ORGANIC CHEMISTRY

FRONTIERS

RESEARCH ARTICLE

Check for updates

Cite this: Org. Chem. Front., 2024, **11**, 5720

Nickel/ligand loading-controlled divergent and selective coupling between redox-active methylenecyclopropanes and ArBr†

Ben Mao,^a Xiao-Yu Zhang,^a Xiu-Fen Bi,^c Xiao-Yin Yang,*^c Min Shi^b*^{a,b} and Yin Wei^b*^b

Divergent and selective synthesis has been widely achieved in transition-metal-catalyzed reactions through a ligand property tuning strategy. However, ligand-loading-controlled divergent synthesis has rarely been reported. Due to changes in ligand loading, different metal complexes should be formed and exhibit diverse catalytic properties. Herein, we disclose a Ni/photoredox-catalyzed divergent and selective coupling between redox-active methylenecyclopropanes and aryl bromides through ligand loading adjustment, providing ranges of alkyne derivatives and dibenzylethylene derivatives. Two different catalytic cycles are proposed to demonstrate the generation of two sets of products, where homopropargyl radicals and nickelacy-clobutane species should be crucial intermediates in the corresponding catalytic cycle, respectively.

Received 22nd July 2024, Accepted 24th August 2024 DOI: 10.1039/d4qo01346h

rsc.li/frontiers-organic

Introduction

A ligand-controlled divergent synthetic strategy has been widely explored in the field of transition-metal catalysis over the past few decades.¹ By tuning the electronic properties and steric hindrance of ligands, the metal–ligand complex as the ligated catalyst can exhibit different catalytic activities and site selectivities.² Hence, broader transformations of the same starting materials can be achieved under almost identical conditions, offering facile access to diverse synthetic applications. Nevertheless, this strategy requires chemists to invest more effort in the process of ligand screening and rational ligand design. Typically, the choice of ligands plays a crucial role in transition-metal-catalyzed reactions, but the ligand loading often does not receive careful attention in many reported protocols. In most cases, the dosage ratios of the ligand to metal are in the range of 1 to 2, and the specific ratios are not strictly controlled, only ensuring that the ligand is more than the metal to form the desired complex. Simple changes of ligand loading may lead to different metal-ligand complexes forming in the reaction system, providing metal centers with different catalytic activities and site selectivities. In particular, when the ligand loading is less than the metal loading, the free metal and ligand-coordinated metal in the system may show different properties, leading to some unexpected transformations of the substrates. Alternatively, when the ligand is far in excess, the multiple coordination of the ligand with the metal would also result in a catalytic center different from the single-coordinated metal-ligand complex. Compared with the differentiation of ligand selection, this scheme can also provide a potential path for divergent synthesis in transition-metal catalytic transformations (Scheme 1a).

In divergent and selective synthetic protocols, different catalytic transformations can be realized by changing the reaction conditions,³ which in turn requires the starting materials to have multiple reactivities to accomplish the corresponding conversions. Hence, chemists usually prefer substrates with multiple reactive sites since the rational design and selection of substrate structures are essential for achieving divergent synthesis.⁴ Methylenecyclopropanes (MCPs), a class of three-membered ring substrates with a strained exocyclic double bond, have been comprehensively explored as versatile C-3 building blocks since the 1970s.⁵ The highly strained structure with multiple active sites makes MCPs excellent candidates in the field of transition-metal-catalyzed carbon-carbon activation.⁶ Among them, ligand-controlled divergent transformations of MCPs have also been explored. For example, in



View Article Online

View Journal | View Issue

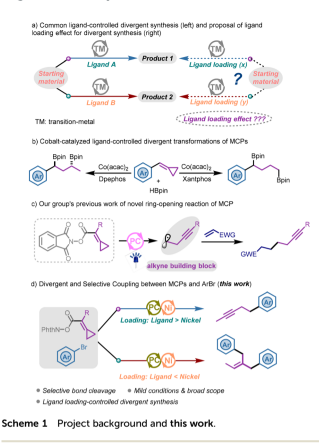
^aKey Laboratory for Advanced Materials and Institute of Fine Chemicals, School of Chemistry & Molecular Engineering, East China University of Science and Technology, 130 Meilong Road, Shanghai 200237, China.

E-mail: mshi@mail.sioc.ac.cn

^bState Key Laboratory of Organometallic Chemistry, Center for Excellence in Molecular Synthesis, Shanghai Institute of Organic Chemistry, University of Chinese Academy of Sciences, Chinese Academy of Sciences, 345 Lingling Road, Shanghai 200032, China. E-mail: weiyin@sioc.ac.cn

^cHuafon Chemical Co. Ltd., 1688 Kaifaqu Road, Ruian Economic Development Zone, Zhejiang 325200, China. E-mail: yang.xiaoyin@huafeng.com

[†]Electronic supplementary information (ESI) available: Experimental procedures and characterization data of new compounds. See DOI: https://doi.org/10.1039/ d4q001346h



2023, Ge's group reported a cobalt-catalyzed ring-opening reaction of MCPs to obtain two sets of skipped diboronates through a ligand-tuning strategy (Scheme 1b).⁷

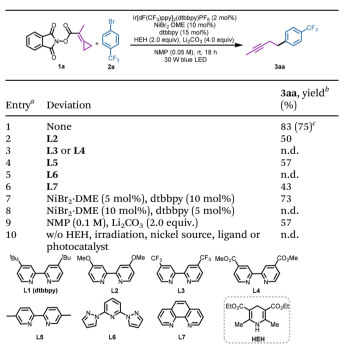
Our group has long been investigating applicable conversions of MCPs through careful substrate design and mild condition selection.8 In 2021, we successfully demonstrated that redox-active MCPs could be applied as alkyne building blocks through a facile radical rearrangement process (Scheme 1c).9 Under mild photocatalytic conditions, the MCP substrate tethered with N-(acyloxy)phthalimide (NHPI ester), a redox-active ester used as an efficient radical precursor,¹⁰ could afford a homopropargyl radical adding to the alkene in the protocol. To broaden the applications of these valuable building blocks, we envisioned that such substrates may be applied to nickel-catalyzed coupling reactions to develop potential transformations. Herein, we wish to report a ligand loading-controlled photoredox/nickel catalyzed divergent and selective coupling between redox-active MCPs and ArBr, delivering a series of alkyne derivatives and dibenzylethylene derivatives (Scheme 1d). Notably, the alkyne product was afforded when the loading of the bidentate ligand exceeded the nickel catalyst, while the dibenzylethylene product was obtained when the nickel catalyst was excessive.

