Electronic Supplementary Information

Efficient and Flexible Approach for Local Distortion: Distortion Distribution Analysis enabled by Fragmentation

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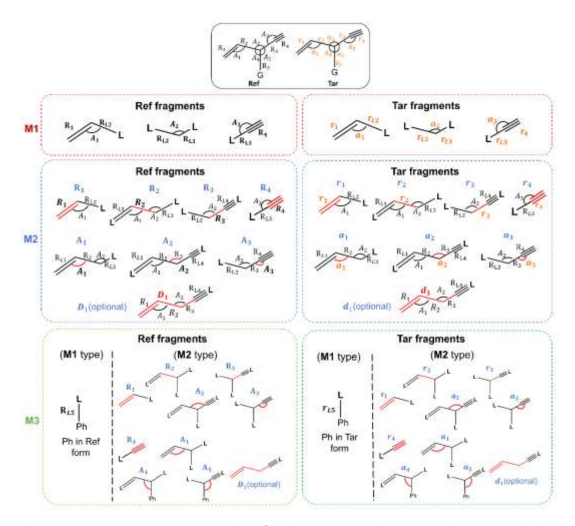
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Methodology details (a) Ref vs. Tar Ref Fragments & Tar Fragments (b)M1: $(r, \theta, \varphi)_{Ref}$ $(r, \theta, \varphi)_{Ref}$ $(r, \theta, \varphi)_{Tar}$ M2: Alter only one internal coordinate Input Structures Tar Alter r, θ or φ Openbabel Fragmentation M3: Hybrid of M1 + M2 $E_{M3} = E_{M1} + E_{M2}$ c) Link atom (d) Calculations ETar & ERej $\mathbf{R}_{\mathrm{LA}} = \mathbf{R}_{\mathrm{A}}$ $E_{Distort,i} = E_i^{Tar} - E$ Gaussian, ORCA, xTB, $+g(\mathbf{R}_{B}-\mathbf{R}_{A})$ TorchANI, MLAtom (e) Distortion Distribution/Map Distribute to each bond: M1M2 Analysis & max $E_{bond,i}$ Visualization $E_{angle,i}/2$ Edihedral, i Pymol

Scheme S1 (a) Our workflow of distortion distribution analysis with fragmentation (D2AF). The input structures contain the targe system (**Tar**) and its reference (**Ref**) form. (b) Schematic fragmentation and coordinate manipulation methods (**M1-M3**). (c) ONIOM-type link atom (LA) treatment to "project" distortion-energy of a given covalent bond into the fragments. (d) Energy calculations of fragments. (e) Distortion distribution analysis and visualization.

As shown in Scheme S1, our fragmentation-based approach consists of three stages: fragmentation, calculation, and analysis. The selected target system (denoted as **Tar**, with more distortion) and its reference configuration (denoted as **Ref**, usually with less distortion) were used as input for D2AF. Distortion energy is the energy difference between the reference configuration and its target counterpart. The local distortion within a molecule can be analyzed by fragmenting the systems using one of three fragmentation and coordinate manipulation approaches, depending on the system's complexity (see below).



Scheme S2 Fragmentation examples for three approaches. **M3** scheme is a hybrid approach combined with **M1** and **M2** schemes, in which the only sole change of the coordinate label for the **M2**-type fragments is shown for clarity.

Method 1 (M1) Fragmentation/Coordinate scheme: In many-body expansion approximation used in prevalent fragmentation schemes, S1, 2 the energy of a specific molecule can be expressed from the inclusion-exclusion principle as follows (eq. S1):

$$E_{Sys} \approx \sum_{i} E_{i} + \sum_{i,j} E_{ij} + \sum_{i,j,k} E_{ijk} + \cdots$$
 (S1)

Where E_i , E_{ij} represent the one-body energy, two-body energy contribution, respectively. The energy difference between the whole **Tar** and **Ref** molecules can be decomposed by many-body expansion approximation as follows (eq. S2):

$$E_{Distort} \approx \sum_{i} (E_{i}^{Tar} - E_{i}^{ref}) + \sum_{i,j} (E_{ij}^{Tar} - E_{ij}^{ref}) + \sum_{i,j,k} (E_{ijk}^{Tar} - E_{ijk}^{ref}) \dots (S2)$$

In this method (M1), the **Tar** and **Ref** molecules were divided into smaller fragments as much and reasonable as possible to get higher "resolution" of distortion map and reduce the computational costs (Scheme S2). As we focus on the distortion energy of each subsystem/fragment (i.e., local distortion), the one-body energy part is considered only. Therefore, local distortion is estimated as the distortion energy of each fragment, i.e., the difference between the homologous fragment of the **Tar** and **Ref** systems (eq. S3). In addition, distortion of all coordinates (e.g., bonds, angles and/or dihedrals) within each fragment are computed collectively in this **M1** method.

$$E_{Distort,i} = E_i^{Tar} - E_i^{ref}$$
 (S3)

To attain a higher resolution of distortion map, each system can be generally fragmentated in the smallest and reasonable species, consisting of a single heavy atom as well its link atoms in most cases. Alternatively, users can customize the system's partition to expand the size of particular fragement(s) to include complex and key interactions (such as delocalization in conjugate aromatic or alkenyl/alkynyl groups, lone pairs repulsions). Consequently, **M1** typically cuts all single bonds of the **Tar** and **Ref** molecules except X-H bonds (X denotes a heavy atom), which serve as boundaries between fragments (Scheme S2). For instance, as shown in Scheme S2, both **Ref** and **Tar** molecules are partitioned into three fragments, each preserving the bonds and angles characteristic of its corresponding state. All boundaries of each fragment containing dangling bonds are also capped by link atoms (L). It should be noted that local distortion energy is partially estimated by LA approximation, when its distorted boundary contains a LA.

When the system contains fused rings (e.g. indole in Figures S30, S60 and S67), in order to maintain the minimum-size ring structure in **M1** scheme, the overlapped atoms of the rings may appear in multiple separated ring fragments. In the visualization, these overlapped/shared atoms are colored according to the last fragment list (including these shared atoms) defined in the input file.

Method 2 (M2) Fragmentation/Coordinate scheme: Inspired by molecular mechanics energy expression (eq. S4), an alternative perspective on decomposing the

distortion energy into three bonding contributions. Within this molecular mechanics framework, the total energy of the system is conceptualized as follows:

$$E_{Sys} = E_{bond} + E_{angle} + E_{dihedral} + E_{elec} + E_{VDW}$$
 (S4)

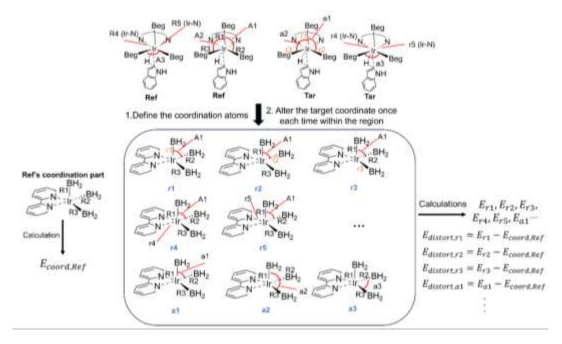
Where E_{bond} , E_{angle} and $E_{dihedral}$ are bonding components contributed from each bond, angle and dihedral energy; E_{elec} and E_{VDW} are non-bonding/noncovalent components contributed from each electrostatic and van der Waals interaction. Only the bonding components were considered in the M2 scheme, as the local distortion energy should mainly be dictated by the three bonding components and focus on the local geometrical effects. Therefore, the (relative) distortion energy of the target system (**Tar**) referring to its **Ref** counterpart can be approximated as follows (eq. S5):

$$\Delta E_{Distort} \approx \Sigma \Delta E_{bond} + \Sigma \Delta E_{angle} + \Sigma \Delta E_{dihedral}$$
 (S5)

In this method (M2), we aim to estimate these above energy changes by altering only one bonding internal coordinate (such as bond, angle and (optionally) dihedral) from its **Ref** to **Tar** state while keeping the other coordinates same as in **Ref** (Scheme S2). To achieve such distortion decomposition, fragmentation including two, three or four heavy atoms (with their link atoms) for one target internal coordinate (bond, angle or dihedral, respectively) taken from the reference form (R_i , bond/ A_i , angle/ D_i , dihedral) is performed to generate the **Ref** fragments. Then, only one target internal coordinate ($r_i/a_i/d_i$) of each generated fragment is altered to be identical as that coordinate value in the target system to set up all combinations of the **Tar** fragments (Scheme S2).

Scheme S2 illustrates all fragments obtained to reveal each individual local distortion energy contribution by altering one specific internal coordinate. Therefore, the **Tar** fragments differ from their corresponding **Ref** fragments by the one altered internal coordinate $(R_i/A_i/D_i \ vs \ r_i/a_i/d_i)$: each coordinate change corresponds to individual local distortion energy contribution for one specific bond, angle or dihedral. Accordingly, in contrast with the **M1** method, local distribution of each bonding coordinate within one fragment is evaluated individually and should not rely on the LA approximation in the **M2** method.

Method 3 (M3) Fragmentation/Coordinate scheme: Using M2 can elucidate the intricate relationship between (relative) energy changes and alterations in bonds/angles (/optional dihedrals). This approach provides an intuitive and high-resolution perspective of (relative) local distortion energy. However, this approach exhibits limitations to apply to cyclic conjugated groups containing delocalized electrons (such as π electrons) in a ring. M2 encounters challenges in strictly decoupling the conjugated moiety into bonds and angles in a ring by altering the only one specific bonding coordinate in the Tar form and keeping its remaining coordinates unchanged as in the Ref form. Therefore, when dealing with such complex systems, a hybridized partition method (refer as M3) was applied for two parts: a conjugated moiety using the M1 method and the rest part(s) using M2 (Scheme S2). As shown in Scheme S2, when the functional group (G) is aromatic Ph, the Ph group is categorized into the M1 part, and the rest part(s) is(are) treated using M2.



Scheme S3 Schematic fragmentations for metal coordination systems by the **M3** scheme.

Moreover, according to ligand field theory, metal coordination compounds exhibit distinctive and complex metal-ligand interactions. The key orbitals for this system are generally influenced and sensitive by ligands (coordinate numbers/elements), directly

reflecting the electronic structure around the transition metal. Therefore, maintaining consistent metal-ligand interactions becomes crucial when analyzing distortions related to the transition metal compounds. To tackle this challenge, we propose a novel feature in M3, in which users can define a minimum and reasonable metal coordination region as one special fragment to capture the key metal-ligand interactions (Scheme S3) using M2. The Ref fragment was generated including all atoms in this special fragment (coordination region with their link atom(s)) taken from the reference form. Then, only one target internal coordinate in the coordination region is altered to be identical as to that coordinate value in the target system to set up all combinations of the Tar fragments. Accordingly, in local distribution distortion of each bonding coordinate within the coordination region is evaluated individually. After calculations to obtain energy of these fragments, (relative) local distortion energy of each individual target M-L bond can be estimated. It should be noted that this coordinate treatment cannot work perfectly if any large coordinate change involving the metal-ligand chelation is involved between the Ref and Tar cooridinate fragment. This approach should help facilitate our analysis of local distortion distribution in metal coordination compounds.

Likewise, for the system contains fused rings (e.g. indole in Figures S30 and S60) in M3 scheme, the overlapped atoms of the rings may appear in multiple separated ring fragments. In the visualization, these overlapped/shared atoms are colored according to the last fragment list (including these shared atoms) defined in the input file.

Link-atom Treatment: When any fragment generated by any one of the fragmentation schemes (M1-M3) contain any dangling bonds, the resulting dangling bonds should be saturated using the ONIOM-type link atom (LA, denoted as L in Scheme S2) approach. Hence, the position of the link atom is determined by the following formula (eq. S6). In M1, such LA approach can project distortion of the bonding boundary to the link atom, whereas local distortion contribution should not rely on the LA approximation in M2.

$$\mathbf{R}_{\mathrm{LA}} = \mathbf{R}_{\mathrm{A}} + g(\mathbf{R}_{\mathrm{B}} - \mathbf{R}_{\mathrm{A}}) \tag{S6}$$

where A is the link-atom connection (LAC, belonging to the fragment), B is the

link-atom host (LAH), g is the scaling factor, \mathbf{R}_{LA} is the position vector of the link atom. Following our benchmark analysis of various molecular models, hydrogen and nitrogen atoms are generally selected to cap single and triple dangling bonds, respectively. Additionally, the carbon atom was generally used for double dangling bond, except only one atom as the double- or triple-bond LAH (without other connection) to maintain a consistent chemical environment.

2. Computational details

For the LA benchmark calculations, the realistic molecules were first fully optimized and then rigid scan calculations (using 'scan' keyword in Gaussian16) or relaxed scan calculations (using 'opt=modredundant" and "scan' keywords in Gaussian16) using M06-2X/6-31G(d) and/or B3LYP-D3/6-31G(d) methods. For each bond's scanning, a total of 14 steps with the step size of 0.05Å were taken. For each angle's bending scanning, a total of 12 steps with the step size of 5° strategy were taken. Moreover, structures from these rigid/relaxed scan calculations of the realistic systems were taken and then performed with the link-atom replacement and single-point energy calculations using the same computational method to obtain the related energy profiles using the LA approximation.

All these distortion calculations were performed using our open-source python package (D2AF, https://github.com/oscarchung-lab/D2AF), which employs Open Babel package to treat internal coordinates of fragments. S3 Several interfaces of calculators such as Gaussian16 for DFT, S4 GFN2-xTB as semi-empirical method (SE), S5 AIQM1, S6 and ANI-series S7, 8 as MLPs, ORCA5.0 for CCSD(T) can also be used for energy computations. S9, 10 Scripts for PyMOL S11 were finally generated by our code to visualize (relative) distortion distribution (or so-called distortion map). The choice of the DFT method and basis sets as well as structures for each application system (unless stated otherwise) were taken from the previous publications, which are discussed in the below sections 4-5.

3. LA Benchmark details

In our study, fragmentation combined with ONIOM-type link atom (LA) treatment was adopted to estimate local distortion energy of a molecule (so-called distortion distribution). Several benchmarks for LA approximation involving typical bonds encountered in this study were comprehensively investigated and these results show that LA approximation can qualitatively give reasonable distortion energy profiles. (Figures S1-S16)

First, the effects of the LA approximation on the local distortion energy in M1 were

examined. For distortion involving a single bond (e.g. C-C), a fragment with the popular hydrogen link atom (H-LA) can give similar energy profiles as the original ethane system with varying its bond length and angle (Figures S1-S2). In addition, for the case of distortion involving a C-C double-bond boundary in M2, uncommon carbon or oxygen link atom (C- or O-LA) was applied to ethene for evaluating these effects of double-bond link atom approximation (Figures S8-S14). Compared to the computed energy profiles for the original ethene system, our M06-2X results show that C-LA exhibits a smaller energy deviation than O-LA for the C-H bond change (e.g., ~0.5 kcal/mol (C-LA) vs. ~10.1 kcal/mol (O-LA) at 1.4 Å; Figure S8d). Regarding the H-C-H angle bending, O-LA and C-LA have an error of ~10 kcal/mol at large angle of 140° (Figure S8c), whereas the former LA has a smaller error than the latter one for small H-C-H bond angles.

Moreover, similar evaluations were conducted for the uncommon link atoms (N-LA or radical-type C-LA) associated with a C-C triple bond boundary of 1-butyne (Figure S16). Our findings indicate that N-LA and C-LA give very similar energy curves for the distortion of altering C-C distance and C-C-C angle. Based on our analysis of these benchmarks, H-LA, C-LA and N-LA were generally employed for the boundary involving single, double and triple bonds, respectively, unless stated otherwise.

A few possible choices of link atoms for single-, double- and triple-bond as well as special cases in our selected systems were systematically investigated and conducted to see the effect of link atoms on energetic profiles.

3.1. Single-bond LA and boundaries

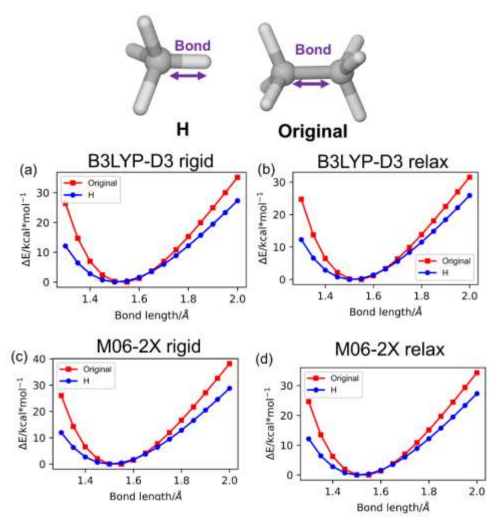


Figure S1 Benchmark of link-atom approximation for C-C single bond. CH₃-CH₃ as real system and its corresponding model (CH₄) with H as the link atom were used to get the energy profile for their varying bond length (C-H vs C-C). (a, c) Rigid scan and (b, d) relaxed scan from 1.3 Å to 2.0 Å. All the energy profiles derived from scan calculations of the real system followed by replacement by the link atom and single-point energy calculations computed at (a, b) B3LYP-D3/6-31G(d) and (c, d) M06-2X/6-31G(d) levels.

