

H-Bonding leading to latent initiators for olefin metathesis polymerization†‡

Artur Brotons-Rufes,^a Naeimeh Bahri-Laleh  ^{*b} and Albert Poater  ^{*a}

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Ruthenium–NHC based catalysts, with a chelated iminium ligand *trans* to the N-heterocyclic carbene (NHC) ligand, that polymerize dicyclopentadiene (DCPD) at different temperatures are monitored using Density Functional Theory calculations to unveil the reaction mechanism, and subsequently how important are the geometrical and electronic features vs. the non-covalent interactions in between. The balance is very fragile and H-bonds are fundamental to explain the different behaviour of latent catalysts. This computational study aims to facilitate future studies of new generations of latent initiators for olefin metathesis polymerization, with the 3D and mainly the 2D Non-Covalent Interaction plots the characterization tool for H-bonds.

Introduction

Among the wide variety of polymers, one property that can make them more special is that they have the ability to withstand aggressive temperatures.¹ Thermoset polymers are a group of highly versatile materials due to their capability to harden in an irreversible curing process,² which changes working materials from soft solids or resins to rigid networks.³ Phase transformation can be triggered by heat and pressure, radiation or when mixed with a catalyst,⁴ in particular by means of olefin metathesis reactions.^{5–12} A typical example of a thermoset polymer is poly(dicyclopentadiene) (PDCPD) which is synthesized through ring-opening metathesis (ROMP) of dicyclopentadiene (DCPD).^{13,14} ROMP changes the soft structure of monomeric DCPD or DCPD-derived linear oligomeric chains to a hard cross-linked polymer array. Derived materials are of great interest because of their thermal and corrosion resistance. Because of their mechanical properties, they also find applications in the materials, petrochemical and energy generating

^a*Institut de Química Computacional i Catàlisi, Departament de Química, Universitat de Girona, c/Maria Aurèlia Capmany 69, 17003 Girona, Catalonia, Spain. E-mail: albert.poater@udg.edu*

^b*Polymerization Engineering Department, Iran Polymer and Petrochemical Institute (IPPI), P.O. Box 14965/115, Tehran, Iran. E-mail: n.bahri@ippi.ac.ir*

† Dedicated to the memory of our colleague and inspiration Prof. Dr Robert H. Grubbs.

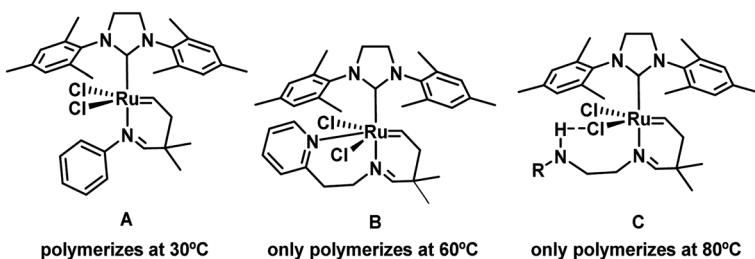
‡ Electronic supplementary information (ESI) available: XYZ coordinates and absolute energies of all species. See DOI: <https://doi.org/10.1039/d2fd00163b>



industries.¹⁵ Effective ROMP reactions of DCPD are successfully achieved *via* the reaction injection moulding (RIM) technique,^{16,17} in which the curing reaction is accomplished by mixing together of the DCPD monomer and a catalyst in the moulding process.^{15,18} Apart from preventing decomposition due to the presence of leaving groups such as phosphines *trans* to the N-heterocyclic carbene (NHC) ligand,^{19–22} or even getting rid of them,^{23–25} to prevent undesired premature polymerization, *i.e.* room temperature (rt) polymerization, the latent catalysts have been proposed as a potential solution,²⁶ and therefore the interest in them has grown in recent years,²⁷ knowing that a drawback of catalysts that initiate ROMP polymerization at rt is the requirement for a pretreatment, which is not suitable for RIM.^{28,29} Anyway, efficient latent catalysts have been described^{30–33} that work through thermal^{34–41} and (photo)chemical activation mechanisms.^{42–49}

