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# Stereochemical and electronic interaction studies of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones

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#### Abstract

Infrared carbonyl band analysis, supported by B3LYP/6-31+G(d,p) and single-point PCM calculations, natural bond orbital (NBO) analysis and X-ray diffraction were carried out for the diastereoisomers of a selection of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones bearing the substituents NO<sub>2</sub> 1a, Br 2a, H 3a, Me 4a and OMe 5a for the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> enantiomeric pair and Br 2b and Me 4b for the C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub> pair. For the gas phase, the theoretical data indicated the existence of three stable conformations for the C<sub>R</sub>S<sub>R</sub> series and only two for the C<sub>R</sub>S<sub>S</sub> series. For the whole C<sub>R</sub>S<sub>R</sub> series 1a-5a, the most stable  $c_1$  and the intermediate  $c_2$  conformers have similar  $v_{CO}$ frequencies, lowered by about 20 cm<sup>-1</sup> with respect to the less stable  $c_3$ . Likewise, the less stable  $c_2$  conformers for the  $C_RS_S$  series **2b** and **4b** exhibit the higher  $v_{CO}$ frequencies. The single-point PCM calculations show that the relative abundance of both the less stable  $c_3$  C<sub>R</sub>S<sub>R</sub> and  $c_2$  C<sub>R</sub>S<sub>S</sub> conformers, in the gas phase, progressively increases as the dielectric constant of the media increases. The balance between the electrostatic and orbital interactions controls the calculated stability for compounds 1a-**5a**  $(C_RS_R/C_SS_S)$ , along with the  $v_{CO}$  frequency order of the three conformers. Conversely, the larger stabilisation of the  $c_1$  conformer with respect to the  $c_2$  one in the 2b and 4b (C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub>) compounds depends mainly on the orbital interactions  $LP_{(S)} \rightarrow \sigma^*_{\text{C-Se}}, \ LP_{O(S\text{-O})} \rightarrow \sigma^*_{\text{C-H(CH2Me)}} \ \text{ and } \ LP_{O(S\text{-O})} \rightarrow \sigma^*_{\text{C-H(o-PhSe)}}, \ \text{ which are present}$ uniquely in the former conformer. The X-ray single crystal analysis indicates that the major component of 3a ( $C_RS_R$ ) adopts the same geometry as the  $c_2$  conformer in the gas phase. Similarly, the solid-state structure of **2b** matches that of the  $c_2$  conformer. The molecules in the solid are linked in centrosymmetrical pairs through  $\pi...\pi$  interactions between Se-bound phenyl rings, along with phenyl-C-H...O(carbonyl) and phenyl-C-H...O(sulphinyl) interactions.

Keywords: Stereochemical isomerism; infrared spectroscopy; theoretical calculations	Keywords: Stereochem	cal isomerism; infrared	spectroscopy; theoretic	al calculations;
olvent effect; X-ray diffraction; 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl) cetophenones.		diffraction; 4'-substitute	ed 2-(phenylselanyl)-2	-(ethylsulfinyl)-

#### 1. Introduction

Previous conformational analysis performed on some  $\beta$ -carbonyl-sulfoxides XC(O)CH<sub>2</sub>S(O)R using (IR, <sup>13</sup>C NMR, UV and UPS) spectroscopic techniques, X-ray diffraction (XRD) and theoretical calculations have shown that the *gauche* conformer is the most stable for  $\alpha$ -sulfinyl-ketones [1–3],  $\alpha$ -sulfinyl-thioesters [4] and sulfinyl-esters [5], while the *cis* conformer is predominant for the  $\alpha$ -sulfinylacetophenones and  $\alpha$ -sulfinylamides [6–10]. The stabilisation of the *gauche* conformers for these compounds has been ascribed to the  $\pi^*_{CO}/\sigma_{C-S}$  and, to a minor extent, the  $\pi_{CO}/\sigma^*_{C-S}$  orbital interactions, while the larger stabilisation of the *cis* conformers has been attributed to the  $\pi_{Ph}/\pi^*_{CO}$  and  $\pi_N/\pi^*_{CO}$  conjugations, which originate the strong electrostatic [S<sup> $\delta$ +</sup>=O<sup> $\delta$ -</sup>] and  $\pi_{O(CO)}/\sigma^*_{SO}$  charge transfer interactions.

Additionally, an *ab initio* and XRD study of the bis-thio-acetophenone:  $\alpha$ -methylthio- $\alpha$ -methylsulfinyl-acetophenone [6] indicated that in the gas and solid phases, the SMe group adopts the *gauche* geometry with respect to the C=O bond (dihedral angle *ca.* 90–93°), while the SOMe group assumes the *quasi-cis* geometry (dihedral angle 31–35°). The stabilisation of these conformations was ascribed to the occurrence of the hyperconjugative  $\sigma_{C-SR}/\pi^*_{CO}$  interaction that, by increasing the negative charge at the carbonyl oxygen atom favours the  $O^{\delta_{-CO}} \rightarrow S^{\delta_{+}}_{(SO)}$  electrostatic and charge transfer interactions.

Furthermore, the  $v_{CO}$  IR,  $\alpha$ -methylene  $C^{13}$  NMR and  $n \rightarrow \pi^*_{CO}$  UV analyses of  $\alpha$ -phenylseleno p-substituted propiophenones [11] have shown that the  $n_{Se}/\pi^*_{CO}$ ,  $\sigma_{C-Se}/\pi^*_{CO}$  and  $\pi^*_{CO}/\sigma_{C-Se}$  orbital interactions strongly stabilise the gauche conformer with respect to the cis one. This behaviour, along with studies of the  $\alpha$ -sulfinylacetophenones and bis-thio-acetophenone, prompted us to investigate, by means of IR spectra, XRD, density functional theory and NBO calculations, some mixed acetophenones bearing in

the α position both the phenylseleno and ethylsulfinyl groups, as both should compete for the *syn-clinal* (*gauche*) and *syn-periplanar* geometries with respect to the carbonyl group. In particular, the 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones 1–5 (Scheme 1) were chosen, taking into account the fact that the orbital and electrostatic interactions that stabilise the conformers might be affected by changes in the conjugation involving the 4'-substituents.

