## SCIENCE CHINA Chemistry



· ARTICLES ·

SPECIAL ISSUE: Celebrating the 100th Anniversary of Chemical Sciences in Nanjing University

https://doi.org/10.1007/s11426-019-9711-x

# Multi-layer 3D chirality: new synthesis, AIE and computational studies

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Received December 30, 2019; accepted February 24, 2020; published online April 2, 2020

New synthesis of multi-layer 3D chiral molecules has been developed under novel conditions to give better outcomes. The aggregation-induced emission (AIE), UV irradiation/excitation, charge transfer (CT) and local excited (LE)  $\pi\pi^*$  transitions have been investigated on a representative individual enantiomer of *pseudo*  $C_2$  asymmetry which was made possibly by differentiating moieties on phosphorous on *N*-phosphonyl ring of chiral sandwich framework. Meanwhile, a new tandem C–N/C–C coupling reaction was unexpectedly rendered providing a novel access to special benzo[a]carbazoles.

multi-layer 3D chirality, organo sandwich chirality, architecture chirality, aggregation-induced emission (AIE), charge transfer (CT)

Citation: Liu Y, Wu G, Yang Z, Rouh H, Katakam N, Ahmed S, Unruh D, Cui Z, Lischka H, Li G. Multi-layer 3D chirality: new synthesis, AIE and computational studies. Sci China Chem, 2020, 63, https://doi.org/10.1007/s11426-019-9711-x

#### 1 Introduction

Chirality has attracted widespread attention in scientific communities for over a century because of its importance in chemistry, biology, material, and interdisciplines [1,2]. An increasingly large number of drugs have been showing chirality in their structures. The asymmetric synthesis and catalysis have thus become an active topic in synthesis and process chemistry in the past several decades, which heavily depend on controlling molecular chirality [3,4]. So far, the most popular and influential chirality has been represented by chiral BINAP/BINOL and  $C_2$  symmetric scaffolds, such as vicinal chiral diols and diamines [5–10]. However, there is

In stereochemistry, the doubly layered chirality has been established and proven to be effective in controlling asymmetric reactions [11–14]. For example, chiral titanocenes and zirconocenes have been successfully applied to asymmetric reduction/hydrogenation reactions and asymmetric polymerization processes [11]. Wilkinson ferrocene-based planar chirality has served well for asymmetric catalysis, such as Staudinger reaction, rearrangement of *O*-acylated enolates, acylation of alcohols and kinetic resolutions [13,14]. Two-layer chiral flavinium salt has been reported on the enantioselective alkyne conjugate addition and oxidation of sulfides to give modest to high enantioselectivity [15]. Very recently, helicene dimers of two-layer chirality have been

very limited work on conceptually new chirality of general interest and extensive potentials for chemical and biological research and material applications.

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synthesized for material research; this molecular framework contains chiral elements on each wing and can be rotated to give both *syn* and *anti* stereoisomers [16].

So far, the design of multi-layer chirality and its chemical synthesis have not been well documented in literature until very recently when we became involved in this research [17]. In this work, we report the new synthesis of individual enantiomers of multi-layer chirality and organic sandwich shape (Figure 1) by performing dual C–N couplings under new catalytic systems. At the same time, we also disclose an unexpected tandem C–N coupling/cyclization reaction of 8-([1,1'-biphenyl]-4-yl)naphthalen-1-amine coupling with 1,2-dibromobenzene (Scheme 1).

As displayed in Figure 1, this new chirality shows unprecedented  $pseudo\ C_2$  symmetry and belongs to a type of multi-layer organic framework (M-LOF) of single organic molecules. Its unique characteristics are represented by three levels of planar units arranging nearly in parallel fashion with one on top and the other one down from the centered planar anchor.

