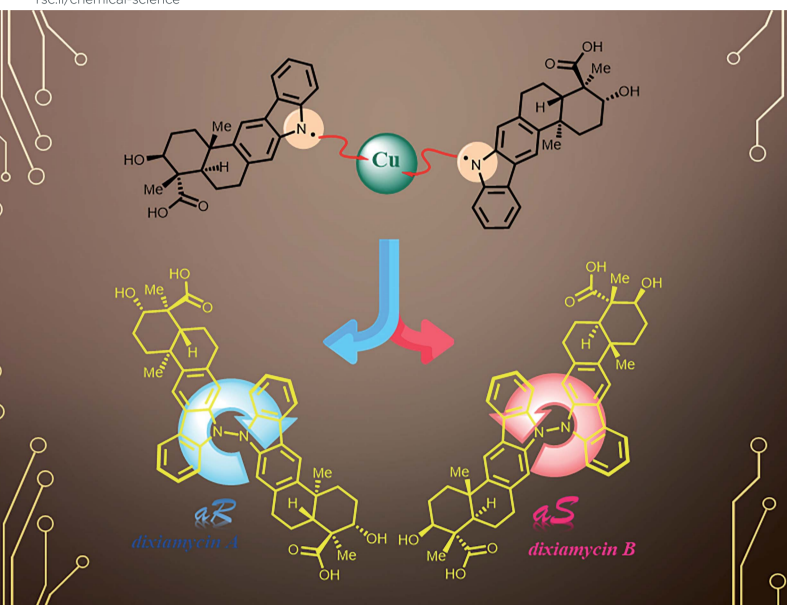
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# Total synthesis of atropisomeric indolosesquiterpenoids via N-N bond formation: dixiamycins A and B†‡

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N-N dimeric indolosesquiterpene alkaloids constitute a class of under-investigated architecturally intriguing natural products. Herein, we report the first chemical oxidation approach to the asymmetric total syntheses of these atropisomeric indoloses quiterpenoids through N-N bond formation. Specifically, dixiamycins A (1a) and B (1b) were prepared through a Cu(ı)-mediated aerobic dehydrogenative dimerization from the naturally occurring monomer xiamycin A methyl ester (2b); this preparation also represents the first total synthesis of dixiamycin A (1a). The monomer xiamycin A methyl ester (2b) was synthesized via a late-stage Buchwald Pd(II)-mediated aerobic dehydrogenative C-N bond formation.

#### Introduction

Pentacyclic indolosesquiterpene alkaloids (e.g., 1-3; Fig. 1), a novel and growing class of architecturally complex secondary metabolites, were first isolated from a range of Streptomyces species in 2010, exhibit important biological activities such as antimicrobial, antiviral, antitumor, immunomodulatory, and enzyme inhibitory activities, and are commonly referred to as the xiamycin family of alkaloids. In 2012, the atropisomeric indolosesquiterpenoid natural products dixiamycins A (1a) and B (1b) were isolated independently by Zhang and Hertweck.<sup>2,3</sup> These indole alkaloids each have a rare N-N linkage that generates an important class of atropisomers, featuring axial chirality about the N-N axis where the N atoms are sp<sup>3</sup>hybridized. Prior to this finding, the monomers of these dimers, i.e., xiamycin A (2a) and its methyl ester (2b), were isolated by Hertweck and coworkers from Streptomyces sp. GT2002/1503 (ref. 3a) and HKI0595,3b which are endophytes from the mangrove plants Bruguiera gymnorrhiza3a and Kandelia candel,3b respectively. Alkaloids 2a-b have been reported to display anti-HIV and antibiotic activities.<sup>3a</sup> Structurally, these alkaloids include a pentacyclic framework with four contiguous

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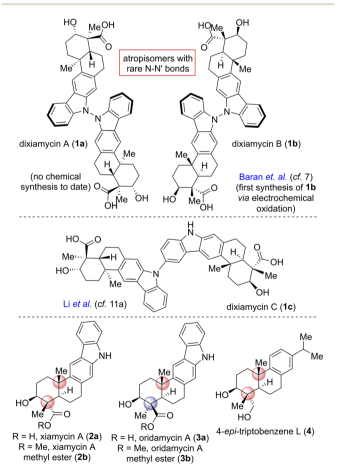


