxanickelacycle intermediate as discussed previously in Jamison's work⁴³ (Scheme 9b2). An alternative feasible pathway for this transformation could be the initial carbonickelation followed by β-hydride elimination to produce an alkyl Ni-H intermediate,^{57,58} which undergoes reductive elimination to deliver the final hydrofunctionalised product (Scheme 9c).

In addition to the common ionic pathways involved in hydrofunctionalisation (illustrated in Scheme 9), there are also reports of radical-based pathways proceeding *via* metal-hydride hydrogen atom transfer (MHAT).⁵⁹ These will be briefly discussed along with relevant literature studies. Intricate mechanistic details governing the regioselectivity (Markovnikov or anti-Markovnikov addition) and/or stereoselectivity of specific reactions will also be analysed in detail.

3.1 Hydroalkenylation of alkenes

Transition-metal catalysed alkene hydroalkenylation/arylation reactions offer an efficient way to construct highly valuable $C(sp^3)-C(sp^2)$ bonds. However, as with most hydrofunctionalisation reactions, there are several key considerations beyond the reaction itself, including regioselectivity (Markovnikov *vs.* Anti-Markovnikov), stereoselectivity, and the catalyst's turnover frequency, all of which can greatly influence the outcome and efficiency of the reaction.

Hydroalkenylation generally refers to the addition of a vinyl/ alkenyl group and a hydrogen atom across an unsaturated C=C bond. The products of a typical hydroalkenylation reaction are like those of a Heck reaction, but there are subtle variations in the mechanistic pathway. This process can also be viewed as a heterodimerisation or cross-coupling reaction between alkenes.

The exploration and utilisation of NHCs as ligands for hydroalkenylation reactions were pioneered and mainly carried out by Ho's group. In 2010, Ho and co-workers reported the Ni/NHC-catalysed intermolecular branched-selective (tail-totail) hetero-hydroalkenylation of styrenes with α -olefins.⁶⁰ The tentative proposed catalytic species is an *in situ* generated [(NHC)NiH]OTf species, prepared from a modified procedure reported by Jamison and co-workers in their [(IPr)NiP(OPh)₃]mediated silyltriflate/alkene/aldehyde coupling reaction (Scheme 5). Building on this initial discovery, subsequent research by Ho's group primarily centered on the chemistry of an NHC–nickel hydride species generated *in situ*, which they identified as active catalysts in a variety of hydroalkenylation reactions (Schemes 11–13).

In their proposed mechanism (Scheme 10), addition of the Ni-H of the in situ generated (NHC)nickel-H species 48 to the vinylarene (chemoselective and regioselective hydronickelation) forms the resonance-stabilised benzylic nickel complex **49.** The incoming α -olefin (with a less bulky substituent than the aryl group of the vinylarene) is then coordinated with 49 in an orientation that minimises steric repulsion between the bulky NHC ligand and the alkene substituent, forming the π -complex 50 (favoured over 50'), which is primed to undergo regioselective migratory insertion with the α -olefin to give the primary nickel intermediate 51. Finally, β -hydride elimination occurs, releasing the final hetero-coupled gem-disubstituted alkene product and regenerating the nickel hydride catalyst 48. It is noteworthy that, despite the thermodynamic driving force for the isomerisation of 1,1-disubstituted alkenes to the more stable internal or conjugated alkenes, only limited isomerisation was observed under those conditions. It is believed that the steric effect of the bulky IPr ligand explains this observation and governs the regioselectivity of the migratory insertion process as well as the chemoselectivity for the desired hetero-coupling reaction.

In 2015, Ho and co-workers developed the asymmetric variant of their previously reported hetero-hydroalkenylation





Scheme 10 Ni/NHC catalysed branched-selective hetero-hydroalkenylation of styrenes and unactivated α -olefins.⁶⁰

of vinylarenes and unactivated α -olefins, with the use of an unsymmetrical chiral NHC ligand (52). This reaction also operates with an analogous mechanism to Scheme 10, where the active species is the *in situ*-generated [(52)NiH]OTf.⁴³ Unsymmetrical *gem*-disubstituted olefins were obtained as the final major product in high yields and high ee (Scheme 11). Aryl-substituted vinylarenes like 3-methoxystyrene (56) and thioanisole (55) were well-tolerated, giving the corresponding product in good yields and ee. Regarding the scope of terminal olefins, the use of allylbenzenes (53) resulted in the highest obtained ee, while vinylcyclohexane (54) was also found to be an effective coupling partner.

The author proposed a tentative stereochemical model to rationalise the enantio-determining vinylarene insertion step



Ho 2015

Scheme 11 Asymmetric Ni/chiral NHC catalysed hetero-hydroalkenylation of styrenes and $\alpha\text{-olefins.}^{61}$

into Ni–H, purely based on steric factors, which minimises the unfavourable steric repulsion between the *N*-aryl substituent of the NHC and the aryl group of the vinylarene substrate (Scheme 11). This is exemplified with the increased ee values when more sterically bulky α -olefins were employed. The asymmetric induction by the Ni/chiral NHC complex can be influenced by the different *N*-aryl substituents on the NHC ligand as

well as the external π -systems of the vinylarenes. However, their proposed stereochemical model has its limitations and does not fully explain how certain electronic factors can affect the enantioselectivity. For example, it is not fully understood why the presence of an *ortho*-methyl substituent on the *N*-aryl ring (of the NHC) is crucial for high enantioselectivity.

Not limited to only coupling with vinylarenes, Ho and coworkers further applied their chemistry on *in situ* generated [(NHC)NiH]OTf⁴³ to design a novel method to access valuable organosilanes. In 2011, they developed the cross-hydroalkenylation reaction of vinylsilanes and α -olefins to afford homoallylsilanes in high chemoselectivity and regioselectivity (Scheme 12).⁶² Unlike their previous work on the cross-hydroalkenylation using vinylarenes,^{60,61} which affords the branched tail-to-tail product, the use of vinylsilanes results in a complete reversal of regioselectivity, governed exclusively by the steric properties of the NHC ligand.



Scheme 12 Ni/NHC catalysed cross-hydroalkenylation of vinylsilanes and α -olefins.

The rationale for the observed chemoselectivity and regioselectivity in the reaction mechanism lies in the intricacies of the reaction mechanism. Specifically, the regioselectivity of Ni-H insertion is primarily directed by the steric influence of the bulky NHC ligand. In this reaction, 1,2-hydronickelation with the vinylsilane reagent occurs preferentially to 2,1-hydronickelation to give the primary alkylnickel intermediate 60, which is less sterically hindered. Unlike previous reports on reactions with vinylarenes,^{60,61} there is little thermodynamic incentive for 2,1-hydronickelation to occur with vinylsilanes due to the absence of adjacent aryl groups for π -stabilisation. Interestingly, β -silvl elimination of **60** is also strongly suppressed, presumably due to the attenuation of orbital overlap of Ni-β Si under the steric environment of the NHC. Coordination of α -olefin with 60 (in an orientation that minimises steric repulsion) forms 61, which subsequently undergoes another regioselective insertion to furnish 62. β-hydride elimination of 62 then regenerates the NHC nickel-hydride species and releases the homoallylsilane product in overall excellent regioselectivity.

Modifications to the steric and electronic properties of the Si centre do not appear to have a large influence on the reaction's efficiency (57,58). The reaction is also tolerant to a variety of substituents on the α -olefins, including enolisable esters (59).

In 2014, Ho and co-workers developed a novel pathway to access seven-/eight-membered medium sized heterocycles *via* a Ni/NHC-catalysed n^{γ} -*exo-trig* selective cycloisomerisation reaction of 1,*n*-heterodienes (n = 7,8).⁶³ Their novel work utilises the synergistic effect between NHC's strong electron-donating and steric properties and γ -heteroatom chelation to selectively form medium-sized heterocycles despite the increased ring strain. Through their optimisation studies and competitive experiments, they found that the tethering length, γ -oxy chelation, and the steric effect of NHC all play important roles in the selective formation of the n^{γ} -*exo-trig* product over the competitive (n - 1)^{γ}-*exo-trig* product. Other than substituted oxepanes, siloxepane (63) and thiepane (64) can also be obtained through this reaction. 1,8-Oxadienes (65) also proved effective, successfully delivering the desired eight-membered oxecane product.

Their proposed mechanism (Scheme 13) postulates that the steric influence of the bulky NHC ligand promotes the initial 1,2-hydronickelation of the allyl ether arm of 66 (from an in situ generated [(NHC)NiH]OTf species), forming a less hindered primary carbocentre on nickel. Chelation assistance from the heteroatom (e.g., oxygen) present in the 1,n-diene also proves to be crucial for the reaction's efficiency and selectivity, possible stabilising the hydrometallated nickel species in the form of an oxanickelacyclopentane 67. The strongly electron-donating NHC ligand also led to improvement in functional group tolerance while mitigating the strength of γ -oxy chelation to the nickel centre. It must be balanced in a way that is strong enough as a stabilising group, yet not too strong that it deactivates the nickel catalyst. The stabilised oxanickelacyclopentane 67 then undergoes a facile intramolecular n^{γ} -exo-trig ring closure reaction to form the cyclised intermediate 68, which releases the desired product after a syn β -H elimination





Ho 2017

Scheme 14 Ni/NHC catalysed cross-hydroalkenylation of endocyclic dienes with $\alpha\text{-olefins.}^{64}$

step. While $(n - 1)^{\gamma}$ -exo-trig favours a product with a lower ring strain, the strong preference for n^{γ} -exo-trig could be due to the less favourable 2,1-hydronickelation on the homoallyl ether terminus and the higher energy barrier for ring closure with a secondary alkylnickel-NHC species. However, it is noteworthy to point out that no in-depth mechanistic studies or calculations on the transition states were performed. It is therefore possible that other competitive insertion pathways may exist. However, due to the reversible nature of hydrometallation reactions, the steric influence of the NHC ligand may direct the mixed intermediates to converge into the lowest energy optimised geometry for ring closure, resulting in the observed selectivity for the n^{γ} -exo-trig product.

In 2014, Ho and co-workers described the first cross-hydroalkenylation of endocyclic 1,3-dienes with α -olefins enabled with neutral and cationic Ni/NHC precatalysts⁶⁴ (Scheme 14). A rare product selectivity was observed under this protocol, where the 4,3-hydroalkenylation (the alkenyl group and H atom are introduced at the C3 and C4 position of the original diene substrate, respectively) skipped diene product was afforded with high chemo-, regio- and diastereo-selectivity. The reaction can be efficiently promoted with an in situ generated [(IPr)Ni-H(OTf)⁴³ or [(69)Ni(allyl)]⁺Bar_F⁻ complex.⁶⁵ Upon reaction with catalytic alkenes (1-octene was used), the precatalyst [(69)Ni-(allyl)]⁺BAr_F⁻ generates a similar [(NHC)NiH] hydride complex in an allylnickelation-β-hydride elimination sequence. Regioselective hydronickelation at the 4,3-position of the endocyclic 1,3-diene (at the least substituted C=C) initially forms the allylic nickel intermediate (nickel at 3-position). The allylic shift of this nickel intermediate to a more substituted carbon of the diene (1- or 2-position) was not observed, possibly due to

the sterically encumbered environment around the bulky NHC, which partly governs the observed regioselectivity for hydronickelation as well. The mechanism for this reaction could be thought of as analogous to that illustrated in Scheme 10.

Another important criterion for desired reactivity is the presence of a coordinating heteroatom, either on the diene or the α -olefin substrate. It is postulated that the electronic effect of the heteroatom suppresses the undesired isomerisation of the alkene substrate. Cyclohepta-1,3-dienes were well tolerated, forming the corresponding skipped diene product in good yield (71). More sterically hindered fused bicyclic 1,3-dienes (73) were also feasible under their system, with the use of a less sterically hindered N-cyclohexyl-N'-mesityl NHC ligand (Scheme 14). Notably, high levels of diastereoselectivity of hydroalkenylation were also observed when chiral endocyclic dienes were employed (72). To minimise the steric strain between the diene substituents and the NHC ligand in the Ni-complex, the insertion of the α -olefin occurs on the face, with the newly formed stereogenic centre oriented anti to the existing chiral centre in the diene.

As an extension of their previous work (Scheme 14), Ho and co-workers developed an enantioselective cross-hydroalkenylation reaction of cyclic 1,3-dienes and hetero-substituted terminal alkenes/ α -olefins to afford endocyclic 1,4 skipped dienes with a chiral allylic framework⁶⁶ (Scheme 15). Similarly, the reaction employs the use of $[(74)Ni(allyl)]^+BAr_F^-$ as the precatalyst, which forms the active nickel–hydride species upon reaction with catalytic amount of octene. A less bulky and structurally flexible C_2 -symmetric chiral NHC ligand was crucial for the balance of reactivity and selectivity of this reaction, considering the undesired steric interactions between the



Scheme 15 Ni/chiral NHC catalysed enantioselective synthesis of 1,4dienes by cross-hydroalkenylation of endocyclic 1,3-dienes and heterosubstituted terminal alkenes.⁶⁶</sup>

substituted dienes and the *N*-aryl substituents on the NHC. Hetero-substituted 1,3-dienes (75) without bulky R1 or R2 substituents proved to be compatible, delivering the product in moderate yield and excellent ee. N- or O-substituted terminal olefins were also efficient (76, 77). The reaction is also highly chemoselective for the insertion of the less hindered terminal alkenes over internal allylic ethers as exemplified with 77. Effective 1,3-allylic shift of the NHC–Ni intermediate can also be achieved without the erosion of enantioselectivity, which was found to be controlled by the steric properties of the substituents on the substrate. The 1,3-allylic shift equilibrium

is also tunable with the electronic properties of the heterosubstituted terminal alkenes. Using a more electron-rich alkene (78) favours the 1,3-allylic shift diene product in excellent ee despite the 1-position being more hindered (adjacent to the gem-dimethyl and silvl ether moiety in 78). From the proposed stereochemical model of the transition state, the enantiodetermining step is likely to be the hydronickelation of the diene, under the chiral environment of the NHC-Ni-H active species. Minimisation of the steric interactions between the substituents of the cyclic 1,3-diene and the cyclohexyl group of the C_2 -symmetric NHC ligand favours a face of nickel-hydride insertion in the conformation illustrated in Scheme 14. 1,3-Allylic shift of the allyl-nickel intermediate (if possible) then proceeds after nickel-hydride insertion. Subsequent regioselective insertion of the hetero-substituted terminal olefin (induced by the steric control of the NHC ligand) followed by β -H elimination affords the final enantioenriched allylic 1,4-diene product

Leveraging the utility of $[(NHC)Ni(allyl)]^+BAr_F^-$ as an efficient precatalyst for hydroalkenylation, Ho and co-workers successfully achieved a novel regiodivergent synthesis of 1,2and 1,3-disubstituted allyl ethers in one step from vinyl ethers and terminal alkenes⁶⁷ (Scheme 16). Under their mild protocol and by careful design of the NHC ligands (1c, 79 and 80 in Scheme 16), unwanted side reactions like cationic polymerisation, β -OR elimination and metal-catalyst deactivation (common for reactions with vinyl ethers)⁶⁸ were effectively suppressed. This enabled electron-rich vinyl ethers to act as acceptors, leading to the formation of a new C(sp²)-C bond at the 1-position which delivers the allylic ether product, opening a new mode of reactivity for vinyl ethers.

