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Introduction

The dearomatization reaction has emerged as an expedient and versatile synthetic strategy to construct three-dimensional complex molecular skeletons from easily available flat aromatic compounds.1 Within this realm, significant attention has been directed toward the dearomatization of arenes and heteroarenes to assemble cyclic and heterocyclic compounds. Despite the high synthetic value of this strategy, the inherent challenge lies in the low reactivity of the aromatic substrates due to their reluctance to lose aromaticity as well as the difficulty in controlling the regio- and stereoselectivity in the course of generating the target three-dimensional products. In view of the salient features of pyridine, quinoline and isoquinoline including ready availability, stability and ease of handling, the straightforward dearomatization reaction with electrophilically improved pyridinium, quinolinium and isoquinolinium salts represents one of the most efficient and robust approaches for the synthesis of hydrogenated azaarenes, which are common molecular architectures of a number of biologically active natural products and pharmaceuticals.²

Regio- and diastereoselective dearomatizations of *N*-alkyl activated azaarenes: the maximization of the reactive sites[†]

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An unprecedented base-promoted multi-component one-pot dearomatization of *N*-alkyl activated azaarenes was developed, which enabled the synthesis of complex and diverse bridged cyclic polycycles with multiple stereocenters in a highly regio- and diastereoselective manner. Besides, we realized the step-controlled dearomative bi- and trifunctionalization of quinolinium salts. These transformations not only achieved the maximization of the reaction sites of pyridinium, quinolinium and isoquinolinium salts to enhance structural complexity and diversity, but also opened up a new reaction mode of these *N*-activated azaarenes. A unique feature of this strategy is the use of easily accessible and bench-stable *N*-alkyl activated azaarenes to provide maximum reactive sites for dearomative cascade cyclizations. In addition, the salient characteristics including high synthetic efficiency, short reaction time, mild conditions and simple operation made this strategy particularly attractive.

A survey of literature revealed that the reaction modes of pyridinium, quinolinium and isoquinolinium salts were mainly divided into two categories (Scheme 1a). The first one was the extensively studied monofunctionalization of pyridinium,3 guinolinium⁴ and isoquinolinium salts⁵ by taking advantage of their electrophilic activity via nucleophilic attack at the more electron-deficient carbons. However, for pyridinium and quinolinium salts, both the C-2 and C-4 positions are active sites. Thus, the regioselectivity of these additions was a main issue of concern. Generally, mixtures of regioisomers were observed in many cases. Introduction of an electron-withdrawing group at the C-3 position could regioselectively favor the C-4 nucleophilic attack.6 The second one was the sparsely reported bifunctionalization of these N-activated azaarenes, where the C-2 and C-4 positions of pyridinium and quinolinium salts and the C-1 and C-3 positions of isoquinolinium salts were skillfully employed to construct interesting bridged cyclic azaheterocycles (Scheme 1b).7 Although these elegant strategies have proved to be quite successful to assemble hydrogenated azaarenes via dearomative nucleophilic additions, only one or two reactive sites were exploited, which largely restricted their application for the construction of complex compounds. Therefore, it is highly important, but an extremely challenging task, to develop new reaction systems that facilitate the exploitation of all of the potential reactive sites of pyridinium, quinolinium and isoquinolinium salts to achieve structural complexity and diversity.

Theoretically, pyridinium salts possess three electrophilic sites at the C-2, C-4 and C-6 positions and two nucleophilic sites at the C-3 and C-5 positions once the first attack by a nucleophile at the C-4 position was initiated (Scheme 2a). Similarly,



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^bInstitute of Chemistry, Henan Academy of Sciences, Zhengzhou 450002, P. R. China [†] Electronic supplementary information (ESI) available: Detailed experimental procedures; characterization data of all the new products; copies of ¹H NMR and ¹³C NMR. CCDC 1944660 (**3a**), 1947688 (**3zb**), 1947689 (**5m**), 1947690 (**6n**), 1947693 (**10g**), 1947691 (**10p**) and 1944631 (**11**). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc04880d



Scheme 1 (a) Previous dearomative monofunctionalization of *N*-activated azaarenes. (b) Sparsely reported bifunctionalization of *N*-activated azaarenes.

quinolinium and isoquinolinium salts also own two electrophilic sites and one nucleophilic site, respectively. Based on this knowledge, we envisioned that the maximization of the reactive sites of *N*-alkyl activated azaarenes could be achieved to enable the construction of complex and diverse bridged cyclic heterocycles by choosing the appropriate binucleophile and electrophile *via* a one-pot three-component dearomative cascade reaction (Scheme 2b). Actually, *N*-alkyl activated azaarenes themselves could be capable of behaving as electrophiles.

