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Advances of well-defined alkali metal complexes in organic synthesis

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Alkali metal reagents in organic synthesis are ubiquitous within academia and industry due to the wide array of synthetic utility as bases, nucleophiles, and Lewis acids. While organolithium chemistry has been the most prominent, the development of analogous technologies with the later alkali metals has garnered recent interest. This review aims to develop an introductory primer on well-defined heavier alkali metal complexes and the corresponding contemporary utility in organic synthesis. Key developments will be highlighted from the past decade to provide direct comparison to early fundamental studies, aspiring to provide a template for future advances in the field of alkali metal-mediated methods and catalysis.

1. Introduction

Organic transformations mediated by alkali metals (AM) represent some of the foundational concepts taught in every introductory organic textbook as characteristic bases and nucleophiles. Although alkali metal-based intermediates have been studied since the early 1900s, the relationship between structure and activity has remained nebulous due to ionicity, aggregation, and instability.^{1–7} Of the alkali metals, organolithium compounds are at the forefront of organic chemistry as potent nucleophiles, Brønsted bases, and productive transmetalation and ligand-exchange agents. The wealth of available organolithium precursors has facilitated detailed mechanistic study of elusive intermediates, granting key insight into its metal-based reactivity.^{8–13} As a result, the usage of organolithiums as organometallic reagents has become commonplace in synthesis. Moving forward, growing environmental and sustainability concerns warrant investigation into alternative avenues.¹⁴

In this vein, the heavier alkali metals (AM = Na–Cs) represent an attractive substitute to lithium, maintaining analogous high nucleophilicity and Brønsted basicity coupled with increased earth abundance and biocompatibility. Contrastingly, the heavier alkali metals have larger ionic radii, enabling expanded coordination spheres (CN = 6–12) resulting in weaker and reactive bonds (Fig. 1).^{15–18} The use of heavier alkali metal salts and alkoxides has been prominent in organic chemistry as the increased basicity and proton affinity engendered

enhanced rate and selectivity. Simultaneously, these attributes make the structural and mechanistic analysis of well-defined organometallic complexes of the alkali metals challenging to study. Considering their versatility, this review seeks to highlight reported systems that provide thorough structural and mechanistic elucidation, extending to well-defined complexes and intermediates whose identity is supported by spectroscopic data, X-ray diffraction, or reactivity studies.

Given the effectiveness of the heavier alkali metal reagents in organic synthesis, initial work was focused on isolation and speciation of reactive intermediates of organosodium analogues, building upon existing studies with organolithium complexes. Aggregation and solvent effects in lithium systems have been well studied, but initial investigations of organosodium complexes were not as straightforward considering the changes in radii and coordination sphere. In this regard, the Collum group has extensively studied both lithium and sodium-based systems, specifically targeting alkali metal enolates to understand aggregation processes and enable entry to study these reactive intermediates.²⁰ By utilizing the method of continuous variation (MCV) and NMR spectroscopy, the structure of metal-based aggregates was determined and demonstrated that solvent plays a key role in aggregate formation. It should be mentioned that although ^{6,7}Li and ²³Na NMR spectroscopy can be used to characterize the formation of *in situ* metal complexes, the ²³Na NMR nuclei is quadrupolar and exhibits broad resonances that make it difficult to extract useful structural and mechanistic information.

Despite these challenges, Collum and coworkers utilized these spectroscopic methods to determine the structures of sodium amides, with NaN(SiMe₃)₂ represented as a titular example below (Fig. 2A). Solvation effects on sodium complexes result in a wider array of different solvent-based aggre-

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Periodic trends	Ionic radii ^a $M^0 / M^+ (\text{Å})$	Common Coordination numbers ^b	Basicity Proton affinity (kcal/mol)
Li	1.28 / 0.92	3	238.9 ^c
Na	1.66 / 1.39	6	256.2 ^c
K	2.03 / 1.64	8-10	263.0 ^c
Rb	2.20 / 1.83	10	282.7 ^d
Cs	2.44 / 1.88	12	269.2 ^c

Fig. 1 General trends of the alkali metals. ^aShannon radii for most common coordination number, ^bmost common coordination numbers, ^cexperimental data, ^dcomputational data calculated using PBE0/CRENBL level of theory with ECP basis set.^{15,16,19}

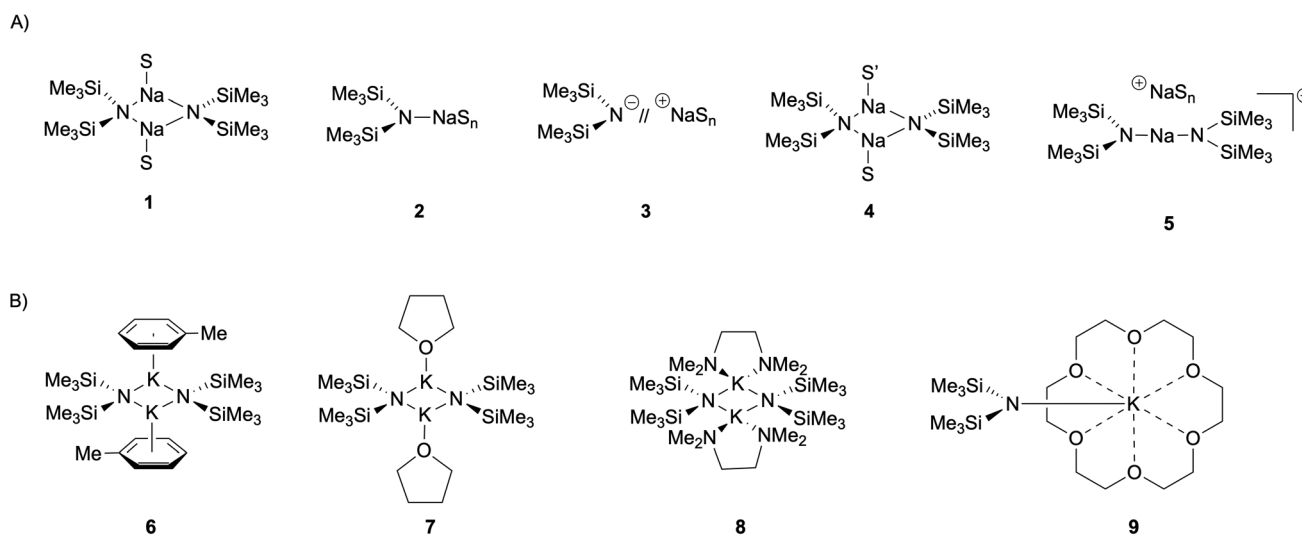


Fig. 2 (A) Common solvated structures of $\text{NaN}(\text{SiMe}_3)_2$, (B) various solution state structures of $\text{KN}(\text{SiMe}_3)_2$.

gates and ion pair complexes (1–5) than are normally observed with lithium counterparts, ultimately emphasizing the stark contrast of solution- and solid-state reactivity.^{21,22} Further studies continued to expand on the structure of alkali metal amides and reported the solution-state structures of $\text{KN}(\text{SiMe}_3)_2$, which exists predominantly as a dimer (6–8) in non-coordinating solvents, but transitions to a monomer (9) with strongly coordinating solvents and ligands (Fig. 2B).²³ Subsequent titration studies, isotopic labelling experiments, and thorough 2D and heteronuclear NMR spectroscopic analysis enabled rigorous solution-state investigation, providing a precise basis for the heavier alkali metal amide chemistry. As a result, $\text{NaN}(\text{SiMe}_3)_2$ and $\text{KN}(\text{SiMe}_3)_2$ based reactivity are the most studied systems and represent the framework for comparison amongst many alkali metal complexes.²⁴

In contrast, rubidium and cesium complexes have been much less explored, though literature reports indicate enhanced reactivity in line with general periodic trends.²⁵ Interestingly, organocesium reagents show increased solubility and decreased aggregation, in comparison to sodium and potassium, resulting in faster reaction rates and chemodivergent activity, coined as the ‘*Cesium effect*’ which has been reviewed previously.^{26,27} As such, recent studies demonstrate new developments for the heavier organoalkali compounds in rising areas of interest in organic chemistry such as C–C bond formation, C–H activation, and hydrofunctionalization. The main challenges for advancement of the later alkali metals are staunchly entranced in the decreased solubility of these reagents in hydrocarbon and ethereal solvents, the increased thermodynamic preference for forming aggregates, and the



incompatibility or deleterious reactivity with certain functionalities or substrates.

With this heightened reactivity, methods to tame these metals have included the use of Lewis donor-based solvents or additives as exogenous supports in solution, and the incorporation of *N*- or *O*-based ligand scaffolds to minimize aggregation and maximize stability at the metal center. In addition to enhanced stability, the incorporation of ligands provides spectroscopic handles by which solution-state reactivity can be deciphered and compared to isolated solid-state complexes. Progress towards the isolation of well-defined AM complexes sets the stage for systematic mechanistic study and demonstrates significant differences in reactivity that can be exploited for new organic transformations.

This review focuses on recent advances of the heavier alkali metals over the last ten years, and the resultant applications of these reagents within organic synthesis, methods, and catalysis. Distinct emphasis on synthetic availability and mechanistic investigations of key transformations is highlighted and contextualized. Within this scope, reports on the use of alkali metal dispersions, hydrides, salts, alkoxides, and carbonates have been omitted, as well as discussion of francium complexes due to their limited syntheses.^{28,29}

2. Donor assisted deprotonation

2.1. Deprotonation

The deprotonation chemistry of AM reagents is expansive and commonly mediated using prototypical AM hydroxides, with other examples including alkoxides, aryloxides and carbonates. Selectivity in deprotonation for these systems is seldom controlled due to a lack of synthetic tunability at the alkali metal center. The addition of exogenous donors, in the form of mono- or multidentate ligands, represents a new avenue to promote site-selective deprotonation. Few well-defined analogues that enable selectivity exist, and current studies have focused on developing structural elucidation of metalated intermediates to understand how to modulate basicity and selectivity in deprotonation. Solvent choice plays a key role in deprotonation reactivity as Lewis basic donor solvents readily engage AM complexes in metalation and aggregation, inducing changes in the Brønsted basicity of the metal center.

2.1.1. Sodium. In alkali metal chemistry, solvent incompatibility results in diminished reactivity, wherein the basicity of the AM center has also been shown to induce decomposition of the solvent or reagent. Within this context, Collum and co-workers studied the metalation/deprotonation of THF with sodium amides, comparing the reactivity to that of established lithium systems.³⁰ Particularly, the comparison of sodium diisopropylamide (NaDA) in solvents such as THF and *N,N*-dimethylethylamine (DMEA) to that of lithium diisopropylamide (LDA) in THF was explored by comparing the relative rates of metalation reactions between NaDA and LDA (k_{rel}). The use of NaDA has been limited due to the scarcity of solvents that allow for high solubility; however, NaDA–DMEA demonstrates

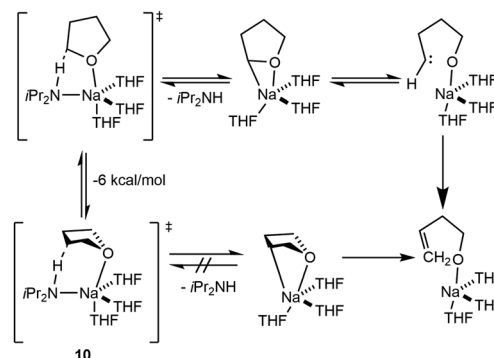
higher metalation rates than those of LDA–THF with comparable yields and selectivity.¹

Further investigation into the sodium mediated decomposition of THF showed that NaDA forms a tetra-solvated monomer in solution, which is then coordinatively primed to perform the α -deprotonation of THF (Scheme 1). The proposed mechanism for the decomposition of THF is shown below where intermediate **10** deprotonates at the α -position, followed by the loss of the diisopropylamide group and formation of the oxacarbenoid-precursor to the carbene–Na structure. Alternatively, an E2-like β -metalation pathway may occur; however, isotopic studies showed that scrambling prevents clear differentiation between the two mechanisms. Comparisons with THF and THF-*d*₄ results in a $k_{\text{H}}/k_{\text{D}} \sim 6$, suggesting that the C–H cleavage is the rate limiting step. The decomposition of DME by NaDA affords sodium methoxide and methyl vinyl ether proceeding through a di-solvated-monomer-based transition state.

