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# Confining Nitrogen Inversion to Yield Enantiopure Quinolino[3,2,1-k]Phenothiazine Derivatives

Cassandre Quinton, Lambert Sicard, Nicolas Vanthuyne, Olivier Jeannin, and Cyril Poriel\*

The first examples of optically pure 9H-quinolino[3,2,1-k]phenothiazine (QPTZ)-based molecules are reported. The inversion of the nitrogen atom, locked in the QPTZ fragment, is confined with a high-energy barrier that has allowed the isolation at room temperature of each stereoisomer. Considering the growing necessity in molecular electronics to introduce chiral characteristics within highly efficient molecular fragments, a molecular strategy to generate enantiopure derivatives constructed on the very promising electron rich core QPTZ is provided. As this work aims to report the foundations of the QPTZ chirality, the present findings may open avenues towards the use of this fragment in its optically pure form.

## 1. Introduction

The incorporation of chirality in highly efficient molecular fragments is one of the chief future directions for all three types of organic electronics devices,<sup>[1]</sup> namely organic light-emitting diodes (OLEDs),<sup>[2]</sup> field effect transistors,<sup>[3]</sup> and solar cells.<sup>[4]</sup> The incorporation of a chiral compound in device can enable the detection and emission of chiral light or a enhance charge transport.<sup>[1]</sup> However and despite recent promising studies, only a few examples have been reported to date. This is partly due to the limited number of chiral fragments displaying a high efficiency when incorporated in devices. Two years ago, a new electron rich core, 9H-quinolino[3,2,1-k]phenothiazine (QPTZ), has appeared in literature displaying its potential in the field of phosphorescent OLEDs.<sup>[5]</sup> This fragment consists of a phenylacridine core<sup>[6]</sup> to which a sulfur bridge was added between the top and side phenyl rings. In this previous work, the crucial role played by this sulfur atom in the electronic properties (i.e., raising of the highest occupied

molecular orbital level and of the spin-orbit coupling) has been demonstrated.<sup>[5]</sup> The QPTZ fragment incorporated in spiroconfigured materials has thus led to highly efficient blue phosphorescent OLEDs, displaying better performance than those obtained with the widely used phenylacridine.<sup>[5]</sup> In the present work, we wish to focus on the dynamic behavior of the nitrogen atom in QPTZ-based molecules. Indeed, *N*-pyramidal amines with three different substituents attached to the nitrogen atom can be optically active if there is a thermodynamically predominant conformer in the conformational

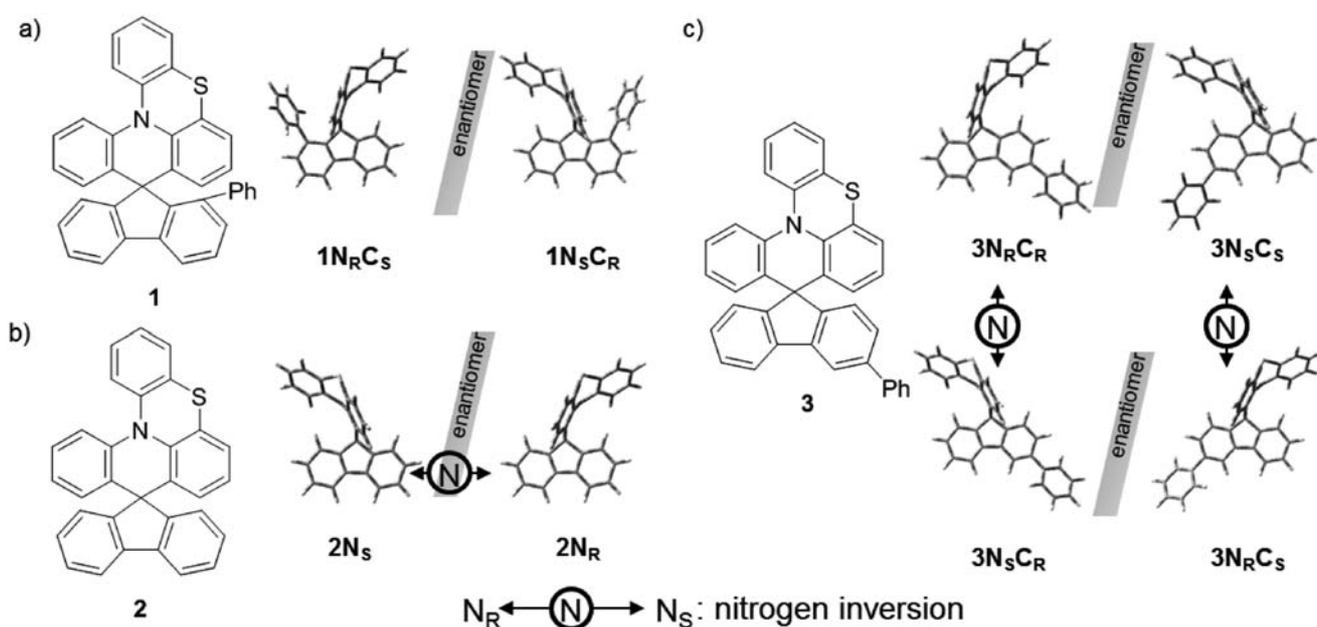
equilibrium, but the resulting conformers often cannot be isolated as the nitrogen inversion barrier is too low.<sup>[7]</sup> In fact, for configurational stability to be observed, the nitrogen inversion energy barrier should be higher than 100 kJ mol<sup>-1</sup> at room temperature.<sup>[8]</sup> This value can be reached by inserting the nitrogen atom into a ring, adding strain in the planar transition state.<sup>[9]</sup> Interestingly, in the rigid QPTZ fragment investigated herein, the confinement of the nitrogen atom within a dissymmetric bridged structure increases the energy of this inversion barrier which therefore enables the isolation of the resulting isomers. Moreover, the sulfur atom seems to contribute to the structural and chiral characteristics described herein. Indeed, the sulfur/carbon linkage rigidifies the system compared to the phenylacridine fragment but retains enough flexibility, unlike indoloacridine,<sup>[10]</sup> to allow the inversion of the nitrogen atom when heated. Note that a structurally related analogue possessing an oxygen atom instead of the sulfur atom has been reported with nevertheless no mention of nitrogen inversion.<sup>[11]</sup> Thus, this work presents our investigations on the nitrogen inversion of the QPTZ fragment and the first examples of optically pure QPTZ-based molecules (**1–3**, **Figure 1**) described in literature. Constraining the nitrogen atom in a pyramidal configuration at room temperature appears hence as an efficient strategy to generate enantiopure compounds. Furthermore, the spiro carbon of spirofluorene-based compounds also allows the introduction of chirality in such systems.<sup>[12]</sup>

