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Hydroalkylation of Olefins to form Quaternary Carbons

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Supporting Information Placeholder

ABSTRACT: Metal-hydride hydrogen atom transfer (MHAT) functionalizes electronically unbiased alkenes with predictable branched (Markovnikov) selectivity. The breadth of these transformations has been confined to π -radical traps; no sp³ electrophiles have been reported. Here we describe a Mn/Ni dual catalytic system that hydroalkylates unactivated olefins with unactivated alkyl halides, yielding aliphatic quaternary carbons.

Olefins represent versatile feedstocks and intermediates for chemical synthesis. Metal-hydride hydrogen atom transfer (MHAT) has emerged as a useful reaction platform for the branched-selective hydrofunctionalization of olefins. Its high chemoselectivity for olefins and mild reaction conditions have allowed its deployment in medicinal chemistry and natural product synthesis.¹ The bulk of these transformations involve carbon-heteroatom bond formation, whereas intermolecular C-C formation has been relatively unexplored and has largely required stoichiometric radical traps by π -electrophiles. Pioneering advances in the formation of C-C bonds are represented by hydrocyanations and hydrooximation from Carreira² and Boger.³ More recently, Baran and coworkers developed a powerful variant of the Giese reaction^{4,7d} as well as a two-step procedure for hydromethylation (Figure 1A).⁵ Finally, our group⁶ and others⁷ have investigated the branched-selective hydroarylation of olefins using MHAT, establishing olefins as progenitors for arylated quaternary centers.

One contributing factor to the limited range of Markovnikov hydro-*alkylations* is a dearth of alkyl radicalophiles. Whereas MHAT has relied historically on stoichiometric radical traps such as O₂,^{1a} our group has become interested in combining MHAT with a second catalytic cycle, thereby expanding the variety of coupling partners.^{1b} Recently, we established a dual catalytic platform to allow for the hydroarylation of unactivated olefins^{6b,c} as well as the addition of carbanion surrogates to aldehydes,⁸ yielding branchedselective products otherwise inaccessible by tradition radical reactions. A dual catalytic approach for the hydro*alkylation* of olefins would allow us to use known alkyl coupling partners (e.g. alkyl halides⁹, carboxylic acids¹⁰) in lieu of alkyl radical traps, and cross a longstanding methodological gap.





Markovnikov hydroalkylation would also provide a new transform to dissect quaternary carbons, which remain challenging motifs in natural products and drug scaffolds. While radical chemistry has emerged as a useful platform for the construction of sterically congested motifs,^{11,12} sp³–sp³ cross-coupling remains an underdeveloped area for quaternary carbon formation.^{9a,13} The use of nickel catalysis to generate and engage open-shell intermediates has been revolutionized by Fu,^{9a,14} but its use in the construction

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of quaternary carbons has only recently been described,^{6c,15,16} with the development of alkylation reactions restricted to stabilized radicals¹⁷ or Giese reactions.¹⁸ Recently, olefins have become viable coupling partners in reductive coupling¹⁹ and nickel catalysis as surrogates for organometallic reagents.^{17,20} Whereas these nickel hydride-mediated methods yield anti-Markovnikov (linear) hydrofunctionalized products, MHAT dual catalysis provides access to branched products, even quaternary carbons, using similarly benign starting materials and conditions (Figure 1B). Herein we describe an approach for the hydro*alkylation* of unactivated olefins using Mn/Ni dual catalysis.

A. Envisioned Catalytic Cylce



Figure 2. (a) Plausible catalytic cycle for the hydroalkylation of olefins. (b) Optimization parameters. ^a0.1 mmol scale, yield determined by GC-FID using 1,3,5-trimethoxybenzene as an internal standard. ^b0.3 mmol scale, isolated yield 15:1 branched (*b*): linear (*l*) product. ^cno HFIP. ^dusing 2-iodoethyl benzoate instead of **1**. dpm=dipivaloylmethane; HFIP=1,1,1,3,3,3-hexafluoro-2-propanol; 1,2-DCE=1,2-dichloroethane; PC=propylene carbonate

Our previous method to form arylated quaternary centers^{6c} led us to hypothesize that our reaction design might translate to alkylation. Namely, an MHAT-generated tertiary radical or organometallic could be intercepted by a low valent nickel species, which could subsequently engage with an alkyl halide (or alkylnickel species) and yield our desired product upon reductive elimination (Figure 2A).^{10b,21} Regeneration of low valent nickel may proceed through formation of Ni–H via silane or Mn–H.

