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Organocatalytic atroposelective synthesis of axially chiral N,N'-pyrrolylindoles *via de novo* indole formation†

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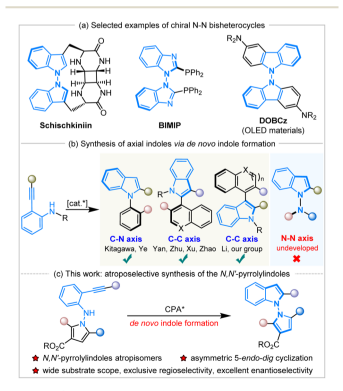
The first organocatalytic atroposelective synthesis of axially chiral N,N'-pyrrolylindoles based on o-alkynylanilines was successfully established via de novo indole formation catalyzed by chiral phosphoric acid (CPA). This new synthetic strategy introduced CPA-catalyzed asymmetric 5-endo-dig cyclization of new well-designed o-alkynylanilines containing a pyrrolyl unit, resulting in a wide range of axially chiral N,N'-pyrrolylindoles in high yields with exclusive regioselectivity and excellent enantioselectivity (up to 99% yield, >20:1 rr, 95:5 er). Considering the potential biological significance of N-N atropisomers, preliminary biological activity studies were performed and revealed that these structurally important N,N'-pyrrolylindoles had a low IC₅₀ value with promising impressive cytotoxicity against several kinds of cancer cell lines. DFT studies reveal that the N-nucleophilic cyclization mediated by CPA is the rate- and stereo-determining step, in which ligand-substrate dispersion interactions facilitate the axial chirality of the target products.

Introduction

N–N atropisomers are widely present as core structures in a variety of important chiral molecules, such as natural products, chiral ligands and functional materials (Scheme 1a).¹ In sharp contrast to the well-developed syntheses of C–C and C–N axially chiral compounds,² the construction of N–N atropisomers has remained in an underdeveloped state in the decades since their discovery in 1931.³ The asymmetric catalytic synthesis of N–N axially chiral compounds was considered a challenging issue until 2021, when Li,⁴a Lu and Houk,⁴b and Liu's⁵a groups developed various synthetic methods to access these fascinating molecules promoted by chiral transition metals or organocatalysis.

Subsequently, a great deal of attention has been paid to the development of efficient synthetic strategies and some progress has been achieved, mainly involving direct N-H

functionalization,⁴ the functionalization of prochiral or racemic bis-heteroaryl rings,⁵ *de novo* ring formation *via* a Paal–Knorr reaction,⁶ or dual-ring formation by cyclization and oxidation.⁷



Scheme 1 Selected examples of N-N atropisomers and asymmetric de novo construction of indole atropisomers.

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In addition, the catalytic atroposelective synthesis of N-N axially chiral molecules via de novo indole formation has emerged as an attractive route.8 The 5-endo-dig cyclization of o-alkynylanilines is a versatile method for the de novo generation of indole derivatives, and a variety of chiral indole atropisomers9 have been well established through this catalytic enantioselective cyclization (Scheme 1b). To our knowledge, the documented protocols rely heavily on asymmetric transition metal catalysis, and there are only two examples using organocatalysis for the construction of axially chiral indoles via the 5-endo-dig cyclization of o-alkynylanilines. In 2019, Yan and workers developed chiral quinine-derived thiourea-catalyzed asymmetric cyclization of o-alkynylanilines to give chiral naphthyl-C2-indoles with a C-C axis via vinylidene orthoquinone methide intermediates with excellent results.9c Meanwhile, Ye and coworkers realized chiral CPA-catalyzed atroposelective 5-endo-dig cyclization of ynamides to access chiral N-arylindoles with a C-N axis in high yields with excellent enantioselectivities.9d However, the approach to accessing axially chiral N-N bis-heteroaryl atropisomers through the de novo formation of chiral indole rings by the organocatalytic annulation of o-alkynylanilines remains undeveloped and unprecedented. During the preparation of this manuscript, Sparr and coworkers reported an asymmetric Pd-catalyzed 5-endo-hydroaminocyclization for the synthesis of N-N atropisomeric bisindoles and indolyl-carbazoles with moderate results.9j

Herein, we have designed and synthesized a new type of molecule, *N*-pyrrole *o*-alkynylanilines, as the substrates, and applied them in Brønsted-acid CPA-catalyzed atroposelective 5-endo-dig cyclization to achieve the de novo formation of axially chiral *N*,*N*'-pyrrolylindoles (Scheme 1c). It should be noted that this is the first organocatalytic annulation of *o*-alkynylanilines to prepare N-N axially chiral compounds in a practical and atom-economic manner. This fascinating protocol also faces a lot of challenges, including: (1) the development of new and efficient synthetic strategies to access N-N axially chiral molecules with excellent enantioselective control, (2) the design and synthesis of new kind of substrates and chiral N-N bisheteroaryl pyrrolylindoles atropisomers, (3) the discovery and application of these highly-valuable molecules in the field of synthetic and medicinal chemistry.

