

Article

Central-to-Helical-to-Axial-to-Central Transfer of Chirality with a Photoresponsive Catalyst

Stefano F. Pizzolato, Peter Stacko, Jos C. M. Kistemaker, Thomas van Leeuwen, Edwin Otten, and Ben L. Feringa

J. Am. Chem. Soc., Just Accepted Manuscript • DOI: 10.1021/jacs.8b10816 • Publication Date (Web): 20 Nov 2018 Downloaded from http://pubs.acs.org on November 20, 2018

Just Accepted

"Just Accepted" manuscripts have been peer-reviewed and accepted for publication. They are posted online prior to technical editing, formatting for publication and author proofing. The American Chemical Society provides "Just Accepted" as a service to the research community to expedite the dissemination of scientific material as soon as possible after acceptance. "Just Accepted" manuscripts appear in full in PDF format accompanied by an HTML abstract. "Just Accepted" manuscripts have been fully peer reviewed, but should not be considered the official version of record. They are citable by the Digital Object Identifier (DOI®). "Just Accepted" is an optional service offered to authors. Therefore, the "Just Accepted" Web site may not include all articles that will be published in the journal. After a manuscript is technically edited and formatted, it will be removed from the "Just Accepted" Web site and published as an ASAP article. Note that technical editing may introduce minor changes to the manuscript text and/or graphics which could affect content, and all legal disclaimers and ethical guidelines that apply to the journal pertain. ACS cannot be held responsible for errors or consequences arising from the use of information contained in these "Just Accepted" manuscripts.



is published by the American Chemical Society. 1155 Sixteenth Street N.W., Washington, DC 20036

Published by American Chemical Society. Copyright © American Chemical Society. However, no copyright claim is made to original U.S. Government works, or works produced by employees of any Commonwealth realm Crown government in the course of their duties.

Central-to-Helical-to-Axial-to-Central Transfer of Chirality with a Photoresponsive Catalyst

Stefano F. Pizzolato, Peter Štacko, Jos C. M. Kistemaker, Thomas van Leeuwen, Edwin Otten and Ben L. Feringa*

Center for Systems Chemistry, Stratingh Institute for Chemistry and Zernike Institute for Advanced Materials, Faculty of Mathematics and Natural Sciences, University of Groningen, Nijenborgh 4, 9747 AG Groningen (The Netherlands)

KEYWORDS. chirality transfer · molecular switches · photoisomerization · atropisomers · photoresponsive catalyst · dynamic selectivity control

ABSTRACT: Recent advances in molecular design have displayed striking examples of dynamic chirality transfer between various elements of chirality, e.g. from central to either helical or axial chirality and vice versa. While considerable

progress in atroposelective synthesis has been made, it is intriguing to design chiral molecular switches able to provide selective and dynamic control of axial chirality with an external stimulus to modulate stereochemical functions. Here, we report the synthesis and characterization of a photoresponsive bis(2-phenol)substituted molecular switch **1**. The unique design exhibits a dynamic hybrid central-helical-axial transfer of chirality. The change of preferential axial chirality in the biaryl motif is coupled to the reversible switching of helicity of the overcrowded alkene core, dictated by the fixed stereogenic center. The potential for dynamic control of axial chirality was demonstrated by using (R)-**1** as switchable catalyst to direct the stereochemical outcome of the catalytic enantioselective addition of diethylzinc to aromatic aldehydes, with successful reversal of enantioselectivity for several substrates.



INTRODUCTION

Chirality plays a fundamental role in a myriad of biological processes, including information storage and transmission, gene expression, energy production and cellular motion.1-4 For instance, life has developed on Earth by optimizing its biological functions using Lamino acids as polypeptide building blocks and D-glucose as chemical energy source. The chirality of D-deoxyribose is amplified to the (almost) exclusively right handed helices of DNA.5 The supreme control of directional movement showcased by biological machine structures like ATP synthase,6 proteasomes,7 ribosomes,8 myosin,9 kinesin¹º and bacterial flagella¹¹ are astonishing demonstrations of how transfer of chiral information leads to accurate control of metabolic functions and motion in cells. None of these processes could take place without precise propagation, amplification and coupling of movement, from the very bottom scale of single molecular chiral entities to the fine interplay of large sub-units. While early research protein on stereochemistry mainly focused on point chirality, other

motifs that feature axial chirality,^{12,13} helical chirality,¹⁴ and planar chirality^{15,16} have been extensively investigated for their potential use in synthesis, in asymmetric catalysis and as chiral dopants. Compared with molecules that feature fixed central chirality (i.e. point chirality), axially chiral compounds may not comprise stereogenic center(s) yet exist as enantiomers.^{17,18} Atropisomers belong to the class of axially chiral compounds: in this case the enantiomers exist due to the restricted rotation around a single bond. The phenomenon of equilibration of stereoisomers about a rotational axis atropisomerization¹⁹ - has become a main topic of investigation in organic,20 materials,21 and medicinal chemistry.²² Despite a number of responsive molecular devices based on reversible cis-trans isomerization of double bonds,23-25 cyclizations,26 redox cycles27 and rotation around single bonds,^{21,28} only scattered examples of stimuli responsive systems featuring elements of axial chirality have been reported.25,29-31 Focused efforts have produced elegant systems displaying unidirectional arylaryl bond rotation of biaryl structures via sequential

ACS Paragon Plus Environment

addition of chemical stimuli, overcoming the atropisomerization energy barrier inherently featured by the open structures via more flexible macrocyclic or intermediates.^{21,28,32-35} tricyclic Solvent-dependent atropisomerism of a flexible 2,2'-biphenol core bridged tricyclic structure was reported by Reichert and Breit.36 However, a catalytic application in which the biphenol acts as a responsive ligand was not shown. Therefore, the major challenge remains to design chiral based molecular switches able to selectively and dynamically generate and harness biaryl axial chirality with an external stimulus.

1

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56 57

58 59

60

Combining dynamic chirality, chirality transfer, and photoswitches, our group has previously achieved control of activity and stereoselectivity by switchable catalysts^{29,37-} ³⁹ based on first generation molecular motors.⁴⁰⁻⁴⁴ These responsive catalysts harness the intra- and intermolecular transfer of chirality to ultimately control the stereochemical outcome of a catalytic transformation via photochemical E-Z isomerization and thermal helix inversions of a functionalized unidirectional four-stage rotary motor based on an overcrowded alkene core. By combining the fixed point chirality originating from the stereocenters in first generation molecular motors with the dynamic alkene configuration and helical chirality, the enantiomeric excess and configuration of the chiral product can be reversibly controlled. However, out of the four possible diastereoisomeric forms only the two pseudoenantiomeric cisoid isomers exert significant asymmetric induction via internal cooperative catalysis while an exclusive light-driven switchable catalyst based on these chiral motor remains elusive.

We anticipated that the development of new bis(2phenol)-functionalized switchable catalysts which harness the pairing of hybrid helical-axial chiralities within chiroptical switchable units could provide unprecedented levels of dual stereoselectivity with noninvasive control and high spatio-temporal resolution. Here we report the photochemical control of axial biaryl chirality in a light-responsive BINOL-type catalyst based on a chiral molecular switch, which displays dual stereocontrol in an asymmetric addition of organozinc reagents to aromatic aldehydes. The bis(2-phenol) unit operates as a chiral flexible bifunctional catalytic unit, the conformation of which is dictated by the dynamic stereochemistry of the central photoswitchable scaffold via internal dynamic transfer of chirality. Moreover, the catalyst described here benefits from a highly thermally stable switch core unit which can be reversibly photoisomerized exclusively between two pseudoenantiomeric forms.

RESULTS AND DISCUSSION

Design. Molecular motors of the second generation are helical-shaped overcrowded alkenes consisting of a symmetric tricyclic lower half and an asymmetric upper half that features a single stereocenter (Scheme 1).⁴⁵⁻⁴⁷ Utilizing the hybrid chirality generated by the stereogenic center and the helical structure during the photochemical

E-Z isomerization (PEZI) and thermal helix inversion (THI) of the central alkene bond allows unidirectional rotary motion controlled by a light- and heat-driven fourstage cycle (Scheme 1). The combinations of an upper half containing a six-membered ring and a lower half featuring a five membered ring, are characterized by a high activation energy for the thermal relaxation process and have been recently reported as a new class of bi-stable photoswitches.^{46–48} Due to the long half-life at room temperature, i.e. high thermal stability, of their photogenerated metastable isomers,⁴⁷ they allow for the design of systems capable of displaying dual stereocontrol while retaining the desired configuration for extended time intervals at elevated temperatures. This property, combined with their unique dynamic helical chirality, is highly desirable in the field of switchable asymmetric catalysis.

