Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2022, 13, 4972

dll publication charges for this article have been paid for by the Royal Society of Chemistry

Received 27th October 2021 Accepted 2nd April 2022

DOI: 10.1039/d1sc05936j

rsc li/chemical-science

Isolable fluorinated triphenylmethyl cation salts of $[HCB_{11}Cl_{11}]^-$: demonstration of remarkable hydride affinity†

S. Olivia Gunther, Chun-I Lee, Ellen Song, Nattamai Bhuvanesh and Oleg V. Ozerov **

Significantly fluorinated triarylmethyl cations have long attracted attention as potentially accessible highly reactive carbocations, but their isolation in a convenient form has proved elusive. We show that abstraction of chloride with a cationic silylium reagent leads to the facile formation of di-, tetra-, and hexafluorinated trityl cations, which could be isolated as analytically pure salts with the $[HCB_{11}Cl_{11}]^-$ counterion and are compatible with (halo)arene solvents. The F_6Tr^+ cation carrying six meta-substituents was computationally predicted to possess up to 20% higher hydride affinity than the parent triphenylmethyl cation Tr^+ . We report that indeed F_6Tr^+ displays reactivity unmatched by Tr^+ . F_6Tr^+ at ambient temperature abstracts hydrides from the C-H bonds in tetraethylsilane, mesitylene, methylcyclohexane, and catalyzes Friedel–Crafts alkylation of arenes with ethylene, while Tr^+ does none of these.

Introduction

The triphenylmethyl or trityl cation (Ph₃C⁺ or Tr⁺) is a textbook example of a carbocation that is isolable owing to the high degree of benzylic conjugation and the steric protection afforded to the central carbon by the three phenyl substituents. ¹ In organometallic chemistry and catalysis, salts of Tr⁺ are frequently used to study the thermodynamics and kinetics of hydride transfer,2,3,4 or to generate reactive main-group and transition-metal cations through hydride or alkyl anion abstraction.5,6,7 Tr⁺ can also serve as a convenient one-electron oxidant.8 Trityl cation derivatives bearing stabilizing electrondonating groups can even exist in aqueous solutions, with a rich history of use as indicators and dyes.9 The trityl cation versions bearing electron-withdrawing substituents have proven more challenging to obtain. Fluorinated trityl cations, up to $(C_6F_5)_3C^+$ (A, Fig. 1), have been of particular fundamental interest, 10-12 including as isoelectronic analogs of the widely used borane (C₆F₅)₃B, ^{13,14} and more recently have been studied by Horn and Mayr¹⁵ and Dutton et al. 16 The more reactive A or other ortho- and/or meta-fluorinated trityl cations were not isolated in those studies, but were generated in situ, or their intermediacy was indicated by kinetic studies. While generation

Department of Chemistry, Texas A&M University, 3255 TAMU, College Station, Texas 77842, USA. E-mail: ozerov@chem.tamu.edu

of fluorinated trityl cations in oleum and other superacidic media, 10-12,16 or by *in situ* abstraction of halides with element halide Lewis acids 15 is possible, these media and counteranions are not fully compatible with either the more electron-deficient trityl cations themselves or with their potential use in the synthesis of other reactive main-group or transition metal cations. Thus, the full extent of the reactivity of the fluorinated trityl cations can only be accessed when paired with more robust weakly coordinating anions in weakly coordinating solvents. 17 The only example of an isolated trityl-type cation

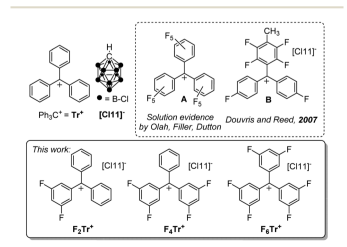


Fig. 1 The parent trityl cation Tr⁺, selected literature examples of fluorinated trityls, and the fluorinated trityl salts prepared and studied in this work.

