

CHEMISTRY

Direct transformation of dinitrogen: synthesis of *N*-containing organic compounds via N−C bond formation

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ABSTRACT

N-containing organic compounds are of vital importance to lives. Practical synthesis of valuable *N*-containing organic compounds directly from dinitrogen (N_2) , not through ammonia (NH_3) , is a holy-grail in chemistry and chemical industry. An essential step for this transformation is the functionalization of the activated N_2 units/ligands to generate N−C bonds. Pioneering works of transition metal-mediated direct conversion of N_2 into organic compounds via N−C bond formation at metal-dinitrogen $[N_2-M]$ complexes have generated diversified coordination modes and laid the foundation of understanding for the N−C bond formation mechanism. This review summarizes those major achievements and is organized by the coordination modes of the $[N_2-M]$ complexes (end-on, side-on, end-on-side-on, etc.) that are involved in the N−C bond formation steps, and each part is arranged in terms of reaction types (*N*-alkylation, *N*-acylation, cycloaddition, insertion, etc.) between [N2-M] complexes and carbon-based substrates. Additionally, earlier works on one-pot synthesis of organic compounds from N_2 via ill-defined intermediates are also briefed. Although almost all of the syntheses of *N*-containing organic compounds via direct transformation of N_2 so far in the literature are realized in homogeneous stoichiometric thermochemical reaction systems and are discussed here in detail, the sporadically reported syntheses involving photochemical, electrochemical, heterogeneous thermo-catalytic reactions, if any, are also mentioned. This review aims to provide readers with an in-depth understanding of the state-of-the-art and perspectives of future research particularly in direct catalytic and efficient conversion of N_2 into *N*-containing organic compounds under mild conditions, and to stimulate more research efforts to tackle this long-standing and grand scientific challenge.

Keywords: dinitrogen transformation, metal-dinitrogen complex, N−C bond formation, *^N*-containing organic compounds

INTRODUCTION

As the most abundant constituent in Earth's atmosphere (atm), dinitrogen (N_2) is the main nitrogen source of *N*-containing compounds on the Earth. Therefore, $N₂$ fixation and activation are essential both for nature and humans. Nevertheless, the high bond dissociation energy (942 kJ/mol) and large highest occupied molecular orbital (HOMO) lowest unoccupied molecular orbital (LUMO) gap (10.82 eV) make N₂ exhibit extremely low reactivity and be regarded as an inert gas. Currently, the N_2 fixation and conversion in nature and industry mainly rely on two pathways, in which ammonia (NH_3) is the product $\begin{bmatrix} 1 \end{bmatrix}$. In nature, nitrogenase metalloenzymes employ iron-sulfur clusters asthe key cofactor (FeMo, FeV or FeFe cofactor) and water as the proton source to transfer N_2 into NH_3 at ambient temperature and pressure $[2]$. This biosynthetic NH₃ is a versatile precursor for the synthesis of *N*-containing organic compounds, such as amino acids and nucleic acids. Although the precise biological N_2 reduction mechanism is still controversial, spectroscopic and computational studies suggested the presence of an interstitial carbon atom at the center of the FeMo and FeV cofactors $\lceil 3-5 \rceil$.

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In industry, more than 170 million metric tons of $NH₃$ is produced from the Haber-Bosch process annually, in which N_2 reacts with dihydrogen $(H₂)$ under the harsh condition in the presence of iron or ruthenium catalysts. This $NH₃$ synthesis process consumes 1–2% of the world's annual energy supply along with the huge $CO₂$ emission, due to the drastic reaction condition and the energy requirement for H_2 production from fossil fuels and water [\[6\]](#page-17-4). As the main route of N_2 fixation and transformation in industry, \sim 20% of NH₃ produced from the Haber-Bosch process is used as the feedstock to produce *N*-containing chemicals, including higher-value *N*-containing organic compounds, like amines, nitriles, nitro and so on. To better understand the reaction mechanism of biological and industrial reduction of N_2 into NH_3 , several catalytic systems including homogeneous molecular systems, electrochemical systems and heterogeneous systems have been studied for decades, and there are comprehensive reviews that readers may refer to [\[7–](#page-17-5)[13\]](#page-17-6).

Compared to NH_3 -based N_2 fixation process, an alternative route of N_2 fixation is the direct conversion of N_2 into *N*-containing organic compounds under mild condition. This approach is always targeted because it provides the potential solution to developing a sustainable system with reduced fossil fuel requirements. The earliest study towards this goal began in the 1960s, when Vol'pin *et al.* discovered that the titanium species, for example, Cp_2TiCl_2 could react with PhLi under N_2 to give aniline after hydrolysis $[14]$. However, further application of this reaction was hindered by the low yields and the lack of reaction details. During the same period, the first metal-dinitrogen (N_2-M) complex $\left[\text{Ru(NH_3)_{5}(N_2)\right]^{2+}}$ was reported in 1965 [\[15\]](#page-18-1). After that, thousands of N_2 -M complexes have been documented $[16]$. The reactivity exploration reveals that the functionalization of the N_2 ligands can also be fulfilled for some N_2 -M com-plexes [\[17\]](#page-18-3). Making N−C bonds from the reactions of transition metal N_2 complexes with carbonbased reagents has received much attention in recent decades, although the catalysis system has not been realized $[18,19]$ $[18,19]$.

This review will focus on the previous works regarding the transformation of N_2 into organic compounds. In almost all of these works, the N−C bond formation steps are fulfilled upon the welldefined N_2 -M complexes with diversified coordination modes. This review is organized by the coordination modes of the N_2 -M complexes (to clarify, the N_2 -derived metal nitrides are also considered as a coordination mode of N_2 -M complexes) that

Figure 1. The classification of this review. The N–C bond formation is reported $(\sqrt{})$ or not reported (\times) .

are involved in the N−C bond formation steps, and each part is arranged in terms of the type of reactions between N_2 -M complexes and carbon-based substrates. The earlier works about one-pot synthesis of organic compounds from N_2 via ill-defined intermediates are also introduced briefly in this review (Fig. [1\)](#page-1-0).

N−**C BOND FORMATION VIA END-ON N2-M COMPLEXES**

End-on bond is the most prevalent bonding mode for N_2 -M complexes and the N_2 -M complexes with this binding mode have been known to assemble N−C bond for a long time. Main works were achieved via the reaction of end-on terminal N_2 -M complexes with alkyl or acyl halides and their analogues. N−C bond formations from the cycloaddition and insertion reactions of end-on-bridged N_2 -M complexes with imido-like N_2 ligands have also been reported.

Scheme 1. *N*-alkylation of end-on terminal N_2 -Mo complexes by electrophiles. (a) *N*-methylation of N2-Mo complex by MeOTs to afford methyldiazenido complex. (b) N-methylation of N₂-Mo complexes by MeOTs and MeOTf to afford methyldiazenido and *N*,*N*-dimethylhydrazido complexes.

*N***-alkylation**

N-alkylation by electrophiles

The strong electrophilic alkyl triflates and their analogues are often employed to functionalize the endon N_2 -M complexes because the N_2 ligands in these complexes feature a nucleophilic character by electron donation from the electron-rich metal centers. Peters *et al.* [\[20\]](#page-18-6) and Greco and Schrock [\[21\]](#page-18-7) reported that the methylation reaction occurs when the anionic end-on N_2 -Mo complexes 1 and 3 are treated with methyl tosylate (MeOTs) to provide methyldiazenido complexes **2** and **4** (Scheme [1a](#page-2-0) and b). Additionally, **⁴** could further undergo N−^C bond formation to furnish *N*,*N*-dimethylhydrazido complexes **5** by reaction with excess methyl triflate (MeOTf) or MeOTs (Scheme [1b](#page-2-0)).

