Transition-state Acidities and the pH Dependence of Drug Stability*

K. Barbara Schowen and Richard L. Schowen**

Departments of Chemistry and Pharmaceutical Chemistry
Higuchi Biosciences Center, University of Kansas,
Lawrence, Kansas 66045-0046, USA

Received May 7, 1996; revised August 6, 1996; accepted August 10, 1996

The pH dependence of hydrolytic reactions of drugs allows some control of their stability through adjustment of the pH of storage. The transition-state acidity concept of J. L. Kurz is shown to apply to the systematics of relevant pH rate profiles. As limiting cases for the hydrolysis of a series of carboxylic acid derivatives (often employed in pro-drug modifications) in neutral and basic solution, two situations are considered: 1) larger structural effects in basic solution, in which case the most stable compound of the series has the highest transition-state $pK_a$, and 2) larger structural effects in neutral solution, in which case the most stable compound will have the lowest transition-state $pK_a$. The former is the expected situation for variations of reactant electronic features because the negatively charged transition state for the hydroxide-promoted reaction in basic solution should respond more sensitively to electronic effects than should the dipolar transition state for the uncatalyzed reaction in neutral solution. Available data for some important substrates in fact do not show the expected behavior, which may be indicative of a concerted reaction (no tetrahedral intermediate) for reactive substrates with hydroxide ion, a mechanism for which others have already provided evidence, and possible reaction through an ion pair for the reaction of reactive substrates with water.

* This paper is dedicated to the memory of Professor Stanko Borčić, 1931–1994.
** Author to whom correspondence should be addressed.
INTRODUCTION

Professor Stanko Borčić contributed greatly to our knowledge of the transition states of complex organic reactions in solution, and his publications will continue to serve not only for their fundamental scientific value but also as paradigms of the rational, precise dissection of difficult and significant questions. As a member of the Faculty of Biochemistry and Pharmacy in Zagreb, Professor Borčić would, we believe, have welcomed the attempt we describe here to make use of fundamental concepts of transition-state chemistry in approaching a practical problem of pharmaceutical science and would without doubt have advanced the solution to a far greater degree than we will have been able to do.

Many pharmaceutical products are stored in aqueous solution and their hydrolytic degradation presents a medical problem. Drug-stability studies commonly include the establishment of a profile of degradative rate data vs. pH since one strategy for minimizing loss of drug is to select a pH for the storage formulation at which the drug is maximally stable, i.e., at which the degradation rate is minimal. Such profiles often exhibit complex features because drugs often exist in various protomeric forms, each of which may be subject to catalytic degradation with acid catalysis, with base catalysis or spontaneously (un catalyzed reaction or water reaction). For drugs without ionizable groups in the common pH range, a frequently encountered pH profile is that shown in Figure 1 (to be discussed below). In spite of the absence of ionizable groups, the profile exhibits breaks similar to those seen in titration curves; this raises the question, what is titrating?

Joseph L. Kurz answered this question in a series of very influential papers (reviewed in Kurz1) which introduced the concept of acid-base chemistry of transition states (most properly, of activated complexes in the transition state). Kurz showed that a reaction system consisting of parallel processes for acid catalysis, uncatalyzed reaction, and base catalysis (Eqs. 1, where SH is the substrate and TH₂⁺, TH and T⁻ are transition states for the acid-catalyzed reaction [rate constant kₜ], the uncatalyzed reaction [rate constant kₜ], and the base-catalyzed reaction [rate constant kₜ]) would change from the acid-catalyzed pathway to an uncatalyzed pathway at a pH equal to the pKₐ of the transition state for acid catalysis, pKₐ, and would change from the uncatalyzed pathway to the base-catalyzed pathway at a pH equal to the pKₐ of the transition state for the neutral reaction, pKₐ₂. (Eqs. 2, where Kₜ is the ion product of water):

\[
\begin{align*}
SH + H^+ &\rightleftharpoons TH_2^+ \quad K_{eq} = k_h/(kT/h) \\
SH &\rightleftharpoons TH \quad K_{eq} = k_u/(kT/h)
\end{align*}
\]
\[
\begin{align*}
SH + \text{HO}^- & \rightleftharpoons T^- & K_{eq} = \frac{k_b}{(kT/h)} \\
\text{TH}_2^+ & \rightleftharpoons \text{TH} + \text{H}^+ & K_{a1}^* = \frac{k_u}{k_h} \\
\text{TH} & \rightleftharpoons T^- + \text{H}^+ & K_{a2}^* = \frac{K_w k_b}{k_u}
\end{align*}
\]

