Walking Metals for Remote Functionalization

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ABSTRACT: The distant and selective activation of unreactive C–H and C–C bonds remains one of the biggest challenges in organic chemistry. In recent years, the development of remote functionalization has received growing interest as it allows for the activation of rather challenging C–H and C–C bonds distant from the initiation point by means of a “metal-walk”. A “metal-walk” or “chain-walk” is defined by an iterative series of consecutive 1,2- or 1,3-hydride shifts of a metal complex along a single hydrocarbon chain. With this approach, simple building blocks or mixtures thereof can be transformed into complex scaffolds in a convergent and unified strategy. A variety of catalytic systems have been developed and refined over the past decade ranging from late-transition-metal complexes to more sustainable iron- and cobalt-based systems. As the possibilities of this field are slowly unfolding, this area of research will contribute considerably to provide solutions to yet unmet synthetic challenges.

Initiated by the pioneering work of Breslow in the early 1970s for biomimetic transformations and later used by Schwarz in gas-phase chemistry, the concept of remote functionalization consists of an indirect activation of a site distant from the initial functional group (Figure 1a). This distant activation of an “unactivated position” strongly modifies our perception of organic transformations which was mostly concentrated on direct transformation of the chemically most reactive position. For instance, the concept of activation of carbon–hydrogen bonds, coined as C−H activation, has the ability to decorate hydrocarbons of all couler from simple arenes to highly functionalized natural products. To achieve control of the selectivity of these transformations, the inherent directionality of the substrate is usually exploited (Figure 1b). This goal is commonly achieved by utilizing directing groups, present or preinstalled, or through the undirected activation of particularly reactive C−H bonds. While the field of directed activation of C−H bonds matured considerably in the past 20 years, the undirected process poses still significant challenges limiting its applicability to C−H bonds “within reach” of the catalyst.

Mechanistic Aspects of Olefin Isomerization. Olefin isomerization can occur via either 1,2- or 1,3-hydride shift mechanism, and both pathways can potentially compete with each other. In the 1,2-hydride shift mechanism, the metal−hydroxide complex undergoes migratory insertion into an olefin, commonly referred to as hydrometalation reaction, giving rise to a well-defined alkyl−metal species (Figure 2a). Subsequent β-hydride elimination furnishes the isomerized olefin π-complex, which undergoes rotation and hydrometallation.

Alternatively, olefin isomerization can occur via a 1,3-hydride shift mechanism through an inner- or outer-sphere mechanism. The inner-sphere mechanism requires two vacant orbitals for this process: one for olefin coordination and the second one for

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the C–H allylic activation to occur (Figure 2b). After formation of the \( \eta^3 \)-allyl complex, reductive elimination provides the isomerized olefin–metal complex. The vacant filled orbital can also be replaced by a strategically placed base within the ligand sphere of the catalyst (Figure 2c). Upon olefin coordination and allylic activation, this base can abstract the allylic proton to furnish the \( \eta^3 \)-allyl complex, which undergoes reductive elimination to give the isomerized olefin. Finally, some \( \pi \)-acidic transition metals such as cationic silver and palladium complexes are able to acidify the allylic position upon olefin coordination (Figure 2d). Intermolecular deprotonation subsequently furnishes the \( \eta^3 \)-allyl complex, which rapidly undergoes protodemetalation resulting in overall olefin migration. Independent of the mechanism, the directionality of the “chain-walking” depends on the terminating event as all steps in the isomerization sequence are in rapid equilibrium. Consequently, a variety of catalytic systems with unique termination steps have been developed and are discussed in the following with representative examples.

Titanium and Zirconium Complexes. The isomerization of branched Grignard reagents to linear compounds via the formation of olefin intermediates under titanium or zirconium catalysis has been reported as early as 1961. But only in the past two decades have efforts been devoted toward the development of protocols that allowed the extension of these findings for the selective isomerization of olefins. Most notably, Schwartz’s reagent (\( \text{Cp}_2\text{Zr(H)Cl} \))\(^{14,15} \) and Negishi reagent (\( \text{Cp}_2\text{Zr(C}_5\text{H}_5) \))\(^3\) exhibit high activity for the isomerization of olefins and dienes. These reagents are easily accessible from zirconocene dichloride (\( \text{Cp}_2\text{ZrCl}_2 \)) and can be readily prepared in situ. “Chain-walking” of linear olefins occurs under mild conditions whereas hindered substrates typically require higher temperatures. For example, simple fatty alcohols underwent a sequence comprising olefin isomerization, elimination, and

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**Figure 1.** “Metal-walk” vs classical modes of activation: (a) general concept of remote functionalization; (b) directed C–H activation at close proximity; (c) metal-walk for remote functionalization.

