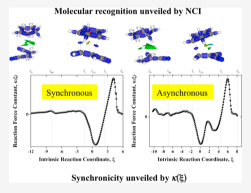
Real-Space Approach to the Reaction Force: Understanding the Origin of Synchronicity/Nonsynchronicity in Multibond Chemical Reactions

Published as part of The Journal of Physical Chemistry virtual special issue "Paul Geerlings Festschrift". Diana Yepes,[¶] Julen Munarriz,[¶] Daniel l'Anson, Julia Contreras-Garcia,* and Pablo Jaque*



ABSTRACT: In this article, we present a complementary analysis based on the reaction force $F(\xi)$ /reaction force constant $\kappa(\xi)$ and noncovalent interactions (NCI) index to characterize the energetics (kinetic and thermodynamics) and mechanistic pathways of two sets of multibond chemical reactions, namely, two double-proton transfer and two Diels-Alder cycloaddition reactions. This approach offers a very straightforward and useful way to delve into a deeper understanding of this type of process. While $F(\boldsymbol{\xi})$ allows the partition of the whole pathway into three regions or phases, $\kappa(\boldsymbol{\xi})$ describes how orchestrated are the bond-breaking and bondformation events. In turn, NCI indicates how the inter- and intramolecular bonds evolve. The most innovative aspect is the inclusion of the formation of the reactant complex along the pathway, which, by means of NCI, unveils the early molecular recognition and the comprehension of its role in determining the degree of the synchronicity/nonsynchronicity of one-step processes. This approach should be a



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useful and alternative tool to characterize the energetics and the mechanism of general chemical reactions.

INTRODUCTION

Multibond chemical reactions are those in which two or more covalent bonds are formed and/or broken. They can proceed either in a single kinetic step without an intermediate or in a stepwise mechanism, where a diradical or zwitterionic species is suggested as intermediate.¹ In the former case the bond breaking and/or formation can occur in unison, that is, synchronously, or through a highly asynchronous mode, currently defined as two-stage one-step mechanism; thus, a time gap between the bond-forming processes appears. Note that a complete gradation in asynchronicities lies between these two extremes, that is, fully synchronous to fully asynchronous processes, or even when the latter converts into a stepwise mechanism, which can be seen as an extreme case of asynchronicity.²⁻⁷ The most widely used criterion to characterize this mechanistic concept, for instance, in Diels-Alder reactions, has been the (a)symmetry of the emerging C-C bond distances at the transition state.⁸⁻¹⁰ However, this approach provides a continuum scale of (a)synchronicities, making it hard to rank the different flavors between synchronous to two-stage one-step mechanisms, or even the transition into a stepwise mechanism. In this context, the features of the fine structure of the reaction force constant, $\kappa(\boldsymbol{\xi})$ ¹¹ along the transition regions defined within the reaction force frame¹² have been proved to be directly related to the formation of the new bonds. Consequently, it has been used to reliably reveal the synchronicity level as well as to show how it is affected by the presence of catalysts¹³ or the solvent polarity in prototypical multibond chemical reactions, such as doubleproton transfer reactions^{14,15} and Diels-Alder cycloadditions.^{4–6,16–18}

The progress of single-step multibond chemical reactions can be characterized by following the motion along the minimum-energy pathway (also known as the intrinsic reaction coordinate (IRC) ξ , expressed in mass-weighted Cartesian coordinates^{19,20}) connecting the transition state (TS) to the reactants (**R**) and products (**P**). Generally, the reaction profiles are referred to the isolated reactants (see Figure 1a). However, different works have drawn attention to the key role the reactant complexes (RC) (also known as van der Waals or preorganized complexes) could be playing in a specific reaction pathway.²¹ This species corresponds to a minimum on the potential energy surface (as shown Figure 1b) and gives rise to

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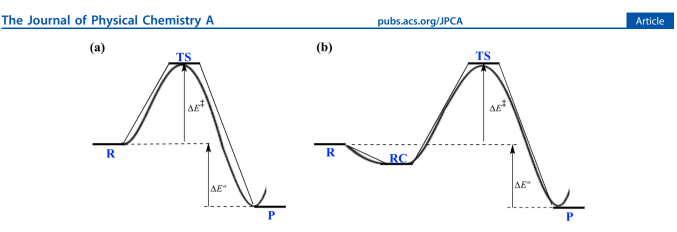
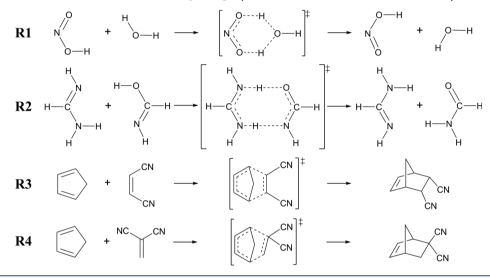


Figure 1. Prototypical potential energy profiles (a) without and (b) with RC.

Scheme 1. Double-Proton Transfer Reactions and [4 + 2] Cycloadditions Considered in This Study



the earliest molecular recognition between the reactants involving both the reactive centers (which will form the new covalent bonds) and any other fragments that could be assisting the initial assembling. Concomitantly, the non-covalent interactions (NCI) at the **RC** are mostly responsible for its stability and can even be strengthened along ξ until the proximities of the transition state. In this last region the covalent interactions become dominant, thus driving the process toward the final products.^{22–24} In this sense, the scope of these interactions in determining the initial orientation mode of the reactants can be responsible for the mechanistic issues that have been exclusively ascribed to the **TS**, such as the stereo- and regioselectivity, as well as the synchronicity/asynchronicity of multibond chemical reactions.^{17,25}

Therefore, a deeper understanding of the reaction mechanisms can be achieved by combining the classical IRCbased analyses with the specific role of noncovalent interactions (the so-called molecular recognition). The synergy between both approaches constitutes a valuable tool to provide important information into the origin of the regio- and/or stereochemical selectivity, paving the way toward the rational design of novel chemical systems (such as catalysts) with improved performance.^{25,26} We believe this approach will complement other methodologies readily available, such as Bonding Evolution Theory (BET), which constitutes a powerful tool to rationalize the electron reorganization along the molecular mechanism for a variety of chemical processes.^{27–33} Such methodology consists of the study of the Electron Localization Function (ELF) along the reaction coordinate, which makes it very useful to study covalent bonds and lone pairs reorganization. This way, we expect to complement it by adding the NCI framework and being able to also study the noncovalent interactions along the reaction pathway.

In the current work, we present a study of two doubleproton transfer reactions and two [4+2]- cycloadditions. They take place in a single kinetic step, but within each set, one reaction is fully synchronous, and the other one has a highly asynchronous mechanism (see Scheme 1). The main goal of this contribution is to scrutinize how the noncovalent/covalent interactions evolve along the reaction pathway. Special attention is paid to the key points defined by the reaction force model, to determine their role in the (a)synchronicity in both sets of reactions, going beyond the **TS**-symmetry based criterion.

THEORETICAL BACKGROUND

The Reaction Force and the Reaction Force Constant. For a conservative physical system, the negative gradient of a potential energy is a force, and the second derivative is a force constant. In analogy, for a chemical process that takes place in a single kinetic step, the reaction force $F(\xi)^{12}$ and the reaction

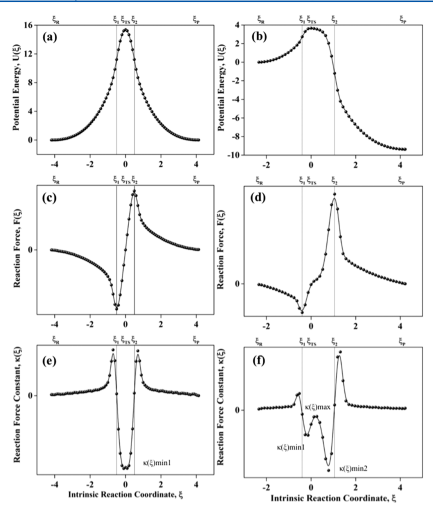


Figure 2. Generic profiles of (a, b) $U(\xi)$, (c, d) $F(\xi)$, and (e, f) $\kappa(\xi)$ along the intrinsic reaction coordinate ξ for a synchronous (left) and an asynchronous (right) one-step mechanism.

force constant $\kappa(\boldsymbol{\xi})^{11}$ can be, respectively, defined assimilating $\boldsymbol{\xi}$ to a displacement, as stated in eqs 1 and 2.