[†] Electronic supplementary information (ESI) available: Experimental details and pictorial NMR spectra, details of the computational studies and the coordinate files. CCDC 2079761 and 2079762. For ESI and crystallographic data in CIF or other electronic format see https://doi.org/10.1039/d1sc05936j

Edge Article Chemical Science

fluorinated in the *ortho-/meta-*positions is **B** (Fig. 1), obtained by Douvris and Reed in an undefined vield, and not studied further.18 The perchlorotrityl cation has also been isolated.19

Our group has been attracted to the highly reactive carboand main-group cations in the context of our work on silylium and alumenium cation-catalyzed activation of aliphatic C-F bonds, 20-23 which permitted exhaustive defluorination of perfluoroalkyl groups under mild conditions. The chemistry of abstraction of fluoride from certain fluoroarenes with silylium cations has led to innovative reactivity, as well.24,25 Trialkylsilylium cations are typically generated by hydride abstraction from trialkylsilanes (R3SiH) using Tr+,26 but our theoretical analysis suggested that the parent Tr⁺ only barely has the thermodynamic hydride affinity (HA) to abstract hydrides from even the relatively electron-rich SiH bonds in trialkylsilanes. Given the perceived challenge10,16 in the isolation of the fully fluorinated $(C_6F_5)_3C^+$, we decided to first explore the partially fluorinated derivatives. Here, we report the isolation of analytically pure di-, tetra-, and hexafluorosubstituted trityl cation salts, and the remarkable contrast in the hydride abstraction reactivity with the parent Tr+.

Results and discussion

Theoretical HA analysis

Wilson and Dutton calculated gas-phase and CH₂Cl₂ solvent continuum HA values for a series of symmetric polychloro- and polyfluorosubstituted trityl cations.27 They discussed the fit to the known experimental values provided by the various computational methods and settled on the use of B3LYP/aug-ccpVTZ//B3LYP/def2-TZVPP.28,29

The Wilson-Dutton calculations showed that replacement of H with F in the para-position has an essentially zero effect on HA, whereas introduction of each ortho- or a meta-fluorine increases HA by ca. 2.4-2.7 kcal mol⁻¹ (CH₂Cl₂ continuum) or ca. 3.5 kcal mol⁻¹ (gas phase). This is in line with the more negative pK_{R+} values for the various *ortho*- and *meta*-fluorinated trityls compared to Tr⁺ or the para-F substituted trityls, determined by Filler et al. 10 The ortho- and para-CF positions are conjugated to the central carbon by resonance and the para-CF has been identified as a site of alternative nucleophilic attack on (C₆F₅)₃C⁺ related to its decomposition pathways. ^{10,16} We decided to avoid fluorination in the ortho- or para-positions and focus on meta-fluorination. The Wilson-Dutton HA values for F₆Tr⁺ $(213.0 \text{ and } 108.3 \text{ kcal mol}^{-1})$ were 11% and 17% higher than for Tr^{+} (191.4 and 92.5 kcal mol^{-1}) in the gas phase and $\mathrm{CH_2Cl_2}$ continuum, respectively.

In 2011,30 we analyzed the HA and FA values for a series of cations relevant to the silylium-catalyzed HDF using the M05-2X functional with the basis sets 6-311+G(d) for F, and 6-31++G(d,p) for C and H.31 Utilizing the DFT approach from our 2011 paper, we calculated the gas-phase and the chlorobenzene solvent continuum HA values for F₆Tr⁺ to be 229.4 and 135.0 kcal mol^{-1} , representing a 13% and a 20% increase vs. Tr⁺. These relative increases are similar to those in the Wilson-Dutton work.27 The substantial increase suggests that the HA of **F**₆**Tr**⁺ is thermodynamically sufficient to abstract a hydride from a range of Si-H containing molecules, and rivals the HA values calculated (also in PhCl) for Me₃C⁺ (126.6 kcal mol⁻¹), PhCH₂⁺ $(137.8 \text{ kcal mol}^{-1})$, and Me_2CH^+ $(138.9 \text{ kcal mol}^{-1})$. Without assessing quantitative accuracy, we nonetheless surmised that **F₆Tr⁺** might be able to abstract hydrides from tertiary and possibly secondary and benzylic C(sp³)-H bonds.