Although many late-transition metal complexes with end-on N_2 ligands have been documented, reports on their reactivity toward electrophiles to make N−C bond are very rare. Peters*et al.* described that the anionic end-on terminal N_2 complexes of Fe **6** and Co **8** react with MeOTs to give *N*-methylation species **7** and **9** (Scheme [2a](#page-2-1)) [\[22\]](#page-18-8). In 2016, the same group found that the modified N_2 -Fe complex **10** bearing monoanionic tetradentate trisphosphinosilyl ligand can also be alkylated to afford *N,N*-dimethylated product **11** (Scheme [2b](#page-2-1)) [\[23\]](#page-18-9).

N-alkylation by *in situ* formed radicals

There are only a few examples of N−C bond formation at N_2 -M complexes by radicals. One ex-

Scheme 2. *N*-alkylation of end-on terminal N_2 -Fe, Co complexes by MeOTf or MeOTs. (a) *N*-methylation of N₂-Fe, Co complexes by MeOTs to afford methyldiazenido complexes. (b) N -methylation of N_2 -Fe complexes by MeOTf to afford *N*,*N*-dimethylhydrazido complex.

ample is the reactions of terminal end-on N_2 -Mo, W complexes **12** with alkyl halides, driven by light (*vide infra*). Mechanism investigation reveals that the radicals in these reactions are generated *in situ* by the homolysis of the alkyl halides within the coordination sphere. The attacking of these alkyl radicals at the N_2 ligands provides 13. Furthermore, dialkylhydrazido complexes **14**, **15** and **16** can also be obtained via alkylation of **13** or one-pot dialkylation of 12 (Scheme [3a](#page-3-0)) $\lceil 24,25 \rceil$. It is noteworthy that if the diphosphine ligands in **12** are replaced by the monophosphine ligands, the corresponding N_2 -Mo, W complexes fail to react with alkyl halides to assemble N−C bond. Another example that involves the radical mechanism is the *N*-functionalization of the terminal end-on N_2 -Mo complex **17**, which possesses higher reactivity than **12** (Scheme [3b](#page-3-0)) [\[26\]](#page-18-12). For instance, the treatment of **17** with BnBr or aryl iodide gives the *N*,*N*dibenzylation product **18** or *N*-arylation complex **19,** the latter of which can also be converted to the organo-hydrazido species **20** by further reaction with MeI. More intriguingly, when **17** is treated with MeI in toluene, the prospective product **21** is formed together with isolation of an unexpected product **22**. A plausible mechanism is raised for the generation of **22** (Scheme [3c](#page-3-0)). The initial reaction between **17** and MeI results in iodine atom abstracting to afford intermediate **A** and the methyl radical, which would abstract an *H*-atom from toluene to yield benzyl radical. The latter reaction between **A** and benzyl radical gives the *N*-benzylation intermediate **B**, which can further react with MeI to afford the final product. The formation of **22** confirms

Scheme 3. *N*-alkylation of end-on terminal N_2 -Mo, W complexes by *in situ* formed radicals. (a) *N*-alkylation of N2- Mo, W complexes supported by diphosphine ligands. (b) *N*alkylation of N_2 -Mo complex supported by tetra-thioether ligand. (c) A plausible mechanism for the generation of **22**.

the radical process of these *N*-alkylation reactions again.

N-alkylation by nucleophiles

For end-on terminal N_2 -M complexes, simple Lewis formulas could be used to depict their structures. As shown in Scheme [4a](#page-3-1), the *N* atom adjacent to the metal atom (N_{α}) features positive charge and could be attacked by nucleophiles in theory. Surprisingly, there is only one example of this reactivity for N_2 -M complexes [\[27,](#page-18-13)[28\]](#page-18-14). Sellman *et al*. found that an end-on terminal N₂-Mn complex 23 reacts with methyl or phenyl lithium reagent at low temperature to give the N_α -functionalize[d](#page-3-2) products 24, which could subsequently react with Meerwein reagent $Me₃OBF₄$ upon N_{β} atom to afford 25. This

Scheme 4. Manganese-promoted direct conversion of N_2 into azomethane via the reaction between nucleophiles and N2-Mn complex. (a) Simple Lewis formulas for end-on terminal $N₂$ -M complexes. (b) A synthetic cycle for synthesis of azo-compound from N_2 .

Scheme 5. *N*-acylation of end-on terminal N_2 -Mo, N_2 -W complexes.

azomethane complex would ultimately liberate free azomethane by pressuring with 100 bar of N_2 along with reforming N2-Mn complex **23**.Thus, a synthetic cycle was raised for synthesis of azo-compound from N_2 .

*N***-acylation**

Besides alkyl halides, acyl chlorides are also used to functionalize end-on N2-M complexes. Chatt *et al.* found that the N_2 -Mo, W complexes 12 supported by bidentate phosphines ligands react with acyl chloride to afford acyldiazenido complexes (Scheme [5\)](#page-3-2) [\[29,](#page-18-15)[30\]](#page-18-16). These *N*-acylation reactions possibly proceed through nucleophilic attacking of the N_2 ligands on the acyl carbons.

Cycloaddition and insertion

For some end-on-bridged N_2 -M complexes with strongly activated N_2 ligands, the imido-like structures make them able to undergo cycloaddition or insertion reactions with carbon-based

Scheme 6. N–C bond formation from cycloaddition of end-on-bridged N₂-Ti complex with phenylallene, ^{*t*}BuNCO and CO₂.

unsaturated substrates to assemble N−C bond. In 2017, the reaction of an end-on bridging binuclear N_2 -Ti complex 27 with phenylallene, *tert*-butyl isocyanate (*^t* BuNCO) and CO2 was investigated by Kawaguchi *et al*., to provide N−C bond formation products (Scheme [6\)](#page-4-0) [\[31\]](#page-18-17). Treatment of **27** with an excess of phenylallene results in the formation of dititanium hydrazido complexes **28** and **29** as a mixture of isomers. The formation of **28** and **29** can be rationalized in terms of an initial $[2+2]$ cycloaddition of phenylallene with $Ti = N$ bond in 27 to give the 4-membered titanacycle intermediates (two isomers), and the subsequent protonolysis of the Ti−C bonds in these intermediates to give the final products. Further studies indicate that the proton source in this reaction could be a second equivalent of phenylallene, the ancillary ligands, or even adventitious impurities present in the reaction mixture. The reaction of **27** with *^t* BuNCO also proceeds through a formal $[2+2]$ cycloaddition reaction to afford **30**. However, when **27** is introduced with an atm of $CO₂$, the insertion of three molecules of CO_2 into Ti = N bonds in 27 is achieved to furnish **31**. By adding an excess amount of TMSCl, **31** could be converted to organic compound $N_2(TMS)(CO_2TMS)_3$, which is unstable under the reaction condition and decomposes to two hydrazine derivatives $[TMS(CO₂TMS)N]_{2}$ and $(TMS)_2NN(CO_2TMS)_2$ via decarboxylation.