Scheme 1. Isomerization processes leading to unidirectional rotation in second generation molecular motor.



Four-stage cycle with only two distinct stereoisomers in case of symmetrically substituted lower half (here R = R'). S = stable isomer, MS = metastable isomer.

We envisioned that merging a flexible 2,2'-biphenol core with the rotor of a rigid second generation overcrowded alkene scaffold would result in transfer of chirality from the helical core of the overcrowded alkene to the biphenyl unit by steric interactions (Scheme 2). We expected the lower phenol group to remain parallel to the fluorene half, as a consequence of the system's tendency to reduce steric hindrance between these two moieties. In this way the distinctive dynamic helicity of the switch unit and the conformational versatility of the substituted biaryl motif are combined. The system described herein features three stereochemical elements (Scheme 2). The first element is the stereogenic center of the switch (highlighted in red), which can exist with either the *R* or *S* configuration. The second element is the helicity of the overcrowded alkene (highlighted in blue), which is

Page 3 of	f 17 Jou	rnal of the American Chemical Society
1	controlled by the configuration at the ster-	eogenic center
2	but can be inverted upon photoisomenzatio	<i>л</i> п.
3		
4		
5		
6		
7		
8		
9		
10		
11		
12		
13		
14		
15		
17		
18		
19		
20		
21		
22		
23		
24 25		
25 26		
20		
28		
29		
30		
31		
32		
33		
34		
35		
30		
38		
39		
40		
41		
42		
43		
44		
45		
40 47		
48		
49		
50		
51		
52		
53		
54		
55		

Page 4 of 17

Scheme 2. Design of photoswitchable 2,2'-biphenolsubstituted overcrowded alkene 1.



The assigned descriptors are based on the structure of compound (*R*)-1 (for explanation of the chiral descriptors, *vide infra*). Axial helicity and chirality (green) of the 2,2⁻ biphenol core are coupled to axial helicity (blue) and point chirality (red) of the molecular switch scaffold. Two diastereomers with opposite coupled helicity can be selectively addressed by irradiation with UV-light: (*R*,*P*,*S*_a)-1 (S); (*R*,*M*,*R*_a)-1 (MS).

More precisely, the more stable diastereoisomer (stable isomer, S) of the R enantiomer will adopt a P helicity, while the photo-generated diastereoisomer with higher energy (metastable isomer, MS) will adopt an *M* helicity. The axial chirality of the biaryl unit (highlighted in green) is the third stereochemical element which can be assigned either R_a or S_a according to the CIP rules.^{49,50} For biphenyls with an average dihedral angle of 90°, such as ortho substituted biphenyls, these stereochemistry descriptors are interchangeably used with M and P, respectively. Depending on the size of the groups and substitution pattern at the ortho positions, the dihedral angle can be smaller than 90°. Each rotamer with either R_a or S_a absolute configuration possesses two conformational helical geometries, also assigned as righthanded (P) or left-handed (M) according to the CIP rules.18

Recently our group reported a study on the tidal locking of an aryl moiety in a molecular motor, showing that among the four theoretically possible conformations of a biaryl unit, only conformations in which the nonannulated aryl group was parallel to the fluorenyl lower half were adopted.⁵¹ The other conformations with the aryl orientated perpendicular with respect to the lower half are expected to induce significant steric strain. With such a diastereotopic constraint, the true helicity (P_a/M_a) of the biaryl is inextricably connected to the helicity ($P_=/M_=$) of the overcrowded alkene chromophore, and is identical to it in each isomer (Figure 1b). Therefore, three stereodescriptors (R/S, P/M and R_a/S_a) will be sufficient for the assignment of any expected isomer reported in this work. So for isomer $(R, P_{=}, P_{a}, S_{a})$ -1: R = configuration of stereogenic center, $P_{=}$ = helicity of alkene, P_{a} = helicity of biaryl, S_{a} = axial chirality of biaryl (Figure 1a).



Figure 1. a) Example of top-down schematic view and front structural view of $(R,P_{=},P_{a},S_{a})$ -**1**. Upper half ring (red, methyl substituent omitted); fluorenyl lower half (blue); biaryl moiety (black). Assigned stereodescriptors based on the structure of compound (R)-**1** (see main text for details). b) Depiction of four conformations of the biaryl moiety as viewed from the top along the central double bond and biaryl single bond. c) Hydrogen-bonding assisted biaryl rotation of 2,2'-biphenol with inversion of stereochemistry. d) Schematic energy vs. biaryl torsional angle profile upon clockwise rotation of lower phenol group around the biaryl single bond in (R)-**1**.

2

3

4

5

6

7

8

9

58 59

60

The asterisks at the stereodescriptors throughout the text denote a racemic mixture of isomers with identical relative stereochemistry (e.g. R^*, P^*, S_a^* means a mixture of R, P, S_a and S, M, R_a). The doubly expressed axial stereodescriptor (R_a/S_a) throughout the text denotes a with identical mixture of rotamers absolute stereochemistry at the stereocenter and configurational helicity but opposite axial chirality (e.g. $R_{,P,S_{a}}/R_{a}$ means a mixture of atropisomers $R_1P_1S_2$ and $R_2P_1R_2$. The inversion of axial chirality in 2,2'-biphenols is likely to take place via a coplanar transition state along the syn-periplanar 10 conformation of the phenol rings taking advantage of the 11 intramolecular hydrogen bonds between the hydroxyl 12 groups ($\Delta^{\ddagger}G^{\circ} = 48.1 \text{ kJ mol}^{-1} \text{ T} = 298.15 \text{ K}$; Figure 1c), based 13 on a DFT study by Fujimura and co-workers.52 These 14 calculations support the proposal of reversible axial 15 chirality when applied to our system, as we expected the 16 syn- and anti-conformers (hydroxyl groups in proximity 17 or pointing away from each other, respectively) to be in 18 equilibrium in solution in the absence of metals or other 19 coordinating species. Recently our group has shown how 20 the behavior of a related biphenol-functionalized 21 molecular motor can be controlled via reversible covalent 22 modifications.53 and non-covalent Α schematic 23 representation of four possible conformations of 1 upon 24 rotation of the aryl-aryl bond is presented in Figure 1b. 25 We expect conformations with matching helicities of 26 biaryl and overcrowded alkene units to be highly favored 27 (A and C), while the two conformers with the aryl 28 perpendicular to the lower half to experience steric 29 hindrance (B and D), as shown in the relative energy vs. 30 torsional angle profile plot (see Figure 1d) based on DFT 31 calculations (vide infra). Scheme 3 illustrates the delicate 32 interplay of dynamic stereochemical elements and the 33 switching process between the stable and metastable 34 isomers of (*R*)-1. The rotamers of the stable state (R,P,R_a) -35 **1** and (R,P,S_a) -**1** interchange via atropisomerization (Scheme 3, left equilibrium) presumably facilitated by 36 37 internal hydrogen bonding between the two phenolic moieties.52 We envisioned that upon irradiation with UV-38 light of a mixture of (R, P, R_a) -1 and (R, P, S_a) -1 brings about 39 isomerization into the corresponding conformers of the 40 metastable state (R,M,R_a) -1 and (R,M,S_a) -1 isomers, 41 respectively, during which the upper half containing the 42 biaryl motif rotates with respect to the fluorenyl lower 43 half yielding isomers with opposite helicity $(P \rightarrow M)$. 44 Notably, the metastable isomer was also expected to 45 display atropisomerization (Scheme 3, right equilibrium). 46 We undertook a theoretical study a priori to verify the 47 design as shown in Figure 1, with particular attention to 48 the barrier for biaryl rotation and the relative energy of 49 the four accessible conformers upon reversible 50 irradiation. The structures of the four ground states were 51 computed via DFT method calculations (see 52 Supplementary material for details), which suggested an 53 energetic preference in both biaryl rotation equilibria for 54 the conformers (R,P,S_a) -1 and (R,M,R_a) -1, respectively 55 (Scheme 3). These conformers are characterized by 56 having the lower hydroxyl substituent pointing away from 57

the central overcrowded alkene in a syn conformation with the upper phenol group.

In summary, our design is based on the following elements: a) the selective and reversible photoisomerization of the overcrowded alkene scaffold between only two states; b) the unique change in helicity of the chiroptical switch governed by the fixed configuration of the stereogenic center; c) the coupled change in axial chirality of the biaryl core achieved via a central-tohelical-to-axial transfer of chirality; d) the introduction of the switchable chiral biphenol functionality with the potential of various applications in catalvtic enantioselective transformations.

