

Particularly, key precedents by Oshima and Kambe showcased the possibility of introducing two differentiable alkyl fragments across conjugated alkenes with alkyl bromides and alkyl Grignard reagents, presumably enabled by the stabilization of resulting π -allyl and π -benzyl nickel species after initial insertion.^{4e,f} Early work by Cárdenas and coworkers demonstrated nickel-catalyzed intramolecular dialkylation.³ In the past year, there have been several reports on three-component 1,2-dicarbofunctionalization using substrate directivity strategies (Scheme 1B).^{6,10} In terms of examples with *alkyl* coupling partners, Nevado and coworkers demonstrated that allyl acetate and related alkenes could react with tertiary alkyl iodides and aryl iodides in a nickel-catalyzed reductive conjunctive cross-coupling.^{6a} Our group developed a 1,2-aryllalkylation reaction using aryl iodides and organozinc reagents under nickel catalysis.^{6c} This reaction employed a removable bidentate directing group¹¹ to control the regiochemical outcome through putative nickelacycle stabilization and to prevent otherwise rapid β -hydride (β -H) elimination (Scheme 1C).

The efficiency of our 1,2-aryllalkylation reaction prompted us to consider whether it would be possible to use alkyl halides and alkyl organometallic reagents with non-conjugated terminal and internal alkenes *via* chelation control (Scheme 1D). At the outset we recognized that successful realization of the proposed transformation would necessitate overcoming a series of obstacles, most notably the fact that alkylmetal species capable of undergoing β -H elimination are present in every step of the catalytic cycle. Additionally, with alkyl halide electrophiles it is known that single-electron oxidative addition pathways are sometimes preferred, particularly with first-row metals,^{8,9} which meant that there was potential for mechanistic divergence compared to our earlier work ($\text{Ni}^{\text{I}}/\text{Ni}^{\text{III}}$ vs. $\text{Ni}^{\text{0}}/\text{Ni}^{\text{II}}$).

Results and discussion

To reduce this idea to practice, we began by exposing an alkenyl carbonyl compound **1a** to reaction conditions using iodoethane as the electrophile and dimethyl zinc as the nucleophile under nickel catalysis (Table 1). To suppress β -H elimination, we hypothesized that a strong, bidentate directing group would be beneficial. Given our group's previous success in employing Daugulis's 8-aminoquinoline (AQ) directing group¹⁰ in 1,2-difunctionalization reactions,^{6e,11} we focused our efforts on this directing group.

In a series of initial experiments, we were delighted to observe formation of the desired 1,2-dialkylated product **2a** in 19% yield along with 14% of dimethylated byproduct **2a'** based on conditions from our previous work using 20 mol% $\text{Ni}(\text{cod})_2$.^{6c} Interestingly, the product was determined to have electrophile incorporation at the β -position, opposite to the results in our previously reported 1,2-aryllalkylation methodology, suggestive of a different underlying mechanism *that is induced by use of an alkyl halide electrophile (vide infra)*. We then moved on to examine nickel(II) precatalysts, which are inexpensive and bench-stable alternatives to air-sensitive nickel(0) precatalysts. With $\text{NiCl}_2 \cdot \text{glyme}$ as catalyst, the procedure could reliably be reproduced without the use of an inert atmosphere

Table 1 Optimization of 1,2-dialkylation^a

Entry	Cat. Ni	Etl (equiv.)	ZnMe ₂ (equiv.)	Solvent (M)	Temp. (°C)	Yield 2a ^b (%)	Yield 2a' ^b (%)
1	20% Ni(cod) ₂	2	2	Dioxane (0.5)	100	19	14
2	20% Ni(cod) ₂	2	2	THF (0.5)	100	23	18
3	10% NiCl ₂ ·glyme	2	2	DMA (0.5)	100	39	24
4	10% NiBr ₂	2	2	DMA (0.5)	100	37	18
5	10% NiCl ₂ ·glyme	2	2	Dioxane (0.5)	RT	n.d.	n.d.
6	10% NiCl ₂ ·glyme	2	2	DMF (0.5)	RT	13	5
7	10% NiCl ₂ ·glyme	2	2	DMA (0.5)	RT	28	17
8	10% NiCl ₂ ·glyme	2	2	DMA (0.1)	50	46	26
9	10% NiCl ₂ ·glyme	2	2	DMA (0.2)	60	53	13
10	10% NiCl ₂ ·glyme	4	4	DMA (0.2)	60	68	14
11	15% NiBr ₂	8	8	DMA (0.2)	60	81	17
12	15% NiBr ₂	8	6	DMA (0.2)	60	83	12

