Tandem Cycloaddition-Cycloreversion of 2-pyrone and 1,4-oxazinone with Acetylene - A DFT Insight

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Abstract: Reaction of either 2-pyrone or 1,4-oxazinone with acetylene follows the sequence of cycloaddition - cycloreversion through concerted mechanism. Transition states for both cycloaddition and cycloreversion pathways have been obtained in both the cases by modelling the reactions at B3LYP/6-31g (d) level. Cycloreversion is faster than cycloaddition in the case of 2-pyrone due to the enhancement of aromaticity resulting the product as benzene. In contrast, oxazinone has rapid cycloaddition. It is ascribed to the presence of nitrogen in this system. Removal of either CO₂ or HCN is plausible in this mechanism to complete the reaction. Even though two pathways are feasible for cycloreversion, CO₂ extrusion is more preferable than HCN elimination. In these two studied molecules, there is an enhancement of aromaticity up to transition states like any other pericyclic reaction and further it diminishes during cycloaddition. Further, aromaticity is specifically augmented in cycloreversion phase during CO₂ elimination resulting to yield pyridine whereas competitive HCN elimination results in the formation of 2-pyrone which is less facile. In both the molecules the aromatic enhancement of the cycloreversion is substantiated through the study of magnetic susceptibility of the ring fragment along the reaction coordinate. Further the study also reveals the effect of halogen substituted at different carbons of 2-pyrone ring.

Keywords: Aromaticity, Cycloaddition-cycloreversion Reactin, DFT Study, Intrinsic Reaction Coordinate

1. Introduction

Tandem cycloaddition-cycloreversion (CA-CR) is commonly known in molecules whose cycloadducts have a potential extruding group. Some of the well-known tandem reactions of this type reported in the literature include cycloaddition of a variety of alkynes to pyrimidines resulting in cycloadducts that spontaneously extrudes nitrile to form pyridines [1]. A similar nitrile expulsion has been reported with oxazoles [2]. Pyrone undergoes cycloaddition with alkene/alkyne to form a bicyclic lactone that spontaneously extrudes CO₂ to form molecules with cyclohexadiene/ benzene unit. First reaction of this kind was reported by Diels and Alder in 1931 [3]. The diene and dienophile in this reaction were 5-carbomethoxy-2-pyrone and maleic anhydride respectively. Such kind of reactions have been recognized as synthetically important and useful reactions [4, 5]. Cycloadditions afford in the first instance bridged bicyclic lactones. These functionally rich bridged cycloadducts are valuable starting materials for the synthesis of highly functionalized six-membered rings found in many natural products [6-12] such as lasalocid A, rufesine, imeluteine, chrysophanol, islandicin, emodin, sendaverine, juncusol, and norketoyobirine [13-17]. Owing to their partial aromatic nature, 2-pyrones participate in Diels-Alder reactions less readily than most cyclic conjugated homodiienes [18]. In general, high temperature and pressure is required for cycloadditions to take place. However, these cycloadducts are labile at such a high temperature and rapidly extrude CO₂. As the cycloadducts formed from olefinic dienophiles are good source of highly funtionalized and stereochemically rich compounds their isolation received the attention of many chemists. It has been found that utilization of
electronically matching partners is one of the strategies used not only for arresting the cycloreversion (loss of CO$_2$) in this reaction but also has the advantage of adding to the diversity of substituents on the cycloadducts [18a, d].

Though there has been extensive experimental study on Diels-Alder cycloadditions of 2-pyrone and its derivatives with alkenes and alkynes, enough theoretical insights have not been attempted at an equal pace. Cycloaddition reaction of 2-pyrone with acetylene has been reported only at HF level. Therefore, the present paper intends to explore the reaction with the following questions in mind: (i) what is the effect of aromaticity in these tandem cycloaddition-cycloreversion reactions? (ii) How does heteroatom (nitrogen) in 1, 4-oxazinone affect this tandem reaction (Does it accelerate or retard cycloaddition /cycloreversion)? (iii) There are two extrusible groups in the cycloadduct formed from oxazinone and acetylene reaction viz., CO$_2$ and nitrile; which of the two is preferentially extruded? In search of these answers the reaction path of the tandem processes have been scanned. Thermochemical activation parameters and magnetic susceptibility isotropy at every point along the Intrinsic Reaction Coordinate have been computed. Another interesting aspect in this work is the variation of aromaticity in the ring fragment of the diene unit during cycloaddition and forming product ring fragment during cycloreversion. This will help to relate the perturbation occurring in the ring moiety in terms of aromaticity during the tandem process.

2. Computational Methods

The computations have been done using Gaussian 98 program [19] at B3LYP [20, 21]/6-31G (d) level. The magnetic susceptibility isotropy has been calculated by computing the NMR shielding tensors at B3LYP/6-311+G (2d, p) using IGAIM method [22, 23] which is a slight variation of CSGT [22-24] method. NBO calculations [25-27] have been done. The variation in magnetic susceptibility isotropy ($\chi_{iso}$) along the reaction coordinate and has been monitored between the range (-1 to +1 (amu)$^{1/2}$ Bohr) for both cycloaddition and cycloreversion process. Frontier orbitals of both diene and dienophiles have also been analysed. Here, apart from this the loss of the aromaticity of the parent ring during cycloaddition and the gain of aromaticity of the newly formed ring during cycloreversion has also been calculated along the reaction coordinate by retaining the desired ring moiety alone.

