Chemical Science

EDGE ARTICLE



Cite this: Chem. Sci., 2016, 7, 1773

Single-chain polybutadiene organometallic nanoparticles: an experimental and theoretical study⁺

Inbal Berkovich,^a Sudheendran Mavila,^a Olga Iliashevsky,^b Sebastian Kozuch^{ac} and N. Gabriel Lemcoff^{*a}

High molecular weight polybutadienes and rhodium complexes were used to produce single chain organometallic nanoparticles. Irradiation of high *cis*-polybutadiene in the presence of a photosensitizer isomerised the double bonds to produce differing *cis/trans* ratios within the polymer. Notably, a higher *cis* percentage of carbon-carbon double bonds within the polymer structure led to faster binding of metal ions, as well as their faster removal by competing phosphine ligands. The experimental results were supported and rationalized by DFT computations.

Received 25th November 2015 Accepted 7th January 2016

DOI: 10.1039/c5sc04535e

www.rsc.org/chemicalscience

Introduction

The creation of new compounds and the study of their properties is one of the most rewarding endeavours for experimental scientists. A new area of macromolecular research deals with the study and preparation of organic nanoparticles (ONPs).1-4 An emerging technique for the facile generation of tailor-made well-defined ONPs is the intramolecular cross-linking of single polymer chains; these ONPs are called single chain nanoparticles (SCNPs).⁵⁻¹¹ Moreover, the insertion of metals to these polymeric structures poses promising prospects for various applications, such as catalysis, drug delivery and sensing.12-18 In part, this potential arises because metallopolymers may combine the physical and chemical properties of both polymer and the embedded metal. Thus, the study of the factors affecting the formation, folding and architecture of metal containing SCNPs is an appealing goal.¹⁹⁻²⁵ The concept of single chain collapse driven by metal coordination has been introduced recently by several research groups. Pomposo and coworkers reported on the preparation of organometallic SCNPs by the intramolecular cross linking of copolymers bearing β ketoester chelating groups with Cu(II).19 The use of these nanoparticles in oxidative coupling reactions revealed an enhanced catalytic specificity compared to the classical catalysts such as $CuCl_2$, $Cu(OAc)_2$ and $Cu(acac)_2$. In another recent

work, Pomposo et al. have extended this methodology for the preparation of water-soluble globular SCNPs with polymerase mimicking activity.²⁰ Paik et al. detailed a cyclotetramerization cross-linking reaction within polymers bearing phthalonitrile pendant groups with CuCl resulting in SCNPs containing copper phthalocyanines.²¹ The Barner-Kowollik group recently used telechelic phosphine ligands to fold polymers with Pd(II).22 Moreover, Pd-SCNPs formed by a similar strategy have shown catalytic activity for a benchmark Sonogashira coupling reaction.²³ The carbon–carbon double bond π -electron coordination to metals is one of the most ubiquitous bonding types in organometallic chemistry.26-28 A recent example by Manners et al. shows how Pt ions may be used to reversibly cross-link a corona of polyisoprene to control the overall architecture of the macrostructure.²⁹ Our group has recently developed an intramolecular chain collapse approach for the synthesis of well-defined organometallic ONPs by a direct exchange of labile ligands of chlorobis(ethylene)rhodium(1) dimer by 1,5-hexadiene units present in poly-1,5-cis-cyclooctadiene (PCOD) produced by ring-opening metathesis polymerization (ROMP).24 The method was also expanded for the preparation of SCNPs with Ir(I) and Ni(0), showing promising applications in catalysis.25 The polymers used were about 50 kDa in size and had 20% cis content; typical values for ROMP reactions using Grubbs initiators.^{30,31} While the production of PCOD by this method is fairly straightforward, we sought for a way to achieve higher molecular weight polymers and to control the cis/trans ratio, as it may widen the scope of applications for this family of SCNPs and in turn afford new insights on the intricacies of metal- π bonding in these materials. 1,4-Polybutadiene (PBD), which has the same repeat unit as PCOD, is the second largest synthetic rubber product by volume and the one with the best performance to price ratio $(1.1 \text{ kg}^{-1}, 18.2 \text{ performance/price})$ ratio).32 In addition, it is available in high molecular weights

View Article Online

View Journal | View Issue

^aDepartment of Chemistry, Ben-Gurion University of the Negev, Beer-Sheva 84105, Israel. E-mail: lemcoff@bgu.ac.il