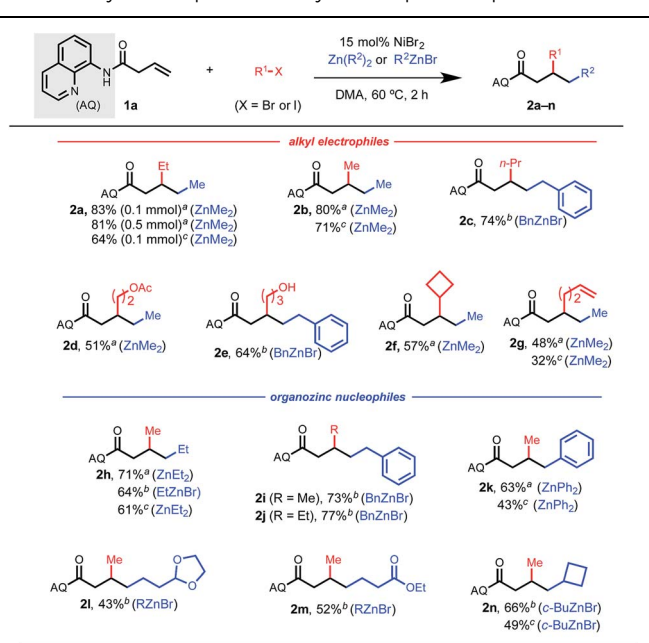
^a Reaction conditions: alkene (0.1 mmol), ZnMe₂ (1.2 M in toluene).

^b Yields determined by ¹H NMR analysis using CH₂Br₂ as internal standard; n.d. = not detected.

glovebox, delivering the desired 1,2-dialkylation product in 39% yield (entry 3). We were also encouraged by the efficaciousness of a range of nickel(II) sources, including NiBr₂ which performed comparably to NiCl₂·glyme (entry 4). The reaction only proceeded in appreciable yield when polar aprotic solvents were used, so we opted to use DMA as the solvent for subsequent optimization. Through systematic screening, we were pleased to observe that the yield increased when the reaction was performed at concentrations lower than 0.5 M. With this change, the temperature could be decreased to 60 °C, and under these conditions, the desired product was formed in 53% yield. Next, we reasoned that the nucleophile and electrophile were being consumed through competitive two-component cross-coupling or homodimerization. Indeed, we observed a dramatic increase in yield proportional to the increase in equivalents of both electrophile and nucleophile that were used (entry 8). The reaction provided 68% yield using 4 equivalents of each (entry 10), and the yield could be further improved to 83% using 6 equivalents of nucleophile and 8 equivalents of electrophile (entry 12).¹² Because the reaction partners in this study were generally inexpensive and commercially available, we performed most of the ensuing experiments using the conditions shown in entry 12, though we also carried out several examples under the conditions shown in entry 10 for comparison.

Having identified optimal conditions, we first explored potential alkyl electrophiles and nucleophiles that were compatible in this 1,2-dialkylation reaction (Table 2). It is important to note that competitive 1,2-dialkylation with the nucleophile was also observed in 5–20% by crude ¹H NMR in most cases. To our satisfaction, a wide range of primary alkyl electrophiles and nucleophiles were tolerated. By simply changing the identity of the nucleophilic and electrophilic components (*i.e.*, iodoethane/dimethylzinc to iodomethane/diethylzinc), both regioisomers could be synthesized in high yields (**2a** and **2h**). In general, secondary and tertiary alkyl reaction partners did not perform well under the optimized conditions, likely due to steric constraints of the resulting chelation-stabilized nickelacycle and/or rapid β -hydride elimination for partners with 4 or more accessible β -hydrogen atoms.



Table 2 Alkyl nucleophile and alkyl electrophile scope^{a,b,c,d}

^a Reaction conditions: alkene **1a** (0.1 mmol or 0.5 mmol), alkyl halide (8 equiv.), dialkyl/diarylzinc (6 equiv.), DMA, 60 °C, 2 h. ^b Organozinc bromide (8 equiv.) in place of dialkyl/diarylzinc. ^c 4 equiv. alkyl halide and 4 equiv. organozinc. ^d Percentages represent isolated yields.