This is illustrated in Figure 1 for a hypothetical example. The \textit{«breaks»} in the profile correspond to the acid dissociation constants of transition states. The transition state for the acid-catalyzed pathway \textit{«titrates»} to the transition state for the uncatalyzed pathway with a pK_a of 2; the transition state for the uncatalyzed pathway \textit{«titrates»} to the transition state for the base-catalyzed pathway with a pK_a of 11.

![Figure 1. Schematic example of a hydrolytic pH-rate profile for a reaction exhibiting an acid-catalyzed pathway, an uncatalyzed pathway, and a base catalyzed pathway, each with the indicated rate constant.](image)

Valuable information could be provided for drug-design purposes if a general relationship could be established between the pH-inflation points of drug-stability pH-rate profiles and structural features of drugs. As the arguments above show, such a relationship would be equivalent to a relationship between the magnitudes of transition-state acidity constants and the structures of transition states. Since the relationship between the acidities
of ordinary stable molecules and their structures is one of the best-established of chemical concepts for complex systems in solution, this could be a promising approach to the practical pharmaceutical problem.

Our purpose in this paper is to illustrate the concept by examining the results available in the literature for the transition from uncatalyzed to base-catalyzed pathways for the hydrolysis of three carboxylic-acid derivatives. This choice can be considered moderately realistic because such structures occur frequently in drugs, and are of particular importance in prodrugs, species that have undergone potentially reversible modification to improve their physical and chemical properties for pharmaceutical use.

RESULTS AND DISCUSSION

Structure and Reactivity for the Transition from Uncatalyzed to Base-Catalyzed Reaction

Consider a series of hydrolyzable carboxylic acid derivatives such as might constitute a collection of candidate molecules for drug use. If the ratio of hydrolytic reactivities along the series is the same for the uncatalyzed reaction at neutral pH as it is for the base-catalyzed reaction at basic pH, i.e., if $k_b/k_u$ is constant for all compounds in the series, then, as Eq. 2b shows, the transition-state pH will be the same for all compounds.

On the other hand, there is no fundamental reason to expect the reactivity patterns for the two pathways to be identical. If they are not, two lim-

![Figure 2. Schematic pH-rate profiles for a series of three compounds, each exhibiting uncatalyzed and base-catalyzed pathways of hydrolysis: two limiting cases.](image-url)
iting cases arise, illustrated in Figure 2. The «break» in the profile for each compound occurs at a pH equal to the acid dissociation constant of the uncatalyzed transition state to generate the base-catalyzed transition state. Figure 2a (left) shows a case where there is little effect of substrate structure on the rate constant for the uncatalyzed reaction and a larger effect on the rate constant for the base-catalyzed reaction. In this case the least stable (most reactive) compound will have the most acidic transition state. Figure 2b (right) shows a case where there is little effect of substrate structure on the rate constant for the base-catalyzed reaction and a larger effect on the rate constant for the uncatalyzed reaction. In this case, the most stable (least reactive) compound will have the most acidic transition state. The figures also make clear that if the base-catalyzed reaction has a larger structural effect (of whatever origin), then the most stable (least reactive) compound will have the least acidic transition state ($pK_{a2} = 7$ in Figure 2a) and if the uncatalyzed reaction has the larger structural effect, then the opposite is true: the most stable compound will have the most acidic transition state ($pK_{a2} = 5$ in Figure 2b).