Results and discussion

We commenced our studies by utilizing NHPI ester **1a** (1.0 equiv.) with ArBr **2a** (1.0 equiv.) as the model substrates. In the

presence of $Ir[dF(CF_3)ppy]_2(dtbbpy)PF_6$ (2 mol%), NiBr₂·DME (10 mol%), 4,4'-di-tert-butyl-2,2'-bipyridine (L1, 15 mol%), Hantzsch ester (HEH, 2.0 equiv.), and Li₂CO₃ (4.0 equiv.) in NMP as the solvent under 30 W blue LED irradiation, the desired alkyne product 3aa could be obtained in 83% yield (Table 1, entry 1) (see ESI Tables S1-S9[†] for more details). Apart from ligand L1, other substituted 2,2'-bipyridine ligands were also screened in the reaction but did not offer better results (Table 1, entries 2-4). Notably, substituted 2,2'-bipyridine ligands containing electron-deficient groups (CF₃, CO_2Me) (Table 1, entry 3) were ineffective in this nickel-catalyzed coupling reaction. The polydentate ligand L6 showed no reactivity (Table 1, entry 5), while ligand 1,10-phenanthroline (L7) provided the alkyne product in 43% yield (Table 1, entry 6). Reducing the loading of the nickel source and ligand resulted in a slight decrease in the yield of 3aa (Table 1, entry 7). Increasing the concentration of the system and reducing the amount of base (Table 1, entry 9) would lower the yield of 3aa (see ESI Table S5[†] for more details). Control experiments revealed that HEH, light irradiation, nickel source, ligand and photocatalyst were indispensable for the generation of the alkyne product (Table 1, entry 10). It should be noted that in the process of nickel catalyst and ligand loading optimization, the excessive addition of a nickel salt over the ligand led to the disappearance of alkyne product 3aa (Table 1, entry 8) surprisingly, but gave a different coupling product 4aa (see ESI Table S9[†] for more details).

Table 1 Optimized conditions for the formation of 3aa



^{*a*} Reactions were performed on a 0.1 mmol scale. ^{*b*} Yields were determined by analyzing ¹H NMR spectroscopic data using 1,3,5-trimethoxybenzene as an internal standard. ^{*c*} Isolated yield on a 0.2 mmol scale.

View Article Online
Organic Chemistry Frontiers

Next, condition screening for the double aryl group coupling product 4aa was also conducted (Table 2). To better compare the differences in catalyst loading for producing two distinct products, we still utilized NHPI ester 1a (1.0 equiv.) with ArBr 2a (1.0 equiv.) as the model substrates under approximate reaction conditions. Product 4aa could be obtained in 81% yield (Table 2, entry 1) only by changing the loading of NiBr₂·DME (15 mol%) and L1 (7.5 mol%) based on the optimized conditions for 3aa. Photocatalysts such as 4CzIPN, Ir(dtbbpy)(ppy)₂PF₆, and Ir[dF(Me)ppy]₂(dtbbpy)PF₆ were all competent in the catalytic process, albeit with moderate yields ranging from 51% to 69% for product 4aa (Table 2, entries 2–4). Lowering the Ni L^{-1} loading resulted in a significant decrease in the yield of 4aa to 50% (Table 2, entry 5), while increasing the Ni L⁻¹ loading provided a similar yield (Table 2, entry 6). Analogous to our previous report,¹¹ extra nickel salts seem to be indispensable for obtaining product 4aa (see ESI Table S9[†] for more details). When a commercially available NiBr₂·dtbbpy complex was used as the sole catalyst, the double aryl group coupling product could also be delivered in 24% yield, probably due to the presence of dissociated nickel salts (Table 2, entry 7). NiCl₂·DME was also an effective nickel source in this catalytic system (Table 2, entry 8), while $Ni(acac)_2$ showed no catalytic activity in the catalytic process (Table 2, entry 9). Similarly, the use of ligands containing electron-deficient groups and polydentate ligands failed to yield the final product 4aa (Table 2, entry 10). The other screened ligands did not afford better results (Table 2, entries 11-13). A

Table 2	Optimized conditions for the formation of 4aa
---------	---

¢	$ \begin{array}{c} & & & \\ & $	4aa
Entry ^a	Deviation	4aa , yield ^b (%)
1	None	$81(72)^{c}$
2	4CzlPN as the photocatalyst	51
3	$Ir(dtbbpy)(ppy)_2PF_6$ as the photocatalyst	69
4	$Ir[dF(Me)ppy]_2(dtbbpy)PF_6$ as the photocatalyst	58
5	NiBr ₂ ·DME (10 mol%), dtbbpy (5 mol%)	50
6	NiBr ₂ ·DME (20 mol%), dtbbpy (10 mol%)	78
7	NiBr ₂ ·dtbbpy (10 mol%)	24
8	NiCl ₂ ·DME instead of NiBr ₂ ·DME	68
9	$Ni(acac)_2$ instead of $NiBr_2 \cdot DME$	n.d.
10	L3, L4 or L6 instead of L1	n.d.
11	L2 instead of L1	75
12	L5 instead of L1	58
13	L7 instead of L1	35
14	1.0 equiv. of HEH	56
15	DMAc instead of NMP	32
16	NMP (0.1 M)	33
17	w/o HEH, irradiation, nickel source, ligand or photocatalyst	n.d

^{*a*} Reactions were performed on a 0.1 mmol scale. ^{*b*} Yields were determined by analyzing ¹H NMR spectroscopic data using 1,3,5-trimethoxybenzene as an internal standard and calculated on a 0.05 mmol. ^{*c*} Isolated yield on a 0.3 mmol scale.