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**Figure 2.** Mechanistic intricacies of olefin isomerization: (a) 1,2-hydride shift mechanism; (b) 1,3-hydride shift; (c) 1,3-hydride shift with a ligand acting as a base; (d) 1,3-proton shift.
electrophilic trapping in the presence of stoichiometric amounts of the Negishi reagent (Figure 3a).\textsuperscript{17} The \textit{in situ} formed alkoxy-zirconocene species serves as a leaving group to yield the required allylzirconocene. As double bond geometry and position in the starting material are inconsequential of the reaction outcome, a convergent synthesis from isomeric mixtures can be realized.

If simple olefins are employed in the presence of Schwartz’s reagent, the resulting terminal alkylzirconium species are suitable reaction partners in asymmetric conjugate additions (Figure 3b).\textsuperscript{18} Recently it was found that strategically positioned cyclopropanes offer the possibility to achieve bisfunctionalization of olefins by means of zirconium “chain-walk”, C–C bond cleavage, and electrophilic trapping.\textsuperscript{19,20} The transiently formed allylzirconacyclobutane selectively reacts first with carbonyl electrophiles at the allylic position. Subsequent electrophilic activation of the remaining alkylzirconium bond then furnishes the product (Figure 3c).\textsuperscript{21} Although currently stoichiometric amounts of zirconium reagent have to be employed, these results might pave the way for the implementation of a catalytic methodology. Furthermore, the development of titanium-based protocols appears desirable under sustainability viewpoints considering the natural abundance of both elements.

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**Ruthenium, Rhodium, and Iridium Catalysis.** In 1974, Wells and co-workers described the isomerization of 1-pentene to 2-pentene under homogeneous ruthenium catalysis.\textsuperscript{21} Following these initial findings, different research groups investigated this transformation in more detail and eventually discovered that internal olefins undergo this transformation as well. Furthermore, it was also found that \(\alpha,\beta\)-unsaturated carbonyls readily underwent deconjugative isomerization, favoring the formation of enol ethers.\textsuperscript{22} The isomerization of olefins over a large number of positions has also been achieved utilizing a bifunctional catalyst via the outer-sphere \(1,3\)-hydride shift mechanism. In the presence of this “alkene-zipper” catalyst, olefin isomerizations over more than 30 positions became possible (Figure 4a).\textsuperscript{23} In combination with an olefin metathesis catalyst (W-1), a tandem olefin isomerization/metathesis sequence has been applied for the synthesis of long-chain olefins (Figure 4b)\textsuperscript{24} or phenylproponoids.\textsuperscript{25} With the finding of Wilkinson and Osborn that PPh\(_3\) modified rhodium catalysts undergo hydroformylation reactions of terminal olefins,\textsuperscript{26,27} this field underwent impressive developments in the past 50 years.\textsuperscript{28} In the past two decades, catalytic systems have been developed that favor an isomerization/hydroformylation sequence, even starting from mixtures of olefins, to furnish linear aldehydes and derivatives thereof (Figure 4c).\textsuperscript{30–32} Furthermore, a rhodium-catalyzed olefin isomerization/Michael addition sequence has been developed based on the Rh/BIPHEPHOS system (Figure 5a).\textsuperscript{33} Based on earlier findings that cationic iridium complexes in the presence of hydrogen readily promote olefin isomerization,\textsuperscript{34,35} a protocol for the tandem isomerization/enantioselective allylation was developed.\textsuperscript{36} Under the same reaction conditions,
A tandem hydroboration/isomerization/allylation sequence was demonstrated as well (Figure 5b).

As can be seen from these examples, the true potential of long-range olefin isomerization can be unleashed when coupled with an ensuing coupling reaction. Furthermore, the catalytic systems employed demonstrate a reasonable chemoselectivity, allowing a variety of tandem reactions to be conducted.

