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Activation of the $S_N2$ Reaction by Adjacent $\pi$ Systems: The Critical Role of Electrostatic Interactions and of Dissociative Character

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Supporting Information Placeholder

ABSTRACT:The activation of the $S_N2$ reaction by $\pi$ systems is well documented in textbooks. It has been shown previously that this is not primarily due to classical (hyper)conjugative effects. Instead, nonconjugated substituents enhance favorable substrate-nucleophile electrostatic interactions, with electron-withdrawing groups on the sp$^2$ system leading to even stronger activation. Herein we report computational and experimental results which show that this activation by sp$^2$EWG-substitution only occurs in fact in a fairly limited number of cases, when the nucleophile involves strong electrostatic interactions (usually strongly basic negatively charged nucleophiles). In other cases, where bond breaking is more advanced than bond making at the TS, electrophilic-nucleophile-electrostatic interactions are less important. In such cases, (hyper)conjugative electronic effects determine the reactivity, and EWG-substitution leads to decreased reactivity. The basicity of the nucleophile as well as solvent effects can help to determine which of these two regimes occurs for a given electrophile.

It is well documented that allylic and benzylic derivatives react faster than corresponding alkyl derivatives in $S_N2$ reactions, and that the activation is greater with electron deficient $\pi$ systems.1-3 In the classical textbook explanation, this is because the $\pi$-symmetric p orbital at the central $\alpha$-carbon becomes more populated in the TS due to donation from the nucleophile, and delocalization into the $\pi$ system results in stabilization of the TS.3Brauman, Allen and Galabov5 and many others6 have however showed that conjugative and hyperconjugative effects are limited and should not be considered as the main origin of “allylic” and “benzylic” effects in $S_N2$ reaction. According to Allen and Galabov5 the critical effect of the (substituted) $\pi$ system is instead to make substrate-nucleophile electrostatic interactions more favorable in the TS (Figure 1). This effect can be probed by calculating the electrostatic potential at the Cα and Cβ nuclei at the transition state (TS) structure. The additional activating effect of electron-withdrawing substituents14,5,6 is then attributed to an increased positive charge at Cα, and the associated strengthened electrostatic interactions with the approaching nucleophile.

Figure 1. Allen and Galabov5 rationale for activation of $S_N2$ reaction by $\pi$ systems.

Figure 2. Substituent effects in intermolecular $S_N2$ reaction and cyclization to epoxides or aziridines (X = O or NSO:Ph).

We were thus surprised when, in the context of the development of sulfur ylide-mediated epoxidation and aziridination reactions, we observed decreased reactivity in the intramolecular nucleophilic displacement step with conjugated EWGs (e.g. R = CON-Me2, CO2Me or electron-poor aryls), whereas electron-rich aryls were found to activate this step (Figure 2).7,9 This unexpected difference in substituent effects between intramolecular (3-exo-tet) and intermolecular $S_N2$ reactions prompted us to investigate the factors governing substituent effects in the elimination step of the ylide-mediated epoxidation and aziridination reactions. We report herein computational and experimental data which show that the conventional EWG-acceleration effect only occurs when electrostatic interactions play a dominant role. Where such interactions are less important due to the nature of nucleophile or to stereoelectronic factors, (hyper)conjugative
electronic effects can determine the order of reactivity. In such cases, EWGs then deactivate the substrate.

We have used density functional theory (DFT) to compute activation barrier heights and to analyze the factors that affect them. The analysis relies in part, as in previous work, on the calculated electrostatic potential $V_{TS}$ at the position of $C\alpha$ and $C\beta$ nuclei within the system, at the structure of the TS. We also use a calculated property of the nucleophile that relates to the extent it will stabilize the TS through electrostatics, which we call the “unrelaxed proton affinity”, $PA_X$. This is calculated as the difference in energy between the bare nucleophile, and the nucleophile to which a proton has been added at the position of $C\alpha$ and $C\beta$ in the TS structure. This is in fact very similar to the electrostatic potential created by the electron density of the nucleophile at the position of $C\alpha$ and $C\beta$ ($V_X$, reported in SI) though $PA_X$ includes electronic relaxation effects. The $V_{TS}$ electrostatic potential property can be calculated on the whole system, including the nucleophile, and so can be treated identically for inter- and intramolecular cases. However, $PA_X$ needs to be calculated for the nucleophile only. For the intramolecular case, we therefore used a truncated system, based on the -CH$_2$X nucleophilic part, which was capped with a hydrogen atom; the structure of the resulting CH$_3$ group was optimized while holding the rest of the system at the TS structure (see SI for details).