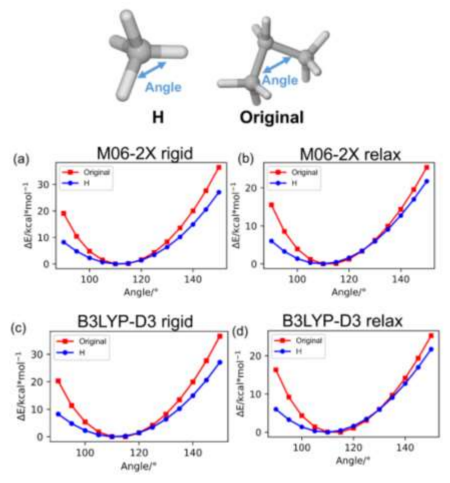


Figure S2 Benchmark of link-atom approximation for C-C single bond. C₃H₈ as real system and its corresponding model (CH₄) with H as the link atoms were used to get the energy profile for their varying angle (H-C-H vs C-C-C). (a, c) Rigid scan and (b, d) relaxed scan from from 90° to 150°. All the energy profiles were derived from scan calculations of the real system followed by replacement by the link atom and single-point energy calculations computed at (c, d) B3LYP-D3/6-31G(d) and (a, b)M06-2X/6-31G(d) levels.

As shown in Figures S1-S2, the common H-LA for the single-bond boundary was first assessed and can qualitatively reproduce the potential energy curve for a realistic C-C bond and C-C-C angle at B3LYP-D3/6-31G(d) and M06-2X/6-31G(d) levels. When the C-C bond changes from equilibrium, the H-LA begins to increasingly underestimate the C-C bond energy curve, (Figure S1: with the error of ~5.6 and ~9.4 kcal/mol at 2.0 Å for rigid and relaxed scans, respectivly.). Similarly, when the C-C-C angle bending from equilibrium, the H-LA begins to increasingly underestimate the C-C-C angle energy curve, (Figure S2: with the error of ~9.4 and ~3.7 kcal/mol at 150° for rigid and relaxed scan, respectivly.). Generally, H-LA can qualitatively replicate the overall behavior of the original potential energy curve.

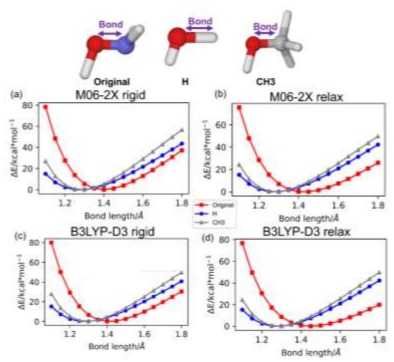


Figure S3 Benchmark of link-atom approximation for the O-N single bond in **S1** case (Figure 3). HONH₂ (a fragment in **M1**) as real model and its scaled model systems (HOH, HOCH₃) with H and CH₃ as the link atom, respectively. (a, c) Rigid scan and (b, d) relaxed scan from 1.1Å to 1.8Å. Energy profiles for the varying bond length derived from (a, c) rigid scan and (b, d) relaxed scan calculations of the real system followed by replacement by the link atom and single-point energy calculations computed at (a, b) M06-2X/6-31G(d) and (c, d) B3LYP-D3/6-31G(d) levels.

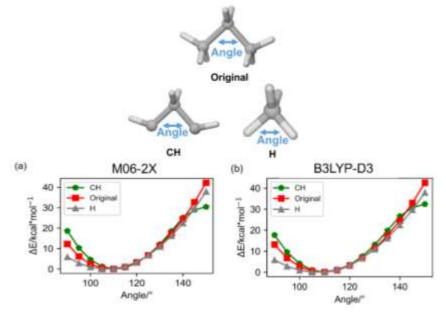


Figure S4 Benchmark of link-atom approximation for C-C single bond in the fragment **I** of the [4+2] Diels-Alder addition reaction (Figure 4). C_3H_8 as the original system and its correspongding models (CH₄ and CH₂(CH)₂) with H and CH as the link atom, respectively. Energy profiles for the varying angle derived from rigid scan calculations of the real system followed by replacement by the link atom and single-point energy calculations computed at (a) M06-2X/6-31G(d) and (b) B3LYP-D3/6-31G(d) levels.

For the LA testing on the C-C-C bending of fragment I of the [4+2] Diels-Alder addition reaction, H or CH as the link atom were applied (Figure S4). The H or CH as the link atom can reasonable reproduce the energy profiles, especially for the increased C-C-C angle bending. However, the H and CH as the link atom modestly under- and over-estimates the energies for the small C-C-C bending.

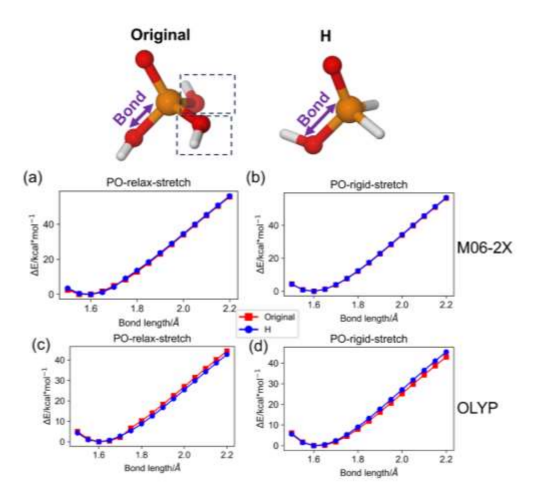


Figure S5 Link atom benchmark results of the P-O single bond from H₃PO₄ (fragment in case **SS2**) as the original system and its corresponding model O=PH₂(OH) with H as the link atom. Energy profiles for the varying bond length derived from relaxed (a, c) and rigid (b, d) scan calculations of the real system followed by replacement by the link atom and single-point energy calculations at (a, b) CPCM M06-2X/def2-TZVP and (c, d) CPCM OLYP/def2-TZVP levels.

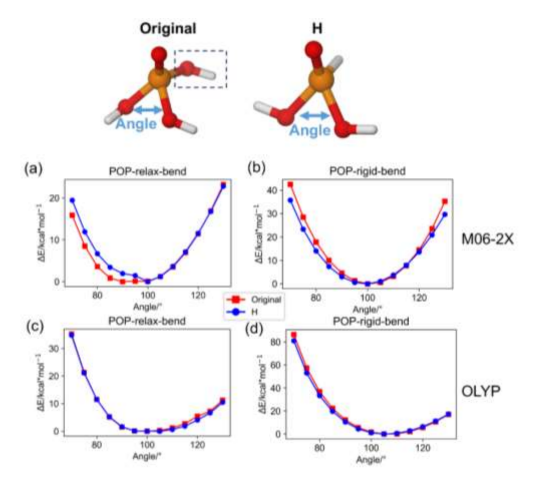


Figure S6 Link atom benchmark results of the P-O single bond from H_3PO_4 (fragment in case **SS2**) as the original system and its corresponding model $O=PH(OH)_2$ with H as the link atom. Energy profiles for the varying bond length were derived from relaxed (a, c) and rigid (b, d) scan calculations of the real system followed by replacement by the link atom and single-point energy calculations at (a, b) CPCM M06-2X/def2-TZVP and (c, d) CPCM OLYP/def2-TZVP levels.

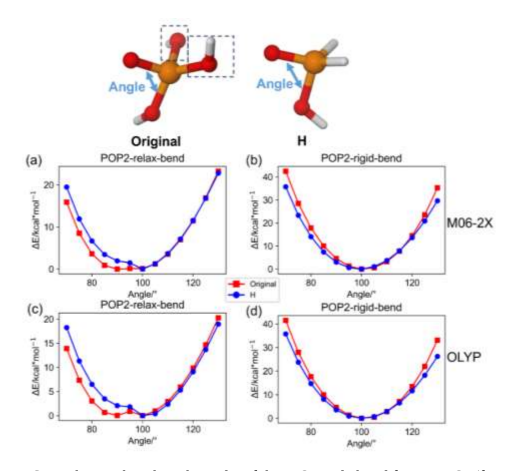


Figure S7 Link atom benchmark results of the P-O single bond from H_3PO_4 (fragment in case **SS2**) as the original system and its corresponding model $O=PH_2(OH)$ with H as the link atom. Energy profiles for the varying angle were derived from relaxed (a, c) and rigid (b, d) scan calculations of the real system followed by replacement by the link atom and single-point energy calculations at (a, b) CPCM M06-2X/def2-TZVP and (c, d) CPCM OLYP/def2-TZVP levels.

Figures S5-S7 show the benchmark results of link-atom approximation for P-O single bond in case of triphosphate hydrolysis via S_N2 (**SS2**, Figure S40). It reveals the energy profile for the varying bond length and angles can be reasonably reproduced using H-LA.

3.2. Double-bond LA and boundaries

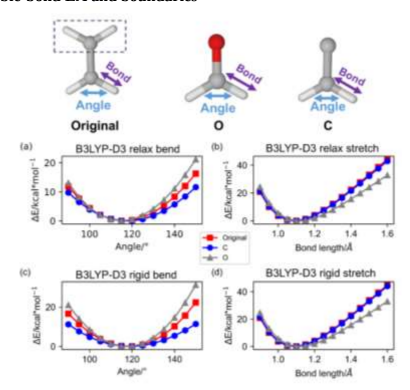


Figure S8 Benchmark of link-atom approximation for C-C double bond. Ethene as the original system and its two corresponding models (CH₂=O, CH₂=C) with either C (radical) or O as the link atom, respectively. Energy profiles for their varying bond length (b, d) and angle (a, c) derived from relaxed (a, b) or rigid (c, d) scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at B3LYP-D3/6-31G(d) level.

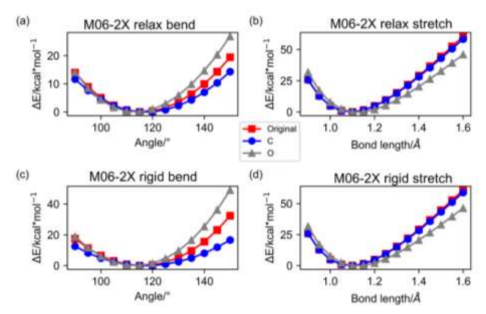


Figure S9 Benchmark of link-atom approximation for C-C double bond. Ethene as the original system and its two corresponding models (CH₂=O, CH₂=C) with either C (radical) or O as the link atom, respectively. Energy profiles for their varying bond length (b, d) and angle (a, c) derived from relaxed (a, b) or rigid (c, d) scan calculations of the real systems followed by replacement by the link atom and single-point energy

calculations computed at M06-2X/6-31G(d) level.

Similar to the results and discussion in Figures S1-S2, our additional results further support that C-LA exhibits a smaller energy deviation than O-LA for the C-H bond change (Figures S8b,d and S9b,d). In addition, O-LA and C-LA give a large error for the large H-C-H angle bending, but error become smaller for small H-C-H bond angles (Figures S8a,c and S9a,c).

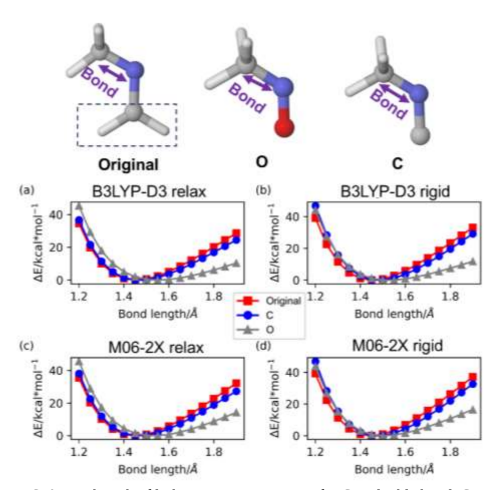


Figure S10 Benchmark of link-atom approximation for C-N double bond. $CH_2=NCH_3$ as the original system and its two corresponding models ($CH_3N=C$, $CH_3N=O$) with either C (radical) or O as the link atom, respectively. Energy profiles for their varying bond length derived from relaxed (a, c) or rigid (b, d) scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at B3LYP-D3/6-31G(d) (a, b) and M06-2X/6-31G(d) (c, d) levels.

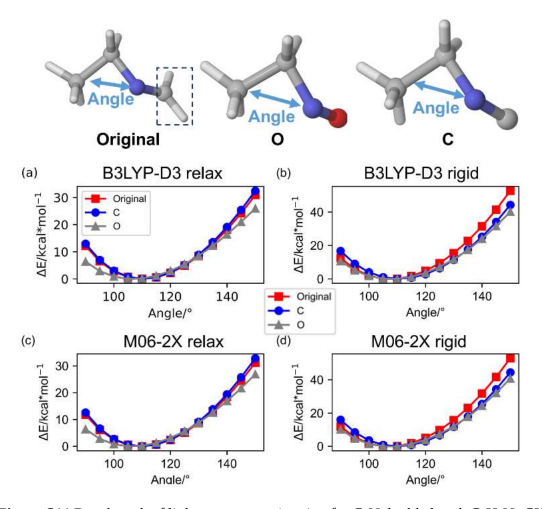


Figure S11 Benchmark of link-atom approximation for C-N double bond. $C_2H_5N=CH_2$ as the original system and its two corresponding models ($C_2H_5N=O$, $C_2H_5N=C$) with either O or C (radical) as the link atom, respectively. Energy profiles for their varying angle derived from (a,c) relax and (b,d) rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at M06-2X/6-31G(d) and B3LYP-D3/6-31G(d) level.

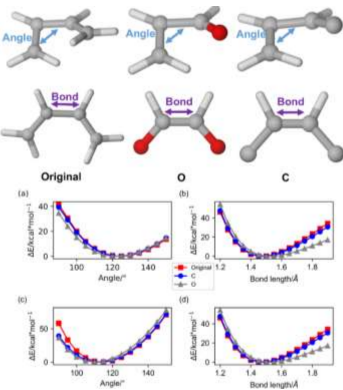


Figure S12 Benchmark of link-atom approximation with two conjugating C-C double bonds as boundaries. CH₂=CH-CH=CH₂ as the original system and its corresponding models (CH₂=CH-CH=O for angle bending/O=CH-CH=O for bond stretching, CH₂=CH-CH=C for angle bending/C=CH-CH=C for bond stretching) with either O or C (radical) as the link atom, respectively. Energy profiles for their varying angle (a, c) and bond length (b, d) derived from (a, b) relaxed and (c, d) rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at B3LYP-D3/6-31G(d) level.

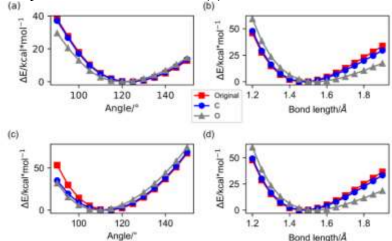


Figure S13 Benchmark of link-atom approximation with two conjugating C-C double bonds as boundaries. CH₂=CH-CH=CH₂ as the original system and its corresponding models (CH₂=CH-CH=O for angle bending/O=CH-CH=O for bond stretching, CH₂=CH-CH=C for angle bending/C=CH-CH=C for bond stretching) with either O or C (radical) as the link atom, respectively. Energy profiles for their varying angle (a, c) and bond length (b, d) are derived from (a, b) relaxed and (c, d) rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at M06-2X /6-31G(d) level

In addition, C-LA can be a good LA for the conjugating C-C double bonds as it generally gives reasonable energetics to approximate changes in the C-C bond stretching (Figures S12b, d and S13b, d) and C-C-C angle bending (Figures S12a, c and S13a, c). Furthermore, C-LA and O-LA can generally give reasonable energetics to approximate changes in the C-C bond (Figures S12b and S13b), but C-LA is a slightly better choice for the large C-C-C angle bending (Figures S12c and S13c).

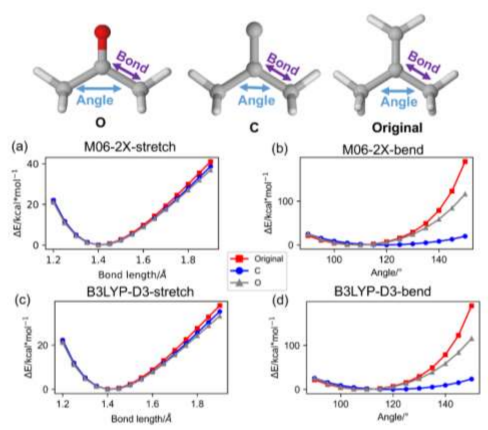


Figure S14 Benchmark of link-atom approximation for C-C double bond. $CH_2=C(CH_3)_2$ as the real system and its two corresponding models $(O=C(CH_3)_2, C=C(CH_3)_2)$ with either O or C (radical) as the link atom, respectively. Energy profiles for their varying bond length (a, c) and angle (b, d) derived from rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at (a, b) M06-2X/6-31G(d) and (c, d) B3LYP-D3/6-31G(d) levels.