Scheme 3. Schematic representation of switching process between the rotamers of stable and metastable isomers of (R)-1.



Proposed ground states of rotamers and transition states (middle) of atropisomerization processes, as viewed from the top along the axis given by the double bond. Top and bottom left: rotamers of stable isomer $(R,P,S_a/R_a)$ -1; top and bottom right: metastable isomer $(R, M, R_a/S_a)$ -1.

Synthesis. Key steps in the synthesis of 1 are the Barton-Kellogg coupling of thioketone 7 and diazoderivative 6, followed by deprotection of the biphenol moiety and chiral resolution of the target molecule 1 as illustrated in Scheme 4. Commercially available 7-Nmethoxy-1-tetralone was brominated with bromosuccinimide in acetonitrile to yield $2 (93\%)^{54}$, followed by Suzuki-Miyaura cross-coupling catalyzed by Pd₂dba₃ and SPhos to provide the dimethoxy-biaryl motif in ketone 3 (94%). α -Methylation provided ketone 4 (86%), which was converted to the corresponding hydrazone 5 (75%) via condensation with hydrazine monohydrate using $Sc(OTf)_3$ as a catalyst. The diazo coupling partner 6 was accessed via in situ oxidation with [bis(trifluoroacetoxy)iodo]-benzene at low temperature. Fluorene-9-thione 7, freshly synthesized by thionation of 9-fluorenone with Lawesson's reagent, was subsequently added to yield a variable mixture of episulfide **8** and overcrowded alkene **9**.

Scheme 4. Synthesis and chiral resolution of 2,2'-biphenol molecular switch 1.



Note on resolution of 1: i) result from first resolution; ii) $(S,M,R_a/S_a)$ -1 obtained by second resolution of the solid fraction: (8S,9R)-(–)-*N*-benzylcinchonidinium chloride 10 (0.9 equiv), 79% yield, >99% ee (solid); $(R,P,S_a/R_a)$ -1 obtained by second resolution of the residue from solution: 10 (0.3 equiv), 81% ee (residue from solution), followed by recrystallization from EtOH/H₂O = 1:1 of the residue from solution, 15% yield, 96% ee.

After separation, the remaining episulfide was desulfurized by treatment with HMPT at elevated temperature to provide **9** (85%, for the 3-step sequence). The use of boron tribromide, widely applied for the deprotection of methoxy-substituents, resulted in partial decomposition of the overcrowded alkene and in an inseparable mixture of target compound and sideproducts. Successful deprotection was accomplished using methyl magnesium iodide at 165 °C55 to afford racemic $(R^*, P^*, S_a/R_a)$ -1 (86%) as a mixture of two atropisomers in their thermodynamic equilibrium ratio of 60:40 (in CDCl₃) according to ¹H NMR analysis. Optical resolution of 1 was accomplished by two-step resolution with (8S, 9R)-(-)-*N*-benzylcinchonidinium chloride (10) in ethyl acetate.55 Both enantiomeric mixtures of conformers were obtained in high optical purity: $(R,P,S_a/R_a)$ -1 (96%) ee, 15%); $(S,M,R_a/S_a)$ -1 (>99% ee, 31%). The structure of 1 was unequivocally proven by NMR spectroscopy (vide infra), HRMS, as well as by single-crystal X-ray structure analysis. By means of a high-brilliance Cu IµS microfocus source (Cu K_{α} radiation wavelength = 1.54178 Å), the absolute configuration of enantiomerically pure (R)-1 was determined despite the absence of atoms that show significant anomalous scattering.56-58 The reconstructed unit cell of the lattice was shown to contain only the synconformer (R,P,S_a) -1 (see Figure 2). The experimental data confirmed the proposed model of coupled helical-to-axial transfer of helicity, demonstrating the most favored

conformation of the lower aryl substituent to be parallel to the fluorenyl lower half of the switch core (synclinal) in the crystal lattice. The dihedral angle over the biaryl motif determined from the X-ray structure in the solid state was found to be $+55.7^{\circ}$ ($+52.7^{\circ}$ by calculation, *vide infra*).



Figure 2. a) X-Ray structure of (R,P,S_a) -1. Left: front view; right: top view. Ellipsoids set at 50% probability. Hydrogen bond lengths (intra: H101–O1 1.874 Å, inter: H100–O1' 1.826 Å) and Oxygen-Oxygen distances (intra: O1–O2 2.629 Å, inter: O1–O2' 2.685 Å). b) Newman projections. Left: top view

through	overcrowded	alkene	bond	Right:	ton	view	through	h
through	overcrowded	aikciic	bonu.	rugint.	top	VICW	unougi	.1

- aryl-aryl bond of biaryl unit. Torsional angles of alkene unit
- , γ--- , απα υπη μ unit (55,7/*) are shown (13.92°) and biaryl unit (55.71°) are shown



Figure 3. a) Schematic representation of the photochemical *E-Z* isomerization of stable atropisomers $(R,P,S_a/R_a)$ -1 to metastable atropisomers $(R,M,R_a/S_a)$ -1. ¹H NMR spectra of (R)-1 (~5.0 mg, toluene- d_8 (0.7 mL), 25 °C): b) stable state $(R,P,S_a/R_a)$ -1 (A:B = 67:33); c) after irradiation with UV light

(365 nm) of (R)-1 to the metastable state (R,M, R_a/S_a)-1 (~65% of MS); d) metastable state (R,M, R_a/S_a)-1 (C:D = 58:42) isolated by preparative HPLC from the irradiated mixture (see Supplementary material for details).

The angles found by calculation agree with those found by X-Ray, and the small deviation between the two can be explained by a packing effect in the crystal structure which is expected to bring about small distortions with respect to a free molecule in solution or gas phase.

Atropisomerization process. The chiral resolution and initial characterization of 1 by ¹H NMR (Figure 3b) disclosed a very interesting yet initially unexpected phenomenon. Compound 1 was obtained from synthesis as pure stable isomer and could be resolved in two enantiomerically pure fractions, which by chiral HPLC analysis appeared as single eluted fractions with sharp symmetric peaks (see Supplementary material for details). However, both racemic and enantiopure fractions comprised two inseparable species, as displayed by ¹H NMR spectroscopy analysis. We observed two sets of absorptions in the ¹H NMR spectra recorded with a solution of stable (*R*)-1 in toluene-*d*₈ (Figure 3b): the relative integration of the best resolved peaks of the Page 8 of 17

atropisomers revealed a ratio of A:B = 67:33 [methyl protons: δ 1.34 ppm (A), 1.22 ppm (B); proton H1 at the stereogenic center: δ 3.95 ppm (A), 3.86 ppm (B); proton H₂ of fluorenyl substituent: δ 7.78 ppm (A), 7.70 ppm (B)]. Based on calculated ¹H NMR spectra (see Table S₃-S₄ and Figure S₃₀ in Supporting material), we assigned the experimental sets of absorptions A and B to the atropisomers (R,P,S_a) -1 and (R,P,R_a) -1, respectively (Figure 3a, see Supporting material for atropisomer assignment study). Similar behavior with minor variation in the ratio was observed in other deuterated solvents (see Supporting material for details). Based on our design, we assumed the syn and anti atropisomers to equilibrate via biaryl rotation (Scheme 3). Initial attempts to determine the rate of the atropisomerization process via dynamic NMR focused on the coalescence of the aforementioned diagnostic absorption peaks in the aromatic region (see Figure 3b).⁵⁹⁻⁶¹ ¹H NMR spectra (400 MHz) of a sample of stable isomer $(R, P, S_a/R_a)$ -1 in toluene- d_8 , were recorded at temperatures ranging from 50 to 100 °C.62 No coalescence of the aforementioned diagnostic absorption peaks was observed, suggesting the activation barrier for the biaryl rotation process to be higher than typical exchange processes usually determined via Dynamic NMR (see Supporting material for details).⁶³ Dynamic HPLC (DHPLC) analysis was also considered, as it was previously reported to allow for the successful determination of rotational barriers for other substituted biphenyl atropisomers.⁶⁴⁻⁶⁷ Despite the screening of temperatures down to o °C, no splitting of the elution peaks was observed, indicative of a relatively fast equilibration process even at lower temperatures. The rotational process was eventually demonstrated and studied by one dimensional exchange spectroscopy (EXSY, ¹H-¹H nuclear Overhauser enhancement spectra). The measurements were conducted in the temperature range of 39.2-60.9 °C, consisting of an arrayed cluster of multiple mixing times per temperature (see Supporting material for details). Exponential curve fitting of the EXSY traces provided the exchange constant (k) and associated standard error (σ_k) for each temperature. Eyring analysis of rate versus temperature afforded the Gibbs free energy of activation for the isomerization of the major into the minor atropisomer ($\Delta^{\pm}G_{\text{BI}}$ at rt = 78.2±1.1 kJ mol⁻¹). The equilibration time at room temperature (20 °C) is extrapolated to be in order of minutes ($t_{\frac{1}{2}} = 1.2 \pm 0.4 \text{ min}$), while the temperature at which t_½ equals one hour is calculated to be equal to -50.5 ± 0.5 °C. This analysis explains why the isolation of atropisomers was not successful, as well as the lack of coalescence in the ¹H NMR spectrum even at higher temperatures and the unresolved elution profile in the analytical HPLC chromatograms. Notably, when the isolated metastable isomer $(R,M,R_a/S_a)$ -1 (via preparative HPLC, vide infra) was subjected to the same EXSY experiments no exchange was observed (measurement temperatures up to 60 °C). This observation is in accordance with the larger elution band of the metastable isomers fraction compared with the elution band of the stable isomers observed in the analytical HPLC analysis (see Supporting material for