Key elements of the reaction include the initial 2,1hydronickelation (H at the terminal 2-position and Ni at the internal 1-position adjacent to the –OR group) of the vinyl ether substrate by the *in situ* generated [(NHC)Ni–H] active species. This suggests that the chemo- (coordination of vinyl ether over terminal alkene) and regio-selective (2,1-addition over 1,2addition) hydronickelation is not controlled by the steric effects of the bulky NHC ligand, as the 2,1-addition of Ni–H leads to the formation of a more hindered 2° alkyl nickel intermediate. Rather, this step can be attributed to electronic control, favoured by the catalyst–substrate interaction. Subsequent regioselective migratory insertion of the terminal alkene was controlled by the steric interactions between the alkene substituent and the *N*-aryl substituent on the NHC (R^s being less bulky than mesityl), as illustrated in Scheme 16.

This allows for regiodivergence, where 1,2 or 1,3-disubstituted allyl ethers can be selectively obtained just by varying the nature of NHC ligands. The use of a bulky NHC ligand 1c allows for the selective formation of the 1,2-product (81, a : b = 91 : 9), while the use of a sterically less demanding 79 and 80 favours the formation of 1,3-disubstituted allylic ether (82-84). 80 was designed to be less electron-rich than 79 while being less sterically bulky than 1c (absence of one methyl group of the *N*-aryl), which proved to be more efficient for the reaction of (*Z*)-configured internal vinyl ethers with terminal alkenes,

X = NMs







Ho 2020

Scheme 17 NHC/Ni-catalysed 1,3- and 1,4-diastereodivergent hetero-cycle synthesis from enynes. 69

delivering **84** with excellent chemo- and regio-selectivity. Even electron-deficient vinylsilanes (**83**) were compatible under their reaction protocol, despite being bulkier than the vinyl ether substrate. This further corroborates that the chemoselectivity for the reaction is largely governed by electronic effects, with cross-coupling being preferred over homo-coupling of the terminal alkenes/vinyl ethers.

In 2020, Ho and co-workers devised a novel synthetic methodology for the synthesis of heterocycles, which involves the NHC/Ni-catalysed 1,3- and 1,4-diastereodivergent intramolecular hydroalkenylation of heteroenynes⁶⁹ (Scheme 17). Although the authors did not explicitly detail the exact mechanism behind the origin of diastereoselectivity, preliminary mechanistic studies suggest the potential involvement of an NHC-nickelacyclopentene intermediate (illustrated in Scheme 17) instead of Ni–H insertion to the enyne. The different X heteroatom of the heteroenyne, in combination with the steric influence of the NHC, could favour a specific conformation leading to the diastereoselective formation of the nickelacyclopentene. Trapping of the NHC-nickelacyclopentene with an alcohol then results in a formal reductive hydroalkenylation process, leading to the expedient synthesis of various N- or O-heterocycles bearing an exocyclic olefin. The reductive hydroalkenylation reaction was shown to be effective with various substituted propargyl- or allyl– X (X = O or N) substrates, producing 1,3-*syn/anti* or 1,4-*syn/anti* products with high diastereoselectivity depending on the nature of X. Not limited to terminal alkenes, reductive hydroalkenylation with internal alkenes (**85**, **87**) also proceeded efficiently. Notably, heterocycles with stereodefined quaternary centres can also be accessed using this strategy.

3.2 Hydroarylation of alkenes

Hydro(hetero)arylation, akin to hydroalkenylation, represents a straightforward method for the construction of $C(sp^3)$ – $C(sp^2)$ bonds and a means to access synthetically valuable alkyl(hetero)arenes from simple olefin feedstock. Traditional methods of hydroarylation involve the classical Friedel-Crafts type alkylation of electron-rich arenes,⁷⁰ palladium-catalysed reductive Heck coupling of alkenes⁷¹ and transition metal-catalysed C–H oxidative addition – hydrometallation – reductive elimination sequence⁵¹ (Scheme 9b). The last of these methods includes examples of Ni/NHC catalysis which will be discussed in this section. The ligand-to-ligand hydrogen transfer (LLHT)-type mechanism and arylnickelation – nickel hydride reductive elimination are also possible pathways suggested for hydroarylation reactions, as supported with DFT calculations of the different transition state energies.

Seminal work in this field was disclosed by Hiyama and coworkers in 2009 with the development of a C6-selective functionalisation reaction of pyridone derivatives – by insertion into unsaturated C=C bonds – enabled with nickel/Lewis acid catalysis.⁷² This reaction also represents a formal hydroheteroarylation reaction of alkenes. While most of their optimisation studies and reaction scope mainly involves the use of a bulky, electron-rich triisopropylphosphine, two successful examples (**89-90**) using IMes **1c** were also highlighted, demonstrating the effectiveness of NHC ligands in this process as well.

The N-heteroaryl substrate is first activated by the Lewis acid catalyst (AlMe₃) upon coordination with the more basic carbonyl oxygen. η^2 -coordination of the activated pyridone ring with (NHC)Ni⁰ primed it for a subsequent oxidative addition of the C6–H bond, forming the heteroaryl-nickel hydride intermediate **92**. Coordination of the 2-vinylnapthalene substrate to the Ni centre in **92**, with the bulky IMes ligand oriented away from the naphthalene group (to minimise steric repulsion) forms **93**, which undergoes a regioselective hydronickelation to furnish the π -allyl nickel intermediate **94**. The stabilisation of the benzylic nickel **94** by the adjacent π -system of the naphthalene is presumably responsible for the observed regioselectivity in migratory insertion. Final reductive elimination of **94** affords the hydroarylated product.

In 2010, Hiyama and co-workers expanded upon their preliminary scope of Ni/Lewis acid co-catalysed hydroarylation of 2-vinylnaphthalene (Scheme 18)⁷² and reported a Ni/IMescatalysed hydroheteroarylation reaction of vinylarenes, to furnish a diverse range of heteroaryl-bearing 1,1-diarylalkanes (Scheme 19).⁷³ In contrast to their previous work, no Lewis acid co-catalyst is required for heteroaryl activation, possibly due to the use of relatively more electron-poor heteroarenes. A variety of heteroarenes were well-tolerated, including an enolisable ketone-containing indole derivative (97) and a benzothiazole (98). Electron-rich vinylarenes were also effective substrates (95). Notably, 1,2-disubstituted internal alkenes (96) also participated well in the reaction.

Deuterium labelling experiments and previous mechanistic investigation into similar systems suggest a plausible catalytic pathway as illustrated (Scheme 19). Coordination of the heteroarene with Ni⁰/IMes forms the η^2 -arenenickel **99** which undergoes reversible oxidative addition of the C(sp²)–H bond of the heteroarene to give the nickel–hydride intermediate **100**. A series of reversible coordination of the vinylarene substrate followed by a regioselective hydronickelation affords the 1-arylethylnickel **102**, stabilised by the adjacent π -system. Lastly, irreversible reductive elimination of **102** regenerates the Ni/IMes species and releases the 1,1-diarylethane product.

In the same year, Hiyama and co-workers reported a (NHC)Ni/Lewis acid co-catalysed C-4-selective addition of



pyridine across alkenes⁷⁴ (Scheme 20). Direct alkylation of the C-4 position of pyridines has been more elusive and challenging to achieve compared to functionalisation at the C-2 position. Their protocol avoids the stoichiometric activation of the pyridine substrate and achieves direct C-4 alkylation under the steric control provided by both the highly bulky NHC ligand (IPr **1f**) and the aluminium-based Lewis acid MAD (MAD = methylaluminium bis(2,6-di-*tert*-butyl-4-methylphenoxide)).

Chemoselective addition to terminal alkenes over internal alkene was demonstrated (103), and no detectable second addition of pyridine was observed. Ester-containing pyridine derivatives like methyl nicotinate (104) also participated well in the reaction. Using styrene as a coupling partner leads to exclusive formation of the branched C-4 alkylated product, instead of the linear product obtained with aliphatic alkenes. This highlights the substrate-controlled regiodivergence in the insertion step, with the selective formation of a stabilised benzylic nickel intermediate (with vinylarene substrates) over the primary nickel intermediate (with an aliphatic alkene substrate).

A plausible reaction mechanism is illustrated in Scheme 20. Coordination of the Ni^0/NHC with the activated pyridine



Scheme 19 Ni/IMes-catalysed hydroheteroarylation of vinylarenes.⁷³

substrate (after N-coordination to MAD) forms η^2 -arenenickel 106. Kinetically favoured oxidative addition of the C(4)-H bond of the activated pyridine substrate (preferred over the C(2)-H or C(3)-H bonds) forms the nickel-hydride intermediate 107. Regioselective hydronickelation into the aliphatic alkenes from the π -complex **108** forms the less sterically hindered primary alkylnickel intermediate 109 (if a vinylarene substrate was used, a π -stabilised benzylic nickel intermediate was preferentially formed instead^{73,74}). The regioselectivity for insertion is solely governed by steric factors, due to the lack of electronic stabilisation of the branched 2° alkylnickel regioisomer (after insertion to aliphatic alkenes), in addition to the increased steric repulsion between the NHC and alkene substituent. Ratedetermining, irreversible reductive elimination of 109 yields the final branched C4-alkylated pyridine product and regenerates 106.

More recently, building upon the pioneering efforts from Nakao and Hiyama,⁷⁴ Shi and co-workers reported the enantioselective hydropyridylation of β -substituted styrene derivatives enabled by a cooperative chiral NHC/Ni–Al bimetallic catalysis.⁷⁵ Similarly, the C(4)–H bond of the pyridine is selectively activated and added across an alkene, to deliver 1,1diarylalkanes with high levels of regio- and enantio-control. Tunable bulky C_2 -symmetric chiral NHC – ANIPE-type ligand – as







Scheme 20 (NHC)Ni/Lewis acid catalysed selective C4-alkylation of pyridine/hydropyridylation of alkenes.⁷⁴

developed by their group⁷⁶ was used, with the bulky 3,4-xylyl substituted ANIPE ligand **110** providing the best regioselectivity and enantioselectivity. The steric environment of bulky NHC is postulated to supress the undesired β -H elimination side reaction which gives the Heck product whilst facilitating reductive elimination.

The use of β -substituted styrenes as acceptors is crucial for efficient hydroarylation, possibly due to their weaker ability to coordinate to the nickel catalyst. Two molecules of unsubstituted styrenes were found to strongly coordinate to the nickel centre, deactivating the catalyst. cis-Substituted alkenes also result in poorer regioselectivity (with a greater proportion of a linear anti-Markovnikov product being formed) and enantioselectivity, likely due to the smaller steric repulsion between them and the NHC ligand. The authors performed detailed DFT first-principles calculations to elucidate the mechanism and the origin of the enantioselectivity. Supported with the results from previously done calculations,⁷⁷ their calculations also suggest a reaction pathway involving a ligand-to-ligand hydrogen transfer (LLHT) mechanism, contrary to the stepwise oxidative addition-hydronickelation pathway as suggested by Hiyama and co-workers.⁷⁴ LLHT involving the ortho or meta-C-H bonds was also less feasible due to a higher energy barrier,



Scheme 21 Enantioselective (NHC)Ni/Lewis acid-catalysed hydroarylation of styrene/C4-alkylation of pyridines.⁷⁵

which supports the observed regioselectivity. Calculated energies of the transition state for the rate- and *enantio*-determining LLHT process (Scheme 21) also revealed the lower energy barrier incurred in the formation of the (*S*)-isomer over the (*R*)-isomer due to the stronger π - π stacking between the NHC aryl ring and the coordinated *trans*-styrene. The computed relative energies align well with the experimental enantiomeric ratio, where the (*S*)-isomer is the dominant product.

 β -Silyl-substituted styrenes (111) and medicinally relevant N-heterocycles (112) were effective substrates, delivering the enantioenriched 1,1-diarylethane product in high yields and enantiomeric ratio. Ferrocene-containing alkene also participated well (114). Utilisation of an *ortho*-substituted pyridine substrate (113) which presents increased steric hindrance led to

greater complexity, potentially attributable to the inhibition of coordination with the aluminium Lewis acid. Switching to a less bulky triethylaluminium Lewis acid was imperative in catalysing the required transformation, albeit with a lower yield due to formation of a competing C6-alkylated side product.

Hiyama and co-workers continued their efforts towards NHC-nickel/Lewis acid co-catalysed hydroarylation of alkenes, this time enabling the selective formation of linear alkylated pyridones⁷⁸ (Scheme 22). This work explores the novel use of unactivated alkenes as viable coupling partners instead of activated vinylarene-type substrates. The optimal NHC ligand of choice is 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene (IPr 1f), in synergy with MAD as the Lewis acid co-catalyst, which efficiently delivers the linear hydroarylation product in good yields and selectivity. Pyridone derivatives like 1-methylquinazolone delivered the desired alkylated product (115) in good yields. Bulky vinylsilanes (116), gem-disubstituted alkenes (117), or cyclohexene (118) also participated well, and the linear products were obtained in high yields. The reaction mechanism is analogous to their previous report,⁷⁴ where the linear selectivity originates from the regioselective insertion to form the less sterically hindered primary alkylnickel intermediate. With no electronic stabilisation possible (for unactivated aliphatic alkene substrates), this step is solely governed by steric control from the bulky IPr NHC.