Ruthenium catalysts bearing NHC and alkylidene ligands are found to be active at room temperature for ROMP of DCPD.³¹ Hejl *et al.* reported tuneable latency activity through alteration of the stereoelectronics of the substituted ligands. In these studies, alkylidene ROMP catalysts were tested, showing latent behaviour when an endocyclic imine moiety was present (Scheme 1A). Other studies reported that the nature of the imine substituents affected the activity of the catalyst, showing potential tuneability.³⁰ All these results also coincide with the current mechanistic descriptions of ruthenium catalysed metathesis, where the removal of a donor ligand provides entrance to the catalytic cycle.^{50,51}

With the idea to favour the latency of the initiators it was not enough with the addition of donor groups such as methoxy or thioether substituents,³⁰ but Kim and co-workers found it was mandatory to introduce a deeper skeletal modification, including pyridine based ligands on the ruthenium centre, facilitating the polymerization at 60 °C.³⁹ Next, the same research group showed that the presence of a donor ligand with an additional potential binding capacity could enhance the process. The results clearly showed that for alkylidene-containing NHC–Ru complexes,⁵² the additional donor atom increases the initiation temperature, avoiding polymerization under 30 °C, but achieving normal reactivity when the temperature was increased up to 60 °C (Scheme 1B).³⁴ Similar results were reported when a hydrogen donor such as carbamate or acetamide was introduced in the imine moiety, in detail preventing the dissociation of the imine nitrogen in the alkylidene moiety. Acting as a latent initiator, the hydrogen donor provides a weak hydrogen-bond interaction with the metal-bound chloride anion of the catalyst (Scheme 1C), which would stabilize sufficiently the



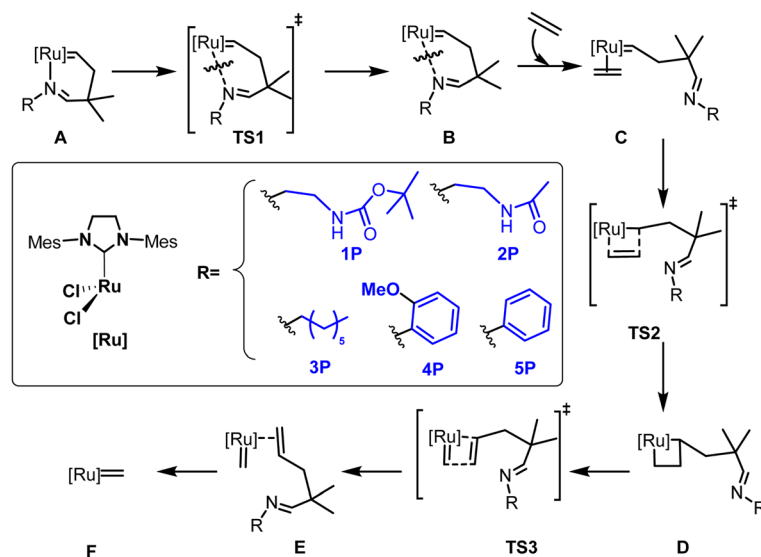
Scheme 1 Different ruthenium–NHC catalysts for DCPD polymerization with their temperature of initiation.

precatalyst,⁵³ and consequently the initiator remains unactive below 80 °C.³⁵ In parallel, Czarnocki and co-workers claimed the enhancement of the latency with the insertion of a carbonyl ester group in a modified Hoveyda–Grubbs catalyst, replacing the oxygen by a nitrogen bonded to the ruthenium.⁵⁴ Nevertheless, the conversion of carbonyl groups to carboxylates, with air and a strong base such as triethanolamine, cannot be extrapolated to any of the systems included in Scheme 1 for their lack of acidic protons to facilitate the intramolecular cyclization, apart from the low chances of a carbonyl group to generate 7-membered rings together with the metal centre and the chelated imine.

Inspired by this novel H-donor based latent strategy of Kim and co-workers,³⁹ computational studies using Density Functional Theory (DFT) calculations and Non-Covalent Interactions (NCI) plots were conducted to delve into the origin of their increased thermal latency, trying to better define how this intramolecular interaction is able to increase the latent behaviour of these systems.