2

3

4

5

6

7

8

9

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51 52

53

54

55

60

details), which suggests a lower atropisomerization rate hence a slightly higher activation barrier for the biaryl rotation process in the metastable isomers. As observed in the X-ray structure analysis and based on the model investigated by Fujimura and co-workers51, we propose a thermodynamically favored cyclic seven-membered ring conformation generated upon internal coordination via hydrogen bonding of the two hydroxyl substituents (see Scheme 3). Experimental evidence and calculation data suggest that such a conformation provides access to a transition state with a relatively low barrier for 10 atropisomerization, allowing for a fast exchange of two 11 conformers in solution at room temperature. In these two 12 calculated transition states $(TS_{BI}-(R,P,Syn)-1 + and TS_{BI}-$ 13 (*R*,*M*,*Syn*)-1[‡], see Figure S25 in Supplementary material) 14 the hydrogen bond between the two phenol moieties is 15 shorter than it is in any other conformation suggesting 16 additional stabilization of the transition state with respect 17 to its corresponding minima explaining the relatively low 18 barrier for atropisomerization. Moreover, the barrier for 19 biaryl rotation is sufficiently low to allow the desired syn 20 atropisomer to act as a thermodynamic sink upon its 21 depletion in a reaction selective for it, for instance by 22 bidentate coordination of the biphenol to a metal center 23 (vide infra, Scheme 5a). Such a complexation would 24 require a syn conformation of the biaryl motif and 25 concordant alkene and biaryl helicity, due to the system's 26 tendency to reduce steric hindrance between the lower 27 phenol moiety and the fluorenyl lower half (Figure 1; vide 28 infra, Scheme 5b). 29

NMR spectroscopy. In order to investigate the photochemical behavior of 1 in more detail, an NMR sample of stable isomer $(R,P,S_a/R_a)$ -1 in toluene- d_8 was irradiated with UV light (365 nm) for 30 min at room temperature. ¹H NMR spectra were taken before (Figure 3b) and after exposure (Figure 3c). Upon irradiation two new sets of absorptions, C and D, with intensities increasing over time were obtained [methyl protons: δ 1.22 ppm (C+D, peaks not resolved); proton H1 at the stereogenic center: δ 3.75 ppm (C), 3.35 ppm (D); proton H2 of fluorenyl substituent: δ 7.61 ppm (C), 7.47 ppm This is indicative of the photo-induced (D)]. isomerization to the metastable isomer $(R_1M_1, R_2/S_2)$ -1 comprising of two distinct atropisomeric species, (R,M,R_a) -1 (C) and (R,M,S_a) -1 (D), respectively. The relative integration revealed a final ratio in toluene- d_8 of $(R,P,S_a/R_a)$ -1 (A+B) : $(R,M,R_a/S_a)$ -1 (C+D) = 35:65, upon irradiation over 30 min. It should be specified that the PSS was not reached in order to avoid degradation upon prolonged exposure. Due to the high thermal stability of the metastable isomers $[(R,M,R_a/S_a)-1]$, their isolation from a crude mixture of an irradiated solution of (R)-1 was achieved by preparative HPLC (see Supplementary material for details). Analysis by 'H NMR revealed the metastable isomer to comprise a mixture of atropisomers (R,M,R_a) -1 (C) and (R,M,S_a) -1 (D) in a ratio of C:D = 58:42 (Figure 3d).

Photochemical isomerization. The switching properties of (*R*)-1 were monitored by UV-vis absorption and circular dichroism (CD) spectroscopy (Figure 4). A schematic representation of the reversible photochemical *E-Z* isomerization process of (R)-1 is shown in Figure 4a. A solution of stable $(R,P,S_a/R_a)$ -1 (toluene, 4.5 10⁻⁵ M) in quartz cuvettes was purged with argon and irradiated at room temperature towards either the metastable isomer using UV light (365 nm, Figure 4b, black to red gradient) or the stable isomer using visible light (420 nm, Figure 4c, red to blue gradient). The reversible photochemical E-Z isomerization was found to be characterized by a clear isosbestic point at 368 nm, indicating the absence of side reactions. A bathochromic shift of the major absorption band ($\pi \rightarrow \pi^*$) of about 40 nm was observed, indicative of an increase in alkene strain and consistent with other second generation motors and switches as is expected for the metastable form $(R,M,R_a/S_a)$ -1.⁴⁸ The sample was subsequently subjected to irradiation cycles (see Figure S6 in Supporting material), displaying non-perfect switching fatigue resistance with minor decomposition, as opposed to the highly resistant unfunctionalized parent compounds recently studied.48 However, upon addition of the radical scavenger TEMPO (10-5 M) to a solution of (R)-1 (toluene, $4.0\cdot10^{-5}$ M), we observed no evidence of degradation after six irradiation cycles (Figure 4d). This observation suggests that radicals may be involved in the decomposition process.⁶⁸ Lastly, a solution of stable $(R,P,S_a/R_a)$ -1 (toluene, 4.5 10⁻⁵ M) was subjected to CD spectroscopy in order to perform a qualitative analysis of the change in its helical structure (Figure 4e). The CD spectrum displayed a strong Cotton effect in the area of 320-370 nm. Upon irradiation with 365 nm light an inversion of the absorption band was observed, which is indicative of an inversion in helicity and shows that the photochemical isomerization of the stable isomers $(R,P,S_a/R_a)$ -1 to the metastable isomers $(R,M,R_a/S_a)$ -1 has occurred. Upon irradiation with 420 nm light, the original absorption band could be partially recovered. Chiral HPLC analysis confirmed the presence of the metastable species and allowed determining the S:MS ratio at the PSS of the irradiated mixtures upon selection of detector wavelength at the isosbestic point (368 nm, see Supplementary material for details). An efficient photoswitching process was observed upon irradiation with 365 nm light, with a high ratio towards the metastable diastereoisomer (S:MS = 17:83) at the PSS₃₆₅. However, the reverse process achieved upon irradiation at 420 nm light was found to be less selective, affording an equimolar mixture of stable and metastable isomers (S:MS = 50:50) at the PSS₄₂₀.

Switchable asymmetric catalysis. Having established the reversible switching process between $(R, P, S_a/R_a)$ -1 and $(R,M,R_a/S_a)$ -1, we investigated their abilities for dual stereocontrol in a model asymmetric catalysis reaction.³⁷



Figure 4. a) Schematic representation of photochemical *E-Z* isomerization of stable isomer (R,P, S_a/R_a)-1 to metastable isomer (R,M, R_a/S_a)-1. b) Experimental UV-vis absorption spectra of stable (R,P, S_a/R_a)-1 (toluene, 4.5·10⁻⁵ M, black) and upon irradiation with UV-light (365 nm) of (R,P, S_a/R_a)-1 towards the metastable isomer affording a PSS₃₆₅ mixture (S/MS = 17:83, red) with an isosbestic point at 368 nm. c) Experimental UV-vis absorption spectra after irradiation of the PSS₃₆₅ sample using visible light (420 nm), resulting in reversed *E-Z* isomerization towards the stable isomer affording a new PSS₄₂₀ mixture (S/MS = 50:50). d) Irradiation cycles of (R)-1 (toluene, ~4.0·10⁻⁵ M) in the presence of TEMPO (~10⁻⁵ M) towards opposite PSS mixtures (red: 365 nm, 4 min; blue: 420 nm, 15 min). e) Experimental and calculated CD spectra of (R)-1 (toluene, 5.0·10⁻¹ M): black, starting stable isomer (R,P, S_a/R_a)-1; red: CD spectra of PSS₃₆₅ mixture; cyan: metastable isomer (R,M, R_a/S_a)-1. Note: PSS ratios determined by HPLC analysis of the irradiated solutions via quantitative analysis with PDA detector wavelength set at the isosbestic point (368 nm).