The sodiation of a series of unactivated arenes was achieved through the application of highly basic 2,2,6,6-tetramethylpiperidide (NaTMP) and a polyamine donor, either *N,N,N',N'*-tetramethylethylenediamine (TMEDA) or *N,N,N',N',N''*-penta-methyldiethylenetriamine (PMDETA), by the Hevia group.^{31,32} Although the arylation step resulted in low isolated yields, subsequent arene borylations done in a one-pot procedure resulted in 80–83% yields (Fig. 3). Mechanistic studies with anisole as a representative substrate for *ortho*-metalation were explored. In the presence of either NaTMP/PMDETA (36% yield) or NaTMP/TMEDA (35% yield), a bimetallic intermediate forms with two sodium centers bridged by one TMP and one anisole, flanked by one donor molecule each. This represents a rare example of a well-defined mixed sodium-aryl/sodium-amide complex, elucidated by NMR spectroscopy and X-ray crystallographic studies (Fig. 3c).

2.2. Metal-directed deprotonation

Falling under the umbrella of general deprotonation, examples of metal-directed deprotonation involve the formation of discrete metal–substrate bonds which are integral for subsequent bond formation or rearrangement.



Scheme 1 Mechanism of THF decomposition by NaDA.



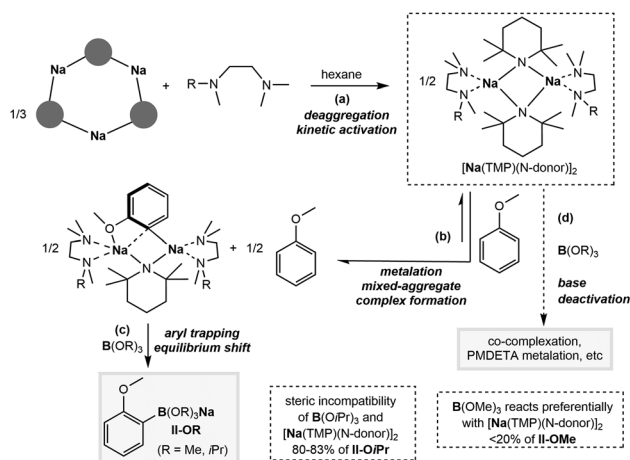
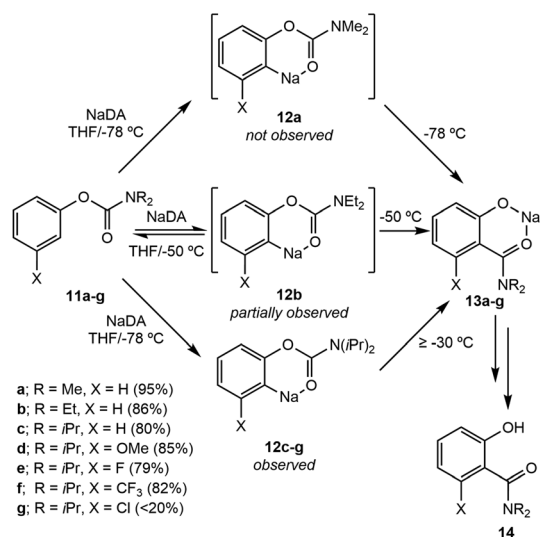


Fig. 3 Proposed pathway for the borylation of anisole by NaTMP.

2.2.1. Sodium. The utility of NaDA extends to the conversion of aryl carbamates (**11**) to *ortho*-acylated phenols in THF.³³ The *ortho*-metalation of the phenols is proposed to form *via* an arylsodium intermediate (**12a–g**) which undergoes a Snieckus-Fries rearrangement (**13a–g**) to form the *ortho*-acylated product (**14**, Scheme 2). DFT calculations suggest that the metalation and rearrangement steps both proceed through THF-solvated, monomeric NaDA pathways. The rearrangement reactions occur in good yields; however, aryl carbamates with halogen substituents may undergo either rearrangement or competitive halide eliminations, depending on the substituents, forming benzyne preferentially.

Based on prior work with lithium Evans enolates, the Collum group generated a series of enolates using NaDA or NaN(SiMe₃)₂ in the presence of TMEDA, *trans*-*N,N,N',N'*-tetramethylcyclohexanediamine (*R,R*)-TMEDA, or (*S,S*)-TMEDA



Scheme 2 The synthesis of *ortho*-acylated phenols (**14a–g**) from aryl carbamates (**11a–g**) using NaDA in THF through arylsodium intermediates (**12a–g**).

(Scheme 3).³⁴ The sodiated Evans enolates formed mixed dimers, each of which was examined for aldol additions, stereo-selective quaternizations, and azaaldol additions. Although similar results were produced using either NaDA or NaN(SiMe₃)₂, it was found that NaN(SiMe₃)₂ was the preferred reagent due to its commercial availability and versatility. The TMEDA-solvated sodiated Evans enolates (**16**) demonstrate improved structural control compared to THF-solvated counterparts and maintain comparable selectivity (Scheme 3). This diverges from lithium enolates, with which THF has been the solvent of choice, and the potential utility of sodium enolates for transformations with imines and aldehydes further widens this divide.

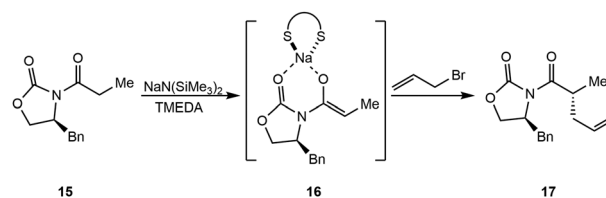
Application of NaTMP to halothiophenes (**18**, X = Cl or Br) by Mori and coworkers results in metalation, forming sodium thiophenes (**19**) as precursors for polymerization.³⁵ Quenching the metalated thiophene product with iodine afforded 2-chloro-3-hexyl-5-iodothiophene (85%) and 2-bromo-3-hexyl-5-iodothiophene (17%, **20**, Scheme 4), with the latter resulting in lower yields due to deleterious reactivity by NaTMP at the C–Br bond.

2.2.2. Potassium, rubidium, cesium. Examples of deprotonation with potassium, rubidium, and cesium alkoxides, aryl-oxides, and carbonates are known, similar to the lithium and sodium analogues. While there are well-defined examples where deprotonation is utilized with these metals, the transformations have been classified under an alternative reaction type (see 3.3.1, 6.1.2 and 6.2.2) within this review.

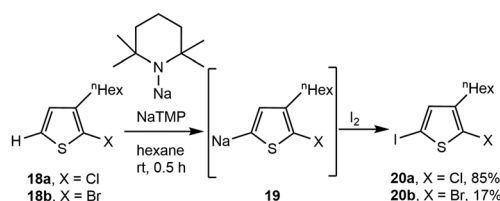
3. Amination, enolization, and epoxide ring-opening

3.1. Amination and Mannich reactions

The formation of new C–N bonds enables incorporation of nitrogen into organic molecules, which is indispensable in



Scheme 3 Synthesis of sodiated Evans enolates using NaN(SiMe₃)₂.



Scheme 4 Deprotonative metalation of halothiophenes (**18**) with NaTMP to yield sodiated halothiophenes (**19**).



pharmaceutical chemistry and total synthesis of natural products. The improved biocompatibility of alkali metals compared to transition metals increases the appeal of pursuing AM-mediated routes for installing nitrogen centers in organic moieties.

3.1.1. Sodium. Solvent-dependent control over reactivity, commercial availability, and heightened tolerance to a variety of conditions render $\text{NaN}(\text{SiMe}_3)_2$ the preferred sodium amide for amination reactions. The Collum group explored $\text{NaN}(\text{SiMe}_3)_2$ as a strong base toward carbon-centered electrophiles for a series of C–N bond-forming reactions.³⁶ The reaction of $\text{NaN}(\text{SiMe}_3)_2$ with electrophiles (ester-substituted arenes, pyridines, and epoxides) in various solvents resulted in pristine products of generally high yields (55%–96%). Discrepancies in reactivity and yield were explored with mechanistic studies in which it was found that the solvent choice or reaction conditions heavily influenced the reaction outcomes, delineated in Table 1. Although THF and DMEA preferentially form monomeric $\text{NaN}(\text{SiMe}_3)_2$ (**1**), reactivity was dominated by dimeric $\text{NaN}(\text{SiMe}_3)_2$ (**2**) in the presence of substrates. Solvent choice dictated chemoselectivity, whereby in THF the release of NaOSiMe_3 results in an imino ether intermediate, affording entry 5 or entry 6. However, in DMEA, MeOSiMe_3 is released instead, forming the carboxamide product preferentially (entry 2), demonstrating the importance of solvent selection for generating the necessary aggregate intermediate.

Diversifying the landscape of sodium reagents through rigorous mechanistic and coordination studies, the Collum group extended similar investigations to include sodium alkylsilazides such as sodium isopropyl(trimethylsilyl)amide (NaPTA) and sodium *tert*-butyl(trimethylsilyl)amide (NaBTA). Reactivity such as *N*-alkylations, epoxide openings, and aminations were achieved with similar efficacy to $\text{NaN}(\text{SiMe}_3)_2$ while maintaining the lability of ethereal solvents demonstrated by NaDA. Despite these enhanced modifications, sodium alkylsilazides are less synthetically accessible than the parent $\text{NaN}(\text{SiMe}_3)_2$.³⁷

3.1.2. Potassium. Proceeding down the group, potassium exhibits a larger ionic radius and increased reactivity compared to sodium, making it suitable for more challenging transformations. Kobayashi and colleagues present a chiral potassium salt catalyst system (K-box) for catalytic, asymmetric Mannich reactions.³⁸ K-box catalysts (**21**) are prepared from $\text{KN}(\text{SiMe}_3)_2$ and chiral bis(oxazoline) ligands, and have demonstrated efficiency in targeting imines (**22**) and weakly acidic amides (**23**) to produce β -amino acid derivatives (**24**) with high diastereo- and enantioselectivity (Scheme 5A). $\text{KN}(\text{SiMe}_3)_2$ interacts with the chiral K-box ligand to form a potassium enolate-K-box complex (**25**), which facilitates imine addition (Scheme 5B). The complex stabilizes the potassium enolate and promotes highly efficient deprotonation of the amide, enabling precise asymmetric control during imine addition. High yields and enantioselectivities (up to 94% ee) were observed, with substrate-specific modifications further enhancing selectivity. Applications included gram-scale reactions and β -lactam synthesis, demonstrating the system's scalability and practical utility with a heavier alkali metal.

3.2. Enolizations

Alkali metal bases have been routinely applied towards enolizations under basic conditions, but the structural conformation of the alkali metal intermediates has been minimally investigated. Comparatively, organic systems using lithium amides (LDA, $\text{LiN}(\text{SiMe}_3)_2$) have demonstrated reactive competency with limited efficiency and selectivity.^{39–42}

3.2.1. Sodium. In comparison, the shortcomings of lithium amides are addressed through the use of $\text{NaN}(\text{SiMe}_3)_2$ for ketone enolization, whereby aggregation and conformation are directly influenced by solvent and substrate, with *E*:*Z* selectivity and kinetic rates varying heavily based on the system. The Collum group has identified several plausible mechanistic pathways, wherein $\text{NaN}(\text{SiMe}_3)_2$ can exist as a monomer (**26**), dimer (**27**), ion pair (**28**), or free ion (**29**) depending on the solvent, which then reacts with the ketone substrate in any variation of these forms to generate the enolate products **30-E** and **30-Z** (Scheme 6).⁴³ The effect on *E*:*Z* selectivity was shown to range from 20:1 (Et_3N /toluene) to 1:90 (THF). Et_3N and methyl-*t*-butyl ether (MTBE) show the highest selectivity towards the *E*-product, with an *E*:*Z* ratio greater than 10:1. Toluene, TMEDA, and PMDETA have an *E*:*Z* ratio less than 10:1 but still exhibit preference for the *E*-enolate. Diglyme is the only example studied with a 1:1 ratio. Both THF and DME (1:20) selectively form the *Z*-product. Small changes in substrate were shown to impact the mechanism drastically, as changing the substrate from 2-methyl-3-pentanone to 2-methylcyclohexanone causes the mechanism to change from a monomer-promoted pathway to one involving a triple-ion pair system. In comparison to lithium reagents, superior selectivity and faster reactivity were reported, emphasizing $\text{NaN}(\text{SiMe}_3)_2$ as a competent, accessible reagent choice.