Results and discussion

The series of imine ylidene ligands **1P–5P** in Scheme 2 was compared in terms of the activation pathway in olefin metathesis, following the mechanism in Scheme 2. Five different substituents were considered as imine derivatives in the calculations. Both methyl carbamate and amide, substituents **1P** and **2P** respectively, include a hydrogen donor moiety that allows a hydrogen bond with one of the chlorides of the metal centre. Substituent **3P** also has a long chain as in the previously mentioned ones, but the lack of any additional functionality makes it unable to undertake additional electronic intramolecular interactions with the metal centre. Finally, substituents **4P** and **5P** were introduced in order to evaluate possible differences due to the length of the imine substituent. Additionally,



Scheme 2 The studied activation pathway of the Ru based catalysts 1–5.

system **4** contains the methoxy group that could electronically not only modify the electron donating capacity of the imine moiety, but also generate a potential hydrogen bond with its oxygen atom.

From Scheme 2, starting with the imine based chelate precatalyst **A**, and finishing with the ylidene **14e** active species **F**, the mechanism is somewhat analogous with respect to the Hoveyda–Grubbs catalysts,⁵⁴ with the replacement of the oxygen atom bonded to ruthenium by nitrogen.⁵⁵ Before a detailed structural and electronic analysis of the set of catalysts under analysis, the energetic values for the reaction pathway of the activation are included in Table 1.

The breaking of the *trans* Ru–N bond marks the beginning of preactivation, with a kinetic cost that involves a range of values that goes from 15.8 for complex **5** to 22.9 kcal mol^{−1} for complex **2**, therefore with values similar to Hoveyda complexes,^{56,57} and even those that simply have a *trans* phosphine in the NHC.⁵⁸ This first dissociative step seems trivial, but it is also framed by a relatively poor thermodynamic stabilization, since the later intermediate **B** shows that by Hammond's principle the transition state is a late transition state, with a destabilization of at least 14.9 kcal mol^{−1} in all cases. And even more is evident by incorporating ethylene into the equation, being *trans* to the NHC, in the coordination intermediate **C**, extending this value to at least 21.6 kcal mol^{−1} and worryingly to 27.8 kcal mol^{−1} for complex **2**, although the correction of the entropic overestimation has been omitted here,⁵⁹ in order to not treat the concentration at 1 M of solutes,⁶⁰ and neither of solvents,⁶¹ which would cause a reduction of 2–6 kcal mol^{−1} depending on the nature of them. From **C**, **TS2** leads to the metallacycle **D** overcoming the energy barriers of little importance, a maximum of 4.8 kcal mol^{−1}, and likewise, the break in the second coordination intermediate **E**, homologous to **C**, is given by energy barriers that in no case exceed 7.8 kcal mol^{−1}. However, this metallacycle opening **TS3** defines the rate determining step (rds) of the precatalyst activation mechanism, which then ends up with the catalytic active species **F**, except for systems **4** and **5**, where the rds switches to the formation of the metallacycle, **TS2**, however only by 0.3 and 1.2 kcal mol^{−1}, respectively. On the other hand, it is also worth noting the relatively low stability of the intermediate **E**, which even when the solvent effect is added in the point energy calculation turns out to be unexpectedly more unstable than the previous **TS3**.

Once the cost of preactivation has been described, to determine the reason for the different behaviour of the five systems under study, it is necessary to put them under the microscope of structural and electronic factors, preferably emphasizing the imine group, and how it interacts with the rest of the system itself. Thus, the optimized geometries of precatalyst **A** in systems **1** and **2** demonstrate that the N–

Table 1 Relative Gibbs energies (in kcal mol^{−1}) calculated at 353.15 K of the olefin metathesis reaction pathway with catalysts **1–5** and ethylene as substrate