As a proof of principle, we envisioned to use compound (R)-1 as a switchable bidentate ligand, which could coordinate a metal center and eventually be applied to an asymmetric transformation acting as a tunable stereoselective catalyst (vide infra, Scheme 5). We anticipated the isomers of 1 having an anti conformation of the biphenol unit (torsion angle = $\pm 90^{\circ}$ -180°; hydroxyl groups pointing away from each other) to be poor bidentate ligands. Therefore only the isomers with syn conformation (torsion angle = $0^{\circ}-\pm 90^{\circ}$; hydroxyl groups in proximity) were expected to efficiently bind metal ions in a bidentate fashions common to binaphthols (BINOL's) and successfully transfer the chirality within a catalytically active complex (Scheme 5a,b). Hence we proposed that the tunable helicity (P or M) of the switch core in turn would dictate the preferential axial configuration (R_a or S_a) of the desirable syn conformation of the biaryl moiety and eventually, for instance, the configuration (R or S) of a newly formed stereogenic center when applied to an enantioselective catalytic event (Scheme 5c). Zn-BINOL-derived complexes have previously been reported to successfully mediate the catalytic asymmetric aldol⁶⁹⁻⁷¹ and hetero-Diels-Alder⁷² reactions. We decided to use compound (R)-1 as a

switchable bidentate ligand in 1,2-additions of diethylzinc to benzaldehydes. Numerous efforts have been devoted in the past decades to develop new effective chiral ligands for asymmetric addition of diethylzinc to benzaldehyde.73-77 However, only few cases have been reported in which dual stereocontrol was achieved by tuning the reaction conditions. The switching of enantioselectivity in the catalytic addition of diethylzinc to aldehydes was obtained by changes in the reaction conditions (e.g. solvent, temperature) while using the same chiral additive.78-81 Alternatively, complementary catalytic systems were developed by the use of distinct structural derivatives from a common chiral catalyst scaffold to access both enantiomers of the desired products.82-85 In the representative reaction (see scheme in Table 1), benzaldehyde 11a was added to a mixture of ligand (R)-1 and a solution of diethylzinc in toluene, yielding secondary alcohol 12a and in an amount of the sideproduct benzyl alcohol 13a. The latter is the product of the aldehyde reduction, a known process occurring in the case of a slow addition process and proposed to derive from the β -hydride elimination of organozinc species and subsequent reduction of the substrate in case of poorly activated zinc complexes.^{86,87} As we anticipated, photo-

60

1	induced switching of ligand reversal of stereoselectivity	(<i>R</i>)-1 allowed successful in the 1,2-addition of	
2	diethylzinc to benzaldehyde.		
3			
4 5			
6			
7			
8			
9			
10			
11			
12			
14			
15			
16			
17			
18			
19			
20			
22			
23			
24			
25			
26 27			
27			
29			
30			
31			
32			
33			
24 35			
36			
37			
38			
39			
40 41			
41 42			
43			
44			
45			
46			
4/ /8			
40 49			
50			
51			
52			
53			
54 55			
55 56			
57			
58			
59			-
60		ACS Paragon Plus Environme	ent

Scheme 5. Schematic representation of mono- and bidentate coordination equilibrium upon reaction of (R)-1 with organozinc reagents.



a) Depiction of the possible mono-and bidentate coordination species upon reaction of stable isomers of **1** with ZnR_2 . b) Only the isomers with a syn conformation (torsion angle = $o^\circ - \pm 9o^\circ$) were expected to efficiently bind a metal center and successfully transfer the chirality within a catalytically active complex. c) Light-assisted dual stereocontrol could be achieved in a catalyzed organometallic reaction upon photoisomerization of (*R*)-**1** and internal transfer of chirality to the coordinated metal site.

The results of the catalysis experiments are presented in Table 1. 'H NMR analysis allowed determining the conversion and selectivity of organozinc addition versus aldehyde reduction to benzylic alcohol. The enantiomeric excess (ee) of chiral secondary alcohols **12a-g** was determined by chiral HPLC or GC analysis. In addition to benzaldehyde, several para- and ortho-substituted

aromatic aldehydes bearing electron-withdrawing or electron-donating groups were tested as substrates. In all cases, when the stable form $(R,P,S_a/R_a)$ -1 was used as a catalyst, the preferred formation of the (*R*)-enantiomer of secondary alcohols 12 was observed (Table 1, odd-number entries), with ee's up to 68% (entry 1).88 In sharp contrast, upon use of the irradiated mixture of catalyst $(R,P,S_a/R_a)$ -1 (365 nm light, PSS ratio S:MS = 17:83), the addition proceeded with reversed enantioselectivity under the same reaction conditions. Preferred formation of the (S)enantiomer of secondary alcohols 12 was also observed in all cases after irradiation (even-number entries), with ee's up to 55% (entry 8). The difference in enantioselectivity (Δee) between non-irradiated and irradiated catalyst solution was up to 113% (from 68% (R) to 45% (S), comparison of entries 1-2). Notably, use of diisopropylzinc led to no enantioselectivity in either case (entries 13-14).75 A control experiment was conducted by performing the addition of diethylzinc in absence of (R)-1, which resulted in a marked decrease in conversion, as well as lower addition vs. reduction selectivity and isolated yield of 12a (entry 15).⁸⁹ Addition of tetrabutylammonium bromide did not improve the catalytic activity, as otherwise observed in previously reported systems.90 It should be emphasized that no decomposition, racemization or significant thermal relaxation of the recovered catalyst (90% average catalyst recovery) was observed, as determined by ¹H NMR and chiral HPLC analysis (see Supplementary material for details). Moreover, in all cases the catalyst was recovered after an experiment conducted using a non-irradiated catalyst solution which was recycled to perform a subsequent experiment with the same substrate after irradiation of catalyst solution without notable loss of catalytic performance (see Supplementary material for further details). Point chirality of 1 governs the dynamic helical chirality, which in turn is coupled to the axial chirality resulting in a syn conformation in the catalytically active ligand species. The inversion of enantioselectivity is an indication of the reversed local chirality around the alkyltransferring zinc center and the coordinated aldehyde, achieved by using a ligand with opposite chiral induction.⁸²⁻⁸⁴ In this system we propose that upon irradiation and subsequent inversion of the biaryl axial chirality, the metastable isomer $(R,M,R_2/S_2)$ -1 resembles the enantiomer of the stable isomer $(R_1P_1S_2/R_2)$ -1 (Scheme 5c). As the flexible biphenol unit is the chiral ligand for zinc, tunable chiral induction can be achieved in the proximity of the zinc- aldehyde substrate complex, providing a novel approach for light-assisted dual stereocontrol in catalysis.

CONCLUSIONS

The design, synthesis and resolution of a photoresponsive molecular switch featuring a versatile 2,2'-biphenol motif in which chirality is transferred across three stereochemical elements is described.

Table 1: Dynamic enantioselective addition of organozinc to aromatic aldehydes with (*R*)-1.