The cycloaddition reactions between group 5 end-on $N₂$ -M complexes and carbon-based unsaturated bonds have also been observed. For example, N_2 -Nb complex 32 and N_2 -Ta complex 33 with diimido bridging N_2 ligands are known to react with aldehyde and acetone to afford the corresponding ketazines (Scheme [7\)](#page-4-1) [\[32,](#page-18-18)[33\]](#page-18-19).

Scheme 7. N−C bond formation from the reactions of end-on-bridged N_2 -Nb. Ta complexes with aldehyde or acetone. (a) The reaction of N_2 -Nb complex with benzaldehyde. (b) The reaction of N_2 -Ta complex with acetone.

*N***-protonation/carbonylation**

An alternative route for making N−C bond is the treatment of carbon-based substrates with the *N*-hydrogenated complexes derived from N_2 because in some cases *N*-hydrogenation are more accessible than *N*-alkylation for end-on N_2 -M complexes. Seminal works about these transformations were finished by Hidai and others [\[25](#page-18-11)[,34\]](#page-18-20). They reported that the N2-Mo, W complexes **34** and **12** supported by monophosphine or diphosphine ligand react with HX ($X = Cl$, Br and I) or HBF₄ to afford the hydrazido complexes **35** and **36**, which can act as the versatile precursors to construct N−C bond [\[25,](#page-18-11)[35\]](#page-18-21) (Scheme [8a](#page-5-0)). For instance, **35** could react with diphenylketene and phthalaldehyde to provide **37** and **38**, while the reaction between **36** and succinyl chloride gives rise to **39**. More intriguingly, these hydrazido complexes **35** and **36** are also reported to undergo a condensation reaction with ketones and aldehydes in the presence of catalytic amounts of acid to afford all kinds of diazoalkane

Scheme 8. N−C bond formation from the reactions of hydrazido Mo, W complexes with carbon-based reagents. (a) The reaction of N_2 -Mo, W complexes with HX (X = CI, Br and I) or HBF_4 to afford the hydrazido complexes. (b) Carbonylation of hydrazido complexes **35** and **36** to assemble N−C bond. (c) A synthetic cycle for synthesis of 1H-pyrrole from N_2 .

complexes **40** and **41** (Scheme [8b](#page-5-0)). The liberation of the *N*-containing organic compounds from these *N*-functionalized complexes has also been explored [\[36\]](#page-18-22). For example, when the cyclic hydrazido complex **42**, produced from the reaction of **36** and the cyclic acetal of succinaldehyde, is treated with LiAlH4, the reductive destruction of **42** is observed to release 1H-pyrrole accompanied by the generation of the tetrahydride complex **43**. Furthermore, this tetrahydride tungsten complex could be converted to the initial N_2-W complex **12** under photolytic conditions to achieve a cycle (Scheme [8c](#page-5-0)) [\[37\]](#page-18-23).

Involvement of photochemistry

Photochemistry is an emerging approach for the transfer of N_2 . The earliest observation of photocatalyzed N–C bond formation of N₂-M complexes

Scheme 9. An electrochemical cycle for synthesis of piperidine direct from N_2 via end-on terminal N_2 -Mo, W complexes.

is of the reactions between end-on terminal N_2 -M complexes **12** and alkyl halides (Scheme [3a](#page-3-0)) [\[25\]](#page-18-11). In the case of N_2 -W complex, visible light or a tungstenlamp is often necessary for these *N*-alkylation reactions. However, for the N_2 -Mo complex, it could react with alkyl bromide slowly in the dark. It is also reported that the N_2 ligands in 12 are not evolved in the absence of the alkyl halides since irradiation of the N_2 -M complexes without organic halide caused no change. These results indicate the possibility of photo engaging in the assistance of alkyl radicals formation in these reactions [\[24\]](#page-18-10).

Involvement of electrochemistry

Besides photochemistry, electrochemistry is another versatile method in the N_2 conversion process. Although the direct involvement of electrochemistry in the N−C bond formation step has not been discovered, the electrochemical reduction of the *N*-alkylated complexes to release the final organic products has been developed. For example, the organohydrazido complexes **16**, which is synthesized from the reaction of N_2 -M complexes 12 with 1,5-dibromopentane, undergoes electrochemical reduction at a Pt electrode in tetrahydrofuran (THF) under N_2 by using [NBu₄][BF₄] as the electrolyte to liberate piperidine accompanied by the regeneration of N2-M complexes **12**. According to the control experiment under the atmosphere of Ar or CO, a M(II) hydrazido intermediate is proposed in this piperidine releasing process. Based on these results, an electrochemical cycle to synthesize dialkylhydrazine from N_2 was reported by Leigh *et al*. (Scheme [9\)](#page-5-1) [\[38\]](#page-18-24).

N−**C BOND FORMATION VIA SIDE-ON N2-M COMPLEXES**

The side-on bonding modes are often observed at group 3 and group 4 transition metal N_2 -M complexes. The N_2 ligands in these side-on N_2 -M

*N***-alkylation**

There are two reports about the reaction of group 4 side-on N_2 -M complexes with alkyl halides or their analogues to make N−C bond. One example was reported by Hirotsu *et al*. in 2007, in which the side-on-bridged N₂-Hf complexes 44 with extremely activated $(N_2)^{4-}$ ligands can react with ethyl bromide (EtBr) to provide the *N*-ethylated products **45** (Scheme [10a](#page-6-0)) [\[39\]](#page-18-25). Controlled experiments indicate that this reaction is remarkably sensitive to the steric effects of the ancillary ligands. For example, when the R' group in **44b** is changed from Et to *ⁱ* Pr, the corresponding *N*-ethylation product could not be obtained. Besides, **45a** and **45b** fail to undergo further *N*-ethylation, even in the presence of excess EtBr. The other work is reported by the reaction of methyl triflate (MeOTf) with a hafnocene complex **46** that also bears side-on bridging $(N_2)^{4-}$ ligand (Scheme [10b](#page-6-0)) [\[40\]](#page-18-26). This reaction offers a mixture of products and one of them is the N_2 ligand monomethylated product **47**, which could be converted to the final organic compound *N*-methylhydrazine by treating with excess HCl. Besides, an unprecedented triflato hafnocene hydrazonato complex **48** is generated via a second N−C bond formation when additional MeOTf is added to **47**.

Compared with the group 4 transition metals, rare-earth metal promoted direct conversion of N_2 into organic compounds attracts less attention. The only example of this topic was reported by Xi, Zhang *et al.* in 2019 (Scheme [11\)](#page-6-1) [\[41\]](#page-18-27). Treatment of the $(N_2)^{3-}$ -bridged discandium complex **49** with MeOTf leads to the formation of *N*,*N* dimethylation discandium complex **50** in 43% yield. The yield of **50** can be improved via the treatment of **49** with MeOTf and potassium several times. Transformation of the $(N_2Me_2)^{2-}$ ligand into organic compounds could be accomplished by treatment of 50 with I₂, HCl, BnBr and acyl chloride to afford azomethane, 1,2-dimethylhydrazine and a series of tetra-substituted hydrazine derivatives, concomitant with the regeneration of the precursors of the N2-Sc complexes. Hence, a three-step synthetic cycle for scandium-mediated direct conversion of N_2 and carbon-based electrophiles to multi-substituted hydrazine derivatives could be realized. The insertion of a CO molecule into the Sc−N bond of **⁵⁰**

Scheme 10. *N*-alkylation of side-on-bridged N₂-Hf complexes by EtBr or MeOTf. (a) *N*-ethylation of N₂-Hf complex by EtBr. (b) N-methylation of N_2 -Hf complex by MeOTf.