Three molecular models based on the fluorene-*spiro*-QPTZ (F-*s*-QPTZ) architecture have been chosen for this study: the unsubstituted F-*s*-QPTZ (**2**) and two substituted F-*s*-QPTZ possessing different substitution patterns (**1** and **3**) in order to investigate the impact of the steric hindrance on the nitrogen inversion. Considering electronic device applications for the QPTZ core, it is important to know whether the inversion of

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**Figure 1.** Molecular structures of the investigated molecules. a) **1**, b) **2**, and c) **3** and their relationships obtained by molecular modelling (optimized structures with b3lyp/6-31g(d)). The view angle chosen to highlight the phenothiazine flip shows the two phenyl rings of the acridine moiety and the two sigma bonds of the spiro bridge as superimposed (different views of **2N<sub>R</sub>** can be found Figure S48 in the Supporting Information). Sulfur atom in yellow and nitrogen atom in blue.

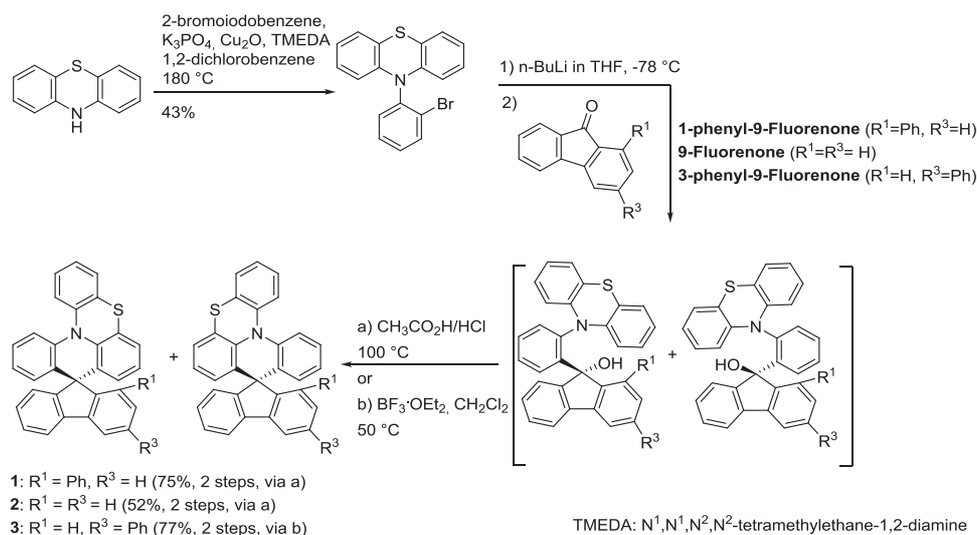
the nitrogen atom can occur or not and, if it can, how quickly and at which temperature.

## 2. Results and Discussion

Compounds **1–3** were readily obtained through a short and efficient synthetic approach by coupling the halogeno-aryl 10-(2-bromophenyl)-10H-phenothiazine (obtained from the copper catalysed Goldberg coupling of phenothiazine and 2-bromiodobenzene), with the corresponding 9-fluorenone:

1-phenyl-9-fluorenone for **1**, 9-fluorenone for **2**, 3-phenyl-9-fluorenone for **3** (Scheme 1). The corresponding fluorenones were not isolated and were involved in an intramolecular electrophilic cyclisation reaction to give the three targets **1–3** in good yields.

After synthesis, the <sup>1</sup>H and <sup>13</sup>C NMR spectra of the three compounds **1–3** provide the first piece of interesting data (see Figures S49–S68 in the Supporting Information). Indeed, **3** displays a complex spectrum, characteristic of a mixture of diastereoisomers, very different to that of the unsubstituted **2** and that of the 1-substituted **1**.