In 2012, Ong and co-workers reported the heteroaromatic C-H activation of benzimidazole derivatives following a vinylarene insertion step, enabled by cooperative (NHC)Ni/Lewis acid catalysis⁶⁶ (Scheme 23). This is, in essence, a hydroheteroarylation reaction of styrene. While the work of Hiyama *et al.*^{59,65} predominantly uses substrate control for the selective formation of branched (with vinylarene as a substrate) or linear (with aliphatic alkenes as a substrate) hydroarylated products,



 $\mbox{Scheme 22} \quad (\mbox{NHC})\mbox{Ni/Lewis acid catalysed formation of linear alkylated pyridones.}^{78}$



Scheme 23 Regioselective switching for the linear/branched hydroarylation product *via* (amino-NHC)/Ni–Al synergistic catalysis.⁷⁹

Ong and co-workers developed a chemical regioselective switch to selectively obtain either the linear or branched product from the same vinylarene substrate. Their system utilises an amino-NHC (with a pendant *tert*-butyl amino arm) ligand, with the chemical switch being the trimethylaluminum AlMe₃ Lewis acid. Addition of the AlMe₃ co-catalyst forms the (NHC)Ni–Al bimetallic complex which enforces a steric environment favouring the formation of a linear insertion product while the absence of AlMe₃ switches the regioselectivity, favouring the branched-selective product probably due to the favourable electronic stabilisation of the benzylic nickel intermediate (**120** *vs.* **121** and **122** *vs.* **123**), which was established in previous reports, as well.^{59,60,62}

In-depth mechanistic studies to probe the nature of this regioselective switch enabled with AlMe₃ were also conducted by the authors,⁶⁷ with the proposed mechanism as illustrated in Scheme 23. There are two distinct pathways differing in the regioselectivity of insertion (in the presence or absence of AlMe₃). Generally, a concerted oxidative addition of the C-H bond of the benzimidazole substrate generates the nickelhydride intermediate Al-124. Facile coordination of the styrene substrate followed by regioselective hydronickelation yields the linear primary nickel species (Al-126), as controlled by the steric environment around the NHC-Ni/Al bimetallic catalyst or an electronically stabilised branched benzylic nickel species (126) in the absence of AlMe₃. Reductive elimination of 126 under electronic control and Al-126 under steric control delivers the branched and linear products, respectively. Notably, the addition of AlMe3 was found to accelerate the hydroarylation reaction, as evident from the increased product yields and lower reaction times (120 vs. 121 and 122 vs. 123). The bidentate amino-NHC with a reversibly coordinating tert-butyl amino pendant group as well as the nature of the R² substituent on the benzimidazole substrate could also play an important role in providing the additional steric pressure around the nickel complex, hence increasing the sterically-controlled linear selectivity.

In 2013, Ong and co-workers reported the first (NHC)Nicatalysed hydroarylation reaction of allylarenes that operated under a tandem alkene isomerisation-heteroaryl $C(sp^2)$ –H bond activation process⁸⁰ (Scheme 24), selectively obtaining the branched 1,1-di(hetero)arylethane product. A linear hydroarylated product can also be selectively achieved by using a chemical toggle (Lewis acidic AlMe₃)⁷⁹ as previously described, which effectively retards alkene isomerisation while promoting the hydroarylation process. Naturally occurring compounds like safrole participated well, delivering the branched (128) and linear (127) products with high selectivity. Other heteroarenes like imidazoles (129) or indoles (130) were also welltolerated.

The criterion for an efficient branched-selective reaction hinges on the alkene isomerisation cycle being relatively faster than the C-H bond activation process, allowing for the two catalytic cycles to operate independently. Preliminary mechanistic studies by the authors and inspiration from previous similar literature studies⁸¹ suggest a mechanism with two operating catalytic cycles (Scheme 24, **cycle A** and **B**), involving a more facile alkene chain-walking mechanism (**cycle A**) and a



Scheme 24 Regioselective hydroarylation of allylarenes via tandem alkene isomerisation and C–H bond activation. 80

hydroarylation mechanism (**cycle B**). In cycle A, oxidative addition of the allylic and benzylic C-H bond forms the η^3 - π allylnickel hydride intermediate **131**, which undergoes a formal 1,3-hydride shift to form the thermodynamic internal alkene **132**.

Cycle B is initiated with the oxidative addition of the C-H bond of the heteroarene to form the arylnickel hydride complex 133. Coordination of the isomerised alkene 132 to the nickel hydride intermediate 133 followed by the regioselective hydronickelation forms the electronically stabilised benzylic nickel intermediate 135. Facile reductive elimination of 135 affords the final branched hydroheteroarylation product, regenerating the active catalyst. The pathway responsible for the formation of the linear product follows an analogous mechanism outlined in Scheme 23 (under steric control). Addition of the chemical trigger AlMe₃ increases the steric bulk around the (NHC)Ni-Al bimetallic active catalyst (NHC = IMes 1c or IPr 1f), which deactivates alkene isomerisation and further enhances the selectivity of migratory insertion to form the less hindered primary alkylnickel intermediate, leading to the formation of the linear product.

Ong and co-workers further expanded on their previous work on bimetallic (NHC)Ni/Al catalysed hydroarylation reaction of allylbenzenes⁸⁰ and achieved convenient access to both linear or branched hydropyridylation products using their regiodivergent protocol⁸² (Scheme 25). The *para*-C-H bond of pyridines was selectively activated and inserted into allylbenzene substrates, following a similar mechanism to that



Scheme 25 (NHC)Ni/Lewis acid-catalysed hydropyridylation of allylarenes with switchable regioselectivity. $^{82}\,$

described in Scheme 20 (for linear selective insertion) and Scheme 24 (tandem alkene isomerisation and branched selective insertion).

Unlike their previous work, this regiodivergence does not arise from the absence or presence of the aluminium Lewis acid as a chemical trigger,⁸⁰ but rather from the size of the NHC ligand/Lewis acid pair. Using a very sterically demanding NHC ligand and a Lewis acid (*i.e.*, condition A with IPr **1f** and MAD) was found to favour the formation of a linear hydropyridylation isomer 137 *via* the less hindered migratory insertion process, with the suppression of alkene isomerisation as well. On the other end, employing a less bulky NHC/Lewis acid pair (*i.e.*, condition B with amino-NHC-2 and AlMe₃) toggles the isomerisation of allylbenzene to the internal alkene before hydroarylation takes place, delivering the final product with high branched selectivity **138**. Subjecting *gem*-disubstituted allylbenzene to the reaction (condition A) also afforded the desired product 139 without diminishment in positional selectivity.

In 2017, Mandal and co-workers reported the use of an abnormal NHC (aNHC) ligand in combination with a nickel(0) precatalyst to promote the hydroheteroarylation of vinylarenes with benzoxazole derivatives, resulting in the exclusive formation of the branched 1,1-diarylalkane product (Scheme 26).⁸³ This is a novel application of a non-Arduengo-type NHC ligand for alkene functionalisation (hydroarylation) despite its emerging utility in organometallic catalysis. Preliminary mechanistic studies led to the observation and isolation of a Ni(II) cyclooctenyl complex 145 (characterised with NMR and X-ray crystallography), formed by the reaction of the free aNHC ligand and Ni(COD)2. Reduction of this Ni^{II}-aNHC precatalyst forms the active Ni⁰-aNHC species, which initiates the reaction by the oxidative addition of the C-H bond (of benzoxazole). Subsequent migratory insertion of aNHCnickel hydride into vinylarene forms an electronically stabilised benzylic nickel intermediate, which then undergoes reductive elimination to form the desired branched hydroheteroarylation product, overall similar to previously postulated hydroarylation mechanisms.^{82,84} While electronic perturbation of the benzoxazole and styrene substrates with the introduction of electron donating or withdrawing groups does not significantly affect the reaction yields (141-144), the scope of the heteroarene is largely limited to derivatives of benzoxazole only.

In 2012, Ananikov and co-workers presented a novel catalytic system for the hydroheteroarylation of alkenes. Their catalytic system utilises a commercially available bench stable Ni(Cp)₂ (bis(cyclopentadienyl)nickel) precatalyst **147**, an NHC salt as a pro-ligand, and sodium formate as a non-toxic reductant to generate the active Ni⁰ catalyst⁸⁵ (Scheme 27, condition A). This approach presents an alternative methodology which obviates the necessity for the utilisation of highly air and moisture sensitive Ni(COD)₂ or (NHC)Ni⁰ catalysts, which are commonly employed in many Ni/NHC-catalysed hydrofunctionalisation reactions. The pre-synthesised well-defined Ni⁰/NHC complex [(IMes)Ni(Cp)Cl] **146** was found to be equally efficacious in their hydroheteroarylation system.

The cyclopentadienyl (Cp) ring of the precatalyst was postulated to serve as an internal mild base for the deprotonation of



the azolium salt (NHC proligand), thereby facilitating Ni-NHC bond formation and circumventing the need for the addition of strong external bases that may cause the decomposition of base-sensitive substrates. Sodium formate was utilised as a mild reducing agent in the *in situ* reduction of the Ni(II)precatalyst to the commonly proposed active species, Ni(0). The widely accepted mechanism for this catalytic activation process⁸⁶ involves the initial ligand exchange of the formate anion with one Cp ligand. Facile decarboxylation yields a nickel-hydride intermediate 152. Reductive elimination of this nickel-hydride intermediate then forms the active NHC-Ni(0) species with concomitant release of cyclopentadiene. The proposed mechanism for the hydroarylation, as put forth by the authors, is in agreement with previous works,^{73,74} proceeding through a sequential oxidative addition of the C(sp²)-H bond, regioselective hydronickelation and reductive elimination.

Activated vinylarenes and norbornenes serve as representative substrates for the scope of alkenes. A wide variety of heterocycles including thiophene **149** and furans **151** were



well-tolerated. Notably, pharmaceutically relevant heterocycles such as theophylline **148** (*n*-propyl derivative of caffeine) and 1,2,4-triazolo[1,5-*a*]pyrimidine derivatives (proposed surrogates of purines) **150** were demonstrated to proceed efficiently under either reaction conditions.

The Hartwig group has made significant contributions to the field of alkene activation *via* hydroarylation, with regard to the utilisation of non-activated aliphatic alkenes. In 2014, they achieved initial success through the reaction of unactivated alkenes with electron-poor trifluoromethyl-substituted arenes, obtaining anti-Markovnikov alkylarene products with remarkable regioselectivity (Scheme 28).⁸⁷ Optimal results were obtained using the bulky IPr **1f** ligand as the NHC ligand in the bis(NHC)Ni⁰ precatalyst employed for reaction. The regioselectivity of the terminal arylated product appears to be primarily driven by steric factors, with the aryl group being introduced at the least hindered position, untethered by any directing moieties.

The reaction proved to be efficacious with a diverse array of substituted olefins like vinvlsilanes (155) and allylic amines (153). Optically pure secondary siloxyethers also reacted smoothly, yielding the product (154) without erosion of stereochemistry. Notably, internal alkenes (such as (E)-2-octene) also underwent hydroarylation to furnish terminal arylated products (156), implying that an alkene isomerisation occurs prior to hydroarylation. Deuterium labelling studies indicates that the isomerisation does not follow a chain-walking pathway involving proton-transfer steps with the arene. Although the precise mechanism for alkene isomerisation remains elusive, it is suggested that the nickel-catalyst selectively coordinates to the more reactive terminal alkene, leading to hydroarylation. As such, isomeric mixtures of internal alkenes could still be utilised in the reaction since they would eventually converge to the terminal alkenes from the equilibrium mixture.

Mechanistic studies and DFT calculations have shed light on a plausible reaction pathway (Scheme 28). Oxidative addition of the aryl C-H bond followed by hydronickelation of the alkene from an unstable, intermediary nickel-hydride species was computed to be much less favourable than a concerted ligandto-ligand hydrogen transfer (LLHT) pathway (Scheme 9a). Coordination of the arene to bis(alkene)-coordinated resting state nickel 157 forms the active complex 158, which undergoes a reversible LLHT to afford the alkylnickel-aryl intermediate 159 that is stabilised by a β -C-H agnostic interaction. **159** subsequently undergoes a two-step isomerisation to form a T-shaped intermediate 160, in which the hydrocarbyl groups are now positioned mutually cis to each other. Turnover-limiting reductive elimination forms the desired $C(sp^2)$ - $C(sp^3)$ bond, yielding the productcoordinated nickel complex 161. Release of the linear hydroarylation product through ligand exchange with incoming substrates regenerates the active nickel species 158. It is noteworthy that the excellent linear selectivity observed results from the lower energy barrier for reductive elimination from the primary alkylnickel complex, rather than the absolute difference in computed free energy between the linear or branched alkylnickel complex.

Later in 2015, Hartwig and co-workers expanded upon their previous findings,⁸⁷ by broadening the substrate scope beyond just trifluoro-substituted arenes. In that work, utilising a more sterically bulky IPr^{Me} **162** or IPr* ligand, they achieved regio-selective hydroheteroarylation of unactivated alkenes with a wide range of heterocycles (Scheme 29)⁸⁸ under solvent-free



Scheme 28 NHC/Ni-catalysed linear-selective hydroarylation of unactivated terminal and internal alkenes with trifluoromethyl arenes.⁸⁷

conditions using the $Ni(COD)_2$ precatalyst and the free carbene ligand. The linear alkylheteroarene product was generally obtained in high yields with high anti-Markovnikov selectivity. Alternatively, the use of a pre-synthesised IPr*OMe-ligated Nibenzene complex 163 was also efficient, sometimes yielding better results. Effective hydroheteroarylation was observed with benzofurans, furans and indoles as heterocyclic coupling partners. Boronate esters (164) and potential coordinating esters (165) were all well-tolerated and led to the formation of the C_2 -selective linear hydroheteroarylation product. With respect to the scope of alkenes, a polar amine group at the allylic position gave moderate yields of the linear product (166). Non-conjugated 1,5-dienes 167 also underwent chemoselective hydroheteroarylation at the less substituted terminal olefin unit, leaving the internal cyclic olefin unreacted. The postulated reaction pathway involves an initial LLHT between the



Scheme 29 Ni/NHC-catalysed anti-Markovnikov hydroheteroarylation of unactivated alkenes with N- and O-heterocycles.⁸⁸

bound arene and alkene facilitated by the NHC/Ni catalyst, followed by isomerisation and *cis*-reductive elimination, as previously described.⁸⁷

Going beyond selective hydroarylation of challenging unreactive aliphatic alkenes, Hartwig and co-workers extended their work by developing a Ni/NHC catalysed linear-selective hydroarylation of unactivated alkenes with unactivated arene substrates, facilitated by non-covalent interactions (Scheme 30).⁸⁹ The use of the benzene-bound NHC-ligated nickel complex [NHC–Ni(η^6 -C₆H₆)], with NHC being ^{m-Xyl}IPr*^{OMe} **168**, as a precatalyst led to excellent yields of the anti-Markovnikov hydroarylation product, with a linear/branched selectivity of >50:1 or no detectable branched isomer. Basic additives, such as NaH and Na(acac), was crucial for high turnover numbers especially at low catalyst loading, presumably removing traces of water which could significantly affect the reaction.