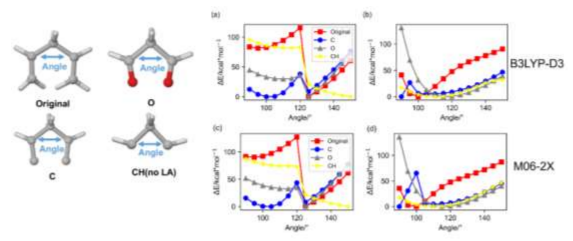


Figure S15 Benchmark of link-atom approximation for C-C double bond of **A1** case in [4+2] Diels-Alder addition reaction (Figure 3). CH_2 =CH- CH_2 -CH= CH_2 as the real system and its four corresponding models (O=CH- CH_2 -CH=O, C=CH- CH_2 -CH=C, CH- CH_2 -CH) with either O or C (radical) as the link atom and no-LA (carbiene), respectively. Energy profiles for their varying angle derived from (a, c) relaxed and (b, d) rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at (a, b) B3LYP-D3/6-31G(d) and (c, d) M06-2X/6-31G(d) levels.

The distortion energy relating to the small C-C-C angle bending (A1) of cyclopentadiene in the S2-[4+2] Diels-Alder addition reaction (Figure 3) is hard to be evaluated, due to the high coupling of a few internal coordinates inside the cyclopentadiene ring. Thus, an additional manual step is necessary to mitigate this coupling effect. As shown in Figure S15, the energy curves of C-C-C show a large difference when C-C-C becomes smaller from the equilibrium position for the O-, C-LA and the original cases. These sudden changes may come from the two too-closed link atoms or CH₂.

To minimize interactions between the links atoms, only the CH group without extra link atom was applied, resulting in a distortion energy of 3.1 kcal/mol for A1. This value is consistent with the value (3.0 kcal/mol) of fragment I (including A1) using the most reasonable link atom (CH) in the M1 scheme (Figure S4).

3.3. Triplet-bond LA and boundaries

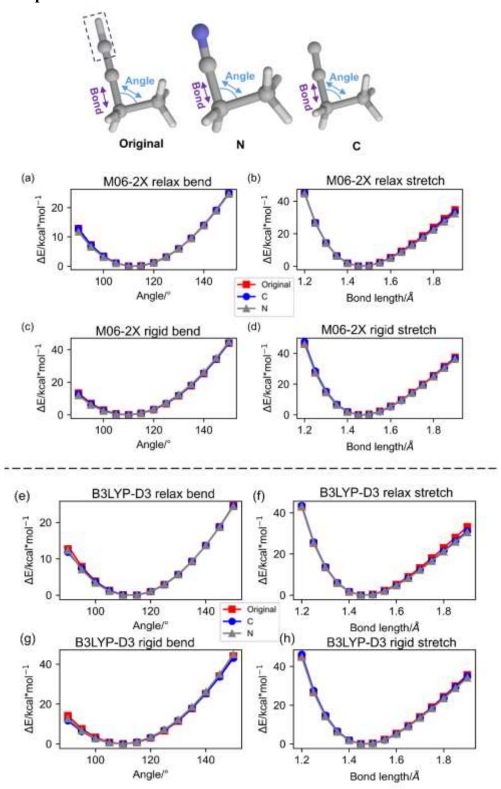


Figure S16 Benchmark of link-atom approximation for C-C triple bonds. 1-butyne as the original system and its two corresponding models (C₃H₅N, C₃H₅C radical) with either N or C (radical) as the link atom, respectively. Energy profiles for their varying (b,d;f,h)bond length and (a,c;e,g)angle derived from rigid scan calculations of the real systems followed by replacement by the link atom and single-point energy calculations computed at (e-h) B3LYP-D3/6-31G(d) and (a-d) M06-2X/6-31G(d) levels.

Similar to results and discussion in Figures S16, our additional results (using B3LYP-D3/6-31G(d)) further support that N-LA and radical-type C-LA can be a reasonable LA associated with a C-C triple bond boundary of 1-butyne (Figure S16).

4. Supplementary figures for the representative systems in the main text

4.1. Reverse Cope Elimination

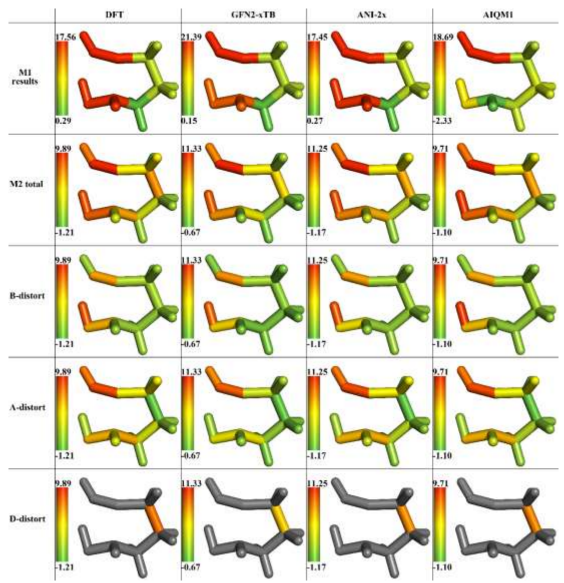


Figure S17 Distortion energy distribution (kcal/mol; coloring in term of log scale) of $\mathbf{S1}^{S12}$ using the $\mathbf{M1}$ scheme and total distortion and its contribution from bond angle and dihedral terms using the $\mathbf{M2}$ scheme. The major local distortion contributed from the dihedral ($\mathbf{D1}$ in Figure 2) was also considered by using an expanded CH_2CH_2 fragment in $\mathbf{M1}$ scheme and adding $\mathbf{D1}$ in $\mathbf{M2}$ scheme). Energies were computed using DFT (B3LYP/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

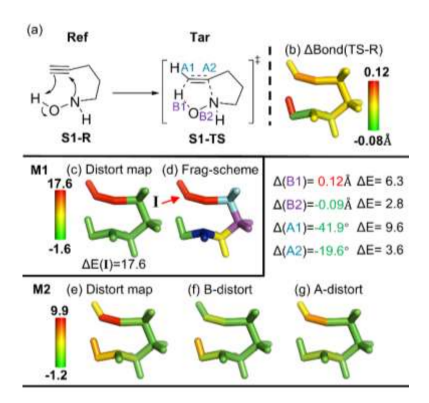


Figure S18 (a) Overview of S1 (structures taken from ref S12). (b) Bond length change (Δ bond in Å) from the reference form (**Ref**; reactant, **S1-R**) to the target form (**Tar**; TS, **S1-TS**). (c) Distortion energy distribution (kcal/mol) and (d) default fragmentation using **M1** scheme using the **default** fragmentation settings. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M2** scheme using the **default** fragmentation settings. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at B3LYP/6-31G(d) level.

Figure S18 shows the **M1** & **M2** results without including one key dihedrals' contribution (**D1**), i.e. the default fragmentation settings for **M1** and **M2** methods. In contrast, a smaller distortion energy of the hydroxylamine part (~1.5 and ~-1.6 kcal/mol, Figure S18†) was shown in the **M1** results than our benchmark for this O-N bond boundary (Figure S3†), when the (default) smallest fragmentation strategy was used (splitting the OH and NH parts). These results indicate that the H-link atom underestimates destabilization of the O/N lone pairs repulsion within the hydroxylamine part. Pleasingly, slight expansion of that fragment by including the OH and NH parts leads to more reliable results (Figures 2c and S12).

4.2. Diels-Alder Addition

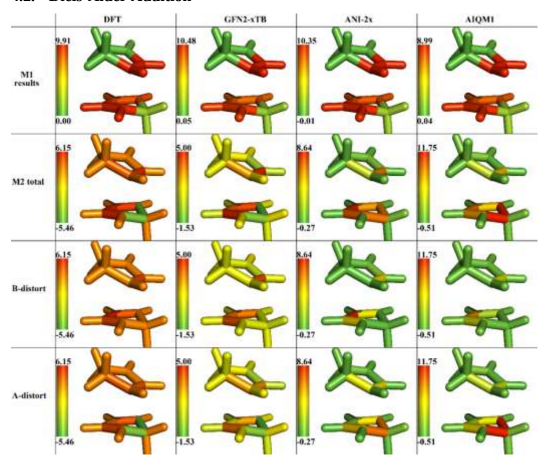


Figure S19 Distortion energy distribution (kcal/mol in log scale) of **S2**^{S13} using **M1** scheme and total distortion and its contribution from bond and angle terms using **M2** scheme. Energy calculations were computed using DFT (B3LYP/6-31G(d)), GFN2-xTB, ANI-2x, AIQM1 methods. These results are obtained from default fragmentation + LA settings (i.e. using default LAs) for **M2** & **M1**.

In addition, our benchmark shows that the distortion energy contributed by the small C-C-C angle A1 cannot be well reproduced by H-LA (Fig. S4†). Accordingly, the fragment I containing the A1 angle results in modest underestimation of its distortion energy (~1.3 kcal/mol) using M1 scheme with H-LA. That distortion energy of fragment I can raise to ~3.0 kcal/mol (Fig. 3c), when a larger CH group was used as the link atom (Fig. S4†). Similarly, the fragment related to the small A1 angle also shows a reasonable energy (~3.1 kcal/mol) in the M2 scheme, only when keeping the CH-CH2-CH group without LAs is adopted (Fig. S15†). As shown in Figure S19 and discussed in the main text, using default fragmentation settings for M1 and M2 schemes can lead to unreasonable results for fragments I and A1 (Figure 3). As shown in the benchmark in Figure S4, H-LA can modestly underestimate the (A1) angle bending

included in fragment **I**. Using CH as link atoms can provide a reasonable value for fragment **I**. Moreover, unreasonable results were also obtained by the default fragmentation settings for the **M2** scheme, (~-5.5 kcal/mol in A-distort (DFT results), Figure S18). To find a better model to describe such cases for **A1**, benchmarks shown in Figures S4 and S15 were carried out and suggest that a few LAs are incapable to well reproduce the distortion energy. Whereas, no introduction of link atoms (CH model) gives the most consistent results with the CH₃-CH₂-CH₃ (Figure S4). This may indicate that such model (CH) is so far the better model to represent distortion relating to the **A1**'s bending.

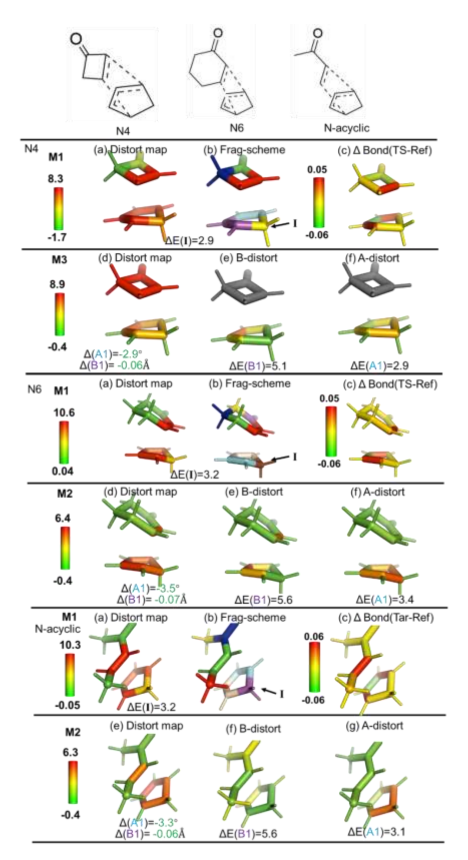


Figure S20 Results for Diels-Alder cycloaddition series (**N4**, **N6**, **N-acyclic** substrates; structures taken from ref S13) Same as the **S2**, the CH was chosen as the link atom for the fragment **I** in **M1** scheme and no link atom was applied to **A1** angle in **M2** scheme (Figure 3). All energies were computed using DFT (B3LYP/6-31G(d)) method.

4.3. Azide-alkyne Cycloaddition

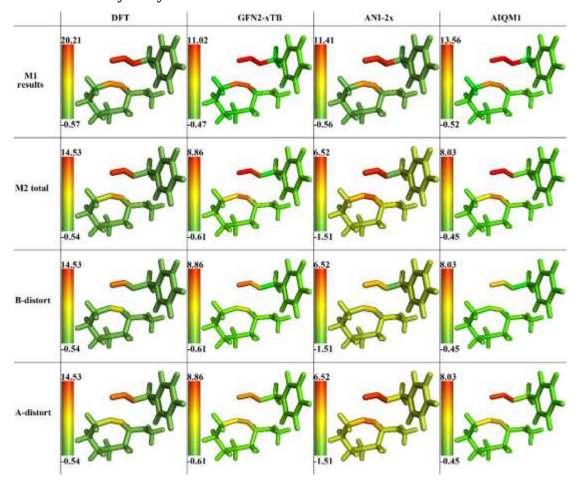


Figure S21 Distortion energy distribution (kcal/mol in log scale) of $S3^{S14}$ using M1 and M2 schemes and its contribution from bond and angle terms. Energy calculations were computed using SCS-MP2/6-31G(d), GFN2-xTB, ANI-2x, AIQM1 methods. SCS-MP2/6-31G(d) calculations were conducted by Gaussian16.

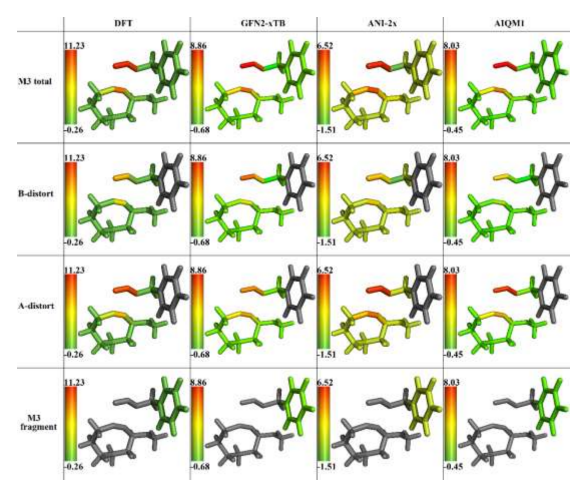
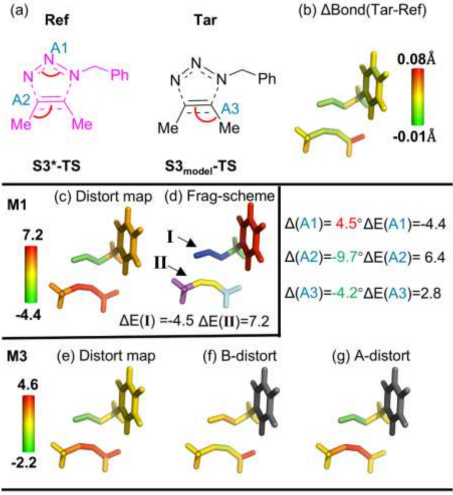


Figure S22 Distortion energy distribution (kcal/mol in log scale) of **S3** using **M3** scheme and its contribution from bond, angle and group fragment terms. Energy calculations were computed using SCS-MP2/6-31G(d), GFN2-xTB, ANI-2x, AIQM1 methods. SCS-MP2/6-31G(d) calculations were conducted by Gaussian16.

To get deeper understanding on this distortion-promoted addition, the same reaction with an acyclic alkyne substrate (2-butyne, S3*-R) was also studied by the same computational method (Figure 4h). Based on S3*-R structure, the potential energy surface (S3*-TS and S3*-P) was calculated. To compared with the strained azide-alkyne cycloaddition (S3), S3_{model}-R, S3_{model}-TS and S3_{model}-P models were generated by truncating S3-R, S3-TS and S3-P (only keeping the alkyne part and the azide) followed by capping with H-LA. Single-point energies on the S3_{model}-R, S3_{model}-TS and S3_{model}-P models were calculated directly to give a rough barrier (~9.6 kcal/mol), which is comparable to the full S3 system (~7.7 kcal/mol).



*ΔE in kcal/mol at SCS-MP2/6-31G(d)

Figure S23 (a) Overview of the truncated $S3_{model}$ -TS (extracting from S3-TS with keeping only the alkyne part) with the reference to $S3^*$ -TS. (b) Distortion distribution (kcal/mol) and (c) fragmentation using M1 scheme. (d) Bond length change (Δ bond in Å) from the reference form ($S3^*$ -TS) to the target form ($S3_{model}$ -TS). (e) Total distortion distribution (kcal/mol) (f) distortion contribution from bond and (g) angle using M3 scheme. The key geometrical changes and their corresponding distortion energy (ΔE) are also given.

In addition, a direct comparison of distortion energy between $S3_{model}$ -TS and S3*-TS (Figure S23) can be easily performed. Our distortion distribution shows that significantly higher distortion ($\Delta E(A2)$: 6.4 kcal/mol, $\Delta E(A3)$: 2.8 kcal/mol) was found in the alkyne moiety of $S3_{model}$ -TS.