^bDepartment of Chemical Engineering, Ben-Gurion University of the Negev, Beer-Sheva 84105, Israel

^cLise Meitner – Minerva Center for Computational Quantum Chemistry, Israel

[†] Electronic supplementary information (ESI) available: Detailed experimental procedures and characterization of ONPs including: ¹H and ¹³C NMR, SEC, DLS, DSC, TGA, UV-vis, GC-MS spectra and computational calculations. See DOI: 10.1039/c5sc04535e

Chemical Science

and various *cis/trans* contents (depending on the catalyst used). Herein, we report on the preparation and characterization of rhodium(I) organometallic nanoparticles prepared by single chain collapse of high molecular weight polybutadienes of differing *cis/trans* ratios and the experimental and computational study of the effect of the double bond stereochemistry on the polymers' metal-binding properties.

Results and discussions

Synthesis of Rh(1)-ONPs from PBD with varying cis content

1,4-Polybutadienes with varying cis/trans content were obtained by irradiating toluene solutions of commercial PBD $(3 \times 10^5 \text{ g})$ mol^{-1} , PDI = 2.5, 95% *cis*, 5% vinyl) with 365 nm UV light in the presence of diphenyl disulfide as a photosensitizer for different periods of time (Fig. S1[†]).³³ The polymers obtained had a final cis content of 95%, 72%, 51% and 20% (determined by ¹H-NMR) without a significant change in their molecular weights. Having obtained the series of PBDs with varying cis/trans ratios, their single chain collapse was attempted via the addition of chlorobis(ethylene)rhodium(1) dimer (1), as rhodium donor, under dilute conditions. To our satisfaction, the PBDs quickly reacted at room temperature to afford polymer bound complexes 2 (Scheme 1), as determined by UV-vis, NMR, DLS and GPC analyses. The size of the organometallic SCNP was controlled by systematically varying the percentage of rhodium(1) introduced onto the polymer chain. As in the PCOD examples, a linear dependence between the amount of rhodium(1) (cross-linker) added and the reduction in the hydrodynamic radius of the Rh(I)-ONP was observed. The degree of chain collapse was



Scheme 1 Schematic illustration of the preparation of Rh(I)-ONPs.



Fig. 1 Overlay of SEC plots obtained for ONPs prepared with varying concentrations of Rh(I).

Table 1 Rh(i)-ONPs with varying rhodium(i) content^a

[III01%]	$[\times 10^5 \text{ g mol}^{-1}]$	PDI ^c	viscosity ^{c} (mL g ⁻¹)	(nm)
0	3.14	2.47	168.4	27.0
2	3.21	1.97	150.6	23.5
5	2.92	1.91	110.8	16.6
8	3.11	1.62	85.5	12.8
	[mol%] ^p 0 2 5 8	$\begin{array}{c c} [mol\%]^b & [\times 10^5 \mathrm{g} \mathrm{mol}^{-1}] \\ \hline 0 & 3.14 \\ 2 & 3.21 \\ 5 & 2.92 \\ 8 & 3.11 \end{array}$	$\begin{array}{c c} [\mathrm{mol}\%]^b & [\times 10^5 \mathrm{g \ mol}^{-1}] & \mathrm{PDI}^c \\ \hline 0 & 3.14 & 2.47 \\ 2 & 3.21 & 1.97 \\ 5 & 2.92 & 1.91 \\ 8 & 3.11 & 1.62 \end{array}$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

^{*a*} Conditions: solvent = THF, [PBD] = 0.5 mg ml⁻¹ (1.5 μ M), *T* = 35 °C. ^{*b*} Condition: Rh(i) = [RhCl(C₂H₄)₂]₂, mol% relative to 1,5-hexadiene units. ^{*c*} Condition: determined by triple-detector SEC in THF. ^{*d*} Condition: determined by DLS in THF.

monitored by both SEC and dynamic light scattering (DLS) in THF (Fig. 1 and Table 1, Fig. S9–S12[†]).

Kinetic study of rhodium(1) complexation

The observation that 1,4-polybutadienes with different *cis* content readily react with 1 to produce well-defined ONPs gave us the opportunity to study the effect of the double bond stereochemistry on the metal binding propensity. Thus, the rate of rhodium complexation was monitored with stopped flow kinetics by observing the change in the characteristic UV-vis absorption spectrum.