Scheme 11. Scandium-promoted direct conversion of N_2 into hydrazine derivatives via the reaction between MeOTf and N_2 -Sc complex.

with further N−C bond formation is also observed to provide **51**[.](#page-6-0)

Cycloaddition and insertion

The group 4 side-on bridging N_2 -M complexes are known to undergo cycloaddition and insertion reactions with carbon-based reagents that contain $C = X$ (X = N, O) bonds or C≡C bond, such as carbon dioxide (CO_2) , isocyanates $(RNCO)$ or alkynes, owing to their imido-like reactivity. Compared with *N*-alkylation and acylation reactions, these cycloaddition and insertion reactions are more atom-efficient for N_2 functionalization,

Scheme 12. N–C bond formation of side-on N₂-Zr complex by reaction with alkynes.

because the formation of transition metal halides and inorganic salts as the by-products is avoided in these reactions. The earliest study of the cycloaddition reactions between group 4 N_2 -M complexes and unsaturated bond to assemble N−C bond was finished by Fryzuk *et al.* via the reaction of side-on bridging N2-Zr complex **⁵³** with arylacetylene (RC≡CH; $R = Ph$, 4-Me-C₆H₄ and 4-^tBu-C₆H₄) (Scheme [12\)](#page-7-0) [\[42\]](#page-18-28). The *N*-functionalization products **54** may result from a sequence of two successive steps: cycloaddition of alkyne across a Zr−N bond in **⁵³** leading to the zircona-aza-cyclobutene intermediate **^D**, which subsequently encounters Zr−C bond cleavage by protonation with another molecular terminal alkyne to yield **54**.

By elegant modulation of the substitutions on the multi-substituted Cp ligands, Chirik *et al*. accomplished a series of reactions of dinuclear N_2 -Zr, Hf complexes with isocyanates or $CO₂$ to assemble ^N−C bond. N2-Hf complex **⁴⁶** bearing tetramethylcyclopentadienyl (Cp^{4Me}) ligand is reported to react with PhNCO to provide the initial product **56** via a possible intermediate **55**. In the solution, **56** also reacts quickly with another molecule of ArNCO $(Ar = Ph \text{ and } p \text{-MeC}_6H_4)$ to afford 57, which could also be prepared directly from **4[6](#page-7-0)** (Scheme [13a](#page-7-1)) [\[43\]](#page-18-29). Besides, further studies indicate that another *N*-functionalization product **58**, in which the same nitrogen atom is di-carboxylated, would be formed predominately when $CO₂$ $CO₂$ $CO₂$ is bubbled into a solution of **46** (Scheme [13b](#page-7-1)) [\[44\]](#page-18-30). Subsequent reaction of **58** with TMSI gives rise to the generation of **59**, which is known to liberate the corresponding hydrazine derivative $(TMS)_2NN(CO_2TMS)_2$ by further reacting with excess TMSI. Unfortunately, the similar *N*-functionalization reactions of PhNCO and CO2 with zirconium congener of **46** are unsuccessful, which is believed to be caused by the deleterious ligand-induced side-on, end-on isomerization of the $(N_2)^{4-}$ ligand. Hence, a [Me₂Si]-bridged ansa-zirconocenes N_2 complex 60 with higher energy barrier for the side-on, end-on isomerization was designed and prepared to investigate the reactivity toward $CO₂$ [\[45\]](#page-18-31). The treatment of 60 with

Scheme 13. N−C bond formation from the reactions of the side-on $N₂$ -Zr, Hf complexes with isocyantes and $CO₂$. (a) The reaction of N_2 -Hf complex with PhNCO. (b) The reaction of N_2 -Hf complex with CO₂. (c) The reaction of N_2 -Zr complex with $CO₂$.

 $CO₂$ leads to the immediate generation of 61, where N2-functionalization takes place at each *N*-atom. Organic compound *N*,*N* -dicarboxylated hydrazine can be released from **61** by reacting with TMSI. Furthermore, a second N−C bond formation occurs when **61** is treated with MeOTf to provide **62**, which is known to liberate region-specific hydrazine

Scheme 14. CO-induced N_2 scission and functionalization at side-on N_2 -Zr and Hf complexes. (a) The reaction of N_2 -Zr, Hf complexes and CO. (b) A plausible mechanism for this CO-induced N_2 scission and functionalization reaction.

 $X(COOMe)NN(COOMe)Me (X = H, TMS) by$ further reacting with H_2O or TMSI (Scheme [13c](#page-7-1)). These results indicate that small modifications of the ligands will change the reactivity of the N_2 -M complexes dramatically.

CO-induced N2 functionalization and cleavage

Being isoelectronic with N_2 , CO is an abundant and cheap diatomic molecule with BDE of 1 079 kJ/mol. Hence, the transformation of CO and N_2 into N−C bond is a challenge but a significant process. Until now, only two systems of CO-induced N_2 ligand scission and functionalization at N_2 -M complexes have been developed, in which all of the N_2 ligands adopt side-on bridging coordination mode.

Following their earlier work on N−C bond formation from N2-Zr and Hf complexes, Chirik *et al*. reported in 2010 the first example that treatment of the *ansa*-zirconocene and hafnocene N_2 complexes **60** and **63** with 4 atm or 1 atm of CO leads to the generation of the dinuclear oxamidide complexes **64** and **65** as two isomers [\[46](#page-18-32)[,47\]](#page-18-33). Besides, when **63** is treated with less CO (1.5 equiv), a new product of imido-bridged dihafnium complex **66** could be isolated, in which the *H*-atom on the bridging imido is derived from the cyclometallation of the *^t* Bu group (Scheme [14a](#page-8-0)). Protonolysis enables these products to release the corresponding *N*-containing organic compounds of free oxamide and isocyanic acid.

Density functional theory (DFT) calculations [\[48\]](#page-18-34) and experimental results $\left[49\right]$ reveal that the formation of **64** and **65** is assumed to be initiated by CO insertion into an Hf−N bond and followed by the retro $[2+2]$ cycloaddition to provide the presumptive μ -nitride species 67. The coordination and insertion of CO to the μ -nitrido intermediate 67 results in the formation of **68**, which was characterized by multinuclear nuclear magnetic resonance (NMR) spectroscopy at low temperature. **68** undergoes C−C bond formation via coupling of the terminal and bridging isocyanate units along with the loss of the terminal carbonyl ligand to give the final products (Scheme [14b](#page-8-0)).

More studies indicate that these CO-induced N2 cleavage and functionalization reactions are also compatible with other zirconocene and hafnocene N2 complexes. Therefore, a tetrametallic hafnocene oxamidide complex **71** could be obtained via a dimeric hafnium intermediate 70 when the N_2 -Hf complex **69** is treated with CO [\[50\]](#page-18-36). The transformations of these oxamidide complexes were also elaborated. Thermolysis of **71** at 110◦C provides a μ -oxo hafnocene complex 72 with both terminal cyanide and isocyanate ligands that undergoes preferential group transfer of the cyanide unit to liberate organonitriles of TMSCN or MeCN along with the generation of **73** by reacting with TMSI or MeOTf (Scheme [15a](#page-9-0)) [\[50\]](#page-18-36). Oxamidide complex 64 reacts with CO_2 and ^tBuNCO to give the formal [2+2] cycloaddition products **⁷⁴** and **75** [\[51\]](#page-19-0) (Scheme [15b](#page-9-0)). Additionally, various free *N*,*N* -dialkyloxamide could be formed via stepwise *N*-alkylation of the oxamidide complex **76** and following protonolysis with HCl or ethanol (Scheme [15c](#page-9-0)).

