See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/6212027

## Kinetic Solvent Effects on the Reaction of an Aromatic Ketone π,π\* Triplet with Phenol. Rate-Retarding and Rate-Accelerating Effects of Hydrogen-Bond Acceptor Solvents

ARTICLE in JOURNAL OF THE AMERICAN CHEMICAL SOCIETY · SEPTEMBER 2007

Impact Factor: 12.11 · DOI: 10.1021/ja071716y · Source: PubMed

CITATIONS READS
16 52

4 AUTHORS, INCLUDING:



Raquel E Galian
University of Valencia

**46** PUBLICATIONS **618** CITATIONS

SEE PROFILE



Julia Pérez-Prieto

University of Valencia

168 PUBLICATIONS 1,697 CITATIONS

SEE PROFILE



Grzegorz Litwinienko

University of Warsaw

**51** PUBLICATIONS **1,655** CITATIONS

SEE PROFILE



Published on Web 07/11/2007

## Kinetic Solvent Effects on the Reaction of an Aromatic Ketone $\pi,\pi^*$ Triplet with Phenol. Rate-Retarding and Rate-Accelerating Effects of Hydrogen-Bond **Acceptor Solvents**

Raquel E. Galian,\*,† Grzegorz Litwinienko,‡ Julia Pérez-Prieto,† and Keith U. Ingold§ Instituto de Ciencia Molecular, Universidad de Valencia, Polígono de la Coma s/n, 46980 Paterna, Spain, Department of Chemistry, Warsaw University, Pasteura 1, 02-093 Warsaw, Poland, and National Research Council, Ottawa, Ontario, Canada K1A 0R6

Received March 12, 2007; E-mail: raquel.galian@uv.es

In a seminal study, Scaiano and co-workers1 reported that the quenching by phenols of two ketone triplets,  $n,\pi^*$  benzophenone and  $\pi,\pi^*$  4-methoxypropiophenone, gave phenoxyl and ketyl radicals efficiently (eqs 1 and 2). For phenol,  $k_2/10^8$  M<sup>-1</sup> s<sup>-1</sup>

$$ArCOR \xrightarrow{h\nu} \stackrel{ISC}{\longrightarrow} {}^{3}[ArCOR]^{*}$$
 (1)

$$PhOH + {}^{3}[ArCOR]^{*} \rightarrow PhO^{\bullet} + ArC(OH)R$$
 (2)

values for these  $n,\pi^*$  and  $\pi,\pi^*$  ketones were, respectively, 13 and 49 in benzene and 0.8 and 1.1 in wet acetonitrile. The lower rates in wet acetonitrile were attributed to formation of a PhOH···solvent hydrogen bond (HB). Carbonyls with lowest  $\pi, \pi^*$  triplets were known to abstract H-atoms from hydrocarbons much more slowly than carbonyls having similar excitation energies with lowest  $n_*\pi^*$ triplets, and Wagner et al.<sup>2</sup> had postulated that the  $\pi,\pi^*$  triplets abstracted hydrogen predominantly via their thermally populated, higher energy  $n\pi^*$  states. Later, Leigh et al.<sup>3</sup> proposed that reactions of phenols with  $\pi,\pi^*$  carbonyl triplets involved an intermediate HB exciplex that gave phenoxyl and ketyl radicals by Electron Transfer Proton Transfer (ETPT) (eq 3).4

$$PhOH + {}^{3}[ArCOR]^{*} \rightleftharpoons {}^{3}[Ar(R)C = O \cdots HOPh]^{*} \rightarrow$$

$${}^{3}[Ar(R)\dot{C} - \bar{O} \cdots HOPh] \rightarrow Ar\dot{C}(OH)R + {}^{\bullet}OPh \qquad (3)$$

Kinetics of the bimolecular reactions of triplet ketones with phenols have usually been measured in a single solvent, there being only a few reports of measurements in two<sup>1,5</sup> and three<sup>6</sup> solvents. This is unfortunate because a systematic study of kinetic solvent effects (KSEs) for any H-atom abstraction,7 eq 4

$$XH + Y^{\bullet} \rightarrow X^{\bullet} + YH$$
 (4)

can provide important insights into the reaction mechanism(s).8 Observed KSEs can be quantitatively accounted for by assuming that *only* the (often small) equilibrium fraction (eq 5)

$$XH + S \xrightarrow{K_{XH/S}^S} XH \cdots S \tag{5}$$

of XH molecules that are not making a HB to a HB acceptor (HBA) solvent molecule, S, can react with Y • (eq 4), with a rate constant  $k^0_{XH/Y^*}$  that is equal to the experimental rate constant in a non-HBA solvent, such as with an alkane. The experimental rate constant for H-abstraction<sup>7</sup> in S,  $k^{S}_{XH/Y}$ , is given by eq I,<sup>8,9</sup>

$$k_{XH/Y_{\bullet}}^{S} = k_{XH/Y_{\bullet}}^{0}/(1 + K_{XH/S}^{S}[S])$$
 (I)