ROMP-derived PCOD (50 kDa, PDI 1.43, 20% *cis*, 80% *trans*) was used as a control under the same conditions. In each experiment, 10 mol% of **1** was mixed with a dilute solution of polymer and a plot of the absorbance at 360 nm *versus* time was recorded. Interestingly, the rate constants increased almost linearly as the *cis* content of the polymer was raised (Fig. 2). In addition, both 20% *cis* PBD and 20% *cis* PCOD showed similar rate constants, suggesting that the rate of the complexation is independent of the polymer size or origin. The influence of the double bond stereochemistry on binding was also studied with two 4,8-dodecadienes, one all *cis* and the other 75% *trans*, as additional model compounds. The reaction of the high *cis*



Fig. 2 Binding rate constants *versus cis* content of PBD and PCOD for the formation of Rh(I)-ONPs.

content 4,8-dodecadiene with 1 was about 7 times faster than the reaction of 75% *trans* 4,8-dodecadiene in accordance with the polymer experiments (Fig. S19†). These results suggest that the intrinsic stereochemistry of the carbon–carbon double bond influences the metal binding, as a similar trend was observed for both the small molecule model compounds and the polymers.

Computational study on the effect of double bond stereochemistry on rhodium binding

Quantum mechanical computations (at the M06/def2tzvpp(THF)//b97d3/def2-svp level,³⁴⁻³⁸ see the ESI†) shed some light on the reason for the faster association of the *cis* conformers of the 1,4-polybutadienes. *cis*- and *trans*-2-butene were used as models for the first step of the complexation process with complex **1**.

The first reaction with 2-butene, as shown in Fig. 3, involves four elementary steps. The first step is the formation of an



Fig. 3 Mechanism for the first ligand substitution of 2-butene with ethene. The numbers are Gibbs relative energies in kJ mol⁻¹ for the *cis*-2-butene, in a THF continuum solvent.

adduct stabilized only by van der Waals interactions (1_b) , which - although exothermic - must pay a high entropy penalty, thus resulting endergonic. The second step is the insertion of the butene to form an $18e^-$ complex (1_c through transition state TS_{bc}). This is followed by the dissociation of the ethene (TS_{cd}) and $\mathbf{1}_d$), which results in the determining transition state for the complete ligand substitution. Finally, the ethene moiety is completely released to form 1_e. The complete reaction should result endergonic (at standard concentrations), but it is driven by the release of ethene as a gas. Several conformers were considered (including the inversion of the bent structure between the two coordination planes),³⁹ but only the most relevant conformers will be discussed here (for higher energy structures see the ESI⁺). Fig. 3 shows the relative Gibbs energies in THF for the pathway including cis-2-butene. All the other pathways have very similar energy profiles, and therefore only the relation between the ΔG of the determining transition states40-42 (TScd) of the various mechanisms is relevant to understand the experimentally observed kinetics for the cis and trans species. The most probable mechanism for the trans-2butene substitution presents a relative ΔG of 76.1 kJ mol⁻¹ for TS_{cd} , only 3.1 kJ mol⁻¹ higher than the equivalent state in the *cis* reaction. At room temperature this corresponds to 3.5 times faster reaction for the ethene substitution with a cis conformer compared to the trans one, agreeing with the experimental results (considering the computational and experimental differences in the reactant models).

The reason for the higher energy of the *trans* pathway is shown in Fig. 4: the repulsion between the methyl group of the *trans* conformer and the ethene neighbour ligand produces a distortion of the verticality of the butene, which translates in a slightly higher energy all along the reaction (see the ESI[†]).

While in the *cis* conformer the Cl–Rh–C–C dihedral angle is virtually perpendicular (89°), the two possible *trans* conformers have angles of 77° and 109° (Fig. 4). Note that the chloride ligand in the opposite side of the ethene also exerts some steric hindrance, pushing out the methyl groups of the butene. However, in the *cis* case the chloride ligand pushes both methyl groups simultaneously, without affecting the dihedral angle of the butene and maintaining the back-bonding character of the metal–olefin bond.

Although the second ligand substitution may be of kinetic importance in the case of 2-butene, for chelating dienes there will be a small entropy penalty on the second substitution, as



Fig. 4 (A) *cis* conformer of the monosubstituted dirhodium complex $\mathbf{1}_{e}$, where the butene double bond is perpendicular to the complex plane, optimizing the back-bonding interaction. (B) *trans* conformers; the curves show steric impediments which diminish the π back-bonding of the metal to the ligand.

the second double bond is already in the vicinity of the metal. In the overall reaction there will be a net entropy gain when a diene species displaces two ethene molecules, and as a result the only critical step of the complete mechanism will be at the first substitution.