3.3. Epoxide ring-opening

Epoxides represent useful chemical feedstocks for polymerizations, alcohol formation, and building larger organic molecules. Epoxide ring-opening can be performed under both basic and acidic conditions, commonly employing alkali salts and alkoxides for base-promoted openings.⁴⁴ In addition, there are few routes utilizing well-defined AM precursors for epoxide ring-opening reactions, though limited examples with sodium complexes have been previously reported (see section 3.1.1).^{36,44,45}

3.3.1. Sodium and potassium. The Shirakawa group employed well-defined glycol-stabilized sodium and potassium salts for the catalytic conversion of epoxides and atmospheric CO_2 into cyclic carbonates (**31**) under mild conditions (Scheme 7A).⁴⁶ The highest performing complexes all consisted of iodide salts and glycols with either three (**33**, 67%) or four (**32**, 68%; **34**, 70%) ethylene linkers (Scheme 7B). Mechanistic investigation using **34** suggests that tetraethylene glycol (tEG) activates epoxides *via* hydrogen bonding, while the iodide anion from KI facilitates nucleophilic attack. This process leads to the formation of cyclic carbonates through a



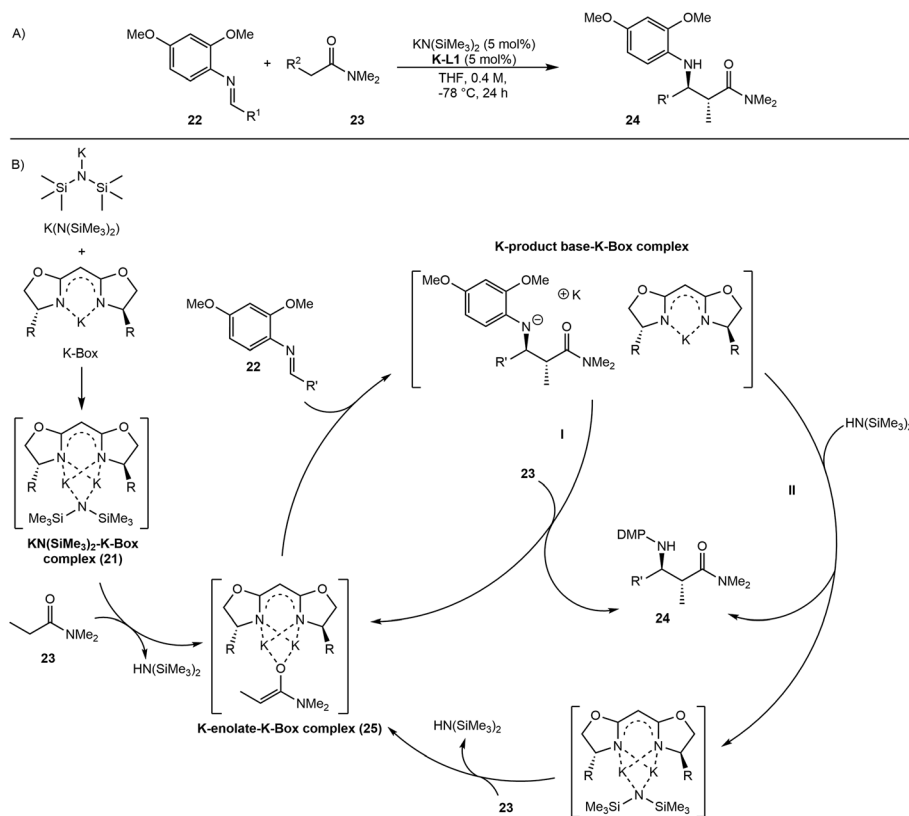
Table 1 Reaction of $\text{NaN}(\text{SiMe}_3)_2$ with electrophiles in various solvents

Reaction of $\text{NaN}(\text{SiMe}_3)_2$ with electrophiles in various solvents									
Entry	Substrate	Conditions	Product	Yield (%)	Entry	Substrate	Conditions	Product	Yield (%)
1		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 5 h Toluene		72	10		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 50 °C, 1 h DMEA		78
2		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 0.3 h DMEA		95	11		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 50 °C, 2 h 25 °C, 24 h THF		92
3		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 0.3 h Toluene		90	12		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 0.05 h DMEA		95
4		1.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 50 °C, 0.1 h THF		86	13		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 2 h Toluene		85
5		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 50 °C, 0.3 h THF		92	14		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 1 h Toluene		77
6		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 70 °C, 1 h Toluene		85	15		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 1 h Toluene		83
7		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 70 °C, 1 h THF		96	16		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 110 °C, 2 h Toluene		55
8		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 70 °C, 1 h Toluene		76	17		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 110 °C, 3 h Toluene		76
9		3.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 70 °C, 1 h THF		95	18		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 60 °C, 24 h Toluene		86 (>50:1)
					19		2.0 equiv. $\text{NaN}(\text{SiMe}_3)_2$ 25 °C, 24 h THF		76

three-step catalytic cycle: epoxide fixation (35) and activation (36), CO_2 fixation (37), and intramolecular ring closure (31, Scheme 7C). The reaction tolerates a broad substrate scope,

including simple and functionalized epoxides, in addition to enantiopure substrates without loss of stereochemical integrity. Large-scale syntheses and catalyst recycling experiments





Scheme 5 A) Optimized reaction conditions for Mannich reaction using K-box catalyst system (21), (B) proposed catalytic cycle.

achieved over 10 catalytic cycles with no loss of efficiency of the glycol-potassium salt pair. Subsequent computational analysis (DFT) from Butera and Detz exploring the mechanism reveals that the rate-determining step involves the nucleophilic attack of iodide on the epoxide, with a calculated energy barrier significantly reduced compared to the uncatalyzed process.⁴⁷ Natural bond order (NBO) and Hirshfeld charge analyses confirm that tEG's coordination with potassium weakens the K-I bond, freeing iodide ions for nucleophilic activity.

4. Isomerization and olefination

4.1. Isomerization

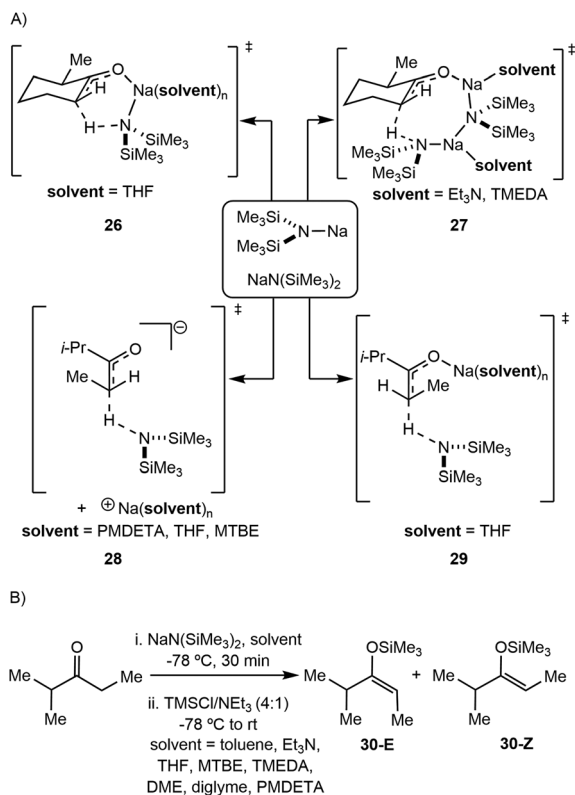
Isomerization is primarily mediated by transition metals that are able to access metal hydride or allyl intermediates.⁴⁸ However, there have been reports of stoichiometric alkali metal bases, such as hydrides, alkoxides, and amides, performing isomerization reactions with both ill- and well-defined systems.^{49–51} Chemical tunability with well-defined systems is part of ongoing efforts to improve AM-isomerization methods, taking advantage of their basicity and reactivity in stoichiometric and catalytic avenues.

4.1.1. Sodium. Sodium diisopropylamide (NaDA) serves as an effective Brønsted base for the isomerization of C(sp²)-C

(sp²) bonds, aided by cation-π interactions. The Collum group has explored the isomerization of alkenes and the metalation of dienes using NaDA in THF (Scheme 8).⁵² The reaction of NaDA with 1,4-dienes afforded the dienyl sodium product *via* metalation (38) while isomerization with alkene substrates demonstrated high selectivity for the *Z*-isomer, with the exception of 1-pentene (39). Treatment of allyloxy ethers with NaDA afforded enol ethers with good selectivity (>50 : 1 *Z* : *E*, 40a–d), while substituted allyl ethers underwent 1,4-elimination rather than isomerization (41a–b). In the case of allyloxytrimethylsilane, the addition of catalytic (6.5 mol%) NaDA resulted in fast conversion to the isomerized product 40b with 80% isolated yield. In most cases, isomerization utilizing dimeric [NaDA(THF)₂]₂ requires lower temperatures.

Super-basic NaTMP has been implemented for alkene isomerization, with the Hevia group exploring the use of catalytic NaTMP in conjunction with the tridentate PMDETA ligand to increase basicity.^{53,54} At 10 mol% loading for both NaTMP and PMDETA, the isomerization of a larger scope of alkenes was observed at room temperature (Fig. 4). The isomerization of allylsilanes yielded predominately *E*-isomer products in good yields at much lower temperatures and durations than comparable reactions using B(C₆F₅)₃. Activated allylbenzenes and internal silyl-substituted olefins showed efficient isomerization to the internal alkene product (43a–43h). Longer and more



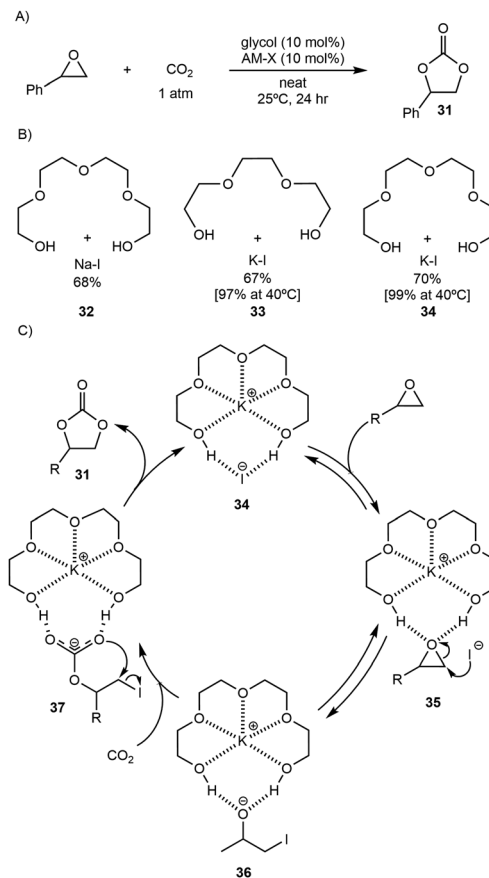


Scheme 6 A) Plausible mechanistic pathways for the enolization of ketones with $\text{NaN}(\text{SiMe}_3)_2$ (B) $E:Z$ selectivity by solvent system.

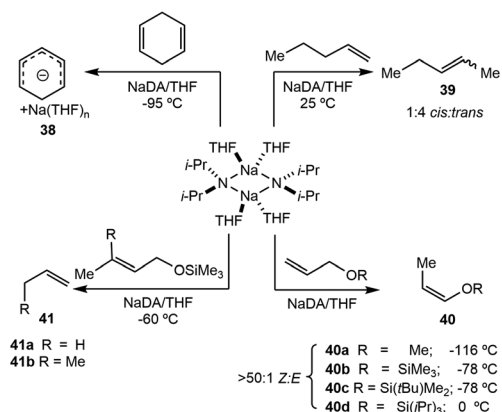
substituted alkyl chains resulted in slightly decreased yields (43g–h), inclusion of both *N*- and *O*- in the allyl unit resulted in good conversion to the target enamines and vinyl ethers (43j–43n). The isomerization of halide-, carbonyl-, and nitrile-containing compounds was unsuccessful, though isomerization of some substituted alkynes and dienes were productive (43o–p). Longer chain α -olefins such as 1-octene lacked selectivity and provided a mixture of isomers (43q). When THF was used instead of hexane, rapid decomposition of the base prevented the isomerization reaction from occurring, in line with similar reports by Collum (see section 2.2.1). In C_6D_6 , the incorporation of deuterium into cycloalkenes was observed in moderate to excellent yields rather than the isomerization exhibited in C_6D_{12} , demonstrating the high basicity of the mixed $\text{NaTMP}/\text{PMDETA}$ system.

4.2. Olefination

Various types of olefinations are mediated by specialized reagents (e.g. Wittig reagent), often in conjunction with an alkali metal base, such as organolithium reagents for Julia olefinations and Shapiro reactions, or alkali metal hydrides and alkoxides for Horner–Wadsworth–Emmons olefinations and Peterson olefinations.^{55–57} In these types of olefinations, the identity of the alkali metal plays a role in the stability of the intermediate and the stereochemistry of the product, ultimately making it challenging to develop a universal system



Scheme 7 A) Conversion of epoxides into cyclic carbonates using atmospheric CO_2 and a catalytic potassium iodide-tetraethylene glycol complex, (B) highest performing glycol and alkali salt combinations, (C) proposed mechanism of cyclic carbonate formation.