Synthesis and characterization of F_rTr⁺ salts

We envisioned the synthesis of fluorinated trityl cations partnered with the exceptionally robust and weakly coordinating $[HCB_{11}Cl_{11}]^-$ anion ([Cl11], Fig. 1)^{18,32-35} via abstraction of a chloride anion from the corresponding F2TrCl, F4TrCl, and F₆TrCl.^{36,37} Na[Cl11] can abstract a chloride from TrCl³⁸ and from F_2 TrCl in o- C_6 H₄Cl₂ at ambient temperature, giving a 97% isolated yield of F2Tr[Cl11] after workup. Attempts to use Na [Cl11] to generate F₄Tr[Cl11] and F₆Tr[Cl11] were unsuccessful and we moved to a more powerful³⁹⁻⁴¹ chloride abstractor $[(Me_3Si)_2OTf][Cl11]\ (\textbf{Si[Cl11]}).^{38,42}$

Indeed, treatment of F₂TrCl with Si[Cl11] in a 2:1 C₆D₆/o- $C_6H_4Cl_2$ solvent mixture at ambient temperature (Fig. 2) resulted in rapid color change. Analysis of the resultant solution by NMR spectroscopy after 10 min revealed the expected formation of equimolar amounts of Me₃SiCl and Me₃SiOTf and 96% yield of $\mathbf{F_2Tr}^+$ (19 F NMR evidence, δ –104.6 ppm). The analogous reactions with F_4 TrCl and F_6 TrCl also proceeded smoothly. The resultant F₄Tr[Cl11] and especially F₆Tr[Cl11] are less soluble than F2Tr[Cl11] or Tr[Cl11], and precipitate readily out of fluorobenzene, allowing isolation of analytically pure solids in 96% and 70% yields.

The ¹³C NMR chemical shifts of the central carbons of F₂Tr [Cl11], F₄Tr[Cl11], and F₆Tr[Cl11] in the 208-210 ppm range, 43 as well as the ¹H and ¹⁹F NMR spectral data did not suggest any significant interaction of the cations with the [Cl11] anion, the

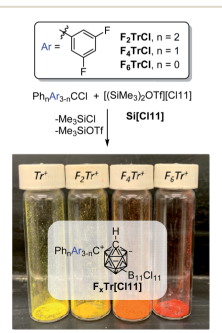


Fig. 2 Synthesis of fluorinated trityl cation salts and their appearance.

arene or $\mathrm{CD_2Cl_2}$ solvents, or the $\mathrm{Me_3SiCl/Me_3SiOTf}$ by-products. Single-crystal X-ray diffractometry (Fig. 3) revealed that the central carbons in $\mathrm{F_2Tr}[\mathrm{Cl11}]$ and $\mathrm{F_6Tr}[\mathrm{Cl11}]$ possessed a planar environment in all the crystallographically independent cations (the sums of C–C–C angles are $ca.~360^\circ$), and the aryl groups splay out in a pinwheel pattern about the central carbon. The closest approach of any chlorine to the central carbon in $\mathrm{F_2Tr}[\mathrm{Cl11}]$ is at least 3.7 Å, and 3.25 Å in $\mathrm{F_6Tr}[\mathrm{Cl11}]$, consistent with the well-separated, ionic nature of the $\mathrm{F_xTr}[\mathrm{Cl11}]$ salts.

Reactivity of F₆Tr⁺ vs. Tr⁺ with Et₃Si-H

It was previously shown that \mathbf{Tr}^+ is not capable of abstracting the full hydride equivalent from Et_3SiH in non-coordinating solvents and that two equivalents of R_3SiH are needed for complete formation of \mathbf{TrH} . Our observations are similar: treatment of either $\mathbf{F_6Tr}[\mathbf{Cl11}]$ or $\mathbf{Tr}[\mathbf{Cl11}]$ with two equivalents of Et_3SiH in a C_6D_6/o - $C_6H_4Cl_2$ solvent mixture led to the quantitative formation of $\mathbf{F_6Tr}H$ or \mathbf{TrH} , respectively. The fate of the " Et_3Si^+ " species in arene solvents is not straightforward, as has been studied in detail by Heinekey and coworkers: the presence of varying amounts of Et_4Si betrays complexity arising from the H/Et redistribution in the Si species and/or reactions with the arenes.