**Iron Catalysis.** Since its disclosure in 1964,47 the iron-catalyzed olefin isomerization has witnessed renewed interest in the past decade.18,39 With continuing efforts being made toward more sustainable catalytic systems, iron catalysts offer a promising alternative to late transition metals. Although to date only a few examples on “chain-walking” have been reported, this field will likely witness important advancements in the upcoming years.

In 2013, a series of mild iron-catalyzed olefin isomerization/hydroborations and hydroisilylations of a variety of internal olefins were reported.40,41 The corresponding iron complexes show similar reactivity although their high sensitivity toward air and moisture render them particularly difficult to handle. Therefore, a protocol for in situ activation of an iron precursor has been developed and applied to a tandem dehydrogenation/isomerization/hydroboration sequence (Figure 6).42

**Cobalt Catalysis.** Prompted by Brookhart’s studies on the hydrosilylation of alkenes with cyclopentadienyl cobalt species,43 a Co-catalyzed isomerization/hydroboration of alkenes using pinacolborane (HBPin) with bis(imino)pyridine ligands was reported (Figure 7a).44 The utilization of aliphatic internal olefins resulted in the formation of hydroboration products.
where the BPin residue was invariably located at the terminal reaction site. The presence of electron-donating amino groups at the para position of the pyridine ring in the ligand culminated in more active catalysts (Co-2), which were able to convert cis- and/or trans-4-octene into 1-octylboronic ester at room temperature with excellent yields and linear selectivity. Labeling experiments showed deuterium incorporation at various positions on the aliphatic chain, suggesting a chain-walking migration via consecutive β-hydride elimination/olefin reinsertion. The reaction was later extended to more challenging hindered tri- and tetrasubstituted alkenes by using redox-active terpyridine and α-diimine ligands. Later on, it was univocally shown that site selectivity could be tuned and controlled depending on the ligand backbone. Specifically, terminal olefins decorated with a pending arene underwent selective C–B bond-forming reaction at the benzylic position with [(PPh3)3CoH(N2)] (Co-3) whereas the employment of Co-1 supported by nitrogen donors gave rise to linear-selective borylation (Figure 7b). The utilization of N-phosphinoamidinate ligands has been shown to be particularly suited in cobalt-catalyzed hydroboration of unactivated olefins via Co-4 (Figure 7c). Unlike other nitrogen donors, the isomerization/hydroboration was not limited to HBpin and could be extended to 1,3-dimethyl-1,3-diaza-2-boracyclopentane or benzo-1,3,2-diazaborolane with equal
ease. The dichotomy exerted by the boron source was further illustrated by the utilization of 2-ethyl-1-butene as counterpart; while HBPin resulted in a mixture of linear alkylboronic esters, clean C−B bond formation at the less-hindered site was achieved with benzo-1,3,2-diazaborolane.

Cobalt complexes have shown a superior performance in olefin isomerization/hydrosilylation when compared to precious metals used for similar means, and intriguing cooperative effects of well-defined Co(I) catalysts supported by β-diketinate ligands were described. In 2014, a dehydrogenative silylation of alkenes by using bis(imino)pyridine cobalt complexes was discovered, ending up in allylsilanes with the C−Si bond formation occurring exclusively at the terminal reaction site (Figure 8). A sequential site-selective dehydrogenative silylation was within reach, offering an opportunity to promote multiple C−Si bond-forming reactions from simple precursors. Stoichiometric experiments revealed the initial formation of Co−SiR₃ active species with concomitant formation of methane upon exposure of the cobalt complex Co-1 to R₃SiH. Preferential formation of the allyl- versus vinylsilane was consistent with a preferred β-hydride elimination distal to the bulky tertiary silyl group.

The involvement of cobalt catalysts in chain-walking scenarios is by no means limited to hydroboration or hydrosilylation reactions. Indeed, C−C bond formations at distal reaction sites could also be achieved via the intermediacy of cobalt hydride species generated by C−H activation in the presence of indole scaffolds decorated with imine directing groups followed by a sequence of olefin insertion/β-hydride elimination (Figure 9). Subtle changes in the reaction conditions resulted in a regio-divergent scenario with unactivated olefins possessing a pending arene on the side chain, accessing either linear or branched products with excellent site selectivities. Specifically, the utilization of N-heterocyclic carbenes at 60 °C allowed for an olefin isomerization en route to the thermodynamically more stable styrene prior to C−C bond formation at the benzylic position, whereas linear selectivity was observed in the absence of N-heterocyclic carbene with tBuCH₂MgBr as promoter. While the role of the ligand still remains unclear, the differences on both tBuCH₂MgBr and CyMgBr are likely attributed to the presence of β-hydrogens in the latter, thus setting the basis for the generation of cobalt hydride species. Recently, the generality of cobalt bis(imino)pyridine complexes in chain-walking scenarios was extended by showing the viability to conduct hydroaluminations of internal olefins via an initial isomerization prior to C−Al bond formation.