Scheme 8 Summary of NaDA reactivity with alkenes.

with common bases.⁵⁸ Despite these challenges, there are contemporary reports of alkali metal complexes used for olefinations.^{59,60} Improved understanding of alkali metal coordination chemistry has led to purposeful design and synthetic control of well-defined systems for general olefinations, broadening the scope of reactivity with common substrates.



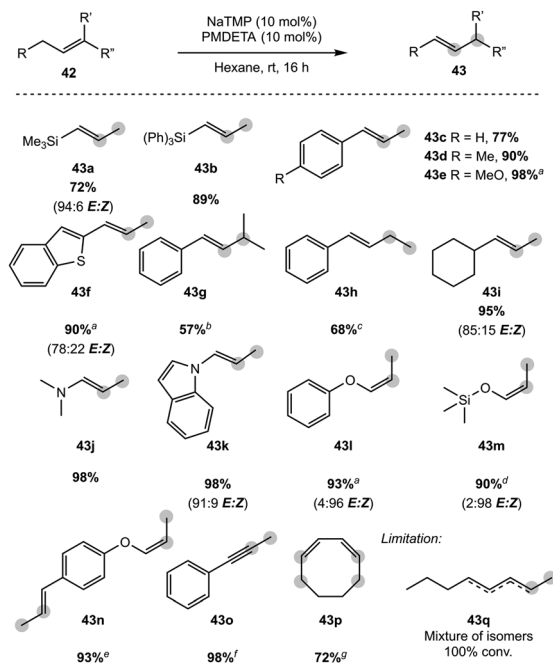


Fig. 4 Substrate scope for the isomerization of alkenes using the catalytic NaTMP/PMDETA system. Grey circles indicate the initial position of the double bond. ^a Isolated yields, ^b 50 °C, 4 h, ^c TMEDA (0.1 mL), 120 h, ^d TMEDA (10 mol%), 0 °C, 15 min, ^e TMEDA (0.1 mL), ^f 80 °C, ^g 96 h.

4.2.1. Sodium. In their rigorous study of NaDA, Collum and coworkers studied the effect of solvent-ligands on sodium amides and competency for olefination.³⁰ NaDA-DMEA reactions exhibited substantially faster k_{rel} values and an increased variety of transformations compared to that of LDA-THF. Dehalogenations (Table 2, entries 1–3) occur at a rate most similar to LDA-THF, with the exception of the equatorial conformation of entry 2, which is highly effective compared to the absence of equatorial elimination with LDA-THF. Entry 6 demonstrates a remarkable deviation from LDA-based reactivity, forming cyclooctenols with a 10:1 *cis:trans* selectivity without requiring additional heating or formation of side products. Facile *ortho*-metalations (entries 7–10) were observed without benzyne formation at low (less than –30 °C) temperatures. Metalation with subsequent silylation (entry 11) and ether rearrangement (entry 12) both proceed as expected and significantly faster than the analogous LDA reactions.

More recently, the utility of NaDA-DMEA for olefinations was demonstrated by the Collum group through the addition of catalytic PMDETA (10 mol%) to NaDA (Table 3).⁶¹ PMDETA was compared to the bidentate TMEDA using both kinetic and computational studies which showed the inherent hemilability of the different ligand scaffolds was crucial for enabling reactivity. Olefin products in most cases were achieved in high yields (82–95%) and experienced rate acceleration upon the addition of PMDETA; however, for entries 16–18, PMDETA did not result in acceleration, with the authors predicting that the tridentate coordination prevents substrate addition to the sodium center.

Metalated benzylsilanes have demonstrated the effective *in situ* olefination of ketones, aldehydes and amides by well-defined lithium (44) and sodium (45) silylbenzyl complexes, affording trisubstituted alkenes in excellent yields.⁶² High conversion (>95%) of both bulky and simple carbonyls under mild conditions in C₆D₆ presents these alkali metal complexes as competitive alternatives in comparison to traditional olefination methods. Moderate *E:Z* selectivity was observed for entries 2, 7, 8, and 9 and suggests the potential use of chiral ligands for manipulating selectivity (Table 4).

Despite the similarity between 44 and 45 for olefination, (Me₆Tren)Li(CH₂SiMe₃) and (Me₆Tren)Na(CH₂SiMe₃) (compounds 46 and 47, respectively) exhibit divergent reactivity for the olefination of ketones as highlighted by the Lu group.⁶³ Treatment of ketones with 46 results in nucleophilic addition at the O-atom upon the addition of the ketone substrate; however, reaction with 47 proceeds through three possible pathways (Scheme 9). Methylenation of the carbonyl was the predominant transformation across 10 substrates, including benzophenone (>95%), electron deficient fluorinated ketones (>90%), and ketones that do not readily undergo enolization: dicyclohexyl ketone (~70%), phenyl cyclohexyl ketone (>95%), phenyl *tert*-butyl ketone (>95%). Nucleophilic addition was only the major reaction path (>95%) when benzaldehyde was the substrate, but after three days, full conversion to the methylenated product was observed. Deprotonation to form the enolate was only the primary path when acetophenone was used (>95%). It was determined that catalytic amounts of the Me₆Tren ligand (5 mol%) could be added *in situ* to stoichiometric equivalents of [Na(CH₂SiMe₃)]_∞ and substrate, resulting in >95% conversion to the methylenated product.

4.2.2. Potassium. Although potassium is commonly used as a base additive in olefinations, specifically as an alkoxide, Essman and Jacobsen report a potassium-isothioureabornate ion pair complex as an efficient chiral Lewis acid catalyst for enantioselective Wittig olefinations (Scheme 10).⁶⁴ The system enables the asymmetric synthesis of axially chiral alkenes *via* olefination of 4-substituted cyclohexanones with non-stabilized phosphorus ylides, achieving high enantioselectivity (up to 92% ee). The reaction proceeds through a Lewis acidic mechanism involving a stepwise cycloaddition, where the potassium center coordinates to the macrocyclic amide-bornate framework (48), stabilizing the intermediate oxaphosphetane in turn facilitating enantioselective bond formation. Steric and electronic tuning of the arylpyrrolidine moiety on the potassium complex significantly impacts selectivity, with the 3-phenanthryl derivative affording the highest enantioselectivity. Upon comparison of the analogous Li, Na, and K complexes, the authors proposed that potassium's unique coordination to the framework, emulating an amide metal bornate instead of an amide metal isothiurea, makes 48 much more effective than its lighter congeners. Electron-deficient aromatic substrates enhanced reaction rates and selectivity, while sterically hindered ylides reduced enantioselectivity.



Table 2 Reaction of substrates using NaDA

Entry	Substrate	Conditions	E ⁺	Product	k _{rel}	Yield (%)
1	<i>n</i> C ₈ H ₁₇ Br	1.2 equiv. NaDA 0 °C	—	<i>n</i> C ₆ H ₁₃ CH=CH ₂	5	87
2		1.2 equiv. NaDA 0 °C	—		>500	80
3		1.2 equiv. NaDA 0 °C	—		5	87
4		1.1 equiv. NaDA -78 °C	TMSCl		N/A	61
				<i>E</i> : <i>Z</i> = 1:4		
5		1.2 equiv. NaDA -78 °C	CH ₃ I		>300	78
6		1.3 equiv. NaDA rt	H ₂ O		>500	80
				<i>cis</i> : <i>trans</i> = 10:1		
7		1.1 equiv. NaDA -78 °C	CO ₂		N/A	60
8		1.2 equiv. NaDA -78 °C	H ₂ O		>200	92
9		1.2 equiv. NaDA -78 °C	CH ₃ OD		>500	94
10		1.2 equiv. NaDA -78 °C	CH ₃ OD		1000	92
				2- <i>d</i> :6- <i>d</i> = 6:1		
11		1.1 equiv. NaDA 0 °C	TIPSCl		>100	82
12		1.1 equiv. NaDA 0 °C	—		>1000	83
	R = Me, Ph, SiMe ₃			<i>Z</i> : <i>E</i> > 50:1		R = Ph



Table 3 Reactivity of NaDA in DMEA and PMDETA using different substrates

Entry	Substrate	Product	substrate $\xrightarrow[\text{10 mol\% PMDETA}]{\text{neat Me}_2\text{NEt (DMEA), } i\text{Pr}_2\text{NNa (1.2 equiv)}} \text{product}$		
			T ($^{\circ}\text{C}$)	Yield (%)	k_{PMDETA}
1			-30	82	300
2			0	86	220
3			25	—	>200
4			0	90	340
5			-30	95	150
6			25	86	230
7			-80	95	350
8			0	93	330
9			25	84	35
10			25	90	40
11			25	89	70
12			25	91	20
13			0	—	20
14			25	85	85
15			-78	—	45



Table 3 (Contd.)

		$i\text{Pr}_2\text{NNa}$ (1.2 equiv) neat Me_2NEt (DMEA) 10 mol% PMDETA			
substrate		→		product	
Entry	Substrate	Product	T (°C)	Yield (%)	k_{PMDETA}
16			-78	—	35
17			-78	92	<1
18			-78	—	15

5. Alkylation, benzylic functionalization, and allylic activation

5.1. Alkylation

Alkali metal reagents, specifically organolithium reagents, have been widely utilized as alkylating reagents since their discovery due to their nucleophilic nature.² However, the availability and commonplace usage of organolithium reagents have overshadowed the study of its heavier congeners (see section 3.2.1 for further discussion).

5.1.1. Sodium. Sodium enolates, also known as Oppolzer enolates, are commonly used in organic syntheses as potent nucleophiles, which exhibit solvent-dependent structural conformation and reactivity. To generate the sodium enolate, the Collum group used solutions of either sodium isopropyl(trimethylsilyl)amide (NaPTA) or $\text{NaN}(\text{SiMe}_3)_2$ in a variety of solvents (Scheme 11A).⁶⁵ The conformation of the sodium amide was explored through *in situ* kinetic studies. It was found that enolate and allyl bromide were first-order suggesting a monomer-based alkylation (50), with second-order dependence on hexamethylphosphoramide (HMPA) further inferring the formation of a complex solvated ion pair (49-HMPA, Scheme 11B). In the absence of HMPA, alkylations were performed in PMDETA and TMEDA without success, while THF resulted in a 100-fold decrease in rate. In the case of MeI, alkylation (51) occurred sufficiently fast to allow for the omission of HMPA (49-THF, Scheme 11C). Further alkylation rate studies suggest the reaction involves $[\text{Na}(\text{THF})_6]^+$ as the cationic half of a hexa-solvated ion pair intermediate *en route* towards alkylation.

The investigation of NaDA continued with the synthesis of pseudoephedrine-derived (52) Myers enolates (54) by forming the disodium salt (53, Scheme 12).⁶⁶ Poor yield and incom-

plete reactions were found to be associated with deleterious aggregation of NaDA and *O*-alkylations in the second step.





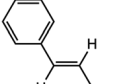




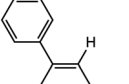






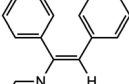


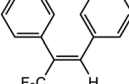


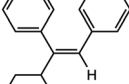
5.2. Benzylic functionalization

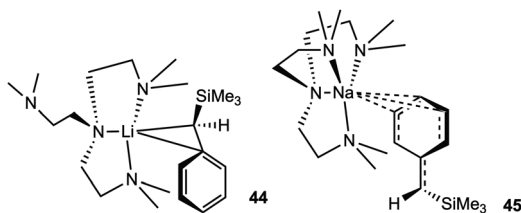
5.2.1. Sodium. The influence of solvent-ligands additionally dictates the aggregation of alkali metal alkyl complexes. The benzylic activation and arylation of toluenes with Weinreb amides by $\text{Na}(\text{CH}_2\text{SiMe}_3)_\infty$ has been demonstrated by the Hevia group using PMDETA to deaggregate polymeric $\text{Na}(\text{CH}_2\text{SiMe}_3)_\infty$ for $\text{C}(\text{sp}^3)\text{-H}$ activation.^{67,68} The addition of stoichiometric toluene, $\text{Na}(\text{CH}_2\text{SiMe}_3)_\infty$, and PMDETA results in the formation of discrete isolated complex (PMDETA)Na(CH₂Ph) (55). Subsequent addition of a Weinreb amide to 55 affords the arylated toluene product 56a-y (Fig. 5). This approach gave reasonable yields for simpler alkyl and aryl substituted toluenes (75–95% yield), and methyl-substituted pyridines were similarly compatible (59–93%). Tolerance towards other functional groups varied widely (28–78%), for some bisaryl substrates, metalated intermediates were not basic enough to react with the Weinreb amides. The efficacy of (PMDETA)Na(CH₂Ph) insertions into C=O, C=N, and C=C bonds were additionally investigated and explored. Under a CO₂ atmosphere, phenylacetic acid (91%) was obtained after aqueous workup. Addition to *N*-benzylideneaniline affords amine *N*-(1,2-diphenylethyl)aniline (88%).

