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PAPER

Thermodynamic parameters of the pedal motion in the crystal structures of two bromomethylated azobenzenes†

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Two bromomethylated azobenzene derivatives were characterized by X-ray crystallography at 100 K after fast cooling, and showed disorder of the central N-atoms. The structures were re-determined over a range of temperatures, providing evidence for dynamic disorder due to pedal motion of the central N=N bond.

Using van’t Hoff plots, thermodynamic parameters for the pedal motion were determined. Computationally very cheap Atom–Atom-Force Field (AA-CLP) calculations were employed, which showed that the differences in dynamic disorder enthalpy between the two compounds are predominantly due to intermolecular interactions. AA-CLP calculations and gas phase electronic structure calculations were employed to show the link between intermolecular interactions and activation energy for pedal motion.

Introduction

Similar to compounds with C=C double bonds, azobenzenes¹ can exist as two different configurational isomers of the N=N bond; the (E)-isomer tends to be planar and about 50 kJ mol⁻¹ more stable than the (Z)-isomer,² which presents an angular geometry for steric reasons. Photochromic properties of azobenzenes originate in the photochemical isomerization between (E)- and (Z)-states.³ This isomerization leads to a considerable reduction in length, making azobenzenes useful building blocks for the introduction of large scale motion on the molecular level,⁴ such as photocontrollable cation binders⁵ and catenanes,⁶ photomechanically controlled π-conjugated systems,⁷ and even Bronsted bases.⁸ Being interested in the construction of azobenzene-tetrathiafulvalene macrocycles,⁹ we have prepared several bromomethylated azobenzene derivatives as precursors. Two of these derivatives, (E)-3,3'-bis(bromomethyl)azobenzene 1 and (E)-4,4'-bis(bromomethyl)azobenzene 2, were shown to display dynamic disorder in the solid state due to the pedal motion of the central double bond (Fig. 1).

Disorder in crystals can either be static or dynamic. When static, the overall population of the disordered moieties is locked in during the crystallization process. When it is dynamic, however, populations change with temperature. Depending on the activation energy of the process, the speed of cooling can influence the temperature at which the equilibrium becomes frozen in, and the population ratio of the two conformers at which this happens. Measurements at several temperatures can allow to find out whether the

![Fig. 1. Plot of the two conformers of 1 (top) and 2 (bottom) in the crystal structure at 90 K. Displacement ellipsoids are at the 50% level.](Image)

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† Electronic supplementary information (ESI) available: X-ray crystallography experimental details (Table S1 and S2), details of electronic structure calculations, and CIF files. CCDC 1063365–1063381. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5ce00905g
equilibrium is static or dynamic, and to determine its thermodynamic parameters.

Dynamic behavior of molecules in the crystal phase has been the subject of considerable interest in the past. Bicycle pedal motion\(^\text{10}\) in double bond compounds, ranging from azobenzenes\(^\text{11}\) over stilbenes\(^\text{12}\) to diphosphines,\(^\text{13}\) has been implicated in causing apparently shorter double bonds,\(^\text{14}\) inducing of concerted motion in crystals,\(^\text{15}\) as well as influencing the gas adsorption process\(^\text{16}\) or solvent exchange in MOFs.\(^\text{17}\) Additionally, it can influence the photodimerisation process of olefins, where it either helps to overcome the usual topochemical constraints,\(^\text{18}\) or, on the contrary, prevents cyclizations where they would be expected to occur.\(^\text{19}\)

### Results and discussion

#### van’t Hoff plots and thermodynamic parameters

In the crystal structures of compounds 1 and 2, the typical disorder due to pedal motion of the N≡N moiety, reported previously for two other azobenzene derivatives,\(^\text{11}\) is observed. Exploratory structure determinations at room temperature and at 100 K with slow cooling yielded quite different results for the populations of the two conformers for 1, whereas for 2 the difference was rather small. This observation prompted us to further study the thermodynamics of the pedal motion in both compounds. There are three examples in the literature on the study of thermodynamics of “pedaling” conformers by means of a van’t Hoff plot analysis\(^\text{20,21}\) (compounds 3–5), which all point to entropic and enthalpic differences between the two conformers being constant throughout the temperature range being studied. In other words, the van’t Hoff plot, which displays \(\ln(K)\) with respect to \(1/T\), is linear, down to a point where the equilibrium freezes in due to the temperature of the \(kT\) bath becoming insufficient to overcome the activation energy for the pedal motion. However, a fourth example is known in the literature (distyrylbenezene (DSB) derivative 6) for which the van’t Hoff plot is not linear, having distinctly different \(\Delta H\) and \(\Delta S\) values at high and low temperatures.\(^\text{22}\)

![Image of molecular structures](image)