We considered first the set of reactions in Table 1, with an initial focus on the typically used DMSO solvent. The trend in DFT free energy barriers for elimination to epoxides (eq. 2) matched the experimental trends. Para substitution of the aromatic ring by an EDG led to a decrease in barrier, while EWGs increased it, as shown in the penultimate column of Table 1. In contrast, for the analogous intermolecular reactions with methoxide (eq. 1), DFT predicted that EWGs lowered the barrier, in agreement with the conventional observation of acceleration by these substituents (see second column in Table 1).

Table 1. Substituent effects in intermolecular and intramolecular (3-exo-tet) $S_N2$ reactions.*

<table>
<thead>
<tr>
<th>$X$</th>
<th>$PA_X(C\alpha)$</th>
<th>$PA_X(C\beta)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>-405.973</td>
<td>-221.9</td>
</tr>
<tr>
<td>O</td>
<td>-404.661</td>
<td>-217.9</td>
</tr>
<tr>
<td>$O$</td>
<td>-405.540</td>
<td>-217.9</td>
</tr>
<tr>
<td>$O$</td>
<td>-403.094</td>
<td>-217.9</td>
</tr>
</tbody>
</table>

* Free energy barrier in kcal/mol. $V_{TS}(C\alpha)$ and $V_{TS}(C\beta)$ are the electrostatic potential (Volt) at, respectively, $C\alpha$ and $C\beta$ in the $S_N2$ TS. $PA_X(C\alpha)$ and $PA_X(C\beta)$ are the unrelaxed proton affinity (kcal/mol) of $X$ at the position of $C\alpha$ and $C\beta$ in the $S_N2$ TS, respectively. $\Delta Q_{CHR} = \text{NBO charge of } C\alpha \text{H}x \text{R at } TS - \text{NBO charge of } C\alpha \text{H}x \text{R in reactant.}$

**Table 2.** Electronic and structural parameters for intermolecular and intramolecular (3-exo-trig) $S_N2$ reactions ($R = Ph$)

<table>
<thead>
<tr>
<th>$X$</th>
<th>$\Delta G$</th>
<th>$V_{TS}(C\alpha)$</th>
<th>$V_{TS}(C\beta)$</th>
<th>$PA_X(C\alpha)$</th>
<th>$PA_X(C\beta)$</th>
<th>$\Delta Q_{CHR}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>26.9</td>
<td>-405.973</td>
<td>-221.9</td>
<td>256.2</td>
<td>205.9</td>
<td>0.075</td>
</tr>
<tr>
<td>$O$</td>
<td>27.0</td>
<td>-404.661</td>
<td>-217.9</td>
<td>251.2</td>
<td>206.0</td>
<td>0.206</td>
</tr>
<tr>
<td>$O$</td>
<td>7.8</td>
<td>-405.540</td>
<td>-217.9</td>
<td>231.1</td>
<td>251.2</td>
<td>0.252</td>
</tr>
<tr>
<td>$O$</td>
<td>7.8</td>
<td>-403.094</td>
<td>-217.9</td>
<td>197.9</td>
<td>251.2</td>
<td>0.315</td>
</tr>
</tbody>
</table>

Figure 3. Transition state structure for inter- (left) and intra- (right) molecular $S_N2$ reactions ($X = O$; $R = Ph$) and electrostatic potential surface of methoxide anion (o and x show the relative positioning of $C\alpha$ in the $S_N2$ transition state for inter- and intramolecular case, respectively).

With the sulfonamide nucleophile ($X = \text{NSO}_3\text{Me}$), EWG substituents are predicted to decrease reactivity in both the inter- and
intra-molecular cases (see Table 1). This could be explained by lesser electrostatic stabilization of the TS by the nucleophile (with its delocalized charge), associated with a more dissociative mechanism, as shown by the PAx metric in Table 2. The subsequent greater increase in positive charge ΔQCHR for the alkyl group part of the substrate at the TS means that substituents capable of stabilizing positive charge (conjugated EDG substituents) lead to relative stabilization of TSs and hence an activation of the reaction, as in the intramolecular epoxide formation.