R 11	$H + ZnR_2 \xrightarrow{(R)-1 (10 \text{ mol}\%)}_{\text{toluene}} + R_{11} \xrightarrow{(R)-1 (10 \text{ mol}\%)}_{\text{toluene}}$				OH HO HO (R Stable	HO HO HO $(R,P,S_a)-1$ Stable isomer (S) $\lambda = 365 \text{ nm}$ S:MS = 17:83 $(R,M,R_a)-1$ Metastable isomer			
Entry ^a	11	R	Catalyst	Conv. (%) ^b	Yield of 12 (%) ^d	ee of 12 (%) ^c	⊿ee of 12 (%) ^c	12:13 ^b	
1		Et	(<i>R</i>)-1	>95	86	68 (R)-12a	212	93:7	
2	11a	Et	(R)-1 + 365 nm	>95	87	45 (S)-12a	113	93:7	
3		Et	(R)-1	94	80	35 (R)-12b		81:19	
4		Et	(<i>R</i>)- 1 + 365 nm	>95	80	24 (S)- 12b	59	81:19	
5		Et	(R)-1	94	87	40 (R)- 12C	87	89:11	
6	11c	Et	(R)-1 + 365 nm	>95	86	42 (S)-12C	02	88:12	
7	O H	Et	(R)-1	66	37	40 (R)- 12d		62:38	
8	∽₀∕√ 11d	Et	(<i>R</i>)-1 + 365 nm	>95	76	55 (S)- 12d	95	97:3	
9	, Charles and Char	Et	(R)-1	>95	58	48 (R)-12e	08	63:37	
10	11e	Et	(<i>R</i>)- 1 + 365 nm	>95	79	50 (S)- 12e	90	85:15	
11		Et	(R)-1	>95	81	46 (R)- 12f		89:11	
12	11f	Et	(R)-1 + 365 nm	>95	72	31 (S)- 12f	//	83:17	
13		<i>i</i> -Pr	(R)-1	95	40	<5 (±)-12g	NA	41:59	
14	11a	<i>i</i> -Pr	(<i>R</i>)-1 + 365 nm	>95	57	<5 (±)-12g	11.21	58:42	
15		Et	/	59	24	N.A.		80:20	

^a **General reaction conditions:** 0.0125 mmol of $(R,P,S_a/R_a)$ -1 in 0.5 mL of dry toluene at 0 ° C; 0.375 mmol of R_2Zn (Et₂Zn, 1.0 M in hexane; *i*-Pr₂Zn, 1.0 M in toluene) added dropwise and stirred over 10 min; 0.125 mmol of 11 added to the mixture. Reaction mixture stirred for 7 d at 0 ° C. **Reaction with irradiated mixture of** (*R*)-1: 0.00125 mmol of $(R,P,S_a/R_a)$ -1 in 15 mL of dry, degassed Et₂O, irradiated with UV-light (365 nm) for 30 min until the PSS was reached (S:MS = 17:83). PSS ratio determined by chiral HPLC analysis. Reaction procedure follows as described above. ^b Determined by 'H NMR analysis of crude. ^c Determined by chiral GC or chiral HPLC analysis of isolated product. ^d Isolated yield. Abbreviations: N.A., Not Applicable.

The comparison of experimental and computational data confirmed the proposed model of coupled central-tohelical-to-axial transfer of chirality, demonstrating the most favored conformation of the lower aryl substituent of the biaryl unit to be parallel to the fluorenyl lower half of the switch core. Compared with previously reported molecular motor based systems, the reduction from four to two isomerization stages featured by the biarylfunctionalized design described herein provides a simple, reusable and efficient dynamic responsive chiral core. Extensive studies based on chiral HPLC analysis, ¹H NMR, UV-vis absorption and CD spectroscopy proved the reversible photoswitching of 1, with no fatigue over multiple cycles in presence of substoichiometric amount of TEMPO. The chirality transfer was successfully applied to creation of another stereogenic element as

demonstrated via dynamic central-to-helical-to-axial-tocentral transfer of chirality by using (R)-1 as switchable catalyst in the enantioselective addition of diethylzinc to various benzaldehydes. Reversal of enantioselectivity was accomplished for each substrate, with ee values for12 up to 68%, $\Delta ee's$ up to 113% and yields up to 87%. These results achieved in switchable asymmetric catalysis highlight the proof-of principle of a two-stage dynamically responsive chiral biaryl-functionalized switch scaffold. The future development of analogous biarylswitch structures combined with the well-established role of biaryls as privileged structures in asymmetric catalysis may lead to the construction of a variety of unprecedented switchable chiral catalysts that could perform multiple enantioselective transformation in a sequential manner. In addition, this chiral switch system has considerable potential as responsive chirality selector for a range of other applications beyond the field of asymmetric catalysis, such as control of supramolecular architecture, host-guest interaction, and polymer or liquid crystal morphology.

ASSOCIATED CONTENT

Experimental procedures, supporting characterization data, crystallographic detail and data, supporting irradiation and EXSY kinetic experiments, computational details, catalysis procedures, copies of the ¹H and ¹³C NMR spectra and HPLC chromatograms are presented in the Supporting material. This material is available free of charge via the Internet at <u>http://pubs.acs.org</u>.

AUTHOR INFORMATION

Corresponding Author

*b.l.feringa@rug.nl

Notes

1

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58 59

60

The authors declare no competing financial interest.

ACKNOWLEDGMENT

Financial support from The Netherlands Organization for Research (NWO-CW), Foundation Scientific for Fundamental Research on Matter (FOM, a subsidiary of NWO), the Zernike Institute for Advanced Materials, The Royal Netherlands Academy of Arts and Sciences (KNAW), The European Research Council (Advanced Investigator Grant no 694345 to BLF), Ministry of Education, Culture and (Gravitation program 024.001.035), Science and the University of Groningen are acknowledged. The authors would like to thank Ing. P. van der Meulen for the technical support during the EXSY experiments.

ABBREVIATIONS

PEZI, photochemical *E-Z* isomerization; THI, thermal helix inversion; S, stable; MS, metastable; PSS, photostationary state; CD, circular dichroism; DFT, density functional theory; ee, enantiomeric excess.

REFERENCES

- Eliel, E. L.; Wilen, S. H. Stereochemistry of organic compounds; Wiley: New York, 1994.
- (2) Yamamoto, H.; Carreira, E. M. *Comprehensive chirality*; Elsevier Science: Oxford, 2012.
- (3) Blackmond, D. G. Cold Spring Harb. Perspect. Biol. 2010, 2 (5), a002147.
- Bentley, R. In Encyclopedia of Molecular Cell Biology and Molecular Medicine; Wiley-VCH Verlag GmbH & Co. KGaA: Weinheim, Germany, 2006.
- (5) Timsit, Y.; Youri. DNA Self-Assembly: From chirality to evolution. *Int. J. Mol. Sci.* 2013, *14* (4), 8252–8270.
- (6) Boyer, P. D. Molecular motors: What makes ATP synthase spin? *Nature* 1999, 402 (6759), 247–249.
- (7) Murata, S.; Yashiroda, H.; Tanaka, K. Molecular mechanisms of proteasome assembly. *Nat. Rev. Mol. Cell Biol.* 2009, 10 (2), 104–115.
- (8) Ramakrishnan, V. Ribosome structure and the mechanism

of translation. *Cell* **2002**, *1*08 (4), 557–572.