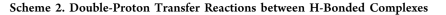
$$\mathbf{F}(\boldsymbol{\xi}) = -\frac{\partial U(\boldsymbol{\xi})}{\partial \boldsymbol{\xi}} \tag{1}$$

$$\kappa(\boldsymbol{\xi}) = \frac{\partial^2 U(\boldsymbol{\xi})}{\partial \boldsymbol{\xi}^2} = -\frac{\partial \mathbf{F}(\boldsymbol{\xi})}{\partial \boldsymbol{\xi}}$$
(2)

Figure 2a-f illustrates the corresponding generic profiles of $U(\boldsymbol{\xi})$, $\mathbf{F}(\boldsymbol{\xi})$, and $\kappa(\boldsymbol{\xi})$ for a fully synchronous (left panels) and a highly asynchronous two-stage (right panels) reaction, respectively. For both processes the potential energy $U(\boldsymbol{\xi})$ is quite similar in the proximities of the maximum, ξ_{TS} (which corresponds to the **TS**). On the contrary, the derivatives of $U(\boldsymbol{\xi})$ exhibit remarkable differences, which have been directly related with the (a)synchronicity degree of the mechanism. $^{5-7,15-18,34,35}$

 $F(\boldsymbol{\xi})$ and $\boldsymbol{\xi}$ are vectors pointing from reactants (**R**) to products (**P**). The reaction force profiles $F(\boldsymbol{\xi})$ (Figure 2c,d) are the result of the forces acting over the system at any point along $\boldsymbol{\xi}$. Notice that $F(\boldsymbol{\xi})$ is negative along the activation process (i.e., from **R** to **TS**) indicating that it is retarding in nature; therefore, energy is required. On the contrary, it is positive along the relaxation process (i.e., from **TS** to **P**) indicating that it is supporting the motion along $\boldsymbol{\xi}$; consequently, energy is released. It can be also noted that $F(\boldsymbol{\xi})$ presents two critical points, a minimum at ξ_1 and a maximum at ξ_2 . They split the process into three segments labeled as reactant (from ξ_R to ξ_1), transition (from ξ_1 to ξ_2), and product regions (from ξ_2 to ξ_P). Different studies have unveiled that, in the middle stage, most of the electronic events (such as bond breaking/forming and charges redistribution) are accentuated, while the geometrical changes (bond lengthening, rotations, etc.) are mainly emphasized during both sided regions.³⁶⁻⁴³

With respect to the reaction force constant profiles (Figure 2e,f), $\kappa(\boldsymbol{\xi})$ is positive in the structurally intensive first and third regions, whereas $\kappa(\boldsymbol{\xi})$ is negative along the electronically intensive regions, that is, transition regions. Therefore, we can establish a clear picture for both $U(\xi)$ profiles studied thus far: $\kappa(\boldsymbol{\xi})$ is negative within the transition region, and for the fully synchronous mechanism (left-panels) there is only one $\kappa(\boldsymbol{\xi})$ minimum $(\kappa(\boldsymbol{\xi})_{\min 1})$ (Figure 2e). On the contrary, for the highly asynchronous one (right panels) two minima ($\kappa(\boldsymbol{\xi})_{\min 1}$ and $\kappa(\boldsymbol{\xi})_{\min 2}$ connected by one negative maximum ($\kappa(\boldsymbol{\xi})_{\max}$) are revealed (Figure 2f). Previous studies have revealed that, for moderate asynchronous processes, $\kappa(\boldsymbol{\xi})$ displays a minimum with a shoulder, while for an extremely asynchronous reaction the negative maximum of $\kappa(\boldsymbol{\xi})$ becomes a positive maximum of $\kappa(\boldsymbol{\xi})$, which is indicative of a mechanism change from a one-step to stepwise one.^{6,7,15}



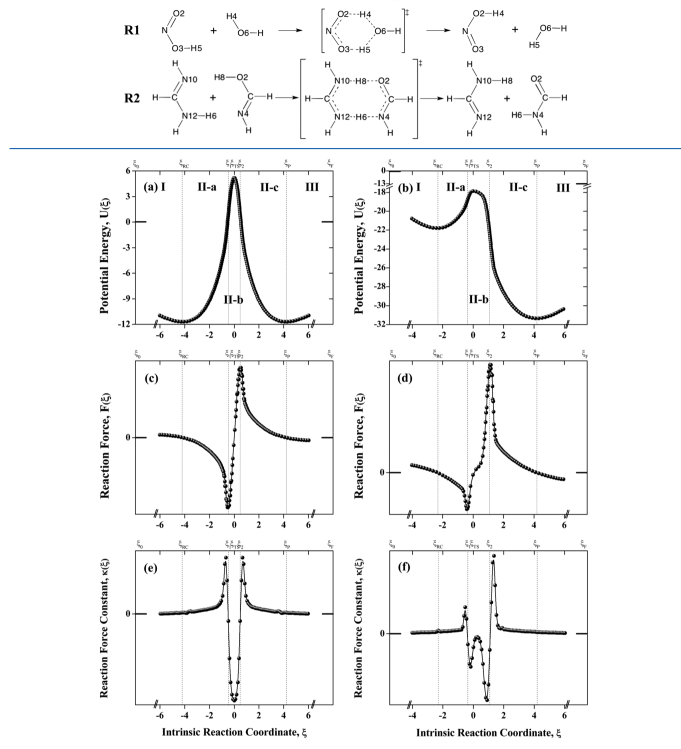


Figure 3. Profiles of $U(\xi)$, $F(\xi)$, and $\kappa(\xi)$ along ξ for synchronous R1 (left) and asynchronous R2 (right) double-proton transfer reactions.

The transition region goes beyond the traditional transition state, and its existence is experimentally supported by the transition-state spectroscopy of Zewail and Polanyi, which defines a continuum of transient, unstable states between perturbed forms of the reactants and products, where all bond breaking and formation processes occur.^{44,45}

Noncovalent Interactions (NCI) Index. During any multibond chemical reaction, namely, double-proton transfer and [4 + 2] cycloaddition reactions, intra- and intermolecular

bonds are expected to change. To follow those changes we resorted to the NCI index. NCI is a method for the characterization and visualization of interactions on the basis of the electron density and its derivatives, enabling the identification and classification of noncovalent interactions by means of the peaks that emerge in the reduced density gradient at low-density regions.⁴⁶ In particular, the procedure localizes minima of the reduced density gradient *s* (see eq 3), which

appear as a consequence of the interference between the different atoms that constitute a given system.

$$s = \frac{1}{2(3\pi^2)^{1/3}} \frac{|\nabla \rho|}{\rho^{4/3}}$$
(3)

Most commonly, these regions are located around electron density critical points (although there are some exceptions).⁴⁷ This enables the reconstruction of a three-dimensional (3D) image of the quantum theory of atoms in molecules (QTAIM),⁴⁸ where the relationship between critical points is recovered visually (i.e., the information commonly conveyed by bifurcation diagrams is retrieved directly by visual inspection of isosurfaces). It has proved to be a useful tool in the study of a broad range of noncovalent interactions, such as hydrogen and halogen bonds, as well as in the design of hybrid materials, among many others.^{49–51} The approach is able to reveal both attractive and repulsive interactions.

A color code is used to differentiate the type of interaction. Namely, strong attractive interactions, such as hydrogen bonds, are depicted in blue; very weak interactions, such as van der Waals, are shown in green, and strong repulsive interactions, like steric clashes, are colored in red. Since these surfaces appear for all bonding types (from strong to weak, from covalent to ionic),^{52,53} they are specially suitable for following reactivity studies in real space and compare them to the insight obtained from energy derivatives (reaction force/reaction force constant).