3. Results and Discussion

The scheme 1 & 2 explain the steps of 2-pyrrone (2P) and 1,4-Oxazinone (OXZ) reacting with acetylene (Ace) respectively. From these schemes one can understand that cycloadditions take place in the first step resulting in bicyclic lactone adducts (A1 & A2). In the 2P-Ace reaction the adduct (A1) tends to eliminate CO$_2$ to form an aromatized product. In OXZ-Ace reaction, the adduct A2 formed has two paths for cycloreversion; one being CO$_2$ extrusion and another being HCN extrusion As similar systems have been proved to pass through a concerted path rather than the polar mechanism, the search here is restricted to concerted path only [28].

![Figure 1. Reaction pathways for addition of 2P with Ace.](image-url)
A. Energetics And FOE Analysis

In the 2P-Ace reaction there are 2 transition states-TSl & TS2; The former is the cycloaddition TS and the latter is cycloreversion TS. The relative free energy profile for 2P-Ace reaction and OXZ-Ace reaction are presented in Figures 3 & 4 respectively. Figure 3 indicates that the barrier to cycloaddition of 2P with Ace is 36.84 kcal.mol\(^{-1}\) while that for OXZ is 30.99 kcal.mol\(^{-1}\) (Figure 4). Both these barriers are much larger than the activation energy for the reaction of acetylene and cyclopentadiene (\(\Delta E^\ddagger=25.7\) kcal.mol\(^{-1}\)) [27]. Such a large difference among the barriers of these embedded dienes compared to cyclopentadiene is due to the partial aromaticity of 2P and OXZ. Nitrogen atom in OXZ has in fact reduced the aromaticity of the ring consequently has its free energy of activation 5.85 kcal.mol\(^{-1}\) lesser than 2P-Ace reaction. The bond distances of the newly forming bonds in TSl are unequal and the observed asynchronicity is due to the asymmetry of 2P. A2 has two extrusible groups viz., CO\(_2\) and HCN (Figure 2 and Figure 4). A2 gives pyridine on CO\(_2\) elimination. On the other hand, on HCN extrusion, it forms 2-pyrone. Energy profile presented in Figure 4 and the data presented in Table 2 show that cycloreversion by CO\(_2\) elimination is more facile than HCN elimination, kinetically and thermodynamically CO\(_2\) elimination yields pyridine an aromatic product, and therefore more stable, while HCN elimination leads to the formation of 2P that is partially aromatic and obviously less stable. Thus gain in aromaticity in CO\(_2\) elimination pathway reduces the activation energy greatly as reported [29]. It is noted that barrier to cycloreversion of A1 (11.97 kcal.mol\(^{-1}\)) and A2 (14.52 kcal.mol\(^{-1}\)) is rather much lower and the reaction is also thermodynamically favoured to a greater extent. In both cycloreversions the cleavage of C6—O1 is faster than C2—C3. But in HCN elimination pathway during cycloreversion of A2 there is larger asynchronicity (\(\Delta r=0.31\)) with N4—C3 cleaving much ahead of C5—C6.

The relative positioning of FOs of 2P with acetylene and OXZ with acetylene is shown in Figure 5. It could be seen that both the cases are inverse electron demand type (IED) in which electrons move from HOMO of acetylene to LUMO of either 2P or OXZ. This is verified from \(Q_{CT}\) presented in the Table 1.
Figure 3. Schematic free energy profile of 2P-Ace reaction computed at B3LYP/6-31G (d) level.

Figure 4. Schematic free energy profile of OXZ-Ace reaction computed at B3LYP/6-31G (d) level.
Figure 5. Frontier Orbital separation in diene and dienophile pairs.

Table 1. Lengths of forming and cleaving bonds during cycloaddition and cleaving during cycloreversion, asynchronicity (a). Quantum of charge transferred during cycloaddition of simple and substituted pyrones with acetylene.

<table>
<thead>
<tr>
<th>System</th>
<th>Cycloaddition</th>
<th>Cycloreversion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C6-C7</td>
<td>C3-C8</td>
</tr>
<tr>
<td>2P</td>
<td>2.15</td>
<td>2.33</td>
</tr>
<tr>
<td>3F2P</td>
<td>2.16</td>
<td>2.30</td>
</tr>
<tr>
<td>3Cl2P</td>
<td>2.11</td>
<td>2.36</td>
</tr>
<tr>
<td>3Br2P</td>
<td>2.12</td>
<td>2.34</td>
</tr>
<tr>
<td>4F2P</td>
<td>2.12</td>
<td>2.34</td>
</tr>
<tr>
<td>4Cl2P</td>
<td>2.12</td>
<td>2.34</td>
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<tr>
<td>4Br2P</td>
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<tr>
<td>5F2P</td>
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<td>5Cl2P</td>
<td>2.13</td>
<td>2.38</td>
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<tr>
<td>5Br2P</td>
<td>2.12</td>
<td>2.39</td>
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<td>6F2P</td>
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<td>6Cl2P</td>
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<td>2.28</td>
</tr>
<tr>
<td>6Br2P</td>
<td>2.16</td>
<td>2.29</td>
</tr>
</tbody>
</table>