Electron-neutral, unactivated arenes such as benzene participated well in the hydroarylation reaction. Sterically hindered alkene bearing a quaternary carbon at the α -position (169) or vinlysiloxanes (170) were also tolerated. DFT calculations provided further insight into the mechanism, consistent with



Scheme 30 Ni/NHC catalysed anti-Markovnikov hydroarylation of unactivated alkenes with unactivated arenes facilitated by non-covalent interactions.⁸⁹

previous findings.⁸⁷ Reversible, direct transfer of the C–H bond of the coordinated arene and ligand in **172** (in a concerted LLHT process) was computationally lower in energy than oxidative addition of the aryl C–H bond. The resulting linear, agostic C β –H stabilised alkylnickel-aryl complex **173** then isomerises to the *cis*-aryl-alkyl nickel complex **174**. Ratedetermining reductive elimination of **174** then forms a new carbon–carbon bond in the linear alkylarene product.

Unexpectedly, the rate control for reductive elimination under their reaction system goes beyond simple steric control within the transition state, as conventionally thought, despite the use of an extremely bulky NHC ligand. Energy decomposition analysis reveals a different reason for rate acceleration in reductive elimination: extensive non-covalent interactions within the coordination sphere of the catalyst. While Pauli's repulsion factor due to steric effects, cannot be ignored, the stabilising electrostatic interactions between the multiple aryl groups of the NHC and favourable London dispersion interactions resulting from the peripheral methyl groups in the ^{m-Xyl}IPr*^{OMe} ligand (Scheme 30) outweigh the energy cost due to steric influence, thus explaining the high activity of the catalytic system. The significant stabilisation by non-covalent interactions was also corroborated by visualisation using a noncovalent interaction (NCI) plot.⁹⁰

Recently, Montgomery and Chen contributed to an underdeveloped area of nickel catalysis, with their report on the Ni/ NHC-catalysed intermolecular enantioselective hydroarylation of alkenes (Scheme 31).⁹¹ This novel approach is distinct from Shi's prior work that employed an SIPE/ANIPE-type ligand⁷⁵ where the stereochemical induction was provided by the chiral 1-phenethyl substituents on the *N*-aryl ring. Instead, the authors used a sterically hindered NHC **176** bearing a



Scheme 31 Ni/NHC-catalysed intermolecular enantioselective hydroheteroarylation of alkenes. $^{91}\,$

2,4,6-tri*iso*propyl *N*-aryl ring, with a chiral backbone derived from chiral diamines. This chiral NHC ligand provided appreciable enantioinduction while displaying high reactivity with a variety of heterocycles. Paired with an alternative condition that employs a discrete C_2 -symmetric 1,5-hexadiene-supported chiral NHC–Ni⁰ complex 177 (instead of generating the active NHC–Ni *in situ*), this combination allowed for increased functional group tolerance and enhanced the catalytic activity with certain heterocyclic substrates.

Most coupling reactions with norbornene as the alkene partner gave the desired product in good yields, as a single exocyclic diastereomer in high enantiomeric ratios. Hydroarylation of norbornene with 1,2,4-triazole (178) surprisingly afforded the C5-alkylated product with high regio- and enantio-selectivity, despite the greater steric hindrance (closer to the benzyl substituent). Notable N-heterocycles like caffeine (179) were also highly efficient in the reaction. With condition B employing the presynthesised NHC–Ni(1,5-hexadiene) catalyst, the use of previously unreactive substrates under condition A could now be feasible, such as chloro-substituted benzoxazole (180) and indoles bearing electron-withdrawing substituents (*i.e.*, esters) (181).

From the preliminary mechanistic studies and analogies with previous computations,^{87–89} the authors proposed a similar reaction pathway. Initiation of the catalytic cycle involves ligand exchange with the alkene and heteroarene substrate to form the active Ni species, which undergoes reversible, *enantio*-determining ligand-to-ligand hydrogen transfer (LLHT). Facile isomerisation to a *cis*-aryl-alkyl T-shaped intermediate preceding the rate-limiting reductive elimination delivers the final enantioenriched *exo*-selective hydroarylation product.

Significant advancements have been made in the field of intramolecular hydroarylation enabled by Ni/NHC catalysts. In 2015, Cramer and Donets reported NHC ligand-controlled regiodivergent 1,6-annulation of 2-pyridone derivatives, through a selective intramolecular olefin hydroarylation (Scheme 32). The simple cyclooctadiene (COD) ligand favours an exo-selective cyclisation while bulky IPr NHC (1f) favours the endo-selective mode via formation of the alkylnickel at the less hindered end of the alkene. This approach offers an alternative way for controlling the cyclisation mode apart from the ring size, which is highly dependent on substrates. Vinylfuran-containing pyridones (184) underwent endocyclisation smoothly (with exo-184' formed preferentially in the absence of NHC). The desired *cis*-fused bicyclic ring (182) was also obtained from a cyclopentene-tethered pyridone in good yields. Medium-sized heterocycles (183) can also be efficiently accessed under their protocol. endo-Selectivity is purely ligand-controlled, presumably due to the favoured formation of the less hindered nickelacycle intermediate (Scheme 32). Initial enantioselective endo-cyclisation attempts using the chiral NHC ligand 185 based on the isoquinoline framework were successful, yielding the desired pyridone (-)-186 in a modest enantiomeric ratio.

With the demonstration of the Ni-catalysed regiodivergent annulation of pyridines by control of the ancillary NHC ligand,⁹² Cramer and co-workers recently improved on the control of enantioselectivity for this class of reactions (Scheme 33).



Scheme 32 NHC-ligand controlled regiodivergent Ni-catalysed annulation of pyridones.⁹²

A modular class of chiral NHC **187**, which mimics the properties of the privileged achiral IPr **1f** ligand, was utilised for the intramolecular enantioselective *endo*-cyclisation of 2-and 4-pyridones with their tethered olefin unit. The most effective chiral ligand was a modified C_2 -symmetric carbene first reported by Gawley,⁹⁴ with a bis(imino)acenaphthene backbone.⁹⁵ Enantiocontrol of the reaction was enhanced through the introduction of 3,5-xylyl groups to the *N*-aryl side arms of the NHC. This class of ligand was termed, in the same year, by Shi as an ANIPE-type⁷⁶ ligand.

The generality of the reaction was assessed using various substituted 2- and 4-pyridones and was found to be tolerant to heterocyclic substituents such as furans (**189**). The reaction was also amenable to the synthesis of medium-sized 7-membered heterocycles (**188**). The chiral and steric influence of the backbone modifications and the *N*-aryl arms of the chiral NHC was studied in-depth, with the analysis of analogous NHC–Ni and Au complexes. Preliminary data suggest a degree of



Scheme 33 Ni/NHC-catalysed enantioselective annulation of 2- and 4-pyridones via intramolecular olefin hydroarylation.⁹³

preorganisation of the flanking bulky *N*-aryl groups by π stacking interactions with the acenaphthene backbone, reducing the overall buried volume $%V_{\text{bur}}$ and enhancing the accessibility of the C_2 -symmetric binding pocket. This, in turn, improves the catalytic performance and enantiocontrol provided by the chiral ligand.

In 2019, Shi and co-workers reported an unprecedented Ni/NHC-catalysed enantioselective C-H alkylation reaction of alkene-tethered polyfluoroarenes through an *endo*-selective intramolecular hydroarylation process (Scheme 34).⁹⁶ Chiral fluorotetralin derivatives–which are important bioisosteric analogues of tetralin for drug design⁹⁷–were efficiently synthesised under this protocol, with chemoselective activation of the C-H bond over the C-F bond. Competitive side reactions such as hydrodefluorination and alkene chain-walking were also suppressed.

The most effective chiral ligands were found to be the bulky 3,5-Xyl ANIPE or SIPE-type NHC ligands, which led to high yields and enantioselectivity of the fluorotetralin product. The large steric bulk of the NHC was proposed to play a role in accelerating the reductive elimination of the nickelacycle **199** and favouring the *endo*-cyclisation pathway. Selective activation of the C-H bond over the C-F bond was also attributed to the monomeric Ni/NHC active catalyst as the bulky NHC disfavours aggregation.

The method was successfully applied to difluro-(196), trifluoro-(197) and tetrafluorobenzenes (193–195), even on a gram scale, showcasing the general practicality of their method. Notably, substituents bearing acidic protons, (phenol in 194),



Scheme 34 Ni/NHC-catalysed asymmetric intramolecular hydroarylation of olefin-tethered fluoroarenes.⁹⁶

pharmaceutically relevant heterocycles like indole **193** and 1,3dienes **195** were all well tolerated, furnishing the product with high levels of enantiocontrol. Preliminary mechanistic studies suggest a direct ligand-to-ligand hydrogen transfer (LLHT) from the fluoroarene to the alkene within the coordinated complex **198**, under the influence of the chiral environment of the NHC– Ni complex. The *endo*-cyclised nickelacycle **199** that was formed underwent reductive elimination to yield the chiral fluorotetralin product.

By employing the same bulky chiral (R,R,R,R)-SIPE NHC ligand, Shi and co-workers, in successive years, developed the

Ni/Lewis acid catalysed asymmetric C–H functionalisation of alkene-tethered pyridines⁹⁸ and pyridones⁹⁹ through an intramolecular hydroarylation process, affording the corresponding annulated chiral N-heterocycles from racemic building blocks (Scheme 35). From their preliminary mechanistic investigations involving deuterium labelling studies, a similar operating mechanism seems to be the case for both reactions. Dual coordination of the substrate with the aluminium-based Lewis acid additive and the Ni/NHC active catalyst activates the substrate, forming a Ni/Al bimetallic complex **206**. Oxidative addition of Ni(0) into the C–H/D bond forms a tentative Ni–H



Scheme 35 Ni/SIPE-catalysed intramolecular enantioselective hydroarylation of alkene-tethered pyridines⁹⁸ and pyridones.⁹⁹

species **207**, which subsequently undergoes enantioselective anti-Markovnikov hydrometallation to form the sevenmembered nickelacycle intermediate **208**. The observed regioselectivity could be influenced by the combined steric effect from the bulky NHC ligand and the Lewis acid. Final reductive elimination yields the enantioenriched (*R*)-N-heterocycle and regenerates the active NHC/Ni⁰ species. The strongly electrondonating SIPE ligand is believed to facilitate the process of C–H oxidative addition, with enantiocontrol arising from the efficient *enantio*-discrimination of one face for alkene insertion, owing to the steric repulsion of the pyridone/pyridine moiety and the phenyl rings of the 1-phenethyl arm on SIPE.

Endo-Selective annulation of C3-tethered alkenes with trisubstituted cyclopentene and heterocyclic enamines successfully results in the formation of the corresponding tricyclic (**200**) and pyrrole-containing (**201**) products with high enantioselectivity. Reaction with pyridine C2-tethered alkenes was also viable, affording enantioenriched 5,6,7,8-tetrahydroquinolines in excellent yield. In the annulation of pyridones, trifluoromethyl substituted pyridone cores (**203**) and heterocyclic functional groups like thiophene (**205**) were well-tolerated. Mediumsized heterocycles (**204**) could also be accessed in good yields and with high enantioselectivity under this regime.

Cramer and co-workers took the concept of previously reported enantioselective C–H functionalisation of various heterocycles *via* hydroarylation,^{92,93,100} and applied it to biologically relevant electron-rich heterocycles like pyrroles and indoles. In 2019, they reported the Ni/NHC-catalysed intermolecular enantioselective hydroarylation of alkene-tethered pyrroles or indoles, with a high anti-Markovnikov selectivity despite the absence of a directing group¹⁰⁰ (Scheme 36). Usage of a highly sterically demanding 3,5-di-*tert*-butylphenyl substituted SIPE-type ligand **209** proved to be superior in delivering the *endo*-cyclised tricyclic tetrahydropyridyl indole scaffolds in high yields and with high enantioselectivities.

Base-sensitive boronate ester (210) functionality was tolerated and remained intact after the reaction. Other pharmaceutically relevant azaheterocycles like pyrrolopyrimidine (211) and pyrroles (213) also participated well in the reaction. Indole bearing tethered vinylsilanes (212) also reacted successfully, albeit with a lower enantioselectivity, presumably due to the steric influence of the bulky trimethylsilyl group at the newly formed stereogenic centre.

In 2021, Ye and co-workers reported the construction of medium sized 7-membered heterocycles *via* an NHC/Ni-Lewis catalysed C7-H bond cyclisation reaction of benzimidazoles with a tethered alkene moiety¹⁰¹ (Scheme 37). Formally an intramolecular hydroarylation process, enantioenriched tricyclic imiazdoles can be achieved, with exclusive *endo*-selectivity in the cyclisation mode, despite the challenging enantiocontrol due to the flexible large ring transition state. The synergistic (NHC)Ni–Al bimetallic system responsible for the success of this reaction comes from the bulky chiral ANIPE type ligand **214** and trimethylaluminum Lewis acid.

The reaction is quite general, where various aryl or alkyl alkenes were well compatible under the reaction conditions,



with the latter resulting in a poorer ee of the seven-membered cyclised product (cf. 215 and 216), presumably due to greater structural flexibility. Racemic version of the reaction could also be easily realised with the use of the achiral IPr 1f ligand, and the utility of the reaction was demonstrated to access bioactive molecules like a translocator protein inhibitor (217) from imdazol-2-one. Preliminary mechanistic studies provide some insight into a possible mechanism (Scheme 37). Activation of the benzimidazole substrate by AlMe₃ and coordination of the activated substrate with NHC-Ni active species forms 218. This facilitates the subsequent rate-determining oxidative addition of the C7-H bond, forming the alkene-coordinated nickelhydride intermediate 219. Governed by the steric and chiral environment of the ANIPE ligand, an irreversible endo-type alkene migratory insertion occurs, forming the 8-membered nickelacycle 220 with the stereochemistry being determined. Lastly, stereoretentive reductive elimination releases the final tricyclic product after dissociation with the aluminium Lewis acid, to initiate the next cycle.