In this scenario, there were three inherent synthetic challenges: (1) the dearomative multifunctionalization of pyridinium, quinolinium and isoquinolinium salts has never been



E¹/E²/E³ represents another one molecule of *N*-activated azaarene. Challenges: (1) finding suitable binucleophiles (2) controlling the regio- and stereoselectivity

Scheme 2 (a) The potential reactivity of N-activated azaarenes. (b) Our synthetic strategy to maximize the reactive sites of N-alkyl activated azaarenes.

studied; (2) the appropriate binucleophile and suitable reaction conditions had to be found to improve the synthetic efficiency and control the regioselectivity and stereoselectivity; (3) the generation of multiple stereocenters embedded into bridged ring systems was a big challenge. As a continuation of our program aiming at the exploration of new cascade reactions for the construction of bridged rings with high synthetic efficiency and excellent stereocontrol,8 herein, we wish to present a basepromoted diastereoselective one-pot multicomponent dearomative cascade reaction of pyridinium, quinolinium and isoquinolinium salts, which realized the maximization of the active reactive sites of these N-alkyl activated azaarenes for dearomatization to construct diverse and complex bridged ring systems with multiple stereocenters. Such skeletons are privileged structural units of numerous natural products and biologically active pharmaceuticals.9

Results and discussion

As an initial study, enaminone **1a** and *N*-benzyl-3nitropyridinium salt 2a were chosen as the model substrates to test the feasibility of our synthetic design. When the reaction was conducted in CH₃CN at 60 °C with 2.0 equivalents of Cs₂CO₃ as the base, gratifyingly, the desired bridged polycycle product 3a was delivered in 12% yield with complete regio- and diastereomeric control, whose structure was unambiguously verified by a comprehensive NMR, HRMS and X-ray analysis (Table 1, entry 1). This preliminary result definitely indicated that the dearomative cascade annulation of pyridinium salts was feasible. The formation of 3a was very appealing from the perspective of the reaction process and product structure. This transformation opened up an unprecedented reaction mode of pyridinium salt, where all of the five active sites were tactfully utilized to simultaneously construct five new bonds, two bridged rings as well as a challenging fused cyclobutane¹⁰ and eight stereocenters in a highly regio- and diastereoselective one-pot fashion. Moreover, the product contained some useful functional groups, such as nitro, carbonyl and a cyclic enamine, which offered versatile platforms for further architectural modifications.

To improve the synthetic efficiency, some inorganic and organic bases were evaluated. Among them, 1,1,3,3-tetramethylguanidine (TMG) was identified to be optimal in terms of yield and reaction time, in which the reaction went to full conversion within 5 min, affording 3a in 87% yield (Table 1, entry 8). Remarkably, in this condition, the product was precipitated from the homogenous reaction mixture. So only a filtration process was needed to purify it, which largely simplified the purification procedure and met the requirement of green chemistry. Then, the reaction media were examined and the results suggested that they all gave inferior yields compared with CH_3CN (Table 1, entries 9–11 vs. 8). Finally, the loadings of N-benzyl-3-nitropyridinium salt 2a were investigated. When 2.2 equivalents of 2a were used, product 3a was precipitated out in a yield of 95%. No increment in yield was observed on further raising the amount of 3a to 2.4 equivalents. To achieve the asymmetric synthesis of 3a, we also examined some chiral bases, including cinchona alkaloids and their