The reaction of Tr[Cl11] with a substoichiometric (0.9 equiv.) amount of Et_3SiH did not lead to the complete disappearance of the Si–H moiety (16% of the original Si–H intensity remained) and only 82% of the possible TrH was observed (Fig. 4). In contrast, the reaction of $F_6Tr[Cl11]$ with substoichiometric (0.75

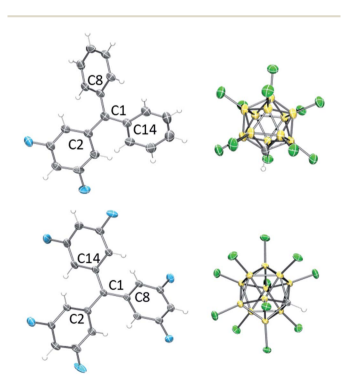


Fig. 3 POV-Ray rendition of the ORTEP (50% probability ellipsoids) drawing of F_2 Tr[Cl11] (top) and F_6 Tr[Cl11] (bottom). Only one cation and one anion from each asymmetric unit is shown. Solvent and disorder are omitted for clarity.

F [CI11]

F
$$0.75 \text{ eq HSiEt}_3$$
 $\text{Et}_3\text{Si[CI11]} + \textbf{F}_6\text{TrH}$

full conversion

[CI11]

 0.90 eq HSiEt_3 $\text{Et}_3\text{Si[CI11]} + \text{TrH} + \text{Et}_4\text{Si} + \text{H}_x\text{SiEt}_{4-x}$

incomplete conversion

Fig. 4 Reactions of Tr^+ and F_6Tr^+ with a substoichiometric amount of $HSiEt_{\bar{3}}$.

equiv.) amount of Et₃SiH led to the production of the expected quantity of F₆TrH, the complete disappearance of the Si-H signals, and without the concomitant observation of Et₄Si.

H-D exchange

In the reactions of **F₆Tr[Cl11**] with Et₃SiH, significant H/D scrambling was observed among the neutral aromatic compounds present in solution: C₆D₆, *o*-C₆H₄Cl₂, and **F₆TrH** (but the C(sp³)–H bond in **F₆TrH** was never deuterated). The extent of H–D exchange was analyzed *via* ¹H, ¹³C, or ¹⁹F⁴⁶ NMR spectroscopy (see ESI† for details). The mechanism of the H/D exchange likely involves the generation of superacidic protonated arenes *in situ*, ⁴⁷ which should enable rapid H/D exchange *via* H⁺/D⁺ shuttling (Fig. S4†). ^{47,48} The product of addition of either Et₃Si⁺ or **F₆Tr**⁺ to a neutral arene can be alternatively viewed as a protonated arene. ⁴⁷ It is also possible that analogous cations are accessed *via* reactions involving the minor components of the mixture. The Oestreich group recently examined this type of H/D exchange catalysis in greater detail. ⁴⁹

Abstraction of hydride from C-H bonds

Given the computational prediction of the enhanced hydride affinity of F_6Tr^+ vs. Tr^+ , we wished to examine their reactivity towards benzylic and aliphatic C-H bonds. As expected, no reaction was observed between Tr[Cl11] and (1) 1 equiv. of mesitylene or (2) 1 equiv. of methylcyclohexane in o-C₆H₄Cl₂ after 1 week at ambient temperature. In contrast, the reaction of $\mathbf{F_6Tr}[\mathbf{Cl11}]$ with mesitylene (as solvent) resulted in 66% yield (NMR evidence) or F₆TrH after 48 h. We propose that hydride abstraction from mesitylene by F6Tr[Cl11] generates a 3,5dimethylbenzyl cation, which rapidly undergoes Friedel-Crafts20,21 addition to mesitylene. GC-MS analysis of the mixture after quenching with water showed the presence of a m/z signal at 238, consistent with compound 4 (Fig. 5). Treatment of F₆Tr [Cl11] in o-C₆H₄Cl₂ with 1 equiv. of methylcyclohexane resulted in the >95% yield (NMR evidence) of F_6TrH after 96 h. The aliphatic region of the 1H NMR spectrum presented a large

F₆TrH 48 h 1 eq SiEt₄ SiEt₃[Cl11] F₆TrH r.t. m/z 352 2. H₂O C₆D₆/o-C₆H₄Cl₂ alkylated arene products r.t., 5-10 days ¹H NMR [CI11] 1 ea excess // F₆TrH o-C₆H₂Cl₂ C₆D₆/o-C₆H₄Cl₂ r.t., 18 h

Fig. 5 Reactions of F_6 Tr[Cl11] resulting in the abstraction of a hydride from $C(sp^3)$ -H bonds.

number of overlapping aliphatic signals, indicating a complex mixture (Fig. 5b).

