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Description						



## Designed Molecular Propellers Based on Tetraarylterephthalamide and their Chiroptical Properties Induced by Biased Helicity through **Transmission of Point Chirality**

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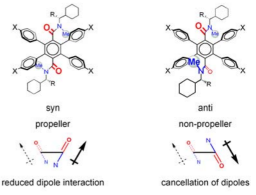
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5 The syn-atropisomers of the title bis(tertiary amide)s were designed as six-bladed molecular propellers based on the "directing effects" of amide dipoles; the helicity of the propeller is biased upon the attachment of point chirality to the amide nitrogens to attain stronger circular-dichroism activity than for 10 the non-propeller-shaped anti-isomers.

Persubstituted benzenes have attracted much attention as promising motifs in the design of molecular rotors, gears, and propellers.1 One of the most-studied classes of molecules is hexaarylbenzenes, 1a,2 which cannot adopt a conformation with 15 all aryl blades lying on the same plane as the central core due to the steric repulsion among neighboring substituents. Instead, substituents are twisted almost perpendicularly, and slight skewing about C<sub>central</sub>-C<sub>aryl</sub> bond in either direction induces numerous conformations, including several chiral 20 ones. The propeller-shaped conformation occurs when all blades are skewed in a conrotatory manner, which is interesting in terms of its mobile chirality with an easy inversion of helicity. While the parent hexaphenylbenzene<sup>2b,e</sup> was shown to adopt a propeller conformation in the crystal, it 25 is difficult to force the molecule to maintain this same structure in solution. The control of propeller helicity is also challenging to bias the (P)/(M)-stereoisomer.

We envisaged that the rational design of persubstituted benzenes would provide a significant preference for the 30 propeller geometry even in solution, and the mobile helicity could be biased to a single handedness through the effective transmission of point chiralities on the periphery. Based on our detailed conformational studies on terephthalamides,<sup>3</sup> we have designed here 2,3,5,6-tetraarylated bis(tertiary amide)s  $_{35}$  1/2 as promising compounds to adopt a propeller geometry. As in the case of hexaarylbenzenes, all of the blades in 1/2 are twisted nearly perpendicularly, which gives two atropisomers (syn and anti) in terms of the relative direction of the two amide groups. Both isomers must prefer the conformation 40 with the amide dipoles reduced/canceled to minimize the electrostatic disadvantage: a conrotatory-skewed  $C_2$ symmetric structure for syn and a centrosymmetric structure for anti (Scheme 1). Such "directing effects" by the amide groups in the former should endow the syn-isomers with a 45 preference for the propeller-shape by forcing conrotatory skewing of the four aryl blades. This is the central point of our concept toward "designed molecular propellers", and we report here the details of a successful demonstration. The

point chirality on the amide nitrogen is transmitted to the 50 helicity-preference of the propellers to attain much stronger chiroptical signals for the syn-isomers of 1/2 than for the nonpropeller-shaped anti-isomers.



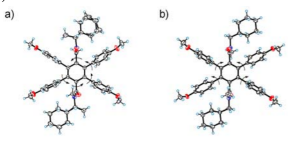
[1 : X = OMe, 2 : X = p-MeO-C<sub>6</sub>H<sub>4</sub>, a : R = H, b : R = Me] Scheme 1

For detailed conformational analyses, 4-methoxyphenyl (1) 55 and 4'-methoxybiphenyl-4-yl (2) groups are selected as the aryl blade to simplify the aryl region of the NMR spectrum. The cyclohexylmethyl (a) or (R)-1-cyclohexylethyl (b) group is attached to each amide nitrogen, which of the latter was anticipated to act as a chiral-handle to control the mobile 60 helicity of the propellers. Further substitution on nitrogens to bis(tertiary amide)s would be necessary to suppress interconversion between syn- and anti-atropisomers. The larger substituent on the amide may preferentially occupy the "lateral" position (s-trans to the benzene core), whereas the 65 smaller one may be located at the "vertical" position (s-cis to the core). The rotational freedom about C<sub>central</sub>-C<sub>aryl</sub> bond depends on the steric bulkiness of the vertical group.