Palladium Catalysis. Despite the rapid development of a range of catalytic systems to induce remote functionalization via “chain-walking”, this area of research is largely dominated by palladium-based catalysts. The ready availability of palladium complexes, mild reaction conditions, and high functional group tolerance are key factors for its popularity. Additionally, most palladium (pre)catalysts can easily be handled outside a glovebox, advocating their robustness and user-friendliness. The strong propensity of alkyl−palladium complexes to undergo
β-hydride elimination is a well-studied phenomenon in organic chemistry with which every practitioner is familiar, and ligands favoring this elimination came recently into the focus of research. “Chain-walking” or “chain-running” palladium (and nickel catalysts) had already made their way into polymerization chemistry but were largely neglected in organic synthesis. The first reports on palladium-catalyzed Heck reaction combined with a metal-walk on ω-alkenols were disclosed in 1976 providing low yields of a mixture of products under harsh reaction conditions. These early examples were subsequently extended to the remote functionalization of longer-chain olefinic alcohols to yield the corresponding carbonyls (Figure 10a). Later, a protocol for the enantioselective remote functionalization of olefinic alcohols was disclosed utilizing chiral PyrOx ligands under mild conditions (Figure 10b). Such a transformation enables the formation of acyclic quaternary stereocenters in high enantioselectivity, a long-standing challenge in organic synthesis, from trisubstituted olefins using different coupling partners such as diazonium salts, boronic acids, or vinyl triflates (Figure 10c). A similar protocol was found to promote the remote functionalization of olefinic alcohols comprising a cyclopropane in the side chain with concomitant selective ring cleavage (Figure 10d). Alternatively, the “chain-walking” can also be initiated by an in situ formed palladium hydride, resulting in an overall long-range redox isomerization of olefinic alcohols. Deconjugative redox isomerization starting

Figure 10. Pd-catalyzed “chain-walking” via C–C bond formations terminated by carbonyl formation: (a) Pd-catalyzed isomerization of alkenyl alcohols; (b) enantioselective Pd-catalyzed isomerization of alkenols; (c) enantioselective Heck reaction on trisubstituted alkenols; (d) palladium-catalyzed Heck isomerization as a trigger for selective ring-opening of cyclopropanes; (e) isomerization of unsaturated alcohols promoted by metal–hydride catalysts.
from α/β-unsaturated carbonyls has been disclosed as well (Figure 10e).74

Instead of relying on carbonyl formation as the terminating event, other strategies exploited the increased stability of allylpalladium species opposed to the initial alkylpalladium. Nucleophilic trapping was accomplished with a variety of carbon and heteroatom nucleophiles giving rise to remotely bisfunctionalized products.75−77 This strategy was also utilized in the total syntheses of pyridine alkaloids (Figure 11a).78 If the substrate contains a strategically positioned olefin, an ensuing cyclization of the palladium intermediate gives rise to various cyclopentane scaffolds. Although this catalytic system has been extensively utilized in the cycloisomerizations of 1,6-dienes,79,80 this methodology was only recently extended to a tandem sequence of "chain-walking"/cyclization (Figure 11b).81,82 Contrary to the examples presented above, it has also been shown that the palladium(II) species generated from oxidative insertion can undergo "chain-walking" prior to reductive elimination. This strategy was successfully applied to the remote arylation of amino acid derivatives.83 Later it was demonstrated that a linear selective Negishi-type cross-coupling was also feasible starting from mixtures of branched alkyl bromides (Figure 11c).84

