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Oxidative Skeletal Rearrangement of 1,1'-Binaphthalene-2,2'-diamines (BINAMs) via C–C Bond Cleavage and Nitrogen Migration: A Versatile Synthesis of U-Shaped Azaacenes

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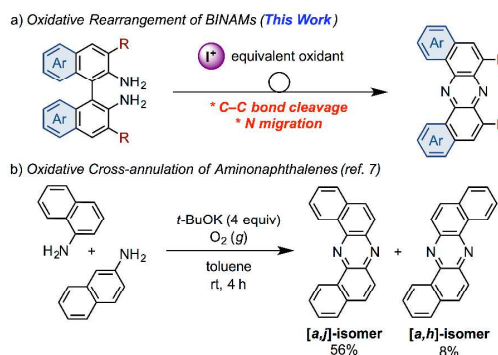
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An oxidative skeletal rearrangement of 1,1'-binaphthalene-2,2'-diamines (BINAMs) that involves the cleavage of a strong C–C single bond of the binaphthalene unit and the nitrogen migration has been discovered. The unprecedented rearrangement enables access to a series of U-shaped azaacenes otherwise difficult to prepare in a selective manner by classical methods. Moreover, physicochemical properties of the unique azaacenes have been comprehensively investigated.

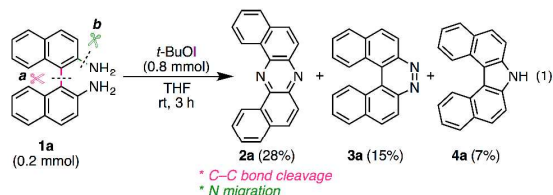
Azaacenes, in which parts of C=CH fragments of acenes are formally replaced with isoelectronic imine (C=N) moieties, have been emerging as promising candidates for electron-transporting (n-type) materials in organic electronics.¹ The introduction of electronegative nitrogen atoms into π -conjugated carbon frameworks allows for increasing electron affinities (EAs) to lower the barriers for electron injection from metal electrodes.² Furthermore, owing to the decrease in the number of C–H $\cdots\pi$ contacts, azaacenes show a high propensity for adopting densely stacked packing-structures in the solid states, which favor carrier-transporting process.³ However, due to limited synthetic accessibility of azaacenes, the exploration of azaacene-based materials has fatally lagged behind those of acenes. Therefore, the development of novel synthetic approaches to azaacenes is fundamentally important from the viewpoints of not only synthetic chemistry but also material sciences. Traditionally, syntheses of azaacenes have exclusively relied on condensation/oxidation protocols starting from *ortho*-arylenediamines and *ortho*-quinone derivatives.⁴ In this context, recently, a new synthetic protocol toward the preparation of azaacenes consisting of a Pd-catalyzed double amination of activated *ortho*-dichloroarenes with *ortho*-arylenediamines and a successive oxidation leading to *linear* azaacenes in an efficient manner, has been reported.⁵ On the other hand, Hiroto and Shinokubo have reported a DDQ-promoted oxidative

annulation of aromatic amines to selectively give *kinked* azaacenes and aza[7]helicenes depending on the conditions applied.⁶ Herein we present a new synthetic strategy to azaacenes based on a discovery of an unprecedented oxidative skeletal rearrangement of 1,1'-binaphthalene-2,2'-diamines (BINAMs) induced by an iodine-containing oxidant, which exclusively provides a series of dibenzo[*a,j*]phenazine-cored U-shaped azaacenes (Scheme 1a). As a relevant work, there is the single report that describes selective preparation of dibenzo[*a,j*]phenazine through an oxidative cross-annulation of α - and β -aminonaphthalenes mediated by a strong base (*t*-BuOK) under O₂ atmosphere (Scheme 1b).⁷ However, this known method intrinsically requires two regioisomers of aminonaphthalenes and suffers from the concomitant production of a constitutional [*a,h*]-isomer, thereby lacking the versatility as a synthetic approach to U-shaped azaacenes.



Scheme 1 Construction of dibenzophenazine-based U-shaped azaacenes

During our studies on the development of oxidative transformations of various amines utilizing iodine-containing oxidants,⁸ we serendipitously found out that the treatment of BINAM (**1a**) with *tert*-butyl hypoiodite (*t*-BuOI)⁹ produced dibenzo[*a,j*]phenazine (**2a**) in 28% yield along with the production of reasonably expected compounds **3a**¹⁰ (15%) and **4a**¹¹ (7%) [eqn (1)].