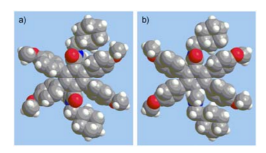
Tetraarylterephthalamides 1a,b were prepared by Suzuki-Miyaura coupling<sup>4</sup> of 2,3,5,6-tetrabromoterephthalamides 70 **3a,b** and 4-methoxyphenylboronic acid followed by Nmethylation of the intermediary secondary amides 4a,b. The tetrakis(biphenylated) derivatives 2a,b were similarly prepared by using 4'-methoxybiphenyl-4-yl-boronic acid' in the Pd-catalyzed coupling step. The syn- and anti-75 atropisomers were formed as mixtures, which were readily separated in pure form by column chromatography since they exhibit quite different  $R_f$  values. The isomers with a smaller  $R_f$  value were tentatively assigned as syn, and this was later confirmed by X-ray analyses on syn- and anti-1 (vide infra). 80 The atropisomers 1/2 were non-interconvertible even after heating at 338 K for hours in CDCl<sub>3</sub>. For the bis(secondary amide)s **4a,b** without vertical methyl substituents, interconversion is much faster than the NMR time-scale, even at room temperature, as predicted.

Single-crystal X-ray analysis<sup>††</sup> demonstrated that (R,R)-syn-**1b** adopts the propeller geometry in crystal, as desired. The aryl blades are skewed in a conrotatory manner, which must be induced by the directing effects of amide dipoles at the para-positions (Fig. 1a). Preliminary investigation indicates this is also the case for achiral syn-**1a** (Fig. S1; P-1, Z = 2). In the latter case, the enantiomeric propellers exist in pair, whereas, in the former with a chiral auxiliary on each amide nitrogen, only propellers with (M)-helicity exist in the crystal, and thus the point chiralities are perfectly transmitted to the mobile helicity, at least in a crystalline state of (R,R)-syn-**1b**.

On the other hand, the electrostatic interaction of the amide "directing effects" must be the major reason why anti-1a adopts a centrosymmetric structure with full cancellation of the dipole, and thus the anti-isomer is not propeller-shaped at all (Fig. 1b). Even with the attachment of chiral auxiliaries, the anti-isomer prefers the pseudo-centrosymmetric geometry to minimize dipole repulsion, as shown by the similarity of the non-propeller structure of (R,R)-anti-1b (Fig. S1; P1, Z=1) and anti-1a.



25 **Fig. 1** X-ray structures of a) (R,R)-syn-**1b** (P1, Z=1) and b) anti-**1a** (P-1, Z=1). One of the aryl blades in a) is almost perpendicular to the central core. Solvated benzene molecule in the crystal in b) is omitted for clarity. Thermal ellipsoids are shown at 50% probability



level

Fig. 2 Energy-minimized structures for a) syn-1a and b) anti-1a according to Monte-Carlo simulations in CHCl<sub>3</sub>.

Conformational searches on achiral 1a using Macromodel software indicated that the propeller conformation is the most stable geometry for the syn-isomer among numerous conformations (Fig. 2a). This theoretical examination suggests that the propeller-shaped geometry must also be the major contributor in solution for the syn-isomers of 1a,b. The

40 centrosymmetric non-propeller-shaped structure was predicted for the anti-isomer in accordance with the observed geometry in crystal (Fig. 2b).

The structures of syn-isomers in solution were experimentally verified by <sup>1</sup>H NMR spectroscopy (Fig. S2). 45 The chemical shifts of their aromatic protons in CDCl<sub>3</sub> at 298 K are summarized in Table 1. The observed spectrum of achiral syn-1a was  $C_{2v}$ -symmetric, which can be accounted for by assuming a rapid inversion of helicity between two energetically equivalent propeller structures. 50 broadening is related to rotational motion about the Ccentral- $C_{arvl}$  bond ( $\Delta G^{\ddagger} = 15.3 \text{ kcal mol}^{-1}$ ,  $T_c = 313 \text{ K}$ ), but there is no indication of  $C_{central}$ - $C_{amide}$  bond rotation. In the case of (R,R)syn-**1b** with chiral auxiliaries, the spectrum is  $C_2$ -symmetric, as expected for rapidly interconverting diastereomeric 55 propellers (Fig. S2). Slight broadening is again induced by  $C_{central}$ - $C_{arvl}$  rotation. There are four sets of resonances (H<sup>C</sup>: 7.18-6.87 ppm in CDCl<sub>3</sub> at 298 K) that correspond to anisyl protons close to the central benzene core. The other four (H<sup>X</sup>: 6.77-6.48 ppm) are ortho to the methoxy group, and are not 60 well separated from each other. The energy barrier for propeller inversion seems to be too low to observe the diastereomeric propellers as two independent sets of signals, even when the temperature is lowered to 223 K. Although the above NMR features might be alternatively rationalized by 65 assuming a single geometry with all substituents twisted perpendicular to the central core, only one of four anisyl signals (H<sup>C</sup>) showed a correlation with the vertical methyl group in an ROE measurement (Fig. S3), which is readily accounted for by assuming a propeller geometry. Furthermore, 70 as shown in Figure S4, the temperature-dependent shift of resonances while maintaining the spectral symmetry in (R,R)syn-1b shows that there are energetically-nonequivalent and rapidly-interconverting (P)- and (M)-propellers, whose diastereomeric ratio is a function of temperature.