In 2021, Koh and co-workers developed a new regime for NHC-Ni catalysed Markovnikov-selective hydroarylation-(alkenylation) of olefins, which operates through a novel pathway involving sequential carbonickelation followed by hydrogenation⁵⁷ (Scheme 38). This reversed order is distinct from the conventional hydroarylation pathway, with hydronickelation preceding carbofunctionalisation, predominantly delivering the anti-Markovnikov product especially in reactions with unactivated aliphatic alkenes due to the absence of the adjacent electronic π -stabilisation from the alkylnickel intermediate. The authors used a dimeric IPr-Ni(1) precatalyst and NaO*i*Pr as the hydride-source, which furnished the branchedselective hydroarylation product in high yields. Excess alkoxide



Scheme 37 (NHC)Ni/Lewis acid catalysed synthesis of tricyclic imidazoles *via* intramolecular enantioselective hydroarylation of alkenetethered benzimidazole.¹⁰¹

base was also found to be crucial for suppressing the undesired Heck-type reaction.

A series of investigations were carried out to gain a deeper understanding of the underlying mechanisms. In a series of radical clock experiments, no cyclopropyl ring-ruptured or radical annulation side-products were observed, supporting the proposition that no long-lived radical intermediate species were present, even if a Ni(1) complex was employed. A plausible non-radical pathway was proposed by the authors (Scheme 38). The active monomeric Ni(1)-isopropoxide intermediate **226** is formed through the alkoxide assisted dissociation of the dimeric Ni(1) precatalyst **25**. Oxidative addition of the aryl/ alkenyl triflate electrophile affords the aryl-Ni(III) species **227**,



Scheme 38 Ni/NHC-catalysed regioselective hydrocarbofunctionalisation of alkenes *via* sequential carbonickelation – hydrogenation.⁵⁷

which undergoes Markovnikov-selective carbonickelation across the incoming olefin substrate, forming the least hindered primary alkylnickel complex 228. The observed regioselectivity in the insertion is likely attributed to the steric influence of the sizeable IPr ligand, where steric repulsion between the IPr ligand and present alkene substituent is minimised. Ligand exchange with excess NaOiPr forms the Ni-bis(isopropoxide) 229. Dehydrogenation of the ligated isopropoxide ligand through β -H elimination as facilitated by the adjacent CB-H agostic interaction yields the nickel-hydride intermediate 230, with concomitant release of acetone as a by-product. Final reductive elimination releases the Markovnikov-selective hydroarylation/alkenylation product, regenerating 226. Although a tentative proposal of a Ni(I)/Ni(III) operating catalytic pathway was presented, the authors acknowledge that a $Ni(0)/Ni(\pi)$ pathway cannot be entirely ruled out. The main emphasis of the mechanism and the observed regiocontrol

lies in the preference for carbonickelation over the adventitious hydronickelation, contrary to many previous reports.¹⁰²

The reaction has been shown to be versatile, accommodating a variety of functionalities, including heterocycles like pyrazoles (222). Challenging quaternary carbon-centres can also be efficiently accessed from 1,1-disubstituted alkenes (224). Notably, the cyclopropane moiety of 224 was not disrupted after the reaction with no traces of radical-induced ring ruptured products. Furthermore, the reaction is effective with unactivated alkenes like 2,5-dihydrofurans (225), and alkenyltriflates could also be utilised as complementary electrophiles to afford the corresponding hydroalkenylation product in good yields and with high selectivity. The utility of the reaction was further demonstrated by the efficient reaction with alkenes derived from bioactive molecules (221).

In continuation of their previous work,⁵⁷ Koh and co-workers have developed a versatile enantioselective nickel-catalysed strategy for olefin hydrocarbofunctionalisation and dicarbofunctionalisation¹⁰³ (Scheme 39). This three-component reaction involves the union of an activated olefin or diene, an organotriflate electrophile and a metal alkoxide as a hydride donor in the presence of a sterically encumbered chiral NHC-Ni(0) catalyst (where NHC = (R,R,R,R)-IPE **231**). This work allows for practical access to the corresponding high value enantioenriched secondary, tertiary, or quaternary stereogenic centres. Substitution of the metal alkoxide with an organometallic reagent leads to the formation of useful dicarbofunctionalisation adducts through the installation of two different carbogenic motifs across the C=C bond.

Electron paramagnetic resonance spectroscopy reveals the absence of paramagnetic radical species in the reaction system, suggesting that it follows a Ni(0)/Ni(II) pathway. Initial oxidative addition of the $C(sp^2)$ aryl or alkenyltriflate electrophile to NHC-Ni(0) forms the chiral arylnickel species 241. Regio- and enantio-selective carbonickelation across the π -complexed alkene substrate forms the less hindered primary alkylnickel intermediate 242. Minimisation of the unfavourable steric interactions between the olefinic substituents and the β arms of the NHC in the chiral environment governs the regio- and enantio-control for the insertion (an unfavoured transition state leading to the minor (S)-enantiomer). Ligand exchange with the excess alkoxide forms the Ni(alkoxide) species 243, which participates in β -H elimination to give the nickelhydride intermediate 244, with the concomitant expulsion of acetone. The formation of the C-H bond occurs following a reductive elimination reaction. Since the enantio-determining C-C bond forming step occurs during carbonickelation, appropriate interception of intermediate 242 with a carbon-based nucleophile instead of an alkoxide led to the formation of dicarbofunctionalisation products.

The catalytic regime is highly general and compatible with even highly hindered substrates like 1,1-disubstituted alkenes, leading to the formation of quaternary stereocentres (233, 236 and 240) with excellent regio- and enantio-selectivity. Heterocyclic substituents, such as Lewis basic indoles (238), benzofurans (240) and pyridines (234) were well-tolerated. Chemoselective





Scheme 39 Generation of tri- and tetra-substituted stereocentres by NHC/Ni-catalysed enantioselective olefin cross-coupling.¹⁰³



Scheme 40 NHC–Ni/Al catalysed *para*-selective C–H alkylation of benzamides and aromatic ketones *via* hydroarylation with alkenes.¹⁰⁴

activation of triflates instead of bromides (233) was also highlighted, providing the potential for orthogonal functionalisation. Medium-sized alkenyltriflates (237) participate well in the hydroalkenylation, and dicarbofunctionalisation adducts can be efficiently converted from various styrene derivatives or dienes with high enantiocontrol, exemplified by various diarylation (238 and 239) products and a notable skipped 1,5-diene arylalkenylation product 240. Utility of the reaction was highlighted with the synthesis of an anti-cancer drug candidate product (232) without the observation of any undesired allylic rearrangement.

Having previously developed the *para*-C–H alkylation of pyridine derivatives using NHC–Ni/Al cooperative catalysis,⁷⁴ Nakao and co-workers applied the concept to the direct arene C–H activation of benzamides and aromatic ketones, in a formal hydroarylation reaction with exogenous alkenes¹⁰⁴ (Scheme 40). The use of a sterically encumbered C_2 -symmetric NHC ligand with 3,5-xylyl substituted *N*-aryl arms in conjunction with the bulky MAD cocatalyst resulted in exceptional *para*-C–H and anti-Markovnikov site-selectivity, aided by the combination of electronic activation and steric repulsive forces.

Bulky olefin substrates such as vinylcyclohexanes (248) and methylbis(trimethylsilyloxy)vinylsilane (247, 249) were well-tolerated, yielding the corresponding *para*-alkylated benza-mides. The 1,2-disubstituted C=C bond of norbornene (246) was also reacted under the protocol even though it was less efficient. Enolisable alkylaryl ketones (249) also underwent reaction smoothly to deliver the C4-alkylated arene.

Theoretical calculations using DFT are in agreement with the previously divulged mechanism,⁸⁷ involving a concerted

LLHT as the key rate-determining C–H activation step. Calculations using the model AlMe₃ reveal the mode of rateacceleration by the Lewis acid. Benzamide coordinating to AlMe₃ was calculated to have a lower LUMO (σ^* of C–H and aromatic ring) energy than the uncoordinated substrate. AlMe₃ also enhances the selectivity for the *para*-position due to the stronger repulsive interactions with *N*-aryl substituents of the NHC in the transition state leading to the formation of the *meta*-functionalised product.

With successful para-C-H alkylation of arenes with selectivity governed by both steric and electronic factors,¹⁰⁴ Nakao and colleagues sought to achieve a site-selective linear C-H functionalisation protocol by variation in the nature of NHC ligands used. As previously established, electron-poor C-H bonds are more reactive due to the ease of activation under a concerted LLHT mechanism. Using anilides as a model substrate, one would expect the meta-C-H bond of anilides to be the most electron poor and electronically activated, assuming negligible steric effects. This presents the possibility for selective and switchable site-functionalisation, which they were able to achieve. The use of a smaller, less sterically demanding IPr^{Me} ligand 162 (condition A, Scheme 41) enables meta-selective functionalisation, while the bulkier ligand 245 (condition B, Scheme 41) sterically favours para-selective functionalisation. Vinylsilanes (251, 253) or vinylcyclohexanes (250, 252) were identified as suitable alkene substrates for this formal hydroarylation reaction. Significant enhancement of para-selectivity was achieved with a bulkier N-alkyl substituent R² on the anilide substrate (R^2 = sec-isopentyl for 252 or isopropyl for 253). Formation of the meta-isomer could be disfavoured as a result of steric hindrance and the strong steric repulsion between R², the sizeable MAD Lewis acid and the NHC ligand.

Recently, Chen and co-workers developed a novel synthetic route to access biologically and pharmaceutically relevant monoterpenoids (Scheme 42).¹⁰⁶ Their atom-economic regime involves an NHC/Ni-catalysed asymmetric heteroarylative cyclotelomerisation of isoprenes (a commercially available bulk chemical), to generate cyclic monoterpene derivatives bearing quaternary carbon stereogenic centres with high enantioselectivity. The best enantioselectivity was achieved with the use of a bulky C_2 -symmetric (R,R,R,R)-ANIPE NHC ligand (254) with a structurally rigid acenaphthene backbone.

Various naturally occurring and medicinally important heterocycles were amendable to the procedure, delivering the telomer in excellent enantioselectivity. N^7 -benzyl theophylline (255), an analogue of caffeine, participated well in the reaction. Adenine, a nucleobase found in nucleic acids DNA and RNA, is also a viable substrate for this asymmetric telomerisation reaction. Dimethylamino-substituted adenine (256) was converted to the corresponding product in excellent yield and with high enantioselectivity. N-Heterocycles such as benzimidazole (257) which can be found in numerous clinical drugs are also tolerated under their regime. In addition to isoprene, other terpenes such as sesquiterpene farnesene with trisubstituted olefin moieties also reacted efficiently with caffeine to deliver the expected product (258).



Scheme 41 NHC–Ni/Al catalysed site-selective linear alkylation of anilides.¹⁰⁵

The key components of their asymmetric heteroarylative cyclotelomerisation reaction include the cyclisation of isoprene and the C–H alkylation of the heterocycle (*via* a hydrohetero-arylation process). Extensive mechanistic investigations were conducted to unveil the sequence of events, with the conclusion that the enantioselective dimerisation of isoprene precedes the C–H bond functionalisation of the heterocyclic coupling partner. Initial coordination of isoprene with the active NHC–Ni⁰ species forms the π -complex **259**, which leads to the formation of the nickelacyclopentene intermediate **260** after oxidative cyclometallation. Subsequent coordination with a second molecule of isoprene forms the complex **261**, which undergoes regio- and enantio-selective carbonickelation (with a depicted transition state) to furnish a seven-membered nickelacycle **263**.



Scheme 42 NHC/Ni-catalysed asymmetric heteroarylative cyclotelomerisation of isoprene. 106

The quaternary carbocentre formed is already stereodefined at this stage. The simultaneous control of the regio- and enantio-selectivity of the migratory insertion is due to the encumbered chiral environment provided by the ANIPE ligand. Reductive elimination of **263** then releases the isoprene-dimer which enters a separate catalytic cycle. In the second cycle, C–H bond activation of the heterocycle *via* oxidative addition forms the nickel(n)-hydride species **264**, which coordinates with the isoprene-dimer to yield a π -complex **265**. **265** then undergoes an anti-Markovnikov selective hydronickelation, under the steric control of the bulky NHC, resulting in the formation of

a primary alkylnickel intermediate **266**. Reductive elimination of **266** affords the desired cyclised terpenoid product and regenerates the active NHC–Ni⁰ catalyst.

3.3 Hydroboration of alkenes

Organoboronates are known for their stability, versatility, and ease of diversification, making them a popular alternative to sensitive organometallic reagents. These compounds can be transformed into a variety of synthetically useful motifs using well-established C–B to C–X (such as X = C, N, O or halide) transformations.^{107,108} A highly efficient method for synthesising aliphatic organoboron compounds involves employing catalytic hydroboration or protoboration of alkenes,¹⁰⁹ which has been achieved with some success under nickel-catalysed systems.¹¹⁰ Although NHCs are widely utilised as ligands in many nickel-catalysed hydro(hetero)arylation and hydroalkenylation reactions, their application in the hydroboration of alkenes is more limited.

In 2016, Schomaker and co-workers first reported the Markovnikov-selective hydroboration of alkenes with a heteroleptic NHC/phosphine nickel catalyst 267.¹¹¹ Benzyl boronic esters were obtained with excellent branched Markovnikov selectivity and in moderate yields (Scheme 43). Both electron-rich (*ortho*-methoxy, 270) and electron-deficient (*meta*-methoxy, 269) styrenes were well-tolerated. Notably, internal 1,2-disubstituted alkenes (268) also performed well under the regime. However, 4-vinylpyridine (271) proved to be ineffective, likely due to the deactivation of the catalyst by the Lewis basic nitrogen atom.