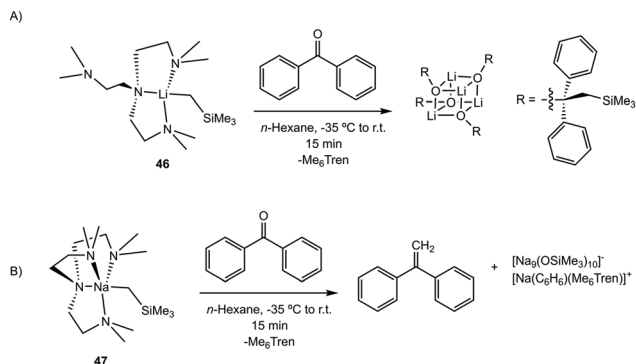
Invoking complementary nucleophilicity and basicity, the Hevia group employs benzylic metal intermediate (57) with stoichiometric $\text{Na}(\text{CH}_2\text{SiMe}_3)_\infty$ and PMDETA with 1.5 equivalents of toluene, inducing subsequent C–H addition to diarylethenes and ketones to form the desired coupled product (Fig. 6).⁶⁹ Initial studies showed that addition of one equivalent of 1,1-diphenylethylene to 57, followed by hydrolysis,



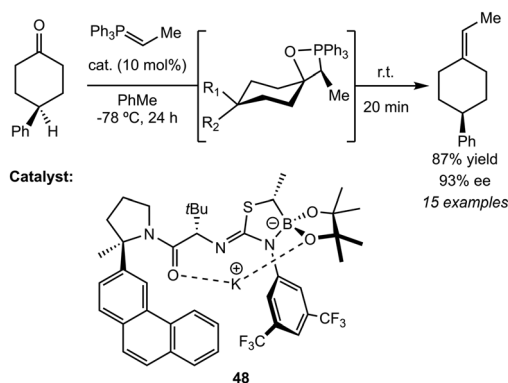
Table 4 Olefination of carbonyls by alkali–metal silylbenzyl complexes

Entry	Starting material	Conditions	Olefination product(s)	44		45	
				Conv. (%); <i>E</i> : <i>Z</i> ratio		Conv. (%); <i>E</i> : <i>Z</i> ratio	
1		60 °C 2 h		>95%		>95%	
2		rt 0.5 h	 	>95% <i>E</i> : <i>Z</i> = 2:3		>95% 1:1	
3		rt 0.5 h		>95%		>95%	
4		rt 0.5 h	 	>95% <i>E</i> : <i>Z</i> = 2:3		>95% 1:1	
5		rt 20 h		>95%		>95%	
6		rt 0.5 h		Intractable mixture		Intractable mixture	
7		rt 0.5 h	 	>95% <i>E</i> : <i>Z</i> = 3:2		>95% 1:1	
8		60 °C 2 h	 	>95% <i>E</i> : <i>Z</i> = 1:2		>95% 1:3	
9		rt 0.5 h	 	>95% <i>E</i> : <i>Z</i> = 2:3		>95% 2:3	

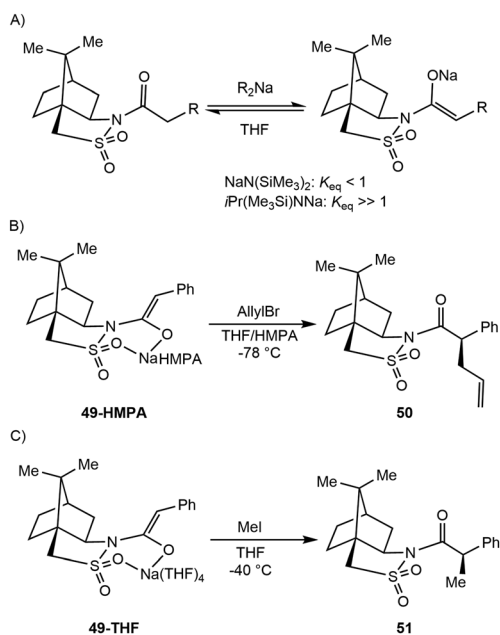




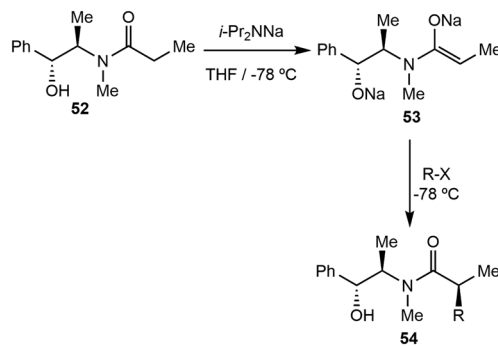
Scheme 9 A) Divergent reactivity of $(\text{Me}_6\text{Tren})\text{Li}(\text{CH}_2\text{SiMe}_3)$ (**46**) with benzophenone, and (B) divergent reactivity of $(\text{Me}_6\text{Tren})\text{Na}(\text{CH}_2\text{SiMe}_3)$ (**47**) with benzophenone.



Scheme 10 Enantioselective Wittig olefinations by chiral potassium-isothiourea-boronate catalyst (**48**).



Scheme 11 A) Comparison of NaDA and $\text{NaN}(\text{SiMe}_3)_2$ (B) alkylation in the presence of HMPA (C) alkylation using MeI in the absence of HMPA.



Scheme 12 Synthesis of Myers enolate from acylated pseudoephedrine (**52**) with NaDA.

afforded the coupled toluene-ethylene product **58**. The crystallographic data shows products from both the sodiation, PMDETA $[\text{Na}(\text{CH}(\text{NMe}_2)_2\text{Ph})]$, and C=C insertion steps prior to hydrolysis $[\{\text{ArCH}_2\text{CH}_2\text{C}(\text{Ph})_2\}\text{Na}(\text{PMDETA})]$. The same approach was additionally applied for the deprotonative coupling of toluenes and aromatic ketones.

5.3. Allylic activation

Allylic activation and functionalization are typically only seen with transition metals, with palladium complexes being the state-of-the-art, due to the ability to access the necessary π -allyl complexes needed for activation.^{70,71}

5.3.1. Sodium. The Schneider group explored the $\text{C}(\text{sp}^3)\text{-H}$ activation of alkenes with $\text{NaN}(\text{SiMe}_3)_2$ for C-C bond formation.⁷² The addition of catalytic $\text{NaN}(\text{SiMe}_3)_2$ (10 mol%) to a terminal alkene and protected imine lead to the desired amine with good yields, selectivity, and tolerance to a variety of functionalities on the imine (Fig. 7). Mechanistic studies indicate that under the reaction conditions, the sodium amide forms a η^3 Na-allyl intermediate *in situ*, akin to that observed with Pd π -allyl complexes, readily observed by ^{23}Na NMR spectroscopy (-5.3 ppm). The observed η^3 Na-allyl intermediate then forms the sodiated product (4.9 ppm) upon the addition of imine, which is considered catalytically active. It was found that both intermediates are essential for the catalysis. Both $\text{LiN}(\text{SiMe}_3)_2$ and $\text{KN}(\text{SiMe}_3)_2$ were also explored, resulting in lower conversions alongside isomerized starting material, indicating that the stability of the AM-allyl intermediate is necessary for the transformation.

6. Hydrogen isotope exchange (HIE) and transfer hydrogenation

6.1. HIE

Hydrogen Isotope Exchange (HIE) is the process of exchanging protons for a heavier isotope of hydrogen ($^2\text{H}/\text{D}$ or $^3\text{H}/\text{T}$), which is important in the pharmaceutical industry for drug design and development.^{73,74} Transition metal complexes have prevailed as typical HIE catalysts, although acid/base catalytic



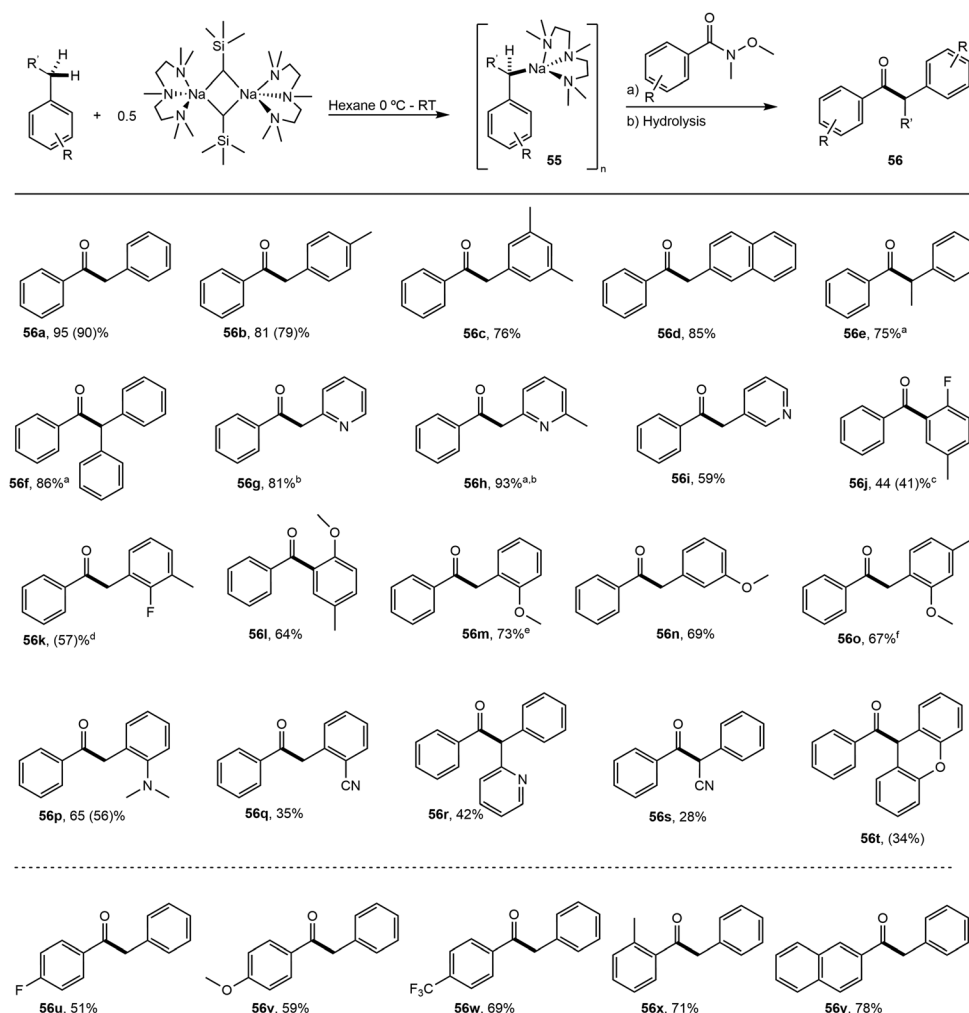


Fig. 5 Substrate scope for the benzylic arylation of toluenes by $\text{NaCH}_2\text{SiMe}_3$. ^a Solvent system: benzene/hexane, ^b mixture of tautomers, ^c $T = -78\text{ }^\circ\text{C}$, ^d mixture of products (single : double addition, 57% : 9%), ^e $T = 0\text{ }^\circ\text{C}$, ^f inseparable mixture of isomers (2 : 1).

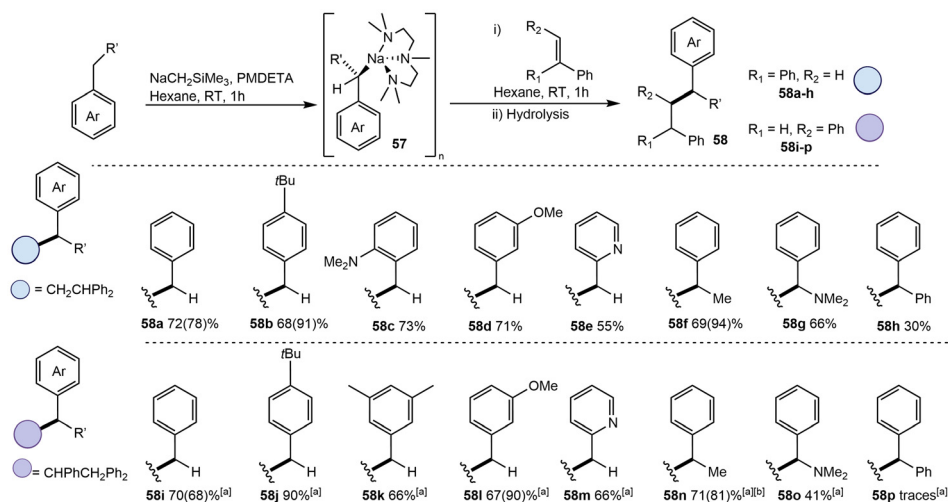


Fig. 6 Substrate scope for the coupling of substituted toluenes and olefins by $\text{NaCH}_2\text{SiMe}_3$. ^a Hexane/benzene 1 : 1 solvent system. ^b A 2 : 1 ratio of two diastereomers observed.



