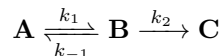


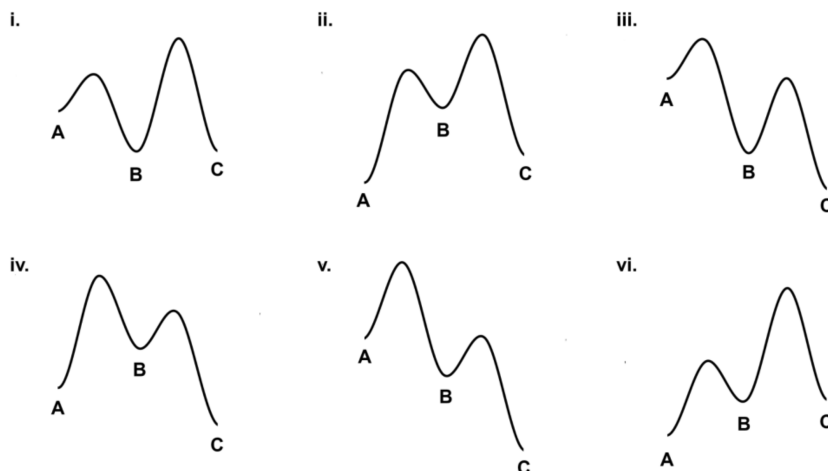
4 Isotope Effects and Kinetics

11/26: The questions pertain to the material covered from Thermodynamic Isotope Effects (Nov 5) to Kinetic Rate Laws (Nov 14).

1. The following schematic illustrates the six possible kinetic scenarios for the reaction



of **A** to **C** via **B**.



- a) In which of these scenarios would the steady-state approximation *not* be valid? Provide a brief explanation for your answers.

Answer. The steady-state approximation would not be valid in **Scenarios i, iii, and vi.**

i: The initial equilibrium between **A** and **B** is exergonic, and there is a significant barrier before conversion to **C**. Symbolically, $k_1 \gg k_{-1} \gg k_2$. Thus, most of the substrate will exist as **B**. Since $[B] \gg [A]$, the steady-state approximation does not hold.

iii: Again, it is easier for **A** to become **B** than for **B** to become **C**, so most of the substrate will sit as the intermediate.

vi: There will be more **A** than **B**, but **B** will accumulate before it converts to **C** □

- b) In which of these scenarios would the quasi-equilibrium assumption *not* be valid? Provide a brief explanation for your answers.

Answer. The quasi-equilibrium assumption would not be valid in **Scenarios iv and v.** In both of these, $k_2 \gg k_1$, so it will be hard for **A** and **B** to effectively equilibrate before some **B** starts getting converted to product. □

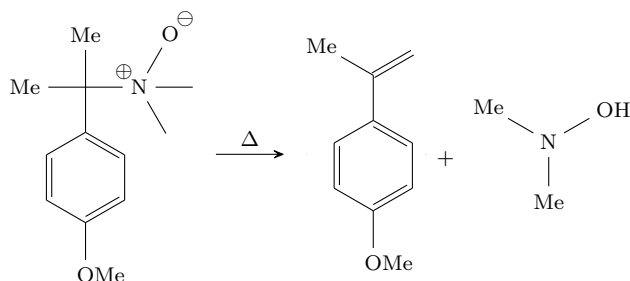
- c) Consider a situation where **A** is isotopically labeled. If isotopic substitution affects only the rate of conversion of **B** to **C** (and not from **A** to **B**), in which scenarios might you expect to observe an independent rate kinetic isotope effect (KIE)? Please explain.

Answer. Under these assumptions, a KIE would be observed when the second step is rate-determining. Recall that the rate-determining step is the step that has the largest energy difference relative either to the starting material or to any previous, lower energy intermediate on the diagram. Thus, we might expect to observe a KIE in **Scenarios i, ii, iii, and vi.** □

- d) In which scenarios might you expect to observe an independent rate KIE if isotopic substitution affects only the step involving formation of **B** from **A** (and not from **B** to **C**)?

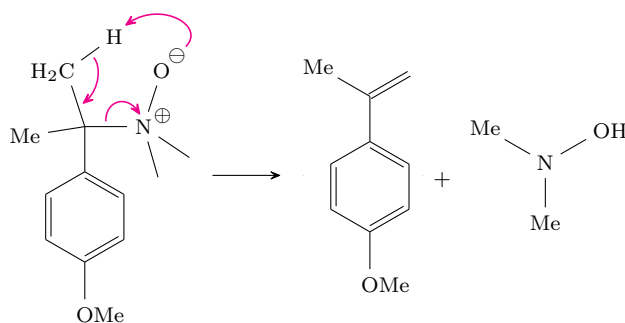
Answer. When the first step is rate-determining, we'll definitely see a KIE. But even when it is not, KIEs will affect $[B]$, altering the rate of the RDS! Thus, **all scenarios** display KIEs. □