Computational Details. All stationary points on the potential energy profiles were fully optimized using the hybrid B3LYP exchange-correlation functional^{54,55} combined with the standard triple- ζ augmented with d- and p-type polarization functions, 6-311G(d,p,) basis set for double-proton transfer reactions. Instead, the range-separated hybrid with semiclassical London dispersion, the ω B97XD functional,⁵⁶ combined with the standard double- ζ augmented with dand p-type polarization and diffuse functions, 6-31+G(d,p)basis set was employed for Diels-Alder cycloadditions. These levels of theory were chosen due to their reported good performance in each type of multibond reaction.^{5,15,57} The nature of the stationary points in the potential energy surface was confirmed through harmonic vibrational analyses: zero imaginary frequency for energy minima and one for transition states. The associated eigenvectors of TSs were validated to correspond to the motion along the minimum energy path by using the IRC procedure^{19,20} with a step size of 0.10 amu^{1/2} bohr. The IRC procedure typically concludes in the RC. Nonetheless, to be able to completely move apart the reacting molecules we added a larger number of steps that were chosen following the projection of the IRC onto the distances between the reactive centers. These calculations were performed using the Gaussian09 suite of programs.⁵⁸

NCI plots were computed by means of the NCIPLOT software,⁵⁹ by using the respective monodeterminantal B3LYP/6-311G(d,p) and ω B97XD/6-31+G(d,p) wave functions for double-proton transfer and Diels–Alder reactions. A cutoff of 1.0 au in the density was chosen for the 3D visualization to ensure that both intra- and intermolecular bonds could be visualized simultaneously.⁶⁰ All *s* isosurfaces were represented, by taking an isovalue of 0.2 au. Since we wanted to study simultaneously both covalent and noncovalent interactions, an unusually large color range (sign(λ_2) ρ from

-0.5 to 0.5 au) was used. Electron density values at the bond critical point (BCP) were obtained with the AIMAll package.⁶¹

RESULTS AND DISCUSSION

The results are divided into two sections, one for each type of multibond chemical reaction. Energetics together with the degree of synchronicity will be first discussed, while the origin of the (a)synchronicity will be thus understood by scrutinizing how interactions evolve along ξ .

Double Proton Transfer: Synchronous Versus Asynchronous Mechanisms. The double-proton transfer processes considered in the present study consist of reactions involving hydrogen-bonded complexes (as RC) between the weak nitrous acid (HNO₂) in its *cis* rotamer and water (R1) and between formamidic acid (FMA) and formamidine (FI) (R2) as displayed in Scheme 2. Notice that the atom labels were included in the scheme so as to facilitate the further discussion on the density at the BCPs.

Notice that, while **R1** proceeds via a symmetrical **TS**, **R2** does so via an asymmetrical **TS**. Figure 3a-f displays the $U(\boldsymbol{\xi})$ (a, b), $\mathbf{F}(\boldsymbol{\xi})$ (c, d), and $\kappa(\boldsymbol{\xi})$ (d-f) profiles for **R1** (left panels) and **R2** (right panels), respectively. In addition to the minimum energy path provided by the IRC calculations, the formation (in reactant side) and breaking (in product side) of H-bonded complexes (i.e., **RC**) is also included as suggested in Figure 1b. Concomitantly, the profile can be divided into three phases: (I) the initial formation of the **RC**, (II) the double-proton exchange reaction, and (III) the separation of the products.

The initial changes, part (I) from ξ_0 to ξ_{RC} , indicate the structural modifications that molecules undergo so as to approach each other through the two H-bonding interactions between the atoms $O_2 \cdots H_4/H_5 \cdots O_6$ in **R1** and $N_{10} \cdots H_8/H_6 \cdots N_4$ in **R2**, thus leading to the formation of the H-bonded complexes located at ξ_{RC} . The formation of the H-bonded complex is a barrier-free process in which energy is released, being more stable in **R2** than the corresponding one in **R1** (21.8 vs 11.7 kcal mol⁻¹); that is, the FMA…FI H-bonded complex is more stable than the HNO₂…H₂O complex.

After the formation of RC, the double-proton exchange reaction takes place and is identified as part (II) in Figure 3, which, in turn, is divided into three regions: II-a, II-b, and II-c. Note that it corresponds to the description of the reaction force analysis given in Figure 2a-f. Therefore, during part II-a (from $\xi_{\rm RC}$ to ξ_1), the so-called reactant region, structural rearrangements are predominant over the electronic ones. The amount of energy $(U(\xi_1) - U(\xi_{RC}))$ needed to overcome the increasing hindrance to the structural changes in part II-a is higher in **R1** than the analogous in **R2** (11.2 vs $2.7 \text{ kcal mol}^{-1}$). With respect to part II-b (i.e., from ξ_1 to ξ_2) it corresponds to the transition region, where the electronic reorganizations are enhanced. The involved energy $(U(\xi_2) - U(\xi_1))$ can be decomposed into two terms: $(U(\xi_{\rm TS}) - U(\xi_1))$ and $(U(\xi_2) - U(\xi_2))$ $U(\xi_{\rm TS}))$, which are, respectively, 5.7 & -5.7 kcal mol⁻¹ for R1 and 1.2 & -5.4 kcal mol⁻¹ for R2. While the first component is positive in accord with a retarding force, the second one is negative, since a positive force is driving the process. Consequently, energy is required and released, respectively. In agreement with this partition, the activation energy referred to RC can be written in terms of two contributions, the amount of energy of part II-a and the first component of part II-b; that is, $\Delta E_{act} = U(\xi_{TS}) - U(\xi_{RC}) = U(\xi_1) - U(\xi_{RC}) + U(\xi_{RC})$ $U(\xi_{\rm TS}) - U(\xi_1)$. The activation energies and contributions are

Key Point along ξ	R1	R2
ξ ₀	H5 N1 O3 H7	N10 N12 N12 H6
ξrc		
ξı		
$κ(ξ)_{min1}$		
к(ξ) _{max}		
$\kappa(\xi)_{min2}$		
ξ2		
ξp		

Table 1. Plots of the NCI Index along the Key Points for R1 and R2

16.9 (11.2 and 5.7) kcal mol⁻¹ for **R1** and 3.9 (2.7 and 1.2) kcal mol⁻¹ for **R2**. We thus noted that **R2** is kinetically more favored than **R1** and that the energy barriers are mainly controlled by structural rearrangements. Moreover, the kinetic feasibility of **R2** is also provided by a lower energy associated with electronic reorganizations. Both contributions to the energy barrier are determined by the strength of the hydrogenbonding interactions unveiled in part I.

Finally, part II-c (ξ_2 to ξ_P) consists of the structural rearrangements that lead toward the formation of the products, the so-called conformational relaxation. The amount of energy released ($U(\xi_P) - U(\xi_2)$) is -11.7 and -8.0 kcal mol⁻¹ for **R1** and **R2**, respectively. Notice that the addition of the last four

terms defines the thermodynamic driving force. Again, the thermodynamic feasibility of **R2** can also be explained on the stronger H-bonding interactions between FMA and FI that promote the proton exchange to form the new H-bonded complex between formamide (FM) and formamidine (FI).

Finally, part III describes the barrier-free dissociation of the formed H-bonded complex. Because **R1** has a perfectly symmetric profile, the energy needed $(U(\xi_F) - U(\xi_P))$ to move apart the interacting fragments is 11.7 kcal mol⁻¹, while for the FM…FI complex of **R2** 18.3 kcal mol⁻¹ are required. Comparing the results in parts I and III, we can conclude that the H-bonded complex between FMA and FI is more stable than that between FM and FI; that is, N…HO/NH…N

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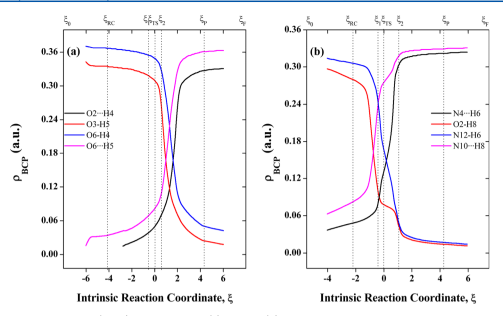


Figure 4. Electron density at the BCP (in au) along ξ for R1 (a) and R2 (b).

interactions are more stable than $NH\cdots O/N\cdots HN$ ones. Consequently, the forward reaction is kinetically more favored than the reverse one.

As it can be seen in Figure 3c-f, the $F(\xi)$ and $\kappa(\xi)$ reveal certain similarity along the structurally intensive regions, that is, reactants and products regions. The differences are noted along the electronically intensive regions, that is, the transition regions. On one hand, a linear and a nonlinear behavior can be noted for R1 and R2, respectively. On the other hand, the barrier width, $\xi_2 - \xi_1$, is broader for R2 than R1 (see Figure 3c,d, respectively). This suggests that the primitive processes associated with the breaking and formation of H-bonding interactions is occurring in a different manner.⁶² The abovementioned observations are confirmed by the $\kappa(\boldsymbol{\xi})$ profiles, which are markedly different in the respective transition regions. This fine structure shows a single minimum of $\kappa(\boldsymbol{\xi})$ within this zone for R1 (see Figure 3e). On the contrary, the shape of $\kappa(\boldsymbol{\xi})$ along the transition region of **R2** exhibits a local negative maximum connecting to two minima of $\kappa(\boldsymbol{\xi})$ located sided (see Figure 3f). Therefore, on the basis of these patterns the double-proton exchange mechanism can be classified as synchronous for R1 and asynchronous for R2, suggesting that the asynchronous character of R2 could also explain its kinetical feasibility.