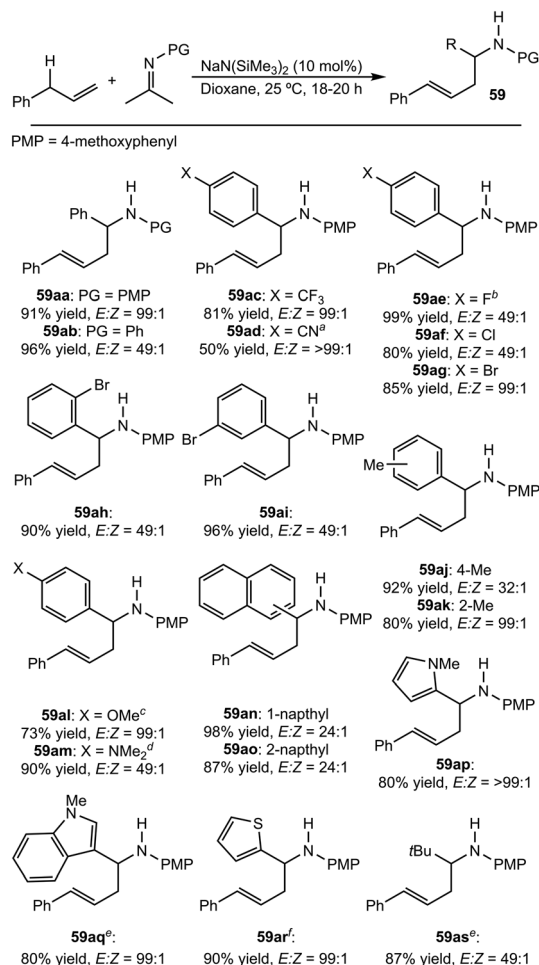


Fig. 7 Substrate scope for the coupling of terminal alkenes and imines by catalytic $\text{NaN}(\text{SiMe}_3)_2$.¹⁰⁹ All yields are isolated yields after preparative thin layer chromatography. ^a Use of 3 equiv. of alkene. ^b The reaction was conducted at 40 °C. ^c Use of 1.8 equiv. of alkene (successive addition). ^d Use of 2.5 equiv. of alkene (successive addition). ^e Use of 2 equiv. of alkene (successive addition). ^f The reaction was conducted with 1.5 equiv. of alkene at 60 °C for 72 h.

systems have also been explored.⁷⁵ The cost effectiveness of the latter makes acid/base mediated HIE and the development of well-defined alkali metal catalysts an engaging and exciting area of research.

6.1.1. Sodium. Taking advantage of the basicity of NaTMP, Tortajada and Hevia illustrate HIE *via* the perdeuteration of arenes. NaTMP, in combination with polyamine PMDETA and C_6D_6 as the deuterium source, showed catalytic deuteration of non-activated arenes (**60**) in $\text{C}(\text{sp}^2)\text{-H}$ and $\text{C}(\text{sp}^3)\text{-H}$ positions (Fig. 8).⁷⁶ Under these conditions, simple arenes (**60a-d**) showed excellent deuterium incorporation, while alkylbenzenes showed a pronounced decrease in deuteration *ortho*- to the alkyl. Symmetric substrates like hexamethylbenzene (95%), ferrocene (94%), and diphenylacetylene (97%) showed equal deuteration at all positions. Substrates that were incompatible with this deuteration system fell into two classes: those with functional groups prone to decomposition from organosodium

reagents, and those with more acidic C-H bonds (diphenylmethane and benzofuran) that rapidly form the metalated product. Further utility for HIE was demonstrated through the addition of ferrocene-*d*₁₀ to C_6H_6 , which afforded the 94% conversion to proteo-ferrocene with <5% deuteration.

The investigation into the structure-activity relationship for sodium amide reagents has led to progress in well-defined HIE reactions using sodium. This area of interest has encouraged the use of more sterically confined, super-basic, sodium amides such as NaTMP, NaNcy_2 , NaNAd_2 , and NaNAdTMS in tandem with TMEDA and PMDETA (Scheme 13).⁷⁷ The synthesis of the sodium amides was performed *via* the addition of *n*BuNa to the appropriate amine in hexanes. Deuteration studies were conducted using anisole as a model compound, with 10 mol% sodium amide and 10 mol% of an amine donor in C_6D_6 over 16 hours. The degree of deuteration by NaNAdTMS was very low, with less than 5% of deuterium incorporation across all positions in anisole using both PMDETA and TMEDA. Both NaTMP and NaNcy_2 , using PMDETA, saw deuterium incorporation of greater than 95% at all positions. NaTMP and TMEDA showed slight preference for the *meta*-position (>95%) compared to the *ortho*- or *para*-positions (82%) alongside selective deuteration at the methoxy group (85%). NaNcy_2 was significantly less effective at all three positions (*meta*:- <5%, *ortho*- or *para*:- 60%, methoxy: 24%).

Deuteration is not limited to extremely bulky amides, as the Hevia group demonstrated the potential for $\text{NaN}(\text{SiMe}_3)_2$ and $\text{NaCH}_2\text{SiMe}_3$ to selectively deuterate heterocycles, N-heterocyclic carbenes (NHCs), fluoroarenes, and other substrates using $\text{DMSO-}d_6$ as the deuterium source (Scheme 14A).^{78,79} When using $\text{NaN}(\text{SiMe}_3)_2$, nitrogen-based heterocycles showed selective deuteration at the more acidic protons (up to 95% exchange in many substrates) and other heterocycles (furan, thiophene) showed selectivity for the α -proton. Characterization of catalytic intermediates utilizing 2-methylpyridine revealed formation of the $\text{C}(\text{sp}^3)\text{-H}$ activated product as a sodiated dimer (**61**), which was predisposed to HIE. Additional crystallographic studies reveal a separated ion pair **62** that showed increased deuteration than both the *in situ* conditions and exclusive $\text{NaN}(\text{SiMe}_3)_2$ (Scheme 14B). Methyl-substituted pyridines preferentially deuterated the $\text{C}(\text{sp}^3)\text{-H}$ bonds of the methyl with more than >95% exchange (Scheme 14C). Faster exchange activity was observed using $\text{NaN}(\text{SiMe}_3)_2$ with toluene (37% compared to 68% after 16 hours), which was then applied to several less-activated substrates, yielding deuteration up to 95% in some cases.

6.1.2. Potassium, rubidium and cesium. Potassium bases ($-\text{OR}$, $-\text{OH}$, $-\text{CO}_3$) have been commonly used for HIE with D_2O and $\text{DMSO-}d_6$ as the deuterium source.⁸⁰ However, any potassium base in DMSO forms what is reported as “superbase media” with no well-defined intermediate, but still demonstrates HIE capabilities.⁸¹

The Guan group showed that both $\text{RbN}(\text{SiMe}_3)_2$ (91% D enrichment, 98% yield) and $\text{CsN}(\text{SiMe}_3)_2$ (97% D enrichment, 98% yield) complexes can selectively deuterate benzylic C-H bonds using D_2 gas at 10 mol% catalyst loading.⁸² The same



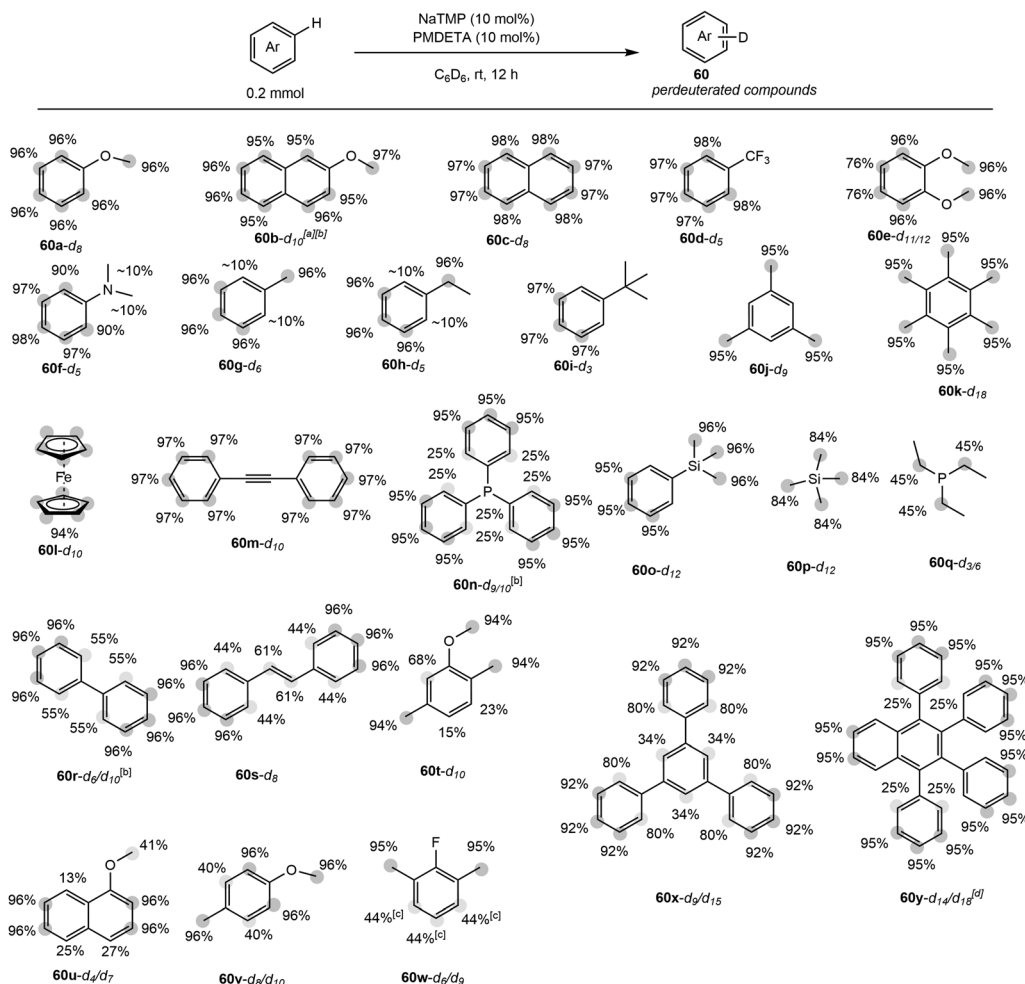
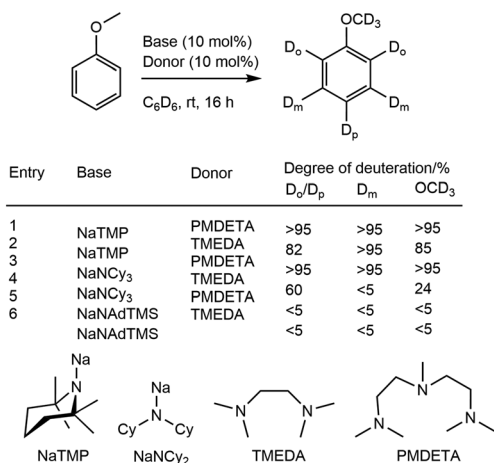


Fig. 8 Substrate scope for the perdeuteration of arenes by NaTMP. ^a 48 h, ^b 0.1 mmol scale, ^c average deuteration reported due to overlapping signals in ¹H NMR spectrum, ^d 0.05 mmol scale.