	A	TS1	B	C	TS2	D	TS3	E	F
1	0.0	21.5	14.9	27.4	25.2	23.2	29.5	26.9	23.9
2	0.0	22.9	21.8	27.8	29.6	25.7	31.9	30.0	25.0
3	0.0	19.4	19.3	25.3	25.8	22.3	30.1	26.5	27.7
4	0.0	17.9	16.0	22.2	27.0	20.3	26.7	27.9	23.6
5	0.0	15.8	15.7	21.6	23.4	19.8	22.2	23.8	21.6



H bond is already oriented toward one of the chlorides of the ruthenium catalyst (Fig. 1). The distances between the polarized hydrogen and the nearest chloride are 2.210 and 2.193 Å, respectively. Thus, with the values obtained, and taking into account that an average hydrogen bond in water has a longer length, the additional intramolecular interaction was considered to be present for the pre-catalytic intermediates with substituents **P1** and **P2**, and enjoying a considerable magnitude. Although the most important weight of H-bonds is installed in the first Ru–N bond cleavage step, they were studied for the rest of the intermediates in the reaction pathway. While for system **1**, the (N)H···Cl interaction disappears for the rest of the mechanism, this is not the case for catalyst **2**, where we still keep a rather weak H-bond in intermediate **B** (2.563 Å).

Surprisingly, after unveiling the activation in Table 1, the above-mentioned hydrogen bond would not affect the overall thermodynamic pathway of the reaction. Although a relatively higher barrier is observed in the first transition state **TS1** of systems **1** and **2**, which belongs to the opening of the chelating ring from the initial precatalytic geometry, it would not affect it dramatically since it does not correspond to the rds. However, a lower kinetic cost is obtained for the

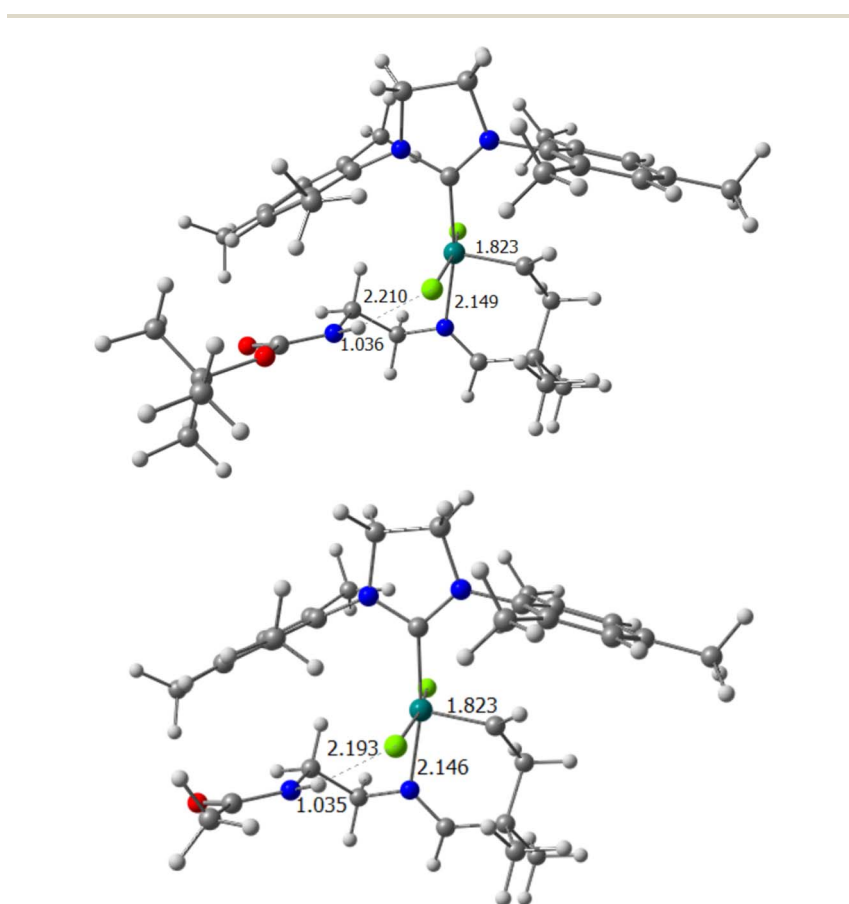


Fig. 1 Precatalyst A for systems **1** (above) and **2** (below), main distances are shown in Å.



systems with no internal hydrogen bond, not only for this **TS1**, but also for the next transition states, including the rds.