Nickel Catalysis. Prompted by the prospective impact of nickel catalysts in the Shell higher olefin process (SHOP) for the homo- and copolymerization of ethylene or α-olefins,85,86 as well as the DuPont process for preparing adiponitrile via hydrocyanation of butadiene,87 the past years have witnessed an increasing interest in nickel-catalyzed olefin isomerization as a vehicle for remote functionalization. In 2015, a mild chemo-selective alkene isomerization/hydrosilylation catalyzed by nickel pincer complexes was reported in which the putative nickel hydride species were generated upon exposure to both metal alkoxides and silanes (Figure 12a).88,89 A significant improvement was observed with nickel nanoparticles, allowing the scope of these reactions to be extended to tertiary silanes such as (EtO)3SiH (Figure 12b). In line with previous findings with both iron and cobalt catalysis, redox-active α-diimine ligands were found to be competent in nickel-catalyzed hydroxylation reactions. Labeling studies with (EtO)3SiD and 1-octene resulted in deuterium incorporation at multiple sites throughout the alkyl side chain.90 This seemingly innocent experiment showed the inherent ability of nickel catalysts to promote an olefin isomerization prior to C−Si bond formation, an observation that could further be corroborated by observing linear hydrosilylation with 4-octene as counterpart.

The first example of a nickel-catalyzed olefin isomerization/hydroarylation of alkenes was disclosed via initial C−H activation or oxidative addition reaction.
C–H bonds and N-heterocyclic carbenes as ligand donors (Figure 13a).90 Intriguingly, the presence of AlMe₃ inhibited olefin isomerization, obtaining linear-selective hydroarylation products. A related hydroarylation reaction but with electron-deficient arenes was also reported (Figure 13b).91 Although this reaction formally falls into the category of a classical “chain walk”, labeling studies and theoretical calculations left a reasonable doubt about the intermediacy of nickel hydride intermediates, arguing against an initial oxidative addition into the C–H bond. The authors suggested that the origin of the linear selectivity arises from the C–C bond-forming reductive elimination rather than the difference in stabilities of the linear versus branched alkyl metal complexes. More recently, a site-selective linear alkylation of anilides using nickel catalysts and aluminum-based Lewis acids was reported with 2-octene as coupling partner.92

Aryl halides can be readily utilized as counterparts in nickel-catalyzed olefin isomerization followed by site-selective C–C bond formation. In early 2017, a site-selective catalytic hydroarylation of alkynes occurring exclusively at benzylic sp³ C–H sites under a Ni/bipyridine regime was reported, using polymethylhydrosiloxane (PMHS) both as reducing agent and as hydride source (Figure 14a).93 Prompted by a recent report that used light alkyl bromides as hydride sources via β-hydride elimination in nickel-catalyzed hydroamidations of alkynes,94 an alternative hydroarylation technique with comparable yields and selectivities to those observed in their preceding protocol based on PMHS was subsequently reported (Figure 14b).95 Such a protocol could be extended to a formal cross-electrophile remote functionalization by using organic halides in lieu of olefin partners.96

Among the various conceivable scenarios, including directing group methodologies, radical-type pathways, or enzymatic protocols, the ability of a metal to cause a dynamic displacement throughout a saturated hydrocarbon chain (“chain-walk”) is particularly appealing.

In the past decade, nickel-catalyzed cross-coupling reactions of unactivated alkyl halides have matured into robust and
reliable technologies for introducing saturated hydrocarbon side chains into organic backbones. These reactions invariably occur at the initial sp3 C–H bond via formal functional group interconversion, contributing to the perception that a nickel-catalyzed “chain-walk” of unactivated alkyl halides initiated by oxidative addition would be particularly problematic. In 2017, a remote sp3 C–H carboxylation of unactivated alkyl bromides under atmospheric pressure of carbon dioxide (CO2) was designed (Figure 15a). Unlike related nickel-catalyzed “chain-walking” hydroarylation events, this transformation occurred selectively at the primary sp3 C–H bond, even in the presence of arenes on the side chain, thus offering an opportunity to activate less-reactive reactive sites in the presence of a priori more reactive ones. A one-pot bromination/carboxylation sequence could be used as a platform for directly converting alkanes or unreified mixtures of olefins into fatty acids as single regioisomers. Site selectivity could be tuned and controlled among different sp3 C–H bonds by a subtle thermal modulation in the presence of carbonyl groups on the side chain. Intriguingly, no significant erosion in enantioselectivity was observed in the presence of stereogenic centers, tacitly suggesting that the nickel catalyst remains bound to the olefin intermediate throughout the “chain-walk” along the hydrocarbon side chain.