A successful Markovnikov olefin coupling would require an override of the inherent anti-Markovnikov migratory insertion found in Liu^{20a} and Fu's¹⁷ Ni–H systems. Unfortunately, initial attempts to utilize our Fe/Ni system^{6c} that provides such override yielded only trace product (Figure 2B). A polar solvent screen, however, indicated propylene carbonate (PC)²² was superior to *N*-methylpyrrolidinone (NMP). Curiously, PC as a co-solvent obviated the need for Mn⁰ and MnO₂ co-reactants, which we proposed to turn over the catalytic cycles. Instead the reaction could be run open to air.²³

Alkyl iodides coupled efficiently, whereas alkyl bromides, redox active esters, and sulfones yielded trace or no product (see SI). A screen of MHAT catalysts indicated that Mn(dpm)_{3²⁴} outperformed Fe(dpm)₃ and Co(dpm)₂. We did observe some product formation in the absence of Ni, but this background reactivity did not prove general and the yield could not be improved without the Ni co-catalyst. Similar to our arylation chemistry, traditional mono-, di- and tridentate ligands on Ni either provided no improvement in yield or ablated reactivity. Preparatively-useful yields were finally obtained with alcoholic additives: isopropanol led to marginal improvement and HFIP was found to almost double the vield. Due to decreased efficiency observed with Ph(*i*PrO)SiH₂, it is unlikely that the improved yield is due to an alcohol-silane complex. However, a noticeable color change from black to rust-red occurs when $HFIP/K_2CO_3$ is added to $Mn(dpm)_3$ in the absence of silane. Attempts to isolate and characterize this complex were unsuccessful. We cannot rule out the formation of a dimeric species bridged by the alcohol additive.7d,25

With optimized conditions in hand we began to investigate the breadth of olefin compatibility (Table 1, **2–28**). The *para*-methoxybenzyl ester of 4-iodobutyric acid (1) allowed the clear identification of products by both UV/VIS and mass spectrometry.²⁶ While our interest was on the formation of quaternary centers, we were pleased to find that all variants of olefin substitution were well-tolerated (**2–10**) and even tetrasusbituted olefins coupled, albeit in diminished

yield. Overall, the transformation exhibited exquisite regiocontrol with tri- and tetrasubstituted olefins yielding products with high branched-to-linear ratios (b:l), highlighting the selectivity of our developed method. Whereas trisubstituted olefins generally afforded higher regioselectivity than their exocyclic counterparts (**3a** vs **3a'**), 5-membered rings with exocyclic alkenes retained the normally high branchedselectivity (15), potentially a result of increased rate of MHAT due to strain release. Interestingly, terminal olefins (8–10) were subject to a background linear reaction, a trend also noted by Carreira with $Mn(dpm)_3$ -mediated transformations.²⁷ Three hypo-

Table 1. Hydroalkylation Olefin Scope^a

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^a0.3 mmol scale, isolated yield, see SI for specific catalyst loading. *b*:*l*= branched/linear ratio. ^breaction run under an air balloon. ^c5 equiv of olefin added in 2 portions (2.5 equiv at start and 2.5 equiv at 24 h).

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theses may explain this aberrant selectivity: a background Ni-only mediated pathway, analogous to reactivity observed by Liu,^{20a} lowered regioselectivity of MHAT itself due to similar potential energies of developing C-centered radicals, or a competitive hydrometallation pathway mediated by low valent manganese.²⁸

The reaction displayed high functional group compatibility in its tolerance of esters (11, 13, 21, 26), phthalimides (14), carbamates (4, 15, 26), silvl enol ethers (16), boronic esters (17), and epoxides (19). Interestingly, the reaction with alkenes proceeded with high chemoselectivity even in the presence of a primary alkyl bromide (12), which did not engage in the reaction or undergo protodehalogenation. Although primary (21) and secondary alcohols (22) required protection under the reaction conditions due to competitive silvlation, tertiary alcohols did not affect catalysis (24). Heteroatom substitution²⁹ on or adjacent to the alkene was well tolerated (16, 17, 25), but, in general, remotely-functionalized alkenes returned the highest yields (e.g. 21) and proximal branching lowered efficiency (22, 28).