A more comparative analysis between the five systems, carried out with a tool capable of analysing the strength of bonds of different nature, such as the Mayer Bond Order (MBO) index,⁶² distils in Table 2 that the Ru–N bond in breaking follows the order of **1** \cong **2** $>$ **3** $>$ **5** $>$ **4**, that is, we would expect it to be more difficult to break the Ru–N bond especially for the first two systems. It is also interesting in Table 2 to confirm the quite strong H-bond between the N–H moiety and the closest Cl ligand, with a MBO over 0.1 for systems **1** and **2**. On the other hand, the MBOs for the metallacycle unveil that once the Ru–N bond has broken the next formation of the metallacycle **D** shows a ruthenacycle with a core nearly identical for the five systems.

Table 2 Mayer bond orders for intermediates **A** and **D** for systems **1**–**5** with ethylene as the substrate (C1 corresponds to the former C_{ylidene}, while C2 and C3 to the carbon atoms of the entering ethylene)

	A		D			
	Ru–N	(N)H···Cl	Ru–C1	C1–C2	C2–C3	C3–Ru
1	0.489	0.117	1.091	0.879	0.955	1.185
2	0.490	0.119	1.093	0.877	0.955	1.183
3	0.480	—	1.092	0.877	0.955	1.184
4	0.465	—	1.093	0.878	0.955	1.184
5	0.474	—	1.093	0.879	0.955	1.185

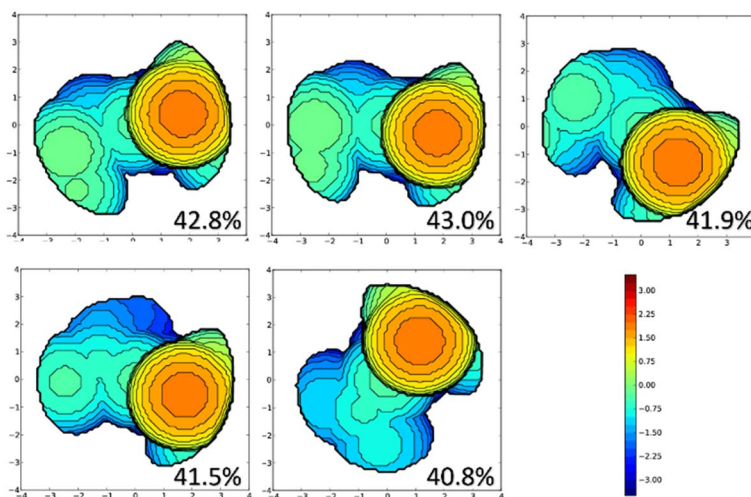


Fig. 2 Topographic steric maps (xy plane) and $\%V_{\text{Bur}}$ of the imine–ylidene ligands of species **A** for systems **1**–**5**. The Ru atom is at the origin and the N_{imine} atom is on the z axis, and the N atoms of the NHC ligand on the xz plane. The isocontour curves of the steric maps are given in Å. The radius of the sphere around the metal centre was set to 3.5 Å, while for the atoms we adopted the Bondi radii scaled by 1.17, and a mesh of 0.1 Å was used to scan the sphere for buried voxels (with a radius of 3.5 Å, the isocontour curves of the steric maps are given in Å).

To assess the lower sterical pressure on the metal centre, as well as to differentiate among the precatalysts **A** for systems **1–5**, we performed steric maps following the scheme developed by Cavallo and co-workers.⁶³ The results in Fig. 2 confirm the larger sterical hindrance on the metal centre due to the imine–ylidene ligands of systems **1** and **2**.⁶⁴ Actually, not only in the precatalyst **A**, but also throughout all the activation pathway (see ESI for further details‡).