**Scheme 1.** Syntheses of compounds **1–3**.

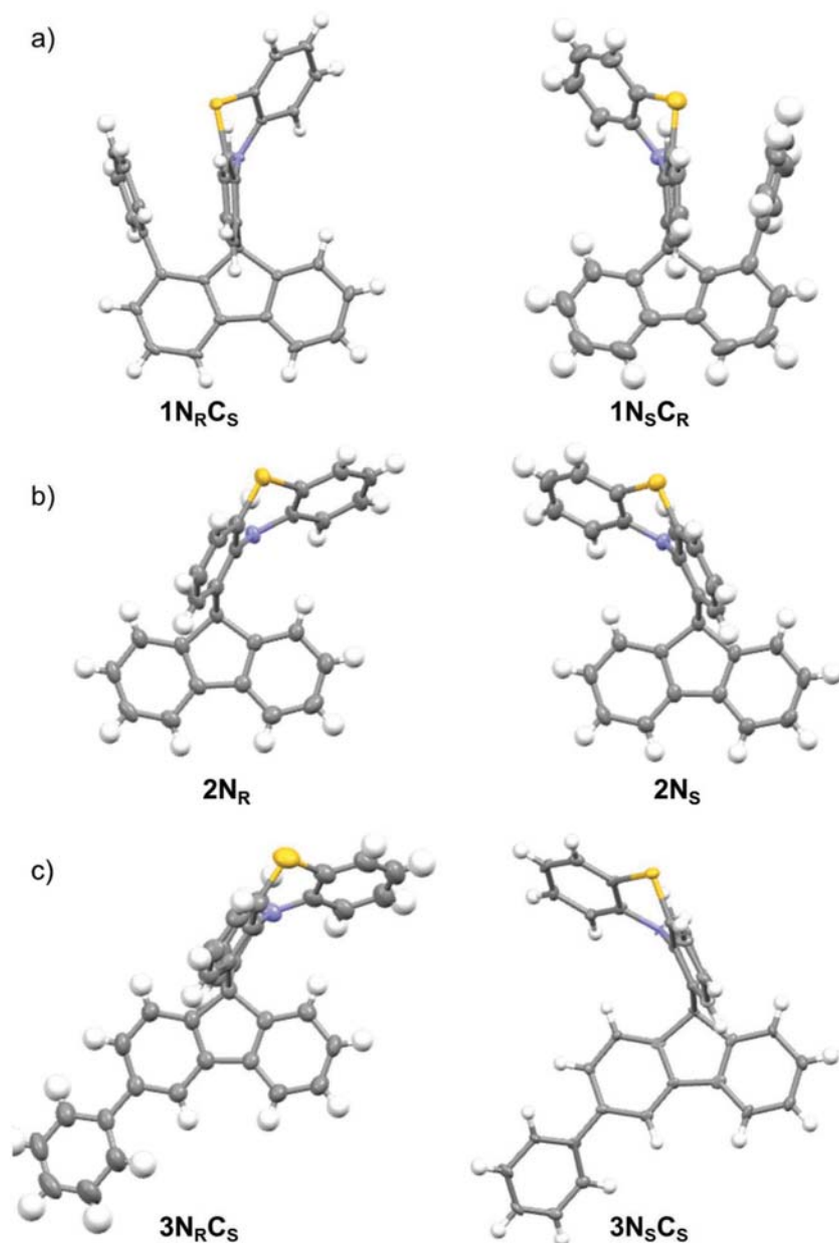
The presence of diastereoisomers in **3** suggests at least two stereogenic centers, identified in our case as the nitrogen atom and the spiro carbon (the fluorene being dissymmetrical). These stereochemical features have been confirmed by chiral high pressure liquid chromatography (HPLC) studies (see Figures S2, S4, and S6 in the Supporting Information), which have allowed to separate and isolate each diastereoisomer of **3**:  $3N_R C_R$ ,  $3N_S C_S$ ,  $3N_R C_S$ , and  $3N_S C_R$  (Figure 1c).

X-Ray crystallography of the two diastereoisomers  $3N_S C_S$  and  $3N_R C_S$  (Figure 2c) clearly confirms the inversion of the nitrogen atom in the QPTZ fragment, which is caused by a flip of the

phenothiazine core. Indeed, the QPTZ moiety is not flat due to the intense deformations induced by the phenothiazine and acridine cores (Figure 2). In fact, the nitrogen atom is pyramidal with values of nitrogen torsion angle found between  $157^\circ$  for  $3N_R C_S$  and  $3N_S C_R$  and  $162^\circ$  for  $3N_S C_S$  (see Figure S17 and Table S3 in the Supporting Information), as expected for nitrogen with one or more aromatic substituents.<sup>[13]</sup> This shows that the trivalent nitrogen in these compounds is in an intermediate hybridization state between  $sp^2$  and  $sp^3$ . Thus, the phenothiazine can move from one side of the fluorene to the other through a structural reorganization of the acridine, much like a wing flapping.

The same kind of stereoisomers would have been expected for **1** which displays the same two stereogenic centres. However, only two stereoisomers (enantiomers) have been isolated by chiral HPLC:  $1N_R C_S$  and  $1N_S C_R$  (Figures 1a and 2a). Indeed, the steric hindrance induced by the pendant phenyl ring in position 1 of the fluorene prevents the flipping of the phenothiazine above discussed and hence the formation of the other pair of enantiomers. This structural feature provides an efficient diastereocontrol and can be easily visualized on the X-ray structures of the two enantiomers  $1N_R C_S$  and  $1N_S C_R$  (Figure 2a). The phenyl ring in C1 displays two important characteristics. First, it is almost orthogonal to the fluorene with a dihedral angle between the mean plane of the phenyl ring in C1 and that of the fluorene of  $83.3^\circ$  for  $1N_R C_S$  and of  $84.1^\circ$  for  $1N_S C_R$  suggesting a strongly sterically hindered environment (see Figures S13 and S14 in the Supporting Information). This structural characteristic is at the origin of the emergence of one-substituted fluorenes recently reported in literature.<sup>[14]</sup> In addition, the pendant phenyl ring is very close to the QPTZ fragment (three C/C intramolecular distances are indeed shorter than the sum of the Van der Waals radii, see Figures S15 and S16 in the Supporting Information).