The methylcyclohexyl cation presumed to be formed initially may undergo isomerization 50 and Friedel–Crafts addition to o- $C_6H_4Cl_2$, with many potential products. Abstraction of a hydride from alkanes, with generation of rearranged tertiary carbocations, was previously reported by the Reed group using Me $[HCB_{11}Me_5Br_6].^{51,52}$ The key difference between Reed's "Me+" reagents and the F_6Tr^{\dagger} reported here is that the latter can be prepared in bulk analytical purity and is stable in haloarene solutions.

Abstraction of a hydride from the β-position in trialkylaluminums with Tr⁺ has been used to generate reactive alumenium (R₂Al⁺) cations.^{6,22,53} The analogous abstraction of βhydride from alkylsilanes by Tr⁺ is not known, and we have confirmed that no reaction takes place between Tr[Cl11] and Et₄Si in C₆D₆/o-C₆H₄Cl₂. However, an analogous reaction of Et₄Si with F₆Tr[Cl11] resulted in the formation of 82% F₆TrH after 96 h (and complete disappearance of Et₄Si after 10 d). The major Si product appeared to be "Et₃Si", but instead of the stoichiometric complement of free ethylene, we observed ethane and other aliphatic resonances. Ethane may result from the protonolysis of Et₄Si by the highly Brønsted acidic cations generated in the reaction (extensive H/D exchange was concomitantly observed), a process reported on by Oestrich and co-workers.54 As a control experiment, we examined the reaction of $\mathbf{F_6Tr}[\mathbf{Cl11}]$ with 6.3 equiv. of ethylene in $\mathbf{C_6D_6/o \cdot C_6H_4Cl_2}$. Within 18 h at ambient temperature, all ethylene had been consumed, with the concomitant generation of ethylbenzene (1.8 equiv.) and other alkylarenes, and quantitative production of F_6 TrH. It is reasonable to propose that F_6 Tr[Cl11] abstracts a hydride from the benzylic positions of ethylbenzene or other alkylarenes generated through Friedel-Crafts alkylation. In complete contrast, no reaction occurred between Tr[Cl11] and ethylene under analogous conditions.

Conclusion

Introduction of six *meta*-F substituents in F_6Tr^+ brought about remarkable contrast with the reactivity of the parent triphenylmethyl (Tr^+) cation, understood primarily through the

greatly enhanced hydride affinity of especially the hexafluorinated $\mathbf{F_6Tr}^+$. Interestingly, while $\mathbf{F_6Tr}^+$ catalyzes the Friedel–Crafts alkylation of arenes with ethylene, and generates alkyl cations via hydride abstraction which then readily engage in Friedel–Crafts addition, $\mathbf{F_6Tr}^+$ itself is stable in combination with (halo)arene solvents and dichloromethane. This shows that fluorinated trityl cations represent a promising class of reagents for achieving the extremes of hydride affinity while minimizing reactivity with other potential substrates.

Data availability

Data for this manuscript are available in the ESI.†

Author contributions

S. O. G. and C. I. L. performed the syntheses and obtained the characterization data. E. S. performed the DFT calculations. N. B. carried out the X-ray diffraction studies on the crystals grown by S. O. G. S. O. G. and O. V. O. wrote the manuscript with assistance of the other co-authors. O. V. O. directed the overall effort

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by the Naval Research Laboratory (grant N00173-20-1-G010 to O. V. O.). We thank Dr Manoj Kolelveetil for insightful discussions. The authors acknowledge the High-Performance Research Computing Center at Texas A&M University for providing access to supercomputing clusters, software, and computing time. We thank R. A. Gholson for assistance with manuscript formatting.