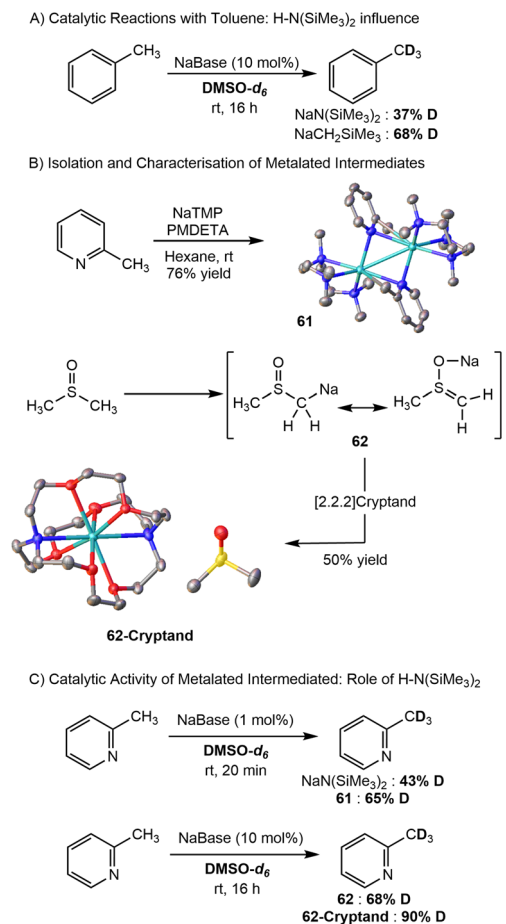


Scheme 13 Summary of deuterium incorporation with different base/donor systems.

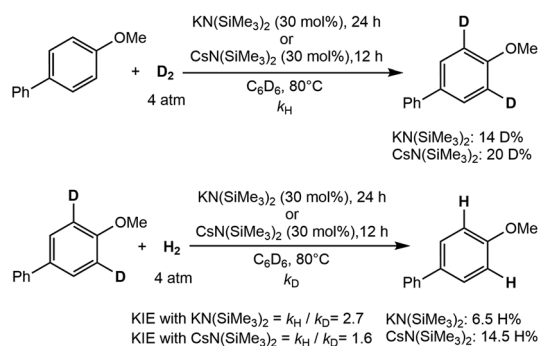
was observed for several drug molecules, with tritiation using T₂ gas performed on Tescmilifene, Imipramine, and Vortioxetine. The authors propose that AMN(SiMe₃)₂ (AM = Rb, Cs) reacts with benzylic substrates through σ-bond metathesis to form the benzylic metal intermediate, which is then reacted with D₂ to afford the deuterated product. Subsequently, a metal hydride intermediate is formed that is protonated again to continue the cycle, although the authors state that the metal hydride intermediate can be accessed first to afford the same deuterated product.

Further extension towards the *ortho*-directed HIE of various aromatic ethers and fluorides was also explored.⁸³ Comparatively, LiN(SiMe₃)₂ and NaN(SiMe₃)₂, along with other potassium bases exhibited no exchange, while KN(SiMe₃)₂ (60 mol%) showed 95% deuterium incorporation to 4-phenylanisole. However, the heavy alkali metals showed enhanced incorporation with CsN(SiMe₃)₂ (30 mol%) affording up to 97% deuterium incorporation (Scheme 15). Mechanistic experiments and DFT calculations reveal AMN(SiMe₃)₂ (AM = Na, K, Cs), in the dimeric form, coordinate three anisole mole-





Scheme 14 Summary of approaches for HIE using various sodium bases in DMSO-*d*₆.

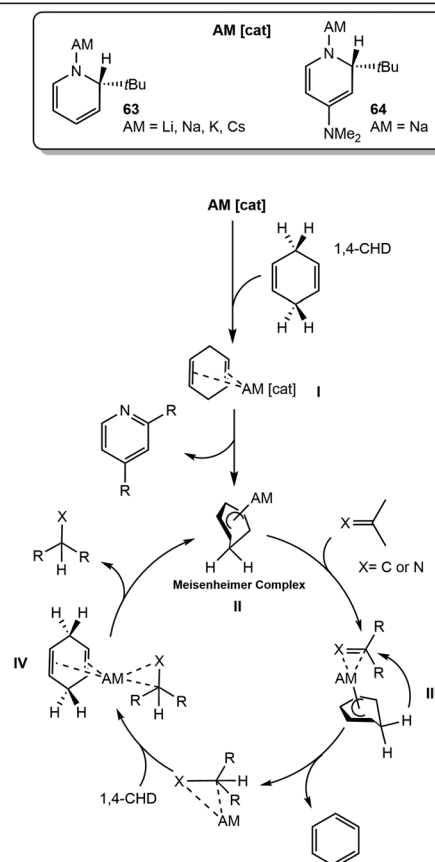
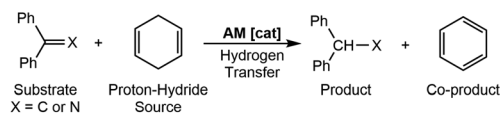


Scheme 15 KIE studies of KN(SiMe₃)₂ and CsN(SiMe₃)₂.

cules, in which one of the three is deprotonated by the metal, leaving an unstable carbanion intermediate that reacts with D₂. The previously released HN(SiMe₃)₂ reacts with the dimer, allowing for regeneration of the catalyst.

6.2. Transfer hydrogenation

Transfer hydrogenation is an alternative to direct hydrogenation, which removes the hazards of using hydrogen gas.



Scheme 16 Transfer hydrogenation of imines and alkenes using 1,4-cyclohexadiene (CHD) and the general mechanism undergoing a Meisenheimer-based intermediate.

There have been few reports of alkali metal bases (–OR, –OH) that have been shown to do transfer hydrogenation on carbonyls.^{84–87}

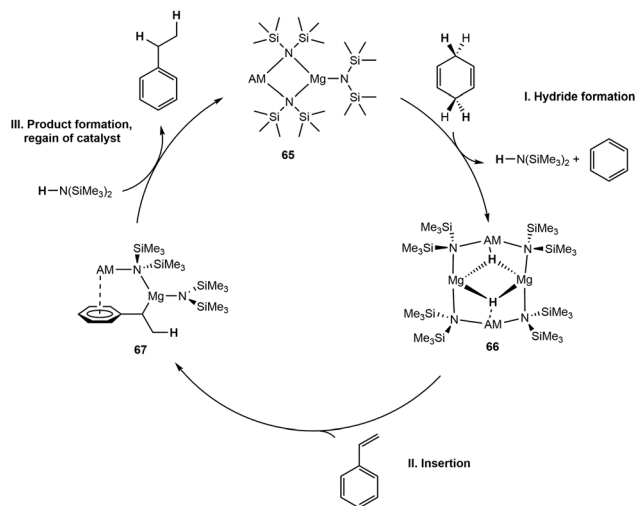
6.2.1. Sodium. Mulvey and coworkers focused on generating a “masked” sodium-hydride analogue *via* the dihydropyridylsodium compound (Na-1,2-*t*Bu-DH(DMAP)) (**64**), and its monomeric variant, [Na-1,2-*t*Bu-DH(DMAP)]Me₆TREN for catalytic transfer hydrogenation (Scheme 16).⁸⁸ Mechanistically, reduction of DPE was found to be facile from Na-1,2-*t*Bu-DH(DMAP) to generate the sodiated diphenylmethane *en route* to the Meisenheimer intermediate. The reactivity of these sodium complexes proved useful in the reduction of alkenes, outperforming those of NaN(SiMe₃)₂, NaTMP, and NaH with similar mechanistic profiles.

6.2.2. Potassium, rubidium and cesium. In follow-up studies, Mulvey and coworkers synthesized a series of mono-metallic alkali metal (AM = Li–Cs) dihydropyridines [AM (*t*BuDHP)] complexes (**63**) and explored their utility for transfer hydrogenation (Scheme 16).^{89,90} Catalytic competency for each



metal complex was tested using 1,4-cyclohexadiene (1,4-CHD) as the hydride source and *N*-benzylideneaniline as an initial substrate. The heavier congeners (Rb and Cs) were found to be higher yielding (92 and 97%) with fast reaction times. Similarly, full conversion and high yields (91%) with Cs(*t*BuDHP) was reported upon changing the substrate to the more polar *N*-benzylidene-*tert*-butylamine, in stark contrast to Li(*t*BuDHP), which showed minimal conversion (15%). Mechanistic studies and DFT allude to the mechanism of transfer hydrogenation for Cs(*t*BuDHP) following a base mediated initiated pathway by generation of a π -complex **I**, followed by deprotonation to form the key Meisenheimer intermediate **II**. This intermediate then coordinates with the substrate for hydride transfer **III**, releasing benzene as a byproduct in the process. Upon coordination of an additional equivalent of the 1,4-CHD to the metal center **IV**, the hydrogenated product is then released, regenerating the active catalyst.

Mulvey and coworkers synthesized a series of bimetallic magnesium and alkali metal complexes for exploring the catalytic transfer hydrogenation of alkenes, utilizing 1,4-CHD as the hydrogen source.⁹¹ A series of alkali metal magnesiate are synthesized by reacting $\text{Mg}(\text{N}(\text{SiMe}_3)_2)_2$ with $\text{AM}(\text{N}(\text{SiMe}_3)_2)$ (AM = Li–Cs) to form $\text{AMMg}(\text{N}(\text{SiMe}_3)_2)_3$ (**65**, Scheme 17). $\text{CsMg}(\text{N}(\text{SiMe}_3)_2)_3$ shows the best conversion for the hydrogenation of styrene to ethylbenzene with >98% conversion with 5 or 10 mol% at 75 °C within 30 minutes. The proposed mechanism for transfer hydrogenation utilizing these bimetallic manifolds (**65**) begins by hydride formation from the sacrificial 1,4-CHD to form the AM–Mg bridging hydride intermediate **66**. Intermediate **66** can then undergo insertion of styrene into the AM-stabilized Mg–H bond, forming **67**, followed by protonation by free $\text{HN}(\text{SiMe}_3)_2$, product release, and regeneration of the active catalyst. Oligomerization was observed as a major side reaction that the authors further investigated by utilizing 1,1-diphenylethylene (DPE) as a substrate. The studies



Scheme 17 Proposed catalytic cycle for the transfer hydrogenation of styrene.

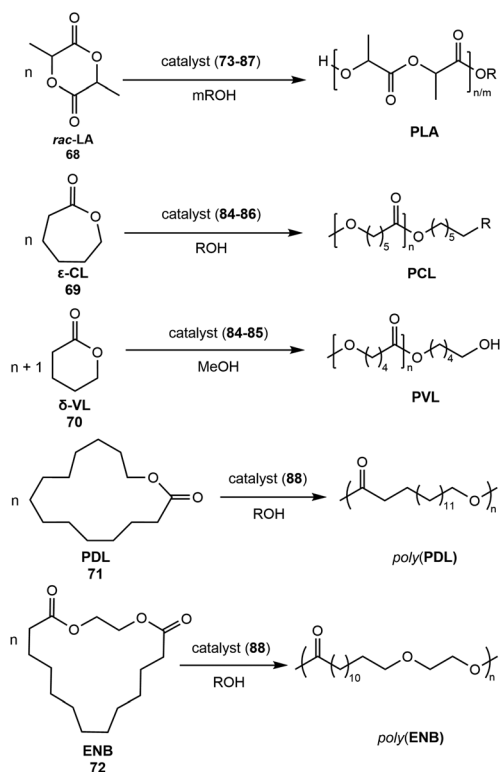
with DPE demonstrated that the potassium analogue was selective for the oligomerization of DPE, with full conversion in a shorter period than its larger congeners, Rb and Cs.

7. Ring opening polymerization

Ring opening polymerization (ROP) is the main method in which to produce industrially relevant polymers from cyclic monomers, such as lactones, lactams, and epoxides, and is mediated by a wide array of organo- and organometallic catalysts. Alkali metal reagents have been incorporated in base-catalyzed ROP, but the use of well-defined alkali systems is relatively new.⁹² The ring opening polymerization of cyclic *rac*-lactide (*rac*-LA, **68**) to form polylactic acid (PLA) is one of the most application-based polymerizations available to alkali metal systems. Extension to other cyclic lactones (**69**, **70**) and strainless macrolactones (**71**, **72**) has also been explored (Scheme 18). The usage of well-defined sodium and potassium metal precatalysts in this area over the last decade is summarized by ligand-supported classes.