4.4. Triplet Excited-state Di- π -methane Rearrangement.

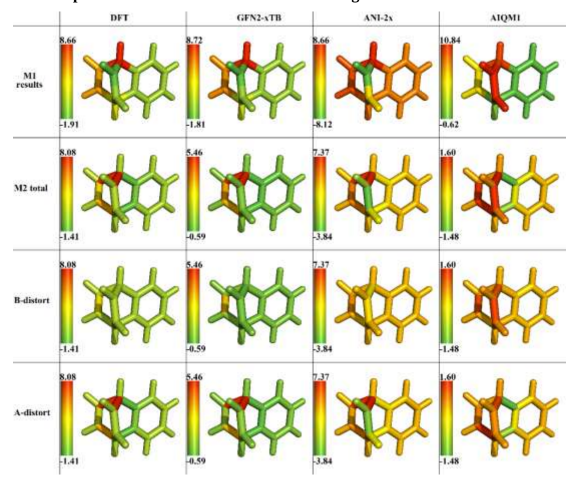


Figure S24 Distortion energy distribution (kcal/mol in log scale) of $S4^{S15}$ using M1 scheme and total distortion and its contribution from bond and angle terms using M2 scheme. Energy calculations were computed using DFT (B3LYP/6-31+G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

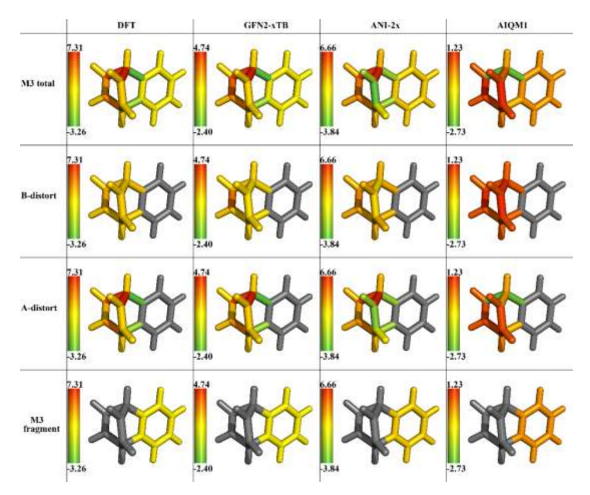


Figure S25 Distortion energy distribution (kcal/mol in log scale) of **S4** using **M3** scheme and its contribution from bond, angle and fragment terms. Energy calculations were computed using DFT (B3LYP/6-31G+(d), GFN2-xTB and ANI-2x and AIQM1 methods

4.5. A Cyclization Reaction of the Fumaramide within [2]Rotaxane.

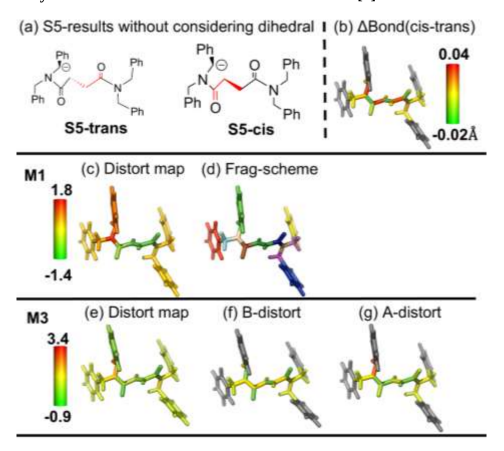
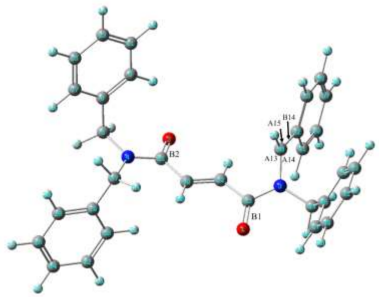


Figure S26 (a) Overview of the cis and trans isomers of an intermediate (**S5**) during the CsOH-promoted cyclization. S16 (structures taken from ref S16) (b) Bond length change (Δbond in Å) from the reference form (**Ref**; **S5-trans**) to the target form (**Tar**; **S5-cis**). (c) Distortion distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Distortion distribution (kcal/mol), distortion contribution from (f) bond and (g) angle using **M3** scheme. **These results are obtained without consideration of the two key dihedral contribution.** All energies were computed using DFT (SMD M06/6-311++G(d,p)) method.

Table S1 Atom indexes and major distortion contributions (>0.5 kcal/mol) from bond and angle using M3 scheme. These results are obtained without consideration of the two key dihedral contribution. All energies were computed using DFT (SMD M06/6-311++G(d,p)) method.



Coordinate ID	Atom indexes	Distortion /kcal*mol ⁻¹	Coordinate change Bond: Å; Angle: °
A13	3,30,31	1.49	-5.7
A14	3,30,32	1.16	-4.4
A15	31,30,32	0.99	-7.5
B14	30,32	0.93	0.02
B1	1,65	-0.56	-0.01
B2	2,61	-0.63	-0.01
Sum	NA	1.0 ^a	NA

a. Summation of local distortion energy of all bonds and angles (including those with <0.5 kcal/mol are not listed above). Distortion of the whole molecule: **5.0 kcal/mol**

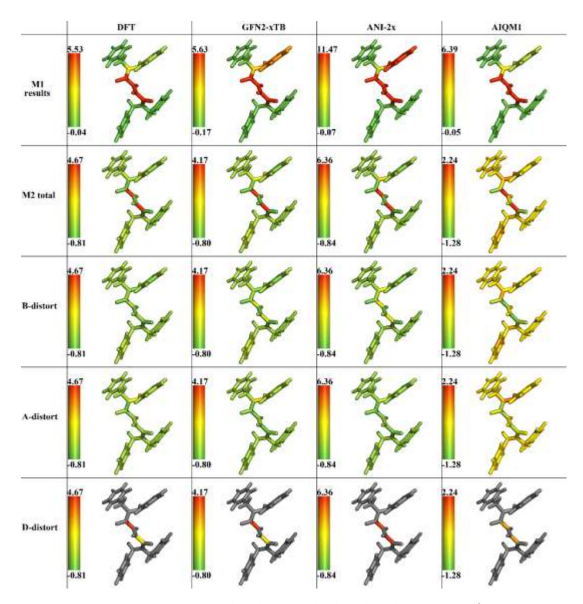


Figure S27 Distortion energy distribution (kcal/mol in log scale) of **S5** using **M1** scheme and total distortion and its contribution from bond, angle and dihedral terms using **M2** scheme. Energy calculations were computed using DFT (SMD M06/6-311++G(d,p)), GFN2-xTB and ANI-2x, AIQM1 methods. The larger fragment **I** (including **D1** and **D2** dihedrals) was further treated using the **M1** scheme and the two key dihedrals were included in the **M2** scheme (Figure 6).

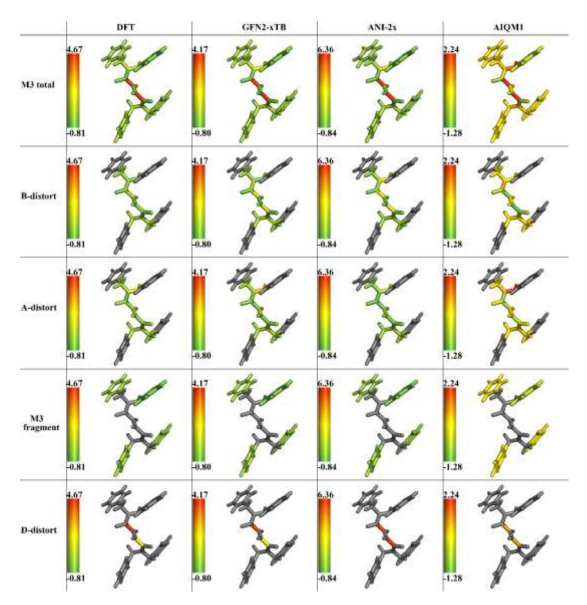


Figure S28 Distortion energy distribution (kcal/mol in log scale) of **S5** using **M3** scheme and its contribution from bond, angle and dihedral and group fragment terms using DFT (SMD M06/6-311++G(d,p)), GFN2-xTB and ANI-2x, AIQM1 methods. The two key dihedrals were included in the **M3** scheme (Figure 6).

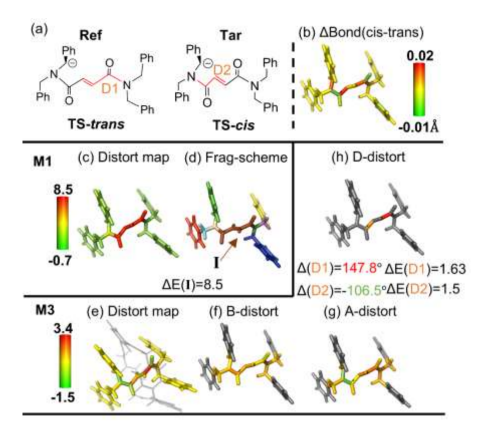


Figure S29 (a) Overview of the cis and trans isomers of the corresponding transition state (**S5**) during the CsOH-promoted cyclization. S16 (structures taken from ref S16) (b) Bond length change (Δ bond in \mathring{A}) from the reference form (**Ref**; **TS-trans**) to the target form (**Tar**; **TS-cis**). (c) Distortion distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Distortion distribution (kcal/mol) (within the rotaxane in grey transparent line representation), distortion contribution from (f) bond and (g) angle and (h) torsion using **M3** scheme. All energies were computed using DFT (SMD M06/6-311++G(d,p)) method.

4.6. Iridium-Catalyzed C-H Borylation

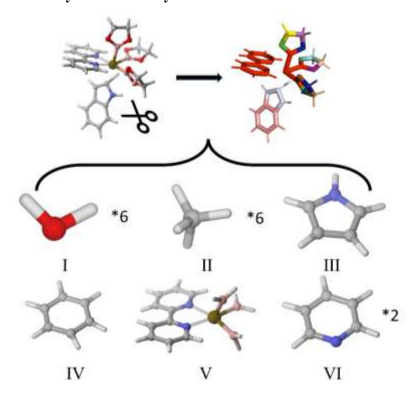
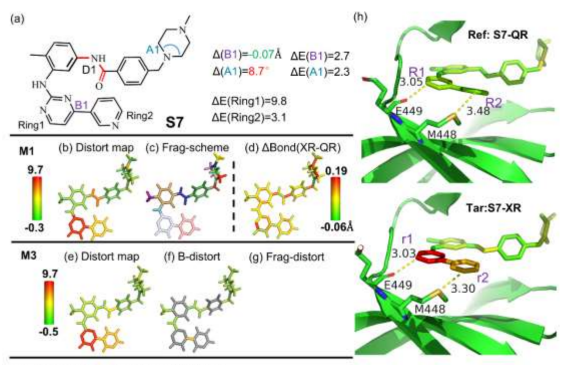


Figure S30 All molecular fragments generated by the M1 scheme from the Ir-indole system (S6)

As shown in Figure S30, the whole system of the Ir-indole system is divided into six kinds of fragments using the M1 scheme. Six fragments I come from six oxygen atoms in the Beg rings. Six fragments II come from the sp3 carbon atoms. Fragments III and IV come from separation of Indole ring in order to localize the distortion. The specialized fragment V, which includes the Ir center, along with its coordination boron atoms and the bpy are set to capture the crucial coordination interactions. Two ring fragments VI were obtained by separating the bpy to calculate the ring distortion.

4.7. Imatinib in Spleen Tyrosine Kinase

The drug Imatinib in spleen tyrosine kinase from the X-ray crystal and QR refined structures were studied, in which the latter one were obtained using ONIOM(ω B97-D3/6-31G(d):GFN2-xTB:Amber) method. S17 As shown in Figure S31b, e, the M1 and M3 results show some difference in distortion energy related to the amide group, possibly due to the mis-counting contribution from D1 (Δ (D1) = ~36.6°). After taking D1 into account, distortion energy in this region by M3 scheme becomes more consistent with that by M1 scheme.



*ΔE in kcal/mol at ωB97X/6-31G(d)

Figure S31 (a) Overview of structures of Imatinib-Spleen tyrosine kinase(**S7**)^{S17} (structures from ref S17). (b) Distortion energy distribution (kcal/mol) and (c) fragmentation using **M1** scheme. (d) Bond length change (Δbond in Å) from quantum refinement (**Ref**, QR, **S7-QR**) structure to X-ray (**Tar**, XR, **S7-XR**) crystal structure. (e) Distortion energy distribution (kcal/mol), distortion contribution from (f) bond and (g) angle using **M3** scheme. (h) Scheme for binding site interaction with neighbouring residues taken from the XR and QR structures. The drug structures are colored based on their relative distortion to that optimized structure in gas phase using **M1** scheme. These **M3** results did not include contribution from the key **D1** dihedral. All energies were computed using DFT (ωB97X-D/6-31G(d)) method.

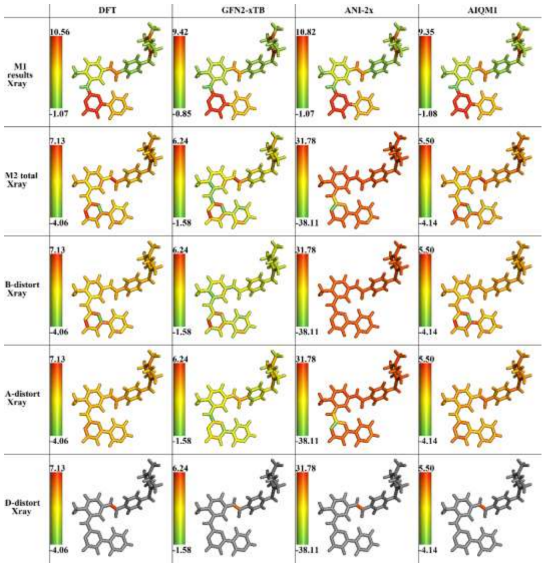


Figure S32 Distortion energy distribution (kcal/mol in log scale) of Imatinib-Spleen tyrosine kinase. The X-ray crystal structure and the optimized structure in the gas phase are set as the target and reference forms. Total distortion distribution using **M1** scheme, total contribution and its contribution from bond, angle and dihedral terms using **M2** scheme. Energy calculations were computed using DFT (ω B97X-D/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

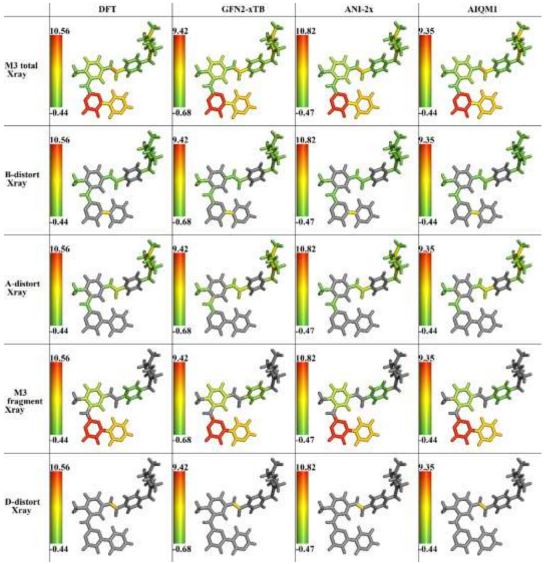


Figure S33 Distortion energy distribution (kcal/mol in log scale) of Imatinib-Spleen tyrosine kinase. The X-ray crystal structure and the optimized structure in the gas phase are set as the target and reference forms. Total contribution and its contribution from bond, angle and dihedral terms using **M3** scheme. Energy calculations were computed using DFT (ωB97X-D/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

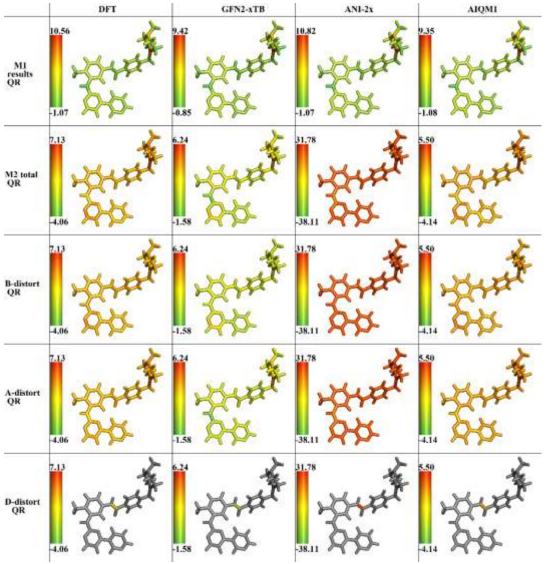


Figure S34 Distortion energy distribution (kcal/mol in log scale) of Imatinib-Spleen tyrosine kinase. The QR refined structure and the optimized structure in the gas phase are set as the target and reference forms. Total distortion distribution using **M1** scheme, total contribution and its contribution from bond, angle and dihedral terms using **M2** scheme. Energy calculations were computed using DFT (ω B97X-D/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

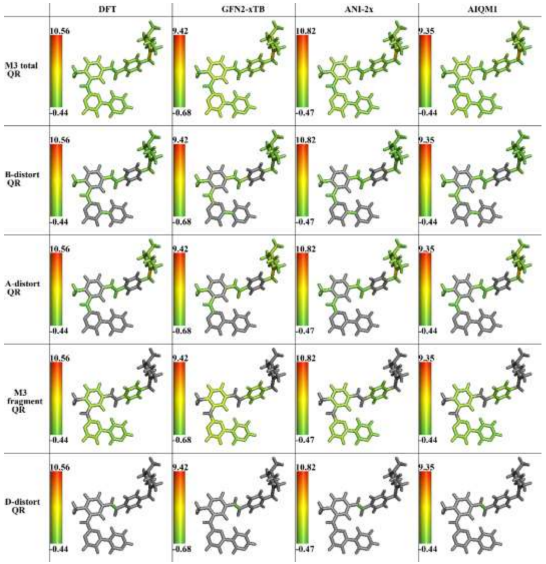


Figure S35 Distortion energy distribution (kcal/mol in log scale) of Imatinib-Spleen tyrosine kinase. The QR refined structure and the optimized structure in the gas phase are set as the target and reference forms. Total contribution and its contribution from bond, angle and dihedral terms using **M3** scheme. Energy calculations were computed using DFT (ωB97X-D/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

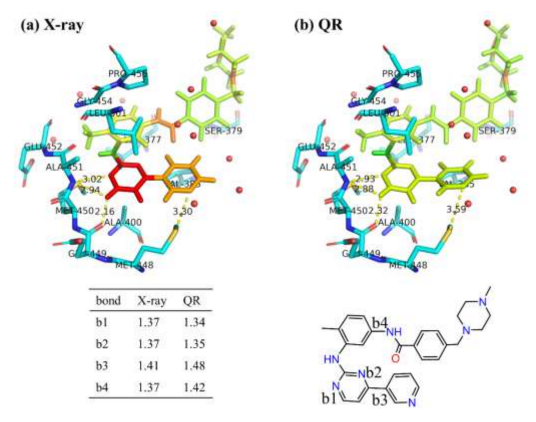


Figure S36 The key distances between Imatinib and neighboring residues in Imatinib-Spleen tyrosine kinase from the X-ray crystal and QR refined structures. The key bond lengths (\mathring{A}) are listed. Imatinib is colored based on distortion distribution using M1 scheme while the optimized structure in the gas phase was set as the reference form.