Kinetic and computational study of rhodium(1) release

Having shown that this type of organometallic ONPs are reversible in nature,24 the release of Rh from PBDs was also studied by addition of a competing ligand, tricyclohexylphosphine (PCy₃, Scheme 2). Thus, excess PCy₃ was added to 10 mol% Rh(I)-ONPs and the absorbance at 360 nm was monitored using stopped flow kinetics. Addition of the phosphine led to a reduction in the absorption at 360 nm, indicating the disruption of the organometallic bond. A plot of the rate constants versus the cis content of the polymer showed that the reaction of 95% cis Rh(1)-ONPs with PCy3 was the fastest among the series of PBDs used; while the slowest rate constants were obtained with 20% cis-PCOD and PBD (Fig. 5). Also rhodium complexes of 4,8-dodecadienes were mixed with PCy₃ and the absorption at 360 nm was monitored (Fig. S20[†]). The control experiment with cis, cis-4,8-dodecadiene-rhodium complex was also faster than that with the trans isomer.

Thus, experiments showed that not only the association, but also the dissociation of the alkenes – driven by substitutions with PCy_3 – are faster in the *cis* system, a fact that may seem to be at odds with the greater stability of the η^2 bond in intermediate complex $\mathbf{1}_e$ with *cis*-2-butene compared to *trans*-2-butene. To gain a computational insight into this occurrence, 2,6-octadiene with a double *cis* or double *trans* configuration was used as a model. Computations showed that the *cis*-2,6-octadiene forms a complex 36 kJ mol⁻¹ higher in energy than the double *trans* case, thus being more reactive to ligand substitution.



Scheme 2 Reaction of 2 with PCy₃.



Fig. 5 Rate constants versus cis content of PBD and PCOD for the disruption of Rh(i)-ONPs.



Fig. 6 *cis* (A) and *trans* (B) 2,6-octadienes as chelating double η^2 ligands. In the *cis* species the double bonds cannot achieve the ideal vertical position that maximizes the back-bonding interaction, and therefore results in a higher energy, more reactive complex.

Again, this is mainly due to strain arising when the olefin double bonds cannot achieve a vertical position (that is, perpendicular to the complex plane where the back-bonding interaction is maximized). As can be seen in Fig. 6, the double *trans* diene can be positioned with both π bonds perpendicular to the Cl–Rh–Cl plane, a preferred geometry, but the double *cis* conformer (Fig. 6A) is forced to rotate one of the double bonds.

In summary, the formation of complexes based on *cis*-dienes is faster than with the *trans* ligands because the first substitution (which determines the rate of association) has less steric impediments in the *cis* case. The dissociation of *cis*-dienes is also faster than *trans*, since the chelating diene in the *cis* conformer cannot achieve a stable geometry with both double bonds perpendicular to the complex plane, partially breaking the back bonding stabilization and making this a more reactive complex prone to a faster ligand substitution, in accordance to the experimental observations.

The effect of the intermolecular cross-linking of PBD on the thermal properties

Having shown that dilute solutions of polymers containing the 1,5-hexadiene unit have the ability to form organometallic complexes, we sought to investigate whether the bulk polymer could also act as a metal sponge and change its properties when ions are absorbed. It has been shown that the coordination of transition metals to olefin containing polymers, such as Pd(n)

Table 2 T_g of 95% cis-PBD with varying amount of Rh(I)

Entry	$Rh(i)^{a}$ [mol%]	$T_{\rm g}$ (°C)
1	0	-101.9
2	5	-88.5
3	10	-71.2
^{<i>a</i>} mol% relative t	o 1.5-hexadiene units.	

with atactic 1,2-polybutadiene, has a significant effect on the glass transition temperature (T_g) of the composite.⁴³ It has also been reported that Rh containing PBD copolymers (SBS) could be prepared by heating a THF or benzene solution in the presence of RhCl₃.⁴⁴ However, the influence of different amounts of metal coordination on the T_g of 1,4-polybutadiene has not been studied. Thus, samples of PBD were soaked overnight in THF solutions with varying concentrations of rhodium. The effect of this procedure was studied by analyzing the change in the glass transition temperature of the polymer sample and the shift in the UV-vis absorption spectrum (Fig. S21†). Indeed, as shown in Table 2, the metal led to a significant change in the polymer's physical properties as expected from the intermolecular cross-linking caused by the metal bridges.