By matching the results, we can conclude that a less stable symmetric H-bonded complex is associated with a higher energy barrier, where protons are synchronously transferred. Such highly stable asymmetric H-bonded complex is thus related with a lower energy barrier indicating that enhanced bonding interactions promote dynamically delocalized protons, and consequently, the protons are asynchronously exchanged.

As previously introduced, the noncovalent interactions were thoroughly studied by means of the NCI index on the key points along ξ defined by the $F(\xi)/\kappa(\xi)$ analysis (see Table 1).

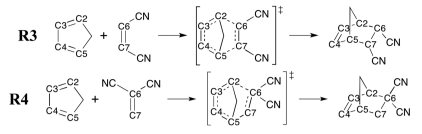
In **R1** the NCI profile of the N–O bonds in HNO_2 shows large asymmetric *s* isosurfaces with a flat shape toward the most electropositive atom (N) and a round one in the proximities of the electronegative one (O).⁴⁷ The character of these bonds is interconverted from single into double along the

transformation, going all of them through an intermediate bond order at the **TS**. Along the protons exchange reaction, the main changes are noted in both the covalent O–H bonds and the H…O intermolecular noncovalent interactions. Namely, at the initial state at ξ_0 , a weak hydrogen bonding is established between H5 and O6 with localized dispersive interactions (see Table 1). The covalent O–H bond involved in the O3–H5 is weaker (red-shift) than the O6–H4 (which is not involved in any H bonding). This feature is reflected in the electron density at the BCP as displayed Figure 4: $\rho_{\rm BCP}(O3-H5) = 0.345$ au versus $\rho_{\rm BCP}(O6-H4) = 0.371$ au.

The formation of the H-bonded complex at $\xi_{\rm RC}$ is guided by new dispersive interactions established between N-O2 double bond and H4. These interactions (depicted in green in Table 1) are highly delocalized. Note that this interaction does not lead to a bond critical point until the proximity of the minimum force at ξ_1 ($\rho_{BCP}(O2-H4) = 0.039$ au at ξ_1), and, as highlighted before, NCI is a useful tool to identify noncovalent interactions at a very much lower value of density.^{46,52} At $\xi_{\rm RC}$, the system has two H-bonding interactions, O6-H5 and O2-H4. Nonetheless, they are quite different in nature. Specifically, from Figure 4a we can see that O6-H5 falls into a classical hydrogen bond (HB) ($\rho_{BCP} = 0.042$ au) and a well-defined disklike shape s isosurface (Table 1), indicative of a localized interaction. On the contrary, O2-H4 corresponds to a delocalized interaction, with an extended and diffuse s isosurface, and a lower density at the BCP ($\rho_{\rm BCP}$ = 0.015 au). This evolves progressively into a more localized interaction achieving a classical HB character within the first part of the transition region. Such changes are not occurring at the expense of the covalent O-H bonds (note the flatness of their BCP densities in Figure 3a).

As it can be seen in Table 1, the TS is characterized by nearly symmetrical NCI profiles around O6, O2, and O3 atoms, unveiling that the strength of both intramolecular and intermolecular interactions are rather similar ($\rho_{\rm BCP}(O6-H4) = 0.349$ au vs $\rho_{\rm BCP}(O3-H5) = 0.310$ au and $\rho_{\rm BCP}(O2-H4) = 0.050$ au vs $\rho_{\rm BCP}(O6-H5) = 0.082$ au). The fact that both chemical events happen at the same time highlights the symmetric nature of the reaction from the electronic

Scheme 3. Diels-Alder Reactions between Cyclopentadiene and Cyanoethylenes



viewpoint. The reaction phase ends with the simultaneous transfer of H4 from O6 to O2 ($\rho_{\rm BCP}({\rm O6-H4}) = 0.050$ au vs $\rho_{\rm BCP}({\rm O2-H4}) = 0.327$ au) and H5 transfer from O3 to O6 ($\rho_{\rm BCP}({\rm O3-H5}) = 0.025$ au vs $\rho_{\rm BCP}({\rm O6-H5}) = 0.361$ au) at $\xi_{\rm P}$, forming a H-bonded complex.

The NCI analysis also reveals the synchronous character of the reaction (see Table 1). The *s* profiles for the main reaction points are symmetrically equivalent. Similarly, the **TS** structure has mirror symmetry. This pattern along ξ is originated in the encounter mode between HNO₂ and H₂O.

The double-proton transfer reactions can be seen as selfneutralization acid—base reactions. Thus, the synchronous or asynchronous character can be understood on the basis of the difference of the respective pK_a of the molecules that form the H-bonded complexes. In this context, the reaction **R1** is given by the H-bonded complexes between HNO₂ and H₂O, whose aqueous pK_a values are 3.29 and 14, respectively.⁶³ On the contrary, **R2** is controlled by the H-bonded complex between FMA and FI, with corresponding pK_a values of 14 and 28.5 (measured in dimethyl sulfoxide (DMSO)).^{64,65}

The reaction **R2** (see Scheme 2) proceeds by means of an asymmetric **TS**. Specifically, within the **TS** an amidinic and a formamidic proton (H6 and H8) migrates to a nitrogen (N4 and N10), respectively. As a consequence of the difference in the pK_a values of these moieties, the two proton migrations occur at different stages of the IRC, leading to a highly asynchronous process. In particular, the formamidic proton H8 migrates sooner (before the **TS**) than the amidinic one H6, whose transference takes place after the **TS** as displayed in the last column of **Table 1**. Note that the mere inspection of the **TS** would lead to the wrong conclusions.

From the NCI profiles included in Table 1, we can observe that strong intermolecular interactions are early recognized at ξ_{00} (though they are different in nature than in **R1**, since there are two localized dispersive interactions). Once again, intramolecular bonds within each backbone present a polar character.

At the **RC**, the strength of intra- (O2–H8 and N12–H6) and intermolecular bonds (N10…H8, N4…H6) shows some differences (see Figure 4b); the density at the BCP of O2–H8 bond is lower than N12–H6 ($\rho_{\rm BCP}$ (O2–H8) = 0.280 au vs $\rho_{\rm BCP}$ (N12–H6) = 0.306 au), which explains the lability of H8. $\rho_{\rm BCP}$ (O2–H8) decreases monotonically as the reaction advances, until the minimum force is reached at ξ_1 . After this, it stays constant along the transition region. From this decoupled movement of H8 arises the first minimum of $\kappa(\xi)$, $\kappa(\xi)_{min1}$.¹⁵

On the contrary, the strength of the N12–H6 bond monotonically decreases, but until the maximum force at ξ_2 , indicating that it breaks in a more advanced point of the IRC, the motion of H6 originates the second minimum of $\kappa(\boldsymbol{\xi})$, $\kappa(\boldsymbol{\xi})_{\min}$.¹⁵ At this degree of progress, the reaction the electron

density of both bonds are equivalent ($\rho_{BCP}(O2-H8) = 0.042$ au vs $\rho_{BCP}(N12-H6) = 0.047$ au), and they remain constant afterward, indicating that the classical HBs are formed.

Like the previous reaction, in the case of R2, there are two hydrogen bonds with different strengths at ξ_{RC} (Figure 4b): while H6 is strongly bonded to N12 ($\rho_{BCP} = 0.306$ au) and just forming a weak HB with N4 ($\rho_{BCP} = 0.048$ au), H8 establishes weaker intramolecular bond to O2 ($\rho_{\rm BCP}$ = 0.280 au) and stronger HB to N10 (ρ_{BCP} = 0.082 au). Thus, H8 is transferred earlier than H6 with a high degree of synchronicity between Hbond breaking and formation within the $N10{\cdots}H8{\cdots}O2$ backbone (see the profile at $\xi = \kappa(\xi)_{\min 1}$ in Table 1). This pattern is slightly modified, but emphasizes that the new N10-H8 bond begins to be formed close to $\kappa(\xi)_{\min 2}$ (see Table 1). Hence, contrarily to R1, the hydrogen transfers occur before (H8) and after (H6) the TS. Then, we can identify a transition region in the range of $[\kappa(\xi)_{\min 1} - \kappa(\xi)_{\min 2}]$ (see Figure 3b). The whole transition region in between corresponds to changes of HBs strength without significant hydrogen movements. However, in this case the TS is separated from the highly symmetric structure, which corresponds to $\kappa(\xi)_{max}$. At $\kappa(\xi)_{\max}$, we can see that both hydrogen atoms are found at a similar electron environment with respect to both fragments (see Table 1). Note that the electron density profiles are different at the reactant side than at the product side. This pattern suggests that the closeness in the former region is indicative that through-bond interactions (delocalization) are predominant, while the separation in the latter side is indicative that *through-space* interactions (localization) are predominant. While this observation explains the kinetics feasibility in the forward direction of R2, the behavior of NCI profiles (strength of hydrogen-bonding interactions) along the transition region explains the degree of asynchronicity in the mechanism.