4.8. S_{N2} (MD)

C1
$$\overset{-}{-}$$
 + CH₃Cl $\longrightarrow \begin{bmatrix} C1 & \overset{H}{-} & \overset{H}{-} & \overset{H}{-} \\ C1 & \overset{\dagger}{-} & \overset{\dagger}{-} & CICH_3 & + CI \end{bmatrix}^{\ddagger}$

R

TS

P

Figure S37 SN2 reaction (S8)

Computational details: the reactant, transition state and product for this reaction were first optimized by HF/3-21G. Then, on-the-fly quasi-classical MD simulations were initiated by normal mode sampling at 298.15 K from this optimized transition state at the HF method using Progdyn code. S18 The trajectory was propagated in both the forward and backward directions for 150 MD steps with a time step of 1 fs. The energy and force of all structures at each step were calculated on-the-fly by the HF method.

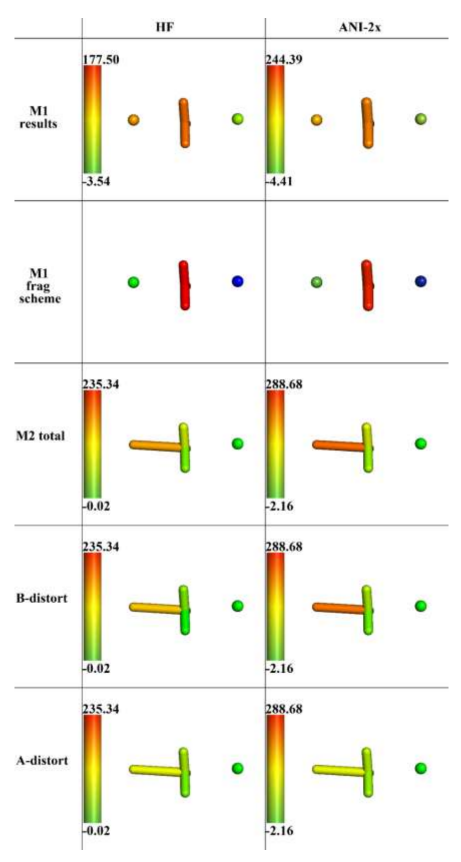


Figure S38 Distortion energy distribution (kcal/mol in log scale) for the transition state (TS) for a S_N2 reaction between CH₃Cl and a Cl⁻ using **M1** scheme, total distortion and its bond and angle contributions using **M2** scheme. All energies were computed using HF/3-21G and ANI-2x methods.

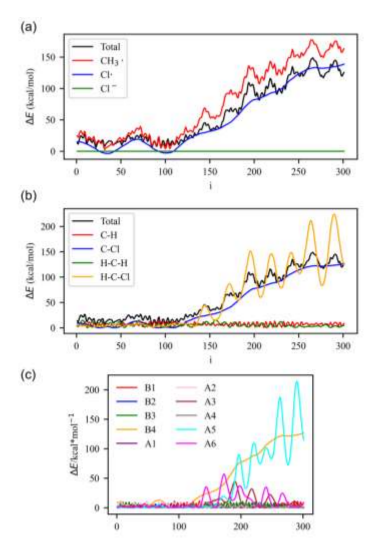


Figure S39 Distortion energy distribution (kcal/mol in log scale) for a MD trajectory of the S_N2 reaction between CH_3Cl and a Cl^- . (a) Distortion distribution using **M1** scheme. (b) Distortion distribution using **M2** scheme (with summing distortion energy belonging to the same kind of chemical bonds/angles) (c) Distortion distribution using **M2** scheme (B1, B2, B3: C-H; B4: C-Cl; A1, A2, A4: H-C-H; A3, A5, A6: H-C-Cl).

5. Supplementary application results

The additional testing systems can be categorized into different types: inorganic systems, organic systems, excited-state organic systems, supramolecular systems, coordination systems, biochemical systems and multiple structures.

5.1. Inorganic systems

5.1.1. Frustrated Lewis-Pair Catalysis

This inorganic system is a geminal aminoborane-based frustrated Lewis pair,^{S19} which activates H₂. All structures in Figure S40 were taken from ref. S19 for our D2AF analysis using DFT (M06-2X/def2-TZVPP) and GFN2-xTB methods.

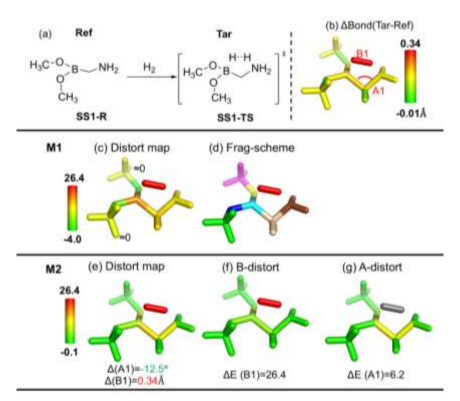


Figure S40 (a) Overview of the system of hydrogen activation with a frustrated Lewis pair. (b) Bond length change (Δ bond in \mathring{A}) from the reference form (**Ref**; R, **SS1-R**) to the target form (**Tar**; TS, **SS1-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M2** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at M06-2X/def2-TZVPP level.

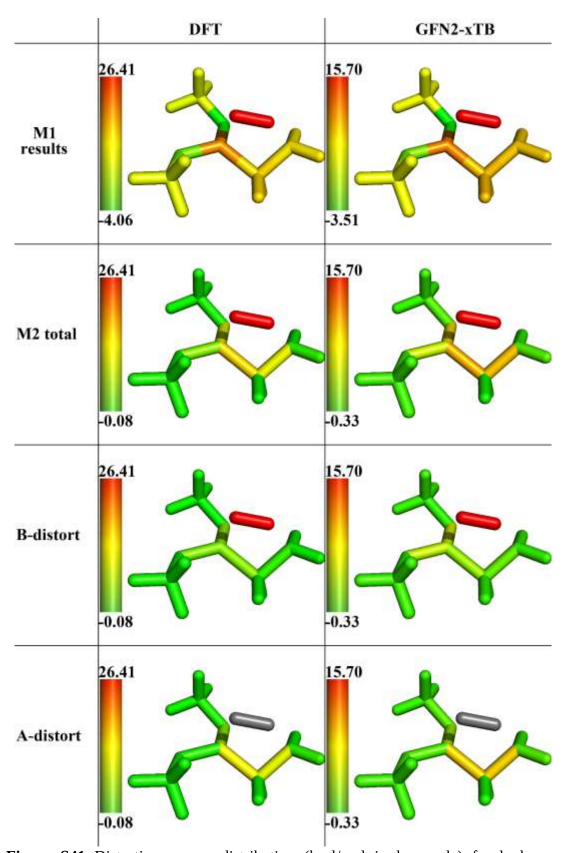


Figure S41 Distortion energy distribution (kcal/mol in log scale) for hydrogen activation by the frustrated Lewis pair using M1 scheme and total distortion and its contribution from bond and angle terms using M2 scheme. All energies were computed using DFT (M06-2X/def2-TZVPP) and GFN2-xTB methods.

5.1.2. Triphosphate Hydrolysis via S_N2

This inorganic system is a Mg-catalyzed triphosphate hydrolysis, with MeO⁻ as the nucleophile and pyrophosphate as the leaving group.^{\$20} All structures in Figure S42 were taken from ref. S20 for our D2AF analysis using OLYP/def2-TZVP and GFN2-xTB methods.

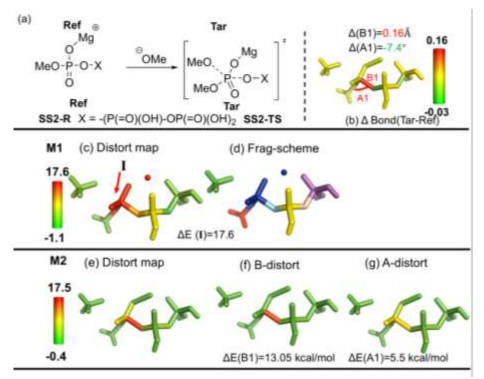


Figure S42 (a) Overview of the Mg-catalyzed triphosphate hydrolysis. (b) Bond length change (Δ bond in Å) from the reference form (**Ref**; R, **SS2-R**) to the target form (**Tar**; TS, **SS2-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M2** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at OLYP/def2-TZVP level

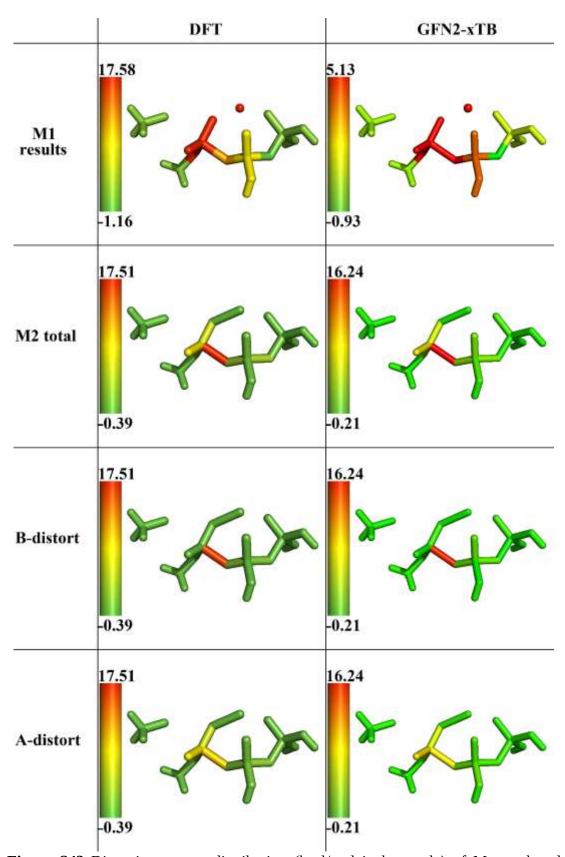


Figure S43 Distortion energy distribution (kcal/mol in log scale) of Mg-catalyzed triphosphate hydrolysis using M1 scheme and total distortion and its contribution from bond and angle terms using M2 scheme. All energies were computed using DFT (OLYP/def2-TZVP) and GFN2-xTB methods.

5.2. Organic systems

5.2.1. HCl-Mediated Prins Reaction (Organocatalysis)

In this case, a transition state (TS) structure as **Tar** and its preceding complex as **Ref** in a Prins reaction were taken from ref. S21 for our D2AF analysis using DFT (CPCM, B3LYP/6-31G(d)), GFN2-xTB and ANI-2x methods.

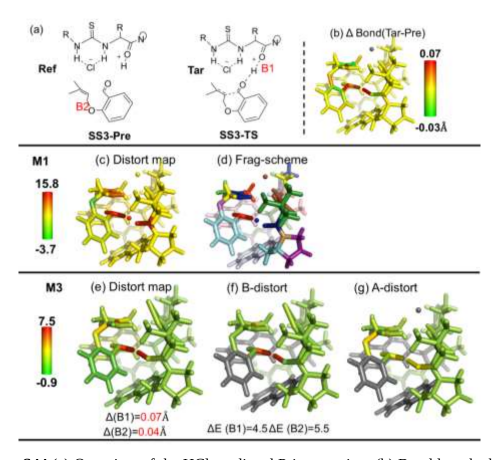


Figure S44 (a) Overview of the HCl-mediated Prins reaction. (b) Bond length change (Δ bond in Å) from the reference form (**Ref**; Pre, **SS3-R**) to the target form (**Tar**; TS, **SS3-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at CPCM B3LYP/6-31G(d) level.

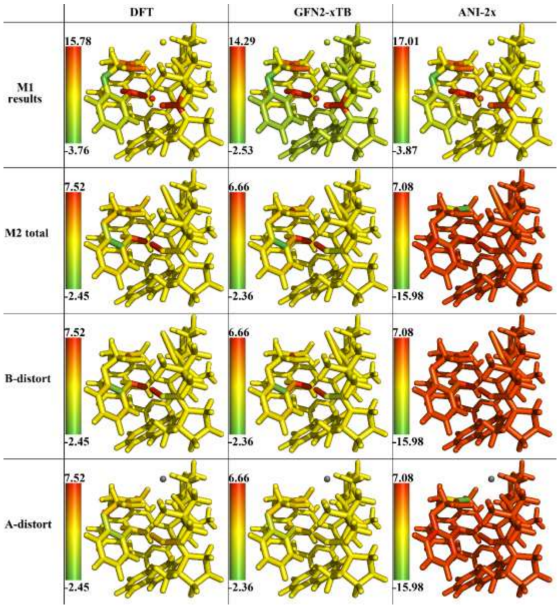


Figure S45 Distortion energy distribution (kcal/mol in log scale) of HCl-mediated Prins reaction using **M1** scheme and total distortion and its contribution from bond and angle terms using **M2** scheme. All energies were computed using DFT (CPCM B3LYP/6-31G(d)), GFN2-xTB and ANI-2x methods.

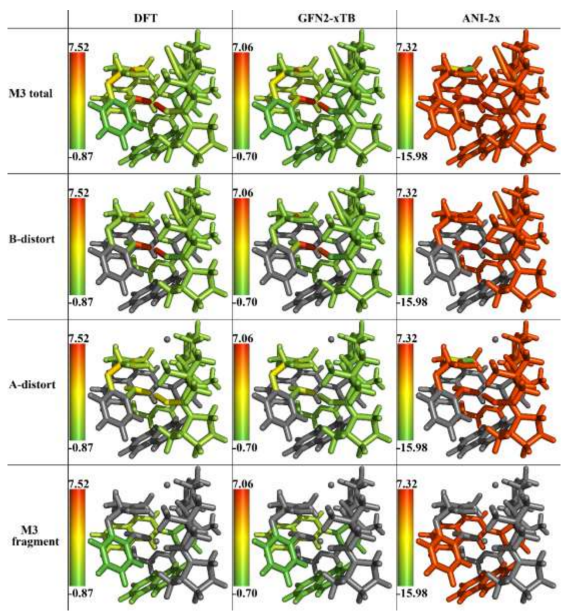


Figure S46 Distortion energy distribution (kcal/mol in log scale) of HCl-mediated Prins reaction using **M3** scheme and its contribution from bond, angle and fragment terms. All energies were computed using DFT (CPCM B3LYP/6-31G(d)), GFN2-xTB and ANI-2x methods

5.2.2. Lewis-base Catalyzed Sulfeno-functionalization

In this system, the transition state (TS) as **Tar** and its precursor as **Ref** for the Lewis-base catalyzed sulfeno-functionalization of alkenes^{S22} were taken from ref. S22 for our D2AF analysis using DFT (SMD B3LYP/6-31G(d)) and GFN2-xTB methods.