Cyclobutyl coupling partners were an exception, however, and the corresponding difunctionalized products **2f** and **2n** could be obtained in high yields. Several synthetically useful functional groups were tolerated in this reaction, allowing for the installation of acetals, esters, and free alcohols in moderate to good yields (**2d**, **2e**, **2l**, and **2m**). Heterocycle-containing reaction partners were generally incompatible, which we hypothesize is due to catalyst coordination. Interestingly, 1,2-aryllkylation products could be accessed using diphenyl zinc (**2k**). Terminal alkene-tethered electrophiles were compatible but delivered the desired product in modest yield (**2g**).

To provide a representative survey of the effects of coupling component equivalents, we conducted several examples using 4 equivalents of both electrophile and nucleophile. The yields were typically within 20% of the optimized yields, providing potential end-users with the opportunity to prioritize yield over reagent equivalents, or *vice versa*. We found that the AQ auxiliary of a representative product could be unmasked to the corresponding carboxylic acid or methyl ester under standard hydrolysis or methanolysis¹³ conditions, respectively (Scheme 2).

Next, we examined internal and α -substituted terminal β,γ -unsaturated alkenes (Table 3). For these substrates, we elected to use iodoethane as the electrophile and dimethyl zinc or benzylzinc bromide as the nucleophile. Internal alkenes provided the desired *syn*-1,2-difunctionalized products in good yields and with high diastereomeric ratios (**3a–3g**). The relative stereochemistry was determined by reacting representative internal alkene **1b** with diphenyl zinc and methyl iodide,



Scheme 2 Removal of the AQ directing group.

Table 3 β,γ -Alkene substrate scope^{a,b,c}

^a Reaction conditions: alkene **1b–1** (0.1 mmol or 0.5 mmol), iodoethane (8 equiv.), dimethylzinc (8 equiv.), DMA, 60 °C, 12 h. ^b Benzylzinc bromide (8 equiv.) in place of dimethylzinc. ^c Percentages represent isolated yields.

yielding a 1,2-aryllkylation product reported in our previous publication (see ESI[†]). Upon subjecting a trisubstituted alkene to the reactions conditions, we were pleased to discover that quaternary centers could be formed at the γ -position (**3h**). We questioned whether a 1,1-disubstituted alkene could react; however, the reaction did not proceed at all with this substrate (**3i**). We hypothesize that the putative chelation-stabilized alkylnickel intermediate is very sensitive to the proximal steric environment.

To investigate the steric constraints of our optimized system, we subjected α -substituted substrates to the reaction conditions (**3j** and **3k**).¹⁴ More sterically bulky groups led to lower yields. In both successful cases, only a single diastereomer was detected, which was assigned as *trans* in analogy to our previous work.¹⁵

It is typically difficult to extend catalytic directed alkene functionalizations to substrates containing more distal functional groups due to the instability of the metalacycles that are six-membered or larger. Recently our group published a tridentate directing group strategy for the hydrofunctionalization of γ,δ -unsaturated alkenes; the tridentate directing group is thought to suppress β -H elimination in six-membered palladacycles.¹⁶

An alternative strategy is to intercept the metalacycle in a rapid subsequent step, such that this downstream reaction



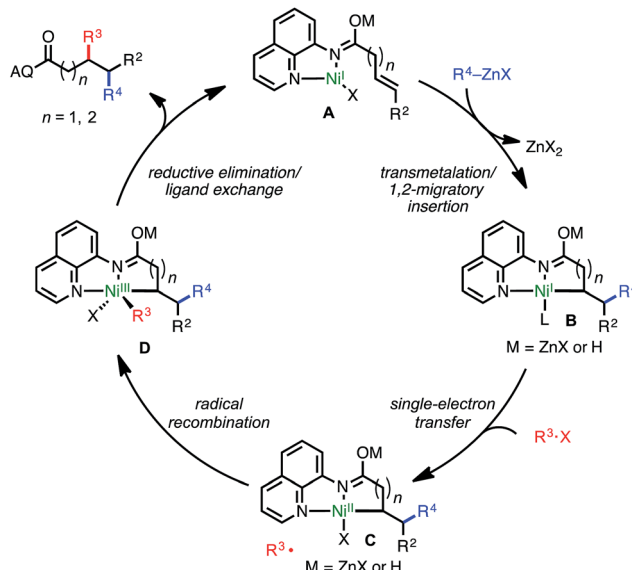
can outcompete undesired β -H elimination. Given the unique reactivity patterns in this 1,2-dialkylation reaction, we thus questioned whether γ,δ -dialkylation would be feasible *via* a six-membered AQ-bound nickelacycle (Table 4). To this end, we employed a terminal γ,δ -unsaturated alkene substrate with iodoethane and benzylzinc bromide. Our standard reaction conditions delivered the desired product in 65% yield (5a), showcasing the ability of this transformation to introduce alkyl fragments at the γ - and δ -positions, which are challenging to functionalize using existing chemistry. We then introduced substituents at various positions to investigate the effects on diastereoselectivity. Interestingly, α -substituted starting materials gave the desired product, but with significantly diminished diastereoselectivity (5b). When the analogous β -substituted starting material was used, the desired product was delivered as a single diastereomer (5c).¹⁵ These results shed light on the influence of proximal substituents on competing nickelacycle formation pathways. Internal alkenes within this substrate class also reacted well to give dialkylated products (5e and 5f). To our delight, the corresponding trisubstituted alkene also proved to be reactive, allowing formation of a quaternary center (5g). We further probed a representative δ,ε -unsaturated substrate but in this case could only detect isomerization byproducts.