Contrary to other studies, where the chemical arrangement occurs closely around the **TS**, in this case, the analysis of second derivatives coincides with the electron density analysis in finding the range of proton transfer (outside the minimum force-maximum force region!) and to identify the **TS** as a region of transition, where all interactions are weak.

Diels–Alder Cycloadditions: Synchronous Versus Asynchronous Mechanisms. Another multibond reaction analyzed in the present study is the Diels–Alder⁶⁶ cycloaddition of an alkene (or dienophile) to a diene. These processes are widely known for being a very useful synthetic tool in organic chemistry to coalesce two pairs of C–C atoms through new single bonds to produce cyclic or bicyclic adducts. The reactions under study consist of the addition of a symmetrically and an asymmetrically disubstituted dienophile to the cyclopentadiene. Specifically, we considered the addition of *cis*-1,2-dicyanoethylene (R3) and 1,1-dicyanoethylene (R4),

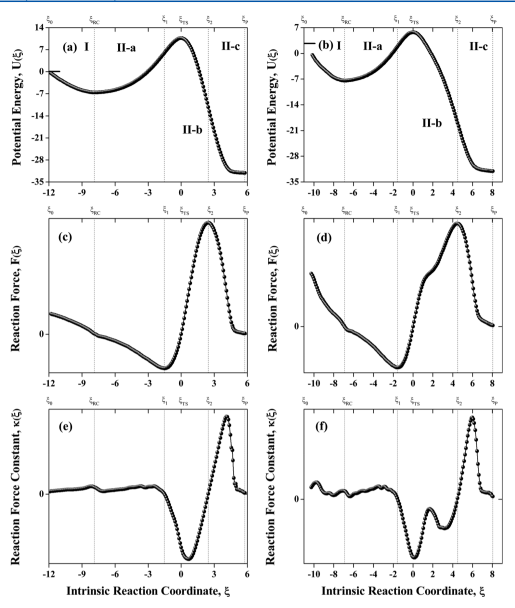


Figure 5. Profiles of $U(\xi)$, $F(\xi)$, and $\kappa(\xi)$ along ξ for synchronous R3 (left) and asynchronous R4 (right) Diels-Alder cycloadditions reactions.

as displayed in Scheme 3. Notice that the atoms labels are included for the ulterior density analysis at the BCP along ξ .

The addition of the dienophiles to cyclopentadiene is suprafacial, being the *endo* isomer generally generated. While **R3** proceeds via symmetrical **TS**, **R4** does so via an asymmetrical one. Figure 5a-f shows the profiles of $U(\xi)$ (a, b), $F(\xi)$ (c, d), and $\kappa(\xi)$ (e, f) for **R3** (left panels) and **R4** (right panels), respectively. As for the previous set of reactions, in addition to the minimum energy path provided by IRC calculations, we included (in the reactant side) the formation of the reactive complexes (i.e., **RC**) as suggested Figure 1b. Concomitantly, the profile can again be divided into two phases: (I) the initial formation of **RC** and (II) the cycloaddition reaction, which leads to the formation of sixmembered carbocyclic products.

As in the case of the double-proton exchange reactions, in the first phase (I), that is, from ξ_0 to ξ_{RC} , the formation of the van der Waals complex between diene and dienophile takes place, in this case through an *endo* approach and interacting C2…C6/C5…C7 atoms in both **R3** and **R4** (see Scheme 3). The formation of **RC** is also a barrier-free process and is slightly more favored in **R4** than in the corresponding **R3** reaction (7.4 vs 6.6 kcal mol^{-1} , respectively).

After the formation of **RC**, the cycloaddition takes place, which is labeled as part (II) in Figure 5, and this region is again partitioned into three zones: II-a, II-b, and II-c. This part corresponds to that provided by the IRC calculations and described in Figures 2a–f.

Therefore, the amount of energy required to overcome the structural distortions in the so-called reactant region (from $\xi_{\rm RC}$ to ξ_1) is higher in **R3** than the analogous in **R4** (12.5 vs 9.2 kcal mol⁻¹). The two energy terms ($U(\xi_{\rm TS}) - U(\xi_1)$) and ($U(\xi_2) - U(\xi_{\rm TS})$) involved along the transition region (part II-b) that are associated with electronic reorganizations are, respectively, 4.8 and -21.2 kcal mol⁻¹ for **R3** and 4.0 and -24.7 kcal mol⁻¹ for **R4**. Consequently, the activation energies and contributions with respect to **RC** are 17.3 (12.5 and 4.8) kcal mol⁻¹ for **R3** and 13.2 (9.2 and 4.0) kcal mol⁻¹ for **R4**. As for the previous set, we noted that the barrier is mainly controlled by structural rearrangements rather than by

Table 2. Plots of the NCI Index along the Key Points for R3 and R4	Table 2. Plo	ts of the NCI	Index along	the Key	Points for	R3 and R4
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Key Point along ξ	R3	R4
ξ ₀	C4 C5 C3 C2 C7 C7 C6	
ξ _{RC}		
ξı		
$κ(ξ)_{min1}$		
κ(ξ) _{max}		
$\kappa(\xi)_{min2}$		
ξ2		
ξp	-	

electronic reorganizations, being **R4** the most kinetically feasible reaction. Also notice that the first part of the relaxation process (i.e., from $\xi_{\rm TS}$ to ξ_2) is more favored in **R4** than in **R3**, by ~3 kcal mol⁻¹. This suggests a difference in the degree of the advance of the new C–C single bonds. This can be seen in Table 2, which reveals that the nonsubstituted C atom of the 1,1-dicyanoethylene reacts earlier than the substituted one.

At the final stage, (part II-c, from ξ_2 to ξ_P), the amount of energy released in the conformational relaxation to finally lead to the formation of the carbocyclic product is more highly favored for **R3** than for **R4** (-20.9 vs -13.1 kcal mol⁻¹). This, again, suggests that the formation of new C–C bonds is proceeding in a more decoupled manner in R4 than in R3. As already mentioned, the sum of the last four terms provides the thermodynamic driving force. In the case of the cycloadditions, it is found that R3 presents a slightly higher thermodynamic feasibility than R4 (-25.5 vs -24.6 kcal mol⁻¹). It is interesting to note that the exothermicity is quite similar for both reactions, while the activation barriers are well-differentiated. Therefore, these reactions cannot be explained in terms of the Bell-Evans-Polanyi principle^{67,68} but by the

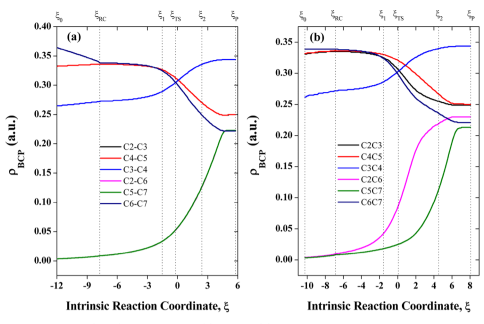


Figure 6. Electron density at the BCP (in au) along ξ for R3 (a) and R4 (b).

mechanistic concept of synchronicity or nonsynchronicity as will be discussed in the next paragraphs.