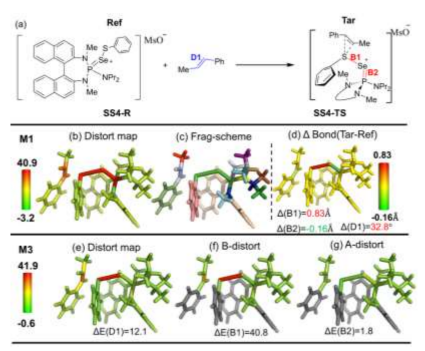


Figure S47 (a) Overview of the Lewis-base catalyzed sulfeno-functionalization. (b) Distortion energy distribution (kcal/mol) and (c) fragmentation using **M1** scheme. (d) Bond length change (Δbond in Å) from the reference form (**Ref**; R, **SS4-R**) to the target form (**Tar**; TS, **SS4-TS**). (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (ΔE) are also given. All energies were computed at SMD B3LYP/6-31G(d) level.

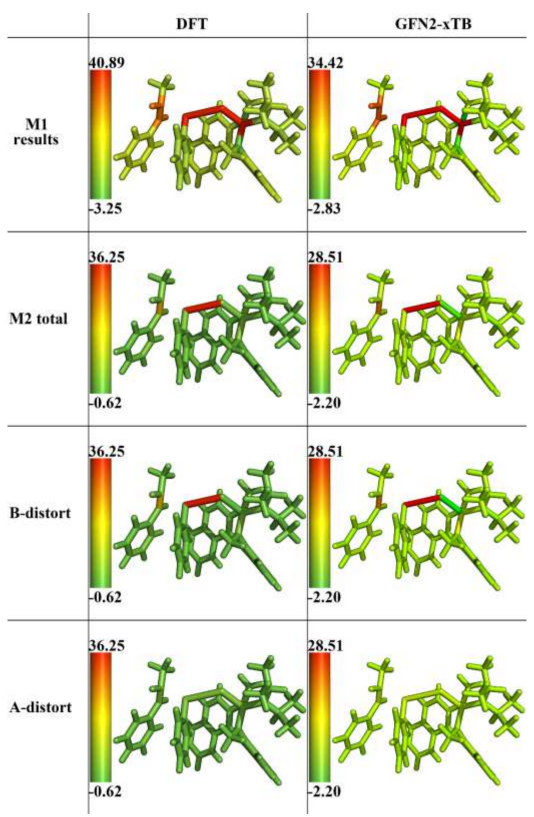


Figure S48 Distortion energy distribution (kcal/mol in log scale) of Lewis-base catalyzed sulfeno-functionalization using **M1** scheme and total distortion and its contribution from bond and angle terms using **M2** scheme. All energies were computed using DFT (SMD B3LYP/6-31G(d)) and GFN2-xTB methods.

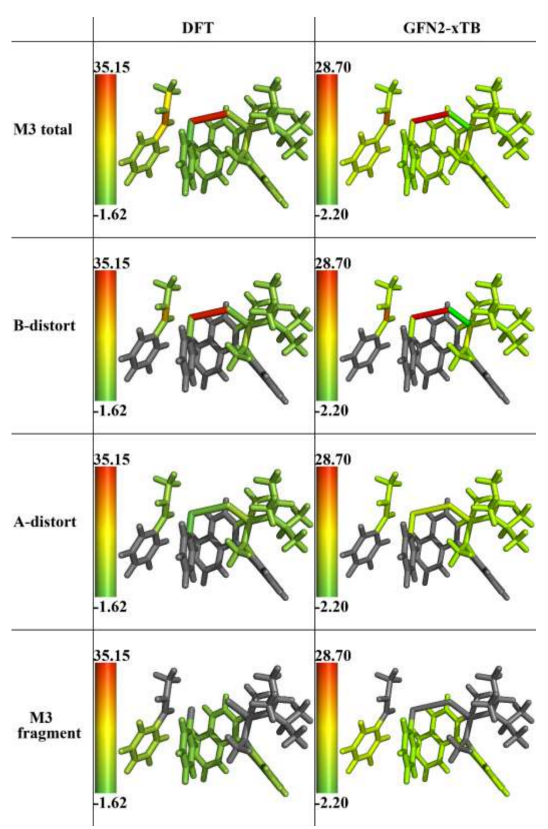


Figure S49 Distortion energy distribution (kcal/mol in log scale) of Lewis-base catalyzed sulfeno-functionalization using **M3** scheme and its contribution from bond, angle and fragment terms. All energies were computed using DFT (SMD B3LYP/6-31G(d)) and GFN2-xTB methods.

5.3. Excited system

5.3.1. Radical Sulfinyl-Smiles Rearrangement

The preceding intermediate and its corresponding TS for an asymmetric radical sulfinyl-Smiles rearrangement were taken from ref. S23 for D2AF analysis using DFT (IEFPCM B3LYP/6-31+G(d, p)), GFN2-xTB and ANI-2x methods.

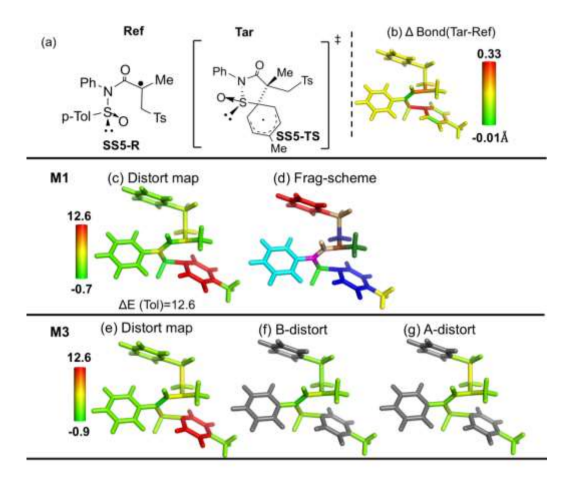


Figure S50 (a) Overview of the radical sulfinyl-Smiles rearrangement. (b) Bond length change (Δ bond in Å) from the reference form (**Ref**; R, **SS5-R**) to the target form (**Tar**; TS, **SS5-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at IEFPCM B3LYP/6-31+G(d, p) level.

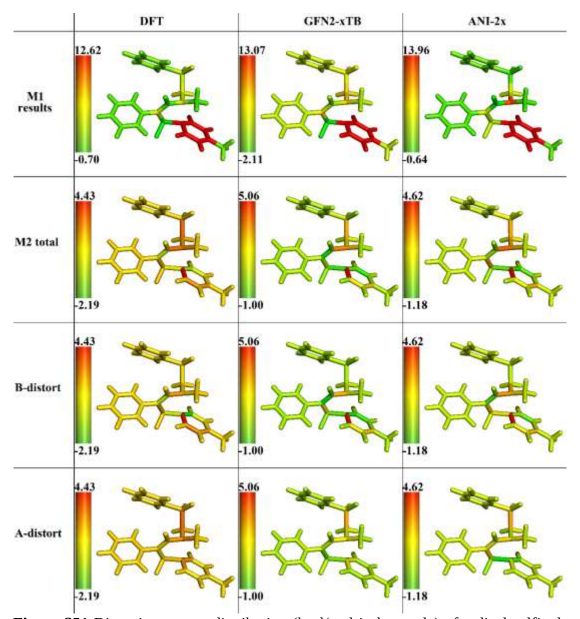


Figure S51 Distortion energy distribution (kcal/mol in log scale) of radical sulfinyl-Smiles rearrangement using **M1** scheme and total distortion and its contribution from bond and angle terms using **M2** scheme. All energies were computed using DFT (IEFPCM B3LYP/6-31+G(d, p)), GFN2-xTB and ANI-2x methods.

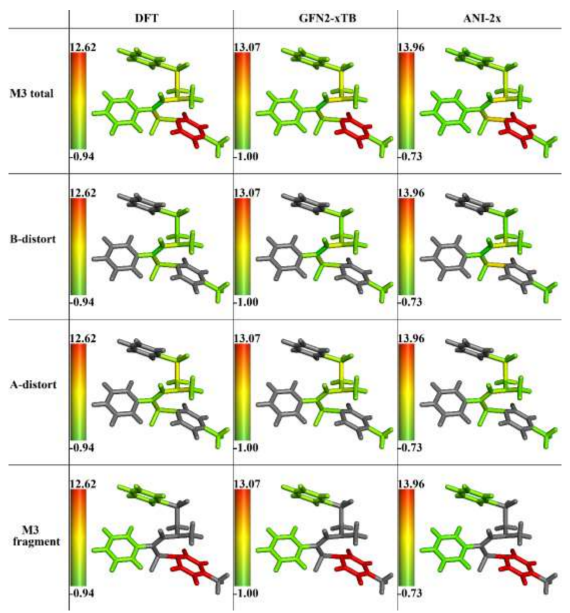


Figure S52 Distortion energy distribution (kcal/mol in log scale) of radical sulfinyl-Smiles rearrangement using **M3** scheme and its contribution from bond, angle and fragment terms. All energies were computed using DFT (IEFPCM B3LYP/6-31G(d)), GFN2-xTB and ANI-2x methods.

5.4. Supramolecular system

5.4.1. Diels-Alder Reaction in Pd-Cage

Self-assembled cages are novel supramolecular platforms to explore certain catalysis. S24 In this section, a Diels-Alder addition of two dienes (denoted as q1, d1 in the ref. S24) within a [Pd(pyridine)₄]²⁺ cage (denoted as C-1 in the in ref. S24) is considered. The reactant structures were taken from the ref. S24 as **Ref** and **Tar** for D2AF analysis using DFT (M06-2X/def2-TZVP) and GFN2-xTB methods.

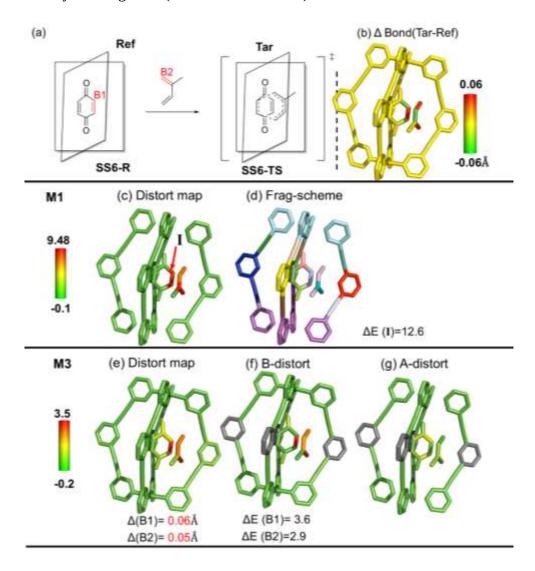


Figure S53 (a) Overview of the Diels-Alder reaction within the Pd-cage. (b) Bond length change (Δ bond in Å) from the reference form (**Ref**; R, **SS6-R**) to the target form (**Tar**; TS, **SS6-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed at M06-2X/def2-TZVP level.

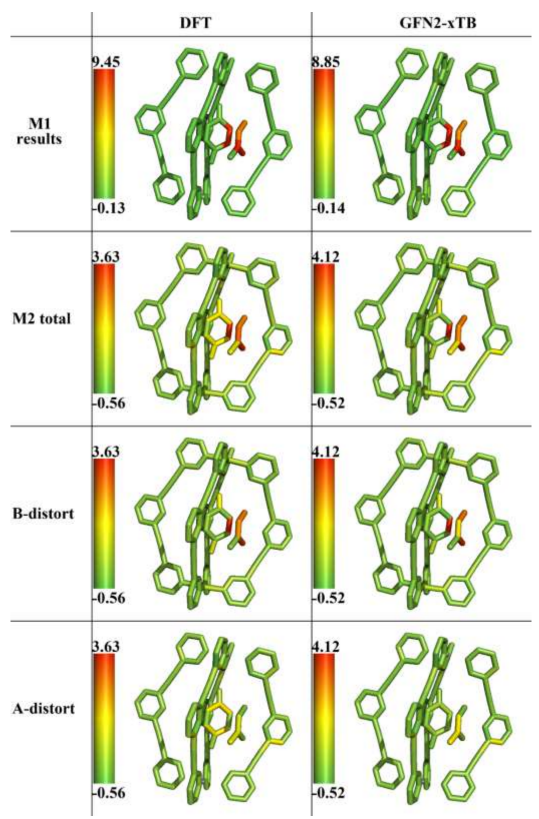


Figure S54 Distortion energy distribution (kcal/mol in log scale) of Diels-Alder reaction inside the Pd-cage using M1 scheme and total distortion and its contribution from bond and angle terms using M2 scheme. All energies were computed using DFT (M06-2X/def2-TZVP) and GFN2-xTB methods.

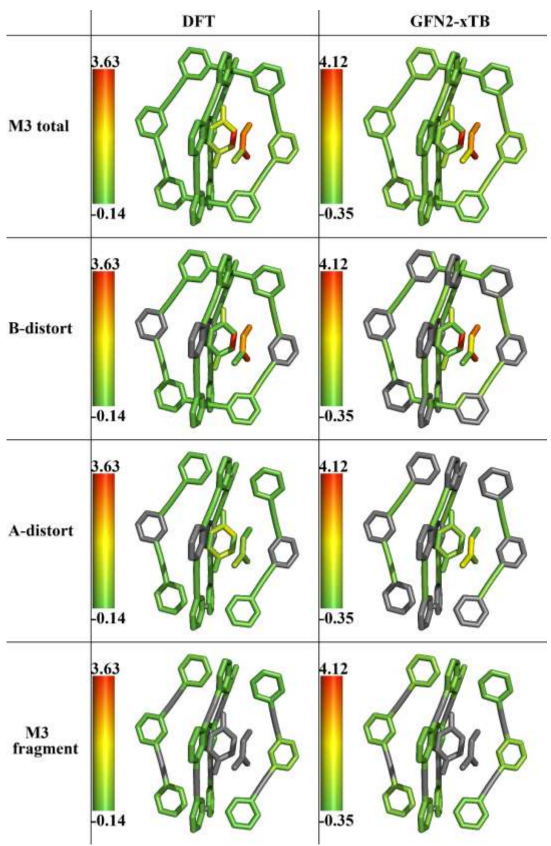


Figure S55 Distortion energy distribution (kcal/mol in log scale) of Diels-Alder reaction in the Pd-cage using **M3** scheme and its contribution from bond, angle and fragment terms. All energies were computed using DFT (M06-2X/def2-TZVP) and GFN2-xTB methods.

5.5. Coordination systems

5.5.1. Palladium-Catalyzed Cross-Coupling Reaction

A famous Pd-catalyzed cross-coupling reaction S25 was taken as a simple example of our coordination systems. The structures of reactant and catalyst ($C_5H_3NCl_2$ and $Pd(PH_3)_2$) and corresponding transition state taken from ref. S25 were treated as **Ref** and **Tar** for our D2AF analysis using DFT (B3LYP/(Lanl2DZ for Pd and 6-31G(d) for other elements)) method.

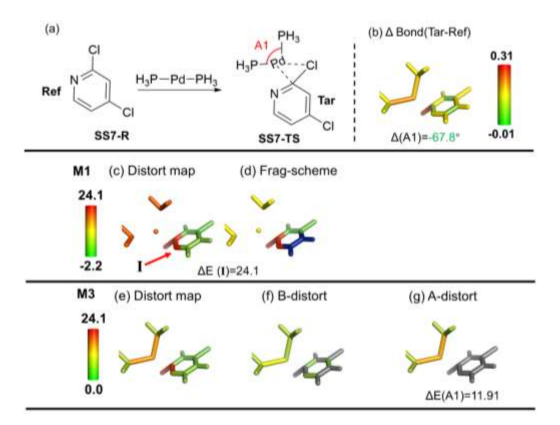


Figure S56 (a) Overview of the Pd-catalyzed cross-coupling reaction. (b) Bond length change (Δbond in Å) from the reference form (**Ref**; R, **SS7-R**) to the target form (**Tar**; TS, **SS7-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (ΔE) are also given. All energies were computed at B3LYP/(Lanl2DZ for Pd and 6-31G(d) for other elements) level.

5.5.2. Al-catalyzed Ring-opening Transesterification

The effects of a series of Al complexes which can catalyze ring-opening transesterification polymerization reactions were studied. Among these Al complexes, one TS and its precomplex were taken from ref. S26 as $\bf Tar$ and $\bf Ref$ for our D2AF analysis using DFT (SMD M06-L/6-31+G(d, p)) and GFN2-xTB methods.

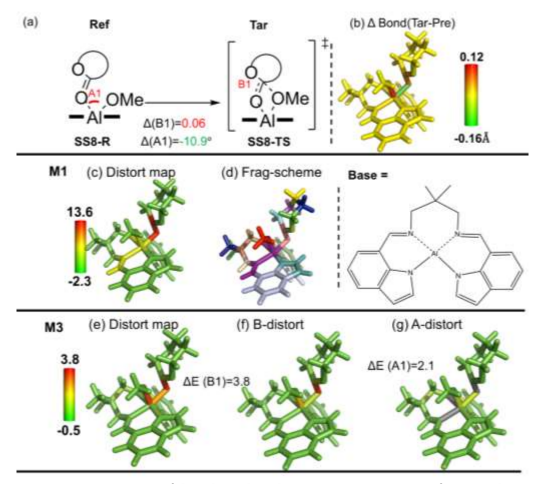


Figure S57 (a) Overview of the Al-catalyzed ring-opening transesterification. (b) Bond length change (Δ bond in \mathring{A}) from the reference form (**Ref**; Pre, **SS8-R**) to the target form (**Tar**; TS, **SS8-TS**). (c) Distortion energy distribution (kcal/mol) and (d) fragmentation using **M1** scheme. (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (Δ E) are also given. All energies were computed using DFT (SMD M06-L/6-31+G(d, p)) method.