Finally, the case of **2** provides a different example. For this molecule, in which the unsubstituted fluorene is symmetrical, the only stereogenic centre is the nitrogen atom. This feature, clearly observed in the X-ray structure of the racemic mixture of **2** leads to two enantiomers  $2N_S$  and  $2N_R$  (Figure 2b), which have been successfully isolated by chiral HPLC (see Figure S2 in the Supporting Information). The assignment of the absolute stereochemistry was performed for each enantiomer of **1** and **3** by crystallography (Figure 2a,c and see Figures S7–S10,S12 and Tables S1–S2 in the Supporting Information) and confirmed by the comparison of experimental and theoretical circular dichroism (CD) spectra (see

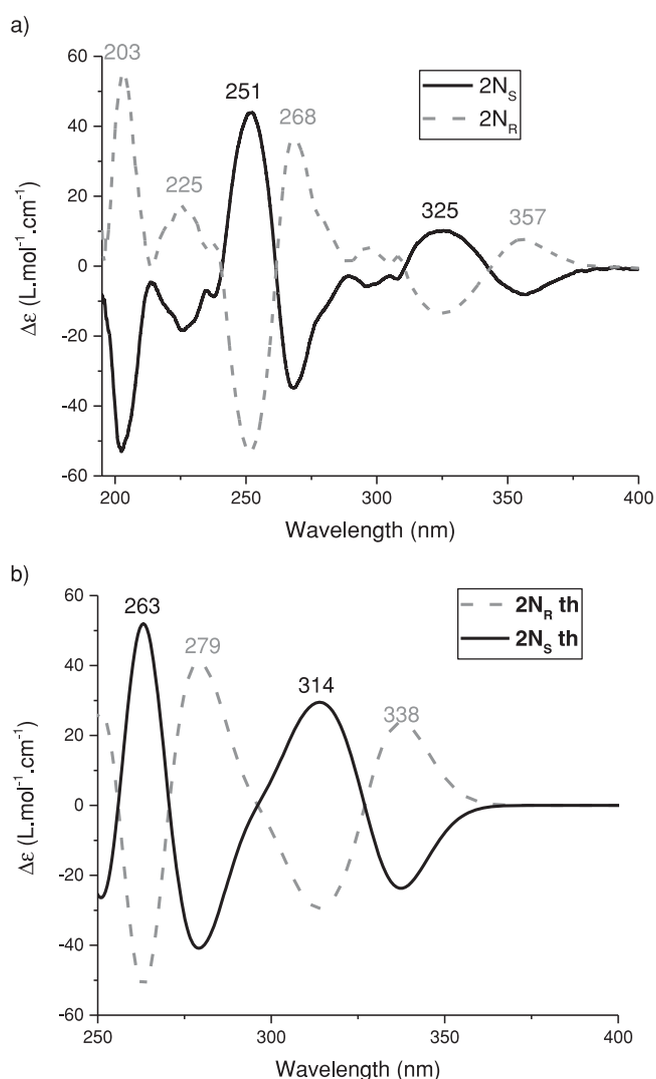


**Figure 2.** X-ray structures of a)  $1N_R C_S$  and  $1N_S C_R$ , b)  $2N_R$  and  $2N_S$ , and c)  $3N_R C_S$  and  $3N_S C_S$ . The view angle chosen to highlight the phenothiazine flip shows the two phenyl rings of the acridine moiety and the two sigma bonds of the spiro bridge as superimposed. Sulfur atom in yellow and nitrogen atom in blue.

Figures S31–S33 in the Supporting Information). In the case of  $2N_S$  and  $2N_R$ , the stereochemistry has been assigned, thanks to their CD spectra (Figure 3).

These spectra are mirror images of each other and in good coherence between experimental and theoretical studies. The CD spectrum of  $2N_S$  ( $2N_R$ ) displays i) a small negative (positive) band at 357 nm, ii) a small positive (negative) band at 325 nm, iii) an intense negative (positive) band at 268 nm, iv) an intense positive (negative) band at 251 nm, v) a small negative (positive) band at 225 nm, and vi) an intense negative (positive) band at 203 nm (Figure 3).

To sum up, compound **2** displays only the nitrogen atom as stereogenic center (and thus two enantiomers), **3** displays both the spiro carbon and the nitrogen atom as stereogenic centers (leading to two pairs of enantiomers and thus four diastereoisomers) and finally **1** also displays both stereogenic centers but leading to only one pair of enantiomers due to the steric



**Figure 3.** a) Experimental ( $\text{CH}_3\text{OH}$  at 298 K) and b) theoretical (time-dependent density functional theory (TD-DFT), PBE0/6-311+g(d,p)) circular dichroism spectra of enantiomers  $2N_S$  and  $2N_R$ .

hindrance induced by the presence of the phenyl ring in position 1. Finally, it should be stressed that **1–3** present the same electronic properties.