The $F(\boldsymbol{\xi})$ unveils a certain resemblance along the structurally intensive zones, that is, II-a and II-c. On the contrary, the differences are evidenced along the transition regions, that is, II-b zone regions. While $F(\boldsymbol{\xi})$ increases linearly along the transition region in R3, it does so nonlinearly in R4. Once again, the energy width, $\xi_2 - \xi_1$, is broader for **R4** than for **R3** (see Figure 5c,d, respectively). The primitive processes associated with the formation of new C-C single bonds are occurring in a different manner, as confirmed by the $\kappa(\boldsymbol{\xi})$ profiles that are markedly distinctive in the respective II-b zone. The fine structure shows a single minimum of $\kappa(\boldsymbol{\xi})$ along the transition region for R3 (see Figure 5e), whereas for R4 it exhibits a local negative maximum connecting the two minima of $\kappa(\boldsymbol{\xi})$ located on each side (see Figure 5f). On the basis of these patterns the formation of new bonds is fully or slightly orchestrated in R3, while it is nonorchestrated in R4 (or twostage one-step mechanism), suggesting that the asynchronous character of R4 could also explain its kinetic feasibility.

The difference with the double-proton transfer reactions discussed above deserves further discussion. For the proton transfer reactions, it was noted that a large difference in the stabilization energies of the H-bonded reactive complexes referred to the isolated molecules took place. This early recognition was proposed as responsible for both the kinetic feasibility and the mechanistic issues as synchronicity or nonsynchronicity. In spite of the fact that, for the case of the Diels–Alder reactions, the difference in the stability of the preorganized complexes referred to the isolated molecules is rather small, this cannot explain the kinetic feasibility, but whether it can give an explanation of the synchronicity patterns in the mechanism, going beyond the (a)symmetry criterion.

The noncovalent interactions were thoroughly studied by means of the NCI index on the key points along ξ defined by $F(\xi)/\kappa(\xi)$ analysis (see Table 2).

The Diels–Alder cycloadducts are built as a consequence of the formation of two new C–C single bonds (C2–C6 & C5–C7) and a C–C double bond (C3–C4) at the expense of

breaking two C-C double bonds (C2-C3 & C4-C5). As it can be seen in Table 2, the NCI profiles for both reactions at the initial stage (from ξ_0 to ξ_{RC}) allow to identify the covalent bonds (single and double character) in both diene and dienophile backbones. This feature is revealed by the electron density analysis, as displayed in Figure 6a,b (note that the pink/green and red/black lines are superposed in Figure 6a), which, as expected, shows higher densities in the C-C double bonds ($\rho_{BCP}(C2-C3) = \rho_{BCP}(C4-C5) = \rho_{BCP}(C6-C7) =$ 0.336 au for both R3 and R4) than in the C-C single bonds $(\rho_{\rm BCP}(\rm C3-C4) = 0.272$ au for both **R3** and **R4**) at $\xi_{\rm RC}$. These moieties are stabilized by weak dispersive van der Waals interactions, as they are brought closer to each other, exhibiting the typical diffuse s profile and low electron densities as displayed in Figure 6a,b ($\rho_{BCP}(C2-C6)$ = $\rho_{\rm BCP}(\rm C5-C7) = 0.009$ au for both R3 and R4) at $\xi_{\rm RC}$. Notice that the s profile is more extended for R4 than for R3 and that a localized interaction is observed between the unsubstituted C atom (C7) of dienophile and C5 of the diene. This pattern, on the one hand, explains the slight stabilization of the initial complex of R4 and, on the other hand, the identified initial recognition between C5 and C7 atoms that can promote the asynchronicity in the formation of new C-C bonds unveiled by the $\kappa(\boldsymbol{\xi})$ analysis. This result deserves to be highlighted, since depending on the strength of this type of interaction, it can reveal the origin of the degree of synchronicity versus nonsynchronicity, beyond a criterion based on the (a)symmetry of the incipient C–C bonds at the TS.⁵

After the formation of the **RC** the electron density variations revealed by the NCI profiles are very subtle until the force minimum is reached at ξ_1 . One of them consists of the symmetrically and asymmetrically strengthening of noncovalent interactions, that is, C2–C6 and C5–C7, for **R3** and **R4**, respectively. Interestingly, the small increase in the electron density at their BCP is produced at the expense of the intramolecular regions, since double bonds are slightly weakened.

Along the transition region (from ξ_1 to ξ_2), weaker bonds start to noticeably become stronger, while the double-bond

character becomes progressively weaker. This is indicative of the electronic reorganization taking place. Here, note that the tendencies are closer to the unison in R3, while the changes are decoupled in R4. Just as in the double-proton exchange, the former can be ascribed to the *through-space* (localization) interactions, whereas the latter can be related to the throughbond (delocalization) interactions. At the beginning of the transition zone, from ξ_1 to ξ_{TS} , both new C–C bonds can be considered to start forming in R3, whereas only one (between the unsubstituted C7 of the dienophile and C5 of the diene) can be considered in R4. This feature is unveiled by the presence of $\kappa(\boldsymbol{\xi})_{\min 1}$ in both reactions, while the new C-C bonds are strengthened along the second part of the transition region, from ξ_{TS} to ξ_2 , in **R3**. On the contrary, just the first new C-C bond makes stronger and the other starts forming in R4 giving rise to the negative maximum of $\kappa(\boldsymbol{\xi})_{\text{max}}$ and the second new C-C bond becomes stronger before leaving the transition region originating the second minima of $\kappa(\boldsymbol{\xi})$, that is, $\kappa(\boldsymbol{\xi})_{\min 2}$. This is in agreement with the energetics analysis aforementioned.

At the end, the formation of the bonds connecting the diene and the dienophile in R3, C2–C6 and C5–C7, progresses identically (i.e., fully symmetrically), and the resulting bonds are also identical. However, the new double bond, C3–C4, donates some extra electron density to the adjacent single bonds (C2–C3 & C4–C5), providing them with a slightly higher electron density than that of a standard single bond. This constitutes a partial measure of conjugation. The main electronic changes are those related to C3–C4, which undertakes an evolution from simple to double character, finally, acquiring the same electron density as the initial C–C double bonds.

A decrease in the C6–C7 electron density is observed as a C–C double bond evolves to a C–C single bond. These observations are the same in **R4** but in a decoupled manner.

As a final discussion, we want to compare previous BETbased studies applied in Diels-Alder reactions with the current approach. These have distinguished several phases (more than three phases as in $F(\xi)$ analysis), which are characterized by a depletion and formation of the double bonds character and formation of C–C single bonds.^{27,28} However, in those studies the degree of synchronicity/nonsychronicity has not received much attention, while these mechanistic concepts together with the evolution of noncovalent and covalent interactions (by means of the NCI) along ξ are the main focus of attention in the current approach. On the one hand, possibly, it could be interesting to combine BET with the $F(\boldsymbol{\xi})/\kappa(\boldsymbol{\xi})$ -based analysis in future investigations. On the other hand, the current approach presents some limitation, since it is based on an static view; therefore, the inclusion of the dynamic effects could change some conclusion as was recently pointed out by Longo and co-workers.69

CONCLUSIONS

In the present study our focus of attention has been to find the origin of an important issue in the reaction mechanism, that is, synchronicity or nonsynchronicity, in prototypical multibond chemical reactions, well beyond the criterion based on the symmetry at the **TS**. We considered two double-proton transfer and two cycloaddition reactions that proceed via a symmetrical and asymmetrical **TS** in each set of reactions. The study was performed by means of combining the reaction force/reaction force constant frame and noncovalent inter-

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actions index. The $\kappa(\boldsymbol{\xi})$ profile along the transition region has been already proposed as a suitable indicator of the degree of (a)synchronicity in multibond reactions depending on its fine structure along the transition region: one minimum of $\kappa(\boldsymbol{\xi})$ unveils a fully or slightly synchronous process, while two minima of $\kappa(\boldsymbol{\xi})$ connected by a negative maximum of $\kappa(\boldsymbol{\xi})$ does for asynchronous or two-stage one-step mechanism. Special attention has been paid on unraveling the role played by the early molecular recognition in determining the degree of (a)synchronicity in one-step mechanism. Our results showed that the early intermolecular interaction in the preorganized complexes can predetermine and solve this mechanistic issue, where a differential early intermolecular recognition between the reactive atomic centers was stressed along the activation process, so determining a highly asynchronous character in the mechanism, the so-called two-stage one step mechanism. Consequently, the real-space approach to the reaction force is a useful tool to characterize the kinetic, thermodynamic driving force, and mechanistic aspects of multibond chemical reactions.

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Author Contributions

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Notes

The authors declare no competing financial interest.

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REFERENCES

(1) Dewar, M. J. S. Multibond Reactions Cannot Normally Be Synchronous. J. Am. Chem. Soc. 1984, 106, 209–219.