2. Consider the following reaction.

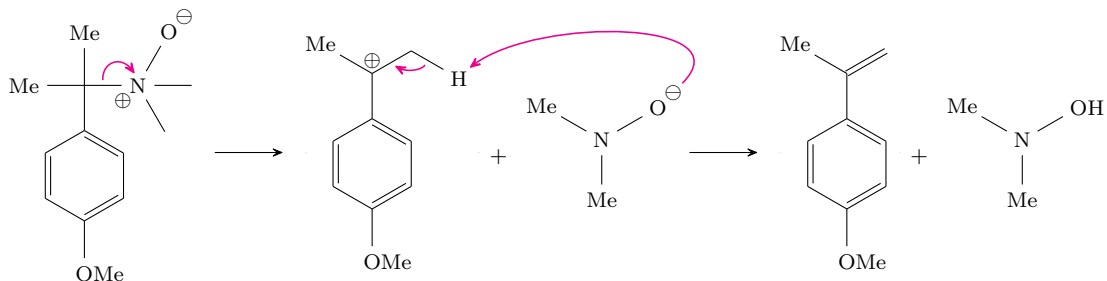


- a) The mechanism for this reaction could proceed via a concerted pathway or a stepwise pathway. Provide arrow-pushing mechanisms for both processes.

Answer. Concerted: This is an E_i mechanism!



Stepwise: This is an E_1 -type mechanism!



□

- b) When the two benzylic methyl groups were deuterated, kinetic isotope effects (KIEs) of 5 and 1.2 were measured in toluene and DMF respectively. Suggest a possible explanation for the observed KIEs.

Answer. A large, likely primary KIE of 5 suggests that C-H/D bond cleavage is part of the RDS. Since the RDS of the concerted mechanism (the only step) involves C-H/D bond cleavage and the RDS of the stepwise mechanism (likely the first step, as is typical of E_1 -type mechanisms) does not involve C-H/D bond cleavage, the concerted mechanism is likely active in toluene.

A small, likely secondary KIE of 1.2 suggests that C-H/D bond cleavage is *not* part of the RDS. Thus, for analogous reasons to the above, the stepwise E_1 -type mechanism is likely active in DMF.

□

- c) When the reaction was run at two different temperatures and in two different solvents, the following rate data were obtained. Compare the activation enthalpy (ΔH^\ddagger) and activation entropy (ΔS^\ddagger) of the reactions in toluene and in DMF. Suggest a possible explanation for the observed differences.

T (°C)	k_{rel} in toluene	k_{rel} in DMF
60	1	10
110	50	500,000

Answer. Although it is not possible to obtain absolute values (e.g., in kcal mol^{-1}) for ΔH^\ddagger and ΔS^\ddagger from this data, a modified Eyring plot can compare the magnitudes of these quantities between the two reactions. For toluene, the line through

$$\left(\frac{1}{273 + 60}, \ln(1)\right) \quad \left(\frac{1}{273 + 110}, \ln(50)\right)$$

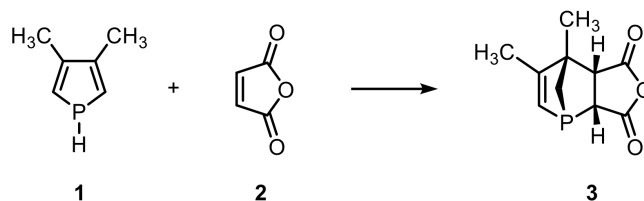
has slope $m = -9979$ and y -intercept $b = 29.97$, so $\Delta H_{rel}^\ddagger = 19.74$ and $\Delta S_{rel}^\ddagger = 0.0593$. For DMF, the line through

$$\left(\frac{1}{273 + 60}, \ln(10)\right) \quad \left(\frac{1}{273 + 110}, \ln(500\,000)\right)$$

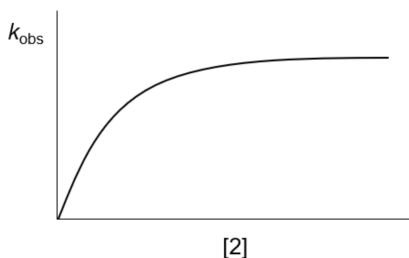
has slope $m = -27600$ and y -intercept $b = 85.18$, so $\Delta H_{rel}^\ddagger = 54.6$ and $\Delta S_{rel}^\ddagger = 0.169$.