Results and discussion

Optimization of reaction conditions

To verify this hypothesis, an initial investigation of the asymmetric 5-endo-dig-cyclization was begun with *N*-pyrroloacetylaniline **1a** as the model substrate promoted by chiral phosphoric acids. A series of BINOL-based chiral phosphoric acids **CPA1-CPA8** as the catalysts were firstly applied to promote this cyclization in CHCl₃. Except for **CPA3** with a 1-naphthyl group, the formation of the expected **2a** with an N-N axis could be obtained in high yields but with poor enantioselectivity (Table 1, entries 1–8). H₈-BINOL-based chiral phosphoric acids **CPA9-CPA10** were also employed but, unfortunately, no further improvement was obtained (Table 1, entries 9 and 10). We found that SPINOL-derived **CPA11-CPA14** could promote

smooth cyclization, and the desired products could be obtained with better results (90-96% yields, 74:26-89:11 er, Table 1, entries 11-14). Pleasingly, SPINOL-derived CPA12 containing a 1-pyrenyl substituted group furnished the best results in terms of reactivity and enantioselectivity (93% yield, 89:11 er, Table 1, entry 12). A further solvent survey in the presence of CPA12 revealed that toluene is the most favorable choice, and the enantioselectivity of product 2a was greatly improved to 91:9 er (Table 1, entry 19). When the reaction temperature was gradually decreased to -10 °C, excellent enantioselectivity was obtained (90% yield, 95:5 er, Table 1, entry 22). It was found that the ee value was maintained without improvement at -40 °C, but lower reactivity was observed with only 50% yield (Table 1, entry 23). Accordingly, it was identified that the optimized the 5-endo-dig-cyclization of N-pyrroloacetylanilines should be conducted under conditions with SPINOL-derived CPA12 as the privileged catalyst in toluene at -10 °C.

Substrate scope study

With the established optimal conditions in hand, we focused on an evaluation of the substrate generality of N-pyrroloacetylanilines for this CPA12-catalyzed 5-endo-dig-cyclization. As summarized in Table 2, a wide range of N-pyrroloacetylanilines with different substituted groups on the phenyl ring and indole ring were applied to prepare chiral pyrrolyindoles containing an N-N axis. It was found that the electronic properties and positions of the substituted groups have a negligible impact on the reactivity and enantioselectivity. The N-pyrrolo-acetylanilines bearing electron-rich (1b-1d) or electron-deficient (1e) groups on the indole ring were examined first, and the expected annulation products (2b-2e) were furnished in 87-97% yields with 91:9-95:5 er. In addition, the Npyrrolo-acetylanilines (1f-1j) containing different substituted groups on the phenyl ring of the aniline motifs could be well accommodated to give the desired products (2f-2j) in good to high yields with excellent enantioselectivities (81-97% yields, 91:9-95:5 er). The absolute configuration of cycloadduct 2b was unambiguously determined as S by X-ray diffraction analysis (CCDC 2279386).10

On the other hand, we also investigated the feasibility of the CPA12-catalyzed 5-endo-dig-cyclization acetylanilines with different substituted groups on the phenyl ring in the pyrrole motif. It was found that a wide range of Npyrrolo-acetylanilines (1k-1t) bearing electron-rich groups or electron-deficient groups on the phenyl ring worked as good reaction partners, and this cyclization proceeded smoothly, resulting in the corresponding chiral products (2k-2t) in good to high yields with excellent enantioselectivities (87-96% yields, 87:13-95:5 er). It is noteworthy that the position of the substituted group did not affect the reactivity or enantioselectivity. Additionally, the heteroaryl-fused substrate 1u was well tolerated to deliver the corresponding product 2u in 90% yield with 90:10 er. Remarkably, the alkyl-substituted substrate 1v also worked well, leading to the expected product 2v in high yield with good enantioselectivity (97% yield, 95:5 er) (Table 3).

