Sizing the role of London dispersion in the dissociation of all-meta tert-butyl hexapheny lethane†

Sören Rösel,a Ciro Balestrieriib and Peter R. Schreinera

The structure and dynamics of enigmatic hexa(3,5-di-tert-butylphenylethane) was characterized via NMR spectroscopy for the first time. Our variable temperature NMR analysis demonstrates an enthalpy–entropy compensation that results in a vanishingly low dissociation energy ($\Delta G_{\text{d}}^{298} = -1.60(6)$ kcal mol$^{-1}$). An in silico study of increasingly larger all-meta alkyl substituted hexapheny lethane derivatives (Me, tPr, tBu, Cy, 1-Ad) reveals a non-intuitive correlation between increased dimer stability with increasing steric crowding. This stabilization originates from London dispersion as expressed through the increasing polarizability of the alkyl substituents. Substitution with conformationally flexible hydrocarbon moieties, e.g., cyclohexyl, introduces large unfavourable entropy contributions. Therefore, using rigid alkyl groups like tert-butyl or adamantyl as dispersion energy donors (DED) is essential to help stabilize extraordinary bonding situations.

Hexapheny lethane (HPE; H-1) was long assumed to have been synthesized by Gomberg in 1900 in the context of preparing the first organic free radical, the triphenylmethyl radical (H-1·) (Scheme 1). However, it was shown in 1968 that H-1· rather corresponded to quinoid H-2. While steric repulsion between the phenyl moieties was readily made responsible for instability of H-1·, the introduction of bulky tert-butyl groups in all meta positions, which should lead to significantly higher steric crowding, resulted in isolable hexa(3,5-di-tert-butylphenyl)ethane (tBu-1) that could even be characterized through X-ray crystal diffraction. Clearly, this questions our (conceptual) understanding of the balance of steric repulsion vs. noncovalent attraction, and a dispersion energy donor (DED) concept employing large alkyl groups may be invoked to rationalize such surprising bonding situations. The classic HPE system therefore seems well suited to probe this concept. Here we report on the challenging re-synthesis and the first NMR-spectroscopic characterization of tBu-1 as well on density functional theory (DFT) computations to gauge the fine balance between Pauli repulsion and London dispersion (LD) attraction.

It is quite remarkable that the structural proposal for H-1· prevailed for almost 70 years because Gomberg himself noted that “Der Körper zeigt in höchstem Maasse die Eigenschaften einer ungesättigten Verbindung” (the substance shows pronounced properties of unsaturation) and readily adds oxygen and halogens. He concluded: “Die oben mitgetheilten experimentellen Ergebnisse zwingen mich zu der Annahme, dass in dem ‘ungesättigten Kohlenwasserstoff’ das Radical Triphenylmethyl vorliegt” (the disclosed experimental findings force me to assume that the “unsaturated hydrocarbon” presents the radical triphenylmethyl). In addition, the neat colourless hydrocarbon was found to be in equilibrium with a coloured form in solution. This triggered an intense discussion regarding the proper structure of triphenylmethyl where, apart from the highly symmetric HPE structure, quinoid structures H-2 and H-3 were suggested (Scheme 1). In an

Scheme 1 The Jacobsen–Nauta structure H-2 is exclusively observed for the parent triaryl radical H-1. All-meta tert-butyl substitution leads to equilibration between the tBu-1· and dimeric tBu-1 in solution.