We wanted to explore a larger number of compounds to see whether another example could be found that would behave in a similarly anomalous manner. The data for the enthalpic and entropic differences obtained can in any case be compared to those derived from the van’t Hoff plots of earlier studies. These enthalpy and entropy differences between the two conformers, calculated from the literature data for the examples previously reported, are presented in Table 1.

| Table 1 Enthalpy and entropy differences for different compound classes |
|---------------------------------------------|-------------|
|                                         | \(\Delta H\) | \(\Delta S\) |
| Stilbene 3\(^\text{20}\)          | 4.1          | −0.5        |
| Azobenzene 4\(^\text{20}\)        | 2.4          | −3.2        |
| 4,4′-Dimethyl-benzylideneaniline 5\(^\text{21}\) | 1.7        | +3.0        |
| 2,4-Dimethoxy-4′nitro DSB 6 (<291 K)\(^\text{22}\) | 7.1        | +16.0       |
| 2,4-Dimethoxy-4′nitro DSB 6 (>291 K)\(^\text{22}\) | 14.9       | +43.1       |

We find that the disorder in 1 is dynamic, with the equilibrium frozen in by cooling to 100 K at 4 K min\(^{-1}\). Upon stepwise reheating it thaws out again at a temperature of around 120 K. At temperatures above 120 K (i.e. \(T^{-1} < 0.00833\) K\(^{-1}\)), the van’t Hoff plot (Fig. 2, blue markers) for 1 shows a single straight line, which indicates that the enthalpic and entropic differences between the conformers remain unchanged throughout the entire temperature trajectory. Since the slope of the van’t Hoff plot is given by \(-\Delta H/\Delta S\) and the intersection with the y-axis yields \(\Delta S/\Delta R\), it is easy to calculate these quantities from the linear regression line (see Table 2 below).

For 2, the crystal structure was determined from room temperature data, and the crystal was also quickly cooled to 100 K. The decrease in the population of the minor conformer on going from 293 K to 100 K was slight. Nevertheless, cooling of the crystal to 90 K at 4 K min\(^{-1}\), and then recording X-ray diffraction data upon reheating it in 50 K steps also afforded a linear van’t Hoff plot (Fig. 2, red markers). This structure thaws out around 200 K. In the same way as for 1, the enthalpic and entropic differences appear to remain constant throughout the temperature range (Table 2).

#### Crystal packing

In the crystal, compound 1 forms layers of herringbone-packed molecules with their long axis oriented along the \(a\)-direction and connected to the next layer by type II halogen interactions,\(^\text{21}\) an arrangement that forms a double layer predominantly comprising bromine atoms, perpendicular to the \(a\) axis. The packing of 2 is different, entailing layers perpendicular to the \(c\) direction, which in turn consist of columns of molecules, which are oriented in the \(b\) direction, and parallel displaced along their long axis. Alternating layers perpendicular to \(c\) have the long axis of the molecule oriented at an angle of nearly 90°. The overall involvement of the bromine atoms in the crystal packing is more or less the same as in 1: perpendicular to the \(c\) direction bromine-rich layers are held together by type II halogen interactions.

#### Influence of crystal packing on thermodynamic parameters

The effect of the crystal packing on the pedal motion can be seen in the slope of the van’t Hoff plots, where we find that the enthalpic difference between the two conformers of 2 is much larger than that of 1. In order to explain this
observations and to chart the intermolecular interaction energies in the crystal, we performed computationally extremely cheap Atom–Atom Coulomb–London–Pauli (AA-CLP) calculations using the CLP program\textsuperscript{24} for both conformers.\textsuperscript{25}

Two approaches were used, the first one by quite simply increasing the population of the minor and major conformer respectively to 100% in the CIF and removing the other conformer. The second approach is somewhat more refined, and involved placing the minor conformer in a supercell so it would be surrounded by only molecules of the major conformer. The results for this minor disorder component can then be directly compared to a CLP calculation on the structure containing major conformer only. These latter calculations confirm that the main stabilizing interactions for 1 are between parallel displaced molecules (44.8 kJ mol\textsuperscript{-1}), followed by molecules perpendicular to each other in the herringbone pattern (25.6 kJ mol\textsuperscript{-1}). Between the herringbone packed layers perpendicular to the a-axis, pairwise interaction energies between the molecules are much lower at 10.4 kJ mol\textsuperscript{-1} and are mainly due to type II halogen interactions. In 2, a similar picture emerges—the largest stabilization is between parallel displaced molecules (49.9 kJ mol\textsuperscript{-1}), followed by molecules interacting mainly via C–H⋯N interactions between the layers (15.7 kJ mol\textsuperscript{-1}) and, finally, the end-to-end contacts with type II halogen interactions contribute an additional 11.2 kJ mol\textsuperscript{-1}. All intermolecular interactions in the two structures are dominated by the dispersion term.