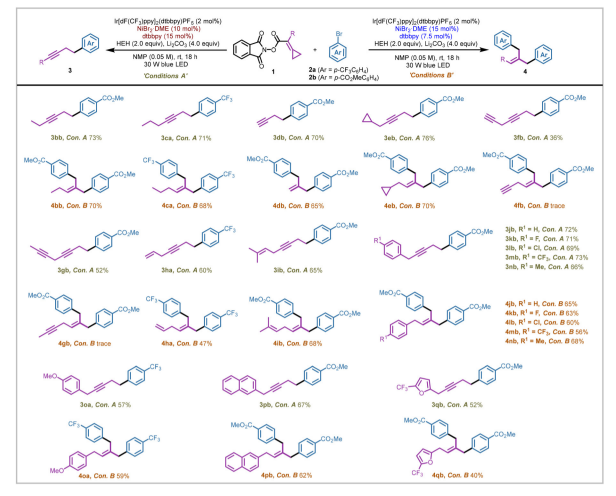
significant decrease was observed upon lowering the amount of HEH to one equivalent (Table 2, entry 14). The use of another aprotic polar solvent DMAc failed to provide **4aa** in a comparable yield to that of NMP (Table 2, entry 15). In addition, such a dual coupling process was quite sensitive to the concentration of the reaction solution. The increase in concentration led to poor reactivity, giving the final product **4aa** with a yield of 33% (Table 2, entry 16). Likewise, HEH, light irradiation, nickel source, ligand and photocatalyst were all indispensable for generating **4aa** (Table 2, entry 17). We also analyzed the internal mixtures of these two reactions (see ESI page S32† for more details).

With the optimized conditions in hand, we then investigated the generality of such a nickel/ligand loading-controlled coupling protocol for producing distinct products 3 and 4. First, a series of NHPI esters bearing methylenecyclopropanes were coupled with ArBr (2a or 2b) under two slightly different conditions (Scheme 2). When the substituent R was an ethyl group (1b) or a propyl group (1c), two modes of coupling reactions could proceed smoothly to afford alkyne products (3bb, 3ca) and double aryl group coupling products (4bb, 4ca) in similar yields to those of substrate 1a. For substrate 1d where the substituent R was a hydrogen atom, the terminal alkyne coupling product 3db was obtained in 70% yield and the 1,1dibenzylethylene product 4db was obtained in 65% yield. The cyclopropane group introduced in substrate 1e was retained in the generation of products 3eb and 4eb. When we introduced a terminal alkyne group in the NHPI ester substrate, the corresponding alkyne product 3fb was obtained in 36% yield, while only a trace amount of 4fb was generated (see ESI page S33[†] for more details). Replacing the terminal alkyne group with an internal alkyne group could increase the yield of alkyne product 3gb to 52%, though the double aryl group coupling product 4gb was still produced in a trace amount. The allyl group (1h) and methyl-substituted allyl group (1i) were well tolerated in this nickel-catalyzed coupling process, affording the corresponding products 3ha and 3ib in 60% and 65% yields, as well as 4ha and 4ib in 47% and 68% yields, respectively. An array of benzyl groups bearing diverse substituents introduced in the substrates (1j-10) were tested under both sets of reaction conditions, affording the alkyne products (3jb-3ob) in yields ranging from 57 to 72% and tribenzylethylene products (4jb-4ob) in yields ranging from 59 to 68%. Notably, substrates containing the fused aromatic ring (1p) and aromatic heterocycle (1q) could also undergo the transformation processes smoothly to give the corresponding products 3pb and 3qb in 67% and 52% yields, as well as 4pb and 4qb in 62% and 40% yields, respectively.

Next, we investigated the scope of the electrophilic coupling partners of aryl bromides under the two sets of reaction conditions (Scheme 3). Aryl bromides bearing electron-withdrawing groups at the *para*-position underwent the two modes of the coupling process smoothly yielding the alkyne products (**3ab-3af**) and 1,1-dibenzylethylene products (**4ab-4ae**) in moderate yields ranging from 56% to 82%. It should be noted that the cyano group was not well tolerated in the double aryl







Scheme 2 Scope of NHPI ester substrates. Reactions were performed on a 0.2 mmol scale for product 3 and on a 0.3 mmol scale for product 4. Isolated yields (yields of product 4 were calculated on a 0.15 mmol scale). See ESI page S12† for more experimental details.

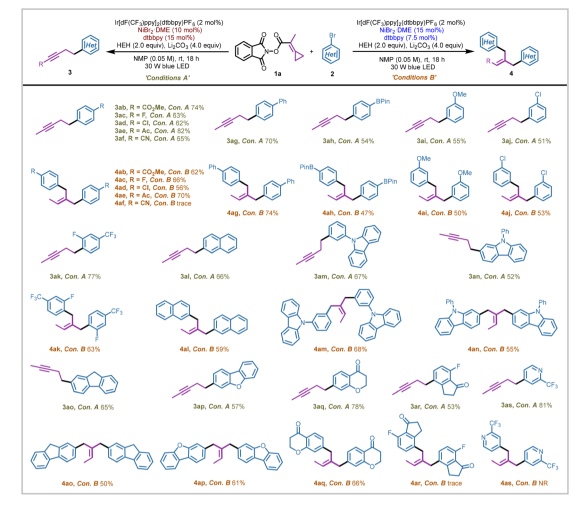
group coupling process, probably owing to coordination with the extra nickel salts resulting in the deactivation of the critical catalytic nickel species. For electron-neutral aryl bromides such as 4-bromobiphenyl, the corresponding coupling products (3ag/4ag) were obtained smoothly in 70% and 74% yields, respectively. Unfortunately, electron-rich aryl bromide namely 4-bromoanisole was not a competent coupling partner in this nickel-catalyzed coupling protocol. Aryl bromides containing meta-substituted groups (OMe, Cl) and ortho-substituted groups (F) were also tested, yielding the corresponding products 3ai-3ak and 4ai-4ak in moderate to good yields ranging from 50% to 77%. Boronic acid pinacol ester was also well tolerated under the two sets of mild conditions, affording the desired products (3ah/4ah) in 54% and 47% yields, respectively. Moreover, diverse polycyclic aryl bromides underwent the coupling processes smoothly to afford the final alkyne products (3al-3ar) and polysubstituted olefinic products (4al-4aq) in yields ranging from 50 to 78%. Notably, product 4ar could not be obtained successfully under the standard conditions B, probably due to the coordination between the nickel species, oxygen atom, and fluorine atom, which restricted the catalytic

ability of the nickel catalysts (see ESI page S33[†] for more details). Furthermore, bromopyridine could not complete the double aryl group coupling process owing to the coordination interactions with the nickel species; however, the corresponding alkyne product **3as** could be furnished in 81% yield.

Furthermore, to investigate the synthetic applicability of the nickel-catalyzed protocol, scale-up experiments of **1a** and synthetic transformations of alkynes were also conducted (see ESI page S36† for more details).