The structure of **2a** was unambiguously confirmed with the X-ray crystallographic analysis of its yellow single crystal (ESI[†]). Most importantly, the discovered transformation leading to **2a** formally involves the cleavage of the strong C(Ar)–C(Ar) bond [**a** in eqn (1)] of the binaphthyl unit (e.g., the bond dissociation energy (BDE) of Ph–Ph is as large as 118 kcal/mol)¹² without the aid of transition metal complexes¹³ and also involves the migration of a nitrogen atom [**b** in eqn (1)] to the adjacent carbon. Encouraged by this finding, we envisioned that the utilization of this unique rearrangement reaction would offer a versatile route to U-shaped azaacenes. To establish a novel approach to U-shaped azaacenes, reaction parameters were scrutinized to identify the conditions to selectively provide **2a** (ESI[†]). Table 1 summarizes the effect of halogen-containing oxidants. To our delight, the use of *N*-iodolactams as an oxidant like NIS, NIPy, and DIH in *t*-BuOH selectively gave **2a** in moderate yields along with trace amounts of **4a** without affording **3a**. The use of 8 mol equivalents of DIH gave **2a** in 77% yield, although the role of excess of DIH is not clear at present. In sharp contrast, when *N*-chloro-containing oxidant (*t*-BuOCl) was used, diaza[5]helicene **3a** was exclusively formed in a high yield (89%). Other types of oxidants like I₂, NBS, DDQ, PhI(OAc)₂, and MnO₂ failed to furnish **2a**.

Table 1 Effect of halogen-containing oxidants^{a,b}

Oxidant	2a (%)	3a (%)	4a (%)
NIPy	36	0	trace
NIS	47	0	4
NIPh	0	0	0
NISac	0	0	0
DIH	49	0	3
TIG	77 ^c	0	0
I ₂	0	0	0
NBS	0	0	0
<i>t</i> -BuOCl	0	89	0

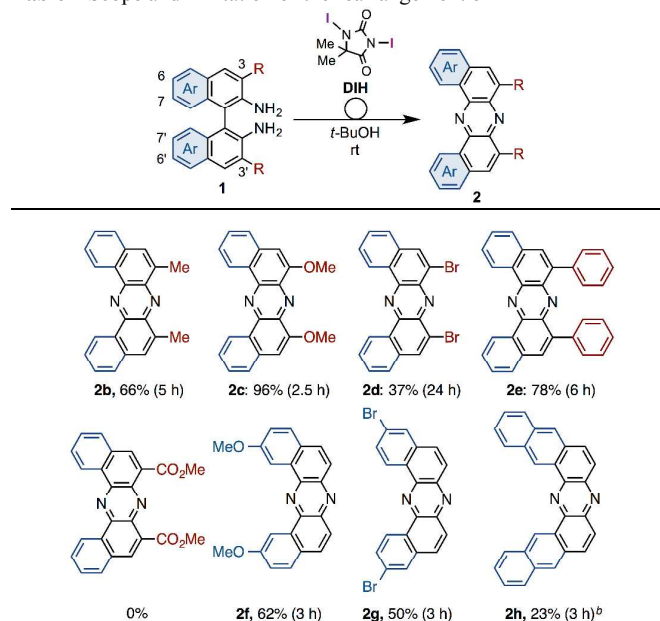
^a 0.2 mmol of DIH was used. ^{**} 1.6 mmol of DIH was used. ^{***} 0.2 mmol of TIG was used.

^a Reaction Conditions: **1a** (0.2 mmol) and oxidant (0.8 mmol) in *t*-BuOH (20 mL) at rt for 3 h. ^b Yields were determined by the integration of ¹H NMR charts of the crude products. ^c Isolated yield.

Having identified the optimal conditions for the rearrangement reaction, the scope and limitation of BINAM substrates were surveyed (Table 2). The treatment of BINAMs bearing two electron-donating substituents (Me and MeO) at the 3,3'-position with DIH successfully provided azaacenes **2b** and **2c** in 66% and 96% yield, respectively. The reaction using 3,3'-dibrominated BINAM **1d** gave desired product **2d** in a low yield, probably because of the poor solubility of the diamine in

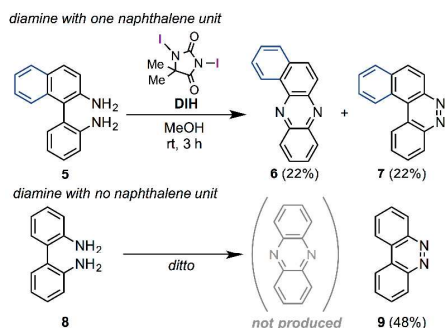
t-BuOH. Notably, the rearrangement of 3,3'-diphenyl diamine **1e** also efficiently proceeded to afford **2e** in a high yield, indicating that steric hindrance around the amino moieties does not affect the reaction efficiency. In sharp contrast, diamine bearing two ester groups (CO₂Me) at the 3,3'-position did not undergo the rearrangement at all. This significant lack of reactivity might be ascribed to the low nucleophilicity of amino moieties, inhibiting the formation of N–I bonds (*vide infra*). Diamines bearing the 7,7'- and 6,6'-disubstituted also underwent rearrangement to selectively provide **2f** and **2g** in moderate yields. Notably, the rearrangement of bianthracene diamine **1h** gave highly conjugated heptacyclic azaacene **2h**, which should be quite difficult to synthesize by conventional organic reactions.