Table 1 Chemical shifts<sup>a</sup> for the aromatic protons in 1/2<sup>b</sup> in CDCl<sub>3</sub> at

	δ / ppm					observed symmetry
syn-1a	7.09 <sup>C</sup> (4H)	6.87 <sup>C</sup> (4H)	6.64 <sup>X</sup> (4H×2)			C <sub>2v</sub>
(R,R)-syn-1b	7.18 <sup>C</sup> (2H)	7.02 <sup>C</sup> (2H)	6.93 <sup>C</sup> (2H)	6.87 <sup>C</sup> (2H)	6.77-6.48 <sup>X</sup> (2H×4)	C <sub>2</sub>
anti-1a	7.55 <sup>C</sup> (4H)	6.79 <sup>X</sup> (4H)	6.66 <sup>C</sup> (4H)	6.48 <sup>X</sup> (4H)		C <sub>2h</sub>
(R,R)-anti-1b	7.59 <sup>C</sup> (2H×2)	6.81 <sup>X</sup> (2H) 6.79 <sup>X</sup> (2H)	6.65 <sup>C</sup> (2H) 6.62 <sup>C</sup> (2H)	6.49 <sup>X</sup> (2H) 6.45 <sup>X</sup> (2H)		C <sub>2</sub>
syn-2a	7.31 <sup>C,X</sup> (4H×3)	7.07 <sup>C</sup> (4H)				C <sub>2v</sub>
(R,R)-syn-2b	-					-
anti-2a	7.74 <sup>C</sup> (4H)	7.50 <sup>X</sup> (4H)	7.17 <sup>X</sup> (4H)	6.86 <sup>C</sup> (4H)		C <sub>2h</sub>
(R,R)-anti-2b	7.80 <sup>C</sup> (2H×2)	7.53 <sup>X</sup> (2H) 7.50 <sup>X</sup> (2H)	7.17 <sup>X</sup> (2H) 7.11 <sup>X</sup> (2H)	6.89 <sup>C</sup> (2H) 6.81 <sup>C</sup> (2H)		C <sub>2</sub>

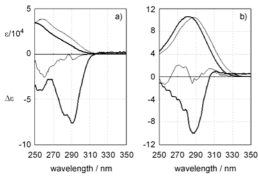
a Assignment is indicated by superscripts C and X. Superscript C denotes the aromatic protons close to the centra benzene ring. Superscript X denotes the aromatic protons close to X  $(X = MeO \text{ for } 1, \text{ and } 4^tMeOC_0H_4 \text{ for } 2)$ . The aromatic protons in  $(R,R) > y \sim 2b$  could not be fully assigned due to peak overlap and broadening. b: Only the protons on four phenylene rings attached on the core are shown.

298 K.

In the case of tetrakis(biphenylated) derivatives syn-2a,b, the NMR spectra show similar characteristics (Table 1; Fig. S2), suggesting that they also prefer the propeller-shape. This idea was supported by the chiroptical properties of (*R*,*R*)-syn-2b, which resemble those of propeller-shaped (*R*,*R*)-syn-1b (vide infra). The NMR data of anti-isomers of 1a,b/2a,b are salso listed in Table 1 and Fig. S2, and can be readily explained by assuming the (pseudo)centrosymmetric

geometries observed in crystal.