This design was initiated by our ongoing project on group-assisted purification (GAP) chemistry which is the first concept consisting of both chemical and physical aspects: reagent, reaction, separation, and purification [18,19]. By taking advantage of GAP chemistry, organic syntheses including their asymmetric versions can be conducted without the use of traditional purification by column chromatography and recrystallization. GAP functional groups can convert oily and sticky products into their solid forms, and often enable single crystals to be formed more easily when compared to classic methods. Essentially, pure GAP products can be obtained simply by washing crude mixtures with common

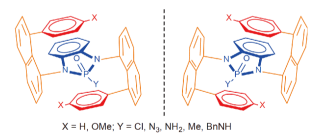
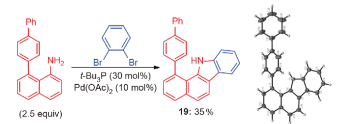


Figure 1 Multi-layer 3D chiral sandwich molecules (color online).



Scheme 1 Tandem catalytic C-N coupling/cyclization reaction (color online).

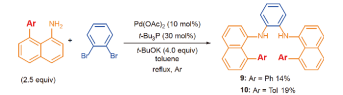
solvents and/or co-solvents.

The search for more efficient GAP reagents has been necessitated for the following objectives: to better meet requirements of solubility, stability and reactivity of starting materials and products; to enhance diastereo-, enantio- and chemoselectivities for asymmetric synthesis and catalysis; to enable GAP protection groups to be cleaved/recycle for reuse more effectively; and to further increase chemical yields for reactions [19c]. GAP chemistry can not only avoid disadvantages for both solution-phase and solid-phase peptide synthesis, but also increase their synthetic efficiency. Moreover, it has benefited establishing solution-phase peptide synthesizer; GAP chemistry has also enabled the Fmoc protection group, which is the most popular in solid-phase peptide synthesis (SPPS), to be utilized in solution-phase peptide synthesis [19a].

#### 2 Results and discussion

During our searching for more efficient GAP groups, we found a new single molecular framework containing multilayered and three-dimensional chirality [17]. One of the key steps of its synthesis involves dual Buchwald-Hartwig couplings [20] of 1,2-dibromobezene with 1-amino-8-arylnaphthalene to get vicinal N,N-bis(8-arylnaphthalen-1-yl) benzene-1,2-diamines, albeit in poor yields (Scheme 2). During this synthesis, we found para-methyl anchored arylnaphthalene substrate led to the coupling product in an improved yield; and we also observed that almost all diamino precursors and multi-layer compounds are highly fluorescence-active (Figure 2; Figure S30, Supporting Information online). These findings led us to attach a stronger electrondonating group, MeO-, onto the arylnaphthalene substrate trying to further enhance chemical yields. In addition, since phenyl rings of two wings of multi-layer 3D chiral products are conjugated by -OMe group, it would result in new photoelectronic properties for future material research. However, under original catalytic systems, this replacement did not improve chemical yields that are indeed on the same level as these of previous couplings. We thus continued searching for new catalytic conditions in order to optimize the key dual couplings.

As shown in Table 1, several factors for this catalytic coupling need to be adjusted carefully. It should be mentioned that in our initial synthesis, the use of 1-bromo-8-phenylnaphthalene to react with vicinal benzene diamine did not result in any diamino product. This lesson suggested us to directly use 1,2-dibromobenzene and 8-(4-methoxyphenyl) naphthalen-1-amine as reactants for present optimizations. The coupling in toluene with 3.0 equiv. of *t*-BuONa at 110 °C only afforded 11% yield. The yield was slightly improved to 15% when the reaction was performed in *p*-xylene.



Scheme 2 Dual Buchwald-Hartwig Diamino couplings [16b] (color online).

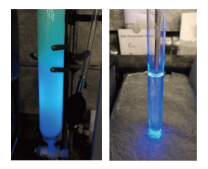


Figure 2 Fluorescence showings of multi-layer 3D chiral amide (16a) (color online).

Replacing *t*-BuONa with *t*-BuOK resulted in a higher yield to 19%. Decreasing the loading of *t*-BuOK to 2.1 equiv. further increased chemical yield to 24%. Shortening reaction time from 24 to 19 and 14 h, chemical yields were increased to 33% and 41%, respectively (Table 1).