7.1. Naphthalenolate catalysts

Crown ether-stabilized sodium and potassium catalysts were synthesized by the Wu group for the ROP of *rac*-lactide. In the case of the naphthalenolate complexes (**73**), the potassium



Scheme 18 General summary of ring-opening polymerizations discussed within.



complex showed higher activity than the sodium counterpart, achieving high isoselectivity ($P_m = 0.73$) and controlled molecular weights at room temperature, with rapid monomer conversion under optimized conditions (Fig. 9A).⁹³

7.2. Phenoxide catalysts

Using phenol ligands treated with different crown ethers and AMN(SiMe₃)₂ (AM = Na and K, 74) allowed access to a series of

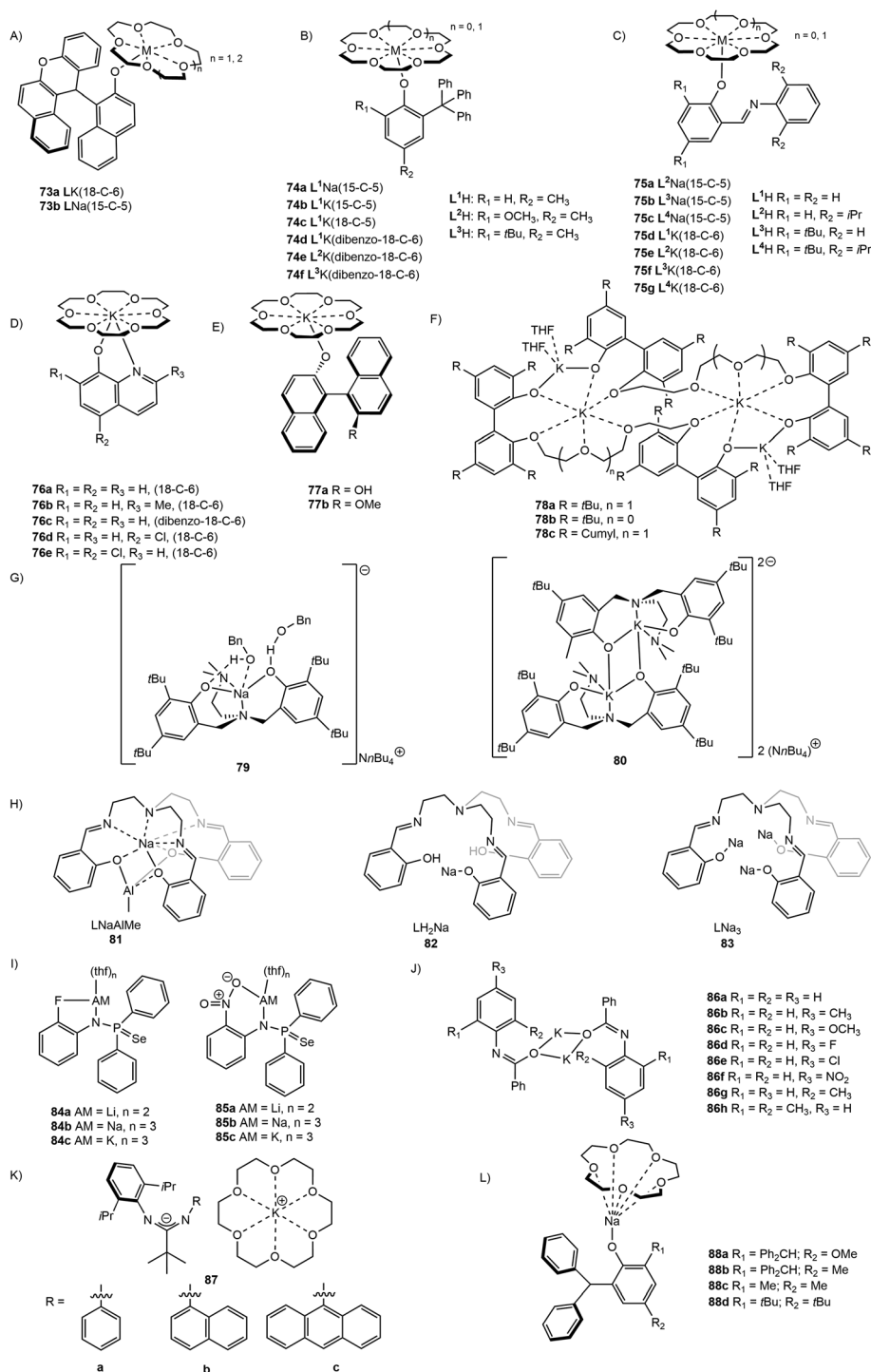


Fig. 9 (A) Naphthalenolate-supported catalysts, (B) phenoxide-supported catalysts, (C) iminophenoxide-supported catalysts, (D) quinoline-8-olate-supported catalysts, and (E) binolate-supported catalysts, (F) potassium phenolate catalysts, (G) aminobisphenolate ion-pair catalysts, (H) TrenSal-supported catalysts, (I) phosphoselenenic amide-supported catalysts, (J) potassium *N*-arylbzimidate catalysts, (K) amidinate-potassium separated ion-pair catalysts, (L) 15-C-5 supported sodium phenolate catalysts.



precatalysts.⁹⁴ The polymerization of *rac*-lactide using these catalysts proceeds rapidly at room temperature, achieving high conversion rates (90–97%) within one minute. Potassium-based systems demonstrated superior performance, with high isoselectivity ($P_m = 0.89$, -60 °C) indicating the efficacy of these alkali-metal catalysts, achieving the formation of multi-block isotactic copolymers with an average isotactic length of nine lactic acid units (Fig. 9B). Single crystal X-ray diffraction studies and experimental results demonstrate the impact steric hindrance and electronic effects have on this transformation, wherein it was determined that larger crown ethers and increased steric hindrance near the active center enhanced isoselectivity, while electron-donating groups diminished it.

Similarly, sodium and potassium iminophenoxide complexes supported by crown ethers have demonstrated successful ROP of *rac*-lactide (Fig. 9C). For both metals, Wang and coworkers observed that the presence of *ortho-tert*-butyl groups on the phenoxy-moiety of the ligand hindered the rate of ROP.⁹⁵ The potassium complex **75d** produced polylactides with the highest isoselectivity ($P_m = 0.75$); however, the sodium complex **75a** proved to be the most active catalyst. Similar potassium quinolin-8-olate complexes (**76**) were investigated by the same group (Fig. 9D).⁹⁶ Unlike **75d**, the **76a–e** exhibit *O,N*-chelation, which restricts rotation of the O–K bond, thereby influencing selectivity.

7.3. Binolate catalysts

Moving away from *O,N*-chelating scaffolds, Pan and coworkers demonstrated the activity of two enantiopure potassium binolate complexes (**77**) for the isoselective ROP of *rac*-lactide using (*R*)-2,2'-dihydroxy-1,1'-dinaphthyl (*R*-BINOL) ligands (Fig. 9E).⁹⁷ The presence of intramolecular hydrogen bonding in **77a** reduces catalytic activity but enhances stereoselectivity ($P_m = 0.80$ at room temperature). Although sodium complexes often show high activity for ROP, transesterification can occur as a side reaction; providing additional stabilization at the metal center reduces this deleterious reactivity. Lin and coworkers utilized *NNO*-tridentate Schiff base ligands with a pendant amino arm to minimize transesterification while also retaining a high catalytic activity at the sodium center for the ROP of *L*-lactide.⁹⁸

7.4. Aminobisphenolate catalysts

Wu and coworkers have also investigated the application and synthesis of bench-stable potassium complexes with multidentate phenolate ligands for the ROP of *rac*-lactide (**78a–c**, Fig. 9F).⁹⁹ Isoselectivity improved significantly at low temperatures due to the suppression of epimerization side reactions, reaching a maximum value of $P_m = 0.83$ at -70 °C. To increase ligand denticity, the tetradentate aminobisphenolate ligand was employed by Wu and coworkers to stabilize sodium and potassium metal centers.¹⁰⁰ Benzyl alcohol coordinates to **79** as both a Lewis donor at the sodium and through hydrogen bonding with the ligand *O*-atoms (Fig. 9G). This coordination does not occur in the case of potassium, which instead forms dimeric **80**. The best molecular weight control and stereoselectivity was observed after 6 hours at -70 °C using sodium catalyst **79** ($P_m = 0.82$, 87% conversion, 0.05 mol% catalyst).

7.5. Salen-based catalysts

In an attempt to balance activity with polymerization control, the Garden group has implemented the salen-derived TrenSal ligand to access three sodium-based catalysts for the ROP of *rac*-lactide (Fig. 9H).¹⁰¹ Of the three catalysts, **81** was the least active but demonstrated the best control (77% conversion in 40 min, $D = 1.25$ – 1.49). Slightly more active (76% conversion in 10 min), **82** provided improved control ($D = 1.75$); however, late-stage transesterification led to dispersity broadening. The improved activity of precatalyst **83** (83% conversion in 4 min) is attributed to the increased number of metal centers, but resulted in poor control over polymerization ($D = 2.00$). In addition to the activated monomer pathway demonstrated by **81** and **82**, MALDI-ToF end group analysis of PLA generated by **83** is consistent with that produced by a coordination-insertion mechanism, suggesting that **83** proceeds *via* two separate mechanisms, resulting in poor control. Tetradentate aminophenolate potassium complexes were applied to the ROP of *rac*-lactide by Ma and coworkers, resulting in high catalytic activity but very little stereoselectivity. Solid state structures were obtained by SC-XRD as dimeric potassium complexes, while DOSY and variable temperature NMR were used to determine monomeric complexes existed in solution.¹⁰²

7.6. Phosphoselenoic amide catalysts

Anionic ROP provides the benefit of improved control. Panda and coworkers applied phosphoselenoic amide alkali metal complexes (**84**, **85**) toward the ROP of *rac*-lactide, ϵ -caprolactone, and δ -valerolactone (Fig. 9I).¹⁰³ The potassium catalysts displayed the highest activity with **85c**, featuring a nitro-functionalized ligand, achieving the highest isoselectivity ($P_m = 0.78$) and converting 1000 equivalents of *rac*-LA to PLA within 30 minutes at room temperature. The steric and electronic properties of the ligands significantly influence the reaction, with nitro-substituted ligands enhancing catalytic activity and stereocontrol.

7.7. Arylbenzimidate and amidinate catalysts

Potassium *N*-arylbenzimidates demonstrate efficient anionic ROP in the presence of benzyl alcohol for ϵ -caprolactone and *L*-lactide.¹⁰⁴ Structural characterization revealed diverse coordination geometries across the eight complexes; some uniquely formed two-dimensional polymeric assemblies (**86a–h**, Fig. 9J), stabilized by π -aryl and σ -donor interactions. For ϵ -caprolactone, conversions exceeded 95% under optimized conditions (0.4 mol% catalyst, 50 °C, 30 minutes), affording polycaprolactones with controlled molecular weights and narrow dispersity. Electron-donating groups improved catalysis, while electron-withdrawing groups hindered it. The polymerization of *L*-lactide showed even greater efficiency, achieving nearly 100% conversion at optimized conditions (0.2 mol% catalyst loading, 60 °C, 30 minutes).

Wu and colleagues report the isoselective anionic ROP of *rac*-lactide using ion-paired potassium amidinate complexes (Fig. 9K).¹⁰⁵ Three potassium amidinate pairs (**87a–c**) were



accessed, with differing degrees of steric hindrance present at the amidinate side arm. Single crystal X-ray diffraction studies were utilized to confirm the formation of separated ion-pair complexes, wherein the potassium is not bound to the amidinate. Reduced temperature ($-30\text{ }^{\circ}\text{C}$ or $-70\text{ }^{\circ}\text{C}$) was found to improve polymer conversion for all three catalysts, which in addition to the presence of BnOH as an initiator, minimized unfavorable side reactions. Under optimized conditions, **87b** showed 92% conversion and low dispersity ($D = 1.05$).

7.8. Phenolate catalysts

Compared to lactide monomers, which can easily undergo ROP, macrolactones are generally harder to open due to reduced ring strain. During the ROP process, transesterification rates are similar to those of chain propagation rates. Liu and coworkers report the ROP of macrolactones (**71**, **72**) using sterically hindered phenoxide sodium complexes with 15-crown-5 (**88a-d**) wherein steric repulsion between the ligand and polymer chain inhibits transesterification, allowing for polymerization to proceed in a controlled manner (Fig. 9L).¹⁰⁶

8. Conclusions and perspectives

From the initial study of organolithiums in the 1930's, organoalkali chemistry has evolved significantly in the context of organic transformations. Simple organoalkali reagents have been extensively studied using modern and updated techniques, such as multinuclear NMR spectroscopy and single crystal X-ray diffraction, allowing these reagents to be used more effectively in a variety of applications. Taken together, these recent advances illustrate the dynamic role of AMs in modern organic chemistry, which are now central to strategies spanning ligand design, small-molecule activation, asymmetric catalysis, and flow chemistry. A recent example is the design of a chiral phenanthroline-potassium catalyst that enables the enantioselective ring opening alcoholysis of biaryl lactams to produce axially chiral δ -amino acid derivatives.^{107,108} Additionally, Hilt and coworkers have demonstrated that organosodium intermediates can also be generated and used under continuous-flow conditions for the synthesis of alkyl aryl ketones from Weinreb amides.¹⁰⁹ As these studies demonstrate, the reactivity of alkali metals is set to inspire even broader applications in organic chemistry.

Conflicts of interest

There are no conflicts to declare.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

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