† Electronic supplementary information (ESI) available: Experimental procedures, spectra and computational details. See DOI: 10.1039/c6sc02727j
early review Gomberg described an equilibrium between an associated and a free radical, the monoquinoidal structure, and HPE.\textsuperscript{12} In the following years an equilibrium between H-1· and H-1\textsuperscript{+} suggested first by Flurscheim\textsuperscript{13,14} was gradually accepted and the quinoidal structures vanished from the discussion, despite Gomberg mentioning later that a quinoid tautomer of H-1· causes the yellow colour of the solution.\textsuperscript{15}

Notwithstanding the fact that some had misgivings regarding the correct structure of HPE,\textsuperscript{16} dimer H-1\textsubscript{2} was generally accepted until Nauta et al. reinvestigated it utilizing 3\textsuperscript{1}H-NMR and IR spectroscopies. These studies demonstrated that HPE adopted Jacobsen’s monoquinoidal dimer H-2.\textsuperscript{3} The first computational investigation of H-1\textsubscript{2} was reported by Mislow et al. utilizing empirical force field (EFF) computations that revealed a very long central R\textsubscript{CC} of 1.64 Å in H-1\textsubscript{2} due to the repulsion between the trityl moieties.\textsuperscript{7} The D\textsubscript{3} symmetric structure was found to be 2.6 kcal mol\textsuperscript{−1} more stable than the S\textsubscript{6} structure. A QM/MM (ONIOM) study of 2002 favoured the S\textsubscript{6} form and gave a BDE = 16.6 kcal mol\textsuperscript{−1} for H-1\textsubscript{1} and a central bond length of 1.72 Å.\textsuperscript{18}

The first isolation of an unbridged HPE derivative was achieved by Rieker in 1978 through the introduction of tert-butyl groups in all 2- and 6-positions onto the Schlenk (tri(4-biphenyl)-methyl) radical.\textsuperscript{4,5} The previously reported introduction of tert-Bu groups onto the trityl radical H-1· yielded tris[3,5-di-tert-butylphenyl]methyl (‘Bu-1·) that was found to be monomeric in benzene\textsuperscript{19} and Rieker concluded that […] steric hindering of formation of the Jacobson–Nauta structure [R-2] by incorporation of bulky groups [in meta and/or para positions], [R-1·] must then either be persistent as the monomer—or dimerize to [R-1,2].\textsuperscript{6,4} Indeed, “poor, colourless crystals” could be grown from the orange cyclohexane solution consisting of the first unbridged HPE derivative 1,1,1,2,2,2-hexakis(2,6-di-tert-butyl-1,1’-biphenyl)-1-yl)ethane (‘Bu-1\textsubscript{2}).\textsuperscript{4,5} The dimer crystallized in S\textsubscript{6} symmetry with an astonishingly short experimental R\textsubscript{CC} of only 1.47 Å. Mislow responded with improved EFF and MNDO SCF-MO computations that gave a R\textsubscript{CC} between 1.62 and 1.68 Å.\textsuperscript{20} Finally, a central bond length of 1.67 Å was determined from a new crystal structure of ‘Bu-1\textsubscript{2};’ magic angle spinning (MAS) nutation experiments gave R\textsubscript{CC} = 1.64–1.65 Å,\textsuperscript{21} confirming the earlier bond distance and refuting Rieker’s structure. Our recent computational reinvestigation of the “HPE riddle” with the modern DFT implementations confirmed these results.\textsuperscript{22}

![Scheme 2](image)

Scheme 2 Synthesis of ‘Bu-1·: (a) ‘BuCl, AlCl\textsubscript{3} (–40) → (–15) °C, 81% (b) Br\textsubscript{2}, Fe, CCl\textsubscript{4}, rt, 86% (c) ‘BuLi, Et\textsubscript{2}O, –78 °C → rt 2. (Et\textsubscript{2}O)\textsubscript{2}CO, 78% (d) AcCl, n-hexane, 95% (e) Zn(Cu), C\textsubscript{6}D\textsubscript{6} or C\textsubscript{6}D\textsubscript{12}, rt (f) only in C\textsubscript{6}D\textsubscript{12}.