The authors proposed a Ni⁰/Ni^{II} catalytic cycle for their hydroboration protocol. Initial reduction of the Ni^{II} precatalyst **267** by HBpin generates the 14 e^{-} Ni⁰ active catalytic species 269. Dissociation of the existing tricyclohexylphosphine ligand (facilitated by the strong trans effect of the electron-donating NHC ligand) followed by the oxidative addition into the B-H bond of pinacolborane yields the boryl-Ni-hydride intermediate 270. Addition of external PCy₃ proved to be highly detrimental to the reaction, supporting the notion for the dissociation of the phosphine ligand. The incoming styrene substrate then proceeds to form a π -complex with the coordinatively unsaturated boryl-Ni-hydride complex 270 to give 271. Markovnikov-selective migratory insertion of the alkene into the Ni-H bond forms the benzylic-Ni intermediate 272, which is presumably stabilised by conjugation with the π -aromatic system. By exploiting the steric properties of NHC ligands, the authors discovered that the use a bulkier IPr* NHC ligand can disfavour the formation of the π -benzylic-Ni intermediate 272, due to a greater steric repulsion between the aryl substrate and bulkier NHC. This results in the production of a higher proportion of linear boronic esters, despite the decreased overall reactivity. Final reductive elimination of 272 then delivers the final Markovnikov product, regenerating the active Ni⁰ catalyst to close the catalytic cycle.

Later in 2020, Ritleg and co-workers developed an anti-Markovnikov hydroboration reaction of alkenes with a halfsandwich type Ni/NHC complex [Ni(Cp*)Cl(IMes)] 273, using catecholborane (HBCat) as the boron source in the presence of catalytic potassium *tert*-butoxide (Scheme 44).





Electron-rich styrenes were all fully converted with high linear, anti-Markovnikov selectivity 275. In comparison, electron-poor styrenes underwent hydroboration with poorer yields and selectivity 276. Surprisingly, a non-aryl derivative like vinylcyclohexane was still effective (274). Limitations of the reported regime include 1,2-disubstituted internal alkenes which showed no conversion (277 was not observed after subjecting (E)-stilbene to their reaction protocol).

Detailed investigation into the mechanism revealed that the nickel-hydride complex [NiCp*H(IMes)] (278) is the key catalytic precursor which would be reduced with concomitant release of H_2 gas to form the active Ni^I species, [NiCp*(IMes)] (279). It was proposed that the reaction of [Ni(Cp*)Cl(IMes)] with



Scheme 44 Anti-Markovnikov selective hydroboration of alkenes with the [Ni(Cp*)Cl(IMes)] complex. 112

KOtBu and HBcat generates the initial nickel-hydride complex **278** *via* a σ -bond metathesis mechanism. The active species **279** forms an initial agostic intermediate **280** with HBCat, which then associates with the styrene substrate to furnish **281**. An energetically favourable homolytic fission of the B–H bond followed by a radical Giese addition to the styrene forms back the nickel-hydride intermediate **278** and the linear borylated benzylic radical **282**. It is noteworthy to point out that the reaction was completely inhibited with the addition of one equivalent of a radical scavenger such as TEMPO. Final hydrogen atom abstraction from **278** releases the final hydroboration product, regenerating the active Ni^I species **279** to close the catalytic cycle.

In 2022, the Kennedy research group developed a class of mononuclear, anionic Ni⁰ complexes **283**, stabilised by a unique bidentate NHC-pyridone ligand which exclusively coordinates to



the Ni centre in a κ^2 -C.N coordination mode¹¹³ (Scheme 45). From single-crystal diffraction analysis, the $[K(18-crown-6)]^+$ counterion of this anionic nickelate was observed to be in a contact ion pair with the pyridone oxygen, allowing for monomeric isolation of the complex instead of multimetallic aggregation and clustering induced by the pyridone.¹¹³ The low-valent Ni⁰ core is highly electron-rich, with an asymmetric electronic and steric coordination environment around the nickel centre. In their work, this class of complexes was demonstrated to exhibit exceptional catalytic activity towards the hydroboration of alkenes (with the evaluation of only styrenes 285 and β methylstyrene 284), affording the desired hydroboration product in good yields and with excellent branched Markovnikov selectivity (Scheme 45). Related stoichiometric experiments reveal that the pyridone ligand plays a role in facilitating the transfer of the Lewis acidic HBpin (pinacolborane) reagent to the alkene through its involvement in the secondary coordination sphere.

3.4 Hydrosilylation of alkenes

Hydrosilylation of alkenes remains a powerful method for the formation of organosilicon compounds, with the addition of silicon hydrides across C—C p-bonds.¹¹⁴ It is widely used in industry to produce silane coupling agents and silicone polymers such as oils, rubbers, and resins. Various phosphine-ligated nickel complexes have been reported as hydrosilylation catalysts, but face the issue of undesirable site-selectivity and activity due to the competing side reactions like dehydrogenative silylation.¹¹⁴ Recently, ligand-free Ni-catalysed electrophilic hydrosilylation of alkenes was also developed.¹¹⁵ However, regioselective nickel-catalysed hydrosilylation of C—C bonds enabled by NHC ligands has been rather elusive and difficult to achieve, with only a few studies having been reported on this topic.

The first attempt of Ni/NHC-catalysed hydrosilylation of alkenes was demonstrated by Valerga and co-workers in 2012¹¹⁶ (Scheme 46). Various Ni^{II}(allyl)–NHC complex bearing bidentate 2-picolyl-imidazolylidene ligands (**286** and **287**) were found to show high activity for the hydrosilylation of simple styrene substrates. Different substituents on the NHC and allyl ligand



Scheme 46 Allyl-Ni^{II}(NHC)-catalysed hydrosilylation of styrenes.¹¹⁶

have a marked impact on the efficiency and selectivity of the hydrosilylation reaction. Only complex **286** was found to display activity when α -methylstyrene was used, giving the hydrosilylated product with complete anti-Markovnikov selectivity albeit in diminished yield. With regular styrene as the substrate, complex **287** affords excellent Markovnikov selectivity whereas complex **286** proved to be much less selective. Preliminary mechanistic studies show the presence of Ni–H and Ni–silyl intermediates, suggesting that the mechanism could proceed as illustrated in Schemes 8b and c, even though a full proposal was not provided.

In 2013, Montgomery and co-workers developed a method for the selective hydrosilylation of monosubstituted allenes using nickel–NHC catalysts.¹¹⁷ Building upon their initial discovery, in 2015 they expanded the scope of their research with the development of a regioselective and stereoselective hydrosilylation protocol for 1,3-disubstituted allenes¹¹⁸ (Scheme 47). The steric bulk of the NHC ligand played a key role in the successful synthesis of vinyl silane **292–294** and (*Z*)-alkenylsilanes **296–297** under the optimised regime, both of which were achieved with exceptional regio- and stereo-selectivity.

Deuterium-labelling studies were performed to further investigate the origin of regioselectivity and the reaction mechanism. The presence of only traces of cross-over products when using deuterated and regular silanes suggests that the mechanism does not involve the formation of nickel-hydride species that are not bonded to a silyl group, or *vice versa*. Extensive computational studies⁸⁴ have provided insight into a plausible mechanism for the reaction (Scheme 47).



Scheme 47 Ni/NHC-catalysed regioselective and stereoselective hydrosilylation of allenes. 117,118

Coordination of the silane and least sterically hindered face of the allene (thereby differentiating the different π -systems of the

unsymmetrical allenes) to the nickel centre forms the putative complex **298**. This is followed by the rate-determining step, in which a concerted Si–H bond oxidative addition – silyl insertion occurs to form the β -silyl nickel–hydride intermediate **299**. This step also determines the regioselectivity of the reaction with silyl-insertion being more thermodynamically favourable than hydride-insertion. The sterically bulky NHC ligand further reinforces this regiocontrol. Facile formation of the η^3 -allyl intermediate (**300**) occurs, with minimisation of allylic **1**,3strain where the R^S and R^L substituents prefer to be *trans* in the π -allyl intermediate. A final C–H reductive elimination, avoiding the formation of a more congested olefin in the transition state,¹¹⁹ delivers the (*Z*)-alkenylsilane product with excellent regioselectivity and (*Z*)-selectivity.

In 2022, Cook *et al.* reported a Ni⁰/NHC-catalysed branchedselective hydrosilylation reaction of styrenes with 2° , 3° and chlorosilanes¹²⁰ (Scheme 48). Industrially relevant silanes, particularly those substituted with chlorine or alkoxy groups, have not been extensively studied in the field of base-metal catalysis. Branched selective hydrosilylation of alkenes remains a challenging task, as it often faces the issue of lower selectivity and a narrower scope.¹²¹ (NHC)Ni⁰(alkene)₂ complexes, which can be easily prepared, have been shown to be effective in catalysing this transformation.

Extensive mechanistic investigations to probe into the reaction mechanism were carried out by the authors. Deuteriumlabelling experiments reveals minimal 2D incorporation at the benzylic position, hence supporting the Chalk-Harrod¹²² mechanism as the primary reaction pathway. In the Chalk-Harrod mechanism, insertion of the alkene into the nickelhydride (Ni-H) bond occurs preferentially to the insertion into the nickel-silyl (Ni-Si) bond. Radical-trapping experiments validate an ionic pathway as expected by the authors, rather than a one-electron radical pathway. In summary, initial alkene dissociation from the resting state catalyst 301 forms the 14electron active complex 306. Oxidative addition of silane into the complex forms the silyl-Ni-hydride intermediate 307. Subsequent migratory insertion of the coordinated styrene into the Ni-H bond affords a benzylic nickel species (if R = aryl) 308 with an open coordination site available for coordination with another alkene substrate. This results in the formation of the 16-electron complex 309 which undergoes facile C-Si reductive elimination, releasing the branched product while regenerating the active coordinatively unsaturated Ni species 306.

Electron-donating amine-containing styrenes (**302**) were well-tolerated, giving the product in moderate yield and with excellent branched selectivity. Terminal aliphatic alkenes are also viable, but with the selectivity being reversed to favour the linear-selective hydrosilylated product (**303**). This demonstrates the importance of an aryl substituent adjacent to the C=C bond for exclusive formation of the branched product, validating the involvement of a benzylic nickel intermediate **308**. Without adjacent π -stabilisation of the nickel intermediate following insertion, the regioselectivity was predominantly governed by the steric interactions between the NHC ligand and alkyl substituent, giving rise to a preference for the linear



Scheme 48 $\,$ Ni/NHC-catalysed hydrosilylation of alkenes with 2°, 3° and chlorosilanes. 120

product resulting from the less hindered transition state. Various electronically altered silanes such as diarylsilanes (**305**), 3° bis(alkoxy)-substituted silanes (**304**) and trichlorosilane (**303**) also reacted well, delivering the product with remarkable branched selectivity and in good yields.

3.5 Hydroamination of alkenes

Nitrogen-containing compounds play a vital role in biological systems and are also widely used in various industries such as bulk chemicals, specialty chemicals, and pharmaceuticals.¹²³ Among the plethora of reported methods for synthesising them, the direct construction of the desired C–N bond through the addition of an amine to an unsaturated C—C bond (known as hydroamination) is still of elegance and significant

importance in industries. This route provides a highly atomeconomical synthetic route starting with inexpensive and readily accessible alkenes. In addition to the challenge of regioselectivity, hydroamination reactions present an additional hurdle as they are only slightly exergonic due to a large entropic cost, particularly for the intermolecular variant.¹²⁴ To the best of our knowledge, the Ni/NHC-catalysed hydroamination reaction remains elusive, with only one reported instance of it.

The only work to date detailing the hydroamination of alkenes with Ni/NHC was reported by Ghosh and co-workers in 2010. Well-defined complexes of nickel with 1,2,4-triazole derived NHC as precatalysts (**310** and **311**) were effective in promoting the hydroamination of activated electron-deficient alkenes with secondary amines¹²⁵ (Scheme 49).



Scheme 49 Ni/NHC-catalysed hydroamination of activated alkenes with 2° amines. 125

Preliminary computational calculations to determine the nature of bonding of the (NHC)₂NiBr₂ complex were performed, which showed that the 1,2,4-triazole based NHC is a strong σ -donating ligand like the common imidazole based NHC ligands (Chart 1). An amine activation route which involves the oxidative addition of the N-H bond to the nickel centre was ruled out as the Ni-N (amido) bond energy is known to be very low. Extensive DFT calculations to probe into the mechanism for this reaction were published separately in 2017.¹²⁶ The putative mechanism with the least energy penalty is illustrated in Scheme 47. Ag-promoted ligand exchange with the bound halide ion forms the triflate coordinated Ni/NHC complex. Initial alkene coordination with the active [(NHC)₂Ni(OTf)]⁺ species forms 316. Anti-Markovnikov type attack of the secondary amine to the less substituted end of the alkene bound to Ni vields intermediate 317 which can be depicted in two resonance forms (α -C bound or N-bound if EWG = CN). Subsequent 1,3proton shuttle of the N-bound complex 317 assisted by another equivalent of the basic amine (via intermediate 318), furnished the final hydroamination product upon dissociation from 319, closing the catalytic cycle.

Under their protocol, pyrrolidine (312), morpholine (314), piperidine (313) and diethylamine (315) are examples of 2° amine substrates that have been successfully employed. Only activated alkenes adjacent to strongly electron-withdrawing groups like nitriles and esters proved to be suitable, delivering the product in moderate yields and with excellent anti-Markovnikov selectivity (312–315).

3.6 Hydroacylation and hydrocarbamoylation of alkenes

The use of transition metal catalysts to formally add an aldehyde C–H bond across unsaturated C=C bonds (termed alkene and alkyne hydroacylation) has become a popular and efficient method for synthesising ketones in recent years.¹²⁷

In the seminal work reported by Ogoshi and co-workers in 2012, they developed a Ni⁰/I^tBu catalysed intramolecular alkene hydroacylation reaction to access five/six-membered benzo-cyclic ketones¹²⁸ (Scheme 50). Mechanistic studies involving stoichiometric experiments¹²⁸ and DFT calculations¹²⁹ provide valuable insights into the plausible reaction mechanism which involves a dioxanickelacycle intermediate in the lowest energy pathway.