Finally, following our continuous search for compounds with anti-inflammatory activity [12] our previous studies of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfonyl)-acetophenones [13] prompted us to determine the crystal and molecular structure of the related compounds 2-(ethanesulfinyl)-1-phenyl-2-(phenylselanyl)ethan-1-ones to be used in molecular docking studies to ascertain the possibility of being a COX-2 inhibitors.

### 2. Experimental

#### 2.1. Materials

All solvents for IR measurements were spectrograde and were used without further purification. The 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones 1 to 5 are new compounds and were obtained following a literature procedure [14]. A THF solution of 2-(ethylsulfinyl)-(4'-substituted)-acetophenone, prepared as previously described [15], was added to a solution of LDA in THF at 195 K. After 20 min, a solution of phenylselanyl bromide in THF was added dropwise to the enolate solution. After the reaction mixture reached room temperature (*ca.* 3 h), water was added and extraction with chloroform was performed. The organic layer was washed with diluted HCl and water, and dried over anhydrous magnesium sulphate. After evaporation of the solvent, the crude solid was purified through flash chromatography with a 1:1 solution

of hexane and ethyl acetate. As the title compounds present two stereogenic centres, *i.e.* the α-methine carbon (C<sub>R</sub> and C<sub>S</sub>) and the sulfinyl sulfur atoms (S<sub>R</sub> and S<sub>S</sub>) (Scheme 1), and the synthesis was performed without asymmetric induction, a racemic mixture of two pairs of diastereomers, [C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub>] and [C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub>], was obtained. The <sup>1</sup>H NMR spectrum of the crude product indicated that the relative abundance of the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> and C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub> diastereomers changes for each 4'-substituent and that the diastereomer corresponding to the higher field double quartet pairs (hydrogens 7 and 8 from Scheme 1) always predominates, as shown in Fig. 1. This diastereomer was isolated by solubilisation in chloroform, followed by the addition of a small amount of n-hexane while maintaining the system at low temperature (around 280 K) until crystal formation occurred. These crystals were collected, washed and dried. For the 4'-bromo and 4'-methyl derivatives, it was also possible to recover the other pure diastereomer with a second selective crystallisation. The purity of the crystallised products (compounds 1a–4a, 2b and 4b) was confirmed by <sup>1</sup>H NMR. Recrystallisation at low temperature in carbon tetrachloride gave pure crystals of compound 5a.

XRD analysis of compound **2b** [16] indicated that the lower field <sup>1</sup>H NMR double quartet pairs (Fig. 1) should be unambiguously assigned to the C<sub>S</sub>S<sub>R</sub>/C<sub>R</sub>S<sub>S</sub> diastereoisomer. Consequently, the more intense signals at higher fields in the <sup>1</sup>H NMR spectra of compounds **1–5** correspond to the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> diastereoisomer. In addition, the <sup>1</sup>H NMR analysis of the **4a** and **4b** methyl derivatives reveals that each C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> and C<sub>S</sub>S<sub>R</sub>/C<sub>R</sub>S<sub>S</sub> diastereomeric pair was purified at about 95% (Fig. 2). Suitable crystals for X-ray analysis for **3a** were obtained by vapour diffusion from chloroform/n-hexane at 283 K. The <sup>1</sup>H and <sup>13</sup>C NMR data and the elemental analysis for compounds **1–5** are presented in Tables 1 and 2.

#### 2.2. IR measurements

The IR spectra for the fundamental carbonyl region (1800–1600 cm<sup>-1</sup>) were recorded using a Michelson Bomem MB100 FTIR spectrometer with a 1.0 cm<sup>-1</sup> resolution in carbon tetrachloride, chloroform, dichloromethane and acetonitrile solutions (1.0 x 10<sup>-2</sup> mol dm<sup>-3</sup>) using a 0.519 mm sodium chloride cell. The spectra of the carbonyl first overtone (3600–3100 cm<sup>-1</sup>) were collected in carbon tetrachloride and dichloromethane solutions with a 1.00 cm quartz cell. The overlapping carbonyl bands (fundamental and first overtones) were deconvoluted by means of the Grams/32 curve fitting program, version 4.04 [17]. The populations of the conformers were estimated from the maximum of each component of the resolved carbonyl doublet, expressed as a percentage of absorbance, on the assumption of equal molar absorptivity coefficients for all the conformers.

#### 2.3. NMR measurements

<sup>1</sup>H and <sup>13</sup>C NMR spectra were recorded on a Bruker DRX 500 spectrometer operating at 500.130 and 125.758 MHz, respectively, for 0.1 mol/dm<sup>3</sup> solutions in CDCl<sub>3</sub>. <sup>1</sup>H and <sup>13</sup>C chemical shifts are reported in ppm relative to TMS as the internal standard.

## 2.4. X-ray measurements

X-ray crystallographic data were collected at 100 K on an Agilent SuperNova diffractometer (Dual source, Cu at zero) fitted with an Atlas detector using graphite-monochromated Cu K $\alpha$  radiation ( $\lambda = 1.54184$  Å). The data were processed with CrysAlisPro [18]. The structure was solved by direct methods with SHEXLS97 [19] and

refined by full-matrix least-squares on F<sup>2</sup> with SHELXL-2014 [20]. All non-hydrogen atoms were refined anisotropically and hydrogen atoms were geometrically placed and refined using the riding model approximation with  $U_{iso}$  (H) = 1.2–1.5  $U_{eq}$  (C). Towards the end of the refinement, additional electron density peaks were apparent. These were modelled as an independent molecule. To be consistent with the <sup>1</sup>H NMR results (see section 2.1), the major component of the disorder was refined with a site occupancy equal to 0.95. Having a small occupancy, constraints and restraints were introduced for the minor component to ensure a stable refinement. Thus, the anisotropic displacement parameters (ADPs) of the atoms were set to those of the major component. The ADPs of the carbon atoms were restrained to be nearly isotropic. Further, the atoms of the Se-C(phenyl) and C-C(phenyl) groups were restrained to be nearly planar. The aliphatic C-C bond distances were restrained with 1.50±0.01 Å and the C=O double-bond distance was set to 1.25±0.01 Å. Finally, for the Se13/S5 and Se31/S5' atoms, pairs of bond distances were restrained to be within 0.01 Å of each other. The key crystallographic data are given in Table 3. The programs ORTEP-3 for Windows [21], PLATON [22], DIAMOND [23] and QMol [24] were also used in the analysis.