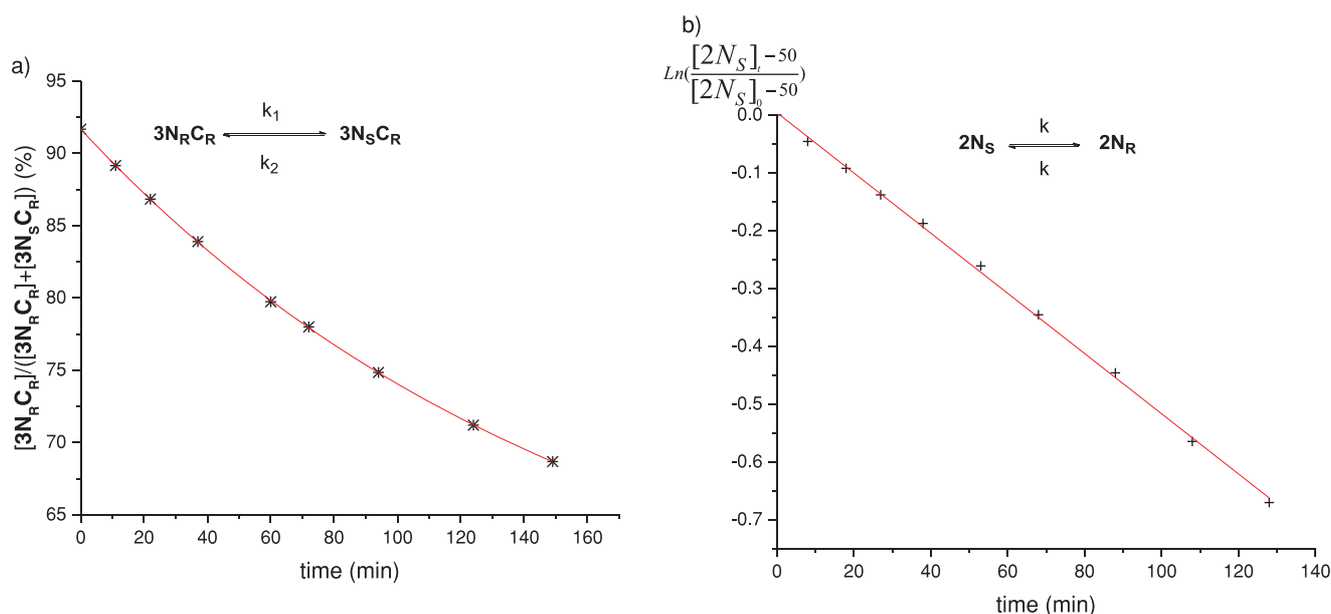
The key feature of this work is related to the nitrogen inversion energy barrier in the constrained structure of the QPTZ fragment, which is high enough to enable the isolation at room temperature of the different stereoisomers of the presented compounds. In order to evaluate this barrier, the mechanism of the nitrogen inversion has been studied by a combination of experimental (NMR and HPLC) and theoretical techniques. Thus, the kinetic investigations of the nitrogen inversion of  $3N_R C_R$  into its diastereoisomer  $3N_S C_R$  have been first performed by measuring through chiral HPLC the evolution of the concentration of a solution of  $3N_R C_R$  in ethanol over time at 351.15 K (Figure 4a).

This study gives a kinetic constant  $k_1$  of  $0.0029 \text{ min}^{-1}$ , a free energy of activation  $\Delta G^\ddagger$  of  $115.5 \text{ kJ mol}^{-1}$  (calculated from the Eyring equation, see Table S7 in the Supporting Information) and a half-life of inversion  $t_{1/2}$  of 112 min for the N inversion from  $3N_R C_R$  to  $3N_S C_R$  (Table 1). Note that the conformer  $3N_R C_R$  is slightly more stable than its diastereoisomer  $3N_S C_R$ , with an energy difference of  $0.4 \text{ kJ mol}^{-1}$ , value in accordance with that found by calculations ( $0.4 \text{ kJ mol}^{-1}$ , see below).

Kinetic investigations have also been performed through  $^1\text{H}$  NMR studies (Figure 5). Indeed, the acridine protons in  $\beta$  positions of the spiro carbon (Ha/Hb for  $3N_R C_R+3N_S C_S$  and Hc/Hd  $3N_S C_R+3N_R C_S$ ) are sensitive to the nitrogen's configuration (Figure 5) and were used as efficient probes to determine the  $[3N_R C_R+3N_S C_S]/[3N_S C_R+3N_R C_S]$  ratio. These signals are detected as doublets of doublets at 6.57 and 6.38 ppm respectively for  $3N_S C_R$  and  $3N_R C_S$  and 6.52 and 6.33 ppm for  $3N_R C_R$  and  $3N_S C_S$  with no overlap.

Thus, the decrease of the concentration  $[3N_S C_R+3N_R C_S]$  in deuterated *N,N*-dimethylsulfoxide ( $\text{DMSO}-d_6$ ) has been measured for the two protons Hc and Hd at five temperatures between 348 and 373 K (see Figures S22–S26 in the Supporting Information). At 348 K, starting with 97% of  $3N_S C_R+3N_R C_S$ , the amount of their corresponding diastereoisomers  $3N_R C_R+3N_S C_S$  gradually increases (15% after 50 min, 35% after 200 min) to reach 48% after 760 min. Indeed, the final state is composed of  $3N_S C_R+3N_R C_S$  at 48% and  $3N_R C_R+3N_S C_S$  at 52% which corresponds to a stabilization of  $0.3 \text{ kJ mol}^{-1}$  for  $3N_R C_R+3N_S C_S$  compared to  $3N_S C_R+3N_R C_S$ . This provides a kinetic constant  $k$  of  $0.00275 \text{ min}^{-1}$ , a free energy of activation  $\Delta G^\ddagger$  of  $114.6 \text{ kJ mol}^{-1}$  at 348 K and a half-life of inversion  $t_{1/2}$  of 132 min (from  $3N_S C_R+3N_R C_S$  to  $3N_R C_R+3N_S C_S$ ).