Both H-2 and ‘Bu-1\textsubscript{2} afford yellowish-orange, EPR active solutions, in line with very low computed dissociation energies.\textsuperscript{22} The equilibrium constants for radical formation form the dimers should therefore be accessible by variable temperature NMR spectroscopy despite Rieker’s notion that “Because of the high degree of dissociation and the sensitivity of the radical towards traces of oxygen no definite proof of the structure of [‘Bu-1·] can be obtained with 1\textsuperscript{H}-NMR, 13\textsuperscript{C}-NMR, and mass spectroscopy, […]”.\textsuperscript{4} The stronger magnetic fields available with current NMR spectrometers together with sophisticated techniques encouraged us to investigate the dissociation/association equilibrium of ‘Bu-1·.

The precursor tris(3,5-di-tert-butylphenyl)methyl chloride (5) was synthesized from benzene (6) via Friedel–Crafts alkylation with ‘BuCl, ipso-bromination followed by lithiation at −78 °C, subsequent triple addition to diethylcarbonate and chlorination of the central methyl carbon with acetyl chloride (Scheme 2). The classic reduction of 5 to ‘Bu-1· with silver in benzene\textsuperscript{21,22} under
meticulous exclusion of oxygen was slow and mainly yielded bis(tri(3,5-di-tert-butylphenyl)methyl) peroxide (7). Even silver derived from reduction of silver nitrate seemed to introduce a significant amount of oxygen. Using the zinc–copper couple instead gave rapid quantitative conversion. The reduction in benzene-\(d_6\) resulted in an orange solution with a \(^1\)H-NMR spectrum that contained, besides several sharp signals, a very broad peak at \(\delta = 1.88\) ppm with a total width of 3 ppm (Fig. 1a, top). The sharp signals were assigned to 7, tris(3,5-di-tert-butylphenyl)methane (8) and tris(3,5-di-tert-butylphenyl)methanol (9) by comparison with the NMR spectra of the pure compounds (cf. ESI†). The sample gave an EPR signal featuring the known spin pattern.\(^{19,24}\) This, together with the known NMR lifetime broadening of organic radicals\(^{25}\) as well as the \(r^{-6}\) line width dependence of the NMR signal with respect to the radical centroid in the molecule,\(^{26}\) made it possible to identify the broad peak as the tert-butyl group hydrogen resonance of \(^{1}\)Bu-1.

The \textit{in situ} radical generation in cyclohexane-\(d_{12}\) through reduction with Zn(Cu) is slower as compared to benzene-\(d_6\) but gave the same orange solution and a similar \(^1\)H-NMR spectrum with five additional peaks at \(\delta = 7.75, 7.19, 6.02, 0.98\) and 0.95 ppm in a ratio of 1 : 1 : 1 : 9 : 9, respectively (Fig. 1a, middle). This indicates a highly symmetric structure that would be incompatible with a less symmetric quinoid structure for which an allylic resonance at 5 ppm and a vinylic resonance at 0.95 ppm in a ratio of 1 : 1 : 1 : 9 : 9, respectively (Fig. 1a, bottom). This first \(^1\)H-NMR spectrum of an unbridged HPE derivative features a significant chemically inequivalence of the \textit{ortho} or \textit{meta} phenyl nuclei (for the \(^{13}\)C-NMR spectrum see the ESI†). The fast inversion of the local helical chirality of an individual trityl group through a two-ring flip mechanism\(^{29,30}\) that makes the two \textit{ortho} or \textit{meta} nuclei appear as a single signal at \(rt\) is blocked in \(^{1}\)Bu-1\(_2\). That is, one edge of the phenyl ring points outside and the other inside towards the \(S_n\) axis. While the phenyl groups in the X-ray crystal structure are indistinguishable due to the superposition of four disordered \(^{1}\)Bu-1\(_2\) arrangements,\(^6\) the computed structure (Fig. 1b) nicely features this structural difference between inside (o') and outside (o) protons (Fig. 1a, bottom). Fortunately, the equilibrium between the \(^{1}\)Bu-1\(_2\) radical and \(^{1}\)Bu-1 revealed a strong temperature dependence (Fig. 2a), allowing us to determine parameters to calculate the equilibrium constant \(K\) at a particular temperature through integration of

\[
K = \frac{x_1^2}{x_1} = \frac{4I_t^2}{I_x(I_x + 2I_t)}
\]

with \(I\) being the \(^1\)H-NMR integrals, \(x\) the molar fraction, \(1\) the radical monomer and \(12\) the dimer (for the derivation see the ESI†).