We have extended this study to a much broader set of nucleophiles (Table 3) and found that, contrary to the received textbook wisdom, EWG deceleration of Sn2 reactions is in fact quite common. Here, only the intrinsic Sn2 barrier is low enough for the reaction to occur or decelerate reaction purely based on the substituent (conjugated EWG or EDG) substituent. We have shown that a reaction's rate constant is proportional to the free energy of activation ΔG‡. For R = Ph

<table>
<thead>
<tr>
<th>Anionic S2: X + RCH2X → RCH2X + X</th>
<th>X-</th>
<th>MeOPh</th>
<th>Ph</th>
<th>p-NO2Ph</th>
<th>PAx/C</th>
<th>ΔQCHR</th>
</tr>
</thead>
<tbody>
<tr>
<td>NMe2</td>
<td>53.2</td>
<td>52.3</td>
<td>48.6</td>
<td>296.5</td>
<td>-0.111</td>
<td></td>
</tr>
<tr>
<td>OMe</td>
<td>57.6</td>
<td>47.2</td>
<td>44.8</td>
<td>273.0</td>
<td>-0.057</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>33.5</td>
<td>33.0</td>
<td>31.1</td>
<td>251.5</td>
<td>-0.025</td>
<td></td>
</tr>
<tr>
<td>PMe2</td>
<td>55.4</td>
<td>50.8</td>
<td>42.5</td>
<td>311.0</td>
<td>0.021</td>
<td></td>
</tr>
<tr>
<td>SMe</td>
<td>38.4</td>
<td>37.9</td>
<td>37.5</td>
<td>275.9</td>
<td>0.120</td>
<td></td>
</tr>
<tr>
<td>Cl</td>
<td>19.1</td>
<td>21.6</td>
<td>23.3</td>
<td>233.6</td>
<td>0.184</td>
<td></td>
</tr>
<tr>
<td>Br</td>
<td>30.5</td>
<td>33.0</td>
<td>33.2</td>
<td>236.1</td>
<td>0.169</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>14.2</td>
<td>14.3</td>
<td>17.1</td>
<td>242.7</td>
<td>0.234</td>
<td></td>
</tr>
<tr>
<td>N(Me)SO2</td>
<td>47.8</td>
<td>48.2</td>
<td>48.6</td>
<td>229.6</td>
<td>0.125</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Neutral S2: X + RCH2X → RCH2X+ + X</th>
<th>X-</th>
<th>MeOPh</th>
<th>Ph</th>
<th>p-NO2Ph</th>
<th>PAx/C</th>
<th>ΔQCHR</th>
</tr>
</thead>
<tbody>
<tr>
<td>NMe3</td>
<td>31.0</td>
<td>31.4</td>
<td>31.8</td>
<td>126.8</td>
<td>0.058</td>
<td></td>
</tr>
<tr>
<td>OMe3</td>
<td>16.8</td>
<td>18.5</td>
<td>21.2</td>
<td>78.4</td>
<td>0.224</td>
<td></td>
</tr>
<tr>
<td>PMe3</td>
<td>52.2</td>
<td>52.2</td>
<td>53.5</td>
<td>146.4</td>
<td>0.246</td>
<td></td>
</tr>
<tr>
<td>SMe2</td>
<td>25.4</td>
<td>26.3</td>
<td>26.9</td>
<td>112.8</td>
<td>0.290</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Influence of substitution on the free energy barrier in identity Sn2 reaction of benzyl derivatives.

For R = Ph

We have shown that the importance of electrostatic interactions in the TS for a given nucleophile depends strongly on PAx. For intermolecular cases, this property is quite well correlated to the basicity of the nucleophile. Accordingly, it is possible to predict whether a conjugated EWG or EDG substituent at Cα will accelerate or decelerate reaction purely based on the pKa of the nucleophile’s conjugate acid. Strongly basic nucleophiles such as dialkylamides, thioalkoxides or fluoride lead to EWG-accelerated Sn2 reactions, whereas for weakly basic nucleophiles such as chloride, bromide, iodide, or neutral nucleophiles, there is an EWG-decelerated trend.