Electronically the Natural Bond Orbital analysis of the charges in Table 3 shows that there are no differences in the metallacycle, and no significant differences in the precatalyst **A** for systems **1** and **2** compared to the other systems, except for the interesting positive charge on the hydrogens of the N–H moieties in systems **1** and **2**.

A deeper analysis with the NBO approach of the NH···Cl interaction for intermediate **A** of system **1** unveiled the importance of the lone pair (LP) corresponding to both chlorides together with the antibonding character owed by the bonding orbital (BD*) for the NH moiety with a total energy of 11.6 kcal mol^{−1}.

Before concluding that the electronic part does not affect the reactivity, taking into account the NBO charges, in Table 4 and Fig. 3, we proceeded to do a simple analysis of the frontier molecular orbitals together with conceptual DFT,⁶⁵ with the surprise that even though it is about similar stability, corroborated by the almost identical chemical hardness for species **A** for all systems, the same does not happen with the chemical potential, nor with electrophilicity, but contrarily to what was predicted recently by Martínez and Trzaskowski with Grubbs–Hoveyda type catalysts, where the higher the electrophilicity the higher the reactivity was.⁶⁶ Thus, a simple NH group modulates the reactivity, making systems **1** and **2** more electrophilic, explained by similar stabilization of both the HOMO and the LUMO.

Table 3 NBO charges for intermediates **A** and **D** for systems **1–5** with acetylene as substrate

	A				D			
	Ru	N	N(H)	(N)H	Ru	C1	C2	C3
1	−0.290	−0.399	−0.691	0.431	−0.300	−0.125	−0.486	−0.322
2	−0.323	−0.403	−0.675	0.416	−0.298	−0.125	−0.487	−0.322
3	−0.303	−0.396	—	—	−0.300	−0.124	−0.486	−0.323
4	−0.276	−0.396	—	—	−0.300	−0.125	−0.487	−0.322
5	−0.275	−0.392	—	—	−0.282	−0.129	−0.489	−0.322

Table 4 Frontier Molecular Orbitals (in a. u.), chemical potential (μ), chemical hardness (η), and Parr electrophilicity (ε) for intermediate **A** for systems **1–5** with acetylene as substrate

	HOMO	LUMO	μ	η	ε
1	−0.155	−0.086	−0.120	0.069	0.105
2	−0.157	−0.088	−0.122	0.069	0.109
3	−0.149	−0.081	−0.115	0.068	0.097
4	−0.146	−0.079	−0.113	0.067	0.095
5	−0.148	−0.081	−0.115	0.067	0.098



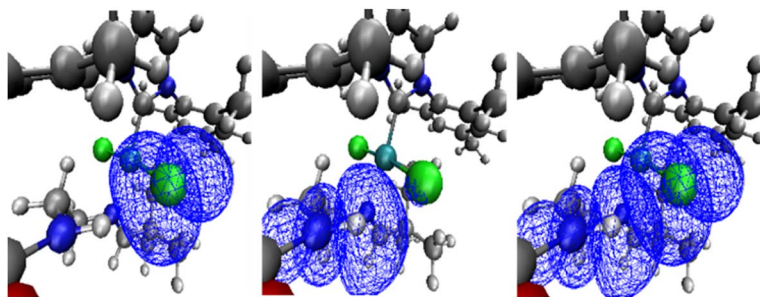


Fig. 3 NBO orbitals for precatalyst A for system **1** involved in the $\text{NH}\cdots\text{Cl}$ interaction: LP (left), BD* (middle) and overlap of the latter orbitals (right), see ESI[‡] for further details.

To unveil how the H-bonds of the halides and the organic moiety of the imine ligand *trans* to the NHC affect the system, 3D and 2D NCI plots in Fig. 4 and 5 were calculated using the program developed by Contreras-García *et al.*^{67,68}

The NCI plots help to understand here the different behaviours between catalysts, however only partially since energetics are still more important since interactions are less important than expected,^{69–71} especially compared to other olefin metathesis catalysts.⁷² Here, NCI interactions might not help to avoid decomposition, but help to influence the activation of the precatalyst. The NCI plots reveal how systems **1** and **2** have the H-bond between the H of the NH group and the nearest chloro ligand, which despite not being present in most of the following intermediates and transition states, does persist in some of them, although not so clearly. In addition, the bidimensional NCI plots in Fig. 5 clarify the difference in the first two systems compared to the other ones,⁷³ *i.e.* comparing systems **2** and **5**, with a remarkable sign (λ_2) ρ contribution for **2** in the region between -0.02 and -0.03 a. u. that for **5** is completely nonexistent.