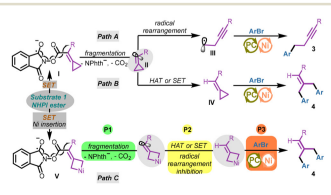
Possible reaction pathways

Our group has recently disclosed that the NHPI ester substrates bearing methylenecyclopropanes could undergo the ring-opening reactions smoothly to deliver the homopropargyl radicals under photoreductive conditions.⁹ Specifically, radical anionic intermediate I formed from substrate 1 (NHPI ester) through a SET (single electron transfer) process could undergo homolytic fragmentation, releasing the phthalimide anion and CO₂. The transiently formed ethylenic radical II underwent a thermodynamically favorable radical rearrangement process to afford homopropargyl radical III. In this protocol, such a



Scheme 3 Scope of aryl bromides. Reactions were performed on a 0.2 mmol scale for product 3 and on a 0.3 mmol scale for product 4. Isolated yields (yields of product 4 were calculated on a 0.15 mmol scale). See ESI page S12[†] for more experimental details.

 $C(sp^3)$ hybridized fragment should engage in the nickel-catalyzed $C(sp^3)$ - $C(sp^2)$ cross-coupling process to give the final alkyne product 3 (Scheme 4, Path A).¹² In addition, the double aryl group coupling product was also obtained in this photoredox/nickel catalyzed protocol. We initially speculated that

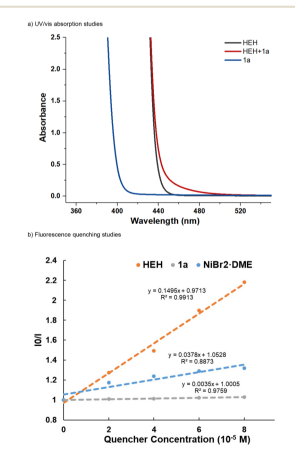


Scheme 4 Possible reaction pathways for the generation of products.

the ethylenic radical II may undergo a HAT (hydrogen atom transfer) process or a SET process to provide the methylenecyclopropane intermediate IV. Then intermediate IV could undergo a Ni-catalyzed coupling process with ArBr via selective distal bond cleavage to afford product 4 (Scheme 4, Path B), a known process recently demonstrated by our group.¹¹ However, considering that the radical rearrangement process is quite thermodynamically favored, Path B could not explain why the alkyne products 3 were not generated during the production of 4. Herein, we provide a tentative hypothesis to explain how the addition of extra nickel salts led to the production of two different products (Scheme 4, Path C). Due to the lack of coordination with bidentate ligands, the additional "free" nickel salts with electron deficiency and less steric hindrance were more inclined to undergo the oxidative addition process with the electron-rich methylenecyclopropane structure in the substrate (cyclopropane insertion). As a result, the nickelacyclobutene intermediate13 would inhibit the radical rearrangement in the homolytic fragmentation process, bypassing the generation of homopropargyl radical III. The fol-

Mechanistic investigations

To investigate the mechanism of NHPI ester activation, we first conducted UV/vis absorption studies of the redox-active substrate **1a** and HEH. Consistent with the previous reports,¹⁴ a bathochromic shift (Scheme 5a, red band) was observed for a mixture of NHPI ester **1a** and HEH, indicating that the formation of an electron donor–acceptor (EDA) complex was probably involved in this nickel-catalyzed protocol. Hence, the EDA-complex could provide a potential pathway for the single electron reduction of the NHPI ester substrate. Next, fluorescence quenching studies of the photocatalyst with HEH, substrate **1a** and NiBr₂·DME were conducted (Scheme 5b). The excited state of Ir[dF(CF₃)ppy]₂(dtbbpy)PF₆ was more strongly quenched by HEH, implying that a reductive quenching cycle¹⁵ was involved in the reaction process and a reducing Ir



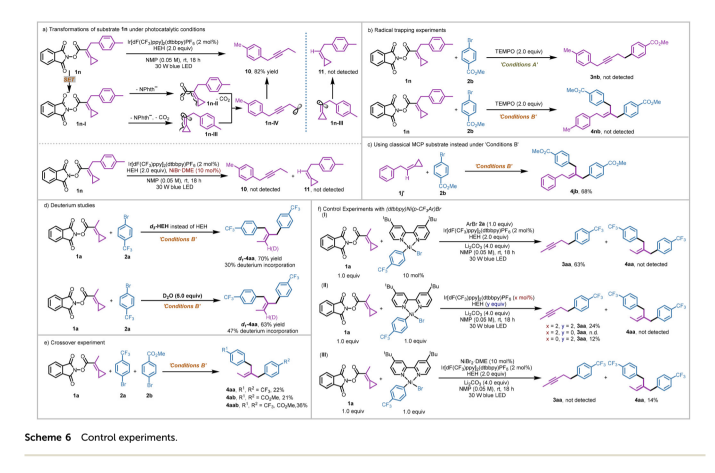
Scheme 5 UV/vis absorption studies and fluorescence quenching studies.

(II) center $(E_{1/2}^{\text{red}} = -1.37 \text{ V vs. SCE in MeCN})^{16}$ was generated. Though substrate **1a** exhibited no quenching effect on the excited photocatalyst, we suggested that the NHPI ester **1a** $(E_{1/2}^{\text{red}} = -1.11 \text{ V vs. SCE in DMAc}$, see ESI Fig. S8[†] for more details) could also receive an electron from the reducing Ir(II) center.⁹ The quantum yields were measured as $\Phi = 0.15$ and $\Phi = 0.04$ (see ESI page S19[†] for more details) in the generation of alkyne product **3aa** and dibenzylethylene product **4aa**, respectively, suggesting that a radical chain process was possibly not involved in the two sets of reactions.