The characterizatio[n](#page-9-1) and reactivity studies of the μ -nitride intermediates were also developed (Scheme [16\)](#page-9-1). Rapid bubbling of CO into N2-Hf complex **69** at a low temperature produces a metastable dihafnocene nitride complex **79**, which is characterized by IR and multinuclear NMR spectroscopy. This base-free μ -nitride can react with various substrates $[52-55]$ $[52-55]$. For instance, the treatment of **79** with TMSI affords silylureate complex **80**. This reaction is involved in the initial iodide ion abstraction to give a transient silyl cation and a formally anionic bridging nitride intermediate, whose nucleophilicity is increased by weakening the Hf−N multiple bonding. Hence, this intermediate could undergo nucleophilic attacking of the nitride group to the terminal isocyanate moiety to form the ureate core, which is then trapped by the silyl cation to yield **80**. Besides, when **79** is treated with cyclooctyne, monosubstituted allenes and isocyanates, the formal $[2+2]$ cycloaddition reactions occur to

Scheme 15. N–C bond formation of CO- and N₂-derived oxamidide complexes. (a) Thermolysis of oxamidide complex **71**. (b) The reaction of oxamidide complex 64 with $CO₂$ and *t* BuNCO. (c) *N*-alkylation of the oxamidide complex **76** to afford *N*,*N* -dialkyloxamides.

afford **81**, **84** and **85**, respectively. The alkyne and isocyanates products are kinetically unstable at elevated temperature and engage in additional ^N−C bond formations to give **⁸²** and **⁸⁶**. Complex **82** could be converted to a binuclear complex **83** with two bridging carbodiimidyl ligands by reacting with TMSCl. The μ -oxo complex 86 can liberate *N*-containing organic compound of carbodiimide along with the generation of dihafnium oxo complex **73** by reacting with MeOTf. In contrast, exposure of the nitride complex **79** to another heterocummulene of $CO₂$ provides μ -oxo bis(isocyanate) complex 87, resulting from deoxygenation of $CO₂$ accompanied by N−C bond formation. Furthermore, the Hf−nitride bond in **⁷⁹** also engages in the insertion of cyclohexylnitrile (CyCN) to provide **88**, which can continue reacting with another molecule of CyCN to afford bridging ureate-type complex **⁸⁹** via additional N−C bond formation $(Scheme 16)$ $(Scheme 16)$.

The other system of CO-induced N_2 scission and functionalization was discovered by Mazzanti *et al*. using uranium complexes. A side-on-bridged binuclear N_2 -U complex 90 with μ -nitride ligands reacts with CO to provide the oxo/cyanate diuranium complex **91** accompanied by releasing of potassium cyanate (KCN), which is formed from the reaction of nitride unit with CO [\[56\]](#page-19-3) (Scheme [17a](#page-10-0)). To understand the role of the bridging nitride in these transformations, a similar N_2 -U complex 92 with bridged μ -oxo ligand was synthesized and its reactivity toward CO was also investigated $\lceil 57 \rceil$. The reaction between **92** and CO immediately results in the generation of cyanamido bridged complex 93 with retaining the μ -oxo moiety via both cleavages of N−N single bond and C≡O triple bond (Scheme [17b](#page-10-0)). DFT calculation indicates that the different reactivity of **90** and **92** is attributed to the different bonding nature of the N_2 ligands, in which the μ -nitride is involved in the binding and resultant activation of N_2 , but the μ -oxo is not.

Scheme 16. N–C bond formation of CO- and N₂-derived Hf-nitride complexes.

Scheme 17. CO-induced N_2 scission and functionalization at side-on N2-U complexes. (a) The reaction between CO and N_2 -U complex **90** with μ -nitride ligand. (b) The reaction between CO and N₂-U complex **92** with μ -oxo ligand.

Scheme 18. *N*-alkylation of side-on-end-on N₂-Ta complex by BnBr.

N−**C BOND FORMATION VIA SIDE-ON-END-ON N2-M COMPLEXES**

The side-on-end-on bound mode is much less common relative to the aforementioned two coordination modes in N_2 -M complexes. All of the work regarding the making of N–C bond from N_2 -M complex with this bonding mode were finished by Fryzuk *et al.* by employing a binuclear N_2 -Ta complex[.](#page-9-1)

*N***-alkylation**

In 2001, the *N*-alkylation of the side-on-end-on bridging binuclear N2-Ta complex **94**was developed to afford *N*-benzylation product **95** in high yield by reaction with benzyl bromide (BnBr) (Scheme [18\)](#page-10-1) [\[58\]](#page-19-5). This reaction was similar to the *N*-alkylation reaction of the side-on N_2 -Zr complexes 44 (Scheme [10a](#page-6-0)).

Cycloaddition and insertion

Besides, this side-on-end-on N_2 -Ta complex 94 was also reported to undergo $[2+2]$ cycloaddition reaction by treating with heteroatom 1,2-cumulenes (Scheme [19a](#page-10-2)) [\[59\]](#page-19-6). For example, the reaction between **94** and *N*,*N* -diphenyl carbodiimide results in the formation of **96**. However, when carbon disulfite or isothiocyanates are added, the *N*-functionalization product **97** is generated

Scheme 19. N−C bond formation from the cycloaddition reactions of side-on-end-on $N₂$ -Ta complexes with heteroatom 1,2-cumulenes. (a) The reaction of $N₂$ -Ta complex **94** with carbodiimide, carbon disulfite and isothiocyanates. (b) A plausible mechanism for the generation of **97** and **98**.

concomitant with the N−N bond scission. In the case of *tert*-butyl isothiocyanate (*^t* BuNCS), the generated intermediate **98** would further undergo N−Si bond formation at elevated temperature to give **99**. As depicted in Scheme [19b](#page-10-2), the formation of **97** and **98** can be rationalized bythe following mechanism. The initial $[2+2]$ cycloaddition reactions between 94 and the $C = S$ bond of the substrates give intermediate **E**, followed by reductive elimination of H2 to provide a transient intermediate **F** that contains a Ta−Ta bond. The Ta−Ta bond in **^F** would trigger the N−N bond cleavage to afford the final products.

N−**C BOND FORMATION VIA METAL NITRIDES**

The complete reduction of N_2 -M complexes might cleave the N−N bond of the N_2 ligands to give metal nitrides. In the most terminal metal nitrides, the strong metal−nitrogen bonds result in these nitrides often exhibiting weak nucleophilicity and just reacting with high-energy species such as alkyl triflates and acyl chlorides to assemble N−C bond. However, some bridging nitrides derived from N_2 can also react with other carbon-based substrates, like MeI and CO, to form N−C bond.

Scheme 20. *N*-methylation of Nb-nitride by reaction with MeI.

Scheme 21. *N*-alkylation of Mo-nitride by reaction with MeI.