After achieving the above optimization, we next re-investigated the original coupling of 1,2-dibromobenzene with 1-amino-8-phenylnaphthalene under this new catalytic system. We found this system resulted in vicinal N,N-bis(8phenylnaphthalen-1-yl)benzene-1,2-diamine in a chemical vield of 27% which almost doubled the original yield of 14% (Scheme 3). It is interesting that as shown in the X-ray structure of this diamino product, the phenyl ring on position  $N^{1}$  of (8-phenylnaphthalen-1-yl)benzene 1,2-diamine is arranged in parallel to the corresponding naphthyl ring on  $N^2$  of this benzene 1,2-diamine, and vice versa. However, as shown in Scheme 4, this conformation has to be adjusted to become those of 9 and 10 so as to enable the next cyclization to occur. The serious steric effects by two phenyl groups of vicinal N, N-bis(8-aryl- naphthalen-1-yl)benzene-1,2-diamines would be responsible not only for low chemical yields of dual couplings (27% and 41% for 9 and 10, respectively), but also for relatively low yields of the next cyclization step (39% and 69% for 11 and 12, respectively).

As shown in Schemes 4 and 5, nine steps were needed to achieve the GAP amide product of 2-amino-1,3-bis(8-(4-methoxyphenyl)naphthalen-1-yl)-1,3-dihydrobenzo[d] [1,3,2]diazaphosphole 2-oxide (16). The synthesis was started from an inexpensive commercial chemical, naphthalene-1,8-diamine, via oxidative cyclization by treating with so-dium nitrite in aqueous media containing acetic acid. The

Table 1 Palladium-catalyzed coupling conditions and results<sup>a)</sup>

Entry	Base	Solvent	Temperature (°C)	Time (h)	Yield (%)
1	t-BuONa (3.0 equiv.)	toluene	110	24	11
2	t-BuONa (3.0 equiv.)	<i>p</i> -xylene	133	24	15
3	<i>t</i> -BuOK (3.0 equiv.)	<i>p</i> -xylene	133	24	19
4	t-BuOK (2.1 equiv.)	<i>p</i> -xylene	133	24	24
5	t-BuOK (2.1 equiv.)	<i>p</i> -xylene	133	19	33
6	t-BuOK (2.1 equiv.)	<i>p</i> -xylene	133	14	41

a) All reactions were performed in tubes which are tightly capped by septum and protected with balloons filled with argon gas; during the reaction, a small amount of hexane employed for preparation of (*t*-Bu)<sub>3</sub>P solution flew into balloon through needle on septum.



**Scheme 3** Diamino coupling using 1-amino-8-phenylnaphthalene (color online).

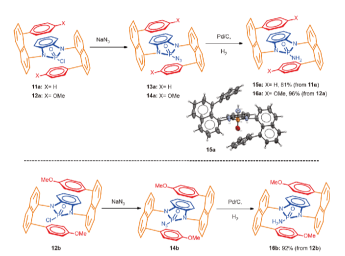
resulting 1H-naphtho[1,8-de][1,2,3]triazine (2) was treated with metal copper in hydrogen bromide to give 8-bromonaphthalen-1-amine (3) which was converted into N-(8bromonaphthalen-1-yl)acetamide (4) by protection with acetyl chloride. Suzuki coupling of 4 with (4-methoxyphenyl)boronic acid gave N-(8-(4-methoxyphenyl)naphthalen-1-yl)acetamide (6) followed by acidic hydrolysis with concentrated aqueous HCl to afford 8-(4-methoxyphenyl) naphthalen-1-amine (8). The optimized conditions of dual Buchwald-Hartwig C-N couplings resulted in vicinal N,Nbis(8-(4-methoxyphenyl)naphthalen-1-yl)benzene-1,2-diamine (10) which exists in an equilibrium of enantiomeric conformers. Two of these conformers are represented, which is based on the conformations of next cyclized N-phosphonyl chlorides (Figure S29) and X-ray structural analysis of its derived phosphoramide. As usual, we made several efforts on obtaining these enantiomeric conformers by chiral high performance liquid chromatography (HPLC), but we fizzled due to their unfixed flexibility. However, after they are cyclized to 2-chloro-1,3-bis(8-(4-methoxyphenyl)naphthalen-1-y1)-1,3-dihydrobenzo[d][1,3,2]diazaphosphole (12), we were able to separate these two enantiomers via chiral prep-HPLC (Scheme 4).