Table 2. Alkyl Halide Scope^a

The abundance and diversity of olefins from commercial sources allowed a rapid survey of alkene scope. We were pleased to observe that a variety of natural product scaffolds (18-28) could be cleanly alkylated. A range of simple to complex terpenoids were successfully employed, which constitutes a new utilization of the chiral pool and potential access to new flavors and fragrances. The transformation of terpenes, whose hallmark features often are electronically unbiased,³⁰ hindered alkenes, have benefited immensely from MHAT methodology,^{1,4,6,7d,29} and served as particularly efficient scaffolds for this methodology. Notably, limonene oxide (19a) and 3-carene (23b) were both hydroalkylated with their scaffolds intact: no retrocyclization of the epoxide or cyclopropane motifs was observed. Pinene (20), on the other hand, predominantly underwent ring opening (3:1 ring opened: closed ratio) and yielded alkylated limonene derivative (20b), which did not undergo further hydrogenation of the resulting trisubstituted olefin.³¹ In some cases, alkylation noticeably altered the odor of these scaffolds, as was the case with rose oxide (25), which underwent alkylation with high branched selectivity. Substrates that contained



^a0.3 mmol scale, isolated yield, see SI for specific catalyst loading. *b:l=* branched/linear ratio. ^b0.1 mmol scale. ^cisolated as a mixture with hydrogenation, yield determined by NMR. Further purified by prep HPLC.

existing stereocenters exhibited modest stereoselectivity, as was the case with terpineol (24b) and carene (23b).

This method allowed the union of diverse metabolic building blocks (terpenes, amino acids, sugars) by strong covalent bonds. In addition to terpenes, ketide-like fragments corresponding to oxygen-polarized carbonchains could be appended. This merger complements Giese reactivity, which yields β -substitution, whereas **24** and **25** correspond to γ - and α -connections relative to a latent carbonyl. Allylglycine (**26**) proved a poor substrate (amino acids could be incorporated efficiently in Table 2, see below), but glucals coupled efficiently and yielded the C-glycoside product (**27**) as a single diasteromer.^{32,33} Furthermore, a single diasteromer of estrone derivative **28b** was observed, remarkably forming vicinal quaternary centers, albeit in reduced yield.

A diverse range of alkyl halides successfully coupled to form quaternary carbon centers (Table 2, 29-48). Sensitive functional groups like acetals (43) and nitrogen containing heterocycles (33) were unaffected by the coupling. Numerous simple alkyl chains could be appended to affect hydromethylation, ethylation, and pentylation reactions with similar efficiency. Ethylation of terpineol (37) resulted in a marked change in fragrance: from the sharp pine parent compound odor to a less-pungent musty, citrus. Methyl-d₃ iodide was also compatible under the reaction conditions, providing the isotopically-mixed geminal dimethylpyrrolidine 35. More complex alkyl iodides also proceeded in good yield, allowing for one-step installation of sugar- and steroid-bearing motifs 45 and 46. Prenyl groups are important motifs found in natural products. While prenvl bromide displayed poor reactivity due to competitive MHAT, we were pleased to find that prenyl surrogate, 4iodo-2-methylbutan-2-ol, yielded unnatural terpene 44 with excellent selectivity.

MHAT dual catalysis provides an orthogonal approach to phthalimide containing compounds **30** and **36**.³⁴ Phthalimide **36** has previously been accessed through disconnection at the quaternary center using a Cu-nanoparticle catalyzed Kumada coupling with the *tert*-alkyl Grignard.^{35,36} Phthalimide **31**, previously accessed in 5 steps from dimedone, has been described in the patent literature in the development of drugs for the treatment of inflammatory disorder and microbial disease.³⁷ Conversely, our method allows direct access to the quaternary center, yielding **30** in two steps after deprotection.

Stereochemistry on the alkyl iodide was found to translate well to the products, with no epimerization observed in the case of the iodoalanine (**39**), proline (**40**) or sugar substrates (**45**). Importantly, the use of iodoalanine provides enantiomerically pure access to unnatural amino acid **39**, providing an orthogonal approach to the racemic conjugate addition product from dehydroalanine.