*N***-alkylation**

In 2007, Kawaguchi *et al*. reported the reaction between MeI and $bis(\mu\text{-nitrido})$ diniobium complex **101**, which is prepared from the tetra(μ -hydride) diniobium precursor **10[0](#page-10-2)** (Scheme [20\)](#page-11-0) [\[60\]](#page-19-7). Stepwise methylation of **101** by MeI yields mono-imido **102** and bi-imido **103**, the latter of which could also react with excess pyridine to give terminal imido **104**, which reacts with $CO₂$ to generate **105** and **106** through further N−C bond formation. A plausible mechanism for this process was raised by the authors. A $[2+2]$ cycloaddition of 104 with $CO₂$ followed by extrusion of methyl isocyanate (MeNCO) results in the formation of a terminal oxo species that dimerizes to give 106. Meanwhile, the generated MeNCO would also undergo $[2+2]$ cycloaddition with another molecule of **104** to form **105** [\[61\]](#page-19-8).

Cummins *et al*. found that the terminal molybdenum nitride **108** synthesized from the three coordination Mo(III) complex **107**, undergoes *N*-alkylation by reacting with MeI to provide **109** (Scheme [21\)](#page-11-1) [\[62\]](#page-19-9)[.](#page-11-1)

Another example of making N−C bond from metal nitrides was fulfilled by Schneider *et al*. in 2016. The reactions between ROTf ($R = Me$, Et and Bn) and a terminal rhenium nitride **111**, which is prepared from the reduction of the dichloride precursor 110 with sodium amalgam or CoCp^{*}₂, give the *N*-alkylation complexes **112** (Scheme [22\)](#page-11-2) [\[63–](#page-19-10)[65\]](#page-19-11). Further studies suggest that the nitriles (RCN) can be liberated by deprotonation and oxidation of **112** viathe ketimido intermediates **113**, accompanied by the generation of trichloride complex **114**, which could also be reduced by sodium amal-

Scheme 22. Re-promoted conversion of N_2 into nitriles via *N*-alkylation of Re-nitride.

Scheme 23. *N*-methylation of Fe-nitride by reaction with MeOTs.

gam to afford the starting material **111**. According to these results, a full synthetic cycle for synthesis of nitriles from N_2 with moderate isolated yields was established (Scheme [22\)](#page-11-2).

Because the group $8-10$ N₂-derived nitrides are rare, the *N*-functionalization of these nitrides is hardly observed. One exception was reported by Holland *et al.*, who employed an unprecedented trinuclear iron nitride **116** to achieve this transformation. This nitride complex **116**, obtained from the reduction of the chloride precursor **115** with precisely equivalent KC_8 under N_2 atmosphere, can react with MeOTs and 18-crown-6 (18-C-6) to give the methylimido complex **118** via a presumptive two-coordinate nitride **117** with higher reactivity $(Scheme 23) [66].$ $(Scheme 23) [66].$ $(Scheme 23) [66].$ $(Scheme 23) [66].$ $(Scheme 23) [66].$

*N***-acylation**

The *N*-acylation of N_2 -derived nitride has also been investigated. For example, the *N*-acylation

Scheme 25. Titanium-promoted direct conversion of N_2 into nitriles via the reaction between acyl chlorides and Ti-nitride.

Scheme 24. Conversion of N₂ into nitriles via *N*-acylation of Mo-nitride.

products **119** are obtained when **108** is treated with acyl chlorides in the presence of additives, such as [TMS(py)][OTf] and *ⁱ* Pr3SiOTf. Furthermore, when the *N*-acylated products **119** reacts with magnesium anthracene $(MgC_{14}H_{10})$ and trimethylsilyl triflate (TMSOTf) in one pot, it would be converted to the trimethylsiloxy-substituted ketimide **121** via the intermediates of **120**. Further reaction of **121** with $SnCl₂$ or $ZnCl₂$ affords the corresponding organic nitriles commitment with the generation of molybdenum chloride complex **122**, a precursor of the trisamide molybdenum complex **107**. In consequence, an efficient synthetic cycle that can directly convert N_2 to nitrile was accomplished (Scheme [24\)](#page-12-0) [\[67\]](#page-19-13).

*N***-acylation/elimination**

In addition to simple *N*-acylated products, the reactions between N_2 -derived metal nitrides and acyl chlorides also afford nitriles proceeded through *N*-acylation and subsequent elimination in formal. Hou *et al*. discovered that the reaction of titanium trialkyl complex 123 with N_2 and H_2 results in a novel diimide/tetrahydride complex **124**. This complex can react with N_2 at elevated temperature to provide a tetranuclear diimide/dinitride complex **125** that can further react with a series of acyl chloride to afford the corresponding nitriles in high yield (Scheme [25\)](#page-12-1) [\[68\]](#page-19-14). Based on the experimental and computational results, the authors think that the functionalization of the imide ligands is prior to the

Scheme 26. Niobium-promoted direct conversion of N₂ into nitriles via the reaction between acyl chlorides and Nb-nitride.

nitride groups in these reactions. Furthermore, by treatment of the crude reaction mixture with HCl, the titanium trichloride complex **127** is isolated, which could be easily converted to **123** by reacting with TMSCH₂Li. Hence, a synthetic cycle of titanium-promoted synthesis of nitriles direct from N2 was proposed (Scheme [25\)](#page-12-1).

Another synthetic cycle for providing organic nitriles from N2 was developed by Cummins *et al*. via a niobium nitride intermediat[e](#page-10-2) (Scheme [26\)](#page-12-2) [\[69\]](#page-19-15). An end-on bridging heterodinuclear N_2 -M complex **129**, prepared from the reaction of the niobium triflate complex 128 and the aforementioned N_2 -Mo

Scheme 27. Vanadium-promoted direct conversion of N_2 into potassium cyanate via the reaction between CO and $N₂$ -V complex.

complex 1, could undergo N_2 ligand scission to form anionic niobium nitride **130** along with the formation of molybdenum nitride **108**, when **129** istreated with sodium amalgam. Treatment of **130** with acyl chloride results in releasing of nitriles accompanied by the generation of the niobium oxo complex **131**. By treating with triflic anhydride, this oxo complex **131** can be converted to a bistriflate complex **132** that could be reduced to the initial compound **128** to finish the cycle.

Redox-coupled *N***-atom transfer**

In comparison with *N*-alkylation or acylation and subsequent reduction, a more efficient route to transfer the nitride into organic compounds is the transformation of nitride-*N* atom into an incoming substrate with concurrent metal reduction. In 2014, Kawaguchi *et al.* reported the redox-coupled *N*-atom transformation of a V-nitrid[e](#page-12-0) (Scheme [27\)](#page-13-0) [\[70\]](#page-19-16). Reduction of the V(III) complex **133** by KH under N_2 results in a split of the N_2 to provide the μ -nitride V(IV) complex 134, which could be oxidized to V(V) nitride compound **135** via reacting with *p*-benzoquinone. When **135** is treated with CO or 2,6-xylylisocyanide in the presence of [2.2.2] cryptand, the *N*-atom transformation of the substrates is observed concomitant with the formation of **136**. The contact-ion-pair complex **137** could also be isolated from the reaction of **135** and CO. Although the extrusion of the cyanate or carbodiimide ligand in **136** are not easy, the contact-ion-pair **137** readily undergoes ligand exchange with 2-butyne to liberate potassium cyanate (KNCO) with the

Scheme 28. N–C bond formation via the reaction of silylimido complexes with $CO₂$.

formation of the alkyne adduct **138**. Additionally, **138** is facilely converted into the starting complex **133** upon dissolving in THF. Hence, a synthetic cycle for direct conversion of N_2 and CO into KNCO was completed. However, achieving the catalytic process of this synthetic cycle remains elusive due to the incompatibility of the individual steps in this cycle, such as the requirement of solvents in the N−C bond formation step and KNCO releasing step being different.