The rate of inversion increases with the temperature. The resulting free energies are gathered in Table S8 in the Supporting Information for the five temperatures. The van't Hoff plot of  $\ln K$  versus  $1/T$  gave us the enthalpy of activation of the inversion  $\Delta H^\ddagger = 106 \text{ kJ mol}^{-1}$  and the entropy of activation of the inversion  $\Delta S^\ddagger = -24 \text{ J}\cdot\text{mol}^{-1}\cdot\text{K}^{-1}$  (Figure 6 and see Figure S27 and Table S9 in the Supporting Information). Thus, the free energy of activation  $\Delta G^\ddagger$  is found to be  $113 \text{ kJ mol}^{-1}$  at room temperature (298 K). This value being higher than  $100 \text{ kJ mol}^{-1}$ ,<sup>[8]</sup> it explains why all the isomers can be isolated at room temperature. Furthermore, we calculated a value of  $\Delta G^\ddagger$  of  $114.9 \text{ kJ mol}^{-1}$  from  $3N_R C_R$  at 315.15 K, slightly lower than the one obtained by chiral HPLC studies in ethanol. The



**Figure 4.** Evolution of the concentration of a)  $3N_R C_R$  or b)  $2N_S$  in ethanol through time at 351.15 K.

nitrogen inversion barrier is therefore found to be higher in ethanol than in DMSO, which could be explained by a stabilization of the ground state through hydrogen bonding, even though this solvent effect seems weaker in our case than the ones reported in the literature.<sup>[7a]</sup> Furthermore, thermogravimetric analysis (TGA) investigations show decomposition temperatures, corresponding to a 5% mass loss, recorded between 572 K (for  $3N_S C_R + 3N_R C_S$ ) and 587 K (for  $3N_R C_S$ ) depending on the sample constitution (Figure S38, Supporting Information). Differential scanning calorimetry (DSC) does not allow us to determine the melting and the glass transition temperatures for each enantiopur form, because the nitrogen inversion begins at lower temperature than these characteristic temperatures (Figures S44–S47, Supporting Information).

Compound **1** displays a drastically different behavior. Indeed, the heating of **1** to 443 K leaves the  $^1\text{H}$  NMR spectrum unchanged, clearly confirming that even with the help of thermal input the nitrogen inversion is impossible due to the steric hindrance induced by the phenyl ring in C1. Furthermore, TGA investigations of **1** and  $1N_R C_S$  show decomposition temperatures of 548 K for the racemic mixture and

579 K for the enantiopur  $1N_R C_S$  (see Figure S37 in the Supporting Information). Thus, the packing mode seems to have an effect on the decomposition temperatures. Moreover, as the mass loss appears total around 633 K for both samples, we believe that the sublimation occurs at this temperature without a real decomposition of the compounds. DSC measurements were performed for both **1** and  $1N_R C_S$  between 253 and 513 K. At the first heating cycle (see Figures S40 and S41 in the Supporting Information), the two compounds present a sharp endothermic peak at 522 and 491 K respectively, associated with the melting of the two compounds ( $T_m$ : 521 and 488 K respectively, determined from the peak onset). When both liquids were cooled down at the same rate from 513 to 253 K, no recrystallization was detected; hence, the cooling leads to amorphous solids, which is highly beneficial for further device incorporation. At the second heating cycle, a glass transition phenomenon was observed at 378 K for **1** and at 377 K for  $1N_R C_S$  ( $T_g$ : 376 and 374 K, respectively determined from the peak onset), which is comfortable high enough for future device application. Finally, the heating of **1** or its enantiopur counterparts does not induce any degradation or nitrogen inversion.

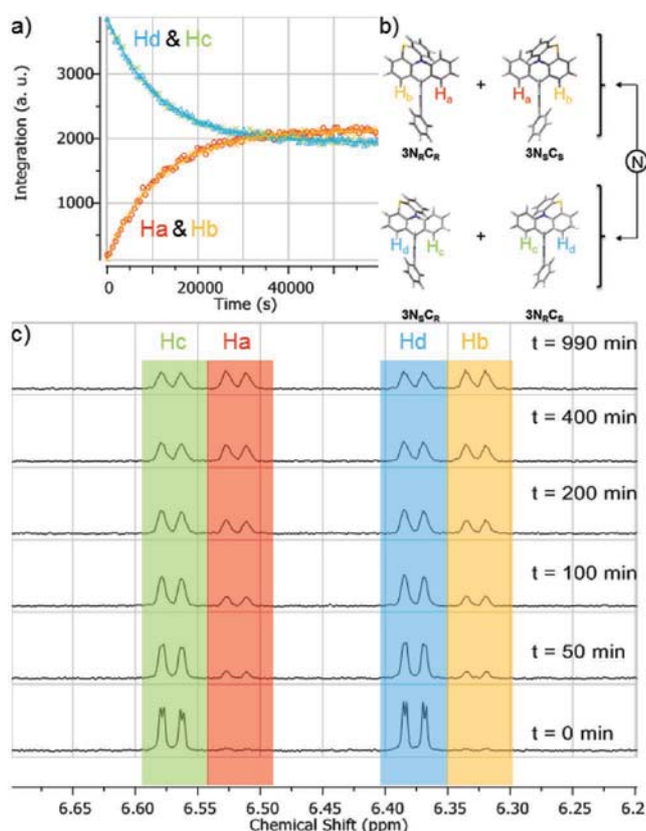
It is important to mention that kinetic investigations through  $^1\text{H}$  NMR study cannot be performed in the case of **2** since the inversion takes place between two enantiomers that are hence indistinguishable from each other through NMR.