The van’t Hoff plot (Fig. 2b) shows a good linear correlation and \(\Delta G_{298}^{\text{ff}} = -1.60(6)\) kcal mol\(^{-1}\) for the homolysis of \(^{1}\)Bu-1 into two radicals reveals a very low dissociation energy. This \(\Delta G_{298}^{\text{ff}}\) value agrees reasonably well with computational predic-
tions of \(\Delta G_{298}^{\text{ff}} = -3\) to +1 kcal mol\(^{-1}\) in cyclohexane with PWPB95-D3/TZV(2d,2p),\(^{32,33}\) and it is in the range of the weakest experimentally determined \(\Delta G_{298}^{\text{ff}}\) of \(-0.2(1)\) kcal mol\(^{-1}\) for a C–C single bond reported for the 2,6-di-tert-butyl-4-methoxyphenoxyl dimer (12, Fig. 3).\(^{34}\) As found for 12, \(\Delta H_{298}^{\text{ff}}\) of 6.1(5) kcal mol\(^{-1}\) is outbalanced by \(\Delta S_{298}^{\text{ff}} = 6.3\) kcal mol\(^{-1}\) (298 K). We find a similar enthalpy–entropy compensation for \(^{1}\)Bu-1, with \(\Delta H_{298}^{\text{ff}} = 7.9(4)\) kcal mol\(^{-1}\) and \(\Delta S_{298}^{\text{ff}} = 9.5(3)\) kcal mol\(^{-1}\) (298 K); this compares well with other weakly bonded hydrocarbons (cf. ESI†).

The dissociation enthalpy of \(^{1}\)Bu-1\(_2\) is 2.8 kcal mol\(^{-1}\) smaller than the 10.7(2) kcal mol\(^{-1}\) \(\Delta H_{298}^{\text{ff}}\) of H-2,\(^{32,33}\) while the entropy contribution in \(^{1}\)Bu-1\(_2\) is significantly larger (\(\Delta S_{298}^{\text{ff}} = 6.0\) kcal mol\(^{-1}\)). This is not surprising because covalent \(^{1}\)Bu-1\(_2\) requires much more ordering of the \(^{1}\)Bu-groups for optimizing the LD interactions, leading to a large, counteracting \(\Delta S\) contribution. This is likely to be a general phenomenon for structures with large LD contributions because the DEDs often require optimal structural alignment for maximizing the LD interactions.

Differences in LD contributions between gas phase computations and solution experiments were rationalized with a di-n-
hexyl substituted Wilcox-type balance by Cockroft and co-workers.\textsuperscript{14,15} Another approach outlined here is the use of the radical dimerization energies to determine the effect of DEDs on chemical equilibrium. While large, polarizable electron donor substituents stabilize the dimers and the free radicals to different degrees, the radicals are stabilized mostly by conjugation. Depending on which radical stability scale one uses, \textit{meta} alkyl substitution of benzyl radicals may be slightly stabilizing\textsuperscript{46-48} or destabilizing,\textsuperscript{49,50} but it is small in all cases. That is, the increased stability of \textit{t}Bu-1 relative to parent H-1\textsubscript{2} must originate from the LD contributions of the \textit{t}Bu groups.