Table 2 Scope and limitation of the rearrangement of **1a**^a

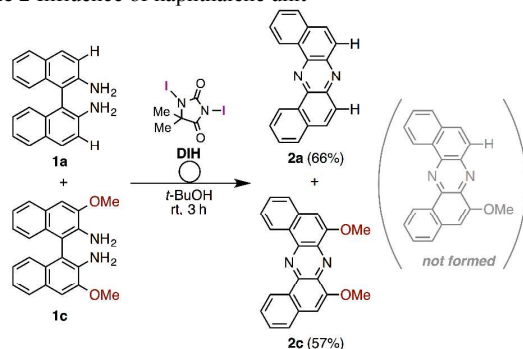


^a Reaction Conditions: **1** (0.2 mmol) and **DIH** (1.6 mmol) in *t*-BuOH (20 mL) at rt for the indicated time. ^b MeOH was used as a solvent.

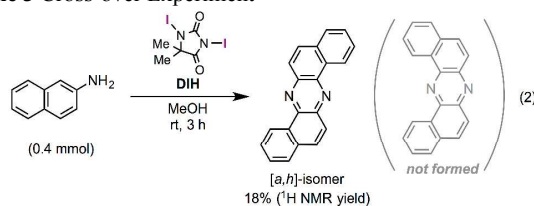
Regarding mechanistic aspects, importantly, the existence of at least one naphthalene unit is indispensable for the unique rearrangement (Scheme 2): when biaryldiamines bearing one (**5**) and none (**8**) of naphthyl unit were subjected to the reaction conditions, the ratios of rearranged product/benzocinnoline product were found to be positively correlated with the number (*n*) of naphthalene unit. These results might indicate the involvement of dearomatization processes in the reaction pathway (*vide infra*), which would be more favorable with naphthalene unit than benzene unit.¹⁴ On the other hand, crossover experiment applying an equimolar mixture of **1a** and **1c** resulted in no production of cross-product at all (Scheme 3), and a control experiment using β-naphthylamine under the optimal conditions led to the production of [*a,h*]-isomer without the formation of **2a** [eqn (2)], suggesting the *intramolecularity* of the rearrangement.



Scheme 2 Influence of naphthalene unit



Scheme 3 Cross-over Experiment



Although precise mechanism should await further investigation, in conjunction with the accumulated knowledge on the reactivities of aromatic amines toward electrophilic iodinating oxidants, a tentative reaction pathway is illustrated in Fig. 1. The reaction would start with exchange of N–H hydrogen(s) with iodine, leading to mono or di N-iodoamine **A**.⁸ Dearomatization process to form azirine **B** could explain the indispensability of naphthyl unit for the rearrangement.¹⁵ Upon **B** is formed, the highly strained intermediate would be attacked by an intramolecular amino nucleophile (Ar–NH₂), and the developing anion would be trapped by extra iodinating reagent to form **C**.¹⁶ Nitrene **D** could be generated from **C** driven by the release of its strain energy, and the subsequent insertion of nitrene to the C(Ar)–C(Ar) bond would give phenazine **2a**. Alternatively, the path from **C** to **2a** could be interpreted as a variant of Stieglitz rearrangement.¹⁷ Carbazole **4a** might be produced through [3,3]-sigmatropic rearrangement of diimine that would be formed by the protonation of **1a** with generating HI, and followed by the elimination of ammonia.¹¹