Fig. 1 Indolosesquiterpene alkaloids 1-3

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stereogenic centers at the periphery of a *trans*-decalin scaffold to which is attached a carbazole unit.<sup>4,5</sup> Oridamycin A (3a) is a diastereomer of xiamycin A (2a) and has been isolated from *Streptomyces* sp. KS84.<sup>4b</sup>

Importantly, two out of the four stereogenic centers are synthetically challenging all-carbon quaternary centers. The new uninvestigated N-N dimeric forms of xiamycin A are nearly an order of magnitude more potent than their monomeric scaffolds.2 The emerging recognition of the biological activity of these indoloses quiterpenoids has drawn considerable attention from the synthetic community, which has produced several syntheses of these alkaloids. The most efficient approach to the atropo-diastereomers of xiamycin A would be to effect a direct oxidation to form an N-N bond from the monomer as per the proposed biosynthesis following a single-electron transfer (SET) mechanism (Scheme 1). However, given that there has been rather limited success in achieving a direct N-N bond formation of two carbazole units with chemical oxidants (such as the use of stoichiometric amounts of oxidants I2,6a KMnO4,6b Ag2O,6c or dichromate<sup>6d</sup>), the synthesis of dixiamycins A (1a) and B (1b), bearing sensitive functionalities, remains a formidable challenge. In this regard, an electrochemical oxidation approach reported by Baran and coworkers elegantly addressed the total synthesis of dixiamycin B (1b).7

Based on the biosynthesis proposal (Scheme 1) and a recent report on aerobic oxidation by Stahl and coworkers,8 we envisioned that a suitably protected xiamycin A might be made to engage, under Cu(1)-catalysis, in an oxidative N-N bond formation to provide both 1a and 1b. Our efforts began with identifying a practical synthesis of xiamycin A methyl ester (2b). Prior elegant approaches to monomeric indolosesquiterpene alkaloids have been independently developed by Baran (cyclization of a carbazole-anchored epoxy ether),7 Krische (TiCl4promoted Friedel-Crafts cyclization),9 Trotta (radical-induced polyene cyclization), 10 Li11 (6π-electrocyclization/aromatization and indole C2-H bond activation/Heck annulation), Sarpong12 (from (R)-carvone, using a photoinduced benzannulation sequence to forge the carbazole core), Dethe<sup>13</sup> (oxidative Heck/ aromatization for carbazole synthesis) and our group (following δCsp<sup>3</sup>-H activation of the pentacyclic skeleton of indolosesquiterpene alkaloid).14

Herein, we report the total synthesis of both N-N atropodiastereomers of indolosesquiterpene alkaloids, namely dixiamycins A (1a) and B (1b), *via* a nature-inspired aerobic oxidation. Impressively, this work represents, to the best of our

Scheme 1 Biosynthesis of dixiamycins A (1a), B (1b) and C (1c).

Scheme 2 Hypothesized accessing of atropisomers of dixiamycins A (1a) and B (1b) and retrosynthetic plan (via 4-epi-triptobenzene L (4)).

knowledge, the first total synthesis of dixiamycin A (1a), clearly consistent with nature's oxidative approach to dixiamycins. Moreover, the currently developed unified practical asymmetric approach to naturally occurring abietane diterpenoids (*via* a key epoxy-ene cyclization<sup>15–18</sup>) and the congeners of the xiamycin family (*via* a late-stage Buchwald oxidative C–N bond formation<sup>19</sup>) would enable access to these natural products in significant quantities.