CCDC 1475836 contains the supplementary crystallographic data for this study. These data can be obtained free of charge from the Cambridge Crystallographic Data Centre (CCDC), 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336 33; e-mail: <a href="mailto:deposit@ccdc.cam.ac.uk">deposit@ccdc.cam.ac.uk</a>.

#### 2.5. Theoretical calculations

A conformational search (HF/STO-3G theory level) was performed with Spartan '06 [25] software for the C<sub>R</sub>S<sub>R</sub> and C<sub>R</sub>S<sub>S</sub> diastereomers for compounds **1a–5a** and **2b** and **4b**, respectively. The obtained conformer geometries were used as initial inputs for all

calculations performed at 298 K with the Gaussian package programs (G03-E01) [26] with a hybrid Hartree-Fock density functional B3LYP method [27(a),27(b),27(c)] and the 6-31+G(d,p) basis set [28]. Full geometry optimisations and analytical vibrational frequency calculations were performed on the more stable conformers. Frequency analyses were carried out to verify the nature of the minimum state of all the stationary points obtained and to calculate the zero-point vibrational energy corrections. To estimate the solvation effects on the relative stability of the most relevant conformers, single-point calculations were conducted on the optimised structures using the polarisable continuum model (PCM) [29(a),29(b)]. The NBO 3.1 program [30] was used as implemented in the Gaussian 03 package and the reported NBO delocalisation energies (E2) were those given by the second-order perturbation theory. The partial atomic charges were calculated using Natural Population Analysis (NPA) [30].

#### 3. Results and discussion

#### 3.1. Enantiomeric $C_RS_R/C_SS_S$ diastereomer pair

The stretching frequencies of the fundamental and first overtone, as well as the absorbance percentage of the analytically resolved carbonyl band of (C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub>) 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl) acetophenones (**1a–5a**) in solvents of increasing relative permittivity [31] are reported in Table 4. The low signal to noise ratio precludes the characterisation of the band profile in the first overtone regions of **1a** (in CCl<sub>4</sub>) and **3a** (in CH<sub>2</sub>Cl<sub>2</sub>). The spectra in the carbonyl fundamental transition region show two components in acetonitrile for all compounds. The intensity of the low frequency doublet component progressively increases as the solvent polarity decreases until it becomes a singlet in compounds **1a** and **3a** in the low relative permittivity solvents chloroform and carbon tetrachloride, and the main doublet component (*ca*.

90%) for compounds **2a**, **4a** and **5a** in CCl<sub>4</sub>. The observed trend is illustrated in Fig. 3 for compound **2a**, chosen as being representative of the series. This trend seems to be mainly determined by the solvent polarity rather than by a 4'-substituent effect related to the different nature of the groups involved, going from the electron-withdrawing nitro group (**1a**) to the electron-donating OMe one (**5a**).

The carbonyl first overtone bands, recorded in CCl<sub>4</sub> and CH<sub>2</sub>Cl<sub>2</sub>, match exactly the behaviour of the corresponding bands in the fundamental and, when a doublet is present, for **1a**, **2a**, **4a** and **5a**, the two components emerge with relative intensities almost equivalent to those previously found (Table 4). Moreover, all the resolved peaks along with the single peak of **3a** (in CCl<sub>4</sub>) have frequencies twice that of the fundamental minus twice the mechanical anharmonicity of  $20 \pm 2$  cm<sup>-1</sup> [32]. The matching of the carbonyl band profiles in the two frequency regions suggests the presence of at least two conformers for the referred compounds, and at least one for **3a** (in CCl<sub>4</sub>), ruling out the existence of any vibrational effects in the fundamental transition of the  $v_{CO}$  mode [33(a),33(b)].

Aiming to determine the geometries and vibrational frequencies of the minimum energy conformations in the gas phase, B3LYP/6-31G+(d,p) calculations were performed for the  $C_RS_R$  diastereomer of compounds **1a–5a.** The calculation results, summarised in Table 5, indicate the existence of three distinct conformations, classified in order of decreasing stability for the whole series as  $c_1$  (82–92%),  $c_2$  (7–17%) and  $c_3$  (< 1%). The dihedral angles  $\alpha$  and  $\alpha$  formed by the carbonyl and the C-S and C-Se groups, respectively, are almost equivalent for all the 4'-derivatives and compel the  $c_1$  conformer to assume *syn-clinal* ( $\alpha \approx 90^{\circ}$ ) and *syn-periplanar* geometries ( $\alpha \approx -30^{\circ}$ ), respectively. Conversely, for both the  $c_2$  and  $c_3$  conformers, the C–S/C=O groups are nearly coplanar ( $\alpha \approx 30^{\circ}$ ), while the C–Se/C=O groups are almost perpendicular ( $\alpha \approx -30^{\circ}$ ), while the C–Se/C=O groups are almost perpendicular ( $\alpha \approx -30^{\circ}$ ).

90°). The major difference between the two conformers in each series consists of the different orientation of both the sulfinyl and the phenyl groups with respect to the carbonyl one, revealed, respectively, by the  $\delta$  and  $\gamma'$  dihedral angles. In particular, the  $c_2$  conformer displays a *quasi-anti* geometry ( $\delta$  *ca.* 164° and  $\gamma'$  *ca.* 130°), while the  $c_3$  conformer presents a *quasi-syn* orientation ( $\delta$  *ca.* -94° and  $\gamma'$  *ca.* -119°). The computed molecular structures of the three conformers for compound **3a**, representative of the whole series, are shown in Fig. 4.

Additionally, it should be noted that the 4'-substituents hardly affect the geometry of each conformer in the series, with the exception of the rotation of the ethyl group indicated by the  $\gamma$  dihedral angle which varies from about 171° in compounds **1a–4a** to 61° ( $c_2$ ) and -75° ( $c_3$ ) in the **5a** derivative.