Coordination of the substrate to the Ni/l^tBu active catalyst forms the (η^2 : η^2 -enal)Ni⁰(I^tBu) complex **326**. Subsequent oxidative cyclisation of **326** forms the monomeric oxanickelacycle intermediate **327** which undergoes facile dimerisation to the dioxanickelacycle **328**. The dimer **328** was isolated in 84% yield in a separate stoichiometric experiment and was characterised with ¹H NMR and X-ray crystallography. A later DFT study¹²⁹ delved deeper into the mechanism and revealed that the facile dimerisation of **327**, which results in the formation of **328**, significantly reduces the free energy required for the subsequent rate-determining β -hydride elimination step that leads to the formation of intermediate **329**. This is followed by a formal reductive elimination step, which releases the final benzocyclic ketone product and regenerates the active Ni/I^tBu species. Their





Scheme 50 Ni/NHC-catalysed intramolecular hydroacylation of alkenes.¹²⁸

protocol presents a significant advantage, as their efficient hydroacylation process takes place through an oxanickelacycle intermediate without the occurrence of any decarbonylation. Decarbonylation of acyl-metal intermediates is a common undesired side reaction that leads to a decrease in atom efficiency and deactivation of the active catalyst through the coordination with carbon monoxide.¹³⁰

Fluorine and the silyl substituted *o*-allylbenzaldehyde substrate delivers the corresponding indanone product in high yields (**320–321**). **1**,1-Dimethyldisubstituted alkenes (**322** and **324**) and **1**,2-disubstituted alkenes (**323**) were also effective substrates. Notably, the reaction can also be extended to the synthesis of six-membered ring systems (324 and 325) with the desired 1-tetralone derivatives furnished in excellent yields.

Ogoshi and co-workers expanded on their previous development of Ni/NHC catalysed intramolecular hydroacylation of alkenes (Scheme 50) and reported the successful intramolecular coupling of alkenes and aldehydes in the presence of hydrosilanes under reducing conditions (Scheme 51). (η^6 toluene)Ni(SIPr) 330 which can be conveniently synthesised in a single step on a gram-scale was used as the precatalyst. This modified approach provides access to a variety of silylprotected 1-indanol derivatives with high syn-diastereoselectivity (between the silyl ether and methyl group), instead of benzocyclic ketones. Although a plausible mechanism has not been described in the literature, some insights into the mechanism can be inferred by drawing parallels to analogous mechanisms involving the reductive coupling of aldehydes and hydrosilanes, which proceed through oxanickelacycle intermediates.^{128,132} The diastereoselective formation of a similar oxanickelacycle (331) is believed to be facilitated by the Lewis acidic hydrosilane. Cleavage of the Ni-O bond of the oxanickelacycle by σ -bond metathesis with the Si-H bond of the hydrosilane forms a nickel-hydride intermediate (332) which undergoes reductive elimination to afford the desired product.

In 2016, Stanley and co-workers reported the intramolecular, *exo*-selective hydroacylation of *N*-allylindole-2-carboxaldehyde enabled with Ni⁰/IAd (IAd = 1,3-bis(1-adamantyl)imidazol-2ylidene)¹³³ (Scheme 52). Their research aimed to expand upon the previous advancements in and scope of Ni/NHC-catalysed hydroacylation, as developed by Ogoshi,¹²⁸ by enabling the formation of heterocyclic ketones (beyond just carbocyclic ketones), which is a key intermediate for a sequential enantioselective α -arylation reaction, thus allowing access to valuable



 $\mbox{Scheme 51}$ Ni/NHC reductive coupling of alkenes and carbonyls with hydrosilanes. 131



Scheme 52 Ni⁰/IAd-catalysed intramolecular hydroacylation of alkenes for synthesis of heterocyclic ketones.¹³³

N-containing heterocyclic ketones with α -chiral quaternary stereocentres.

Various indole moieties bearing electron-donating (333), or electron-withdrawing groups (334) were successfully tolerated under the protocol giving six-membered heterocyclic ketones in moderate yield. However, the scope of the substrates used is primarily limited to monosubstituted alkenes. The presence of additional β -substituents on the alkene was found to be detrimental to the hydroacylation reaction, resulting in poor yields (335). Likewise, 1,1-disubstituted alkenes also performed poorly (336) under their protocol.

Hydrocarbamoylation refers to the formal insertion of unsaturated C=C bonds into the C-H bond of formamide. It provides yet another alternative to access alkanamides, circumventing the need for the introduction of toxic carbon monoxide, which is commonly used in the aminocarbonylation of electrophiles¹³⁴ to access amides.

To date, the only study of Ni/NHC-catalysed hydrocarbamoylation of alkenes was reported by Hiyama and co-workers¹³⁵ in 2012. Using [Ni(COD)₂] and IAd together with triethylaluminium as a co-catalyst, they could obtain linear alkanamides in high yields with excellent anti-Markovnikov selectivity (Scheme 53).

Various *N*-alkyl substituted formamides (substituted with Me or Bn) proved to be effective in their hydrocarbamoylation regime. When an enantioenriched formamide is used, the corresponding optically pure linear alkanamide is likewise formed without the erosion of chirality ((*S*)-337). However, formamides derived from primary amines (with an acidic N–H) were incompatible under the Lewis acidic conditions. Monosubstituted alkenes bearing internal C—C linkages (339), silyl groups (340) and alkoxycarbonyl groups (338) were also compatible. Limitations in the scope include 1,1-disubstituted or 1,2-disubstituted internal alkenes, which were unsuccessful in delivering the desired product.

In a tentative mechanism provided by the authors (Scheme 53), the activated formamide substrate (*via* O-coordination with



triethylaluminium Lewis acid) forms an η^2 -formamidenickel intermediate **341**. Subsequent oxidative addition of the Ni⁰ centre into the C(sp²)–H bond forms the nickel–hydride intermediate **342**. Coordination of the alkene substrate forms **343**, which participates in regioselective migratory insertion to deliver a primary alkyl-nickel intermediate **344**. The less hindered primary alkyl-nickel intermediate is formed preferentially over the secondary alkyl-nickel intermediate which results in excellent linear-selectivity. Lastly, reductive elimination of **344** forms the crucial C(sp³)–C(sp²) bond, with the release of the linear alkanamide product. The presence of a highly bulky NHC ligand is believed to govern the regioselectivity of insertion and to improve the efficiency of the reaction through the facilitation of reductive elimination of **344**.

3.7 Dicarbofunctionalisation of alkenes

Dicarbofunctionalisation of alkenes, involving the incorporation of two carbogenic moieties across the carbon–carbon double bond, has emerged as an attractive method for elevating molecular complexity from coupling of simple alkene precursors.³² Complex sp³-carbon frameworks can be easily accessed in this manner. Nickel-catalysed dicarbofunctionalisation has gained significant traction over the years, and with appropriate adjustment of the catalyst's electronic and steric features, β -H elimination leading to the undesired Heck-type product can be suppressed. Two major pathways typically proceed, the two-electron ionic pathway and the radical pathway which involves a single electron transfer (SET) process. The former pathway is more frequently documented with the utilisation of a robust σ -donating NHC ligand. Another proposed mechanism involving oxidative cyclisation has also been put forth.^{136,137} Often cited in literature studies, redoxneutral dicarbofunctionalisation refers to the coupling of alkene with a nucleophilic (e.g., organometallics) and electrophilic (e.g., halides or pseudohalides) reagent, while reductive dicarbofunctionalisation refers to the coupling of alkene with two electrophilic partners in the presence of a reductant.^{32,138}

The reaction is initiated either through transmetalation with an organometallic compound or through the oxidative addition of an electrophile (e.g., halides and pseudohalides), resulting in the formation of a carbon-nickel species. This species undergoes a site-selective insertion with an alkene, forming a regiodefined alkyl-nickel intermediate. Regioselectivity of the insertion is primarily dictated by steric factors, minimising unfavourable steric repulsions between the olefinic substituent and the NHC. The second carbogenic component can then be trapped and incorporated through cross-coupling with another organometallic compound or electrophile. The challenges associated with dicarbofunctionalisation, such as the control of regio- and stereo-selectivities, competitive β -H elimination and undesired coupling of nucleophilic and electrophilic reagents, have been addressed in recent studies, through the utilisation of N-heterocyclic carbene ligands and their distinctive properties translated to the NHC-Ni complex (Scheme 54).

One of the earlier reports on NHC/Ni-catalysed dicarbofunctionalisation of alkenes was that of Ogoshi's work, who described a selective three-component coupling reaction of tetrafluoroethene, ethylene and aldehydes to furnish fluorinecontaining ketone derivatives (Scheme 55).¹³⁹ This reaction can be formally conceptualised as 1,2-alkylacylation of ethene or hydroalkylation of tetrafluoroethene. The optimal ligand found for this reaction was IPr **1f**, which enabled the formation of the



Scheme 54 Generalised two-electron pathway of NHC/Ni-catalysed dicarbofunctionalisation.



desired cross-trimerisation product while suppressing the formation of the side product originating from the dimerisation of aldehyde. A plausible reaction mechanism was described by the authors. Coordination of ethylene and tetrafluoroethene with the active IPr-Ni⁰ species forms the $\eta^2:\eta^2$ -Ni(IPr) complex **349**, which undergoes an oxidative cyclisation with the bound alkenes, to form a 2,2,3,3-tetrafluoronickelacyclopentane intermediate **350**. Chemoselective nucleophilic addition from the more electron-rich alkylnickel (-CH₂Ni) bond to the aldehyde produces the seven-membered oxanickelacycle species **351**. β -Hydride elimination of **351** precedes the final reductive elimination of **352** to afford the tetrafluorinated ketone product, while regenerating the active IPr-Ni⁰ species.

While olefinic substituents were not varied, different electronic environments of the aldehydes were tested under their regime. Benzaldehyde derivatives such as *p*-anisaldehyde (345), 4-formylbenzoate (346) or boronic ester-substituted aldehydes (347) were well tolerated. Aliphatic aldehydes, such as, cyclohexyl carboxaldehyde (348), also gave the corresponding aliphatic ketone product in moderate yields.

In 2016, Cramer and co-workers reported an *enantio*- and diastereo-selective NHC/Ni-catalysed three-component coupling reaction of aldehydes, silanes and norbornenes, to deliver enantioenriched silyl-protected indanols¹⁴⁰ (Scheme 56). Survey of a variety of chiral NHC ligands led to the most effective C_2 -symmetric NHC ligand **353** with a 1,2-di(napthalen-1-yl)-ethylene diamine backbone scaffold and an *ortho*-isopropyl-phenyl group as the *N*-aryl arm.



Scheme 56 NHC/Ni-catalysed enantioselective three-component reaction to access silylated indanols.¹⁴⁰

A plausible reaction mechanism commences with the η^2 coordination of the norbornene and aldehyde to the chiral NHC–Ni⁰ complex. The favoured orientation mode of the aldehyde group is proposed to be 357, where the aryl substituent of the aldehyde group is positioned away from both the methylene bridge of norbornene and the flanking bulky *N*-aryl substituents of the NHC to minimise unfavourable steric interactions. Enantio-determining oxidative cyclisation of 357 leads to the formation of the fused oxanickelacycle intermediate **358**. Reaction of the nickel-alkoxide with silane breaks the Ni–O bond of the oxanickelacycle, rupturing the ring to form intermediate **359**, which consists of a silyl-protected alcohol and a nickel–hydride moiety. Another molecule of norbornene enters the cycle and reacts with **359** in a sequence, hydronickelation, $C(sp^2)$ -H activation (of the aldehyde substrate) and reductive elimination, to afford the cyclonickelated species **360**, with concomitant release of the hydrogenated norbornane species. Final reductive elimination leads to the formation of the second $C(sp^3)$ - $C(sp^2)$ bond, releasing the silylated indanol product and regenerating the active NHC-Ni⁰ species.

Reaction with electron-deficient benzaldehyde **354** (with a *para*substituted ester moiety) resulted in slightly reduced yields but enantioselectivity remained high. Norbornene derivatives, such as the *exo*-methylenealkoxy-substituted derivative **355** or benzonorbornadiene **356**, were effective substrates, delivering the annulated products in good yields and enantioselectivity. Certain limitations of the catalytic regime were highlighted. A minor hydroalkylated side product was observed for some substrates, possibly formed from the competing reductive elimination of nickel–hydride **359**. No appreciable reaction with unstrained alkenes like cyclopentenes was observed under the reaction conditions.

In 2017, Stanley and co-workers described an NHC/Nicatalysed carboacylation reaction of *o*-allylbenzamides with aryl boronic pinacol esters to access a variety of 2-benzyl-2,3dihydro-1*H*-inden-1-ones¹⁴¹ (Scheme 57). Their methodology was noteworthy as it involves the key NHC–Ni-mediated amide C–N bond activation step and obviates the requirement for strained cyclic ketones or directing groups, which had been conventionally used to facilitate carboacylation.

The carboacylation methodology enabled with the strongly electron-donating SIPr **2b** (1,3-bis(2,6- diisopropylphenyl)-4,5dihydroimidazol-2-ylidine) ligand was evaluated using various substituted arylboronic pinacol esters (ArBpin) and *o*-allylbenzamides. The regime was compatible with heteroarylboronic acid pinacol esters featuring a thiophene **361** group. Electrondeficient *para*-chlorinated ArBpin **362** was well-tolerated, without competing activation of the C–Cl bond. Carboacylation of *o*-allylbenzamide with electron-withdrawing (**363**) or electrondonating (**364**) substituents afforded the corresponding cyclised product in excellent yields. However, the reaction had certain substrate limitations which include *o*-allylbenzamides with a 1,1disubstituted alkene moiety, or acyclic 5-hexenamide derivatives.

The authors proposed a possible reaction pathway, initiated by the $\eta^2(\text{alkene}):\eta^2(\text{amide})$ coordination of the *o*-allylbenzamide substrate with the NHC-Ni⁰ catalyst. Subsequent oxidative addition of the C-N amide bond generates the acyl-Ni(π)-amido complex **366**. Base-assisted transmetalation with the arylboronic pinacol ester produced the acyl-Ni(π)-aryl intermediate **367**, followed by regioselective insertion of the acyl-nickel bond into the tethered alkene, to form the primary alkyl-Ni(π)-aryl complex **368**. Reductive elimination releases the indanone product and regenerates the active species **365** through coordination with another *o*-allylbenzamide molecule. Although the proposed pathway involves transmetalation followed by migratory insertion, the reverse order of events with migratory insertion preceding transmetalation cannot be completely ruled out.