#### Result and discussion

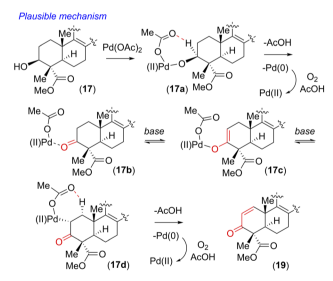
On the basis of their structural similarity to naturally occurring diterpenoids, such as 4-epi-triptobenzene L (4), we envisioned a unified approach to the xiamycins (Scheme 2). Retrosynthetically, we imagined accessing highly functionalized obromo nitroarene 7 as an advanced intermediate for our synthesis. Thus, a late-stage Buchwald oxidative C-N bond formation of 2-phenyl acetanilide 6 could construct the

**Scheme 3** Total syntheses of 4-epi-triptobenzene L (4) via polyene cyclization.

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10 mol%  
Pd(OAc)<sub>2</sub>  
O<sub>2</sub> (1 atm.)  
DMSO  
120 °C, 18 h  
[O] Me O (+)-17 NAC  
Me O (+)-18  
Me O (+)-18  
$$\Delta^{2.3}$$
; (+)-19

Scheme 4 Synthesis of a pentacyclic framework under oxidative C-N bond formation



Scheme 5 Plausible mechanism involving overoxidation

carbazole moiety of the xiamycins. o-Bromo nitroarene 7 could be synthesized from o-bromo isopropylarene 8 using an ipsonitration, which in turn could be synthesized from 13 using Sharpless asymmetric epoxidation followed by Lewis-acidassisted epoxy-ene cyclization (via intermediate 4).

Our synthesis commenced with a Cu-catalyzed Csp<sup>3</sup>-Csp<sup>3</sup> bond formation between functionalized geranyl acetate 10 and 3-isopropylbenzyl bromide 11, followed by TBS deprotection using TBAF to access allylic alcohol 13 (Scheme 3). Sharpless

Scheme 6 Total synthesis of xiamycin A (2a) via Buchwald's oxidative C-N bond formation.

Table 1 Attempts towards oxidative N-N bond formation

dixiamvcin A methyl ester (24a) conditions xiamycin A dixiamycin B methyl ester (2b) methyl ester (24b) Results Entry Conditions NaH, THF, O<sub>2</sub> (1 atm) 1. 92% SM recovered -78 °C to 25 °C O2 (1 atm), DCE 2. 89% SM recovered 25 °C to 70 °C multiple spots + 3. I<sub>2</sub>, KI, K<sub>2</sub>CO<sub>3</sub>, O<sub>2</sub> (1 atm) 73% SM recovered DMSO, 25 °C to 100 °C decomposition + PIDA, HFIP, 40 °C 4. multiple spots Nal, NalO<sub>4</sub> 5. multiple spots CH<sub>3</sub>CN, 25 °C KI, KIO₄ no conversion 6. CH<sub>3</sub>CN, 25 °C Me <sup>n</sup>BuLi, THF -78 °C several Mé conditions Me MeÓ xiamycin A MeÓ methyl ester (2b) I<sub>2</sub>, THF -78 °C multiple spots 14% dixiamycin A & B methyl ester mixture 37% SM recovered

asymmetric epoxidation15 of 13 with TBHP in the presence of Ti(OiPr)<sub>4</sub> and (+)-DET afforded epoxy-alcohol 14 with 95% ee in 92% yield. Treatment of 14 under Lewis-acid-assisted epoxy-ene cyclization completed the total synthesis of naturally occurring

**Table 2** Optimization of Cu(i)-catalyzed oxidative N-N bond formation a,b

S. no.	Condition	Solvent	Time	23a/23b	22
1	30 mol% Cu(ı)I:L1	DCE	48 h	19% (~1:1)	62%
2	20 mol% Cu(ı)Br:L1	DCE	48 h	42% (~1.1:1)	39%
3	20 mol% Cu(1)Br:L2	DCE	48 h	$51\% (\sim 1.2:1)$	31%
4	20 mol% Cu(1)Br:L3	DCE	48 h	23% (~1.1:1)	56%
5	20 mol% Cu(ı)Br:L4	DCE	48 h	46% (~1.2:1)	32%
6	20 mol% Cu(ı)Br	DCE	17 h	68% (~1.6:1)	_
	40 mol% DMAP			` ′	
7	20 mol% Cu(ı)Br	MeCN	17 h	49% (1.6:1)	_
	40 mol% DMAP			, ,	
	R = H, L1 N R = t-Bu, L2				

 $^a$  All the reactions were carried out in the presence of an  $\rm O_2$  balloon (1 atm).  $^b$  Yields are reported as isolated yields after column chromatography.