Entry	Base	Solvent	Time	Yield ^b (%)
1	Cs_2CO_3	CH ₃ CN	58 h	12
2	K ₂ CO ₃	CH ₃ CN	18 h	15
3	Na_2CO_3	CH ₃ CN	18 h	12
4	NaOH	CH ₃ CN	18 h	30
5	$DABCO \cdot 6H_2O$	CH ₃ CN	27 h	16
6	DBU	CH ₃ CN	33 h	45
7	NEt ₃	CH ₃ CN	3 h	32
8	TMG	CH ₃ CN	5 min	87 ^c
9	TMG	$CHCl_3$	4 h	82
10	TMG	Toluene	4 h	74^c
11	TMG	EtOAc	20 min	58
12^d	TMG	CH ₃ CN	5 min	95 ^c
13^e	TMG	CH ₃ CN	5 min	80^{c}

^{*a*} Unless otherwise noted, the reactions were conducted with 0.15 mmol **1a** with 2.0 equivalents of **2a** in the presence of 2.0 equivalents of base in 0.8 mL of specified solvent at 60 °C. ^{*b*} Isolated yields obtained by column chromatography. ^{*c*} Isolated yields obtained by filtration of the precipitate. ^{*d*} 2.2 equivalents of **2a** were used. ^{*c*} 2.4 equivalents of **2a** were used. DABCO = 1,4-diazabicyclo[2.2.2]octane; DBU = 1,8-diazabicyclo[5.4.0]undec-7-ene; TMG = 1,1,3,3-tetramethylguanidine.

derivatives and chiral guanidine (for details, see Table S1 on page S19 of the ESI†). However, the results with respect to yields and enantioselectivity were quite disappointing. Consequently, the optimal conditions for the formation of complex bridged cyclic polycycle **3a** were identified to be: 0.15 mmol of **1a** and 2.2 equivalents of **2a** with 0.30 mmol of TMG in CH₃CN at 60 °C for 5 min (Table 1, entry 12).

With the optimized reaction conditions in hand, the substrate scope with respect to different enaminones 1 was first explored and the results are highlighted in Scheme 3. These transformations had a wide tolerance of arylamine-derived enaminones with different substitution patterns and electron properties, affording the corresponding products 3a-p in 51-97% yields within 5 minutes. Notably, when R¹ was located at the ortho-position of arylamine, new kinds of atropisomers 3mp were delivered in 51-88% yields, albeit with low diastereoselectivities. This promising finding not only provides a robust and facile method to access new atropisomers, but also represents a valuable complement to the existing appealing protocols.¹¹ Other than arylamine-derived enaminones, aliphatic amine-derived ones were also compatible, affording 3q-s in 24-41% yields with complete diastereomeric control. Particularly, furfuryl amine-derived enaminone also turned out to be a suitable reaction partner. Besides, acetylacetone and ethyl benzoylacetate derived acyclic enaminones could participate in this reaction successfully, thus facilitating the synthesis of 3u and 3v in 42% and 43% yields, respectively. An evaluation



Scheme 3 Scope of substrates. ^aIsolated yields obtained by filtration of the precipitate. ^bIsolated yields obtained by column chromatography.

of the substituent effect of R^2 on 3-nitropyridinium salts 2 revealed that not only were the methyl and ethyl groups accommodated well in this dearomative cascade cyclization, more hindered allyl, *n*-propyl and *n*-butyl groups were also amenable, enabling the construction of **3w-za** in good to excellent yields with complete diastereomeric control.

Later on, we investigated the effect of the substituents on the C3-position of pyridinium salts (Scheme 4). In sharp contrast to 3-nitropyridinium salt **2a**, replacement of the nitro group with cyano and trifluoromethyl resulted in different reactivities, in which only bifunctionalization occurred for **2g** and **2h** by regio-selective attack at C-2 and C-6 positions, affording bridged polycyclic *N*,*N*-ketals **3zb** and **3zc** in 50% and 41% yields, respectively.