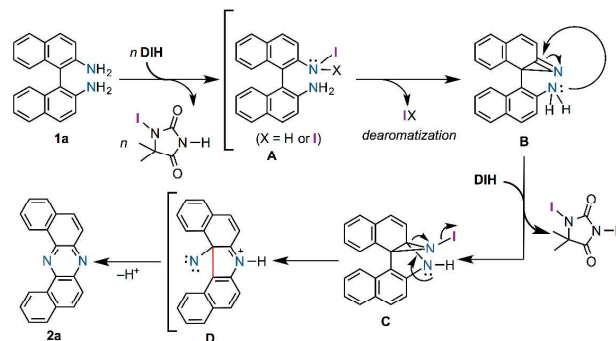
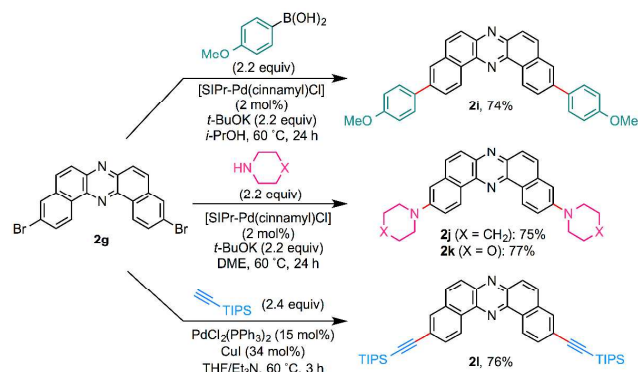


Fig. 1 A tentative pathway of the rearrangement

In light of bromo functionality of **2d** and **2g**, these compounds should serve as useful building blocks for functionalized azaacenes. Synthetic versatility of **2g** was clearly demonstrated by the preparation of **2i–2l** via Pd catalysis (Scheme 4). With slight modifications of the Nolan's original conditions,¹⁸ NHC/Pd-catalyzed cross-coupling of **2g** with an arylboronic acid and cyclic amines gave the corresponding coupled products **2i–2k** in good yields. Furthermore, TIPS-acetylene was efficiently cross-coupled with **2g** to afford conjugation-extended azaacene **2l** in 76% yield.

Scheme 4 Pd-catalyzed functionalization of **2g**

Basic physicochemical properties of U-shaped azaacenes **2** were investigated. UV/Vis and fluorescence (FL) spectra as well as a cyclic voltammogram for **2a** are shown in Fig. 2 as a representative example (for other compounds, see ESI[†]). The mirror-image form UV and FL spectra with fine vibrational structures and small Stokes shift (9 nm) reflect the rigid structure of **2a** like typical (aza-)acenes (Fig. 2a). Likewise, diluted CH₂Cl₂ solutions of other azaacenes **2** emit fluorescence ranging from blue ($\lambda_{em} = 425$ nm) to yellow ($\lambda_{em} = 561$ nm) depending on the substituents on the conjugated core. Especially, the introduction of sterically demanding and strongly electron-donating groups like piperidino and morpholino functionalities (i.e., **2j** and **2k**) resulted in significant Stokes shifts and greatly enhanced quantum yields (**2a**: $\lambda_{em} = 425$ nm, Φ_f 0.14; **2j**: $\lambda_{em} = 561$ nm, Φ_f 0.47; **2k**: $\lambda_{em} = 543$ nm, Φ_f 0.42), probably ascribed to intramolecular charge-transfer (ICT) emission. Notably, most of azaacenes **2** showed one pair of reversible redox waves at the potentials ranging from –1.76 to –1.98 V against the Fc/Fc⁺ redox couple

as shown in Fig. 2b, indicating the good electron-accepting abilities of **2**. These values are compatible to those of n-type organic materials used for organic luminescence diodes (OLEDs) devices like diarylanthrazolines¹⁹ and Alq₃.²⁰

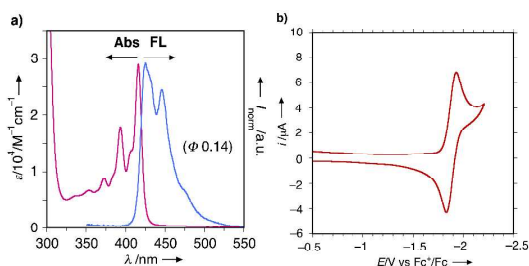


Fig. 2 a) UV/Vis absorption and FL spectra of CH₂Cl₂ solution of **2a** (10⁻⁵ M order). b) Cyclic voltammogram of CH₂Cl₂ solution of **2a** (10⁻⁴ M order) with nBu₄NPF₆ as an electrolyte.

In conclusion, we have discovered an unprecedented oxidative skeletal rearrangement of BINAMs and established an approach to constructing novel type of azaacenes that are otherwise difficult to synthesize by conventional methodologies. Investigation into the creation of functional conjugated materials based on these unique azaacenes are undergoing in our laboratory.

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† Electronic Supplementary Information (ESI) available: Synthetic procedures, spectroscopic data, NMR spectra, and physicochemical properties. CCDC-1004407. For ESI and crystallographic data in CIF or other electronic format, see DOI: 10.1039/b000000x/

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