4-epi-triptobenzene L (4) bearing a *trans*-decalin motif and four contiguous stereogenic centers in 63% yield (Scheme 3).<sup>16</sup> The synthesized 4-epi-triptobenzene L (4) contained all the requisite stereogenic centers for the indolosesquiterpene alkaloids in place.<sup>17</sup> Furthermore, a chemoselective oxidation of the 1,3-diol in the presence of PIDA and TEMPO followed by Pinnick oxidation and methylation with dimethyl sulfate (Me<sub>2</sub>SO<sub>4</sub>) furnished ester 9 in 76% yield (Scheme 3). Next, using the protocol described above, a mass of more than 10 g of compound 9 was prepared.

Next, aromatic electrophilic bromination of **9** with *N*-bromo succinimide (NBS) in acetonitrile afforded **8** in 92% yield

Scheme 7 Total syntheses of dimeric indolosesquiterpene alkaloids 1a-b.

$$\begin{array}{c} \text{TMSOTf, DCE, 0 °C, 1 h} \\ \text{then 20 mol% Cu(l)Br.Me}_2S \\ 40 \text{ mol% DMAP, DCE} \\ O_2 \text{ (1 atm.), 70 °C, 18 h} \\ \text{then K}_2\text{CO}_3, \text{MeOH, 2 h} \\ \text{ester (2b)} \\ \hline \\ \text{20 mol% Cu(l)Br.Me}_2S \\ 40 \text{ mol% DMAP, DCE} \\ \text{40 mol% DMAP, DCE} \\ O_2 \text{ (1 atm.), 70 °C, 72 h} \\ \text{methyl ester (2b)} \\ \hline \\ \text{42% (ca. 1.6:1)} \\ \hline \\ \text{42% (ca. 1.6:1)} \\ \text{(24a) + (24b)} \\ \hline \\ \text{(24b)} \\ \hline \\ \text{(24a) + (24b)} \\ \hline \\ \text{(24b)} \\ \hline \\ \text{(24a) + (24b)} \\ \hline \\ \text{(24b)} \\ \hline \\ \text{(24$$

Scheme 8 A direct conversion of xiamycin A methyl ester (2b) into dimeric indolosesquiterpene alkaloids 1a-b.

(Scheme 4). Thus, the stage was set for the ipso-nitration,18 which was investigated under various conditions. Gratifyingly, the ipso-nitration of the isopropyl group of 8 was realized with fuming  $HNO_3$  at -40 °C. Carrying out a nitration of the hydroxyl group afforded nitrite derivative 15 in 69% yield, with the identity of this derivative confirmed by X-ray analysis of its Suzuki-Miyaura coupling product (after recrystallization). Reduction of the nitro groups of 16 (10 mol% Pd-C under 1 atmosphere of H<sub>2</sub> gas) followed by the treatment with Ac<sub>2</sub>O afforded acetanilide 6 in 92% yield over 2 steps (Scheme 4). At this point we were in a position to test the Buchwald<sup>19</sup> oxidative C-N bond formation of acetanilide 6 to craft the pentacyclic core of xiamycin A (Scheme 4). Since molecular oxygen is a quintessential oxidizing agent that performs several dehydrogenative processes in the presence of substoichiometric amounts of high-valence transition-metal complexes and has important implications for the large-scale preparation of **Edge Article Chemical Science** 

pharmaceuticals and fine chemicals, we shifted our attention to a transition-metal-free process under an oxygen atmosphere.