Subsequently, it has recently been shown that water—typically used as proton source—can be harnessed as a safe, inexpensive hydride source in a remote carboxylation at sp3 C–H sites of unreified mixtures of olefins under atmospheric pressure of CO2 (Figure 15b). Such a method offered an opportunity to formally repurpose three renewable chemical feedstocks (olefins, CO2, and water) while reducing the chemical footprint of the hydride source in “chain-walking” scenarios by using water in lieu of high molecular weight silanes or organometallic reagents.

■ SUMMARY AND OUTLOOK

The functionalization of distal reaction sites has recently gained considerable momentum in the community. This popularity is largely due to the notion that prefunctionalization is not required at the targeted reaction site, thus changing prevailing perceptions...
in synthetic organic chemistry when building up molecular complexity. Among the various conceivable scenarios, including directing group methodologies, radical-type pathways, or enzy-
matric protocols, the ability of a metal to cause a dynamic displacement throughout a saturated hydrocarbon chain (“chain-walking”) is particularly appealing. This interest arises from the utilization of simple hydrocarbons as starting precursors, renew-
able feedstocks of utmost relevance for our chemical industry, and the avoidance of directing groups that are not particularly trivial to modify. The key aspect of “chain-walking” scenarios relies on the utilization of competent organometallic species that trigger a rapid olefin isomerization prior to the subsequent C–C or C–heteroatom bond formation at a previously unfunc-
tionalized reaction site.

The wealth of recent literature on catalytic “chain-walking” reactions suggests that these rather unconventional methodologies represent a powerful alternative to existing protocols for promoting functionalization of distal reaction sites. As for other cross-coupling reactions, the role exerted by the supporting ligand to fine-tune the properties of putative reaction intermediates is rather critical. Indeed, such modulation can result in a tunable and controllable multidirectional “chain-walking”, enabling regiodivergent scenarios via site-selective discrim-
ination among differently substituted remote reaction sites on a hydrocarbon chain. A wide number of daunting challenges remain, however. Among these, the development of enantio-
selective chain-walking scenarios is still at its infancy, with the available portfolio indicating that the chiral center should be introduced at a reaction site that originally contained an olefin residue. From both a conceptual and practical standpoint, it would be particularly attractive to set the chirality at a previ-
ously unfunctionalized reaction site or the desymmetrization of a prochiral substrate through the “chain-walking event”. Likewise, the discovery of new chemical knowledge occurring at the boundaries between classical organometallic catalysis and radical pathways or enzyme catalysis will likely invigorate the field by borrowing the best features associated with these disciplines.

Despite the elegant empirical discoveries realized in “chain-
walking” reactions, however, the mechanisms by which many of these reactions operate remain, in most instances, rather speculative and are based on indirect evidence by isotope labeling. This is probably attributed to the in situ formation of short-lived, yet exceptionally sensitive, organometallic entities that render their isolation and characterization a rather challenging task. Undoubtedly, efforts toward this goal will have a significant impact on the field of “chain-walking”, as under-
standing the mechanistic intricacies at a molecular level will enable the design of conceptually novel technologies not apparent at first sight. In view of the recent meteoric develop-
ment in “chain-walking” strategies, it is inevitable to predict spectacular progress in the years to come, providing a techn-
ological push that will change prevailing dogmas in retrosynthetic analysis.