Figure 3 shows the UV and circular dichroism (CD) spectra of chiral (*R*,*R*)-1b/2b in CH<sub>2</sub>Cl<sub>2</sub> at room temperature. The UV spectra of achiral 1a/2a are quite similar to those of (*R*,*R*)-5 1b/2b, respectively (Table S1). For the atropisomers of (*R*,*R*)-1b/2b, the amplitude of the Cotton effect in the CD spectra differs significantly: CD signals for syn-isomers are much stronger than those for the corresponding anti-isomers. An almost 10-fold increase in Δε at the absorption shoulder (291 nm for 1b)/maximum (286 nm for 2b) is noteworthy (Table 2). Such differences in chiroptical properties must be due to the propeller-shape preference seen for syn-isomers. This is a successful demonstration of the effective transmission of chirality from point asymmetry to the mobile helicity of the 15 propeller structure.



**Fig. 3** UV and CD spectra of a) (R,R)-syn-**1b** (bold line), (R,R)-anti-**1b** (thin line) , and b) (R,R)-syn-**2b** (bold line), (R,R)-anti-**2b** (thin line) in  $CH_2Cl_2$  at room temperature, respectively.

The fluorophoric properties of the tetrakis(biphenylated) derivatives **2** are also interesting. Thus, (R,R)-syn-**2b** represents a new entry into the less-well developed class of chiral fluorophores. The large Stokes-shifts (ca. 100 nm) as well as structureless emission are characteristic of two-25 dimensional  $\pi$ -conjugated oligoarylenes (Table 2, Fig. S5).

**Table 2** UV, CD and fluorescence spectral data of (R,R)-**1b**/**2b** in  $CH_2Cl_2$  at room temperature.

	syn- <b>1b</b>	anti- <b>1b</b>	syn- <b>2b</b>	anti- <b>2b</b>
λ <sub>max</sub> / nm	249 (sh.)	259	282	287
$(\log \epsilon)$	(4.54)	(4.59)	(5.02)	(5.02)
λ <sub>max</sub> / nm	254, 291	261, 291	286, 312	256, 272, 286, 304
$(\Delta \epsilon)$	(-4.06, -7.67)	(-2.56, -0.608)	(-10.0, 1.04)	(-1.38, 2.11, -1.18, 1.43)
λ <sub>max, em</sub> <sup>a</sup> / nm	-	=	379	381

a: Excited at each absorption maximum.

In conclusion, we have demonstrated "designed" six-bladed molecular propellers based on syn-terephthalamides derivatives thanks to the "directing effects" of amide dipoles. A chiral auxiliary on the amide nitrogen gave a preference for skewing the direction of the propeller-helicity to realize 35 chiroptical enhancement. We are now studying the complexation properties of the present amides and hydrogenbonding guests with special focus on guest-induced conformational changes.3b-d A preliminary examination shows that the secondary amides 4a,b prefer a non-propeller 40 anti-conformation, but can undergo rapid syn-anti interconversion to attain the propeller structure upon complexation. Further details will be reported in due course.

## **Notes and references**

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- <sup>50</sup> † Electronic Supplementary Information (ESI) available: X-ray and NMR data; preparation detail of new compounds. See DOI: 10.1039/b0000000x/ ††Crystal data of (R,R)-syn-**1b**: MF C<sub>54</sub>H<sub>64</sub>N<sub>2</sub>O<sub>6</sub>, FW 837.11, Triclinic P1 (No. 1), a=6.305(2), b=13.323(5), c=13.959(5) Å, α = 86.986(10), β = 81.092(9), γ = 85.727(9)°, V = 1154.1(7) ų, ρ = (Z = 1) 1.204 g cm³, T = 155.727(9)°, Z = 10.000 cm², Z = 10.00
- 55 153 K, 7239 independent reflections (*R*int = 0.010), R = 4.3% (6810 data with  $F > 2\sigma F$ ), CCDC 680696. Crystal data of anti-1a: MF  $C_{58}H_{66}N_2O_6$ , FW 887.17, Triclinic P-1 (No. 2), a = 9.668(3), b = 11.144(4), c = 11.778(4),  $\alpha = 106.388(5)$ ,  $\beta = 96.224(5)$ ,  $\gamma = 100.015(5)^\circ$ , V = 1182.0(7) ų,  $\rho = (Z = 1)$  1.246 g cm³, T = 153 K, 5029 independent reflections
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