Computational details

All the DFT static calculations were performed with the Gaussian16 set of programs.⁷⁴ All geometry optimizations were performed in gas phase using the BP86 functional of Becke and Perdew,^{75–77} corresponding to the GGA level, including corrections due to dispersion through Grimme's method (GD3 keyword in Gaussian16).^{78,79} The electronic configuration of the molecular systems was described with the double- ζ basis set with the polarization of Ahlrichs for main group atoms from Karlsruhe (Def2SVP keyword in Gaussian),^{80,81} whereas for ruthenium, the small-core quasirelativistic Stuttgart-Dresden effective core potential was introduced, altogether with an associated valence basis set (standard SDD keyword in Gaussian).^{82–84} The geometry optimizations were performed without symmetry constraints, with analytical frequency calculations for the characterization of the located stationary points. These frequencies were used to calculate unscaled zero-point energies (ZPEs) as well as thermal corrections and entropy effects at 353.15 K. Energies were obtained through single-point calculations on the optimized geometries from the BP86/Def2SVP~ssdd employing the B3LYP functional,^{85,86} and the cc-pVTZ basis set for main atoms,⁸⁷ while for ruthenium the same basis set as before. Also,



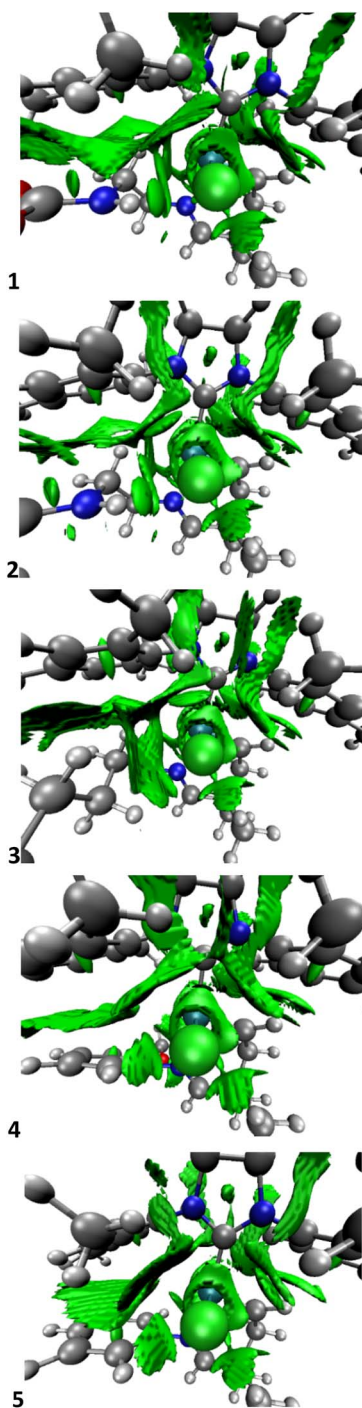


Fig. 4 NCI plots of the intermediate A for systems 1–5 (orientated like Fig. 1, and the whole picture is truncated for the sake of clarity of the metal centre, see ESI† for the NCI plots of the whole molecule).