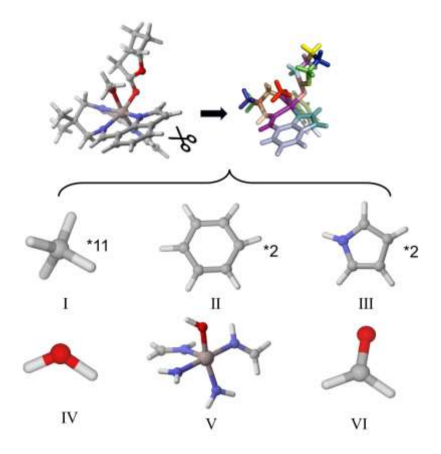


Figure S58 All molecular fragments generated by the M1 scheme from the Alcatalyzed ring-opening transesterification (SS8)

As shown in Figure S58, the whole system of the Al-catalyzed system is divided into six kinds of fragments using the M1 scheme. Eleven fragments I come from eleven sp³ carbon atoms. Two fragments II and two fragments II come from separation of two indoles in order to localize the distortion. The fragments IV and VI come from the sp³ oxygen and carbonyl within hexano-6-lactone. The specialized fragment V, which includes the Al center, along with its coordination nitrogen atoms and N=CH₂ are set to capture the crucial coordination interactions.

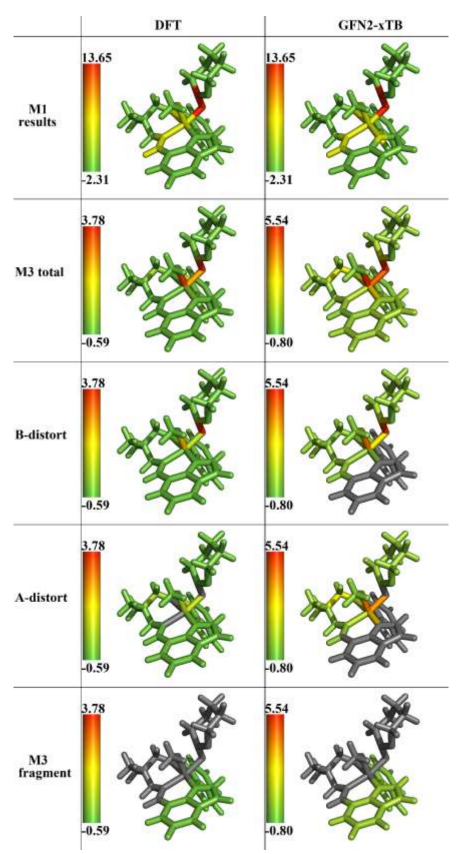


Figure S59 Distortion energy distribution (kcal/mol in log scale) of Al-catalyzed ring-opening transesterification using M1 scheme and total distortion and its contribution from bond, angle and fragment terms using M3 scheme. All energies were using DFT (SMD M06-L/6-31+G(d, p)) and GFN2-xTB methods.

5.6. Biochemical systems

5.6.1. Ni-Dependent Lactate Racemase

A Ni-dependent lactate racemase (LarA) and its unique tethered NAD-like cofactor was studied. S27 The structure of TS and its corresponding reactant (denoted TS_{C4-LS} and RC_{LS} in the ref. S27) were taken from the ref. S27 as Tar and Ref for our D2AF analysis using DFT (SMD M06/6-311G+(d, p)) and GFN2-xTB methods.

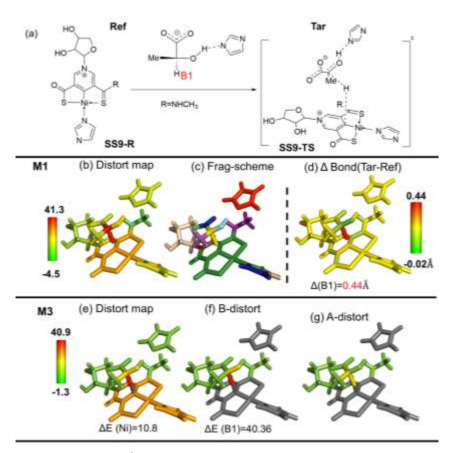


Figure S60 (a) Overview of the Ni-dependent lactate racemase. (b) Distortion energy distribution (kcal/mol) and (c) fragmentation using **M1** scheme. (d) Bond length change (Δbond in Å) from the reference form (**Ref**; R, **SS9-R**) to the target form (**Tar**; TS, **SS9-TS**). (e) Total distortion distribution (kcal/mol), its distortion contribution from (f) bond and (g) angle terms using **M3** scheme. The key geometrical changes and their corresponding distortion energy (ΔE) are also given. All energies were computed at SMD M06/6-311G+(d, p) level.

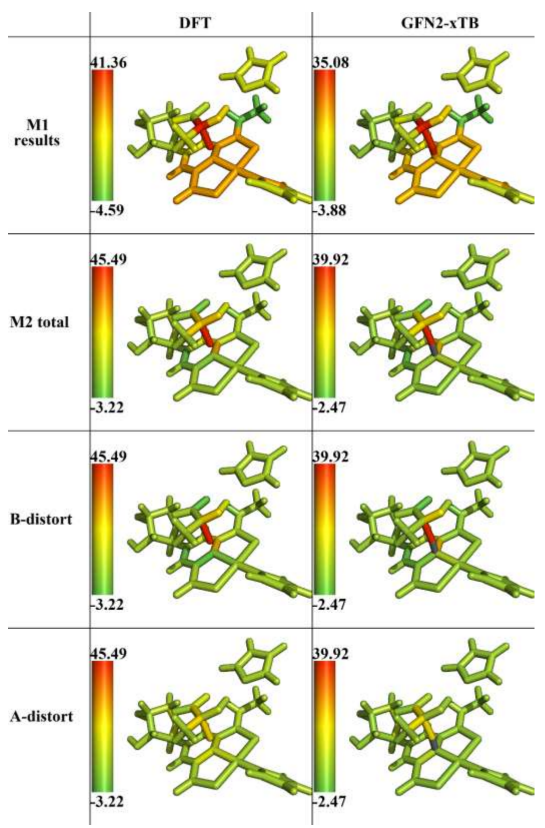


Figure S61 Distortion energy distribution (kcal/mol in log scale) of Ni-dependent lactate racemase using M1 scheme and total distortion and its contribution from bond and angle terms using M2 scheme. All energies were computed using DFT (SMD M06/6-311G+(d, p)) and GFN2-xTB methods.

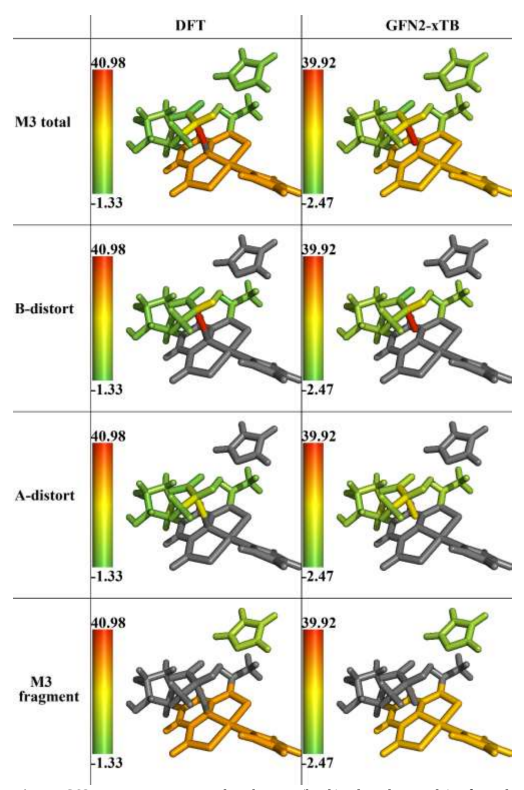


Figure S62 Distortion energy distribution (kcal/mol in log scale) of Ni-dependent lactate racemase using **M3** scheme and its contribution from bond, angle and fragment terms. All energies were computed using DFT (SMD M06/6-311G+(d, p)) and GFN2-xTB methods.

5.6.2. Oseltamivir in Influenza Neuraminidase

The drug Oseltamivir in influenza neuraminidase from the X-ray and QR refined structures were studied, in which the latter one were obtained using ONIOM(ω B97-D3/6-31G(d):GFN2-xTB:Amber) method. The optimized drug structure in the gas phase using ω B97X-D3/6-31G(d) was set as **Tar**, and X-ray and QR refined structures were set as two **Tar** for D2AF analysis using DFT (ω B97-D3/6-31G(d)) GFN2-xTB, ANI-2x and AIQM1 methods.

Figure S63 Oseltamivir structure



Figure S64 (a) Fragmentation scheme of Oseltamivir using different colors. Structures (optimized structure in the gas phase using $\omega B97X-D3/6-31G(d)$, X-ray structure and QR refined structure) taken from the ref. S28. Difference of bond length of (b) X-ray structure and (c) QR-refined structure relative to the optimized structure in the gas phase (color bar green to red: red positive, green negative, yellow: no change).

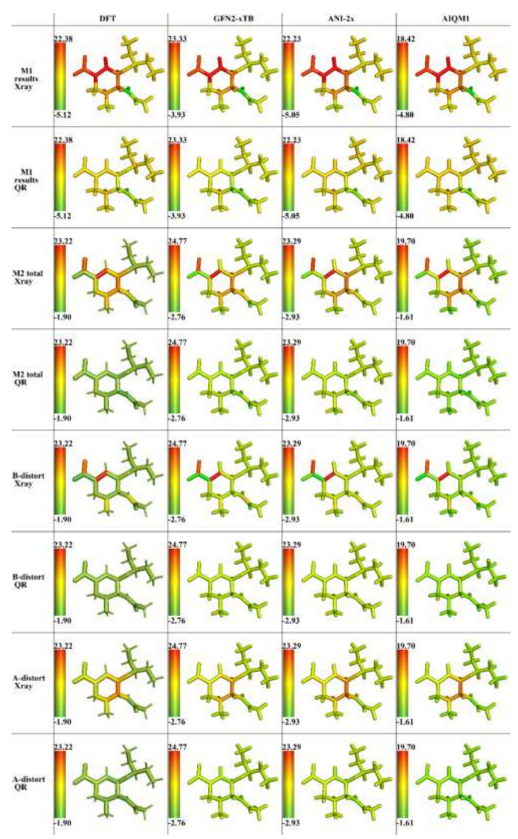


Figure S65 Distortion energy distribution (kcal/mol in log scale) of Oseltamivir's two different **Tar** forms (X-Ray and QR refined structures) including **M1** and **M2** distortion map. Bond and angle contribution of two forms in **M2** are also presented. All energies were computed using DFT (ω B97X-D3/6-31G(d)), GFN2-xTB, ANI-2x and AIQM1 methods.

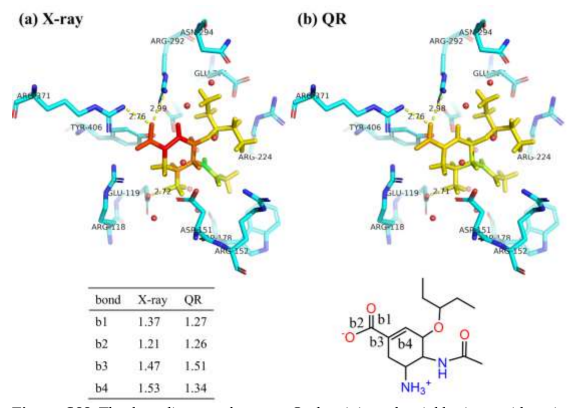


Figure S66 The key distances between Oseltamivir and neighboring residues in influenza neuraminidase from the X-ray and QR refined structures. The key bond lengths (\mathring{A}) are listed. Oseltamivir is colored based on distortion distribution using M1 scheme while the optimized structure in the gas phase were set as the reference form.

5.6.3. Si-H Insertion enabled by Artificial Heme Enzyme

Directed-evolution artificial heme enzymes pioneered by the Arnold group play a critical role in new biocatalysis. S29 They recently developed an artificial heme protein to catalyze novel Si-H insertion (Figure S67a), which is selected as an example of our biological systems. S30 As proposed previously, Figure S67a shows the simplified coordination region/fragment as the core heme part. In our work, the specialized coordination region includes the heme region (the red part in Figure S67b), the ligated C atom of the ester and the truncation part (NHCH₂) of the ligated imidazole to study the heme protein catalyzed Si-H insertion. This region keeps coordinating interactions around central Fe atom.

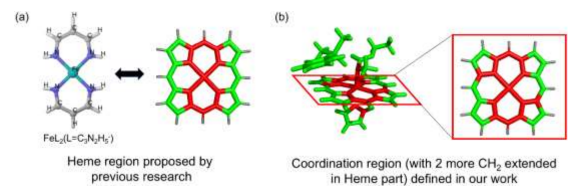
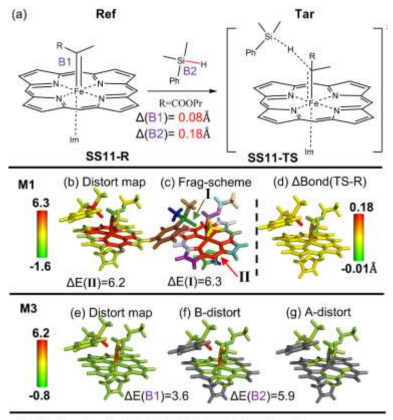


Figure S67 (a) Heme region proposed by previous research. (b) The specialized coordination region defined in our work (with two more CH₂ extension from the previous model (a) in the heme part as well as the truncated ligated parts) for the hemecatalyzed Si-H insertion. The remaining heme parts (green) are treated by the **M1** scheme with each fragment consisting of two sp2 carbon atoms and their hydrogen atoms.

To reduce computational cost and get a higher "resolution" distortion map, a simplified heme region (red fragment \mathbf{II} in Figure S68c; see Figure S67 for the detailed fragments) is utilized as one fragment which was previously proposed to represent the heme's coordination interaction. Our $\mathbf{M1}$ analysis results unveil that the reacting Si-H part (fragment \mathbf{II}) and the heme coordination region (fragment \mathbf{II}) mainly contribute to distortion energy (~6.3 and ~6.2 kcal/mol, respectively, Figures S68b-S68c). Furthermore, as shown in Figure S67d, our $\mathbf{M3}$ results illustrate a comparable distortion energy of ~3.6 and ~5.9 kcal/mol for the elongation of the Fe-C bond ($\Delta(\mathbf{B1})$: 0.08 Å) and Si-H bond ($\Delta(\mathbf{B2})$: 0.18 Å), respectively. Consequently, $\mathbf{M1}$ exhibits distortion of the larger coordination region, while $\mathbf{M3}$ further "localizes" the main distortion into the Fe-C and Si-H bonds.



*ΔE in kcal/mol at B3LYP/def2-TZVP, CPCM

Figure S68 (a) Overview of the heme protein catalyzed Si-H insertion (structure from ref S30). (b) Distortion distribution (kcal/mol) and (d) fragmentation using M1 scheme. (d) Bond length change (Δbond in Å) from the reference form (Ref; reactant, SS11-R) to the target form (Tar; TS, SS11-TS). (e) Total distortion distribution (kcal/mol), distortion contribution from (f) bond and (g) angle using M3 scheme. The key geometrical changes and their corresponding distortion energy (ΔE) are also given.