It is important to note that solvation effects are expected to influence the magnitude of stabilization by electrostatic interactions as well as the associative/dissociative character of the TS. The limit between EWG-decelerated and accelerated nucleophiles may thus well vary with the nature of the solvent. Chloride nucleophile, for instance, was computed to follow the EWG-decelerated trend in DMSO (see Table 3) whereas, in the gas phase, the opposite trend was predicted (see SI).

Our observations are important for a series of reactions, one example being formation of onium salts. Contrary to expectation based on the classic model, we have found that alkylation of tertiary amines or sulfides with electron-poor benzylic derivatives are slower with non- or EDG-substituted analogues as revealed by competition experiments (Table 4). Therefore, the results fit with the new model proposed. In the case of the anionic nucleophile, MeSNa, acceleration due to the EWG was predicted and observed.

<table>
<thead>
<tr>
<th>Nucl.</th>
<th>Conv. (%)</th>
<th>A/B</th>
<th>Exp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>OMe</td>
<td>28</td>
<td>84/16</td>
<td>69/31</td>
</tr>
<tr>
<td>MeSNa</td>
<td>21</td>
<td>&gt;99/1</td>
<td>70/30</td>
</tr>
</tbody>
</table>

Table 4. Competition experiments

If we consider the alternative reaction pathways, we can see that the more reactive chlorides are less prone to deceleration due to the presence of the EWG. This effect is consistent with the competition experiments, where the nucleophile that is more reactive to the chloride is also more reactive to the onium salt.

In summary, we have shown that the conventional received wisdom whereby EWGs at Cα accelerate Sn2 reactions is actually only applicable in a limited number of circumstances: those that involve highly basic nucleophiles which can lead to strong electrostatic stabilization of the TS. In such cases, incipient bond-making effects dominate over bond-breaking ones. In the case of neutral or delocalized anionic nucleophiles or where strained rings are created, electrophile-nucleophile electrostatic interactions are less important and bond-breaking is more advanced at the TS than bond-making. Hence EWGs decelerate Sn2 reactions, in line with the expected (hyper)conjugative electronic effects at what is a partially positively charged carbon center in the TS.

ASSOCIATED CONTENT

Supporting Information
Details of computational methods. Cartesian coordinates and energies for all structures. Additional data on onium salt formation reactions as well as on identity reactions. Relative energy of Sx1 intermediates. This material is available free of charge via the Internet at http://pubs.acs.org.

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Corresponding Author

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A
Notes
The authors declare no competing financial interests.

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REFERENCES


(8) As an example, computed energy barrier for cyclization to epoxide is ~4 and 6–9 kcal/mol for R = Ph and CONMe2, respectively. See ref 7a.


(10) Computations have been carried out at the B3LYP-D3/aug-cc-pVQZ//B3LYP/6-31G(d,p) level of theory, including a continuum description of DMSO solvent for both the geometry optimization and the single point energy calculation using the Jaguar 6.5 and 8.5 program package (Jaguar, version 6.5, Schrodinger, LLC, New York, NY, 2005 and Jaguar, version 8.5, Schrodinger, Inc., New York, NY, 2014). See SI for full computational details and data.

(11) The increase in conjugative effects with electron-donating character of the sp”substriuent can be observed by the variation of the C=C–C bond length according to substitution (see SI).

(12) It was shown that substituent effects in benzylc S22 reactions are the result of variations in the intrinsic barrier. See, for instance, ref. 1a, 4a and 5. This is also supported by our calculations showing the total absence of correlation between the trend of substituent effects on the free energy barrier and the reaction free energy (see SI).


(15) The EWG accelerated trend for chloride nucleophile in the gas phase was also predicted by Streitwieser (see ref. 1a) and Uggerud (see ref. 6b).
For Table of Contents Only

\[ \text{Nu} \xrightarrow{R} \text{LG} \quad \text{Nu} \xrightarrow{R} \text{LG} \quad \text{Nu} \xrightarrow{R} \text{LG} \]

- Strongly basic nucleophiles
  - \( R = \text{EDG} \)
  - Slower
  - Faster

- Weakly basic nucleophiles
  - Faster
  - Slower

- All nucleophiles
  - Faster
  - Slower

Believed general trend so far: Dominant trend