Does the common measure for bond strengths, the bond dissociation energy (BDE), still apply here? BDEs were initially computed in the gas phase (13.7 kcal mol\textsuperscript{−1}) and the experimental \( \Delta G_d \textsuperscript{298} \) value measured at potentials caused by the rotations 

\[
\Delta G_d \textsuperscript{298} \approx \text{−1.6 kcal mol}^{-1}
\]

can be taken as a rough measure of the attenuation of the LD interactions in solution. As the corrections to account for LD effects are large (ca. 60 kcal mol\textsuperscript{−1}), a significant LD contribution must therefore persist also in solution. In light of such large LD contributions (cf. Table S3\textsuperscript{†}), the BDE does not seem to be a good measure for the covalent bond strength of the central bond.

As large-scale computations including properly computed entropy terms can reproduce \( \Delta G_d \textsuperscript{298} \) reasonably well,\textsuperscript{22} we used more approximate DFT computations to make some qualitative predictions regarding other DEDs in R-1\textsubscript{2} as model (Fig. 4). We employed the commonly used alkyl groups Me, \textit{t}Pr, \textit{t}Bu, Cy, and 1-Ad as DEDs. With up to 356 atoms in Ad-1\textsubscript{2}, only DFT methods are currently feasible; these computations still take a very long time because of the very flat potentials caused by the rotations of these groups. The B3LYP\textsuperscript{45,46} functional with the Becke–Johnson damped dispersion correction D3(BJ) introduced by Grimme\textsuperscript{44,45} as well as the hybrid GGA functional M06-2X\textsuperscript{46} of Truhlar et al. were used in conjunction with a cc-pVDZ basis set (Table S3\textsuperscript{†}). In the HPE derivatives the symmetry was restricted to the \( S_6 \) point group (as experimentally observed for \textit{t}Bu-1\textsubscript{2}) while the radical monomers \textit{t}Bu-1\textsubscript{2} were assumed to have \( C_1 \) symmetry. The C-PCM model was employed to account for bulk solvent effects.\textsuperscript{48}

First of all, the B3LYP-D3(BJ) and M06-2X results with a cc-pVDZ basis set show the same general trends but diverge significantly as the size of the DED increases (Table S3\textsuperscript{†}). The M06-2X functional is in good agreement for \textit{t}Bu-1\textsubscript{2} but the trend is not as smooth as with B3LYP-D3(BJ).\textsuperscript{48} Still, the LD contributions are not merely minor corrections but are quite large, exceeding the magnitude of carbon–carbon bond energies for \( R = 1\text{-Ad} \) (cf. ESI\textsuperscript{†}). There is a large dispersion correction even in the hitherto unobserved parent system H-1\textsubscript{2} that is predicted to dissociate with a \( \Delta G_d \textsuperscript{298} \) of less than \( −5 \) kcal mol\textsuperscript{−1}.

Non-covalent interactions can be visualized by plotting the reduced density gradient in regions of low electron density (NCI plot).\textsuperscript{30} This enables a qualitative analysis of the balance between repulsive and attractive contact areas. The NCI plots (Fig. 5) of H-1\textsubscript{2}, \textit{t}Bu-1\textsubscript{2}, and Ad-1\textsubscript{2} visually support the general trend obtained for \( \Delta G_d \textsuperscript{298} \). Parent H-1\textsubscript{2} features repulsive (red) areas between phenyl moieties within and between the trityl.

![Fig. 4](image1.png)  
**Fig. 4** The computed free dissociation energies \( \Delta G_d \textsuperscript{298} \) including dispersion corrections (○) correlate very well with the polarizability \( \alpha \) and therefore the LD contributions of the R groups \( (R = \text{H, Me, } \text{iPr, } \text{tBu, Cy, Ad}) \) and gain stability with size. Neglecting LD leads to negative \( \Delta G_d \textsuperscript{298} \) (●), bearing the opposite, negative trend. Basis set: cc-pVDZ.