The least stable  $c_3$  conformer exhibits the highest  $v_{CO}$  frequency, while those of the  $c_1$  and  $c_2$  conformers differ by less than 3 cm<sup>-1</sup>. As expected, the  $v_{CO}$  frequency of all conformers increases progressively in the series 1a–5a, going from the electron-donor methoxy derivative 5a to the electron-attracting nitro derivative 1a. Therefore, in line with the experimental findings, the more intense component at the lower frequency of the carbonyl doublet in solution may be ascribed to both the  $c_1$  and  $c_2$  conformers, and the less intense higher frequency one can be ascribed to the less stable  $c_3$  conformer (Table 4). PCM single-point calculations performed for compounds 1a–5a (Table 6) confirm these assignments. In fact, the sum of the relative abundances of the most stable  $c_1$  and  $c_2$  conformers decreases progressively as the solvent polarity increases and concomitantly the population of the  $c_3$  conformer significantly increases. An important role in the observed larger solvation effect on the  $c_3$  conformer with respect to the  $c_1$  and  $c_2$  ones seems to be played by the opposite geometry assumed by the two  $C^{\delta_+}$ = $O^{\delta_-}$  and  $S^{\delta_+}$ = $O^{\delta_-}$  dipoles, namely syn in the former conformer and anti in the latter ones. As a

matter of fact, the *syn* orientation forces the two dipoles to be closer, thus leading to a larger local dipole moment. Moreover, the larger Repulsive Field Effect (RFE) [33(a)] between the two dipoles in the  $c_3$  conformer, with respect to the  $c_1$  and  $c_2$  ones, increases its carbonyl bond order and consequently its  $v_{CO}$  frequency.

In order to determine the nature of the orbital interactions that affect the stability of the different conformers of compounds **1a**, **3a** and **5a**, some selected NBO interactions, NPA charges and interatomic distances, calculated at the B3LYP/6-31+G(d,p) level [31], are reported in Tables 7, 8 and 9, respectively.

The strongest interactions acting on the phenacyl group are the  $\pi_{C25C26} \to \pi^*_{C2C1}$  conjugation and the LP<sub>O1</sub> $\to \sigma^*_{C2C3}$  and LP<sub>O1</sub> $\to \sigma^*_{C2C25}$  through bond coupling interactions [34]. For all the conformers in the whole series, the latter interactions are almost constant (ca. 20 kcal mol<sup>-1</sup>), while the former increases progressively going from the 4'-electron-attracting nitro derivative **1a** (ca. 18 kcal mol<sup>-1</sup>) to the 4'-electron-donating methoxy derivative (ca. 23 kcal mol<sup>-1</sup>).

The suitable  $\phi$ ' torsion angles for the  $c_1$  (ca. 78°) and  $c_3$  conformers (ca. -88°) allow the occurrence of the LP<sub>Se13</sub> $\rightarrow \sigma^*_{C3S5}$  interaction (ca. 6 kcal mol<sup>-1</sup>), which is weakened to ca. 3.5 kcal mol<sup>-1</sup> in the  $c_2$  conformer. In addition, the favourable  $\phi$  angle (ca. -77.5°) gives rise to the LP<sub>S5</sub> $\rightarrow \sigma^*_{C3Se13}$  interaction (3.4 kcal mol<sup>-1</sup>) uniquely in the  $c_3$  conformer.

The Y-C-C=O fragments (Y = S or Se) exhibit four additional orbital interactions that are maximised as the  $\alpha$  or  $\alpha$ ' torsional angles approach 90°, that is the LP<sub>Y</sub>  $\rightarrow \pi^*_{CO}$ ,  $\pi_{CO} \rightarrow \sigma^*_{C-Y}$  and  $\sigma_{C-Y} \rightarrow \pi^*_{CO}$  interactions, as well as the unusual  $\pi^*_{CO} \rightarrow \sigma^*_{C-Y}$  interaction [35]. Therefore, as suggested by the conformer geometries in Table 5 and confirmed by the NBO energies in Table 7, these interactions almost exclusively stabilise the  $c_I$  conformer when the sulfur atom is involved ( $\alpha$  of around 90° for all

derivatives), and solely the  $c_2$  and  $c_3$  conformers as concerns the selenium fragment ( $\alpha$ ' of around -90° for the whole series **1a–5a**).

The suitable geometry of the  $c_I$  conformers leads to contacts shorter than the sum of the van der Waals ( $\Sigma$ vdW) radii between the oppositely charged  $O^{\delta_{-(35)SO}}$  and  $H^{\delta_{+}}(27)_{o-Ph(CO)}$  atoms, as well as the  $O^{\delta_{-(1)CO}}$  and  $H^{\delta_{+}}(7)CH2(EI)$  atoms (Tables 8 and 9), which contribute electrostatically to their stabilisation. Moreover, the short distance  $O_{(35)}...H_{(27)}$  is also responsible for the considerable  $LP_{O35} \rightarrow \sigma^*_{C26H27}$  stabilising orbital interaction (hydrogen bond) (mean energy of ca. 3.0 kcal mol<sup>-1</sup>). Analogously, the proper values of the  $\delta$  and  $\gamma'$  dihedral angles in the  $c_2$  and  $c_3$  conformers allow the contacts, shorter than the  $\Sigma$ vdW radii, between the negative  $O_{(35)SO}$  and positive  $H_{(16)o-PhSe}$  atoms to give raise to the  $LP_{O35} \rightarrow \sigma^*_{C15H16}$  stabilising orbital interactions (hydrogen bonds). These hydrogen bonds in the molecular structure of the  $c_1$ ,  $c_2$  and  $c_3$  conformers of the representative compound 2a are illustrated in Fig. 4. In addition, the  $c_1$  and  $c_3$  conformers are slightly stabilised by the weak electrostatic interactions originated by the contacts  $Se^{\delta_{-(13)}} ... H^{\delta_{+(7)CH2(Ei)}}$  close to the  $\Sigma$ vdW radii, as well as by the related weak  $LP_{Se13} \rightarrow \sigma^*_{C6H7}$  orbital interaction.