It follows that the reaction in DMF — which proceeds via the stepwise mechanism, per part (b) — has both a higher enthalpy and entropy of activation. It makes sense that it should have a higher enthalpy of activation since the RDS involves the formation of a high-energy carbocation intermediate. It also makes sense that it should have a higher entropy of activation because the introduction of charged species into a polar aprotic solvent will disrupt the random bonding of the solvent to itself and introduce highly ordered ion cages. \square

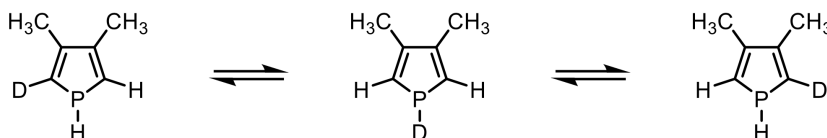
3. In nonpolar solvents, compound **1** reacts thermally with **2** to give only *endo* **3**.



Using small amounts of **1** and 'flooding' with **2**, it has been established that the rate is pseudo-first-order in [**1**]. Measuring the pseudo-first-order rate constant k_{obs} at several different concentrations of excess **2** gives a plot of the type shown below.

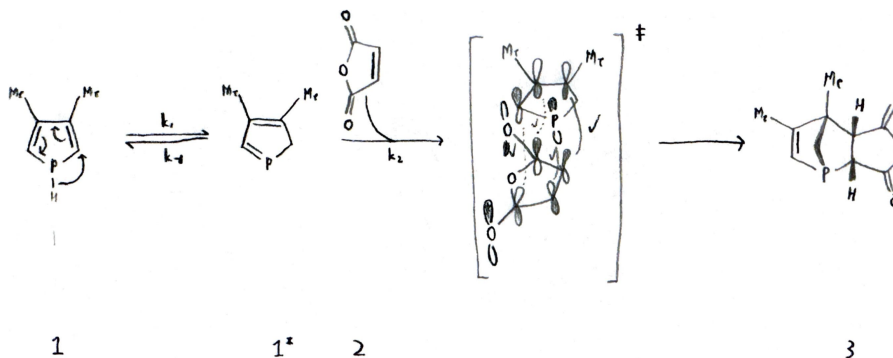


Compound **1**, deuterated at phosphorus, undergoes exchange of D with both α -hydrogens, as shown below. At low concentrations of **2**, the H/D exchange rate is rapid compared to the rate of formation of **3**. At 'saturating' concentrations of **2**, the formation of **3** occurs faster than H/D exchange.



- a) Write a full mechanism and a rate law for the formation of **3** that is consistent with these observations.

Answer. A full mechanism consistent with these observations is



A rate law consistent with these observations is

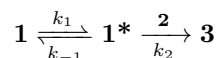
$$\text{rate} = \frac{k_1 k_2 [\mathbf{1}] [\mathbf{2}]}{k_{-1} + k_2 [\mathbf{2}]}$$

□

- b) Explain how your mechanism accounts for the labeling and kinetic behavior.

Answer. We begin by developing a plausible arrow-pushing mechanism for the overall indicated transformation. Nonpolar solvents, thermal reactivity, a starting material (**2**) with two strong EWGs, and *endo*-selectivity suggests that **3** was ultimately formed by a $[4 + 2]$ cycloaddition. Taking the six-membered ring in **3** to be a full retron for a $[4 + 2]$ cycloaddition, we can predict **1*** and **2** as starting materials. Indeed, **1*** and **2** would react in the forward direction via the indicated transition state, with bonds being formed between the HOMO of **1*** and the full LUMO of **2**; *endo*-selectivity is secured by the two secondary orbital interactions indicated with dashed lines. This just leaves the question of where **1*** could have come from. One possible option is from **1** via a suprafacial $[1, 5]$ -sigmatropic H-atom shift. Such pathways are very common for such doubly unsaturated systems, so this is a reasonable hypothesis.

Let's now see why this plausible mechanism is defensible based on the given data, and in the process derive the above rate law. To begin, a simplified chemical equation is



Based on this equation, we would predict that

$$\text{rate} = \frac{d[\mathbf{3}]}{dt} = k_2[\mathbf{1^*}][\mathbf{2}]$$

To (analytically) express $[\mathbf{1^*}]$ in terms of $[\mathbf{1}]$, we may choose between four possible simplifying assumptions:

- The steady-state approximation.
- The steady-state approximation, accounting for $[\mathbf{1}]_T$.
- The quasi-equilibrium assumption.
- The quasi-equilibrium assumption, accounting for $[\mathbf{1}]_T$.