The resulting total calculated intermolecular stabilization energies can be found in Table 3.

It is clear from these numbers that compound 1 displays a much larger energy difference between the two conformers in the crystal, where the values for 1 (17.8 vs. 9.6 kJ mol\textsuperscript{-1}) are, for both models, somewhat more than double those of 2 (7.1 vs. 3.5 kJ mol\textsuperscript{-1}). These results of the calculations are in accordance with the relative size of the experimentally observed enthalpy differences between the conformers ($\Delta H_{\text{obs}}$ in Table 3), with the values for the second approach, quite reasonably, also being closer in value to the experimental observations. That the observed values themselves are smaller than the calculated ones is also reasonable, since the molecules in the matrix used for the calculation are unable to adapt their coordinates to the presence of a molecule of minor conformer.

In addition, the calculations clearly pinpoint the differences between the high and low occupancy conformers in the crystal structures. Discussion of the relevant calculated energies will be done for the low occupancy conformer in a matrix of high occupancy conformer only, but essentially the conclusions are identical for the simpler model in which all molecules are replaced with low occupancy conformer. For 2, there is no particularly striking change in the intermolecular interactions on switching conformers—they all become somewhat less attractive in the dispersion term, and/or a bit more repulsive in the repulsion term. This is most obvious for symmetry equivalents $-x, y + 1/2, -z + 1/2$, and $-x, y - 1/2, -z + 1/2$, for which the dispersion term decreases most. For 1, however, there are two clear culprits for the much larger difference in intermolecular energy in the crystal for high and low occupancy conformers: the interactions between molecules at $x, y, z$ and $x + 1, y, z$ (the end-to-end type II halogen interaction) become less attractive by both coulomb interactions and dispersion (20.3 kJ mol\textsuperscript{-1}) and at the same time become less repulsive by only $-14$ kJ mol\textsuperscript{-1}.

Fig. 3 clearly shows why: as the $-\text{CH}_2\text{Br}$ substituent is oriented meta with respect to the pedaling N==N fragment, there is a sizeable reduction in distance between two opposite $-\text{CH}_2\text{Br}$ substituents upon switching to the minor conformer, which in turn causes considerable strain in the structure.

At the same time, the interaction between molecules $x, y, z$ and $x + 1, y + 1, z$ becomes on the one hand $-32.1$ kJ mol\textsuperscript{-1} more attractive by coulomb interactions and dispersion, but on the other hand more repulsive by 36.4 kJ mol\textsuperscript{-1}.

This is a clear demonstration of why geometric arguments in discussing crystal structures can be misleading. Based on geometry, a contact below the van der Waals radius appears between the molecules along $b$, and this would usually be interpreted as attractive. Some relevant distances are shown in Fig. 4. From the intermolecular energies, we know, however, that in this case the close contact is serendipitous and repulsive.

**Activation energy**

A different effect of the crystal structure on the pedal motion can be found in its activation energy, which is jointly determined by the electronic energy of a single molecule, and by the intermolecular interactions with the rest of the crystal.\textsuperscript{26}

| Table 2 Enthalpy and entropy differences for compounds 1 and 2 |
|---|---|---|
| $\Delta H$ (kJ mol\textsuperscript{-1}) | $\Delta S$ (J K\textsuperscript{-1} mol\textsuperscript{-1}) |
| 1 | 4.3 | +6.8 |
| 2 | 2.6 | +2.6 |
This activation energy determines at what temperature the pedal motion thaws out, which can be observed in the van’t Hoff plots (Fig. 2) as the end of the horizontal plateau at low temperatures. Naively, one would expect a low activation energy to be determined by electronic energies only, with very little interference from the surrounding structure – but the picture is more nuanced than that, as will be seen below.

In order to quantify the effect of the crystal packing on the activation energy, we calculated non-equilibrium gas phase electronic energies for 1 and 2, where all bond distances, angles and torsion angles were relaxed, except the torsion angles around the central N-N bond, which were given fixed values of $0^\circ$ to $180^\circ$ with $30^\circ$ increments.

The calculated energy profiles are very similar for both compounds, and are almost perfectly symmetrical around the planar conformations. The energies for all calculated points can be found in the ESI.$^\dagger$ Energies for the respective low and high occupancy conformers of 1 and 2 at $0^\circ$ and $180^\circ$ are almost identical, and the conformations with the central double bond at $90^\circ$ have also quasi-identical relative energies of approximately $+51 \text{ kJ mol}^{-1}$ for both compounds. In order to determine the (de)stabilization of the transition state through interactions with the rest of the crystal structure, we continued our computationally cheap approach of using the CLP program, and generated coordinates for a transition-state-like molecule. This geometry is the average of the geometries of the major and minor conformer (which yields reasonable