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Table 1 Optimization of reaction conditions^a

| Entry | CPA | Solvent | T (°C) | Yield ^b (%) | ee ^c (%) |
|-------|-------------------|---------------------|--------|------------------------|---------------------|
| 1 | (R)-C PA1 | CHCl_3 | RT | 96 | 50:50 |
| 2 | (R)-CPA2 | $CHCl_3$ | RT | 95 | 54:45 |
| 3 | (R)-CPA3 | $CHCl_3$ | RT | Trace | _ |
| 4 | (R)-C PA4 | $CHCl_3$ | RT | 93 | 50:50 |
| 5 | (R)-C PA5 | $CHCl_3$ | RT | 95 | 50:50 |
| 6 | (R)-C PA6 | $CHCl_3$ | RT | 96 | 61:39 |
| 7 | (R)-C PA 7 | $CHCl_3$ | RT | 90 | 50:50 |
| 8 | (R)-C PA8 | $CHCl_3$ | RT | 90 | 69:31 |
| 9 | (R)-C PA9 | $CHCl_3$ | RT | 95 | 65:35 |
| 10 | (R)-CPA10 | $CHCl_3$ | RT | Trace | _ |
| 11 | (S)-CPA11 | CHCl_3 | RT | 90 | 76:24 |
| 12 | (S)-CPA12 | $CHCl_3$ | RT | 93 | 89:11 |
| 13 | (S)-CPA13 | $CHCl_3$ | RT | 94 | 74:26 |
| 14 | (S)-CPA14 | CHCl_3 | RT | 96 | 84:16 |
| 15 | (S)-CPA12 | EtOAc | RT | Trace | _ |
| 16 | (S)-CPA12 | THF | RT | Trace | _ |
| 17 | (S)-CPA12 | $\mathrm{CH_{3}CN}$ | RT | 92 | 60:40 |
| 18 | (S)-CPA12 | Acetone | RT | Trace | _ |
| 19 | (S)-CPA12 | Toluene | RT | 95 | 91:9 |
| 20 | (S)-CPA12 | Toluene | 10 | 95 | 92:8 |
| 21 | (S)-CPA12 | Toluene | 0 | 93 | 93:7 |
| 22 | (S)-CPA12 | Toluene | -10 | 90 | 95:5 |
| 23 | (S)-CPA12 | Toluene | -40 | 50 | 95:5 |

^a All reactions were carried out with 0.2 mmol 1a in 2 mL of solvent catalyzed by CPA (10 mol%). ^b Isolated yield of the two steps overall. ^c The er value was determined by chiral HPLC analysis.

Scale-up experiments and synthetic applications

The configurational stability of the novel axially chiral N-N biheteroaryl pyrrolylindoles was investigated, and no erosion of enantiopurity was observed when the product 2a was heated in toluene at 130 °C for 48 h. Therefore, the configurational stability of these compounds could certainly show good tolerance for their further application in organic synthesis. To further highlight the application of this protocol for the synthesis of novel axially chiral biheteroaryl pyrrolylindoles, as shown in Scheme 2, the scale-up annulation reaction with 1 mmol substrate 1a was carried out under standard reaction conditions, and product 2a was easily accessible with high yield

and maintained good enantioselectivity (96% yield, 95:5 er). In addition, compound 2a could undergo a condensation-cyclization reaction with 2,2-diethoxy-N,N-dimethylethan-1-amine to furnish product 3 containing an indolo[3,2-a]carbazole skeleton in 80% yield without erosion of the er value, which forms the key core structure in natural indolo[3,2-a]carbazole alkaloids from deep-water sponges of the genus Asteropus.11

Investigation of biological activities

In view of N-N atropisomers being core units in some natural products and bioactive molecules, we are interested in the potential biological activities of these new axially chiral N,N'pyrrolylindoles. Consequently, the preliminary cytotoxic effects **Chemical Science Edge Article**

of randomly selected N,N'-pyrrolylindole products (2a, ent-2a, 2b, 2o, 2i and 2k) and transformation products (3 and ent-3) with an indolo[3,2-a]carbazole skeleton were then examined against some kinds of cancer cell lines, such as HL-60 cancer cells, A549 lung carcinoma cells, SMMC-7721 hepatoma cells, MDA-MB-231 breast adenocarcinoma cells, and SW480 colon cancer cells by MTS assay (MTS, 3-(4,5-dimethylthiazol-2-yl)-5(3carboxymethoxy-phenyl)-2-(4-sulfopheny)-2H-tetrazolium). As shown in Fig. 1a, preliminary evaluation of the results showed that these structurally-important molecules generally showed a high cell inhibitory rate against these cancer cells at a concentration of 40 µM. Then, we turned our attention to testing the corresponding cytotoxicity with half maximal inhibitory concentration (IC₅₀) values of these three molecules in the low micromolar range, with the widely used anticancer drug cisplatin used as a comparison group (Fig. 1b). It is worth