At the cyclization step, the reaction of diamines with phosphoryl trichloride in triethylamine or pyridine did not

**Scheme 4** Synthesis of *Pseudo C*<sub>2</sub> & 3D N,N-phosphonyl chlorides (color online).

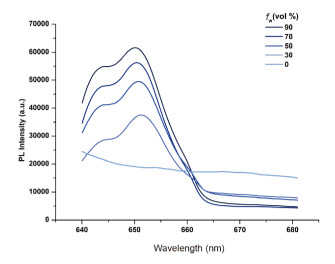
give any cyclization product, although they worked well for cases of (1R,2R)-cyclohexane-1,2-diamine and (1R,2R)-1,2diphenylethane [18,19]. This reaction has to be performed by pre-deprotonating N,N-bis(8-(4-methoxyphenyl)naphthalen-1-yl)benzene-1,2-diamine (10) with *n*-butyl lithium followed by the treatment with phosphorus oxychloride at -78 °C in dried tetrahydrofuran (THF). In our original cyclization of N,N-bis(8-phenylnaphthalen-1-yl)benzene-1,2-diamine, yield of 39% was obtained. However, in the present case, the corresponding yield was substantially enhanced to 69%, which is likely due to the strong-electron donating effect of the methoxy group making diamino moieties more nucleophilically reactive toward the N-phosphonyl protection in both intramolecular and intermolecular manners at this step. This observation is consistent with the case of the methyl derivative under the same conditions which gave a higher chemical yield of 45% than that of the unsubstituted case (39%) [19b]. The resulting azide of this para-methoxy case was found to be not so stable and showed complex unknown species after being isolated and stored for a short period of time at room temperature. It was thus directly subjected to Pd/C catalyzed hydrogenation with the crude azide (14a) to give the final product of 2-amino-1,3-bis(8-(4-methoxvphenyl)naphthalen-1-yl)-1,3-dihydrobenzo[d][1,3,2]diazaphosphole 2-oxide (Scheme 5). Similarly, the multi-layer 3D compounds also showed strong fluorescence activity inside column and in solution (Figure 2). 16a was selected at random for further aggregation-induced emission (AIE) study [21]. The photoluminescence (PL) spectra of 16a in THF/ water mixture with different water fractions  $(f_w)$  and the corresponding plot of emission enhancement versus  $f_{\rm w}$  demonstrate the AIE activity (Figure 3). It is non-fluorescent when dissolved in THF solution but becomes emissive at 651 nm with a 37,500 a.u. intensity when  $f_w=30$  vol% in THF/water mixture. While more water is gradiently added  $(f_w=50 \text{ vol}\%, 70 \text{ vol}\% \text{ and } 90 \text{ vol}\%)$ , the emission intensities are significant increase to 49,500, 56,100 and 61,800, respectively.

We also made several efforts on obtaining single crystals of individual enantiomers of 2-chloro-1,3-bis(8-(4-methox-

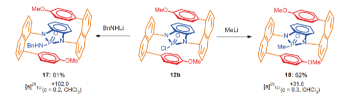


**Scheme 5** Synthesis of *pseudo*  $C_2$  & 3D N,N-phosphonyl amides (color online).

yphenyl)naphthalen-1-yl)-1,3-dihydrobenzo[d][1,3,2]diazaphosphole 2-oxide (12) and the corresponding final amide product (16), but failed in both cases. Fortunately, the absolute stereochemistry can be assigned based on the analogous structure of 2-amino-1,3-bis(8-phenylnaphthalen-1-yl)-1,3-dihydrobenzo[d][1,3,2]diazaphosphole 2-oxide, which has been confirmed by X-ray diffraction analysis [19b]. The enantiomerically pure isomer of 2-chloro-1,3-bis(8-(4methoxyphenyl)naphthalen-1-yl)-1,3-dihydrobenzo[d] [1,3,2]diazaphosphole 2-oxide (12b) was further reacted with lithium benzylamine which was pre-generated by treating benzylamine with butyl lithium in THF at -78 °C (Scheme 6). The reaction was performed at -78 °C for 30 min followed by gradually increasing temperature from -78 °C to room temperature. The reaction proceeded to completion by stirring at room temperature for 6 h to give 2-(benzylamino)-1,3-bis(8-(4-methoxyphenyl)naphthalen-1yl)-1,3-dihydrobenzo[d][1,3,2]diazaphosphole 2-oxide (17) with an 81% yield. It was also subjected to the reaction with methyl lithium under the same conditions to afford 1,3-bis(8-(4-methoxyphenyl naphthalen-1-yl)-2-me thyl-1,3-dihydrobenzo [d][1,3,2] diazaphosphole 2-oxide (18) in a yield of