Whereas benzyl iodides were poorly tolerated under the reaction conditions, we were please to find that benzyl bromides coupled in moderate yield (**47**, **48**). Furthermore, benzyl electrophiles provide an intriguing disconnection. An sp²–sp³ bond scission would transform the product to an arene and neopentyl crosscoupling partner, which retains structural complexity and oftentimes requires an organometallic reagent. Consequently, similar scaffolds have been accessed by formation of a mixed ketone, alkylation, and Wolff-Kishner deoxygenation—overall a 7-step sequence.³⁸ Disconnection to the benzyl electrophile and alkene allows scission of the quaternary carbon in a logical and simplifying transform.

While this method makes significant progress in the formation of sterically congested aliphatic centers, the transformation is sensitive to the steric environment on the alkyl halide. α -Branching (49), neopentyl (50) and secondary alkyl iodides (51) were found to proceed in low yield, predominantly lost to competitive protodehalogenation. This could imply that oxidative addition or a more sterically congested Ni center impedes productive reductive elimination.

In summary, we have reported a Markovnikovselective hydroalkylation of unbiased olefins³⁰ using diverse alkyl iodides and benzyl bromides. The combination of Mn-mediated MHAT catalysis and Ni catalysis enable an unprecedented synthesis of quaternary carbons. The mild reaction conditions and robust functional group compatibility support its utility for late stage modification of small molecules. Efforts are underway to expand this chemistry to more sterically congested centers and complex natural products.

Supporting Information.

The Supporting Information is available free of charge on the ACS Publications website.

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Notes

The authors declare no competing financial interest.

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REFERENCES

(1) (a) Crossley, S. W. M.; Martinez, R. M.; Obradors, C.; Shenvi, R. A. Mn, Fe, and Co-catalyzed radical hydrofunctionalization of olefins. *Chem. Rev.* **2016**, *116*, 8912. (b) Green, S. A.; Crossley, S. W. M.; Matos, J. L. M.; Vásquez-Céspedes, S.; Shevick, S. L.; Shenvi, R. A. The high chemofidelity of metal-catalyzed hydrogen atom transfer *Acc. Chem. Res.* **2018**, *51*, 2628.

(2) (a) Gaspar, B.; Carreira, E. M. Mild cobaltcatalyzed hydrocyanation of olefins with tosyl cyanide. *Angew. Chem. Int. Ed.* **2007**, *46*, 4519. (b) Gaspar, B.; Carreira, E. M. Cobalt catalyzed functionalization of unactivated alkenes: Regioselective reductive C-C bond forming reactions. *J. Am. Chem. Soc.* **2009**, *131*, 13214–13215.

(3) Leggans, E. K.; Barker, T. J.; Duncan, K. K.; Boger D. L. Iron(III)/NaBH₄-mediated additions to unactivated alkenes: synthesis of novel 20'-vinblastine analogues. *Org. Lett.* **2012**, *14*, 1428.

(4) (a) Lo, J. C.; Yabe, Y.; Baran, P. S. A practical and catalytic reductive olefin coupling. *J. Am. Chem. Soc.* **2014**, *136*, 1304. (b) Lo, J. C.; Gui, J.; Yabe, Y.; Pan, C-M.; Baran, P. S. Functionalized olefin cross-coupling to construct carbon–carbon bonds. *Nature* **2014**, *516*, 343.

(5) Dao, H. T.; Li, C.; Michaudel, Q.; Maxwell, B. D.; Baran, P. S. Hydromethylation of unactivated olefins. *J. Am. Chem. Soc.* **2015**, *137*, 8046.

(6) (a) Crossley, S. W. M.; Martinez, R. M.; Zuluaga,
S. G.; Shenvi, R. A. Synthesis of the privileged 8arylmenthol class by radical arylation of isopulegol. *Org. Lett.* 2016, 18, 2620. (b) Green, S. A.; Matos, J. L.
M.; Yagi, A.; Shenvi, R. A. Branch-selective hydroarylation: Iodoarene-olefin cross coupling. J. Am. *Chem. Soc.* 2016, 138, 12779. (c) Green, S. A.; Vásquez-Céspedes, S.; Shenvi, R. A. Iron-nickel dualcatalysis: A new engine for olefin functionalization. J. Am. Chem. Soc. 2018, 140, 11317.

