### Conformational analysis of biphenyls: an upside-down view

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One of the (not a few) pleasures of working in a university is the occasional opportunity that arises to give a new lecture course to students. New is not quite the correct word, since the topic I have acquired is **Conformational analysis**. The original course at Imperial College was delivered by Derek Barton himself about 50 years ago (for articles

written by him on the topic, see DOI 10.1126/science.169.3945.539 or the original 10.1039/QR9561000044), and so I have had an opportunity to see how the topic has evolved since then, and perhaps apply some quantitative quantum mechanical interpretations unavailable to Barton himself.

The example I have chosen to focus on here is biphenyl (a derivative of which also happens to be the first structure shown by Barton in his 1970 Science article noted above), but modified with isoelectronic B/N substitution for carbon for a particular reason.



Four hydrogen atoms are

highlighted in the above drawings by virtue of how close they might approach each-other, and what impact this will have on the conformation of each species. Such close approaches are normally defined with reference to the so-called van der Waals radius of the element concerned. For hydrogen, this radius is either 1.2Å (if the contact is to another hydrogen) or 1.1Å (if its to any other element, see DOI: 10.1021/jp8111556). An interpretation of this value is that the van der Waals attraction due to to dispersion or long range correlation effects reaches a maximum for two non-bonded hydrogen atoms at

~2.4Å. Significantly, a slightly closer approach than this value might still be mildly attractive, but it would be generally agreed that any distance less than ~2.1Å now represents a genuine repulsion between the hydrogens (see also this post). This represents a somewhat more quantitative judgement on what used to be called *steric interactions*.

With the scene set, let me introduce the results of a calculation (wB97XD/6-31G(d,p), a DFT method selected because it treats the long range correlation effects with a specific correction)



Conformational analysis of biphenyl 1

One can see here minima at ~45, 135, 225 and 315° for**1** (see DOI 10042/to-4853). Due to symmetry, the first and last are identical as are 2nd and 3rd, and the 1st and 2nd minima are in fact enantiomers of each other (the symmetry is D2, which is chiral). Two different transition states connect these minima, one with angles of 0/180 and the other slightly lower energy at 90/270°.

The non-bonded H...H distance are as follows:  $1.95\text{\AA}(00^\circ, 2.39\text{\AA}(045^\circ)$  and  $3.54\text{\AA}(090^\circ)$ . We may conclude that the first of these is repulsive, the second attractive and the third non interacting. Counterbalancing this effect is of course resonance due to  $\pi$ - $\pi$ -overlaps across the central bond, which decreases to zero as the angle moves to  $90^\circ$ . The conformational minimum  $(045^\circ)$  is such because of the maximal H...H dispersion attraction and the still significant  $\pi$ - $\pi$ -overlap. This brief analysis suggests however that these two effects are finely balanced, and so the next question is whether one might be able to perturb the system to distort the balance. The perturbation chosen is to replace one or two pairs of carbon atoms with the iso-electronic combination B+N.

The first perturbation is to replace the central rotating bond by a B-N combination **2** (DOI: 10042/to-4854).



Rotation about the B-N bond in 2

For this species, the H...H distances are 2.02Å@0°, 2.36Å@45° and 3.61Å@90°, the only significant difference with **1** emerging as the 0° conformation being around 1 kcal/mol lower relative to the other two. It is tempting to attribute this to the longer H...H separation for this rotamer in **2** due to the B-N bond being longer (1.562Å) than the C-C bond it replaced (1.496Å)

The next perturbation is to relocate the N/B pair as in 3 (DOI: 10042/to-4855). If one imagines that this will be a minor perturbation, take a look at the profile below.



Rotation about central C-C bond in 3.

The world has been turned upside down. What were transition states @0° and @180° are now minima and the reason is easy to find. The central C-C bond is now only 1.400Å long, having acquired substantial double bond character, and being accordingly very much more difficult to twist (the barrier being ~30 kcal/mol). The  $\pi$ - $\pi$ -overlap has won out completely, and in the process has forced the H...H distance down to a presumably repulsive 1.918Å. The penalty for this is that the overall energy of **3** is some 22.8 kcal/mol higher than **2**.

Added in proof (as the expression goes): If the above profile is conducted with full geometry optimization in a solvent field (water), which helps stabilise charge separations, the profile changes to the below. The solvation reduces the barrier to rotation considerably, the energy maxima now reveal a proper stationary point (rather than the cusp), the minima are very slightly non-planar, but the basic inversion of the potential energy surface compared to 1 or 2 is still observed.



Rotation about the C-C bond for 3, with solvation correction

The final perturbation is 4 (DOI: 10042/to-4856) with the following rotational profile. Another surprise:



Rotation about the central C-C bond in 4.

The H...H distances are  $1.930\text{ Å}@0^\circ$ ,  $1.789/2.275\text{ Å}@180^\circ$ . The difference from **1** is that the hydrogens now have opposite polarity for the N-H (which is positive) and the B-H (which is negative). At the rotation angle of  $0^\circ$ , two H(+)...(-)H style dihydrogen bonds (see also this post) are established (these are presumed to be very attractive); at an angle of  $180^\circ$ , the H(+)...(+)H and H(-)...(-)H interactions are presumed to be very repulsive. The difference between the two is ~18 kcal/mol.

We have learnt that conformational analysis for molecules such as these is a fight between  $\pi$ - $\pi$ -overlaps, which themselves can have unexpected outcomes, weak van der Waals dispersion interactions between "neutral" non-bonded hydrogen atoms, and strong electrostatic attractions and repulsions between "ionic" hydrogens. Now perhaps the reason for the choice of the wB97XD DFT method can be seen; it is capable (at least in theory) of balancing these forces properly.

So the world of conformational analysis can be turned upside down, and analysing what happens from this topsy-turvy viewpoint can teach a lot!