Conclusions

Polybutadiene, an abundant starting material, was used for the first time as the precursor single polymer chain for the formation of organometallic SCNPs. Because these polymers are much larger than those previously used, they may bind many more metal ions per nanoparticle; an important requisite for future applications (e.g. templates for inorganic nanoparticles). Interestingly, the introduction of rhodium ions to neat PBD produced increased glass transition temperatures as a consequence of the restricted local chain mobility caused by the metal bridges. By investigating polymers with differing cis/trans ratios, we have experimentally and computationally shown that there is a stark difference in the rate of metal sequestration and release when polymers with different microstructures are used to form organometallic nanoparticles. Indeed, polymers with higher ratio of *cis* double bonds bind rhodium ions faster and, when competitive ligands are added, also release them faster. To put this in perspective, under our experimental conditions the rhodium ions could be bound in less than ten seconds by a 300 kDa all cis-PBD and they could be completely removed in about seven minutes by the addition of excess phosphine, while with a high trans-PBD it would take thrice as long. The availability of cis-selective olefin metathesis catalysts^{45,46} may allow the formation of novel 1,5-cyclooctadiene polymers or copolymers with high cis/trans ratios which may be useful for quick binding and release of selected metal ions for future applications of this method. DFT based computational studies supported the experimental results and revealed the mechanistic and steric reasons for the behaviour observed. Taking into account the great importance of PBD cross-linking in the rubber industry, the study of soft, reversible, cross-linking

methodologies by metals and the way the polymer structure affects the metal intake is of significant value.

Acknowledgements

The United States – Israel Binational Science Foundation (Grant No. 2014116) is gratefully acknowledged for partial funding of this research.

Notes and references

- 1 J. P. Rao and K. E. Geckeler, *Prog. Polym. Sci.*, 2011, **36**, 887–913.
- 2 K. Li and B. Liu, Chem. Soc. Rev., 2014, 43, 6570-6597.
- 3 M. Elsabahy and K. L. Wooley, J. Polym. Sci., Part A: Polym. Chem., 2012, 50, 1869–1880.
- 4 M. Elsabahy, G. S. Heo, S.-M. Lim, G. Sun and K. L. Wooley, *Chem. Rev.*, 2015, **115**, 10967–11011.
- 5 S. Mavila, O. Eivgi, I. Berkovich and N. G. Lemcoff, *Chem. Rev.*, 2015, DOI: 10.1021/acs.chemrev.5b00290.
- 6 C. K. Lyon, A. Prasher, A. M. Hanlon, B. T. Tuten, C. A. Tooley, P. G. Frank and E. B. Berda, *Polym. Chem.*, 2015, 6, 181–197.
- 7 O. Altintas and C. Barner-Kowollik, *Macromol. Rapid Commun.*, 2012, 33, 958–971.
- 8 M. Gonzalez-Burgos, A. Latorre-Sanchez and J. A. Pomposo, *Chem. Soc. Rev.*, 2015, 44, 6122–6142.
- 9 M. Aiertza, I. Odriozola, G. Cabañero, H.-J. Grande and I. Loinaz, *Cell. Mol. Life Sci.*, 2012, **69**, 337–346.
- 10 T. Mes, R. van der Weegen, A. R. A. Palmans and E. W. Meijer, *Angew. Chem., Int. Ed.*, 2011, **50**, 5085–5089.
- 11 P. J. M. Stals, M. A. J. Gillissen, T. F. E. Paffen, T. F. A. de Greef, P. Lindner, E. W. Meijer, A. R. A. Palmans and I. K. Voets, *Macromolecules*, 2014, 47, 2947–2954.
- 12 K. A. Williams, A. J. Boydston and C. W. Bielawski, *Chem. Soc. Rev.*, 2007, **36**, 729–744.
- 13 M. Ramanathan and S. B. Darling, *Polym. Int.*, 2013, **62**, 1123–1134.
- 14 G. R. Whittell, M. D. Hager, U. S. Schubert and I. Manners, *Nat. Mater.*, 2011, **10**, 176–188.
- 15 J. Xiang, C.-L. Ho and W.-Y. Wong, *Polym. Chem.*, 2015, 6, 6905–6930.
- 16 X. Wang and R. McHale, *Macromol. Rapid Commun.*, 2010, **31**, 331–350.
- 17 Y. Liu, T. Pauloehrl, S. I. Presolski, L. Albertazzi,
 A. R. A. Palmans and E. W. Meijer, *J. Am. Chem. Soc.*, 2015, 137, 13096–13105.
- 18 M. Artar, E. R. J. Souren, T. Terashima, E. W. Meijer and A. R. A. Palmans, ACS Macro Lett., 2015, 4, 1099–1103.
- 19 A. Sanchez-Sanchez, A. Arbe, J. Colmenero and J. A. Pomposo, *ACS Macro Lett.*, 2014, **3**, 439–443.
- 20 A. Sanchez-Sanchez, A. Arbe, J. Kohlbrecher, J. Colmenero and J. A. Pomposo, *Macromol. Rapid Commun.*, 2015, **36**, 1592–1597.
- 21 J. Jeong, Y.-J. Lee, B. Kim, B. Kim, K.-S. Jung and H.-J. Paik, *Polym. Chem.*, 2015, **6**, 3392–3397.