A series of control experiments were also conducted to gain more insights into the two sets of reaction mechanisms. We first investigated the transformations of the NHPI ester substrate under the photocatalytic conditions. Rearrangement product 10 could be isolated in 82% yield from substrate 1n as a single product (Scheme 6a, above). This result suggests that such a rearrangement product was formed from the homopropargyl radical 1n-IV. In particular, substrate 1n underwent a SET process to form the radical anionic intermediate 1n-I, which could release the phthalimide anion and CO₂ along with a rearrangement process to give intermediate 1n-IV through intermediate 1n-II or 1n-III. Though the classical MCP substrate 1j' could also deliver the corresponding product **4jb** in 68% yield under conditions B (Scheme 6c),¹¹ we did not observe the cyclopropane-maintained product 11, which could be generated from the possible vinyl radical 1n-III through a HAT (or SET) process. In addition, considering that the nickel species may promote the hydrogenation of vinyl radical 1n-III, we added extra NiBr2·DME (10 mol%) into the reaction (Scheme 6a, below). However, we still did not observe any cyclopropane-maintained product 11. The rearrangement product 10 was not detected owing to the destruction of the MCP moiety by the nickel species (see ESI page S14[†] for more details). These results implied that the possible intermediate IV proposed in Scheme 4 (Path B) was possibly not involved in the reaction process of generating product 4 and path B was unlikely. Next, 2,2,6,6-tetramethylpiperidinooxy (TEMPO) used as the radical scavenger was added to the two sets of reactions, both reactions were inhibited completely and the corresponding products were not detected (Scheme 6b). These two experiments suggest that radical reaction pathways were possibly involved in the two sets of coupling reactions. Deuterium labeling studies using d_3 -HEH and D_2O were conducted, and product d_1 -4aa was obtained with 30% and 47% deuterium incorporation, respectively (Scheme 6d). These results indicate that the olefinic hydrogen atom in the double aryl group coupling product 4aa may originate from HEH or a trace amount of water in the reaction system through SET and HAT processes. A crossover experiment was carried out using substrates 1a, 2a, and 2b under the conditions B. Apart from the production of the corresponding products 4aa and 4ab, a mixed coupling product 4aab was also isolated in 36% yield, presumably indicating that the two aryl group coupling process involved two discrete steps (Scheme 6e).

When a catalytic amount of $(dtbbpy)Ni(p-CF_3Ar)Br$ (10 mol%) was applied in the reaction, **3aa** could be smoothly

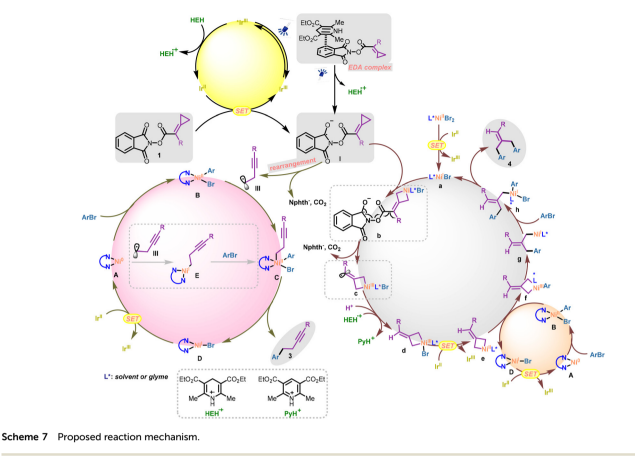
Organic Chemistry Frontiers



obtained in 63% yield while 4aa was not detected, indicating that the Ni complex (dtbbpy)NiArBr was a possible catalytically active species in the generation of product 3 but not responsible for the cleavage of the cyclopropane ring in the NHPI ester substrate (Scheme 6f, eqn (I)). Stoichiometric experiments with (dtbbpy)Ni(p-CF₃Ar)Br as a possible Ni(II) intermediate were next conducted (Scheme 6f, eqn (II)), where alkyne product 3aa could be obtained in 24% yield in the presence of the photocatalyst (x = 2) and HEH (y = 2). In addition, when the sacrificial reductant HEH was omitted (y = 0), no coupling product (3aa or 4aa) was detected. However, in the absence of a photocatalyst (x = 0), product 3aa could be generated in 12% yield. Such stoichiometric control experiments revealed that activating the NHPI ester mediated by the formation of an EDA complex was a feasible pathway.^{14a} Notably, product 4aa was not detected in these stoichiometric experiments, highlighting the indispensable role of "free" nickel salts in generating the dibenzylethylene product 4. When extra NiBr₂·DME was added, 4aa was obtained in 14% yield, while 3aa was not detected (Scheme 6f, eqn (III)). All these results suggest that (i) (dtbbpy)NiArBr should be a catalytically active species in the nickel catalytic cycle for the generation of product 3; (ii) the "free" nickel species, rather than the ligandcoordinated nickel species, should play the role of activating the cyclopropane ring in the NHPI ester substrate for the generation of product 4; and (iii) the SET activation of the NHPI

ester substrate *via* the EDA complex was also feasible in this protocol.

On the basis of the above mechanistic studies and previous reports, we propose a preliminary plausible mechanism for this nickel/ligand loading controlled divergent synthetic protocol (Scheme 7). Initially, the excited photocatalyst Ir(III) (* $E_{1/2}^{\text{red}}$ = +1.21 V vs. SCE in MeCN)¹⁶ was quenched by the sacrificial reductant HEH ($E_{1/2}^{\text{ox}}$ = +0.89 V vs. SCE in DMF),¹⁷ delivering a strongly reducing center Ir(π) ($E_{1/2}^{red} = -1.37$ V vs. SCE in MeCN),¹⁶ which is a competent reducing center for the reduction of the NHPI ester substrate and nickel species. With regard to the single-electron reduction of the NHPI ester substrate 1, two pathways should be responsible for the generation of a radical anionic intermediate (I).^{14a} On the one hand, substrate 1 could be directly reduced by the reducing Ir(II) center to afford the radical anionic intermediate I; on the other hand, an EDA complex formed by the electron-deficient NHPI ester and the electron-rich HEH would also result in the single electron reduction of substrate 1 to generate intermediate I. Subsequently, when there were no "free" nickel salts present in the reaction system, the entity intermediate I could undergo radical fragmentation and a thermodynamically favored radical rearrangement process smoothly to give the homopropargyl radical III. According to the reported detailed computational studies,¹⁸ such a C(sp³) radical fragment may go through two reaction pathways: (i) captured by the $L_n Ni^{II} ArBr$ ($L_n Ni$: nickel



species with bidentate ligand coordination) species **B**, which was formed from the oxidative addition of the aryl halides to Ni⁰ species **A**, to form the high-valent Ni^{III} species **C**; (ii) combined with the initial Ni⁰L_n species **A** to form the Ni^I intermediate **E**. In the first reaction pathway, the Ni^{III} species **C** underwent a reductive elimination process to yield the final $C(sp^2)-C(sp^3)$ coupling product 3 and the corresponding Ni^I species **D**. The Ni⁰L_n species **A** could be regenerated by the reduction of the Ni^I species **D** $(E_p^{red}(Ni^IL_n/Ni^0L_n) = -1.17 V vs. SCE in THF)^{19}$ through a SET event with the Ir(II) reducing center. In the second reaction pathway, subsequent oxidative addition of ArBr to the Ni^I