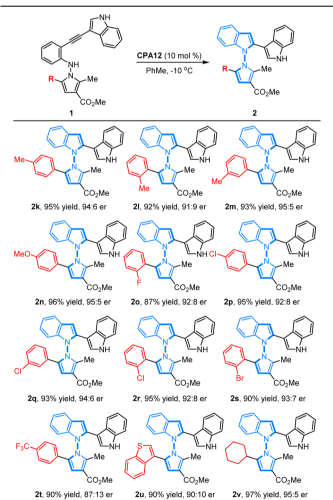
Table 2 Substrate scope of N-pyrrolo-acetylanilines^a

noting that compounds 2a and 2i exhibited impressive cytotoxicity against these five kinds of cancer cells with low IC₅₀ values, which were generally better than the drug cisplatin. Compound 3 also showed favorable anticancer activities against A549 lung carcinoma cells, SMMC-7721 hepatoma cells and MDA-MB-231 breast adenocarcinoma cells. The distinctive preliminary results of these high-value axially chiral N,N'-pyrrolylindoles could to some extent provide potentiality in the drug discovery and development process.

Mechanistic studies

Having established the synthesis of axially chiral N,N'-pyrrolylindoles based on o-alkynylanilines via de novo indole formation, we were interested in the reaction mechanism. To explore the mechanistic insights for this protocol, as depicted in Scheme 3a, further investigations and control experiments were then carried out. Substrates 1w and 1x without an indole group and the N-Bn protected substrate 1y did not work under

Table 3 Substrate scope of N-pyrrolo-acetylanilines^a



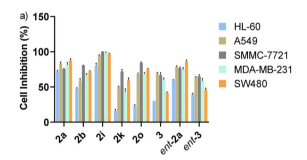
^a All reactions were carried out with 0.2 mmol 1 in 2 mL of toluene catalyzed by CPA12 (10 mol%) at -10 °C. The yield is isolated yield. The er value was determined by chiral HPLC analysis.

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Scheme 2 Scale-up experiment and synthetic transformation

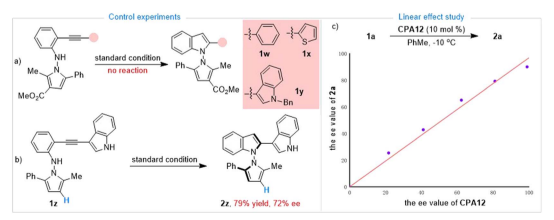
standard reaction conditions, leading to there being no desired product, which indicated that the indole group must be involved in the reaction pathway. In addition, substrate 1z in the absence of an ester group was also utilized to give the desired product 2z in 79% yield with 72% ee, which demonstrated that the ester group may play an important role in determining stereoselective control (Scheme 3b). Then, we paid attention to an investigation of the nonlinear effect between the enantioselectivity of 2a and the ee value of the CPA12 catalyst. As shown in Scheme 3c, linear correlation was observed, indicating that one active catalyst species was involved in the stereodetermination process.

To gain mechanistic insights into the CPA-dependent stereoselectivity, density functional theory (DFT) calculations were performed with the Gaussian 09 package.¹² As shown in Fig. 2a, the reaction starts with the hydrophosphoryloxylation of the alkyne moiety in the CPA-substrate adduct **IM1** *via* **TS1** with an energy barrier of 16.9 kcal mol^{-1} , leading to allene-type intermediate $\text{IM2}.^{13}$ Subsequently, the nucleophilic cyclization of the N atom in the aniline moiety on the central carbon of the allene moiety occurs *via* **TS2** ($\Delta G^{\ddagger} = 21.3 \text{ kcal mol}^{-1}$), forming protonated indole complex **IM3**. Finally, the deprotonation *via* **TS3** delivers product 2a and regenerates **IM1** through product/substrate exchange. Herein, the transition state of proton transfer **TS3** could not be located, which may be because proton transfer from ammonium to the phosphate anion could be regarded as an extremely facile process (see ESI† for more details). Overall, *N*-nucleophilic cyclization *via* **TS2** is the rate-



| b) | IC50 (μM) | | | | | |
|------------|-------------|-------------|-------------|-------------|--|--|
| | 2a | 2i | 3 | cisplatin | | |
| HL-60 | 17.36± 0.37 | 11.72± 0.42 | - | 10.25± 0.73 | | |
| A549 | 6.79± 0.22 | 10.15± 0.38 | 13.89± 1.37 | 25.03± 1.40 | | |
| SMMC-7721 | 13.45± 0.94 | 11.35± 0.51 | 20.90± 1.85 | 23.86± 1.49 | | |
| MDA-MB-231 | 18.22± 1.11 | 12.58± 0.49 | 23.57± 0.74 | 21.31± 1.74 | | |
| SW480 | 15.91± 0.75 | 12.41± 0.53 | -: | 19.95± 1.71 | | |