**Figure 3** Photoluminescence (PL) spectra of **16a** in THF/water mixtures with different water fractions ( $f_w$ ).  $c=1 \mu M$ ; excitation wavelength ( $\lambda_{ex}$ ): 532 nm (color online).



**Scheme 6** Reactions of *N*,*N*-phosphonyl chloride with lithium reagents (color online).

62%. As revealed by chiral HPLC, there is no racemization observed in these two reactions.

Interestingly, when we utilized a phenyl group to replace para MeO- for the coupling, the reaction underwent toward another direction, i.e., intermolecular C-N bond formation was followed by intramolecular C-C cyclizing (Scheme 1). The desired product was barely formed in the resulting mixture. Instead, 1-([1,1'-biphenyl]-4-yl)-11H-benzo[a]carbazole (19) was generated as the major product in a 35% yield, and its structure was determined by X-ray diffraction analysis (Scheme 1). Product 19 appeared as thin snowflakeshaped solids, which benefited to generating single crystals for X-ray analysis. This product is anticipated to serve as a structural lead for material and medicinal chemistry because similar structural units showed applications in these fields [22]. As shown in its structure, the N–H bond on the indole ring is nearly perpendicular to the phenyl ring on position-8 of phenylnaphthalene; this unique arrangement will also have applications for catalyst design by introducing functional groups onto nitrogen atoms.

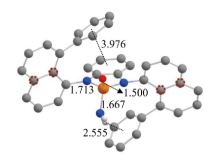
To further understand the electronic behavior of these new chiral molecules, we conducted computational studies by selecting **15a** as a representative model which has been unambiguously confirmed by X-ray structural analysis. In our computational methods, density functional theory (DFT) in

combination with the PBE0-D3 functional [23,24] and the cc-pVDZ basis set [25] was used to optimize the geometry of the complex (Figure 4). To preserve the main features of the crystal structure, selected atoms indicated in this figure were fixed at their positions in the crystal structure during the optimization process. Second-order algebraic diagrammatic construction (ADC(2)) calculations [26–28] were performed using the TVZP basis set [29] to compute the excited-state spectrum. Natural transition orbitals (NTOs) [30] were used to characterize the electronic transition at orbital level. Environmental effects were considered by the conductor-like screening model (COSMO) [31], based on the state specific approach [32]. For the dielectric constant  $\varepsilon$  and the refractive index n, respectively, the values  $\varepsilon$ =8.93 and n=1.42 were used, which correspond to the weakly polar dichloromethane. For the calculation of the vertical emission transition from  $S_1$ , water was chosen as solvent ( $\varepsilon$ =78.4 and n=1.33). The calculations were carried out using the program package TURBOMOLE [33].

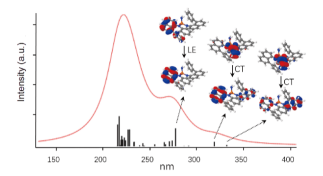
The selected bond and interring distances are displayed in Figure 4. Compared to the crystal structure, the calculated bond distances of the aromatic systems do not show significant differences, whereas the four bonds associated with the phosphorus atom are slightly longer by 0.03 Å. The two interring distances given in Figure 4 are 0.2–0.3 Å shorter than in the crystal, demonstrating the crystal packing effect on these weak nonbonded interactions.