Our initial attempt using a combination of 10 mol% Pd(OAc)<sub>2</sub> and 10 mol% Cu(OAc)<sub>2</sub> in toluene at 110 °C under an oxygen atmosphere was disappointing as acetanilide 6 provided a multitude of spots on the corresponding TLC with no expected product. We did isolate a 32% yield of pentacyclic ketone 18 and 19% yield of enone 19 along with 31% yield of the starting material acetanilide 6 (Scheme 4). Thus, a quick optimization using different solvents, oxidants and Pd(II) source was carried out; it revealed that DMSO was a good choice, where complete consumption of starting acetanilide 6 was observed. Interestingly, we observed a three-fold oxidative process of 6 in the presence of 10 mol% Pd(OAc)2 in dimethylsulfoxide (DMSO) under an oxygen atmosphere (1 atm), without the use of a transition-metal based oxidant,20 to afford an 83% isolated yield of enone 19 (Scheme 4).19,21 The same efficiency was observed when the oxidative reaction was carried out using 10 mol% Pd(TFA)<sub>2</sub> under an O<sub>2</sub> atmosphere (1 atm) in DMSO (85% yield of isolated enone 19). In the presence of Pd and  $O_2$ , the secondary alcohol of 17 was proposed to have oxidized to ketone 18 (Scheme 5). Next, in a one-pot reaction, ketone 18 oxidized to corresponding  $\alpha,\beta$ -unsaturated enone 19. Here, the reaction may have proceeded via a C- and/or O-bound Pdenolate (see, 17a-c in Scheme 5). A β-hydride elimination was thought to have occurred via C-bound enolate to result in enone 19 (Scheme 5).

Since the secondary alcohol was responsible for the overoxidation under Pd(II)-catalysis, it was protected with an acetate group. Hence, compound 16, having an aromatic nitro group as well as a nitrite functionality, was reduced under hydrogenation conditions followed by a reaction with acetic anhydride to afford a 97% yield of acetanilide 20 (Scheme 6). Gratifyingly, 2-phenyl acetanilide 20 in the presence of 10 mol% Pd(OAc)2 in dimethylsulfoxide (DMSO) under an oxygen atmosphere afforded N,Obis-acetylated pentacyclic core 21 in 87% isolated yield, without the use of a transition-metal-based oxidant (Scheme 6).20

Next, deacetylation of compound 21 upon treatment with K<sub>2</sub>CO<sub>3</sub> in MeOH completed the synthesis of xiamycin A methyl ester (2b) in 96% yield. Furthermore, saponification of 2b using a mixture of KOH and LiOH in a MeOH/H2O mixture at reflux completed the synthesis of xiamycin A (2a), thus setting the stage for the dimerization via the key dehydrogenative N-Nbond-forming reaction.<sup>22</sup>

To investigate the chemical feasibility of the biosynthetic hypothesis, we next pursued the key dehydrogenative N-Nbond-forming dimerization of xiamycin A (2a) and xiamycin A methylester (2b) using various oxidants such as I<sub>2</sub>, <sup>6a</sup> KMnO<sub>4</sub>, <sup>6b</sup> Ag<sub>2</sub>O,6c dichromate,6d PIDA, KI or atmospheric O<sub>2</sub>. Each of these conditions, however, simply led to decomposition or no conversion. Interestingly, while using  $I_2$  and <sup>n</sup>BuLi at -78 °C to 25 °C for 72 h,23 xiamycin A methylester (2b) afforded 14% yields of dixiamycin A and B methyl esters (24a and 24b) (dr ca. 1:1), 37% yield of recovered starting material along with multiple TLC spots, probably arising from oxidative C-N and C-C bond formations (Table 1). This result was indeed encouraging, and thus a variety of other oxidative conditions were tested to effect

a chemoselective N-N bond formation for dixiamycins A (1a) and B (1b).

After testing various conditions for this goal of chemoselective N-N bond formation, we turned our attention to aerobic oxidation employing Cu(1)-catalysis as reported by Stahl and coworkers.8 In this regard, our attempt at using natural product xiamycin A (2a) under the Stahl conditions (20 mol% Cu(1) Br·Me<sub>2</sub>S and 40 mol% DMAP in dichloroethane at reflux for 17 h)8 led to decomposition. We found compound 2a to be a challenging substrate due to its several functional groups. To meet this challenge, we decided to use xiamycin A methyl ester TMS ether (22) as a starting material where the carboxylic acid and secondary alcohol of xiamycin A 2a were each in a protected form. Using the Cu(1)-catalyzed oxidative coupling conditions for 22 afforded promising N-N dimeric compounds 23a and 23b.