After successfully maximizing the active sites of pyridinium salts by the dearomative cascade strategy, we further extend this reaction system to quinolinium salts. With the first attempt of the reaction between enaminone 1a and N-benzyl quinolinium salt 4a in CH₃CN at 60 °C, the trifunctionalized dearomatization product 5a, featured by an interesting fusion of bioactive bridged N,O-ketal and 1,2-dihydroquinoline, was precipitated out from the reaction mixture, and was obtained in 98% yield in a highly diastereoselective manner (Scheme 5). Although the product 5a contained four contiguous tertiary stereocenters including two bridgehead stereocenters, only one diastereoisomer was obtained. After flash silica gel column chromatography, 5a was partially decomposed into bifunctionalized product 6a through a ring-opening/closure sequence. This result indicated that 5a was not very stable and it might be sensitive to acid. So, we conducted a one-pot two-step reaction of 1a and 4a. After the first TMG-promoted three-component dearomative trifunctionalization went to completion, one



Scheme 4 Scope of substrates with respect to pyridinium salts 2 with different substituents at the C-3 position.

equivalent of trifluoroacetic acid was added to the reaction mixture, and the desired bifunctionalized bridged N,N-ketal product 6a was generated smoothly in 99% yield with excellent diastereoselectivity.

This appealing step-controlled diverse synthesis prompted us to further investigate the substrate scope for the diastereoselective dearomative bi- and trifunctionalization of quinolinium salts. As demonstrated in Scheme 6, all of the reactions went to full conversion within 5 minutes to afford a series of trifunctionalized products 5a-s in 61-99% yields. Moreover, the workup procedure was extremely simple. All of the products were precipitated out from the reaction mixture and only an easy filtration process was needed to purify them. These prominent merits including high synthetic efficiency, short reaction time and simple operation make this transformation very attractive.

When phenanthrolinium salt 4k and 3-methyl quinolinium salt 4j were used as substrates, only dearomative bifunctionalizations took place to afford 7 and 8 in 87% and 51% yields, respectively (Scheme 7). Next, we turned our attention to the one-pot two-step dearomative bifunctionalization of quinolinium salts. As summarized in Scheme 8, both enaminones 1 and quinolinium salts 4, irrespective of their substitution patterns and electron character, were tolerable, and participated in these transformations efficiently in a highly diastereomeric fashion, thus furnishing the corresponding products 6a-t in 79-99% yields.



Scheme 5 Step-controlled dearomative cascade cyclization of quinolinium salt 4a.



Scheme 6 Diastereoselective dearomative trifunctionalization of quinolinium salts 4



Scheme 7 Diastereoselective dearomative bifunctionalization of phenanthrolinium salt 4k and 3-methyl quinolinium salt 4j.

Finally, we moved on to examine the reactivity of isoquinolinium salt 9a. Likewise, the highly regio- and diastereomeric trifunctionalized product 10a was generated in 36% yield when 2.0 equivalents of TMG were used as the base (Table S2,† entry 1). To benefit the yield, we investigated the effect of various bases and their loadings (for detailed condition optimization, see Table S2[†] on the page S48 of the ESI[†]), and we identified that 3.6 equivalents of DBU were the best of choice. Subsequently, the substrate scope was explored by using a variety of enaminones 1 and isoquinolinium salts 9 (Scheme 9). Gratifyingly, all of them could participate in this transformation successfully to afford 10a-o in 82-97% yields. Remarkably, use of ortho-fluoro-substituted aniline-derived enaminone as the substrate led to the synthesis of atropisomer 10l in 90% yield with 1.6:1 dr. When 4-bromoisoquinolinium salt 9d was employed, the highly diastereoselective bifunctionalization occurred instead of the trifunctionalization to deliver 10p as a single diastereomer in 54% yield (Scheme 10). The difference in reactivities of 9d and 9a-c may be caused by the steric hindrance, which disfavored the subsequent addition with another isoquinolinium salt.

To demonstrate the synthetic robustness, a preparative-scale experiment was conducted with 2.0 mmol of enaminone 1d and



Scheme 8 Diastereoselective dearomative bifunctionalization of quinolinium salts 4.



Scheme 9 Diastereoselective dearomative trifunctionalization of isoquinolinium salts 9. ^aIsolated yields obtained by filtration of the precipitate. ^bIsolated yields obtained by column chromatography.

4.4 mmol of *N*-benzyl-3-nitropyridinium salt **2a** (Scheme 11). Delightfully, the reaction proceeded smoothly to give **3d** in diminished but acceptable yield within 5 minutes (0.89 g, 69%



Scheme 10 Diastereoselective dearomative bifunctionalization of 4-bromoisoquinolinium salt 9d.