(7) (a) Ma, X.; Herzon, S. B. Intermolecular hydropyridylation of unactivated alkenes. *J. Am. Chem. Soc.* **2016**, *138*, 8718. (b) Ma, X.; Dang, H.; Rose, J. A.;

Rablen P.; Herzon, S. B. Hydroheteroarylation of unactivated alkenes using *N*-methoxyheteroarenium salts. *J. Am. Chem. Soc.* **2017**, *139*, 5998. (c) Shigehisa, H.; Ano, T.; Honma, H.; Ebisawa, K.; Hiroya, K. Cocatalyzed hydroarylation of unactivated olefins. *Org. Lett.* **2016**, *18*, 3622. (d) Lo, J. C.; Kim, D.; Pan, C–M.; Edwards, J. T.; Yabe, Y.; Gui, J.; Qin, T.; Gutiérrez, S.; Giacoboni, J.; Smith, M. W.; Holland, P. L.; Baran, P. S. Fe-catalyzed C–C bond construction from olefins via radicals. *J. Am. Chem. Soc*, **2017**, *139*, 2484.

(8) Matos, J. L. M.; Vásquez-Céspedes, S.; Gu, J.; Oguma, T.; Shenvi, R. A. Branch-selective addition of unactivated olefins into imines and aldehydes. *J. Am. Chem. Soc.* **2018**, *140*, 16976.

(9) For representative examples of intermolecular Nicatalyzed sp³–sp³ alkylation reactions using alkyl halides see: (a) Choi, J.; Fu, G.C. Transition metal–catalyzed alkyl-alkyl bond formation: Another dimension in crosscoupling chemistry. *Science* **2017**, *356*, eaaf7230. (b) Yu, X.; Yang, T.; Wang, S.; Xu, H.; Gong, H. Nickelcatalyzed reductive cross-coupling of unactivated alkyl halides. *Org. Lett.* **2011**, *13*, 2138. (c) Smith, R. T.; Zhang, X.; Rincón, J. A.; Agejas, J.; Mateos, C.; Barberis, M.; García-Cerrada, S.; de Frutos, O.; MacMillan, D. W. C. Matallaphotoredox-catalyzed cross-electrophile Csp³–Csp³ coupling of aliphatic bromides. *J. Am. Chem. Soc. 140*, **2018**, 17433.

(10) For representative examples of intermolecular Nicatalyzed sp³–sp³ alkylation reactions using carboxylic acids see: (a) Qin, T.; Cornella, J.; Li, C.; Malins, L. R.; Edwards, J. T.; Kawamura, S.; Maxwell, B. D.; Eastgate, M. D.; Baran, P. S. A general alkyl-alkyl cross-coupling enabled by redox-active esters and alkylzinc reagents. *Science* **2016**, *352*, 801. (b) Johnston, C. P.; Smith, R. T.; Allmendinger, S.; MacMillan, D. W. C. Metallaphotoredox-catalysed sp³–sp³ cross-coupling of carboxylic acids with alkyl halides. *Nature* **2016**, *536*, 322.

(11) (a) Yan, M.; Lo, J. C.; Edwards, J. T.; Baran, P. S. Radicals: Reactive intermediates with translational potential. *J. Am. Chem. Soc.* **2016**, *138*, 12692. (b) Smith, J. M.; Harwood, S. J.; Baran, P. S. Radical retrosynthesis. *Acc. Chem. Res.* **2018**, *51*, 1807.

(12) Povie, G.; Suravarapu, S. R.; Bircher, M. P.; Mojzes, M. M.; Rieder, S.; Renaud, P. Radical chain repair: The hydroalkylation of polysubstituted unactivated alkenes. *Sci. Adv.* **2018**, *4*, eaat6031.

(13) Pitre, S. P.; Weires, N. A.; Overman, L. E. Forging $C(sp^3)$ - $C(sp^3)$ bonds with carbon-centered radicals in the synthesis of complex molecules. *J. Am. Chem. Soc.* **2019**, *141*, 2800.

(14) Tasker, S. Z.; Standley, E. A.; Jamison, T. F. Recent advances in homogeneous nickel catalysis.

Nature 2014, 509, 299.