A detailed optimization of the Cu(1)-catalyzed aerobic oxidation of 22 is shown in Table 2. Following optimization, we found that treatment of 22 with 20 mol% Cu(1)Br and 40 mol% DMAP in DCE at reflux gave a 68% yield (ca.  $\sim$ 1.6:1 mixture) of the TMS ethers of dixiamycins A methyl ester (23a) and B methyl ester (23b) (entry 6). Other bidentate ligands (L1-L4) were found to be inferior to the DMAP ligand, affording the products in only 23-51% yields along with decomposition side products (entries 2-5). Note that the TMS protecting group in 22 survived under the reaction condition at 70 °C. These N-N atropodiastereomers were found to be separable using column chromatography. Following treatment with K2CO3 in MeOH, the atropo-diastereomers were smoothly converted to the corresponding methyl esters of dixiamycins A and B, i.e., 24ab (Scheme 7). Furthermore, saponification of 24a-b with KOH and LiOH in MeOH/H<sub>2</sub>O at 80 °C for 12-16 h completed the total syntheses of dixiamycins A (1a) and B (1b) without an event (Scheme 7). The atropo-diastereomeric natures of dixiamycins A (1a) and B (1b) were confirmed using a detailed HPLC analysis and their retention times were found to differ  $[^tR]$  of dixiamycin  $A(1a) = 8.8 \text{ min and } {}^{t}R \text{ of dixiamycin B} (1b) = 9.8 \text{ min}] (\text{see ESI}^{\ddagger}_{+})$ for detailed HPLC analysis).

To demonstrate the versatility of our synthesis, TMS protection was used as an orthogonal protecting group. In a one-pot operation (Scheme 8), 2b was treated with TMSOTf in DCE followed by application of our optimal N-N-bond-forming conditions and subsequent TMS deprotection using K2CO3 in MeOH; this procedure afforded a direct approach to dixiamycins A and B methyl esters in 66% yield in a ca. 1.6:1 ratio (Scheme 8). In addition, Cu(1)-catalyzed aerobic oxidation of xiamycin A methyl ester (2b) afforded atropo-diastereomers 24a-b in 42% yield upon prolonged heating (72 h) along with 31% recovered 2b, consistent with the hypothesis of nature's oxidative approach to the dixiamycins.

#### Conclusions

In conclusion, a total synthesis of N-N atropo-diastereomers of indolosesquiterpene alkaloids, namely dixiamycins A (1a) and B (1b), was developed. This synthesis involved a key Cu(1)-catalyzed aerobic oxidation of xiamycin A methyl ester (2b), culminating in the first total synthesis of dixiamycin A (1a) bearing

the rare N–N bond. The monomer of this approach, xiamycin A methyl ester (2b), was synthesized from a naturally occurring diterpenoid, namely 4-*epi*-triptobenzene L (4), following key Buchwald oxidative C–N-bond-forming reactions to craft the pentacyclic core of indolosesquiterpene alkaloid 2b. Thus, this synthesis featured the demonstration of two key aerobic oxidations, namely Pd(II)-(oxidative C–N bond formation) and Cu(I)-catalyzed (oxidative N–N bond formation) processes for the syntheses of, respectively, monomeric and dimeric indolosesquiterpene alkaloids. Our study also confirmed the high stability of the dixiamycins A (1a) and B (1b) under elevated temperature and basic conditions, suggesting that N–N atropodiastereomers of the xiamycins could represent viable starting points for potential use in pharmaceuticals and agrochemicals.

### Data availability

Data supporting this article have been uploaded as ESI.‡

#### Author contributions

A. B. conceived and supervised this project. R. N. investigated the key oxidative N–N bond-formation leading to dixiamycins A and B. R. N., S. N., and S. K. investigated the catalytic polyene cyclization and synthesized all the starting materials. S. N., M. M, V. R. G, and R. M. investigated the oxidative C–N bond-formation to craft the pentacyclic core of xiamycin A methyl ester. A. B. and R. N. wrote the original draft of the manuscript which was edited by all authors.

## Conflicts of interest

There are no conflicts to declare.

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