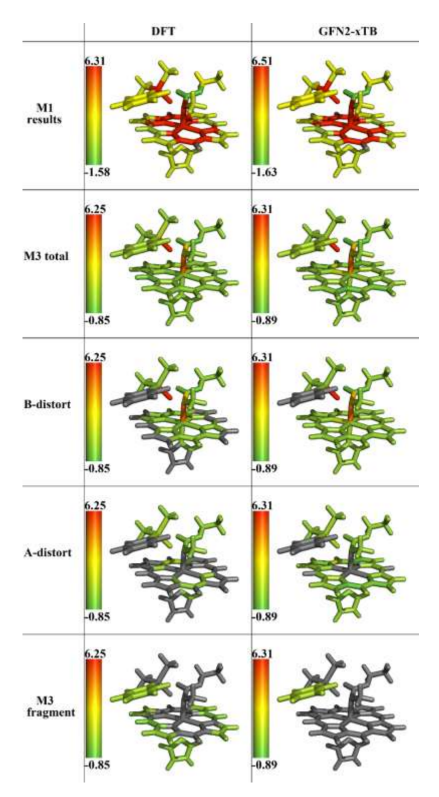


Figure S69 Distortion energy distribution (kcal/mol in log scale) of the Si-H insertion enabled by artificial Heme enzyme (**SS11**) using **M1** scheme and total distortion and its contribution from bond, angle and group fragment terms using **M3** scheme. Energy calculations were computed using DFT (CPCM B3LYP/def2-TZVP) and GFN2-xTB methods

5.7. Multiple-structures System

5.7.1. S_N2 (IRC)

An IRC calculation of one S_N2 reaction (NC⁻ and CH₃Cl) was also applied to show the change in (relative) distortion distribution during the reaction process. The structures (without external electric fields case) were taken from the ref. S15 as the target states for our D2AF analysis throughout the reaction process using DFT(PCM M06-2X/6-31+G(d)), GFN2-xTB and ANI-2x methods.

$$NC^{-} + CH_{3}C1 \longrightarrow \left[NC^{-}C^{-} - C1\right]^{\ddagger} \longrightarrow NCCH_{3} + C1^{-}$$

$$R \qquad TS \qquad P$$

Figure S70 Scheme of a S_N2 reaction

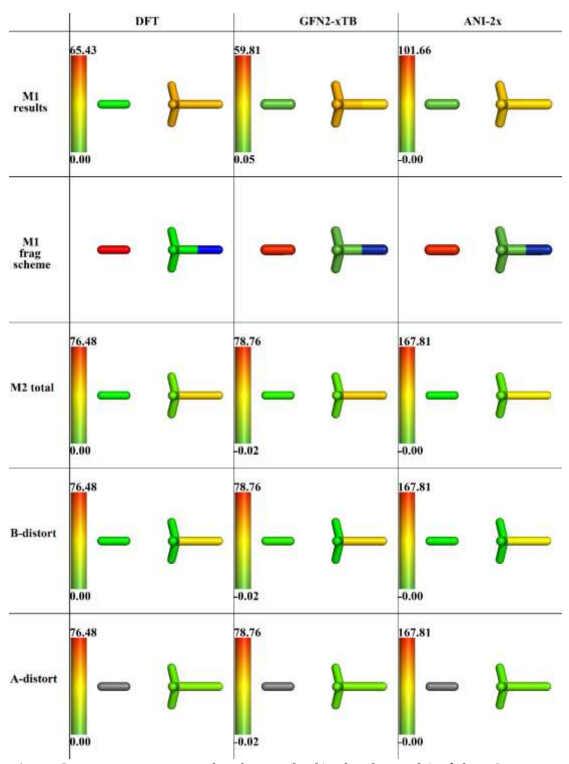


Figure S71 Distortion energy distribution (kcal/mol in log scale) of the TS structure for a $S_{\rm N}2$ reaction using **M1** scheme, total distortion and its bond and angle contributions using **M2** scheme. All energies were computed using DFT (PCM M06-2X/6-31+G(d)), GFN2-xTB and ANI-2x methods.

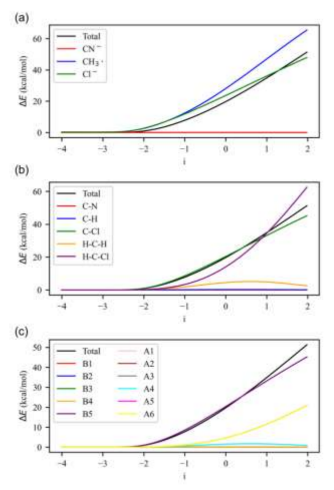


Figure S72 Minimum energy path (MEP) for the S_N2 reaction from the IRC calculations. (a) Distortion distribution changes using **M1** scheme. (b) Distortion distribution changes using **M2** scheme (with summed distortion from the same kind coordinates) (c) Distortion distribution changes using **M2** (B1: C-N; B2, B3, B4: C-H; B5: C-Cl; A1, A2, A4: H-C-H; A3, A5, A6: H-C-Cl). All energies were computed using DFT (PCM M06-2X/6-31+G(d)) method.

5.8. Different Computational Methods

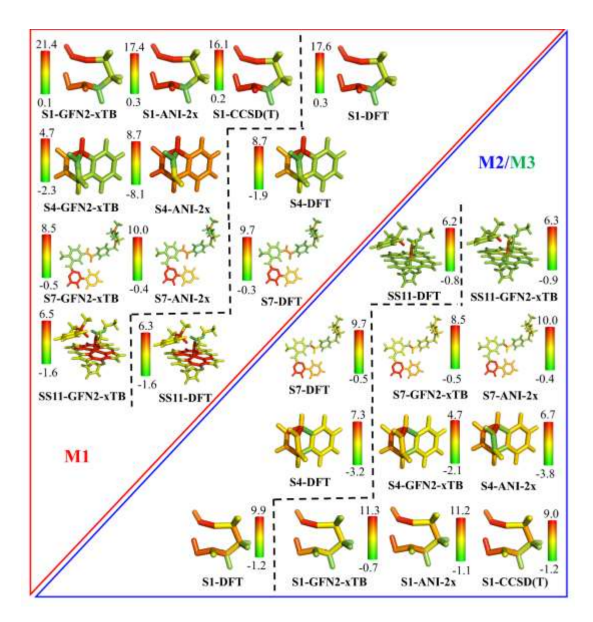


Figure S73 Distortion distribution (kcal/mol) of the four selected systems calculated by DFT, SE (GFN2-xTB), MLP (ANI-2x) and CCSD(T) (only for **S1**) methods

Our flexible local distortion analysis can also be applied to different computational chemistry methods and/or software packages to estimate fragments' energies. A few methods (highly-accurate CCSD(T), cost-effective semi-empirical GFN2-xTB or machine learning potentials (MLPs, e.g., ANI-2x) methods) were employed to evaluate the distortion maps of our four representative systems (Figure S73). To the best of our knowledge, this is the first study using CCSD(T) or MLP to evaluate distortion energy. Generally, GFN2-xTB shows consistent distortion maps

with the DFT- or CCSD(T)-computed results. The ANI-2x method performs well except for the challenging triplet-state system (S4), due to no spin systems used for ANI-2x training. Therefore, the efficient GFN2-xTB method is potentially used for a wider range of systems, while MLPs could be applied to some neutral ground-state systems.

Reference:

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6. Cartesian coordinates and key absolute energy of the new optimized structures

Strı	uctures for benchmark:	O -0.714904 -0.000813 -0.141154
The following structures are optimized		H -1.115593 0.004269 0.738466
	M06-2X/6-31G(d) level.	N 0.680407 0.000720 0.156000
C_2F	I ₆ (Benchmark, Fig. S1)	H 1.036236 -0.812337 -0.347016
$\mathbf{E} =$	-79.83222 AU	H 1.035735 0.809530 -0.354219
C	0.000000 0.000000 0.765684	
Н	-0.510282 0.884734 1.164342	H ₃ P ₂ O ₄ (Benchmark, Fig. S5-7)
Н	1.021343 -0.000450 1.164342	E = -644.234007 AU
Н	-0.511061 -0.884284 1.164342	O -0.013386 -0.030175 -1.571957
C	0.000000 0.000000 -0.765684	P -0.000734 -0.002017 -0.109174
Н	0.510282	O -1.257511 -0.636653 0.598796
Н	0.511061 -0.884284 -1.164342	O 1.197297 -0.757850 0.581337
Н	-1.021343 -0.000450 -1.164342	O 0.076895 1.427118 0.550237
		H -1.607481 -1.419724 0.154190
C ₃ F	I ₈ (Benchmark, Fig. S2, S4)	H 2.040265 -0.670355 0.117490
$\mathbf{E} =$	-119.148581 AU	H -0.448134 2.100820 0.098625
C	1.275119 -0.260775 0.000000	
Н	2.176122 0.363318 -0.000001	C ₂ H ₄ (Benchmark, Fig. S8)
Н	1.315335 -0.908297 -0.884815	E = -78.536838 AU
Н	1.315335 -0.908296 0.884816	C 0.000021 -0.663656 0.000000
C	-0.000157 0.589409 0.000000	Н 0.923927 -1.234268 0.000000
Н	0.000075 1.249746 -0.878170	H -0.923937 -1.234282 0.000000
Н	0.000075 1.249747 0.878169	C 0.000021 0.663586 0.000000
C	-1.274882 -0.260638 0.000000	H -0.923956 1.234302 0.000000
Н	-1.315481 -0.908505 0.884715	H 0.923716 1.234670 0.000000
Н	-2.176462 0.362812 0.000002	
Н	-1.315483 -0.908503 -0.884716	CH ₂ =NCH ₃ (Benchmark, Fig. S11, S12)
		E = -133.868779 AU

C -1.173590 0.184240 -0.000007

N -0.140505 -0.540887 0.000003

NH₂OH(Benchmark, Fig. S3)

E = -131.644293 AU

C	1.141609	0.135855	-0.000003	
Н	-2.156790	-0.286757	0.000002	
Н	-1.144910	1.284022	0.000021	
Н	1.707547	-0.182489	0.880409	
Н	1.707640	-0.182646	-0.880296	
Н	1.061932	1.233512	-0.000100	

CH₂=CH-CH=CH₂(Benchmark, Fig. S13,S14)

E = -155.899155 AUC 1.523110 -0.493068 0.088557 1.133676 -1.426160 0.486821 2.586688 -0.449542 -0.121140 0.726761 0.556885 -0.116569 1.161693 1.492798 -0.464433 C -0.726761 0.556885 0.116569 H -1.161693 1.492798 0.464433 C -1.523110 -0.493068 -0.088557 H -1.133676 -1.426160 -0.486821 H -2.586688 -0.449542 0.121140

CH₂=C(CH₃)₂ (Benchmark, Fig. S15)

E = -157.133604 AUC 0.000012 0.124582 -0.000022 C -1.273105 -0.678192 0.000003 H -1.320240 -1.330943 0.880022 H -2.156747 -0.035539 -0.000076 H -1.320174 -1.331044 -0.879939 1.272947 -0.678456 -0.000001 1.319909 -1.331212 0.880020 1.319923 -1.331286 -0.879958 2.156724 -0.035977 -0.000006 0.000168 1.456531 0.000015

H 0.926179 2.024493 -0.000025 H -0.925704 2.024724 -0.000008 C5H8(Benchmark, Fig. S16)

E = -194.900537 AU

C 0.000000 0.000000 0.000000 0.000000 0.000000 1.070000 1.008806 0.000000 -0.356667 C -0.889119 -1.088945 -0.628702 H -0.664527 -2.121923 -0.463131 C -0.889119 1.088944 -0.628703 H -0.714669 2.118964 -0.397346 C -1.891735 0.747129 -1.473979 H -1.668246 0.357601 -2.445174 H -2.910337 0.867188 -1.169126 C -1.955024 -0.743396 -1.390943 H -1.889943 -0.784264 -2.458180 H -2.864813 -0.428619 -0.923931

1-butyne (Benchmark, Fig. S16)

E = -155.883820 AU

C -0.169625 0.045234 -0.122600 H -0.164117 0.046092 0.973172 0.879547 0.047460 -0.438750 C -0.816286 1.271196 -0.599463 C -1.366620 2.263150 -1.005261 H -1.849979 3.144881 -1.361538 C -0.869062 -1.217358 -0.640814 Н -0.859437 -1.240936 -1.733132 H -0.364043 -2.112743 -0.269010 H -1.910182 -1.242835 -0.310352

Structures for S3:

The following single point energies are calculated at SCS-MP2/6-31G(d) level by Gaussian16.

S3*-Reactant-C2Me2 (Fig. 4)

C	0.604438	0.000003	-0.000063
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S3*-Reactant-PhCH₂N₃ (Fig. 4)

E = -580.920661 AU

C -0.604450 0.000119 -39.999677

S3_{model}-Reactant- C₂Me₂ (Fig. 4)

E = -153.271224 AU

H -2.641366 -0.833435 -1.050950

S3_{model}-TS (Fig. 4)

E = -580.879329 AU

C -2.065989 -0.000029 -40.000091

N	-0.046953	-2.521043	-0.964919	Н	-0.846511	2.452315	-0.915660
C	-1.959480	0.334571	-1.605097	C	-1.546852	0.595107	-0.058162
Н	-1.650165	1.377907	-1.689160	C	-1.166546	-0.708916	0.279984
Н	-2.498258	0.072890	-2.525295	C	-2.902299	0.873830	-0.271455
C	-2.855664	0.136584	-0.398069	C	-2.128980	-1.712774	0.406030
C	-2.462304	0.605153	0.863905	Н	-0.116773	-0.940771	0.432649
C	-4.081736	-0.524469	-0.525465	C	-3.864117	-0.128034	-0.141666
C	-3.285483	0.413318	1.972387	Н	-3.207061	1.882310	-0.544713
Н	-1.508025	1.114877	0.964911	C	-3.479083	-1.426857	0.198350
C	-4.908111	-0.714582	0.584597	Н	-1.819801	-2.722307	0.664273
Н	-4.393737	-0.893144	-1.500556	Н	-4.912049	0.103474	-0.313827
C	-4.510623	-0.246073	1.836587	Н	-4.225837	-2.210457	0.294573
Н	-2.972206	0.780968	2.946353	C	3.823244	-1.543991	-0.338519
Н	-5.858456	-1.229447	0.469572	Н	4.513165	-1.110514	-1.070949
Н	-5.150487	-0.393184	2.702821	Н	4.409748	-1.935094	0.501371
C	2.519099	-2.333933	0.376089	Н	3.319527	-2.382132	-0.832244
Н	3.570307	-1.959774	0.428031	C	1.839444	0.809725	2.195784
Н	2.482657	-3.259910	-0.209253	Н	0.783721	0.609882	2.416109
Н	2.169814	-2.599387	1.384373	Н	2.437593	0.418766	3.028417
C	1.609821	1.291962	-0.479130	Н	1.971706	1.896720	2.161840
Н	2.392394	1.525255	0.287523				
Н	2.004885	1.533473	-1.481410	$S3_1$	model-P (Fig	g. 4)	
Н	0.641053	1.934767	-0.302558	E =	-581.039	276 AU	
				С	2.815363	-1.449917	1.203873
S3 ²	*-TS (Fig.	4)		C	1.193882	1.210024	-0.323396
E =	= -580.887	864 AU		C	0.878727	-0.245571	-0.153391
C	2.247220	0.182674	0.929575	C	1.519588	-1.314724	0.454338
C	2.843937	-0.540508	0.118292	N	-0.257727	-0.801049	-0.676947
N	0.851203	1.310451	-0.401869	N	-0.317094	-2.120182	-0.417583
N	1.127736	0.660492	-1.453222	N	0.754488	-2.432603	0.257598
N	1.928692	-0.122006	-1.796685	C	-1.377644	-0.165498	-1.372512
C	-0.528893	1.721043	-0.159810	Н	-1.061621	0.847511	-1.631702
Н	-0.483295	2.267461	0.790180	Н	-1.537470	-0.716934	-2.303111

C	-2.651853	-0.135942	-0.550793
C	-2.678223	0.483020	0.705822
C	-3.819746	-0.714363	-1.054375
C	-3.861294	0.519854	1.442015
Н	-1.767989	0.925011	1.098701
C	-5.006464	-0.671098	-0.319192
Н	-3.801067	-1.208941	-2.022781
C	-5.029264	-0.053735	0.931290
Н	-3.872460	0.997511	2.418360
Н	-5.907358	-1.127296	-0.720755
Н	-5.949818	-0.023428	1.508189
Н	3.737327	-1.428085	0.569113
Н	2.759673	-2.423746	1.701153
Н	2.891963	-0.699661	2.000401
Н	2.226086	1.481588	-0.022833
Н	1.072605	1.468603	-1.391131

H 0.391242 1.812233 0.295391

S3*-P (Fig. 4)

E = -581.048768 AU

L = -301.040700 AU					
C	3.801952	-0.407883	-1.351129		
C	1.129452	-2.045973	-0.159460		
C	1.632328	-0.660136	0.071213		
C	2.728072	0.035605	-0.408187		
N	1.006681	0.239406	0.882112		
N	1.663185	1.424861	0.893637		
N	2.704473	1.297129	0.118268		
C	-0.259524	0.101011	1.586532		
Н	-0.239533	-0.822882	2.174512		
Н	-0.296243	0.942731	2.284379		
C	-1.449678	0.113638	0.645423		
C	-2.440578	-0.868099	0.731230		
C	-1.564397	1.122957	-0.319236		

C	-3.541195	-0.840139	-0.129102
Н	-2.352681	-1.658975	1.473245
C	-2.659906	1.149094	-1.180345
Н	-0.786764	1.879010	-0.391929
C	-3.652024	0.168304	-1.086588
Н	-4.305428	-1.609034	-0.053910
Н	-2.740533	1.935361	-1.926056
Н	-4.504974	0.189795	-1.759460
Н	3.944794	-1.493162	-1.314715
Н	4.750176	0.072815	-1.092785
Н	3.566886	-0.137527	-2.388460
Н	1.814171	-2.590127	-0.815178
Н	1.044407	-2.608964	0.778933

Structures of $S_N 2$ MD:

The following energy is calculated at HF/3-21G level.

H 0.138339 -2.039101 -0.628810

S_N 2-TS (MD, Fig. 9)

E = -954.059827 AU

 C
 0.000000
 0.000000
 0.000000

 H
 0.000000
 1.060526
 0.000000

 H
 0.918442
 -0.530263
 0.000000

 H
 -0.918442
 -0.530263
 0.000000

 CI
 0.000000
 0.000000
 2.395711

 CI
 0.000000
 0.000000
 -2.395711