Notes and references

1 G. A. Olah and G. K. S. Prakash, *Carbocation Chemistry*; John Wiley & Sons, Inc., 2004.

2 F. A. Carey and H. S. Tremper, *J. Am. Chem. Soc.*, 1968, **90**, 2578–2583.

Chemical Science

- 3 T.-Y. Cheng and M. R. Bullock, Organometallics, 1995, 14, 4031-4033.
- 4 E. S. Wiedner, M. B. Chambers, C. L. Pitman, R. M. Bullock, A. J. M. Miller and A. M. Appel, *Chem. Rev.*, 2016, **116**, 8655–8692.
- 5 E. Y.-X. Chen and T. J. Marks, *Chem. Rev.*, 2000, **100**, 1391–1432.
- 6 K.-C. Kim, C. A. Reed, G. S. Long and A. Sen, *J. Am. Chem. Soc.*, 2002, **124**, 7662–7663.
- 7 K.-C. Kim, C. A. Reed, D. W. Elliott, L. J. Mueller, F. Tham, L. Lin and J. B. Lambert, *Science*, 2002, 297, 825–827.
- 8 P. M. C. MacInnis, J. C. DeMott, E. M. Zolnhofer, J. Zhou, K. Meyer, R. P. Hughes and O. V. Ozerov, *Chem*, 2016, 1, 902–920.
- 9 T. Gessner and U. Mayer, Triarylmethane and Diarylmethane Dyes, in *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley-VCH, 2002.
- 10 R. Filler, C. Wang, M. A. McKinney and F. N. Miller, *J. Am. Chem. Soc.*, 1967, **89**, 1026.
- 11 G. A. Olah and M. B. Comisarow, *J. Am. Chem. Soc.*, 1967, **89**, 1027–1028.
- 12 V. Shrikant, R. Schure and R. Filler, *J. Am. Chem. Soc.*, 1973, **95**, 1859–1864.
- 13 J. R. Lawson and R. L. Melen, *Inorg. Chem.*, 2017, **56**, 8627–8643.
- 14 G. Erker, Dalton Trans., 2005, 11, 1883-1890.
- 15 M. Horn and H. Mayr, J. Phys. Org. Chem., 2012, 25, 979-988.
- 16 E. G. Delany, S. Kaur, S. Cummings, K. Basse, J. D. Wilson and J. L. Dutton, *Chem.-Eur. J.*, 2019, 25, 5298–5302.
- 17 I. M. Riddlestone, A. Kraft, J. Schaefer and I. Krossing, *Angew. Chem., Int. Ed.*, 2018, 57, 13982–14024.
- 18 C. Douvris, E. S. Stoyanov, F. S. Tham and C. A. Reed, *Chem. Commun.*, 2007, **40**, 145–1147.
- 19 E. Molins, M. Mas, W. Manjukiewicz, M. Ballester and J. Castaner, *Acta Crystallogr. C*, 1996, **52**, 2412–2414.
- 20 C. Douvris and O. V. Ozerov, Science, 2008, 321, 1188-1190.
- 21 C. Douvris, C. M. Nagaraja, C.-H. Chen, B. M. Foxman and O. V. Ozerov, *J. Am. Chem. Soc.*, 2010, 132, 4946–4953.
- 22 W. Gu, M. R. Haneline, C. Douvris and O. V. Ozerov, *J. Am. Chem. Soc.*, 2009, **131**, 11203–11212.
- 23 J. C. L. Walker, H. F. T. Klare and M. Oestreich, *Nat. Rev. Chem.*, 2020, **4**, 54–62.
- 24 J. S. Siegel, O. Allemann, S. Duttwyler, P. Romanato and K. K. Baldridge, *Science*, 2011, 332, 574–577.
- 25 B. Shao, A. L. Bagdasarian, S. Popov and H. M. Nelson, *Science*, 2017, 355, 1403–1407.
- 26 H. F. T. Klare, L. Albers, L. Süsse, S. Keess, T. Müller and M. Oestrich, *Chem. Rev.*, 2021, **121**, 5889–5985.
- 27 S. A. Couchman, D. J. D. Wilson and J. L. Dutton, *Eur. J. Org. Chem.*, 2014, 3902–3908.
- 28 (a) A. D. Becke, *Phys. Rev. A*, 1988, 38, 3098–3100; (b) C. Lee,
 W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1988, 37, 785–789.