Similarly to the 2-sulfinylacetophenones [8] in the geometries of the  $c_2$  ( $\delta$  ca. 164°) and  $c_3$  ( $\delta$  ca. -94°) conformers, the negatively charged carbonyl oxygen and positive sulfinyl sulfur atoms, separated by intramolecular contacts shorter than the  $\Sigma$ vdW radii, are responsible for an electrostatic stabilising interaction that is stronger in the  $c_3$  conformers. It is interesting to note that the *anti-periplanar* configuration of the sulfinyl sulfur atom with respect to the carbonyl group evidenced in the  $c_2$  conformer favours the weak LP<sub>O1</sub> $\rightarrow$  $\sigma$ \*<sub>S5O35</sub> charge transfer interaction, which is absent in the  $c_3$  conformer.

Additional stabilising electrostatic and charge transfer interactions caused by short contacts between oppositely charged atoms take place in the  $c_1$  conformers, as a

consequence of the *quasi-syn-periplanar* orientation of the  $C^{\delta+}=O^{\delta-}$  and  $C^{\delta-}-Se^{\delta+}$  groups, along with the *quasi-anti-periplanar* geometry of the C=O and Se-C<sub>Ph</sub> groups ( $\beta$ ' ca. - 156°) and to a lesser extent by the  $O^{\delta-}_{(35)SO}....C^{\delta+}_{(2)CO}$  contact with a distance slightly larger than the  $\Sigma$ vdW radii. Moreover, the *quasi-syn-periplanar* orientation is also responsible for the weak  $LP_{O1} \rightarrow \sigma^*_{Se13C14}$  orbital interaction.

As previously discussed, the repulsion between the  $C^{\delta_+}=O^{\delta_-}_{(1)}$  and  $S^{\delta_+}=O^{\delta_-}_{(35)}$  dipoles is significantly larger in the  $c_3$  conformers, favoured by their syn orientation, opposite to the anti geometry assumed by the  $c_1$  and  $c_2$  conformers. Therefore, the sizable RFE between the two dipoles should destabilise, to a greater extent, the  $c_3$  conformers, thus increasing the carbonyl bond order and consequently the  $v_{CO}$  frequencies. Both the computed and experimental findings fully support this assertion as the carbonyl frequencies of the  $c_3$  conformers were found to be higher than those of the  $c_1$  and  $c_2$  conformers by ca. 20 cm<sup>-1</sup> (theoretical values) and ca. 15 cm<sup>-1</sup> (experimental values).

The sum of the NBO orbital interactions ( $\Sigma E$ ) for compounds **1a**, **3a** and **5a** indicates that the  $c_3$  conformer should be the most stable by ca. 93.7 kcal mol<sup>-1</sup>, followed by the  $c_1$  (ca. 90.4 kcal mol<sup>-1</sup>) and  $c_2$  (ca. 88.1 kcal mol<sup>-1</sup>) conformers. Moreover, the sum of the NBO energies increases progressively for each conformer, going from electroattracting (**1a**) to hydrogen (**3a**) to electron-donating (**5a**) substituents.

Conversely, as noted above, the coulombic repulsion between the  $C^{\delta+}=O^{\delta-}$  and  $S^{\delta+}=O^{\delta-}$  dipoles significantly destabilises the  $c_3$  conformer with respect to the  $c_1$  and  $c_2$  ones. Therefore, it is reasonable to infer that this effect likely prevails over the sum of NBO orbital interactions to determine the lowest computed stability of the  $c_3$  conformer. Alternatively, the  $c_1$  conformer is more stable than the  $c_2$  one by a mean value of  $c_3$ . 2.3 kcal mol<sup>-1</sup>. It should be stated that this value is in agreement with the computed larger stabilisation of the  $c_1$  conformer with respect to the  $c_2$  conformer (Table 5). This trend

may be attributed mainly to both the large coulombic stabilisation between  $O^{\delta_{-CO}}...Se^{\delta_{+}}$  atoms and the  $LP_{O1} \rightarrow \sigma^{*}_{Se13C14}$  orbital interaction. Therefore, the computed order of stability of the three conformers follows from a balance of electrostatic and orbital interactions.

Crystals of **3a** were resolved as a whole molecule disorder model. The molecule of the major component of the disorder, with a site occupancy = 0.95, is illustrated in Fig. 5(a), and that of the minor component is in the Supplementary Materials as Fig. S1. An overlay diagram of the two components of the disorder is shown in Fig. 5(b). From this, it is evident that the major difference between the molecules is related to the chirality of the sulfinyl-S5/S5' atoms. Minor conformational differences are apparent, as seen in the dihedral angle between the two aromatic rings of 3.14(15)° (major component) and 9.1(12)° (minor). The large standard uncertainty for the minor component notwithstanding, this difference is consistent with a greater deviation from a parallel disposition of the rings in the latter. The other difference relates to the relative orientation of the terminal ethyl groups, as seen in the C3–S5–C6–C9 torsion angles of 175.2(2) and -141(7)° for the major and minor components, respectively.

X-ray analysis of 3a indicates that this compound assumes, in the solid state, a conformation where the geometry is close to that of the  $c_2$  conformer found in the gas phase, as evidenced by the almost coincident values of the torsional  $\alpha$ - $\phi$ ,  $\alpha$ ' angles except for  $\beta$ '- $\gamma$ ' which differ significantly in the solid (Table 5). In fact, in the gas phase, the sulfinyl and phenyl groups display a *quasi-syn-periplanar* geometry, while in the crystal, they lie in the opposite direction. It should be noted that the single molecule in the solid is stabilised by the same intramolecular attractive electrostatic interactions that occur in the gas phase between the oppositely charged carbonyl oxygen and sulfinyl

sulfur atoms, connected by a distance shorter than the  $\Sigma vdW$  radii ( $\Delta l = 0.43$  Å), but no longer than the  $O^{\delta_-}...H^{\delta_+}$  contacts (Table 9).

In order to obtain the larger energy gain, the molecular packing (Fig. 6), defined by the major component of the disorder, features readily identifiable C–H…O, C–H…S,  $\pi...\pi$  and Se… $\pi$  interactions that cooperate to stabilise the three-dimensional structure. The most prominent interactions are the tight methine-C-H…O(sulfinyl) interactions that lead to centrosymmetric, eight-membered {...OSCH}2 synthons. Dimeric aggregates are connected by phenyl-C–H…O(carbonyl), phenyl-C–H…O(sulfinyl) and  $\pi...\pi$  interactions, with the latter being between centrosymmetric related Se-bound phenyl rings. Geometrical details are given in Table 10. In addition, a close Se(lp)... $\pi$ (phenyl) contact is noted; such interactions and the supramolecular aggregates they sustain have been reviewed recently [36]. The Se…ring centroid separation is 3.72 Å, so the contact in 3 is most likely an example of a semi-localised Se(lp)... $\pi$ (phenyl) contact as the Se atom is closer to the C19<sup>i</sup> [3.399(3) Å] and C21<sup>i</sup> [3.516(3) Å] atoms, rather than to the centroid (a delocalised interaction).