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(15) For methods pertaining to formation of arylated quaternary centers using Ni see: (a) Zultanski, S. L.; Fu, G. C. Nickel-catalyzed carbon-carbon bond-forming reactions of unactivated tertiary alkyl halides: Suzuki arylations J. Am. Chem. Soc. 2014, 135, 624. (b) Wang, X.; Wang, S.; Xue, W.; Gong, H. Nickel-catalyzed reductive croupling of aryl bromides with tertiary alkyl halides. J. Am. Chem. Soc. 2015, 137, 11562. (c) Wang, X.; Ma, G.; Peng, Y.; Pitsch, C. E.; Moll, B. J.; Ly, T. D.; Wang, X.; Gong, H. Ni-catalyzed reductive coupling of electron-rich aryl iodides with tertiary alkyl halides. J. Am. Chem. Soc. 2018, 140, 14490. (d) Primer, D. N.; Molander, G. A. Enabling the cross-coupling of tertiary organoboron nucleophiles through radical-mediated alkyl transfer. J. Am. Chem. Soc. 2017, 139, 9847. (e) Chen, T. -G.; Zhang, H.; Mykhailiuk P. K.; Merchant, R. R.; Smith, C. A.; Qin, T.; Baran, P. S. Quaternary centers via Ni-catalyzed cross-coupling of tertiary carboxylic acids and aryl zinc reagents. Angew. Chem. Int. Ed. 2019, 58, 2454.

(16) For the formation of allylated quaternary centers using Ni see: Chen, H.; Jia, X.; Yu, Y.; Qian, Q.; Gong, H. Nickel-catalyzed reductive allylation of tertiary alkyl halides with allylic carbonates. *Angew. Chem. Int. Ed.*, **2017**, *56*, 13103.

(17) Wang, Z.; Yin, H.; Fu, G. C. Catalytic enantioconvergent coupling of secondary and tertiary electrophiles with olefins. *Nature* **2018**, *563*, 379.

(18) (a) Qin, T.; Malins, L. R.; Edwards, J. T.; Merchant, R.R.; Novak, A. J.; E.; Zhong, J. Z.; Mills, R. B.; Yan, M.; Yuan, C.; Eastgate, M. D.; Baran, P. S. Nickel-catalyzed Barton decarboxylation and Giese reactions: A practical take on classic transforms. *Angew. Chem., Int. Ed.* 2017, 56, 260. (b) Ye, Y.; Haifeng, C.; Sessler, J. L.; Gong, H. Zn-mediated fragmentation of tertiary alkyl oxalates enabling formation of alkylated and arylated quaternary carbon centers. *J. Am. Chem. Soc.* 2019, *141*, 820.

(19) Nguyen, K. D.; Park, B. Y.; Luong, T.; Sato, H.; Garza, V. J.; Krische, M. J. Metal-catalyzed reductive coupling of olefin-derived nucleophiles: Reinventing carbonyl addition. *Science*, **2016**, *354*, 300.

(20) For representative examples of olefins used in Nicatalyzed cross coupling see: (a) Lu, X.; Xiao, B.; Zhang, Z.; Gong, T.; Su, W.; Yi, J.; Fu, Y.; Liu L. Practical carbon-carbon bond formation from olefins through nickel-catalyzed reductive olefin hydrocarbonation. Nat. Comm. 2016, 7. Article number 11129. (b) Lu, X.; Xiao, B.; Liu, L.; Fu, Y. Formation of through $C(sp^3)-C(sp^3)$ bonds nickel-catalyzed decarboxylative olefin hydroalkylation reactions. Chem. Eur. J. 2016, 22, 11161. (c) Sun, S-Z.; Börjesson, M.;

Martin-Montero, R.; Martin, R. Site-selective Nicatalyzed reductive coupling of α -haloboranes with unactivated olefins. *J. Am. Chem. Soc.* **2018**, *140*, 12765.

(21) Anderson, T. J.; Jones, G. D.; Vicic, D. A. Evidence for a Ni¹ active species in the catalytic cross-coupling of alkyl electrophiles. *J. Am. Chem. Soc.* **2004**, *126*, 8100.

(22) Anka-Lufford, L. L.; Huihui, K. M. M.; Gower, N. J.; Ackerman, L. K. G.; Weix, D. J. Nickel-catalyzed cross-electrophile coupling with organic reductants in non-amide solvents. *Chem. Eur. J.* **2016**, *22*, 11564.