Scheme 11 Scalable preparation of 3d.

yield), thus suggesting that this protocol has the potential for scale-up preparation.

To further highlight the synthetic applicability, some chemical transformations of 3g were carried out (Scheme 12). By subjecting 3g to the NaBH₄–NiCl₂ reductive system, the two nitro groups in 3g were reduced into hydroxylamine and oxime respectively, accompanied by four-membered all-carbon ring opening. This transformation went to completion within 10 min with the formation of the corresponding functionalized product **11** in 62% yield with a ratio of 10 : 1 Z/E. In addition, the presence of a bromine atom in 3g offers a convenient platform for further elaboration to generate molecules with more diverse and complex structures. Thus, two palladium-catalyzed Suzuki couplings of 3g with 4-chlorophenzyl boronic acid and 2-indolyl boronic acid were performed, which afforded **12** and **13** in 99% and 73% yields, respectively.

The structures and the relative configurations of **3a**, **3zb**, **5m**, **6n**, **10g**, **10p** and **11** were unequivocally verified by X-ray analysis.¹² The relative configurations of other products **3**, **5**, **6** and **10** were assigned by analogy.

Based on the experimental results, plausible mechanisms were proposed to rationalize the reaction pathways. As shown in



Scheme 12 Chemical conversions of 3g.

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Scheme 13 (a) Plausible reaction pathway for dearomative multifunctionalization of pyridinium salt 2a. (b) Possible mechanism for dearomative tri- and bifunctionalization of quinolinium salt 4a. (c) Proposed mechanism for the trifunctionalization of isoquinolinium salt 9a.

Scheme 13a, a Michael addition between enaminone 1a and 3nitropyridinium salt 2a occurred first to generate the bienamine intermediate A. Nucleophilic Michael attack at the C-4 position of another 3-nitropyridinium salt molecule 2a resulted in the formation of active iminium ion intermediate B containing a trienamine moiety, followed by an intramolecular Mannich reaction to yield the intermediate C. A final [2 + 2] cycloaddition led to four-membered all-carbon cyclobutane formation. As for the regio- and diastereoselective dearomative tri- and bifunctionalization of quinolinium salt 4a (Scheme 13b), similarly, a 1,4addition took place first to deliver intermediate D, which contained an enamine motif. Then, a Mannich reaction proceeded by regioselective nucleophilic attack at the C-2 position of another quinolinium salt 4a to generate intermediate E with an imine ion skeleton. Through electron transfer, the oxygen atom served as an active nucleophilic site to attack the imine ion via an intramolecular oxa-Mannich reaction, thus forming the

trifunctionalized bridged cyclic product **5a**. In the presence of TFA, **5a** was susceptible to transform into bifunctionalized product **8a** through an acid-catalyzed sequential ring-opening/ enamine–imine tautomerization/aza-Mannich process. For the construction of trifunctionalized tetrahydroisoquinoline product **10a**, a triple Mannich reaction of enaminone **1a** and *N*-benzyl isoquinolinium salt **9a** proceeded, which is shown in Scheme 13c.

Conclusions

In summary, we have successfully developed an unprecedented base-promoted multi-component one-pot dearomatization of pyridinium, quinolinium and isoquinolinium salts, which enabled the synthesis of complex and diverse bridged cyclic polycycles with multiple stereocenters in a highly regio- and diastereoselective manner with high synthetic efficiency. Besides, we realized the step-controlled dearomative bi- and trifunctionalization of guinolinium salts. These transformations not only achieved the maximization of the reactive sites of pyridinium, quinolinium and isoquinolinium salts to enhance structural complexity and diversity, but also opened up a new reaction mode of these N-activated azaarenes, which may largely benefit the prospective reaction designs. A unique feature of this strategy is the use of easily accessible and bench-stable N-alkyl activated azaarenes to provide maximum reactive sites for dearomative multi-component one-pot cascade cyclizations. In addition, the salient characteristics including high synthetic efficiency, short reaction time, mild conditions and simple operation made this strategy particularly attractive. We expect that this fascinating strategy will stimulate the design of new related reactions for the facile construction of natural products and biologically interesting compounds.

Conflicts of interest

There are no conflicts to declare.

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