#### 3.2. Enantiomeric C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub> diastereomer pair

The relevant experimental IR data for the Br (2b) and Me derivatives (4b) are collected in Table 11, to compare with the analogous Table 4 for the  $C_RS_R/C_SS_S$  diastereomer.

The carbonyl stretching band shows two components in the non-polar solvent carbon tetrachloride for both compounds, with the low frequency one being the most prominent. Its relative intensity decreases at different extents for both derivatives as the solvent polarity increases, until it vanishes for compound **2b** in acetonitrile, as evidenced in Figs. 7 and 8 for **2b** and **4b**, respectively.

The carbonyl first overtone region shows two components for **4b** in CCl<sub>4</sub> and for **2b** in CH<sub>2</sub>Cl<sub>2</sub>, at frequencies twice those of the fundamental minus twice the mechanical anharmonicity of ca. 20 cm<sup>-1</sup> [32], and relative intensities that match those of the corresponding fundamental ones. These findings, in agreement with the results previously described for the  $C_RS_R/C_SS_S$  diastereomer, indicate the existence of at least two conformers and exclude the occurrence of any vibrational effect [33(a),33(b)].

The B3LYP/ 6-31G+(d,p) calculation results, summarised in Table 12, indicate, at variance with those performed for the enantiomeric pair  $C_RS_R/C_SS_S$ , the existence in the gas phase of only two *cis* conformers. For both the **2b** and **4b** compounds, the  $c_1$  conformer is the most stable and the less polar and, moreover, presents the lowest  $v_{CO}$  frequency. All the dihedral angles of the  $c_1$  conformers assume quite similar values for both derivatives. The same occurs for the  $c_2$  conformers, with the significant exception of the  $\gamma$  torsion angle, related to the rotation of the ethyl group, that changes from about  $163.7^{\circ}$  in **2b** to  $-59.8^{\circ}$  in **4b**.

Moreover, the  $\alpha$  dihedral angles, as well as the  $\alpha'$  ones, are almost coincident for all the conformers in all compounds. This implies that the carbonyl group in the  $c_1$  and  $c_2$  conformers of both derivatives adopts a *quasi-syn-periplanar* geometry with respect to the C-S group ( $\alpha \approx 39^{\circ}$ ) and a *syn-clinal* orientation with respect to the C-Se one ( $\alpha' \approx 90^{\circ}$ ). On the contrary, the carbonyl and sulfinyl groups have opposite geometries, that is *anti-periplanar* ( $\delta \approx 171^{\circ}$ ) in the  $c_1$  conformers and *syn-clinal* ( $\delta \approx 69^{\circ}$ ) in the  $c_2$  ones. The computed molecular structures of the two stable conformers are shown in Fig. 9 for **2b** and are representative of both compounds.

As evidenced by the PCM single-point calculations reported in Table 13, the relative abundances of the less stable  $c_2$  conformers at higher frequencies decrease noticeably when moving from the gas phase towards solvents of increasing relative permittivity,

and to a major extent for the **2b** derivative. The trend shown by the  $c_2$  conformers for both compounds agrees reasonably well with the experimental IR findings observed for the high frequency and less intense carbonyl doublet components in CCl<sub>4</sub>. Therefore, these components should be assigned to the  $c_2$  conformers and, analogously, the low frequency more intense ones can be assigned to  $c_1$ . The observed larger solvation effect on the  $c_2$  conformer for both **2b** and **4b** is very similar to that previously described for the  $c_3$  conformer of the  $C_RS_R/C_SS_S$  diastereomer and can be explained with the same arguments, that is the syn geometry (about 97°) adopted by the  $C^{\delta+}=O^{\delta-}$  and  $S^{\delta+}=O^{\delta-}$  dipoles with respect to the *anti*-relationship assumed in the  $c_1$  conformer. Moreover, the  $S^{\delta+}=O^{\delta-}$  dipole solvation is partially hindered in the **4b** compound by the closeness of the methyl group to the sulfinyl oxygen atom, as evidenced by the smaller value of the  $\gamma$  dihedral angle of -59.8° in comparison to -163.7° in the **2b** derivative. As a consequence, the solvation effect should be larger for **2b** with respect to **4b**, in agreement with the calculated and experimental IR results.

The comparison between the experimental IR intensities of the higher carbonyl frequency doublet component reveals a more pronounced solvent effect for the (RR/SS)  $c_3$  conformer of 1a-5a with respect to the (RS/SR)  $c_2$  conformer of 2b and 4b (see Tables 11 and 4). This behaviour is likely a consequence of the presence of the short contact  $O_{(35)SO}...H_{(16)}$  (o-Ph) (hydrogen bond) in the former and, on the contrary, of its absence in the latter. Furthermore, both the (RR/SS)  $c_2$  and (RS/SR)  $c_1$  conformers display the same  $O_{(35)SO}...H_{(16)}$  (o-Ph) interaction, which partially inhibits to about the same extent as the solvent approach.

In order to analyse the main factors that determine the relative stabilities of the  $c_1$  and  $c_2$  conformers for compounds **2b** and **4b**, selected NBO interactions, NPA charges

and interatomic distances, calculated at the B3LYP/6-31+G(d,p) level [31], are reported in Tables 14, 15 and 16, respectively.

The phenacyl group of both the  $c_1$  and  $c_2$  conformers is affected by the same strong orbital interactions (about 20 kcal mol<sup>-1</sup>) that take place in the enantiomeric pair  $C_RS_R/C_SS_S$  diastereomer, that is the  $\pi_{C25C26}\rightarrow\pi^*_{C2O1}$  conjugation and the  $LP_{O1}\rightarrow\sigma^*_{C2C3}$  and  $LP_{O1}\rightarrow\sigma^*_{C2C25}$  through bond coupling interactions [34]. The former, like in the  $C_RS_R/C_SS_S$  compounds, was found to be slightly stronger for the 4'-electron-donating derivative **4b**.