- (9) Lodish, H.; Berk, A.; Zipursky, S. L.; Matsudaira, P.; Baltimore, D.; Darnell, J. In *Molecular Cell Biology. 4th edition, Section 18.3*; W. H. Freeman: New York, 2000.
- (10) Berg, J.; Tymoczko, J.; Stryer, L. In *Biochemistry. 5th edition,* Section 34.3; W H Freeman: New York, 2002.
- (11) Berg, H. C.; Anderson, R. A. Bacteria swim by rotating their flagellar filaments. *Nature* **1973**, *245* (5425), 380–382.
- Bringmann, G.; Price Mortimer, A. J.; Keller, P. A.; Gresser, M. J.; Garner, J.; Breuning, M. Atroposelective synthesis of axially chiral biaryl compounds. *Angew. Chem., Int. Ed.* 2005, 44 (34), 5384-5427.
- (13) Bringmann, G.; Gulder, T.; Gulder, T. A. M.; Breuning, M. Atroposelective total synthesis of axially chiral biaryl natural products. *Chem. Rev.* 2010, *11* (2), 563–639.
- Rickhaus, M.; Mayor, M.; Juríček, M. Strain-induced helical chirality in polyaromatic systems. *Chem. Soc. Rev.* 2016, 45 (6), 1542–1556.
- (15) Bolm, C.; Muñiz, K. Asymmetric Claisen rearrangement. Chem. Soc. Rev. 1999, 28 (1), 51–59.
- (16) Sokolov, V. I. Chirality and Optical Activity in Organometallic Compounds; Gordon and Breach Science Publishers: New York, 1990.
- (17) Mislow, K. Introduction to Stereochemistry; Dover Publications: Dover, 2003.
- (18) Nguyen, T. Giving atropisomers another chance. C. & En. News, 2018, 96, 33.
- (19) Oki, M. Recent Advances in atropisomerism. Top. Stereochem. 1983, 14, 1–81.
- (20) Leroux, F. Atropisomerism, biphenyls, and fluorine: a comparison of rotational barriers and twist angles. *Chem. Bio. Chem.* 2004, 5 (5), 644 – 649.
- (21) Collins, B. S. L.; Kistemaker, J. C. M.; Otten, E.; Feringa, B. L. A chemically powered unidirectional rotary molecular motor based on a palladium redox cycle. *Nat. Chem.* 2016, 8 (9), 860–866.
- LaPlante, S. R.; D. Fader, L.; Fandrick, K. R.; Fandrick, D. R.; Hucke, O.; Kemper, R.; Miller, S. P. F.; Edwards, P. J. Assessing atropisomer axial chirality in drug discovery and development. J. Med. Chem. 2011, 54 (20), 7005-7022.
- (23) Bandara, H. M. D.; Burdette, S. C. Photoisomerization in different classes of azobenzene. *Chem. Soc. Rev.* 2012, 41 (5), 1809-1825.
- Feringa, B. L.; van Delden, R. A.; Koumura, N.; Geertsema,
 E. M. Chiroptical molecular switches. *Chem. Rev.* 2000, 100 (5), 1789-1816.
- (25) Erbas-Cakmak, S.; Leigh, D. A.; McTernan, C. T.; Nussbaumer, A. L. Artificial molecular machines. *Chem. Rev.* 2015, *115* (18), 10081–10206.
- (26) Irie, M.; Fukaminato, T.; Matsuda, K.; Kobatake, S. Photochromism of diarylethene molecules and crystals: memories, switches, and actuators. *Chem. Rev.* 2014, 114 (24), 12174–12277.
- (27) Canary, J. W. Redox-triggered chiroptical molecular switches. *Chem. Soc. Rev.* **2009**, 38 (3), 747–756.
- (28) Kelly, T. R.; De Silva, H.; Silva, R. A. Unidirectional rotary motion in a molecular system. *Nature* 1999, 401 (6749), 150– 152.
- (29) Blanco, V.; Leigh, D. A.; Marcos, V. Artificial switchable catalysts. *Chem. Soc. Rev.* 2015, 44 (15), 5341–5370.
- (30) Kottas, G. S.; Clarke, L. I.; Horinek, D.; Michl, J. Artificial molecular rotors. *Chem. Rev.* 2005, 105 (4), 1281–1376.
- (31) De Bo, G.; Leigh, D. A.; McTernan, C. T.; Wang, S. A complementary pair of enantioselective switchable organocatalysts. *Chem. Sci.* 2017, 8 (10), 7077–7081.
- (32) Fletcher, S. P.; Dumur, F.; Pollard, M. M.; Feringa, B. L. A reversible, unidirectional molecular rotary motor driven by chemical energy. *Science* 2005, 310 (5745), 80–82.
- (33) Lin, Y.; Dahl, B. J.; Branchaud, B. P. Net directed 180° arylaryl bond rotation in a prototypical achiral biaryl lactone synthetic molecular motor. *Tetrahedron Lett.* 2005, *46* (48),

2

8359-8362.

- (34) Dahl, B. J.; Branchaud, B. P. 180° Unidirectional bond rotation in a biaryl lactone artificial molecular motor prototype. Org. Lett. 2006, 8 (25), 5841–5844.
- (35) Anyika, M.; Gholami, H.; Ashtekar, K. D.; Acho, R.; Borhan, B. Point-to-axial chirality transfer—a new probe for "sensing" the absolute configurations of monoamines. J. Am. Chem. Soc. 2014, 136 (2), 550–553.
- (36) Reichert, S.; Breit, B. Development of an axial chirality switch. *Org. Lett.* 2007, *9* (5), 899–902.
- (37) Escorihuela, J.; Burguete, M. I.; Luis, S. V. New advances in dual stereocontrol for asymmetric reactions. *Chem. Soc. Rev.* 2013, 42 (12), 5595-5617.
- (38) Vlatkovi ć, M.; Collins, B. S. L.; Feringa, B. L. Dynamic responsive systems for catalytic function. *Chem. - Eur. J.* 2016, 22 (48), 17080-17111.
- (39) Romanazzi, G.; Degennaro, L.; Mastrorilli, P.; Luisi, R. Chiral switchable catalysts for dynamic control of enantioselectivity. ACS Catal. 2017, 7 (6), 4100-4114.
- (40) Wang, J.; Feringa, B. L. Dynamic control of chiral space in a catalytic asymmetric reaction using a molecular motor. *Science* 2011, 331 (6023), 1429-1432.
- (41) Vlatković, M.; Bernardi, L.; Otten, E.; Feringa, B. L. Dual stereocontrol over the Henry reaction using a light- and heat-triggered organocatalyst. *Chem. Commun.* 2014, 50 (58), 7773–7775.
- (42) Zhao, D.; Neubauer, T. M.; Feringa, B. L. Dynamic control of chirality in phosphine ligands for enantioselective catalysis. *Nat. Commun.* **2015**, *6*, 6652.
- (43) Zhao, D.; van Leeuwen, T.; Cheng, J.; Feringa, B. L. Dynamic control of chirality and self-assembly of double-stranded helicates with light. *Nat. Chem.* 2016, 9 (3), 250-256.
- (44) Vlatković, M.; Volarić, J.; Collins, B. S. L.; Bernardi, L.;
 Feringa, B. L. Org. Biomol. Chem. 2017, 15 (39), 8285-8294.
- (45) Koumura, N.; Geertsema, E. M.; van Gelder, M. B.; Meetsma, A.; Feringa, B. L. Second generation light-driven molecular motors. unidirectional rotation controlled by a single stereogenic center with near-perfect photoequilibria and acceleration of the speed of rotation by structural modification. J. Am. Chem. Soc. 2002, 124 (18), 5037-5051.
- (46) Klok, M.; Walko, M.; Geertsema, E. M.; Ruangsupapichat, N.; Kistemaker, J. C. M.; Meetsma, A.; Feringa, B. L. New mechanistic insight in the thermal helix inversion of second-generation molecular motors. *Chem. Eur. J.* 2008, 14 (35), 1183-1193.
- (47) Vicario, J.; Walko, M.; Meetsma, A.; Feringa, B. L. Fine tuning of the rotary motion by structural modification in light-driven unidirectional molecular motors. *J. Am. Chem. Soc.* 2006, *128* (15), 5127-5135.
- (48) Kistemaker, J. C. M.; Pizzolato, S. F.; van Leeuwen, T.; Pijper, T. C.; Feringa, B. L. Spectroscopic and theoretical identification of two thermal isomerization pathways for bistable chiral overcrowded alkenes. *Chem. - Eur. J.* 2016, 22 (38), 13478-13487.
- (49) Cahn, R. S.; Ingold, C.; Prelog, V. Specification of molecular chirality. Angew. Chem., Int. Ed. 1966, 5 (4), 385-415.
- (50) Prelog, V.; Helmchen, G. Basic principles of the CIP-System and proposals for a revision. *Angew. Chem., Int. Ed.* 1982, 21 (8), 567-583.
- (51) Š tacko, P.; Kistemaker, J. C. M.; van Leeuwen, T.; Chang, M.-C.; Otten, E.; Feringa, B. L. Locked synchronous rotor motion in a molecular motor. *Science* 2017, 356 (6341), 964-968.
- (52) Sahnoun, R.; Koseki, S.; Fujimura, Y. Density functional theoretical study on enantiomerization of 2,2'-Biphenol. J. Phys. Chem. A 2006, 110 (7), 2440-2447.
- (53) van Leeuwen, T.; Danowski, W.; Pizzolato, S. F.; Štacko, P.;
 Wezenberg, S. J.; Feringa, B. L. Braking of a light-driven molecular rotary motor by chemical stimuli. *Chem. Eur. J.*

2018, 24, 81-84.