*N***-silylation/imido transfer**

Besides *N*-hydrogenated intermediates, N₂-derived *N*-silylated complexes are also good precursors to make N−C bond. For instance, a cycle of Moand W-promoted synthesis of isocyanates from N_2 via a silyl-imido intermediate was established (Scheme 28) [\[71\]](#page-19-17). Photolysis of the end-on bridging N2-Mo, W complexes **140** leads to the generation of nitride intermediates via N−N bond cleavage (*vide infra*), which would be trapped *in situ* by TMSCl to afford silyl-imido complexes **141**. When TMSCl is replaced by Ph₃SiCl, Me₃CCl or Me₃GeCl, the similar reaction could also take place to provide the corresponding imido complexes. Besides, the organic compound TMSNCO could be obtained concomitant with the formation of the mono-nuclear oxo complexes 142 by treatment of 141 with $CO₂$. These oxo complexes **142** are known to react with additional TMSCl to regenerate the dichloride complexes 139 that are the precursors of the N_2 -M complexes **140**[.](#page-11-1)

Metal-ligand cooperative *N***-atom transfer**

Metal-ligand cooperative *N*-atom transfer is also an efficient strategy because of the avoiding of extra protons and electrons. Recently, a

Scheme 29. Metal-ligand cooperative *N*-atom transfer of a Re-nitride.

metal-ligand cooperative *N*-atom transfer of a Re-nitride was reported by a cooperative *2 H*+*/2 e*[−] transfer of the pincer ligan[d](#page-12-2) (Scheme 29) [\[72\]](#page-19-18). This Re-nitride complex **144**, which is generated from photo-promoted cleaving of the end-on bridging binuclear N2-Re complex **143** (*vide infra*), could react with benzoyl chloride to afford benzamide $(PhCONH₂)$, benzonitrile $(PhCN)$ and benzoic acid (PhCOOH) along with the formation of trichloride Re complexes **145**, in which the pincer ligand is oxidized to an imine-type ligand. The producing of PhCN and PhCOOH is caused from the reaction of the initially formed benzamide with excess benzoylchloride in the crude.

Involvement of photochemistry

Besides photo-promoted radical generation, another pathway of the photochemistry participating in N_2 transformation is the direct photolytic splitting of N_2 ligands into nitrides, which could further engage in *N*-atom transfer. Two examples of this method have been reported. In the first example, irradiation of the above-mentioned end-on bridging N2-Mo, W complexes **140** over several days by medium-pressure Hg lamps leads to the generation of two metal nitrides **146** and **147** (Scheme [30a](#page-14-1)). Furthermore, when **140** are photolyzed in the presence of excess TMSCl, the terminal silylimido complexes **141** are obtained in moderate yield with the formation of dichloride complexes **139** (Scheme [29\)](#page-14-0) [\[71\]](#page-19-17). The other example involves the photolysis of an end-on-bridged N_2 -Re complex 143 that has abnormal thermal-stability, to provide the aforementioned Re-nitride **144** (Scheme [30b](#page-14-1)) [\[72\]](#page-19-18). It is noteworthy that the photo source of this reaction could be $Xe(Hg)$ lamp ($\lambda > 305$ nm) or a 390 nm LED lamp[.](#page-12-0)

Involvement of electrochemistry

Electrochemical N_2 reduction is an alternative to chemical N_2 reduction for the synthesis of N_2 -M complexes. This approach has been utilized to regain the N_2 -Re complex 143 to achieve a cycle (Scheme [31\)](#page-14-2) [\[72\]](#page-19-18). Schneider *et al.* found that in the controlled potential electrolysis experiment, the

Scheme 30. Photolytic cleavage of end-on bridging N₂-Mo, W and Re complexes into nitrides. (a) Photolytic cleavage of end-on bridging N2-Mo, W complexes **140**. (b) Photolytic cleavage of end-on bridging N2-Re complexes **143**.

Scheme 31. Electrochemical reduction involved synthetic cycle of direct conversion of $N₂$ into benzamide and benzonitrile.

trichloride Re complexes **145** formed from the reaction of Re-nitride with PhCOCl, could be converted to the N_2 -Re complex 143 via electrolyzing at $E = -1.65$ V for 8 h in the presence of proton source of 2,6-dichlorophenol (DCP) and subsequently electrolyzing at $E = -1.85$ V for 5 h under N_2 . Thus, a three-step cycle for the synthesis of $PhCONH_2/PhCN$ from N_2 was established, in which the creative approaches of metal-ligand cooperation and photo- and electrochemistry were all used.

N−**C BOND FORMATION VIA UNCHARACTERIZED N2-M INTERMEDIATES**

Compared to the above works, the earlier reports about the conversion of N_2 into organic compounds were achieved by one-pot reactions of ill-defined

 $XTI = N TMS$

 $\sqrt{5}$ Ti $-M$ (TMS)

 $N₂$ (1 atm)

TMSC

 N_2 -complexes or their derivatives with carbonbased substrates and followed by hydrolysis. In this section, some examples of this method were introduced briefly.

The initial works of transition metal promoted direct conversion of N_2 into organic compounds were reported more than 50 years ago, when Vol'pin and Shur*et al*. developed two systems for transformation of N_2 into aromatic amines mediated by titanium spe[c](#page-13-0)ies $[14,73]$ $[14,73]$. In the first system, several aromatic amines are obtained when Cp_2TiCl_2 is treated with excess of aryllithium (aryl = Ph, m - and p -MeC₆H₄) reagents under N_2 pressure of 80-100 atm and followed by hydrolysis (Scheme [32a](#page-15-0)). When the aryllithium in these reactions is replaced by alkyllithium reagents, the corresponding alkylamines could not be obtained. The other system, which also gives arylamines by subsequent hydrolysis, involves the reaction of diaryltitanocenes Cp_2TiAr_2 (Ar = Ph, m - and p -Me C_6H_4) with alkali or alkaline metal (Li, Na and Mg) and N_2 (100 atm) (Scheme [32b](#page-15-0)). Although plenty of effort has been made, the detailed mechanisms of these works are yet unclear.

Meanwhile, a related work was reported in 1970 by van Tamelen and Rudler, who succeeded in the synthesis of organic amines from the reaction between ketones and a supposed titanium nitride species prepared through the reaction of Cp_2TiCl_2 with magnesium under N_2 (Scheme [32c](#page-15-0)) [\[74\]](#page-19-20).

Additionally, Mori *et al*. achieved incorporation of N2 into organic compounds via the *N*-silylation titanium complexes **148**, which were prepared from the one-pot reaction of titanium species (TiCl₄ or $Ti(O^{i}Pr)_{4}$), Li and TMSCl under N₂ or dry air (1 atm) [\[75\]](#page-19-21). Although the precise components and structures of **148** have not been determined so far, they are considered to contain $XTi = NTMS$, X_2 TiN(TMS)₂ (X = Cl, O^{*i*}Pr) and N(TMS)₃

(a)

TiX.