entering into the catalytic cycle (Scheme 7, bottom left). In contrast, when the nickel salts were excessive in the reactions, the cyclopropane ring in the radical anionic intermediate I could not be maintained for the following radical rearrangement process. In this scenario (Scheme 7, bottom right), the electron-deficient nickel center of L*Ni^IBr a ("free" nickel, L*: solvent or glyme) with less steric hindrance, speculatively reduced by the Ir(II) center ($E_{1/2}^{red}$ (Ni^{II}/Ni^I) = -1.70 V vs. Fc/Fc⁺ in DMF),²⁰ could insert into the cyclopropane ring forming the nickelacyclobutane species **b** that inhibited the radical rearrangement. In this case, intermediate **b** possibly undergoes the radical fragmentation process to release the phthalimide anion and CO₂, affording the vinyl radical intermediate **c**. After the SET and HAT processes, the free MCP

intermediate E would occur, affording the same Ni^{III} species C

intermediate d with nickel insertion was generated. The following transformations of such nickel(m)acyclobutane d were previously investigated by our group.11 In particular, the Ni^{III} species **d** could be reduced to Ni^{II} species **e** by Ir(II) through a SET event. Meanwhile, oxidative addition of ArBr to the electron-rich $Ni^{0}L_{n}$ species **A** provided the $L_{n}Ni^{II}ArBr$ complex **B**. An aryl group transfer (disproportionation) event was considered to take place between nickelacyclobutane species e and L_nNi^{II}ArBr **B**,²¹ affording the corresponding aryl-nickelacyclobutane species **f** and $L_n Ni^I Br$ species **D**. The $L_n Ni^I Br$ species **D** could be reduced by the reducing Ir(n) center to regenerate the Ni⁰L_n species. Furthermore, the first reductive elimination from the aryl-nickelacyclobutane species f delivered the Ni^{1} intermediate g, which could undergo oxidative addition with ArBr or potentially another aryl ligand transfer process to deliver intermediate h. Intermediate h then underwent a second reductive elimination, yielding the final product 4 and regenerating the L*Ni^IBr species a. Notably, the aryl group transfer event occurring between nickel(III)acyclobutane **d** and $L_n Ni^{II} ArBr B$ may also be feasible^{21a} (see ESI page S35[†] for more details).

Conclusions

In summary, we have developed a nickel/ligand loading-controlled divergent synthesis protocol, offering an array of alkyne

Research Article

derivatives and dibenzylethylene derivatives in moderate to good yields under mild conditions along with a broad substrate scope. Selective cleavage of carbon-carbon bonds on redox-active MCP substrates can be achieved efficiently by adjusting the loading of the nickel catalyst and bidentate ligand. Specifically, alkyne products can be obtained when the nickel-to-ligand ratio is 2 to 3, while dibenzylethylene products can be obtained when the ratio is 2 to 1 under almost identical conditions. Temporary mechanistic studies demonstrate that the generation of alkyne products should involve homopropargyl radicals stemming from the rearrangement of the MCP structure in the substrates, while the generation of dibenzylethylene products should involve nickelacyclobutane intermediates. Further investigations into this interesting loadingcontrolled catalytic mode and mechanistic studies are ongoing in our group. We hope such a ligand loading-controlled scheme could raise attention to ligand loading effects in transition-metal catalysis and bring more unexpected but applicable transformations in synthetic chemistry.

Author contributions

B. Mao contributed to the experimental work. B. Mao, X.-Y. Zhang, X.-F. Bi, X.-Y. Yang, M. Shi and Y. Wei contributed to conceptualizations and writing of the paper.

Data availability

Experimental and computational data have been made available as the ESI. \dagger

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We are grateful for the financial support from the National Key R & D Program of China (2023YFA1506700), the National Natural Science Foundation of China (21372250, 21121062, 21302203, 21772037, 21772226, 21861132014, 91956115 and 22171078) and the Fundamental Research Funds for the Central Universities 222201717003.

References

 (a) Y. Ping, X. Li, Q. Pan and W. Kong, Ni-Catalyzed Divergent Synthesis of 2-Benzazepine Derivatives via Tunable Cyclization and 1,4-Acyl Transfer Triggered by Amide N-C Bond Cleavage, *Angew. Chem., Int. Ed.*, 2022, 61, e202201574; (b) Y. Ke, W. Li, W. Liu and W. Kong, Ni-catalyzed ligand-controlled divergent and selective synthesis, Sci. China: Chem., 2023, **66**, 2951–2976; (c) Q. Pan, Y. Ping and W. Kong, Nickel-Catalyzed Ligand-Controlled Selective Reductive Cyclization/Cross-Couplings, *Acc. Chem. Res.*, 2023, **56**, 515–535; (d) Y. Wang, J. Feng, E.-Q. Li, Z. Jia and T.-P. Loh, Recent advances in ligand-enabled palladiumcatalyzed divergent synthesis, *Org. Biomol. Chem.*, 2024, **22**, 37–54; (e) I. P. Beletskaya, C. Nájera and M. Yus, Chemodivergent reactions, *Chem. Soc. Rev.*, 2020, **49**, 7101– 7166; (f) T. Long, C. Zhu, L. Li, L. Shao, S. Zhu, M. Rueping and L. Chu, Ligand-controlled stereodivergent alkenylation of alkynes to access functionalized trans- and cis-1,3dienes, *Nat. Commun.*, 2023, **14**, 55.