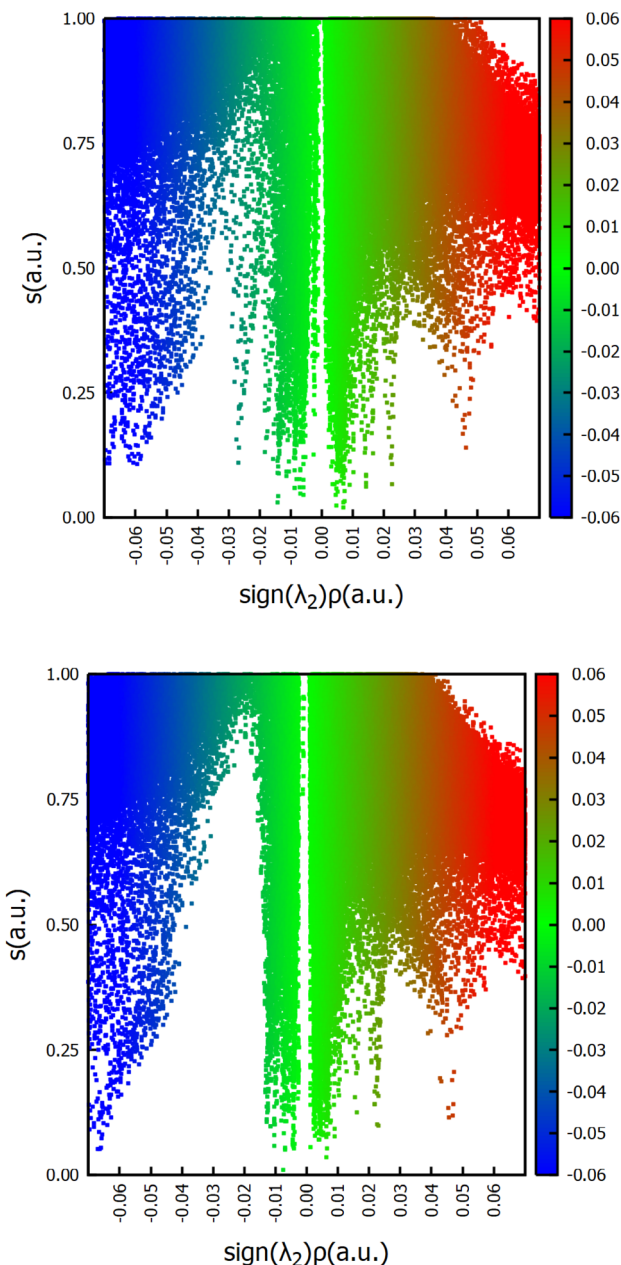


Fig. 5 2D-NCI plots of reduced density gradient (s) vs. $\text{sign}(\lambda_2)\rho$, in a.u., for the initial intermediate A for systems 2 (above) and 5 (below).

dispersion effects were introduced with the Grimme D3 correction term to the electronic energy. Furthermore, the solvent effects were estimated with the PCM model,^{88–90} and Gibbs free energies have been computed at 353.15 K and atmospheric pressure reaction conditions.

Conclusions

In a set of latent Ru based catalysts, the Ru–N bond cleavage is more favoured kinetically for systems **1** and **2**, but this does not correspond to the rds, but the opening of the metallacycle. These results might seem somewhat in disagreement with experiments because kinetically systems **1** and **2** have larger energy barriers, and not only in the Ru–N cleavage, but also in the next transition states, including the rds. Actually, this is what really favours the latent activity in polymerization at over 353.15 K for both systems, whereas it starts at 303.15 K for the other systems of the set of catalysts tested experimentally. In the phase of characterisation, the non-covalent interactions play a key role between the NH moiety and the closest chloride ligand. And probably the potential H-bonds of the NH fragment with any other atom during the reaction pathway may also slow down the olefin metathesis process in the following steps. However, this was not confirmed for the described intermediates. Further studies are ongoing to confirm the validity of the results extending the analysis towards the catalytic cycle, and with the aim to check if the solvent energy calculations at the B3LYP/cc-pVTZ~sdd level might either underestimate the kinetic Ru–N bond cleavage cost or overestimate the opening of the metallacycle.⁹¹

Author contributions

Artur Brotons-Rufes performed all the calculations, the visualization and wrote the original draft together with Albert Poater. Naeimeh Bahri-Laleh and Albert Poater performed the analysis, supervision, review and editing of the text.

Conflicts of interest

There are no conflicts to declare.

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