- 29 A. Schäfer, C. Huber and R. Ahlrichs, J. Chem. Phys., 1994, 100, 5829–5835.
- 30 D. G. Gusev and O. V. Ozerov, *Chem.-Eur. J.*, 2011, **17**, 634–640.
- 31 Y. Zhao, N. E. Schultz and D. G. Truhlar, *J. Chem. Theory Comput.*, 2006, 2, 364–382.
- 32 E. S. Stoyanov, K.-C. Kim and C. A. Reed, *J. Am. Chem. Soc.*, 2006, **128**, 8500–8508.
- 33 S. P. Fisher, A. W. Tomich, S. O. Lovera, J. F. Kleinasser, J. Guo, H. M. Asay and V. Lavallo, *Chem. Rev.*, 2019, **119**, 8262–8290.
- 34 C. Douvris and J. Michl, *Chem. Rev.*, 2013, **113**, PR179-PR233.
- 35 W. Gu, B. J. McCulloch, J. H. Reibenspies and O. V. Ozerov, *Chem. Commun.*, 2010, **46**, 282–2822.
- 36 J. R. Bour, J. C. Green, V. J. Winton and J. B. Johnson, J. Org. Chem., 2013, 78, 1665–1669.
- 37 M. Horn, C. Metz and H. Mayr, Eur. J. Org. Chem., 2011, 6476–6485.
- 38 L. P. Press, B. J. McCulloch, W. Gu, C.-H. Chen, B. M. Foxman and O. V. Ozerov, *Chem. Commun.*, 2015, 51, 14034–14037.
- 39 D. J. Liston, Y. J. Lee, W. R. Scheidt and C. A. Reed, *J. Am. Chem. Soc.*, 1989, **111**, 6643–6648.
- 40 A. Martens, P. Weis, M. C. Krummer, M. Kreuzer, A. Meierhöfer, S. C. Meier, J. Bohnenberger, H. Scherer, I. Riddlestone and I. Krossing, *Chem. Sci.*, 2018, 9, 7058–7068.
- 41 C. A. Reed, Acc. Chem. Res., 1998, 31, 133-139.
- 42 A. Schulz, J. Thomas and A. Villinger, *Chem. Commun.*, 2010, **46**, 3696–3698.
- 43 G. A. Olah, A. L. Berrier and G. K. S. Prakash, *Proc. Natl. Acad. Sci. U. S. A.*, 1981, 78, 1998–2002.
- 44 M. Nava and C. A. Reed, *Organometallics*, 2011, **30**, 4798–4800.
- 45 S. J. Connelly, W. Kaminsky and D. M. Heinekey, *Organometallics*, 2013, 32, 7478–7748.
- 46 V. Salamanca and A. C. Albeniz, Eur. J. Org. Chem., 2020, 3206–3212.
- 47 H. F. T. Klare and M. Oestreich, *J. Am. Chem. Soc.*, 2021, **143**, 15490–15507.
- 48 D. Munz, M. Webster-Gardiner, R. Fu, T. Strassner, W. A. Goddard III and T. B. Gunnoe, *ACS Catal.*, 2015, 5, 769–775.
- 49 T. He, H. F. T. Klare and M. Oestreich, *J. Am. Chem. Soc.*, 2022, **144**(11), 4734–4738.
- 50 I. D. Mackie and J. Govindhakannan, J. Mol. Struct., 2010, 939, 53–58.
- 51 T. Kato and C. A. Reed, *Angew. Chem.*, *Int. Ed.*, 2004, **43**, 2908–2911.
- 52 C. A. Reed, Acc. Chem. Res., 2010, 43, 121-128.
- 53 M. Khandelwal and R. J. Wehmschulte, *Angew. Chem., Int. Ed.*, 2021, **51**, 7323–7326.
- 54 Q. Wu, Z. W. Qu, L. Omann, E. Irran, H. F. T. Klare and M. Oestreich, *Angew. Chem., Int. Ed.*, 2018, 57, 9176–9179.