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REFERENCES

(1) Breslow, R. Centenary Lecture. Biomimetic chemistry. Chem. Soc. Rev. 1972, 1 (4), 553.
(2) Breslow, R. Biomimetic control of chemical selectivity. Acc. Chem. Res. 1980, 13 (6), 170.
(3) Schwarz, H. Remote functionalization of C–H and C–C bonds by ‘naked’ transition-metal ions (Cosi Fan Tutte). Acc. Chem. Res. 1989, 22 (8), 282.
(4) Labinger, J. A.; Bercaw, J. E. Understanding and exploiting C–H bond activation. Nature 2002, 417, 507.
(5) Bergman, R. G. C–H activation. Nature 2007, 446, 391.
(6) Gensch, T.; Hopkinson, M. N.; Glorius, F.; Wencel-Delord, J. Mild metal-catalyzed C–H activation: examples and concepts. Chem. Soc. Rev. 2016, 45 (10), 2900.
(7) Davies, H. M. L.; Morton, D. Collective Approach to Advancing C–H Functionalization. ACS Cent. Sci. 2017, 3 (9), 936.
(8) Hartwig, J. F.; Larsen, M. A. Undirected, Homogeneous C–H Bond Functionalization: Challenges and Opportunities. ACS Cent. Sci. 2016, 2 (5), 281.
(9) Biswas, S. Mechanistic Understanding of Transition-Metal-Catalyzed Olefin Isomerization: Metal-Hydride Insertion-Elmination vs. π-Allyl Pathways. Comments Inorg. Chem. 2015, 35 (6), 300.
(10) Crabtree, R. H. In The Organometallic Chemistry of the Transition Metals; John Wiley & Sons, Inc.: New Haven, CT, 2014; p 224.
(11) Finkbeiner, H.; Cooper, G. Titanium-Catalyzed Isomerization and Olefin-Exchange Reactions of Alkylmagnesium Halides: A Novel Method for Preparation of the Grignard Reagent. J. Org. Chem. 1961, 26 (4), 4779.
(12) Cooper, G. D.; Finkbeiner, H. L. Titanium-Catalyzed Rearrangement and Olefin-Exchange of Grignard Reagents. J. Org. Chem. 1962, 27 (5), 1493.
(13) Prosenc, M.-H.; Brintringer, H.-H. Zirconium–Alkyl Isomer-
izations in Zirconocene-Catalyzed Olefin Polymerization: A Density Functional Study. Organometallics 1997, 16 (18), 3889.
(14) Carr, D. B.; Schwartz, J. Preparation of organoaluminum compounds by hydrozirconation-transmetalation. J. Am. Chem. Soc. 1979, 101 (13), 3521.
(15) Buchwald, S. L.; LaMaire, S. J.; Nielsen, R. B.; Watson, B. T.; King, S. M. A Modified Procedure for the Preparation of Cp2Zr(H)Cl (Schwartz’s Reagent). Tetrahedron Lett. 1987, 28 (34), 3895.
(16) Negishi, E.-I.; Takahashi, T. Patterns of Stoichiometric and Catalytic Reactions of Organozirconium and Related Complexes of Synthetic Interest. Acc. Chem. Res. 1994, 27 (5), 124.
(17) Marek, I.; Chinkov, N.; Levin, A. A Zirconium Promenade - An Efficient Tool in Organic Synthesis. Synlett 2006, 2006 (04), 0501.
(18) Mola, L.; Sidera, M.; Fletcher, S. P. Asymmetric Remote C–H Functionalization: Use of Internal Olefins in Tandem Hydro-
metallation–Isomerization–Asymmetric Conjugate Addition Sequences. Aust. J. Chem. 2015, 68 (3), 401.
(19) Masarwa, A.; Didier, D.; Zabrodski, T.; Schinkel, M.; Ackermann, L.; Marek, I. Merging allylc carbon-hydrogen and selective carbon-carbon bond activation. Nature 2014, 505 (7482), 199.
(61) Werner, E. W.; Mei, T.-S.; Burckle, A. J.; Sigman, M. S. Enantioselective Heck Arylations of Acrylic Alkenyl Alcohols Using a Redox-Relay Strategy. J. Org. Chem. 2012, 78 (13), 4855.

(62) Mei, T.-S.; Werner, E. W.; Burckle, A. J.; Sigman, M. S. Enantioselective Redox-Relay Oxidative Heck Arylations of Acrylic Alkenyl Alcohols using Boronic Acids. J. Am. Chem. Soc. 2013, 135 (16), 5830.

(63) Hilton, M. J.; Xu, L.-P.; Norrby, P.-O.; Wu, Y.-D.; Wiest, O.; Sigman, M. S. Investigating the Nature of Palladium Chain-Walking in the Enantioselective Redox-Relay Heck Reaction of Alkenyl Alcohols. J. Org. Chem. 2014, 79 (24), 11841.

(64) Mei, T.-S.; Patel, H. H.; Sigman, M. S. Enantioselective construction of remote quaternary stereocentres. Nature 2014, 508, 340.