Τi

Scheme 33. Ti-promoted N−C bond formation via illdefined *N*-silylation titanium species. (a) Preparation of *N*-silylation titanium complexes **148** from one-pot reaction of TiCl₄ or Ti(O^{*i*}Pr)₄ with Li and TMSCI under N₂ or dry air. (b) The reaction between **148** and keto-carbonyl compounds to afford nitric heterocycles. (c) Palladium-catalyzed synthesis of aryl- or allyl- amines and amide derivatives from **148** and aryl or allyl halides in the absence or the presence of CO.

[\[76–](#page-19-22)[78\]](#page-19-23). **148** could serve as a nitrogenation reagent to react with a series of keto-carbonyl compounds to provide kinds of nitric heterocycles, such as indole, quinoline, pyrrole, pyrrolizine, lactams and indolizine derivatives. Besides, when **148** is treated with palladium complexes, the transmetalation of *N*-atom from **148** to the palladium center occurs. Hence, palladium-catalyzed synthesis of aryl- or allyl- amines and amide derivatives from aryl or allyl halides and **148** could be fulfilled, inthe absence or the presence of C[O](#page-12-2) (Scheme [33\)](#page-15-1).

CONCLUSION AND OUTLOOK

Direct transformation of N_2 into *N*-containing organic compounds is of fundamental and practical significance. In the past 60 years, the area of direct incorporation of N2 into N−C bond effectuated

many great achievements. Relative to the traditional methods of assembling N−C bond via *^N*-alkylation of N_2 -M complexes, more atom-efficient approaches, such as cycloaddition, insertion and redox-coupled *^N*-atom transfer for making N−^C bond have been developed and received more attention in recent years. By the delicate design, some synthetic cycles about direct conversion of N_2 into organic compounds have also been developed. In these cycles[,](#page-14-1) photo- and electrochemistry are sometimes used to prepare the N2-M complexes, cleave the N−N bond or release the final products. However, all of these reactions are stoichiometric and the catalytic system for the direct introduction of N_2 into organic compounds has not been realized yet. The main factors that prevent these complete synthetic cycles from becoming a catalytic process are the rigorous reaction conditions of the N−C bond formation and *N*-containing organic compounds releasing steps in these cycles, which are incompatible with the preparation steps for N_2 -M complexes. Hence, developing milder systems are imperative. Besides, new reaction types also need to be explored. In this context, we think the following fields can be considered in the future.

New reaction systems

So far, most N−C bond formation occurs at N2-M complexes of group 4–6 transition metals. Exploring other metal promoted N−C bond formation is an attractive topic. Besides, the multi-metal synergistically promoted N_2 activation and functionalization also need to be studied. Toward this end, the design of new types of ligands should be considered.

New reaction types

Reductive elimination, an essential step in catalytic amination reactions, has not been found to take place at N_2 -derived *N*-containing transition metal complexes. This process should be explored in future because it provides an efficient approach to assemble N−C bond concomitant with regaining the N_2 -M complexes or their precursors. Additionally, other intriguing reaction modes, such as the $[4+2]$ cycloadditions of N₂ ligands and insertion of N2 into metal**−**carbon bond should also be investigated.

Polynuclear metal species cooperative N2 scission and functionalization

Stimulated by the previous works on multinuclear Ti, Fe complexes-promoted N_2 cleavage and N−C bond formation $[66,68]$ $[66,68]$, the strategy to realize synergistic N_2 -splitting and subsequent functionalization using polynuclear metal complexes should be further explored. Additionally, recent reports on gasphase polynuclear metal clusters-mediated N_2 scission and subsequent N−C bond formation deserve further attention [\[79](#page-19-24)[,80\]](#page-19-25).

Main group elements promoted N−**C bond formation**

The recent report about N_2 reduction by borylenes from Braunschweig *et al*. suggests the potential of boron mediated formation of N−C bond from N2 [\[81\]](#page-19-26). Besides, some calculation results indicate that the direct reactions of boron or carbene with N_2 are also permitted in some cases. For example, Li and Schaefer *et al.* designed a new molecular system for nitrogen reduction, involving a 2,3 -bipyridineanchored, end-on-bridging dinitrogen complex of the $Me₂B-BMe₂$ intermediate by theoretical calculations [\[82\]](#page-19-27). Zhu *et al.* designed a metal-free dinitrogen activation system based on the boron and NHC carbene system $\lceil 83 \rceil$. These results offer inspiration for future work on p-block elements promoting or catalytic conversion of N_2 into organic compounds.

Analogue of PCET: lessons from N2-to-NH3 catalysis system

Very recently, Nishibayashi *et al*. achieved a remarkable N_2 -to-NH₃ process via a molybdenum-catalysis system. By using the samarium diiodide $(Sml₂)$ as the reductant and alcohol or water as the proton sources, the total turnover number (TON) of this reaction reaches up to 4350 with 91% yield of NH3. Further studies reveal that a proton-coupled electron-transfer (PCET) process, in which O−H bonds in water or alcohols are weakened by coordination to $SmI₂$, is the key to this high reactivity [\[84\]](#page-19-29). Inspired by this, a similar process of C−X $(X = 0, Cl, Br, I)$ bonds coordinated to a relevant reductant to weaken the C−X bonds, which could be named as carbocation-coupled electron-transfer (CCET) should also be studied.This CCET process may offer a path forward for developing catalysis systems that incorporate N_2 into amines via successive alkylation of a N_2 -derived nitride[.](#page-14-1)

Photochemistry

Limited complexes capable of N_2 photoactivation are currently known, and the underlying photophysical and photochemical processes after light absorption are largely unresolved. Light can induce the

split of N \equiv N bond in the M-N₂ complexes and the resulting nitride complexes are typically reactive. Hence, the following N−C formation reaction should be possible. Besides, the excitation into $N-N$ π^* orbitals is also possible, which can lead to a weakened π -bond, and hence a following N–C formation directly from $M-N_2^*$ and carbon-based substrates including CO , $CO₂$ might be possible.

Electrochemistry

Previous work indicates that electrochemical reduction could release final organic compounds along with regeneration of N_2 -M complexes. This stimulates us to create an electrochemical reduction system, in which the electrolyzation step is compatible with the N−C bond formation step. Besides, the recent report of N_2 and CO_2 coupling to produce urea, which was conducted by an electrocatalyst consisting of PdCu alloy nanoparticles on $TiO₂$ nanosheets, suggests that designing a new solid catalyst to incorporate N_2 into high-value *N*-containing product beyond NH₃ should also be attractive $\lceil 85 \rceil$ [.](#page-14-2)

Heterogeneous catalysis systems

Although the industrial Haber-Bosch process produces NH₃ over the surface of heterogeneous solid-state catalysts, a similar process, in which a heterogeneous catalyst catalyzes or promotes direct transformation of N_2 into organic compounds, has not been reported in literature. Development of the new systems, where the merits of the homogeneous molecular systems and the heterogeneous systems are rationally combined, is a promising approach toward the goal.

Via *N***-protonation or silylation intermediates**

The strategy of conversion of N_2 into organic compounds via the *N*-silylation and *N*-protonation complexes can be further extended. As an example, converting the less active $M-N_2$ complexes into M-N−Si/H species *in situ* followed by catalytic reaction with carbon-based substrates, might result in various valuable organic compounds being synthesized.

In a word, with combined efforts from crossdisciplines the dream of direct catalytic and efficient conversion of N_2 into *N*-containing organic compounds under mild conditions is believed to be attainable in the future.

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