In 2019, Newman's group reported a NHC/Ni-catalysed intramolecular carboacylation and hydroacylation reaction of methyl *ortho*-allylbenzoate derivatives (Scheme 58), *via*



Scheme 57 NHC/Ni-catalysed carboacylation of alkenes via amide C–N bond activation.¹⁴¹

C(acyl)-O bond activation, building upon Stanley's prior work.141 Despite the strong C(acyl)–O bond of methyl esters¹⁴² and the high energy cost incurred in the generation of acyl-Ni intermediates, this challenging transformation was rendered feasible through an intramolecular cyclisation step to trap the unstable acyl-Ni intermediate.^{141,143} The most effective proligand was determined to be the saturated SIPr salt, and a combination of CsF and KF in the reaction media was found to be crucial in enhancing the yield and ensuring the reproducibility of the reaction. Both electron-withdrawing (enolisable para-acetyl group) 370 and electron-donating substituents (para-methoxy) 371 on the arylboronic acid substrate were well tolerated. In their attempt to realise an alkylacylation reaction with the formation of challenging $C(sp^3)-C(sp^3)$ bonds, alkylboronic acid was used but instead produced a significant amount of the reduced indanone product. With this serendipitous finding, their group proceeded to optimise this reductive hydroacylation reaction, with 1-phenylethanol being the most effective hydride source and reductant. A monodeuterated



Scheme 58 NHC/Ni-catalysed intramolecular carboacylation and hydroacylation of alkenes *via* ester C–O bond activation.⁵⁸

derivative of the alcohol also led to the formation of the corresponding indanone product **372** with high (85%) deuterium incorporation in the methyl group, opening opportunities for pharmaceutical applications.

A plausible mechanism for the two distinct transformations is depicted in Scheme 57. Reversible oxidative addition of the C(acyl)–O bond of the methyl ester forms the acyl-Ni intermediate **373**. Regioselective migratory insertion into the tethered alkene yields the primary alkylnickel intermediate **374**. The selectivity is likely governed by the steric influence of the bulky SIPr ligand. In the carboacylation pathway, fluoride-assisted transmetalation with the arylboronic acid results in the formation of the alkyl-Ni-aryl complex 375, which furnishes the desired product upon reductive elimination. In the reductive hydroacylation pathway, ligand exchange of the methoxy ligand with 1-phenylethanol (or the monodeuterated derivative) initially occurs, giving the alkoxy-nickel species **376**. β -Helimination from the alkoxide moiety results in the formation of the nickel-hydride (or deuteride) intermediate **377**, accompanied with the release of the acetophenone by-product. A possible minor pathway involving the direct β -H-elimination from the methoxy group, albeit slow, could also produce the same **377** species, without the intervention of the exogenous alcohol. Importantly, no intramolecular Heck-type enone product was observed under their protocol.

In their ongoing pursuit of developing regio- and stereoselective olefin functionalisation protocols enabled with Ni–NHC catalysts, Koh and co-workers reported a regioselective 1,2diarylation reaction of the less electronically activated aliphatic 1,3-dienes with site-selectivity fully controlled by the bulky IPE ligand¹⁴⁴ (Scheme 59). This redox-neutral reaction system facilitated the effective three-component combination of a diene, an aryltriflate electrophile and an aryl organometallic nucleophile, resulting in the formation of the desired diarylation products consisting of tertiary and quaternary carbon centres. Notably, minimal Heck-type or homo-/cross-coupling biaryl by-products were observed. Previous challenges of allylic rearrangement from the dissymmetrical η^3 - π -allyl-metallointermediate leading to mixtures of 1,2- and 1,4-addition regioisomers,¹⁴⁵ and stereoisomeric *E/Z* mixtures were not an issue under this protocol.

Naturally occurring and bioactive metabolites such as (+)4cholesten-3-one (**378**) and (+)-nootkatone (**379**), consisting of a 1,3,5-triene moiety, performed well under the reaction protocol, delivering the products in a single regioisomer in a high diastereomeric ratio (5:1-10:1). Besides using a Grignard reagent as the organometallic coupling partner, the milder arylzinc nucleophile could also be employed, allowing for the tolerance of more sensitive functionalities such as esters (**380**). Arylmagnesium bromide with an electron-withdrawing *meta*fluoro substituent (**381**) also successfully delivered the adduct.

Preliminary mechanistic studies led to a postulated mechanism depicted in Scheme 59. Oxidative addition of the aryltriflate with the active NHC-Ni⁰ species forms the aryl-Ni^{II} intermediate **382**, which selectively inserts into the less substituted C—C bond of the 1,3-diene, producing the aryl-branched primary alkylnickel species **383**. The generation of a rearrangement- and stereoerosion-prone allylnickel intermediate was precluded with this branched-selective carbonickelation. The observed regioselectivity is solely determined from steric effects, proceeding *via* the lower energy transition state that minimises steric repulsions between the sizeable NHC ligand and the diene substituent. Transmetalation with the aryl organometallic reagent led to the formation of an alkyl-Ni-aryl intermediate **384**, which underwent reductive elimination to turn over the catalytic cycle and release the desired 1,2-diarylation product.

In 2022, Tobisu and co-workers developed a NHC/Nicatalysed 1,2-carboaminocarbonylation reaction of norbornene *via* C–C bond cleavage of arylamide derivatives¹⁴⁶ (Scheme 60).





This reaction does not depend on the prerequisite of ring strain nor directing groups for C–C bond cleavage and resembles the net atom economical addition of a C–C bond of an arylamide across the carbon–carbon double bond.

The use of NHC ligands, particularly IMes **1c**, led to efficient addition reaction to yield the corresponding 1,2-carboaminocarbonylation



Scheme 60 NHC/Ni-catalysed 1,2-carboaminocarbonylation of strained alkenes via C–C bond activation of amides. $^{\rm 146}$

products while phosphine ligands failed to deliver any desired product. Heteroaromatic groups of the arylamide such as pyridine (385) or benzofuran (386) were tolerated, but resulted in a lowered yield possibly due to decarbonylation side reactions. Interestingly, benzofuran-substituted arylamide (386) or the introduction of an ester group (387) led to the exclusive formation of the trans isomer. Alternative strained alkenes such as dicyclopentadiene (388) reacted readily to deliver the cis-product as the sole isomer. For the catalytic regime, certain structural requirements for the substrate are essential for effective transformation. The lack of reaction from simple amides derived from aliphatic or aromatic amines indicates the crucial requirement of an indole-type amide for effective reaction. Electronic perturbation of the aryl component of the arylamide substrate has a significant impact on the reaction's efficiency, where higher levels of electron deficiency led to improved yields.

A catalytic cycle proposed by the authors is depicted in Scheme 60. Oxidative addition of the C(acyl)-N bond of the arylamide generates the acyl-Ni^{II}(1-indolyl) complex 389, which undergoes decarbonylation to form the aryl-nickel intermediate 390 with a bound CO ligand. Subsequent carbonickelation of the norbornene substrate forms the secondary alkylnickel species 391. Re-insertion of CO into the C(sp³)-Ni bond results in the formation of an acyl-Ni complex 392, which releases the final cis-product after reductive elimination to form the desired C(acyl)-N bond. The desired 1,2-carboaminocarbonylation product can be efficiently formed despite the reversible nature of CO insertion/de-insertion, which is attributed to the energetically feasible C(acyl)-N bond forming reductive elimination process, in contrast to the $C(sp^3)$ -N bond forming process from 391, which explains the absence of the 1,2-carboamination product. In-depth experimental studies also found that the cis-isomer is the kinetically favoured product, which spontaneously isomerises to the more thermodynamically stable transisomer (from theoretical calculations) in the presence of a Lewis base, such as the IMes 1c free carbene.

In 2022, Koh and co-workers successfully achieved a directing group free approach for direct addition of carbogenic aryl groups across unactivated C–C π -bonds in an NHC/Ni-catalysed three-component reaction¹⁴⁷ (Scheme 61). This novel approach solely relies on catalyst and ligand control for the efficient and highly regioselective 1,2-diarylation of unactivated aliphatic alkenes, eliminating the need for pre-installation of pendant directing auxiliaries or substrate control which hinges on the favourable intramolecular formation of a cyclic intermediate. An extensive survey of catalysts and ligands led to the selection of a bulky dimeric IPr-Ni(1) precatalyst, which was found to be more efficient than the conventional Ni(0) or Ni(1) precatalysts.

The significance of a sufficiently bulky NHC ligand stems from several reasons. The steric influence exerted by the NHC ligand must be substantial enough to provide good regiocontrol in the carbonickelation (migratory insertion) step. Without the presence of a directing group, the vacant coordination site of the nickel complex could trigger adventitious β -H elimination, leading to the formation of Heck-type by-products.¹⁴⁸ Having a sizeable NHC ligand could also preclude agostic C_{β}-H interactions which productively leads to β -H elimination. Lastly, reductive elimination would be facilitated with a sterically encumbered metal complex, outcompeting, and suppressing any undesired side reactions.

With support from DFT calculations, a catalyst-controlled reaction pathway was suggested by the authors. Substrate or solvent assisted dissociation of the dimeric Ni(1) precatalyst **25** forms the monomeric IPr-Ni(1) active species **396**. Oxidative insertion of the aryltriflate electrophile affords the arylnickel(m) intermediate **397**, which undergoes a branched-selective **1**,2carbonickelation reaction across the alkene substrate, resulting in the formation of the primary alkylnickel intermediate **398**. Theoretical calculations showed that the transition state leading to the linear-selective insertion intermediate was comparatively higher in energy and unfavourable due to the severe steric repulsions between the bulky IPr ligand and the olefinic substituent. Irreversible transmetalation with the organometallic



Scheme 61 Directing group free NHC/Ni-catalysed dicarbofunctionalisation of unactivated alkenes.¹⁴⁷

nucleophile followed, resulting in an arylalkylnickel species **399**. Reductive elimination of **399** furnishes the desired 1,2-diarylation product with high regioselectivity and regenerates the active Ni(1) species **396**.

The generality of the reaction has been demonstrated through its application to various functionalised terminal and 1,1disubstituted alkenes. Notably, reactions with the latter class of alkenes allowed for the formation of diarylation adducts which incorporate fully-substituted quaternary carbon centres (**395** and **396**). In addition, alkynylsilanes (**394**) and Lewis basic Nheterocycles like pyridine (**395**) were amenable substrates under the optimised reaction conditions. Coupling with alkene-derived bioactive molecules such as gemfibrozil **393** (a lipid-lowering agent) proceeded smoothly, thereby highlighting its utility for the synthesis of complex, pharmaceutically relevant molecules.

4. Conclusion

In this review, we have underscored the advancements made in nickel-catalysed alkene functionalisation reactions, through

the utilisation of N-heterocyclic carbene (NHC) ligands and their unique associated properties. Three main reaction classes of alkene functionalisation have been presented: Heck-type reactions, hydrofunctionalisation and dicarbofunctionalisation. The reaction methodologies presented have illustrated the potential of the NHC-Ni catalytic system as a valuable tool for the efficient and selective construction of bioactive and pharmaceutically relevant molecular architectures from inexpensive and readily available olefinic precursors. The key features of NHC ligands include their strong σ-donating capacity and their large buried volume ($%V_{bur}$) because of their steric bulkiness. Modular synthesis of structurally diverse NHC precursors¹⁴⁹ allows for convenient tuning of electronic and steric properties, which can considerably impact the efficiency, as well as the chemo-, regio- and stereo-selectivity of a reaction. The multifaceted design of NHC scaffolds permits the incorporation of stereogenicity at different positions of the NHCs, leading to the emergence of chiral NHC ligands employed in numerous enantioselective transformations facilitated with transition-metal catalysts.¹⁵⁰ Many mono- or multi-dentate chiral NHC ligands demonstrated exceptional capacity to induce high levels of enantioselectivity, attributed to their steric influence and attractive secondary non-covalent interactions in the stereodetermining transition state.

Despite the substantial progress made in the field of nickelcatalysed alkene functionalisation over the years, major challenges still remain. For instance, the regio- and enantioselective carbofunctionalisation of unactivated alkenes and light olefins, which have weak electronic and steric bias, is yet to be achieved in the absence of directing auxiliaries. Selective hydro/carbofunctionalisation of fluorinated alkenes, which are prone to defluorinative side reactions, is also a difficult task. These obstacles could potentially be resolved via the use of nickel catalysis in combination with judiciously designed chiral NHC ligands, such as the more elusive chiral CAAC-type or mesoionic NHC ligands that have found limited applications to date.¹⁵⁰ Another exciting prospect in this field could be the integration of NHCs as ligands for nickel/photoredox dual catalysis, which have remained largely unexplored since the seminal work for indoline synthesis by Jamison and Tasker.¹⁵¹ This represents a new disconnection approach to enable access to new chemical space and a border assortment of molecular scaffolds, including prized heterocyclic and hetero-substituted molecules, from simple olefin feedstocks.

Abbreviations

1,2-DCE	1,2-Dichloroethane
Ad	Adamantyl
BAr _F	Tetrakis[3,5-bis(trifluoromethyl)phenyl]borate
Bn	Benzyl
Вру	Bipyridine
Bu	Butyl
COD	1,5-Cyclooctadiene
Ср	Cyclopentadienyl

Ме	Methyl
CAAC	Cyclic(alkyl/aryl)(amino)carbene
Cy	Cyclohexyl
DFT	Density functional theory
ee	Enantiomeric excess
er	Enantiomeric ratio
HBCat	Catecholborane
HBpin	Pinacolborane
^h IEtPyO	(1-(2,6-Diethylphenyl)-3-(6-oxidopyridin-2-yl)- imidazol-2-ylidene)
iPr	Iso-propyl
IAd	1,3-Bis(1-adamantyl)imidazol-2-ylidene
IPr	1,3-Bis(2,6-diisopropylphenyl)-1,3-dihydro-2H-
	imidazol-2-ylidene
IPr*	1,3-Bis(2,6-bis(diphenylmethyl)-
	4methylphenyl)imidazol-2-ylidene
I ^t Bu	1,3-Di- <i>tert</i> -butylimidazol-2-ylidene
LLHT	Ligand-to-ligand hydrogen transfer
MAD	Methylaluminium bis(2,6-di- <i>tert</i> -butyl-4-
	methylphenoxide)
Mes	Mesityl
MTBE	Methyl <i>tert</i> -butyl ether
NHC	N-Heterocyclic carbene
Ph	Phenyl
r.r.	Regioisomeric ratio
<i>t</i> -Am	<i>tert</i> -Amyl
TEMPO	(2,2,6,6-Tetramethylpiperidin-1-yl)oxyl
TIPS	Triisopropylsilyl
<i>t</i> Bu	<i>tert</i> -Butyl
TBAB	Tetrabutylammonium bromide

Conflicts of interest

The authors declare no conflicts of interest.

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