(2) Black, K.; Liu, P.; Xu, L.; Doubleday, C.; Houk, K. N. Dynamics, transition states, and timing of bond formation in Diels–Alder reactions. *Proc. Natl. Acad. Sci. U. S. A.* **2012**, *109*, 12860–12865.

(3) Xu, L.; Doubleday, C. E.; Houk, K. N. Dynamics of 1,3-Dipolar Cycloadditions: Energy Partitioning of Reactants and Quantitation of Synchronicity. J. Am. Chem. Soc. **2010**, *132*, 3029–3037.

(4) Yepes, D.; Murray, J. S.; Pérez, P.; Domingo, L. R.; Politzer, P.; Jaque, P. Complementarity of reaction force and electron localization function analyses of asynchronicity in bond formation in Diels–Alder reactions. *Phys. Chem. Chem. Phys.* **2014**, *16*, 6726–6734.

(5) Yepes, D.; Valenzuela, J.; Martínez-Araya, J. I.; Pérez, P.; Jaque, P. Effect of the exchange–correlation functional on the synchronicity/ nonsynchronicity in bond formation in Diels–Alder reactions: a reaction force constant analysis. *Phys. Chem. Chem. Phys.* **2019**, *21*, 7412–7428.

(6) Yepes, D.; Donoso-Tauda, O.; Pérez, P.; Murray, J. S.; Politzer, P.; Jaque, P. The reaction force constant as an indicator of synchronicity/nonsynchronicity in [4 + 2] cycloaddition processes. *Phys. Chem. Chem. Phys.* **2013**, *15*, 7311–7320.

(7) Politzer, P.; Murray, J. S.; Jaque, P. Perspectives on the reaction force constant. *J. Mol. Model.* **2013**, *19*, 4111–4118.

(8) Caramella, P.; Houk, K. N.; Domelsmith, L. N. The dichotomy between cycloaddition transition states calculated by semiempirical and ab initio techniques. *J. Am. Chem. Soc.* **1977**, *99*, 4511–4514.

(9) Domingo, L. R.; Aurell, M. J.; Perez, P.; Contreras, R. Origin of the synchronicity on the transition structures of polar Diels-Alder reactions. Are these reactions 4 + 2 processes? *J. Org. Chem.* 2003, 68, 3884–3890.

(10) Linder, M.; Brinck, T. On the method-dependence of transition state asynchronicity in Diels–Alder reactions. *Phys. Chem. Chem. Phys.* **2013**, *15*, 5108–5114.

(11) Jaque, P.; Toro-Labbé, A.; Politzer, P.; Geerlings, P. Reaction force constant and projected force constants of vibrational modes along the path of an intramolecular proton transfer reaction. *Chem. Phys. Lett.* **2008**, 456, 135–140.

(12) Toro-Labbé, A. Characterization of chemical reactions from the profiles of energy, chemical potential and hardness. *J. Phys. Chem. A* **1999**, *103*, 4398–4403.

(13) Paredes-Gil, K.; Mendizábal, F.; Jaque, P. Further understanding of the Ru-centered [2 + 2] cycloreversion/cycloaddition involved into the interconversion of ruthenacyclobutane using the Grubbs catalysts from a reaction force analysis. *J. Mol. Model.* **2019**, 25, 305.

(14) Yepes, D.; Murray, J. S.; Santos, J. C.; Toro-Labbe, A.; Politzer, P.; Jaque, P. J. Mol. Model. **2013**, *19*, 2689–2697.

(15) Yepes, D.; Murray, J. S.; Politzer, P.; Jaque, P. The reaction force constant: an indicator of the synchronicity in double proton transfer reactions. *Phys. Chem. Chem. Phys.* **2012**, *14*, 11125–11134.

(16) Politzer, P.; Murray, J. S.; Yepes, D.; Jaque, P. Driving and retarding forces in a chemical reaction. *J. Mol. Model.* 2014, 20, 2351.
(17) Murray, J. S.; Yepes, D.; Jaque, P.; Politzer, P. Insights into

some Diels–Alder cycloadditions via the electrostatic potential and the reaction force constant. *Comput. Theor. Chem.* **2015**, *1053*, 270–280.

(18) Yepes, D.; Martínez-Araya, J. I.; Jaque, P. Solvent effect on the degree of (a)synchronicity in polar Diels-Alder reactions from the perspective of the reaction force constant analysis. *J. Mol. Model.* **2018**, *24*, 33.

(19) Fukui, K. The path of chemical-reaction - The IRC approach. *Acc. Chem. Res.* **1981**, *14*, 363–368.

(20) Gonzalez, C.; Schlegel, H. B. Reaction-Path Following in Mass-Weighted Internal Coordinate. J. Phys. Chem. **1990**, *94*, 5523–5527.

(21) Suárez, D.; Sordo, J. A. On the origin of the endo/exo selectivity in Diels-Alder reactions. *Chem. Commun.* **1998**, 385-386.

(22) Domingo, L. R.; Perez, P.; Saez, J. A. Origin of the synchronicity in bond formation in polar Diels-Alder reactions: an ELF analysis of the reaction between cyclopentadiene and tetracyanoethylene. *Org. Biomol. Chem.* **2012**, *10*, 3841–3851. (23) Domingo, L. R.; Ríos-Gutiérrez, M.; Pérez, P. A Molecular Electron Density Theory Study of the Reactivity and Selectivities in [3 + 2] Cycloaddition Reactions of C,N-Dialkyl Nitrones with Ethylene Derivatives. J. Org. Chem. 2018, 83, 2182–2197.

(24) Andrés, J.; González-Navarrete, P.; Safont, V. S.; Silvi, B. Curly arrows, electron flow, and reaction mechanisms from the perspective of the bonding evolution theory. *Phys. Chem. Chem. Phys.* **2017**, *19*, 29031–29046.

(25) Yepes, D.; Pérez, P.; Jaque, P.; Fernández, I. Effect of Lewis acid bulkiness on the stereoselectivity of Diels–Alder reactions between acyclic dienes and $\alpha_{,\beta}$ -enals. Org. Chem. Front. 2017, 4, 1390–1399.

(26) Poree, C.; Schoenebeck, F. A Holy Grail in Chemistry: Computational Catalyst Design: Feasible or Fiction? *Acc. Chem. Res.* 2017, 50, 605–608.

(27) Berski, S.; Andrés, J.; Silvi, B.; Domingo, L. R. The Joint Use of Catastrophe Theory and Electron Localization Function to Characterize Molecular Mechanisms. A Density Functional Study of the Diels– Alder Reaction between Ethylene and 1,3-Butadiene. *J. Phys. Chem. A* **2003**, *107*, 6014–6024.

(28) Berski, S.; Andrés, J.; Silvi, B.; Domingo, L. R. New Findings on the Diels–Alder Reactions. An Analysis Based on the Bonding Evolution Theory. J. Phys. Chem. A **2006**, 110, 13939–13947.

(29) Polo, V.; Andres, J.; Berski, S.; Domingo, L. R.; Silvi, B. Understanding Reaction Mechanisms in Organic Chemistry from Catastrophe Theory Applied to the Electron Localization Function Topology. *J. Phys. Chem. A* **2008**, *112*, 7128–7136.

(30) Andres, J.; Berski, S.; Domingo, L. R.; Polo, V.; Silvi, B. Describing the Molecular Mechanism of Organic Reactions by Using Topological Analysis of Electronic Localization Function. *Curr. Org. Chem.* **2011**, *15*, 3566–3575.

(31) Andrés, J.; Gracia, L.; González-Navarrete, P.; Safont, V. S. Chemical structure and reactivity by means of quantum chemical topology analysis. *Comput. Theor. Chem.* **2015**, *1053*, 17–30.

(32) Munarriz, J.; Velez, E.; Casado, M. A.; Polo, V. Understanding the reaction mechanism of the oxidative addition of ammonia by (PXP)Ir(i) complexes: the role of the X group. *Phys. Chem. Chem. Phys.* **2018**, *20*, 1105–1113.

(33) Munárriz, J.; Laplaza, R.; Polo, V. A bonding evolution theory study on the catalytic Noyori hydrogenation reaction. *Mol. Phys.* **2019**, *117*, 1315–1324.

(34) Yepes, D.; Murray, J.; Santos, J.; Toro-Labbé, A.; Politzer, P.; Jaque, P. Fine structure in the transition region: reaction force analyses of water-assisted proton transfers. *J. Mol. Model.* **2013**, *19*, 2689–2697.