Using these assumptions, we may derive the following four rate laws, respectively.

$$\text{rate} = \frac{k_1 k_2 [\mathbf{1}][\mathbf{2}]}{k_{-1} + k_2 [\mathbf{2}]} \quad \text{rate} = \frac{k_1 k_2 [\mathbf{1}]_T [\mathbf{2}]}{k_1 + k_{-1} + k_2 [\mathbf{2}]} \quad \text{rate} = \frac{k_1 k_2}{k_{-1}} [\mathbf{1}][\mathbf{2}] \quad \text{rate} = \frac{k_1 k_2}{k_1 + k_{-1}} [\mathbf{1}][\mathbf{2}]$$

Both equations derived under the quasi-equilibrium assumption fail to account for the saturation kinetics on display in the k_{obs} vs. $[\mathbf{2}]$ plot, so we can rule them out. Thus, this reaction must occur in a regime subject to the steady-state approximation. It follows that $k_{-1} \gg k_1$, so we may additionally neglect the k_1 term in the denominator of the fourth option, leaving the indicated rate law as our choice.

Additionally, observe that in the limiting case of 'flooding' with **2**, $k_2[\mathbf{2}] \gg k_{-1}$. This allows the rate law to collapse to

$$\text{rate} = k_1[\mathbf{1}] = k_{\text{obs}}[\mathbf{1}]$$

corresponding to the observation of that the reaction is pseudo-first-order in **1** under these conditions!

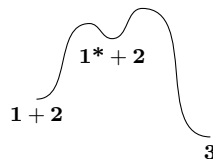
Furthermore, the proposed reversible first step would account for the hydrogen-deuterium exchange at the symmetric α -carbons (even in the absence of **2**): D could migrate from the phosphorus center to either α -carbon, and then either D or H could migrate back.

The final observation regarding the relative rates of H/D exchange vs. the rate of formation of **3** will be useful in part (c). \square

c) Draw reaction coordinate energy diagrams at the limiting cases where **[2]** is...

i) Very low;

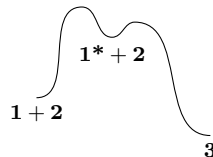
Answer.



Since this reaction obeys the steady-state approximation as discussed in part (b), not much **1*** is ever formed. Thus, **1* + 2** should always be higher in energy than **1 + 2**. Additionally, since the reaction proceeds, it should always be exergonic. Finally, the difference in the heights of the transition states comes from the data that H/D exchange is faster than formation of **3** at low concentrations of **2**; this observation implies a lower barrier for **1* + 2** to return to the reactants than proceed on to the products. \square

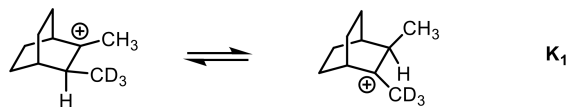
ii) Very high.

Answer.



The analysis is identical to the part (c.i), except that conversion to the products now occurs faster, so its barrier should be lower. \square

4. a) Estimate the magnitudes of the equilibrium constants (K_1 , K_2) for the following reactions. Provide explanations.



Answer. C–D bonds have a slightly greater BDE than C–H bonds. Thus, they hold onto their electrons more tightly and are less likely to participate in stabilizing interactions like hyperconjugation. Essentially, carbocations subject to $\sigma_{CH} \rightarrow p_C$ hyperconjugation will be more stable than those subject to $\sigma_{CD} \rightarrow p_C$ hyperconjugation, all else being equal. Thus, we should have

$$K_1 > 1$$

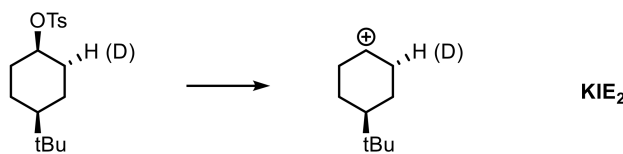
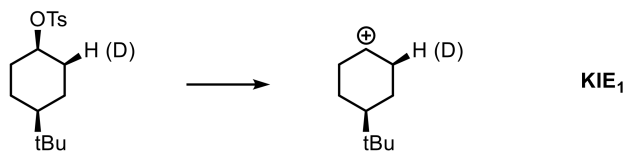
Note that K_1 will likely be only slightly greater than one: When we compared diaxial interactions between methyl groups and perdeuterated methyl groups in class, K_{eq} was 1.042.

For K_2 , the σ_{CD} -orbital is orthogonal to the carbocation's p_C orbital, so it cannot stabilize it anyway. Thus, we should have

$$K_2 \approx 1$$

□

- b) Estimate the magnitudes of KIE_1 and KIE_2 for the following solvolysis reactions. Provide explanations.



Answer. The same effects at play in part (a) are also in effect here, except that the type of hyperconjugation is now anomeric $\sigma_{C-H/D} \rightarrow \sigma_{C-O}^*$ donation. A lack of orbital alignment for the top reaction means that

$$KIE_1 \approx 1$$

Orbital alignment in the bottom case will predict a small normal secondary KIE:

$$KIE_2 \in (1, 1.5)$$

□