- 2 (a) A. M. Allgeier and C. A. Mirkin, Ligand Design for Electrochemically Controlling Stoichiometric and Catalytic Reactivity of Transition Metals, Angew. Chem., Int. Ed., 1998, 37, 894-908; (b) R. Martin and S. L. Buchwald, Palladium-Catalyzed Suzuki-Miyaura Cross-Coupling Reactions Employing Dialkylbiaryl Phosphine Ligands, Acc. Chem. Res., 2008, 41, 1461-1473; (c) C. C. Chintawar, A. K. Yadav, A. Kumar, S. P. Sancheti and N. T. Patil, Divergent Gold Catalysis: Unlocking Molecular Diversity through Catalyst Control, Chem. Rev., 2021, 121, 8478-8558; (d) Y. Himeda, Conversion of CO_2 into Formate by Homogeneously Catalyzed Hydrogenation in Water: Tuning Catalytic Activity and Water Solubility through the Acid-Base Equilibrium of the Ligand, Eur. J. Inorg. Chem., 2007, 2007, 3927-3941; (e) M. D. Bairagya, R. J. Bujol and N. Elgrishi, Fighting Deactivation: Classical and Emerging Strategies for Efficient Stabilization of Molecular Electrocatalysts, Chem. - Eur. J., 2020, 26, 3991-4000; (f) T. Piou and T. Rovis, Electronic and Steric Tuning of a Prototypical Piano Stool Complex: Rh(III) Catalysis for C-H Functionalization, Acc. Chem. Res., 2018, 51, 170-180.
- 3 (a) Y. Sakakibara and K. Murakami, Switchable Divergent Synthesis Using Photocatalysis, ACS Catal., 2022, **12**, 1857– 1878; (b) L. Li, Z. Chen, X. Zhang and Y. Jia, Divergent Strategy in Natural Product Total Synthesis, Chem. Rev., 2018, **118**, 3752–3832; (c) G. Zhan, W. Du and Y.-C. Chen, Switchable divergent asymmetric synthesis via organocatalysis, Chem. Soc. Rev., 2017, **46**, 1675–1692.
- 4 (a) Y.-C. Lee, K. Kumar and H. Waldmann, Ligand-Directed Divergent Synthesis of Carbo- and Heterocyclic Ring Systems, Angew. Chem., Int. Ed., 2018, 57, 5212-5226; (b) Y.-C. Lee, S. Patil, C. Golz, C. Strohmann, S. Ziegler, K. Kumar and H. Waldmann, A ligand-directed divergent catalytic approach to establish structural and functional scaffold diversity, Nat. Commun., 2017, 8, 14043; (c) Y. Shi, T. Huang, T. Wang, J. Chen, X. Liu, Z. Wu, X. Huang, Y. Zheng, Z. Yang and Y. Wu, Divergent Construction of Diverse Scaffolds through Catalyst-Controlled C-H Activation Cascades of Quinazolinones and Cyclopropenones, Chem. - Eur. J., 2021, 27, 13346-13351; (d) T. N. Nguyen and J. A. May, Branched Amine Synthesis via Aziridine or Azetidine Opening with Organotrifluoroborates by Cooperative Brønsted/Lewis Acid Catalysis: An Acid-Dependent Divergent Mechanism, Org.

Lett., 2018, **20**, 3618–3621; (*e*) G. Chen and B. Xu, Divergent Synthesis of Sulfonyl Quinolines, Formyl Indoles, and Quinolones from Ethynyl Benzoxazinanones via AuI Catalysis, AuI-ArI Co-Catalysis, and Silver Catalysis, *ACS Catal.*, 2022, **12**, 7134–7141.

- 5 (a) P. Binger, Cyclodimerization of Methylenecyclopropane on a Ni(0) Catalyst, Angew. Chem., Int. Ed. Engl., 1972, 11, 309-310; (b) L. Yu, M. Liu, F. Chen and Q. Xu, Heterocycles from methylenecyclopropanes, Org. Biomol. Chem., 2015, 13, 8379-8392; (c) L.-X. Shao and M. Shi, Lewis and Bronsted Acid Mediated Ring-Opening Reactions of Methylenecyclopropanes and Further Transformation of the Ring-Opened Products, Curr. Org. Chem., 2007, 11, 1135-1153; (d) H. Pellissier, Recent developments in the reactivity of methylene- and alkylidenecyclopropane derivatives, Tetrahedron, 2010, 66, 8341-8375; (e) H. Cao, F. Chen, C. Su and L. Yu, Construction of Carbocycles from Methylenecyclopropanes, Adv. Synth. Catal., 2020, 362, 438-461; (f) H.-Z. Wei, M. Shi and Y. Wei, Visible-lightinduced reactions of methylenecyclopropanes (MCPs), Chem. Commun., 2023, 59, 2726-2738.
- 6 (a) A. Masarwa and I. Marek, Selectivity in Metal-Catalyzed Carbon-Carbon Bond Cleavage of Alkylidenecyclopropanes, *Chem. Eur. J.*, 2010, 16, 9712–9721; (b) G. Fumagalli, S. Stanton and J. F. Bower, Recent Methodologies That Exploit C-C Single-Bond Cleavage of Strained Ring Systems by Transition Metal Complexes, *Chem. Rev.*, 2017, 117, 9404–9432; (c) I. Nakamura and Y. Yamamoto, Transition Metal-Catalyzed Reactions of Methylenecyclopropanes, *Adv. Synth. Catal.*, 2002, 344, 111–129; (d) P.-h. Chen, B. A. Billett, T. Tsukamoto and G. Dong, "Cut and Sew" Transformations via Transition-Metal-Catalyzed Carbon-Carbon Bond Activation, *ACS Catal.*, 2017, 7, 1340–1360; (e) D. Li, W. Zang, M. J. Bird, C. J. T. Hyland and M. Shi, Gold-Catalyzed Conversion of Highly Strained Compounds, *Chem. Rev.*, 2021, 121, 8685–8755.
- 7 B. B. Tan, M. Hu and S. Ge, Cobalt-Catalyzed Regiodivergent Ring-Opening Dihydroboration of Arylidenecyclopropanes to Access Skipped Diboronates, *Angew. Chem., Int. Ed.*, 2023, **62**, e202307176.
- 8 (a) M. Shi, J.-M. Lu, Y. Wei and L.-X. Shao, Rapid Generation of Molecular Complexity in the Lewis or Brønsted Acid-Mediated Reactions of Methylenecyclopropanes, Acc. Chem. Res., 2012, 45, 641-652; (b) L.-Z. Yu, Q. Xu, X.-Y. Tang and M. Shi, Iron- or Copper-Catalyzed Trifluoromethylation of Acrylamide-Tethered Alkylidenecyclopropanes: Facile Synthesis of CF₃-Containing Polycyclic Benzazepine Derivatives, ACS Catal., 2016, 6, 526-531; (c) M. Chen, Y. Wei and M. Shi, Cascade cyclization reactions of alkylidenecyclopropanes for the construction of polycyclic lactams and lactones by visible light photoredox catalysis, Org. Chem. Front., 2020, 7, 374-379; (d) X.-Y. Zhang, X.-Y. Wu, B. Zhang, Y. Wei and M. Shi, Silyl Radical-Mediated Carbocyclization of Acrylamide-/ Vinyl Sulfonamide-Attached Alkylidenecyclopropanes via Photoredox Catalysis with a Catalytic Amount of Silane

Reagent, ACS Catal., 2021, **11**, 4372–4380; (e) B. Mao, X.-Y. Zhang, Y. Wei and M. Shi, Visible-light-mediated intramolecular radical cyclization of α -brominated amidetethered alkylidenecyclopropanes, *Chem. Commun.*, 2022, **58**, 3653–3656.