![Fig. 5](image2.png)  
**Fig. 5** Non-covalent interaction (NCI) plots (\( s = 0.5 \) au/−0.01 < \( ρ < +0.01 \) au) are depicted separately on the left from the molecular structure on the right for clarity. Repulsion is colour-coded red, ‘strong’ attraction blue and weak interactions in green.
groups and few attractive (C–H⋯π) interactions (blue). Substitution with 1-Bu groups leads to significant growth of both attractive and repulsive regions in the periphery of the HPE core structure but scattered weak interactions (green) also appear. Finally, Ad-1 clearly shows a dense, weakly interacting region between the substituted trityl moieties. Hence, although the 1-adamantane moieties are even larger than the 1-Bu groups, they exhibit a huge noncovalent contact area that maximizes attractive LD interactions visually outweighing the repulsive regions.

Conclusions

We re-synthesized 1-Bu-1,2, a member of the elusive class of HPE derivatives, and were able to obtain its dynamic NMR spectrum in equilibrium with the corresponding free radical 1-Bu-1. The dissociation free energy of ΔGd298° (exp.) = −1.60(6) kcal mol−1 confirms computational pre-dictions and is composed of nearly equally large enthalpy and entropy contributions at 298 K. Computations suggest that substitutions with isopropyl and cyclohexyl DEDs also are accompanied by large unfavourable entropy contributions so that it is imperative to use rigid hydrocarbon moieties as DEDs. As a consequence, the introduction of sterically even more demanding 1-adamantyl substituents predicts an even further reduction the central bond length and a significant increase in stability as compared to 1-Bu-1. Counterintuitively, Ad-1 should therefore be an isolable HPE derivative despite the use of even bulkier groups. Hence, large, highly polarizable and rigid hydrocarbon moieties are the most effective DEDs.

Acknowledgements

We thank Bart Kahr (NY University) for making his PhD thesis available and H. Hausmann (JLU Giessen) for support with the NMR spectroscopic measurements. This work was supported by the Deutsche Forschungsgemeinschaft, Priority Program “Dispersion” (SPP 1807, Schr 597/27-1).