- (54) Cabrera, E. V.; Sanchez, J. L.; Banerjee, A. K. A practical synthesis of 7,8-dimethoxy-2-tetralone. Org. Prep. Proced. Int. 2011, 43 (4), 364-367.
- (55) van Leeuwen, T.; Gan, J.; Kistemaker, J. C. M.; Pizzolato, S. F.; Chang, M.-C.; Feringa, B. L. Enantiopure functional molecular motors obtained by a switchable chiral-resolution process. *Chem. Eur. J.* 2016, 22 (21), 7054-7058.
- (56) Sheldrick, G. M. *Acta Crystallogr. Sect. A, Found. Adv.* 2015, 71 (Pt 1), 3–8.
- (57) Sheldrick, G. M. Acta Crystallogr. A. 2008, 64 (Pt 1), 112-122.
- Hooft, R. W. W.; Straver, L. H.; Spek, A. L. Using the *t*-distribution to improve the absolute structure assignment with likelihood calculations. *J. Appl. Crystallogr.* 2010, *43* (4), 665-668.
- Lunazzi, L.; Mancinelli, M.; Mazzanti, A. Correct values of the rotation barriers of 1,8-ditolylanthracenes. J. Org. Chem. 2007, 72 (26), 10045-10050.
- (60) Mazzanti, A.; Lunazzi, L.; Minzoni, M.; Anderson, J. E. Rotation in biphenyls with a single ortho-substituent. J. Org. Chem. 2006, 71 (15), 5474-5481.
- (61) Casarini, D.; Lunazzi, L.; Mancinelli, M.; Mazzanti, A.; Rosini, C. Structure, conformation, stereodynamics, dimer formation, and absolute configuration of axially chiral atropisomers of hindered biphenyl carbinols. *J. Org. Chem.* 2007, 72 (20), 7667–7676.
- (62) Claridge, T. D. W. High-Resolution NMR Techniques in Organic Chemistry; Newnes, 2008; Vol. 19.
- (63) It should be emphasized that compared with compound 1, the methoxy-protected precursor 9 could be obtained and isolated as variable mixtures of non-equilibrating atropisomers (see Supplementary material for data). Notably no isomerization or coalescence of sets of absorption was detected by 'H NMR spectroscopy even after prolonged heating of a sample of 9 (toluene solution, 110 °C, 12 h), confirming the important role of the hydroxyl groups and internal hydrogen-bonding in the atropisomerization process.
- (64) Ceccacci, F.; Mancini, G.; Mencarelli, P.; Villani, C. Determination of the rotational barrier of a chiral biphenyl: Comparison of theoretical and experimental data. *Tetrahedron: Asymmetry* **2003**, *14* (20), 3117-3122.
- (65) Trapp, O.; Trapp, G.; Schurig, V. Direct calculation and computer simulation of the enantiomerization barrier of oxazepam in dynamic HPLC experiments—a comparative study. J. Biochem. Biophys. Methods 2002, 54 (1), 301–313.
- (66) Cabrera, K.; Jung, M.; Fluck, M.; Schurig, V. Determination of enantiomerization barriers by computer simulation of experimental elution profiles obtained by high-performance liquid chromatography on a chiral stationary phase. *J. Chromatogr. A* **1996**, 731 (1 2), 315-321.
- (67) Andreani, R.; Bombelli, C.; Borocci, S.; Lah, J.; Mancini, G.; Mencarelli, P.; Vesnaver, G.; Villani, C. New biphenylic derivatives: synthesis, characterisation and enantiodiscrimination in chiral aggregates. *Tetrahedron: Asymmetry* 2004, 15 (6), 987-994.
- (68) Degradation of compound 1 was observed in presence of strong bases and radical species, which led to addition of lower phenolic hydroxyl group to overcrowded alkene motif and loss of switching properties.
- (69) Yoshikawa, N.; Kumagai, N.; Matsunaga, S.; Moll, G.; Ohshima, T.; Suzuki, T.; Shibasaki, M. Direct catalytic asymmetric aldol reaction: synthesis of either *syn-* or *anti-* α,β -dihydroxy ketones. *J. Am. Chem. Soc.* **2001**, *123* (10), 2466–2467.
- Kumagai, N.; Matsunaga, S.; Yoshikawa, N.; Ohshima, T.; Shibasaki, M. Direct catalytic enantio- and diastereoselective aldol reaction using a Zn–Zn-Linked-BINOL complex: A practical synthesis of syn-1,2-Diols. Org. Lett. 2001, 3 (10), 1539–1542.

56

57

58 59

- (71) Kumagai, N.; Matsunaga, S.; Kinoshita, T.; Harada, S.; Okada, S.; Sakamoto, S.; Yamaguchi, K.; Shibasaki, M. Direct catalytic asymmetric aldol reaction of hydroxyketones: asymmetric Zn catalysis with a Et₂Zn/linked-BINOL complex. *J. Am. Chem. Soc.* 2003, 125 (8), 2169–2178.
- (72) Du, H.; Long, J.; Hu, J.; Li, X.; Ding, K. 3,3'-Br2-BINOL-Zn complex: a highly efficient catalyst for the enantioselective hetero-Diels–Alder reaction. *Org. Lett.* 2002, *4* (24), 4349–4352.
- (73) Noyori, R. *Asymmetric catalysis in organic synthesis*; Wiley: Chichester, 1994.
- (74) Pu, L.; Yu, H.-B. Catalytic asymmetric organozinc additions to carbonyl compounds. *Chem. Rev.* **2001**, *101* (3), 757–824.
- (75) Soai, K.; Niwa, S. Enantioselective addition of organozinc reagents to aldehydes. *Chem. Rev.* **1992**, *92* (5), 833–856.
- (76) Noyori, R.; Kitamura, M. Enantioselective addition of organometallic reagents to carbonyl compounds: chirality transfer, multiplication, and amplification. *Angew. Chem., Int. Ed.* **1991**, 30 (1), 49–69.
 - (77) Hirose, T.; Kodama, K. In *Comprehensive Organic Synthesis II*; 2014; pp. 204–266.
- (78) Schmidt, B.; Seebach, D. Catalytic and stoichiometric enantioselective addition of diethylzinc to aldehydes using a novel chiral spirotitanate. *Angew. Chem., Int. Ed.* 1991, 30 (1), 99–101.
- Song, T.; Zheng, L.-S.; Ye, F.; Deng, W.-H.; Wei, Y.-L.; Jiang, K.-Z.; Xu, L.-W. Modular synthesis of Ar-BINMOL-Phos for catalytic asymmetric alkynylation of aromatic aldehydes with unexpected reversal of enantioselectivity. *Adv. Synth. Catal.* 2014, 356 (8), 1708–1718.
- (80) Dean, M. A.; Hitchcock, S. R. Divergent enantioselective pathways in the catalytic asymmetric addition of diethylzinc to aldehydes in the presence and absence of titanium tetraisopropoxide. *Tetrahedron: Asymmetry* 2008, 19 (22), 2563–2567.
- (81) Shibata, T.; Tarumi, H.; Kawasaki, T.; Soai, K. Reversal in enantioselectivity induced by achiral alcohols in asymmetric autocatalysis in the presence of a chiral diol as

a chiral initiator. *Tetrahedron Asymmetry* **2012**, 23 (14), 1023–1027.

- (82) Hirose, T.; Sugawara, K.; Kodama, K. Switching of enantioselectivity in the catalytic addition of diethylzinc to aldehydes by regioisomeric chiral 1, 3-amino sulfonamide ligands. J. Org. Chem. 2011, 76 (13), 5413–5428.
- (83) Yue, H.; Huang, H.; Bian, G.; Zong, H.; Li, F.; Song, L. Enantioselectivity switch controlled by N,N'-di- or N,N,N',N'-tetra-substituted chiral thiophosphorodiamide ligands, structural relatives of thioureas, in catalytic additions of diethylzinc to aldehydes. *Tetrahedron: Asymmetry* 2014, 25 (2), 170–180.
- (84) Mao, J.; Wan, B.; Zhang, Z.; Wang, R.; Wu, F.; Lu, S. Reversal of enantioselectivity by adding Ti (OiPr) 4: novel sulfamide-amine alcohol ligands for the catalytic asymmetric addition of diethylzinc to aldehydes. J. Mol. Catal. A Chem. 2005, 225 (1), 33-37.
- (85) Costa, A. M.; Jimeno, C.; Gavenonis, J.; Carroll, P. J.; Walsh,
 P. J. Optimization of catalyst enantioselectivity and activity using achiral and meso ligands. J. Am. Chem. Soc. 2002, 124 (24), 6929-6941.
- (86) Erdik, E. In Organozinc Reagents in Organic Synthesis; CRC Press: Boca Raton, FL, 1996; pp. 215–216.
- (87) Weber, B.; Seebach, D. Ti-TADDOLate-catalyzed, highly enantioselective addition of alkyl-and aryl-titanum derivatives to aldehydes. *Tetrahedron* **1994**, 50 (25), 7473– 7484.
- (88) The same reaction catalyzed by (*P*)-3,3'-diphenyl BINOL was reported to yield the product in 44% ee (see Ref. 82).
- (89) It should be noted that the quality of the commercial diethylzinc solution had a large impact on the results of the catalytic reactions, affording lower yields and selectivities as degradation of the reagent progressed.
- (90) Zong, H.; Huang, H.; Bian, G.; Song, L. Synergistic effect of achiral quaternary ammonium salt on asymmetric additions of diethylzinc to aldehydes. *Tetrahedron Lett.* 2013, 54 (21), 2722–2725.

59

Insert Table of Contents artwork here