The computed UV/Vis spectrum is displayed in Figure 5. The spectrum is characterized by two low-lying charge transfer (CT) bands at 332 and 319 nm. The CT involved in these transitions is quite significant and amounts to 0.8e. It is directed from the central ring to the naphthalene units (see insets in Figure 5). The location of these two weak transitions clearly indicates that the broadened absorption observed at around 320-330 nm can be ascribed to CT transitions. The next group of transitions around 280 nm are responsible for the next band and represent local excited (LE)  $\pi\pi^*$  transitions within the naphthalene ring. The strongest band is formed by a series of closely-spaced transitions between 216 to 228 nm, which are characterized as LE in naphthalene coupled with CT character. The UV/Vis spectrum predicted by ADC(2) is consistent with the experimental data (Figure S31), which shows the broadened absorption at 330 nm (CT transitions) and very pronounced absorption at around 290 nm owing to the LE transitions. The sharp absorption at around 240 nm in computed spectrum can be ascribed to the closely-spaced transitions with LE coupled with CT character.

The luminescence properties of the model 15a structure (Figure 4) have been investigated at ADC(2) level. For that purpose, the structure of 15a has been optimized for the  $S_1$  state in gas phase without geometry restrictions. Structural changes in comparison to the ground state and the vertical emission energy for aqueous solution have been calculated.



**Figure 4** Optimized structures with selected bond distances in Å computed at the PBE0-D3/cc-pVDZ level. Hydrogen atoms connected to carbon are not shown and the carbon atoms fixed in the optimized process are marked by circles (color online).



**Figure 5** Simulated UV/Vis spectrum at the ADC(2)/cc-pVDZ level in dichloromethane, and pictures of the NTOs are given for  $S_1$ ,  $S_2$  (CT transitions) and  $S_5$  (LE) states (color online).

The CT in S<sub>1</sub> has the effect of equalizing the C–C bond distances in the naphthalene subunit. The C–N bond lengths in the heterocycle decrease between 0.03 and 0.05 Å. The vertical emission wavelength is computed at 594 nm for a monomer unit. The comparison of the computed emission wavelength with the experimental luminescence band maximum for 16a at 650 (aggregated state) gives an estimate for the effect of the aggregation on the emission wavelength of about 60 nm. Moreover, a pronounced Stokes shift of 249 nm, computed as the difference of vertical excitation and emission energies in water for the non-aggregated systems, is found.

### 3 Conclusions

In summary, we developed more efficient synthesis of organic sandwich molecules with unique *pseudo*  $C_2$ -symmetry by searching for new dual catalytic couplings. The *pseudo*  $C_2$  design was made possibly by differentiating two moieties on phosphorous on the centred five-membered N-phosphonyl ring. Meanwhile, a new catalytic reaction was established for the formation of 1-([1,1'-biphenyl]-4-yl)-11H-benzo[a]carbazole as confirmed by X-ray diffraction analysis. The AIE, UV irradiation/excitation, CT and LE  $\pi\pi^*$  transitions have

been investigated on a representative individual enantiomer. We have found that the UV/vis spectrum predicted by ADC (2) is very well consistent with the experimental data, which show the broadened absorption at 330 nm (CT transitions) and very pronounced absorption at around 290 nm owing to the LE transitions. The sharp absorption at around 240 nm in experimental spectrum can be ascribed to the closely-spaced transitions with LE coupled with CT character. The comparison of the calculated emission energy for the monomer unit with the experimental band maximum for the aggregated state shows a substantial influence of the aggregation of about 60 nm.

Acknowledgements This work was supported by the National Natural Science Foundation of China (91956110, 21672100) and Robert A. Welch Foundation (D-1361, USA). Z. C. acknowledges financial support from Beijing National Laboratory for Molecular Sciences (BNLMS201910). H. L. are grateful for support from the School of Pharmaceutical Science and Technology (SPST), Tianjin University, Tianjin, China, including computer time on the SPST computer cluster Arran. We thank Dr. Kaz Surowiec for the analytic assistance discussion.

**Conflict of interest** The authors declare that they have no conflict of interest.

**Supporting information** The supporting information is available online at <a href="http://chem.scichina.com">http://chem.scichina.com</a> and <a href="http://link.springer.com/journal/11426">http://link.springer.com/journal/11426</a>. The supporting materials are published as submitted, without typesetting or editing. The responsibility for scientific accuracy and content remains entirely with the authors.

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