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Calculated intermolecular energies by the AA-CLP method for compounds 1 and 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta_{E_{\text{int,calc}}} \text{ [kJ mol}^{-1}\text{]}$</td>
<td>Low occupancy conformer</td>
</tr>
<tr>
<td>1</td>
<td>$-116.4$</td>
</tr>
<tr>
<td>2</td>
<td>$-132.8$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$\Delta E_{\text{int,calc}} \text{ [kJ mol}^{-1}\text{]}$</th>
<th>$\Delta H_{\text{obs}} \text{ [kJ mol}^{-1}\text{]}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>17.8</td>
</tr>
<tr>
<td>2</td>
<td>7.1</td>
</tr>
</tbody>
</table>

Experimental

Synthesis

(1E)-1,2-Bis[3-(bromomethyl)phenyl]-diazene 1. Prepared as described previously.$^7$ Mp: 146–147 °C (lit. 141–143). $R_f = 0.41 (\text{CH}_2\text{Cl}_2/\text{cyclohexane}, 1:2)$. $\delta_{\text{H}}$ (200 MHz, CDCl$_3$, TMS): 4.59 (s, 4H), 7.47–7.56 (m, 4H), 7.85–7.90 (m, 2H), 7.95–7.96 (m, 2H). MS (EI, 70 eV): $m/z$ (%): 366 (70) [M$^+$], 287 (20), 197 (50), 169 (100).

(1E)-1,2-Bis[4-(bromomethyl)phenyl]-diazene 2. Prepared as described previously.$^7$ Mp: 219–225 °C (decomp.) (lit. 217–218). $R_f = 0.34 (\text{CH}_2\text{Cl}_2/\text{cyclohexane}, 1:2)$. $\delta_{\text{H}}$ (200 MHz, CDCl$_3$, TMS): 4.56 (s, 4H), 7.52–7.56 (m, 4H), 7.88–7.92 (m, 2H), 7.92–7.96 (m, 2H).
Table 4 Calculated activation energies (ΔE_{act,calc}) by summing gas phase electronic energies (ΔE_{gas,calc}) with CLP molecular energies in the crystal (ΔE_{m,CLP}) for ground and pseudo-transition states. All energies in kJ mol⁻¹.

<table>
<thead>
<tr>
<th>Angle</th>
<th>0°</th>
<th>90°</th>
<th>180°</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔE_{m,CLP}</td>
<td>-124.6</td>
<td>-134.3</td>
<td>-134.2</td>
</tr>
<tr>
<td>ΔE_{electr}</td>
<td>0</td>
<td>50.9</td>
<td>0</td>
</tr>
<tr>
<td>ΔE_{total}</td>
<td>-124.6</td>
<td>-83.4</td>
<td>-134.2</td>
</tr>
<tr>
<td>ΔE_{act,calc}</td>
<td>41.2</td>
<td>50.8</td>
<td></td>
</tr>
</tbody>
</table>

4H). MS (EI, 70 eV): m/z (I%): 366 (60) [M⁺], 287 (100), 208 (90), 197 (20), 169 (50).

X-ray diffraction

Data sets were collected on a Bruker platform goniometer with Smart Apex detector, using Mo Kα radiation (0.71073 Å) from a sealed tube, monochromated with pyrolithic graphite, and collimated with a pinhole collimator. Crystals were mounted on Mitegen micromesh mounts with a trace of glue and cooled or heated with an Oxford Cryostream cryostat. Crystals were mounted on Mitegen micromesh mounts with a trace of glue and cooled or heated with an Oxford Cryostream cryostat.

Temperature was ramped down from room temperature to 90(2) K at a speed of 4 K min⁻¹ and data collections were carried out on heating in 30 K intervals, with a final dataset at 375(2) K for 1. Compound 2 was cooled at 4 K min⁻¹ to 90(2) K, and then reheated in 50 K intervals, with a final dataset also at 375(2) K. Data were reduced with the Bruker APEX2 software suite and scaled and corrected for absorption effects using SADABS-2008/1. The structures were solved by direct methods with the help of SHELXS97 and refined using SHELXL2013/2, SHELXL2014/7 and shelXle. A model was constructed for each of the structures in which the 1,2- and 1,3-distances for the minor conformer were restrained to those in the major conformer. In addition, the ADPs of corresponding atoms were kept identical. These initial models were used for the refinements at all temperatures, in order to ensure comparability between the resulting populations of the disorder. H-atoms were refined as riding. The experimental data on the crystallographic experiments in this paper can be found in Tables S1 and S2 in the ESI.

Acknowledgements

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Notes and references

25 The gas phase electronic energies of both conformers for both compounds are equal up to the accuracy of DFT calculations at the B3LYP/6-31+G* level (\(<0.25 \text{kJ mol}^{-1}\)).