For derivatives of the same carboxylic acid in which electronic but not steric features of the leaving group are varied, there is reason to believe that the structural effects will be greater on the base-catalyzed reaction. As Figure 3 shows, the charge development in the transition state, which should give rise to the electronic structural effect, is dipolar in the uncatalyzed transition state but negative in the base-catalyzed transition state. In this figure, the traditional two-step mechanism through a tetrahedral interme-

![Figure 3. Hypothetical transition-state structures for the uncatalyzed hydrolysis (top, $k_u$) and base-catalyzed hydrolysis (bottom, $K_{wkb}$) of carboxylic-acid derivatives with variable structure in the group R.](image.png)
diate is assumed and the addition step is taken to be rate-limiting. For the uncatalyzed reaction, the substituent R «sees» the development of a dipole so that variations in the electronic character of R may be expected to produce canceling energetic effects on the positive and negative poles and thus little overall effect on the rate constant. For the base-catalyzed reaction, the substituent R «sees» the development of net negative charge and this transition state should thus be stabilized by electron-withdrawal at R. The acidity constant $K_a$ should therefore be largest for the substrate with the most electron-withdrawing R group, which should also be the most reactive (least stable) substrate. On this model, the expected behavior of a series of substrates is therefore that shown in Figure 2a.

In either reaction, the elimination step might also be wholly or partially rate-limiting. For the uncatalyzed transition state, this might give rise to positive charge on the leaving group and some changes in the argument, but the charge development should still be dipolar and at least some cancellation of effects should come about. For the base-catalyzed transition state, rate-limiting elimination would place negative charge directly in the leaving group and would intensify the argument given above.

The simplest prediction therefore is that the most electron-withdrawing leaving group, which will produce the least stable reactant, will give rise to the most acidic transition state (Figure 2a). This is also in accord with the elementary expectation that transition-state acidity, like reactant-state acidity, should increase with electron withdrawal.

Results for Three Derivatives

As a preliminary test of this model, we examined the data of Skrabal\textsuperscript{2} for ethyl acetate and of Kirsch and Jencks\textsuperscript{3} for 2,4-dinitrophenyl acetate and acetic anhydride. The data are plotted in Figure 4. Note that the transition state $pK_a$ is 5.1 (most acidic) for the least reactive ethyl acetate\textsuperscript{4}, 7.3 for the more reactive 2,4-dinitrophenyl acetate, and 8.5 (least acidic) for the most reactive acetic anhydride. Thus for this series of compounds, the facts correspond to Figure 2b, in contrast to expectations on the basis of the straightforward (tetrahedral) mechanistic model just presented and shown in Figure 3. As is apparent from Figure 4, these three compounds exhibit a larger dependence of structure on reactivity for the neutral than for the basic reaction (the situation depicted in Figure 2b). Kirsch and Jencks\textsuperscript{3} have shown that the underlying structure-reactivity relationship (larger substituent effect on the uncatalyzed than on the base-catalyzed reaction) is quite general for the structural class of substituted phenyl acetates and noted that it applies to a still wider range of compounds as well. They discussed several explanations, chiefly a difference in rate-limiting step between the two pathways, but it was not possible at that time to offer a conclusive interpretation.
Figure 4. Experimental pH-rate profiles derived from the work of Skrabal (ethyl acetate) and of Kirsch and Jencks (acetic anhydride and 2,4-dinitrophenyl acetate).

Concerted Pathways

One possible route to an explanatory model is the increasing evidence that fast acyl-transfer reactions (ones with strong nucleophiles and good leaving groups) may not pass through a tetrahedral intermediate but instead through a single transition state for both bond-formation to the nucleophile and bond-fission to the leaving group. This is unlikely to be true for either the base-catalyzed or the uncatalyzed reaction of ethyl acetate, but would be expected for the reaction of hydroxide ion with 2,4-dinitrophenyl acetate and with acetic anhydride. In itself, this model would not solve the problem but only make it more difficult: for the base-catalyzed reaction a concerted transition state would tend to have more charge on the leaving group than would the transition state for formation of a tetrahedral intermediate (the likely rate-limiting transition state for a two-step model of the hydroxide-ion reaction). This would make the expected substituent effect for the basic reaction still larger, while the data (Figure 4) show that the effect on the neutral reaction is dominant and produces the observed sequence of transition-state $pK_a$s.