(35) Yepes, D.; Murray, J. S.; Perez, P.; Domingo, L. R.; Politzer, P.; Jaque, P. Complementarity of reaction force and electron localization function analyses of asynchronicity in bond formation in Diels-Alder reactions. *Phys. Chem. Chem. Phys.* **2014**, *16* (14), 6726–6734.

(36) Jaque, P.; Toro-Labbé, A. Theoretical study of the double proton transfer in the CHX-XH···CHX-XH (X = O, S) complexes. *J. Phys. Chem. A* **2000**, *104*, 995–1003.

(37) Politzer, P.; Toro-Labbé, A.; Gutiérrez-Oliva, S.; Herrera, B.; Jaque, P.; Concha, M. C.; Murray, J. S. The reaction force: Three key points along an intrinsic reaction coordinate. *Proc. - Indian Acad. Sci., Chem. Sci.* **2005**, *117*, 467–472.

(38) Politzer, P.; Toro-Labbé, A.; Gutiérrez-Oliva, S.; Murray, J. S. Perspectives on the reaction force. In *Advances in Quantum Chemistry;* Sabin, J. R., Brandas, E. J., Eds.; 2012; Vol. *64*, pp 189–209.

(39) Jaque, P.; Toro-Labbé, A.; Geerlings, P.; De Proft, F. Theoretical Study of the Regioselectivity of [2 + 2] Photocycloaddition Reactions of Acrolein with Olefins. *J. Phys. Chem. A* **2009**, *113*, 332–344.

(40) Politzer, P.; Burda, J. V.; Concha, M. C.; Lane, P.; Murray, J. S. Analysis of the reaction force for a gas phase S(N)2 process: CH3Cl +H2O -> CH3OH+HCl. J. Phys. Chem. A **2006**, 110, 756–761.

(41) Politzer, P.; Murray, J. S. The Position-Dependent Reaction Force Constant in Bond Dissociation/Formation. *Collect. Czech. Chem. Commun.* **2008**, *73*, 822–830.

(42) Rincón, E.; Jaque, P.; Toro-Labbé, A. Reaction force analysis of the effect of Mg(II) on the 1,3 intramolecular hydrogen transfer in thymine. *J. Phys. Chem. A* **2006**, *110*, 9478–9485.

(43) Toro-Labbe, A.; Gutierrez-Oliva, S.; Murray, J. S.; Politzer, P. A new perspective on chemical and physical processes: the reaction force. *Mol. Phys.* **2007**, *105*, 2619–2625.

(44) Polanyi, J. C.; Zewail, A. H. Direct Observation of the Transition-State. Acc. Chem. Res. **1995**, 28, 119–132.

(45) Zewail, A. H. Femtochemistry: Atomic-scale dynamics of the chemical bond. J. Phys. Chem. A 2000, 104, 5660-5694.

(46) Johnson, E. R.; Keinan, S.; Mori-Sánchez, P.; Contreras-García, J.; Cohen, A. J.; Yang, W. Revealing Noncovalent Interactions. *J. Am. Chem. Soc.* **2010**, *132*, 6498–6506.

(47) Lane, J. R.; Contreras-García, J.; Piquemal, J.-P.; Miller, B. J.; Kjaergaard, H. G. Are Bond Critical Points Really Critical for Hydrogen Bonding? J. Chem. Theory Comput. **2013**, *9*, 3263–3266.

(48) Bader, R. F. W. A quantum theory of molecular structure and its applications. *Chem. Rev.* **1991**, *91*, 893–928.

(49) González, L.; Tejedor, R. M.; Royo, E.; Gaspar, B.; Munárriz, J.; Chanthapally, A.; Serrano, J. L.; Vittal, J. J.; Uriel, S. Two-Dimensional Arrangements of Bis(haloethynyl)benzenes Combining Halogen and Hydrogen Interactions. *Cryst. Growth Des.* **2017**, *17*, 6212–6223.

(50) Munárriz, J.; Rabuffetti, F. A.; Contreras-García, J. Building Fluorinated Hybrid Crystals: Understanding the Role of Noncovalent Interactions. *Cryst. Growth Des.* **2018**, *18*, 6901–6910.

(51) Lee, J.-H.; Bristowe, N. C.; Bristowe, P. D.; Cheetham, A. K. Role of hydrogen-bonding and its interplay with octahedral tilting in CH3NH3PbI3. *Chem. Commun.* **2015**, *51*, 6434–6437.

(52) Contreras-García, J.; Calatayud, M.; Piquemal, J.-P.; Recio, J. M. Ionic interactions: Comparative topological approach. *Comput. Theor. Chem.* **2012**, *998*, 193–201.

(53) Boto, R. A.; Contreras-García, J.; Tierny, J.; Piquemal, J.-P. Interpretation of the reduced density gradient. *Mol. Phys.* 2016, 114, 1406–1414.

(54) Becke, A. D. Density-Functional Thermochemistry 0.3. the Role of Exact Exchange. J. Chem. Phys. **1993**, 98, 5648-5652.

(55) Stephens, P. J.; Devlin, F. J.; Chabalowski, C. F.; Frisch, M. J. Ab Initio Calculation of Vibrational Absorption and Circular Dichroism Spectra Using Density Functional Force Fields. *J. Phys. Chem.* **1994**, *98*, 11623–11627.

(56) Chai, J.-D.; Head-Gordon, M. Long-range corrected hybrid density functionals with damped atom-atom dispersion corrections. *Phys. Chem. Chem. Phys.* **2008**, *10*, 6615–6620.

(57) González-Rivas, N.; Cedillo, A. Performance of density functional theory methods to describe intramolecular hydrogen shifts. *Proc. - Indian Acad. Sci., Chem. Sci.* **2005**, *117*, 555–560.

(58) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; et al. *Gaussian 09*, Revision B.01; Gaussian, Inc.: Wallingford, CT, 2009.

(59) Contreras-García, J.; Johnson, E. R.; Keinan, S.; Chaudret, R.; Piquemal, J.-P.; Beratan, D. N.; Yang, W. NCIPLOT: A Program for Plotting Noncovalent Interaction Regions. *J. Chem. Theory Comput.* **2011**, *7*, 625–632.

(60) Andrés, J.; Berski, S.; Contreras-García, J.; González-Navarrete, P. Following the Molecular Mechanism for the NH3 + LiH \rightarrow LiNH2 + H2 Chemical Reaction: A Study Based on the Joint Use of the Quantum Theory of Atoms in Molecules (QTAIM) and Noncovalent Interaction (NCI) Index. J. Phys. Chem. A **2014**, 118, 1663–1672.

(61) Keith, T. A. *AIMAll*, Version 19.10.12; TK Gristmill Software: Overland Park, KS, 2019. aim.tkgristmill.com.

(62) Labet, V.; Morell, C.; Toro-Labbe, A.; Grand, A. Is an elementary reaction step really elementary? Theoretical decomposition of asynchronous concerted mechanisms. *Phys. Chem. Chem. Phys.* **2010**, *12*, 4142–4151.

(63) Kolthoff, I. M.; Elving, P. J. Treatise on analytical chemistry; Interscience Encyclopedia: New York, 1959. (64) Bordwell, F. G.; Ji, G. Z. Effects of structural changes on acidities and homolytic bond dissociation energies of the hydrogennitrogen bonds in amidines, carboxamides, and thiocarboxamides. *J. Am. Chem. Soc.* **1991**, *113*, 8398–8401.

(65) Bordwell, F. G.; Bartmess, J. E.; Hautala, J. A. Alkyl effects on equilibrium acidities of carbon acids in protic and dipolar aprotic media and the gas phase. *J. Org. Chem.* **1978**, *43*, 3095–3101.

(66) Diels, O.; Alder, K. Synthesis in the hydro-aromatic tier. Justus Liebigs Ann. Chem. **1928**, 460, 98–122.

(67) Bell, R. P.; Hinshelwood, C. N. The theory of reactions involving proton transfers. *Proc. R. Soc. A* **1936**, *154*, 414–429.

(68) Evans, M. G.; Polanyi, M. Inertia and driving force of chemical reactions. *Trans. Faraday Soc.* **1938**, *34*, 11–24.

(69) de Souza, M. A. F.; Ventura, E.; do Monte, S. A.; Riveros, J. M.; Longo, R. L. Revisiting the concept of the (a)synchronicity of dielsalder reactions based on the dynamics of quasiclassical trajectories. *J. Comput. Chem.* **2016**, *37*, 701–711.