(23) For more optimization results, see SI.

(24) (a) Inoki, S.; Kato, K.; Isayama, S.; Mukaiyama, T. A New and Facile Method for the Direct Preparation of α -Hydroxycarboxylic Acid Esters from α , β -Unsaturated Carboxylic Acid Esters with Molecular Oxygen and Phenylsilane Catalyzed by Bis(dipivaloylmethanato) manganese(II) Complex. Chem. Lett. 1990, 19, 1869. (b) Magnus, P.; Payne, A. H.; Waring, M. J.; Scott, D. A.; Lynch, V. Conversion of α , β -Unsaturated Ketones into α -Hydroxy Ketones using an Mn^{III} Catalyst, Phenvlsilane and of Dioxygen: Acceleration Conjugate Hydride Reduction by Dioxygen. Tetrahedron Lett. 2000, 41, 9725.

(25) For further discussion on the role of HFIP, see SI.

(26) The products of **1** with olefins **24a** and **27a** were inseparable from residual alkyl iodide. Protected

iodoethanol substrates were used for ease of purification. (27) Waser, J.; Carreira, E. M. Catalytic hydrohydrazination of a wide range of alkenes with a simple Mn complex. *Angew. Chem. Int. Ed.* **2004**, *31*, 4191.

(28) Carney, J. R.; Dillon, B. R.; Campbell, L.; Thomas, S. P. *Angew. Chem. Int. Ed.* **2018**, *57*, 10620.

(29) Iwasaki, K.; Wan, K. K.; Oppedisano, A.; Crossley, S. W. M.; Shenvi, R. A. Simple, Chemoselective Hydrogenation with Thermodynamic Stereocontrol *J. Am. Chem. Soc.* **2014**, *136*, 1300.

(30) 'Bias' refers, in this context, to electronic bias by conjugation to another pi-system or heteroatom.

(31) Using (–)-limonene as the olefin gave a complex mixture of products with reactivity observed at both olefins. Overall, compounds containing multiple olefins did not provide selectivity. For other examples, see SI.

(32) The protecting group on the glucal was found to be important for high regioselectivity. The use of acetate in place of benzyl resulted in a mixture of regioisomers (C2 vs C3 alkylation), resulting from an acetate-directed Ni-catalyzed alkylation at C3. A similar directing effect

was observed using esters in ref 17.

(33) Abe, H.; Shuto, S.; Matsuda, A. Highly α - and β selective radical C-glycosylation reactions using a controlling anomeric effect based on the conformational restriction strategy. A study on the Conformation– Anomeric Effect–stereoselectivity relationship in anomeric radical reactions. J. Am. Chem. Soc. 2001, 123, 11870.

(34) For an alternative disconnection to this molecule see: Lardy, S. W.; Schmidt, V. A. Intermolecular radical mediated anti-Markovnikov alkene hydroamination using *N*-hydroxyphthalimide. *J. Am. Chem. Soc.* **2018**, *140*, 12318.

(35) Kim, J. H.; Cung, Y. K. Copper Nanoparticlecatalyzed cross-coupling of alkyl halides with Grgnard reagents. *Chem. Commun.* **2013**, *49*, 11101.

(36) Similar Kumada couplings have been described by Kombe, but the transformation remains largely restricted to the installation of a *t*-butyl group. Iwasaki, T.; Takagawa, H.; Singh, S. P.; Kuniyasu, H.; Kambe, N. Co-catalyzed cross-coupling of alkyl halides with tertiary alkyl Grignard reagents using a 1,3-butadiene additive. J. Am. Chem. Soc. **2013**, 135, 9604.

(37) Siddiqui, M. A.; Mansoor, U. F.; Reddy, P. A. P.; Madison, V. S. Compounds for the treatment of inflammatory disorders and microbial diseases. US patent US2007/0167426 A1. **2017**.

(38) Becknell, N. C.; Dandu, R. R.; Dorsey, B. D.; Gotchev, D. B.; Hudkins, R. L.; Weinberg, L.; Zificsak, C. A.; Substituted 4-benzyl and 4-benzyl piperidine derivatives. International patent: WO2016/205590 A1. .

TOC Graphic