Fig. 1 (a) 40 μ M of eight randomly selected chiral N,N'-pyrrolylindoles were incubated individually with HL-60 cancer cells, A549 lung carcinoma cells, SMMC-7721 hepatoma cells, MDA-MB-231 breast adenocarcinoma cells, and SW480 colon cancer cells. (b) The cell viability after incubation with different concentrations of three chiral N,N'-pyrrolylindoles and their IC₅₀ values were determined in comparison to cisplatin.



Scheme 3 Control experiments and linear effect study.

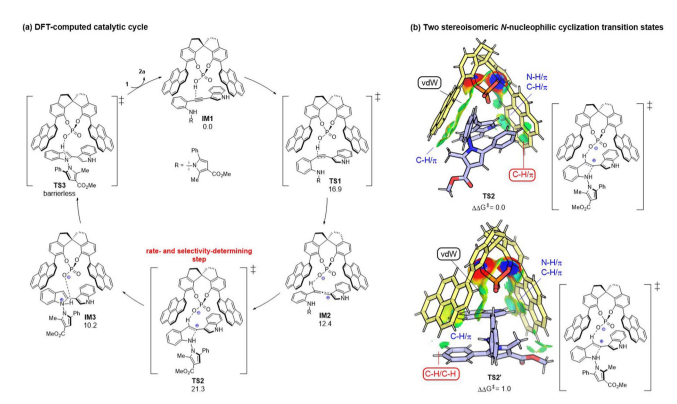


Fig. 2 (a) DFT-computed catalytic cycle with free energy given in kcal mol^{-1} . (b) Origin of the stereoselectivity between TS2 and TS2'. Comparison of key NCIs by IGMH analysis.

and stereo-determining step that controls the axial chirality of the formed product. **TS2** (leading to the major product) is found to be lower than **TS2**′ (leading to the minor product) by 1.0 kcal mol^{-1} , whose computed result agrees with the experimental observations. Noncovalent interactions (NCIs) using IGMH analysis have been utilized to compare the two competing transition states (**TS2** νs . **TS2**′). As shown in Fig. 2b, the main difference between the two transition states is that **TS2** has favorable $\mathrm{C-H}\cdots\pi$ dispersion interactions between the phenyl group of the pyrrole moiety and the pyrenyl substituent in CPA; while only weak $\mathrm{C-H/C-H}$ ligand–substrate interactions can be detected in **TS2**′ of the analogous parts. Accordingly, the ligand–substrate dispersion interactions serve as the key factor controlling the axial chirality of the N,N'-pyrrolylindole products.

Conclusions

In summary, we developed the first organocatalytic atropose-lective synthesis of axially chiral N,N'-pyrrolylindoles based on o-alkynylanilines through CPA-catalyzed asymmetric 5-endo-dig cyclization via de novo indole formation. This elegant protocol provided a facile synthetic strategy, and a broad range of intriguing N,N'-pyrrolylindoles were obtained in high yields with exclusive regioselectivity and excellent atroposelectivities (up to 99% yield, >20:1 rr, 95:5 er). The scale-up synthesis and the synthetic transformation for the construction of an indolo [3,2-a]carbazole skeleton as a key core structure in natural product asteropusazoles exhibited potential utilization.

Preliminary studies of biological activity suggested that these structurally important N,N'-pyrrolylindoles had a low IC₅₀ value, which could offer an encouraging opportunity for drug discovery. DFT mechanistic explorations indicated that the stereochemistry was regulated by the CPA-promoted N-nucleophilic cyclization step, wherein the favored transition state is stabilized by more ligand–substrate C–H··· π dispersion forces.

Data availability

All experimental procedures, characterisation data, mechanistic investigations, NMR spectra and HPLC spectra can be found in the ESI.†

Author contributions

C. J. W. conceptualized the project. C. J. W. and X. Q. D. supervised the investigation. C. S. W., Q. X., and H. R. Y. performed the research. Y. D. directed the DFT calculation, and H. X. performed the DFT calculation research. C. J. W., X. Q. D. and Y. D. co-wrote the paper. All authors analyzed the data, discussed the results, and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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