The kinetic investigations of the nitrogen inversion of  $2N_R$  into its enantiomer  $2N_S$  have therefore been performed by chiral HPLC in similar conditions than those above mentioned (in ethanol at 351.15 K, Figure 4b). The inversion between the two enantiomers (from  $2N_R$  to  $2N_S$ ) provides very similar values ( $k = 0.0026 \text{ min}^{-1}$ ,  $\Delta G^\ddagger = 115.9 \text{ kJ mol}^{-1}$  and a half-life of inversion of 133 min) than those exposed above for compound **3**. Indeed, the phenyl ring in position 3 is too far from the nitrogen atom to have an influence on the kinetics of its inversion. Furthermore,

**Table 1.** Experimental thermodynamical data for the N inversion at 351.15 K.

Compounds	$\Delta G^\ddagger$ [kJ mol <sup>-1</sup> ]	$k$ [min <sup>-1</sup> ]	$t_{1/2}$ [min]	Solvent
$2N_R$	115.9 <sup>a)</sup>	0.00260	133	Ethanol
$3N_R C_R$	115.5 <sup>a)</sup>	0.00288	112	Ethanol
	114.9 <sup>b)</sup>	0.00356	97	DMSO- <i>d</i> <sub>6</sub>
$3N_S C_R$	115.1 <sup>a)</sup>	0.00331	112	Ethanol
	114.6 <sup>b)</sup>	0.00392	88	DMSO- <i>d</i> <sub>6</sub>

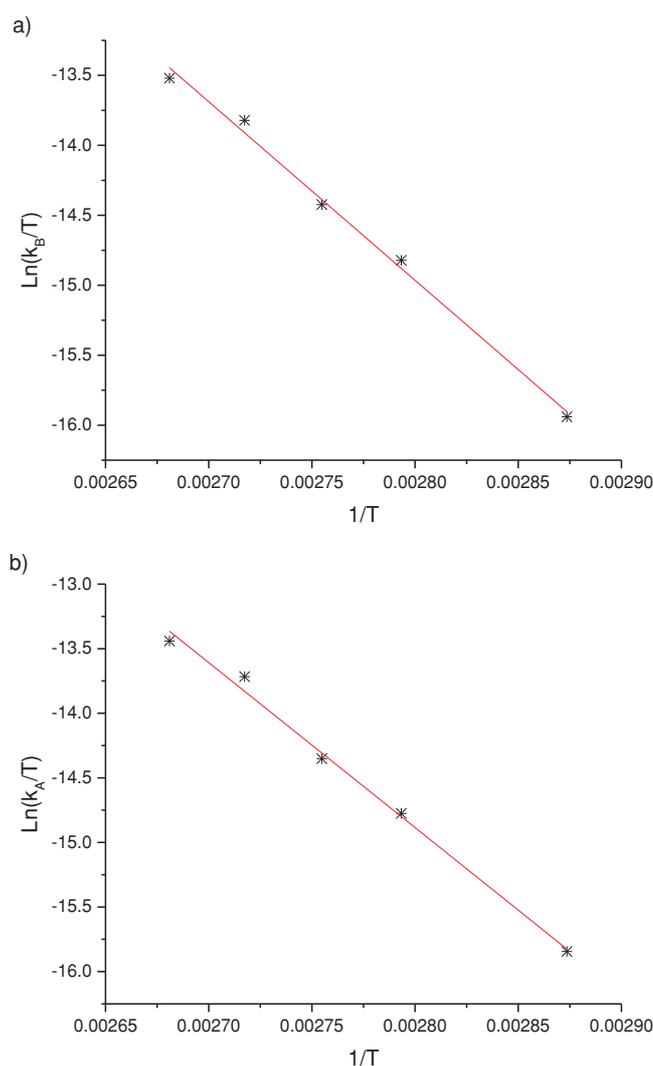
<sup>a)</sup>HPLC measurements (in ethanol,  $T = 351.15 \text{ K}$ ); <sup>b)</sup>Calculated  $\Delta G^\ddagger$  at 351.15 K from the values of  $\Delta H^\ddagger$  and  $\Delta S^\ddagger$  obtained by NMR measurements in DMSO-*d*<sub>6</sub>.



**Figure 5.** a) Evolution of the integration of Ha, Hb, Hc, and Hd through time. b) Nomenclature of the investigated protons Ha, Hb, Hc, and Hd, c) Evolution of the  $^1\text{H}$  NMR spectra of  $3\text{N}_5\text{C}_\text{R}+3\text{N}_\text{R}\text{C}_5$  in  $\text{DMSO-}d_6$  at 348 K.

TGA investigations of **2** and  $2\text{N}_\text{R}$  show decomposition temperatures recorded at 548 K for the racemic mixture and 500 K for the enantiopur  $2\text{N}_\text{R}$  (see Figure S39 in the Supporting Information), again showing the effect of the packing mode on the stability. One can also note that these decomposition temperatures are lower than those exposed above for compounds **1** and **3**, showing the influence of the pendant phenyl ring in raising the thermal stability. As for compound **3**, the melting and glass transition temperatures could not be determined by DSC for each enantiopur form because of the nitrogen inversion (Figures S42 and S43, Supporting Information).