Elimination-Addition Pathway Involving a Transient Ion Pair

However, a model which is consistent with the data of Figure 4 is that shown in Figure 5. This model proposes a concerted pathway for the hydro-
oxide-ion reaction and an elimination-addition mechanism, the key feature of which is a transient ion pair (consisting of an acylum ion and the leaving group), for the reaction with water. If this ion pair is trapped by a nearby solvent water molecule, the resulting transition state for the neutral reaction (upper right in Figure 5) is one for which the leaving group has a full negative charge. Reactions proceeding through such a transition state would be expected to exhibit large variations in rate depending on the electronic structure of the leaving group. Therefore the neutral «uncatalyzed» mechanism is expected to show the larger structural effect just as observed (Figure 4). Such a route is presumed to be too slow for the bimolecular reaction with hydroxide ion, which, it seems reasonable to suppose, proceeds instead through reaction of hydroxide ion directly with the intact, un-ionized reactant. As a reaction involving a strong nucleophile and a good leaving group it is likely, as indicated by several lines of earlier evidence, that the reaction proceeds in a concerted fashion involving both bond formation to the nucleophile and bond fission to the leaving group.

Displacement of the stable acetate or 2,4-dinitrophenyloxide by the reactive hydroxide ion is assumed to pass through a reactant-like transition state with little negative charge on the leaving group. In this case, the effect of structural variation is expected to be much less, again as observed. This unexpected circumstance in which a larger negative charge is present near the substituent R (whose structure is being varied) in the conjugate acid of

Figure 5. Mechanistic model for the two more reactive compounds (acetic anhydride and 2,4-dinitrophenyl acetate) for which data are shown in Figure 4. (The reaction of ethyl acetate is presumed to follow the model of Figure 3.)
the transition state than in the conjugate base, results in the unusual conclusion that increasing electron withdrawal at R leads to a decrease in the acidity of the transition state.

CONCLUSION

The conclusion to be drawn from this presentation is that structure-reactivity relationships as a function of pH are not straightforward and cannot be predicted from simplistic models or generalizations. For practical applications, it will be necessary to obtain experimental data for structural effects and pH-dependence behavior for each particular system and class of compounds.

REFERENCES

4. The data reveal that ethyl acetate shows the incursion of an acid-catalyzed pathway for which the transition state has $pK_a^* = 5.7$, actually less acidic than the transition state for the uncatalyzed pathway.

SAŽETAK

Kiselost prijelaznog stanja stabilnosti lijekova u ovisnosti o pH

K. Barbara Schowen i Richard L. Schowen

Ovisnost hidrolitičkih reakcija lijekova o pH omogućuje kontrolu njihove stabilnosti uz ugodbu pH-vrijednosti uskladištenja. J. L. Kurzov koncept kiselosti prijelaznog stanja primijenjen je kod pH-profila brzina hidrolize. Kao granični slučajevi hidrolize u seriji derivata karboksilnih kiselina (često korištenih u modifikacijama proljekova) u neutralnoj i lužnatoj otopini, razmotrena su dva slučaja: 1) veći struk-
turni efekti u lužnatoj otopini pri čemu najstabilniji spoj u seriji ima najviši $pK_a$ prijelaznog stanja i 2) veći strukturni efekti u neutralnoj otopini pri čemu najstabilniji spoj u seriji ima najniži $pK_a$ prijelaznog stanja. Pri promjenama elektronskih svojstava reaktanata prvi je slučaj očekivan, jer će negativno nabijeno prijelazno stanje u reakciji kataliziranoj hidroksidnim ionom biti više podložno elektronskim efektima nego dipolarno prijelazno stanje za "nekataliziranu" reakciju u neutralnoj otopini. Dostupni podaci za neke važne supstrate ne odgovaraju očekivanom ponašanju, već možda upućuju na uskladenu reakciju (bez tetraedarskog intermedijara) za reaktivne supstrate s hidroksidnim ionom (mehanizam koji su drugi već dokazali) i moguću reakciju preko ionskog para za reaktivne supstrate s vodom.