The LP<sub>Se13</sub> $\rightarrow \sigma^*_{C3S5}$  interaction, found in the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> series also, stabilises to a similar extent both conformers by less than 6.0 kcal mol<sup>-1</sup>, while the LP<sub>S5</sub> $\rightarrow \sigma^*_{C3Se13}$  one, that is absent in the  $c_2$  conformer due to the unfavourable  $\delta$  (ca. 68.2 °) and  $\phi$  (-59.0 °) dihedral angles, stabilises only the  $c_1$  conformer by ca. 4.2 kcal mol<sup>-1</sup>.

As in the (RR/SS) series, the extent of some interactions depends on the values assumed by specific dihedral angles. For instance, in the case of  $\alpha$  and  $\alpha'$ , the unsuitable *synclinal* geometry of the  $c_I$  and  $c_2$  conformers ( $\alpha$  of ca. 39.8° for both compounds) weakens the hyperconjugative interaction  $\sigma_{C-S} \to \pi^*_{CO}$  (about 1 kcal mol<sup>-1</sup>) and the unusual  $\pi^*_{CO} \to \sigma^*_{C-S}$  orbital interaction (ca. 2.3 kcal mol<sup>-1</sup>). On the contrary, the appropriate  $\alpha'$  value of the O=C-C-Se moiety close to -90° for both conformers of **2b** and **4b** favours the  $\sigma_{C-Se} \to \pi^*_{CO}$  hyperconjugation, as well as the  $\pi^*_{CO} \to \sigma^*_{C-Se}$  orbital interaction. The former stabilises the  $c_I$  conformer by ca. 7.2 kcal mol<sup>-1</sup> and the  $c_2$  conformer by ca. 5.8 kcal mol<sup>-1</sup>, while the latter contributes to a further stabilisation of ca. 4.9 kcal mol<sup>-1</sup> for the  $c_I$  and 5.4 kcal mol<sup>-1</sup> for the  $c_I$ . In addition, the  $\alpha$  angles allow the contacts  $O^{\delta_{-(1)CO}}$ ... $S^{\delta_{+(5)SO}}$  to be shorter than the  $\Sigma$ vdW radii by -0.26 Å in  $c_I$  and -0.36 Å in  $c_I$ .

Similarly, the proper values of the  $\beta$ ,  $\beta'$  and  $\gamma'$  angles contribute to uniquely stabilising the  $c_I$  conformers, as they constrain the  $H^{\delta_+}_{(16)o\text{-PhSe}}$  and  $O^{\delta_-}_{(35)SO}$  atoms, as well as the  $O^{\delta_-}_{(1)CO}$  and  $H^{\delta_+}_{(7)CH2(Et)}$  atoms, to contacts significantly shorter than the  $\Sigma$ vdW radii ( $\Delta$ l ca. -0.39 and -0.30 Å respectively). Moreover, the geometry of the  $c_I$  conformers gives rise to the  $LP_{O35} \rightarrow \sigma^*_{C15H16}$  and  $LP_{O1} \rightarrow \sigma^*_{C6H7}$  orbital interactions (hydrogen bonds) of mean energies ca. 3.8 and 1.3 kcal mol<sup>-1</sup>, respectively, absent in the  $c_2$  conformers. This is evidenced in the molecular structures of the two conformers of **2b**, as reported in Fig. 9.

Overall, the computed and experimental order of stability of the two conformers for compounds **2b** and **4b** is in agreement with the sum of all the relevant NBO orbital interactions, which indicates that the  $c_1$  conformer is the more stable by ca. 9 kcal mol<sup>-1</sup>.

The X-ray single crystal analysis indicates that the geometry in the solid state of compound 2b is strictly correlated to that of the  $c_2$  conformer in the gas phase, as evidenced by the similar values of the torsional angles, except for the  $\gamma$  and  $\gamma'$  ones corresponding to the orientation of the ethyl and (Se)-phenyl groups (Table 12). The solid is stabilised by the same intramolecular attractive electrostatic interactions that occur in the gas phase for the  $c_2$  conformer, as suggested by the relevant short contacts reported in Table 16. A more detailed view on the molecular packing and the stabilising intermolecular interaction that occur for 2b in the solid state can be found in the literature [16]. It should be noted that the conformation of the minor component of the 3a crystal structure with the  $C_RS_S$  configuration (Table 5) is similar to that of the 2b single crystal structure and, therefore, to that of the  $c_2$  conformer of compounds 2b and 4b in the gas phase.

#### 4. Conclusions

A conformational study of a selection of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones bearing the substituents NO<sub>2</sub> **1a**, Br **2a**, H **3a**, Me **4a** and OMe **5a** for the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> enantiomeric pair and Br **2b** and Me **4b** for the C<sub>R</sub>S<sub>S</sub>/C<sub>S</sub>S<sub>R</sub> pair was performed through the analysis of their IR carbonyl bands, supported by B3LYP/6-31+G(d,p) and single-point PCM calculations along with NBO analysis for **1a**, **3a**, **5a**, **2b** and **4b** and X-ray diffraction of **3**.