Finally, the mechanism of the nitrogen inversion has been studied by theoretical calculations. Thus, the molecular structures of the two enantiomers of **2** in their ground state and planar-shaped transition state, corresponding to the nitrogen inversion process, were modelled (Figure 7a). Nitrogen torsion angles were calculated at  $165^\circ$  for  $2\text{N}_\text{R}$ ,  $180^\circ$  for the transition state and  $195^\circ$  for  $2\text{N}_\text{S}$ . The nitrogen inversion leaves the fluorene unchanged, whereas the curvature of the acridine part is reversed and the phenyl of the phenothiazine unit moves from below to above the mean plane of the acridine. Thus, the theoretical-free energy change along the nitrogen inversion of **2** was evaluated at  $121.0 \text{ kJ mol}^{-1}$  (see Figure S20 in the Supporting Information for the plot of the theoretical-free energy change against the nitrogen torsion angle). This feature being

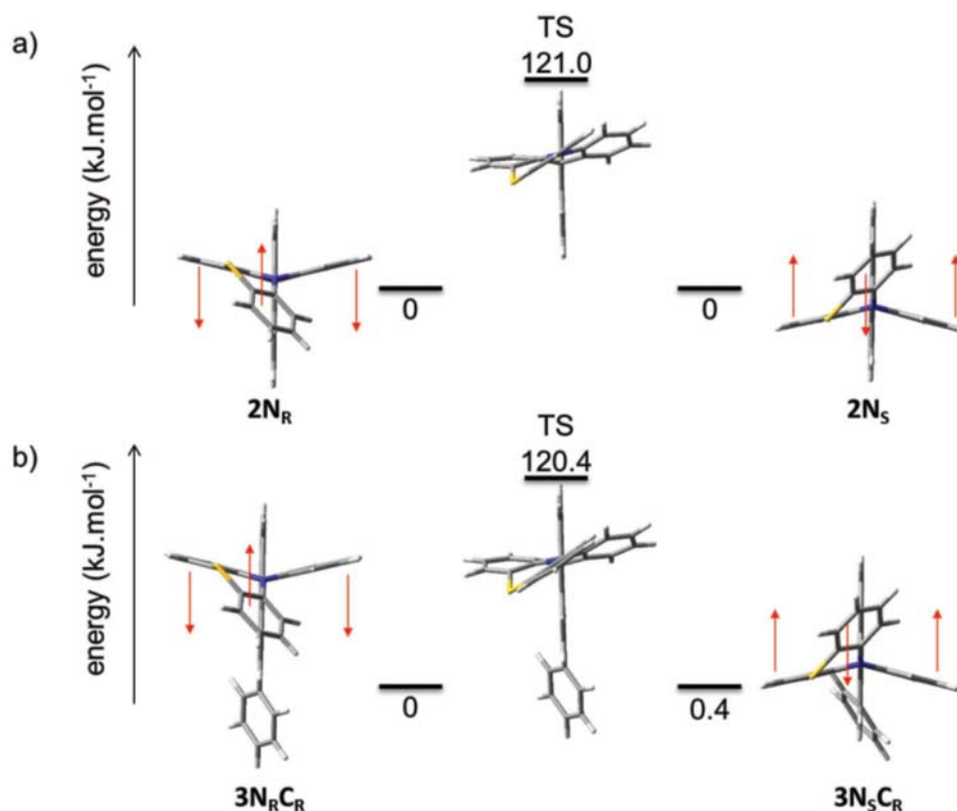


**Figure 6.** Evolution of  $\text{Ln}(k_\text{B}/T)$  in function of  $1/T$  a) for  $3\text{N}_5\text{C}_\text{R}+3\text{N}_\text{R}\text{C}_5$  and b) for  $3\text{N}_\text{R}\text{C}_\text{R}+3\text{N}_5\text{C}_5$ .

at the origin of the chirality of QPTZ, the transition state corresponding to the conversion of  $3\text{N}_\text{R}\text{C}_\text{R}$  (nitrogen torsion angle of  $194^\circ$ ) into  $3\text{N}_5\text{C}_\text{R}$  (nitrogen torsion angle of  $166^\circ$ ) has also been modelled according to the same procedure (Figure 7b and see Figure S28 and S29 in the Supporting Information). The nitrogen inversion barrier is thus evaluated at  $120.4 \text{ kJ mol}^{-1}$  from  $3\text{N}_\text{R}\text{C}_\text{R}$  and  $120.0 \text{ kJ mol}^{-1}$  from  $3\text{N}_5\text{C}_\text{R}$  ( $3\text{N}_\text{R}\text{C}_\text{R}$  being stabilized by  $0.4 \text{ kJ mol}^{-1}$  compared to  $3\text{N}_5\text{C}_\text{R}$  as experimentally observed). Therefore, experimental and theoretical data are in good agreement and provide similar values for both **2** and **3**, highlighting that the presence of the phenyl ring on the fluorene backbone in position 3 has no influence on the nitrogen inversion but induces the formations of the diastereoisomers.

### 3. Conclusion

To conclude, this work presents the first examples of optically pure QPTZ-based molecules. In these systems, the flipping



**Figure 7.** Calculated structures of a)  $2N_R$  and  $2N_S$  and b)  $3N_R C_R$  and  $3N_S C_R$  and their transition states of N inversion in ethanol and the variation of free energy at 351.15 K under 1 atm (B3LYP/6-31g(d,p)). TS = transition state.

of the phenothiazine unit leads to a nitrogen inversion with a high barrier energy ( $\approx 115 \text{ kJ mol}^{-1}$  at 351.15 K), which allows to successfully isolate each isomer at room temperature. Constraining the nitrogen atom in a pyramidal configuration appears hence as an efficient strategy to generate enantiopure compounds. In addition, the introduction of a phenyl ring in position 1 of the fluorene completely prevents the flipping of the phenothiazine and hence the formation of the other pair of enantiomers providing an efficient diastereocontrol. As the efficiency of the QPTZ fragment has recently been highlighted in phosphorescent OLEDs (and is therefore an interesting candidate for other organic electronics applications),<sup>[5]</sup> we believe that the present work may open new avenues in the use of this electron-rich fragment in its chiral form.

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