In an attempt to gain insight regarding the mechanism of the oxidation addition step of this 1,2-dialkylation reaction, we conducted a radical clock experiment using (bromomethyl)cyclopropane as an electrophile (Scheme 3). Upon treating standard substrate 1a with 2 equivalents of electrophile and 2 equivalents of dimethyl zinc, we observed the formation of ring-opened product 2g in 27% yield. This result is consistent with a radical oxidation step involving single-electron transfer (SET). Such SET processes have been previously reported to occur from organonickel(i) species that are formed *via* transmetalation.^{3b,8}

Though a detailed mechanistic investigation of this transformation remains outside of the scope of the present manuscript, a plausible catalytic cycle consistent with experimental data is shown in Scheme 4. The cycle follows a Ni^I/Ni^{III} redox



Scheme 3 Mechanistic experiment involving (bromomethyl)cyclopropane as an electrophile.



Scheme 4 Proposed catalytic cycle.

manifold, in line with other alkyl-alkyl cross-coupling processes.^{3b,8d,e,9h} Initially, a Ni^I catalyst (presumably bound as shown in A) undergoes a transmetalation/*syn*-1,2-migratory insertion sequence to yield putative nickelacycle B. This species then performs SET with the alkyl halide electrophile to generate Ni^{III}-bound species C and the corresponding alkyl radical. After radical recombination with the resulting alkyl radical, Ni^{III} intermediate D undergoes reductive elimination to yield the desired product and regenerate the active catalyst.

An alternative mechanism could also be envisioned in which oxidative addition of the alkyl halide occurs first, followed by *syn*-insertion to deliver the electrophile alkyl fragment proximal to the directing group and generate the 6- or 7-membered nickelacycle for substrates classes 1 and 4, respectively. Though we have not conclusively ruled out this pathway at this stage, several observations stand counter to this mechanism. First, assuming this alternative pathway were operative, the observed regiochemical outcome when utilizing γ,δ -unsaturated alkenes (4) would indicate that the catalyst selectively forms a 7-membered alkyl nickelacycle in preference to a 6-membered alkyl nickelacycle, which is inconsistent with established trends in alkyl metalacycle formation.¹⁶ Second, assuming the former point were true, it would stand to reason that δ,ε -unsaturated carbonyl compounds would also be compatible; however, in practice this substrate class was not tolerated. Third, formation of products such as 5g following this alternative mechanism would require the intermediacy of a 7-membered tertiary-alkyl nickelacycle, a species that would be expected to be highly unstable.

Table 4 γ,δ -Alkene substrate scope^{a,b,c}



^a Reaction conditions: alkene 4a-g (0.1 mmol), iodoethane (8 equiv.), dimethylzinc (8 equiv.), DMA, 60 °C, 12 h. ^b Benzylzinc bromide (8 equiv.) in place of dimethylzinc. ^c Percentages represent isolated yields.



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- 14 To examine whether epimerizable α -stereocenters underwent racemization under the reaction conditions, we performed an additive study in which a protected amino acid derivatives was added to the reaction and reisolated after the standard time. In this experiment racemization was negligible (>99% ee to 97% ee) (see ESI† for details).
- 15 The relative configuration was assigned as *trans* by analogy to compounds prepared in ref. 6c. The preference for *trans* product formation is believed to arise from more facile formation of the *trans* nickelacycle.
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