For the C<sub>R</sub>S<sub>R</sub>/C<sub>S</sub>S<sub>S</sub> compounds, the theoretical results indicated the existence, in the gas phase, of three conformers whose relative abundances are ca. 82–92% (c<sub>1</sub>), ca. 7– 17% ( $c_2$ ) and ca. 1% ( $c_3$ ). The computed  $v_{CO}$  frequencies for the  $c_1$  and  $c_2$  conformers differ by less than 3 cm<sup>-1</sup>, while the  $c_3$  one is about 20 cm<sup>-1</sup> higher. PCM single-point calculations indicate that the relative population of the  $c_3$  conformer increases as the solvent relative permittivity increases, going from CCl<sub>4</sub> to CH<sub>3</sub>CN. The larger solvation effect and the concomitant frequency increase of the  $c_3$  conformers with respect to the  $c_1$ and  $c_2$  ones are likely related to the closeness of the  $C^{\delta+}=O^{\delta-}$  and  $S^{\delta+}=O^{\delta-}$  dipoles, which are syn oriented in the former and anti in the latter ones. This trend, compared to the experimental findings, allows to ascribe the  $c_1$  and  $c_2$  conformers to the low frequency more intense component of the  $v_{CO}$  doublet and the  $c_3$  conformer to the high frequency less intense one. All the conformers are electrostatically stabilised by short contacts (hydrogen bonds) between the sulfoxide oxygen  $O^{\delta_{-(35)SO}}$  and one o-Ph-hydrogen  $H^{\delta_{+}}$  o-Ph atoms. For the  $c_2$  and  $c_3$  conformers, these contacts involve the Se-Phenyl group ohydrogen, while for the  $c_1$  one, it is the phenacyl group o-hydrogen. The sum of the NBO energies suggests that the  $c_3$  conformer should be the most stable one, in contrast with the computed stability order previously described. However, the effect of the strong Repulsive Field Effect between the syn oriented  $C^{\delta+}=O^{\delta-}$  and  $S^+=O^-$  dipoles overcomes the sum of the NBO energies, thus destabilising at a larger extent this conformer with respect to the  $c_1$  and  $c_2$  ones. X-ray data of the major component of the disorder (compound **3a**) have shown that in the solid state, a single molecule assumes a conformation very similar to that of the  $c_2$  conformer in gas phase. Furthermore, in the crystal, molecules are linked in centrosymmetrical pairs through  $\pi...\pi$  interactions between Se-bound phenyl rings, along with phenyl-C-H...O(carbonyl) and phenyl-C-H...O(sulphinyl) interactions.

Conversely, the computational results for the  $C_RS_s/C_SS_R$  compounds **2b** and **4b** indicate the existence of two conformers  $c_I$  and  $c_2$ , with the former being the most abundant (ca. 99%), the less polar and with the lowest  $v_{CO}$  frequency. The good agreement between the PCM single-point calculations and the experimental solvent effect trend of the  $v_{CO}$  doublet components allows to assign the higher frequency  $c_2$  conformer to the higher  $v_{CO}$  component in the IR spectra. The  $c_I$  conformer is stabilised by the  $LP_{(S)} \rightarrow \sigma^*_{C-Se}$  and  $LP_{O(S-O)} \rightarrow \sigma^*_{C6-H7}$  orbital interactions, along with the intramolecular hydrogen bond between the sulfinyl oxygen  $O^{\delta_{-(35)SO}}$  and Se-Phenyl o-hydrogen  $H^{\delta_{+}}_{o-Ph(16)}$  atoms. The absence of the latter interaction in the  $c_2$  conformer leaves the sulfinyl oxygen atom accessible to the solvent molecules and thus accounts for its lower relative stability in the gas phase and its stronger solvation.

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#### Figure and scheme captions

**Fig. 1.** NMR signals of methylenic hydrogens of the crude mixture of diastereoisomers of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones 1-5 with preferential formation of diastereomer  $C_SS_S/C_RS_R$  as a racemate.

**Fig. 2.** <sup>1</sup>H-NMR signals (double quartet pairs) of diastereotopic methylene hydrogen atoms [CH<sub>2</sub>S(O)] of the purified ( $C_RS_R/C_SS_S$ ) and ( $C_SS_R/C_RS_S$ ) diastereoisomers of 2-(phenylselanyl)-2-(ethylsulfinyl)-4'-methylacetophenone (**4a** and **4b**).

**Fig. 3.** IR spectra of  $(C_RS_R/C_SS_S)$  2-(phenylselanyl)-2-(ethylsulfinyl)-4'-bromoacetophenone (2a) showing the analytically resolved carbonyl stretching band, in: carbon tetrachloride [fundamental (a) and first overtone (b)], chloroform (c), dichloromethane (d) and acetonitrile (e).

**Fig. 4.** Molecular structures of the conformers of  $(C_RS_R)$  4'-Br-PhC(O)CH[S(O)Et][SePh] (2a) obtained at the B3LYP/6-31+G(d,p) level. Adopted colours: H = white, C = grey, O = red, S = yellow and Se = orange.

**Fig. 5.** (a) Molecular structure of the major component comprising the crystal structure of **3a**, showing atom-labelling and displacement ellipsoids at the 70% probability level for non-H atoms, (b) overlay diagram of the two components of the structure of **3**. The major component is shown in the red image, and the molecules have been aligned so that the C=O and Se atoms are coincident.

**Fig. 6**. Molecular packing in **3a** (major component only) viewed in projection down the c-axis. The C–H…O, C–H…S and  $\pi$ … $\pi$  interactions are shown as orange, blue and purple dashed lines, respectively.

**Fig. 7.** IR spectra of  $(C_SS_R/C_RS_S)$  2-(phenylselanyl)-2-(ethylsulfinyl)-4'-methylacetophenone (**4b**) showing the analytically resolved carbonyl stretching band, in: carbon tetrachloride [fundamental (a) and first overtone (b)], chloroform (c), dichloromethane (d) and acetonitrile (e).

**Fig. 8.** IR spectra of  $(C_SS_R/C_RS_S)$  2-(phenylselanyl)-2-(ethylsulfinyl)-4'-bromoacetophenone (**2b**) showing the analytically resolved carbonyl stretching band, in: carbon tetrachloride (a), chloroform (b), dichloromethane [fundamental (c) and first overtone (d)] and acetonitrile (e).

**Fig. 9.** Molecular structures of the conformers of  $(C_RS_S)$  4'-Br-PhC(O)CH[S(O)Et][SePh] (**2b**) obtained at the B3LYP/6-31+G(d,p) level. Adopted colours: H = white, C = grey, O = red, S = yellow and Se = orange.

**Figure S1.** Molecular structure of the minor component comprising the crystal structure of **3b**, showing atom-labelling and displacements ellipsoids at the 70% probability level for non-H atoms.

**Scheme 1.** Atom labelling of 4'-substituted 2-(phenylselanyl)-2-(ethylsulfinyl)-acetophenones, 4'-Y-PhC(O)CH[S(O)Et][SePh] and definition of the relevant dihedral angles for the  $C_SS_S/C_RS_R$  (a) and  $C_RS_S/C_SS_R$  (b) series.