Notes and references

1 M. Gomberg, Ber. Dtsch. Chem. Ges., 1900, 33, 3150–3263.
2 M. Gomberg, J. Am. Chem. Soc., 1900, 22, 757–771.
3 H. Lankamp, W. T. Nauta and C. MacLean, Tetrahedron Lett., 1968, 9, 249–254.
4 M. Stein, W. Winter and A. Rieker, Angew. Chem., Int. Ed. Engl., 1978, 17, 692–694.
5 M. Stein, W. Winter and A. Rieker, Angew. Chem., 1978, 90, 737–738.
6 B. Kahr, D. V. Engen and K. Mislow, J. Am. Chem. Soc., 1986, 108, 8305–8307.
7 S. Grimm, R. Huenerbein and S. Ehrlich, ChemPhysChem, 2011, 12, 1258.
8 J. Schmidlin, Ber. Dtsch. Chem. Ges., 1908, 41, 2471–2479.
9 A. E. Tschitschibabin, Ber. Dtsch. Chem. Ges., 1904, 37, 4709–4715.
10 P. Jacobsen, Ber. Dtsch. Chem. Ges., 1905, 38, 196–199.
11 E. Heintschel, Ber. Dtsch. Chem. Ges., 1903, 36, 320–322.
12 M. Gomberg, J. Am. Chem. Soc., 1914, 36, 1144–1170.
13 B. Flürsheim, J. Appl. Chem., 1905, 1, 497–539.
14 B. Flürsheim, Ber. Dtsch. Chem. Ges., 1908, 41, 2746–2747.
15 M. Gomberg, Chem. Rev., 1924, 1, 91–141.
16 J. M. McBride, Tetrahedron, 1974, 30, 2009–2022.
17 W. D. Hounshell, D. A. Dougherty, J. P. Hummel and K. Mislow, J. Am. Chem. Soc., 1977, 99, 1916–1924.
18 T. Vreven and K. Morokuma, J. Phys. Chem. A, 2002, 106, 6167–6170.
19 K. Schreiner, A. Berndt and F. Baer, Mol. Phys., 1973, 26, 929–939.
20 E. Osawa, Y. Onuki and K. Mislow, J. Am. Chem. Soc., 1981, 103, 7475–7479.
21 N. Yannoni, B. Kahr and K. Mislow, J. Am. Chem. Soc., 1988, 110, 6670–6672.
22 S. Grimme and P. R. Schreiner, Angew. Chem., Int. Ed., 2011, 50, 1263–1264.
23 M. Stein and A. Rieker, Tetrahedron Lett., 1975, 16, 2123–2126.
24 H. Sakurai, H. Umino and H. Sugiyama, J. Am. Chem. Soc., 1980, 102, 6837.
25 K. H. Hausser, H. Brunner and J. C. Jochims, Mol. Phys., 1966, 10, 253–260.
26 F. Rastrelli and A. Bagno, Chem.–Eur. J., 2009, 15, 7990–8004.
27 C. Chen, H. Lee and R. F. Jordan, Organometallics, 2010, 29, 5373–5381.
28 R. J. Kurland, I. I. Schuster and A. K. Colter, J. Am. Chem. Soc., 1965, 87, 2279–2281.
29 D. Gust and K. Mislow, J. Am. Chem. Soc., 1973, 95, 1535–1547.
30 K. Mislow, Acc. Chem. Res., 1976, 9, 26–33.
31 J. M. Wittman, R. Hayoun, W. Kaminsky, M. K. Coggins and J. M. Mayer, J. Am. Chem. Soc., 2013, 135, 12956–12959.
32 W. P. Neumann, W. Uizik and A. K. Zarkadis, J. Am. Chem. Soc., 1986, 108, 3762–3770.
33 N. S. Blom, G. Roelofs and J. A. Kanter, Cryst. Struct. Commun., 1982, 11, 297–304.
34 S. Paliwal, S. Geib and C. S. Wilcox, J. Am. Chem. Soc., 1994, 116, 4497–4498.
35 L. X. Yang, C. Adam, G. S. Nichol and S. L. Cockroft, Nat. Chem., 2013, 5, 1006–1010.
36 X. Creary, Acc. Chem. Res., 2006, 39, 761–771.
37 T. H. Fisher, S. M. Dershem and M. L. Prewitt, J. Org. Chem., 1990, 55, 1040–1043.
38 R. A. Jackson and R. Moosavi, J. Chem. Soc., Perkin Trans. 2, 1992, 885–888.
39 W. Adam, H. M. Harrer, F. Kita, H.-G. Korth and W. M. Nau, J. Org. Chem., 1997, 62, 1419–1426.
40 A. M. D. P. Nicholas and D. R. Arnold, Can. J. Chem., 1986, 64, 270–276.
41 R. M. Parrish, J. F. Gonthier, C. Corminboeuf and C. D. Sherrill, J. Chem. Phys., 2015, 143, 051103/1–5.
42 A. D. Becke, J. Chem. Phys., 1993, 98, 5648–5652.
43 C. T. Lee, W. T. Yang and R. G. Parr, Phys. Rev. B, 1988, 37, 785–789.
44 S. Grimme, S. Ehrlich and L. Goerigk, J. Comput. Chem., 2011, 32, 1456–1465.
45 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, J. Chem. Phys., 2010, 132, 154104–154119.
46 Y. Zhao and D. G. Truhlar, Acc. Chem. Res., 2008, 41, 157–167.
47 T. H. Dunning, J. Chem. Phys., 1989, 90, 1007–1023.
48 V. Barone and M. Cossi, J. Phys. Chem. A, 1998, 102, 1995–2001.
49 S. Grimme, A. Hansen, J. G. Brandenburg and C. Bannwarth, Chem. Rev., 2016, 116, 5105–5154.
50 J. Contreras-García, E. R. Johnson, S. Keinan, R. Chaudret, J.-P. Piquemal, D. N. Beratan and W. Yang, J. Chem. Theory Comput., 2011, 7, 625–632.