% α = |(Δr/(r1+r2))| x 100 where r1 & r2 are the newly forming or cleaving bonds in the TSs

Table 2. Thermochemical activation and reaction parameters for the cycloaddition and cycloreversion of 2P, halosubstituted 2P and OXZ with Ace. (Free energy of enthalpy is in kcal mol⁻¹).

<table>
<thead>
<tr>
<th>System</th>
<th>Cycloaddition</th>
<th>Cycloreversion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Activation parameters</td>
<td>Reaction parameters</td>
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<tr>
<td></td>
<td>ΔG°</td>
<td>ΔH°</td>
</tr>
<tr>
<td>2P</td>
<td>36.84</td>
<td>25.35</td>
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<tr>
<td>3F2P</td>
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<td>4Cl2P</td>
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<td>5F2P</td>
<td>34.81</td>
<td>18.68</td>
</tr>
<tr>
<td>5Cl2P</td>
<td>37.28</td>
<td>25.88</td>
</tr>
</tbody>
</table>

The values in the parenthesis are stand for the HCN elimination pathway.
Figure 6. Variation of $\Delta G^\ddagger$ of cycloaddition and cycloreversion of Ace with various halopyrones as dienophiles.

Figure 7. Variation of Magnetic susceptibility along the reaction coordinate of (a) TS1 (b) Pyrone ring of TS1 (c) TS2 (d) Benzene ring of TS2 in 2P-Ace reaction.
B. Aromaticity along the CA-CR pathway

During the first step the 2-pyrone loses its partial aromatic character and forms the cycloadduct. Two electronic phenomena are happening viz., loss of partial aromaticity in
the pyrone ring and gain in the aromaticity in the pericyclic ring along the reaction coordinate. Now the $\chi_{iso}$ of the whole system and of the reacting 2-pyrene fragment as a function of reaction coordinate have been monitored and presented in the Figure 6 (a) and (b) respectively. The nature of change in $\chi_{iso}$ along IRC, as shown in Figure 7 (a), reveals that the aromatic gain in the pericyclic ring is dominant and not have been upset by any loss of aromaticity in the reactant as given in Figure 7 (b). A completely opposite phenomenon has been observed in the second step. The reactant system shows continuous increase m negative $\chi_{iso}$ and this trend parallels the trend shown by the product fragment along the IRC and as shown in Figure 7 (c) and (d). A sweeping change in aromatic game by the formation of benzene ring as the product offsets the changes in $\chi_{iso}$ in the pencyclic ring. This case is similar to that reported by Manoharan et al., [30] wherein he demonstrates that the usual 6e cyclic ($\sigma$, $\pi$) delocalization of the pericyclic TS will be accompanied by the partial delocalization of the peripheral ring of o-quinodimethane (diene). On account of this aromatic gain, the cycloaddition between o-quinodimethane and ethylene occurs with relatively low barrier than the prototype butadiene and ethylene addition. In the present case cycloreversion is accelerated on account of aromaticity developed in the newly forming ring.

C. OXZ-Ace Reaction

OXZ-Ace reaction resembles 2P-Ace reaction in that it forms a bicyclic lactone first and then the lactone decomposes to give the final product. The above reactions differ in that the latter has two paths to cyclorevert; (i) CO$_2$ elimination (ii) HCN elimination. The trend in aromaticity changes as observed through changes in $\chi_{iso}$ are presented in Figures 7 and 8. They are quite similar for both the reactions as far as cycloaddition and CO$_2$ elimination steps are concerned. Two changes can be noted in OXZ-Ace reaction, (i) TS3 is less aromatic than TSI as reflected by lower $\chi_{iso}$ values and this is due to the presence of nitrogen atom in the ring. (ii) The change in aromaticity in HCN elimination path, during cycloreversion is dominated by the aromaticity of the pericyclic ring in TS5, as the final product formed is 2-pyrene that is only partially aromatic. One can observe this from the Figure 5 (c) and (d). This clearly explains the reluctance of the bicyclic lactone adduct (A2) to choose the nitrile elimination path where there is lesser aromaticity gain than in the alternative CO$_2$ elimination path.

4. Conclusion

Tandem cycloaddition-cycloreversion of 2-pyrene and its aza analogue, 1,4-oxazinone with acetylene follows concerted mechanism with asynchronous transition states. 2P-Ace cycloadduct rapidly cycloreverts due to the aromatic enhancement. In OXZ-Ace adduct, two extrusable fragments (viz., CO$_2$ & HCN) compete and the barriers indicate that CO$_2$ extrusion is facile due to the excellent gain in aromaticity during the process.

References