(65) Xu, L.; Hilton, M. J.; Zhang, X.; Norrby, P.-O.; Wu, Y.-D.; Sigman, M. S.; Wiest, O. Mechanism, Reactivity, and Selectivity in Palladium-Catalyzed Redox-Relay Heck Arylations of Alkenyl Alcohols. J. Org. Chem. 2014, 79 (33), 13750.

(66) Patel, H. H.; Sigman, M. S. Palladium-Catalyzed Enantioselective Heck Alkenylation of Alkyl Alkenyl Alcohols Using a Redox-Relay Strategy. J. Am. Chem. Soc. 2015, 137 (10), 3462.

(67) Zhang, C.; Santiago, C. B.; Crawford, J. M.; Sigman, M. S. Enantioselective Dehydrogenative Heck Arylations of Trisubstituted Alkenes with Indoles to Construct Quaternary Stereocentres. J. Am. Chem. Soc. 2015, 137 (50), 15668.

(68) Zhang, C.; Santiago, C. B.; Kou, L.; Sigman, M. S. Enal Carbyl Derivatives in Enantioselective Redox Relay Heck Reactions: Accessing αβ-Unsaturated Systems. J. Am. Chem. Soc. 2015, 137 (23), 72920.

(69) Chen, Z.-M.; Hilton, M. J.; Sigman, M. S. Palladium-Catalyzed Enantioselective Redox-Relay Heck Arylation of 1,1-Disubstituted Homoolylic Alcohols. J. Am. Chem. Soc. 2016, 138 (36), 11461.

(70) Patel, H. H.; Sigman, M. S. Enantioselective Palladium-Catalyzed Alkenylation of Trisubstituted Alkenols To Form Allylic Quaternary Centers. J. Am. Chem. Soc. 2016, 138 (43), 14226.

(71) Chen, Z.-M.; Nervig, C. S.; DeLuca, R. J.; Sigman, M. S. Palladium-Catalyzed Enantioselective Redox-Relay Heck Alkenylation of Alkenols To Access Propargylic Stereocentres. Angew. Chem., Int. Ed. 2017, 56 (23), 6651.

(72) Singh, S.; Bruilaerts, J.; Vasseur, A.; Marek, I. A unique Pt-catalyzed Heck arylaion as a remote trigger for cyclopropane selective ring-opening. Nat. Commun. 2017, 8, 14200.

(73) Larionov, E.; Lin, L.; Guéneau, S.; Mazet, C. Scope and Mechanism in Palladium-Catalyzed Isomerizations of Highly Substituted Allylic, Homoolylic, and Alkenyl Alcohols. J. Am. Chem. Soc. 2014, 136 (48), 16882.

(74) Lin, L.; Romano, C.; Mazet, C. Palladium-Catalyzed Long-Range Deconjugative Isomerization of Highly Substituted αβ-Unsaturated Carbonyl Compounds. J. Am. Chem. Soc. 2016, 138 (32), 10344.

(75) Larock, R. C.; Lu, Y. D.; Bain, A. C.; Russell, C. E. Palladium-catalyzed coupling of aryl iodides, nonconjugated dienes and carbon nucleophiles by palladium migration. J. Org. Chem. 1991, 56 (15), 4589.

(76) Larock, R. C.; Wang, Y.; Lu, Y.; Russell, C. A. Synthesis of Aryl-Substituted Allylic Amines via Palladium-Catalyzed Coupling of Aryl Iodides, Nonconjugated Dienes, and Amines. J. Org. Chem. 1994, 59 (26), 8107.

(77) Albéniz, A. C.; Espinet, P.; Lin, Y.-S. Palladium Migration along Linear Carbon Chains: The Detection of ν1ν2-Enyl Intermediates and the Study of Their Rearrangement. Organometallics 1997, 16 (19), 4138.

(78) Wang, Y.; Dong, X.; Larock, R. C. Synthesis of Naturally Occurring Pyridine Alkaloids via Palladium-Catalyzed Coupling/Migration Chemistry. J. Org. Chem. 2003, 68 (8), 3090.

(79) Goj, L. A.; Widenhoefer, R. A. Mechanistic Studies of the Cycloisomerization of Dimethyl Diallylmalonate Catalyzed by a Cationic Palladium Phenanthroline Complex. J. Am. Chem. Soc. 2001, 123 (45), 11133.