- 22 J. Willenbacher, O. Altintas, P. W. Roesky and C. Barner-Kowollik, *Macromol. Rapid Commun.*, 2014, **35**, 45–51.
- 23 J. Willenbacher, O. Altintas, V. Trouillet, N. Knofel, M. J. Monteiro, P. W. Roesky and C. Barner-Kowollik, *Polym. Chem.*, 2015, 6, 4358–4365.
- 24 S. Mavila, C. E. Diesendruck, S. Linde, L. Amir, R. Shikler and N. G. Lemcoff, *Angew. Chem., Int. Ed.*, 2013, **52**, 5767– 5770.
- 25 S. Mavila, I. Rozenberg and N. G. Lemcoff, *Chem. Sci.*, 2014, 5, 4196–4203.
- 26 C. Huber, F. Bangerter, W. R. Caseri and C. Weder, J. Am. Chem. Soc., 2001, **123**, 3857–3863.
- 27 J. B. Johnson and T. Rovis, *Angew. Chem., Int. Ed.*, 2008, 47, 840–871.
- 28 M. A. Bennett, Chem. Rev., 1962, 62, 611-652.
- 29 P. A. Rupar, G. Cambridge, M. A. Winnik and I. Manners, *J. Am. Chem. Soc.*, 2011, **133**, 16947–16957.
- 30 C. W. Bielawski and R. H. Grubbs, Angew. Chem., Int. Ed., 2000, 39, 2903–2906.
- 31 B. Allaert, N. Ledoux, N. Dieltiens, H. V. Mierde, C. V. Stevens, P. van Der Voort and F. Verpoort, *Catal. Commun.*, 2008, 9, 1054–1059.
- 32 H. Brandt, Ullmann's encyclopedia of industrial chemistry, Wiley-VCH, 2012.
- 33 M. A. Golub, J. Polym. Sci., 1957, 25, 373-377.

- 34 Y. Zhao and D. Truhlar, *Theor. Chem. Acc.*, 2008, **120**, 215-241.
- 35 S. Grimme, J. Comput. Chem., 2006, 27, 1787-1799.
- 36 S. Grimme, S. Ehrlich and L. Goerigk, *J. Comput. Chem.*, 2011, **32**, 1456–1465.
- 37 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. B*, 2009, **113**, 6378–6396.
- 38 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, 7, 3297–3305.
- 39 G. Aullón, G. Ujaque, A. Lledós, S. Alvarez and P. Alemany, *Inorg. Chem.*, 1998, 37, 804–813.
- 40 J. R. Murdoch, J. Chem. Educ., 1981, 58, 32-36.
- 41 S. Kozuch and S. Shaik, Acc. Chem. Res., 2011, 44, 101-110.
- 42 S. Kozuch, in Understanding Organometallic Reaction Mechanisms and Catalysis, Wiley-VCH Verlag GmbH & Co. KGaA, 2014, pp. 217–248, DOI: 10.1002/9783527678211.ch9.
- 43 L. A. Belfiore and M. P. McCurdie, J. Polym. Sci., Part B: Polym. Phys., 1995, 33, 105–124.
- 44 L. M. Bronstein and P. M. Valetsky, J. Inorg. Organomet. Polym., 1994, 4, 415-424.
- 45 B. K. Keitz, A. Fedorov and R. H. Grubbs, *J. Am. Chem. Soc.*, 2012, **134**, 2040–2043.
- 46 M. B. Herbert and R. H. Grubbs, *Angew. Chem., Int. Ed.*, 2015, 54, 5018–5024.