- 9 X.-Y. Zhang, C. Ning, B. Mao, Y. Wei and M. Shi, A visiblelight mediated ring opening reaction of alkylidenecyclopropanes for the generation of homopropargyl radicals, *Chem. Sci.*, 2021, **12**, 9088–9095.
- 10 (a) S. Murarka, N-(Acyloxy)phthalimides as Redox-Active Esters in Cross-Coupling Reactions, Adv. Synth. Catal., 2018, 360, 1735–1753; (b) P. Niu, J. Li, Y. Zhang and C. Huo, One-Electron Reduction of Redox-Active Esters to Generate Carbon-Centered Radicals, Eur. J. Org. Chem., 2020, 5801–5814; (c) S. K. Parida, T. Mandal, S. Das, S. K. Hota, S. De Sarkar and S. Murarka, Single Electron Transfer-Induced Redox Processes Involving N-(Acyloxy) phthalimides, ACS Catal., 2021, 11, 1640–1683.
- 11 B. Mao, M. Shi and Y. Wei, Ni/Photoredox-catalyzed coupling of aryl bromides and methylenecyclopropanes via selective distal bond cleavage, *Org. Chem. Front.*, 2024, DOI: 10.1039/D4Q001072H.
- 12 (a) J. C. Tellis, C. B. Kelly, D. N. Primer, M. Jouffroy, N. R. Patel and G. A. Molander, Single-Electron Transmetalation via Photoredox/Nickel Dual Catalysis: Unlocking a New Paradigm for sp³-sp² Cross-Coupling, *Acc. Chem. Res.*, 2016, 49, 1429-1439; (b) J. Twilton, C. Le, P. Zhang, M. H. Shaw, R. W. Evans and D. W. C. MacMillan, The merger of transition metal and photocatalysis, *Nat. Rev. Chem.*, 2017, 1, 0052.
- 13 M. L. G. Sansores-Paredes, P. M. Pérez-García and M.-E. Moret, Nickelacyclobutanes: Versatile Reactivity and Role as Catalytic Intermediates, *Eur. J. Inorg. Chem.*, 2023, 26, e202300192.
- 14 (*a*) L. M. Kammer, S. O. Badir, R. M. Hu and G. A. Molander, Photoactive electron donor-acceptor complex platform for Ni-mediated $C(sp^3)$ - $C(sp^2)$ bond formation, *Chem. Sci.*, 2021, **12**, 5450–5457; (*b*) F. Cong, X. Y. Lv, C. S. Day and R. Martin, Dual Catalytic Strategy for Forging sp^2 - sp^3 and sp^3 - sp^3 Architectures via β -Scission of Aliphatic Alcohol Derivatives, *J. Am. Chem. Soc.*, 2020, **142**, 20594–20599.
- 15 (a) X. Xi, Y. Luo, W. Li, M. Xu, H. Zhao, Y. Chen, S. Zheng, X. Qi and W. Yuan, From Esters to Ketones via a Photoredox-Assisted Reductive Acyl Cross-Coupling Strategy, Angew. Chem., Int. Ed., 2022, 61, e202114731; (b) L. J. Rono, H. G. Yayla, D. Y. Wang, M. F. Armstrong and R. R. Knowles, Enantioselective Photoredox Catalysis Proton-Coupled Transfer: Enabled by Electron Development of an Asymmetric Aza-Pinacol Cyclization, J. Am. Chem. Soc., 2013, 135, 17735-17738.
- 16 C. K. Prier, D. A. Rankic and D. W. C. MacMillan, Visible Light Photoredox Catalysis with Transition Metal Complexes: Applications in Organic Synthesis, *Chem. Rev.*, 2013, **113**, 5322–5363.
- 17 (a) G. Park, S. Y. Yi, J. Jung, E. J. Cho and Y. You, Mechanism and Applications of the Photoredox Catalytic

Coupling of Benzyl Bromides, *Chem. – Eur. J.*, 2016, 22, 17790–17799; (*b*) K. N. Lee, Z. Lei and M.-Y. Ngai, β -Selective Reductive Coupling of Alkenylpyridines with Aldehydes and Imines via Synergistic Lewis Acid/ Photoredox Catalysis, *J. Am. Chem. Soc.*, 2017, **139**, 5003–5006.

- 18 O. Gutierrez, J. C. Tellis, D. N. Primer, G. A. Molander and M. C. Kozlowski, Nickel-Catalyzed Cross-Coupling of Photoredox-Generated Radicals: Uncovering a General Manifold for Stereoconvergence in Nickel-Catalyzed Cross-Couplings, J. Am. Chem. Soc., 2015, 137, 4896–4899.
- 19 T. J. Steiman, J. Liu, A. Mengiste and A. G. Doyle, Synthesis of β -Phenethylamines via Ni/Photoredox Cross-Electrophile

Coupling of Aliphatic Aziridines and Aryl Iodides, J. Am. Chem. Soc., 2020, 142, 7598–7605.

- 20 Q.-Y. Meng, S. Wang and B. König, Carboxylation of Aromatic and Aliphatic Bromides and Triflates with CO₂ by Dual Visible-Light-Nickel Catalysis, *Angew. Chem., Int. Ed.*, 2017, **56**, 13426–13430.
- 21 (a) K. Osakada and T. Yamamoto, Transmetallation of alkynyl and aryl complexes of Group 10 transition metals, *Coord. Chem. Rev.*, 2000, **198**, 379–399; (b) G. S. Kumar, C. Zhu, R. Kancherla, P. S. Shinde and M. Rueping, Metal Cations from Sacrificial Anodes Act as a Lewis Acid Co-Catalyst in Electrochemical Cross-Coupling of Aryl Bromides and Aziridines, *ACS Catal.*, 2023, **13**, 8813–8820.