which also indicates that the ratio of experimental rate constants in any pair of solvents will be independent of the reactivity of Y.8 This independence has been verified many times.<sup>8,9</sup> Such KSEs can be *quantitatively* described by eq II, 8,9 where  $\alpha_2^H$  and  $\beta_2^H$  are the Abraham et al.'s10,11 thermodynamically based constants representing, respectively, the relative HB donor (HBD) ability of solute XH in CCl<sub>4</sub> [range = 0.00 (alkanes) to  $\sim$ 1.0 (strong organic acids)]10 and the relative HB acceptor (HBA) ability of solute S in CCl<sub>4</sub> [range = 0.00 (alkanes) to 1.00 (hexamethylphosphortria-

$$\log(k_{\rm XH/Y}^{\rm S} {}^{\bullet} {}^{\prime} {\rm M}^{-1} \, {\rm s}^{-1}) = \log(k_{\rm XH/Y}^{\rm 0} {}^{\prime} {\rm M}^{-1} \, {\rm s}^{-1}) - 8.3 \, \alpha_2^{\rm H} \, \beta_2^{\rm H} \quad ({\rm II})$$

Equation II correlates KSEs for H-abstractions7 by a variety of Y radicals from hydrocarbons, aniline, tert-butyl hydroperoxide, and numerous phenols<sup>8,9</sup> (and other substrates).<sup>12</sup> However, in ionizing solvents (e.g., methanol), ionizable substrates (e.g., phenol) react with electron-deficient radicals more rapidly than eq II would predict.8 This is due to fast electron transfer to Y from the (generally low) concentration of X<sup>-</sup> anion present in equilibrium with XH (eq 6). This Sequential Proton Loss Electron Transfer

$$XH + Y^{\bullet} \rightleftharpoons H^{+} + X^{-} + Y^{\bullet} \rightarrow H^{+} + X^{\bullet} + Y^{-}$$
 (6)

(SPLET)8,13 occurs in parallel with the "normal" H-abstraction7 (eq 4). SPLET can usually be completely suppressed by the addition of low concentrations of acetic acid.8,13

The quenching of the  $\pi,\pi^*$  triplet ketone, 2-benzoylthiophene (3BT\*), by a variety of substrates has been previously studied by laser flash photolysis (LFP).5a,6a It was concluded that the phenol + 3BT\* reaction involved a HB exciplex and gave the phenoxyl (PhO•) and BT ketyl (BTH•) radicals by a (concerted) ETPT mechanism with a quantum yield close to unity.6a Herein, we address the question: Are the KSEs for this PhOH + 3BT\* HB exciplex reaction correctly described by eq II?

A pulsed Nd:YAG laser ( $\lambda_{exc}$ = 355 nm, 10 ns pulse, energy 15-17 mJ) was employed with [BT] chosen to give an absorbance of 0.35 ([BT] = 1.9-2.5 mM). The solvents (Table 1) were of the highest purity available and were used as received. LFP of deaerated solutions of BT gives the 3BT\* absorption (maxima at 350 and 600 nm). The  ${}^3BT^*$  decay was accelerated by the addition of PhOH (in a dose-dependent manner) with the appearance of absorbancies due to the ketyl (BTH•,  $\lambda_{max} = 350$  and 580 nm) and PhO• ( $\lambda_{max} =$ 380 and 410 nm) radicals, both being formed with a high efficiency (Supporting Information). A plot of the logarithms of the rate constants for  ${}^{3}\mathbf{BT}^{*}$  quenching by phenol,  $k_{q}$  (Table 1), in the solvents against the solvents'  $\beta_2^H$  values shows an excellent linear correlation provided the point for solvent 2 is ignored (as it should be  $^{14}$  Figure 1, solid line). The slope of this line is -3.9, which is significantly lower than the slope calculated from eq II and phenol's  $\alpha_2^{\rm H}$  value of 0.59<sup>9</sup> [i.e.,  $-8.3 \times 0.59 = -4.9$  (Figure 1, dashed

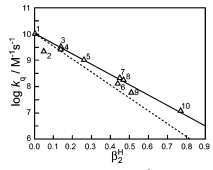
<sup>†</sup> Universidad de Valencia.

Warsaw University.
National Research Council.

**Table 1.** Rate Constants for Quenching of  ${}^3\text{BT}^*$  by Phenol in Various Solvents with  $\beta_2^H$  Values for the Solvents

	•		
no.	solvent	$eta_2^{H\;a}$	$k_{\rm q}/10^8~{\rm M}^{-1}~{\rm s}^{-1}$
1	n-C <sub>8</sub> H <sub>18</sub>	0.00	105 ± 5
2	$CH_2Cl_2$	0.05	$23^{b}$
3	PhH	0.14	$28 \pm 3$
4	$PhCH_3$	0.14	$35 \pm 2$
5	PhOCH <sub>3</sub>	0.26	$11.3 \pm 0.5$
6	CH <sub>3</sub> CN	0.44	$1.4^{b}$
7	$CH_3C(O)OC_2H_5$	0.45	$2.3 \pm 0.2$
8	1,4-dioxane	$0.47^{c}$	$1.9 \pm 0.1$
9	tetrahydrofuran	0.51	$0.62 \pm 0.05$
10	$(C_2H_5O)_3PO$	0.77	$0.13 \pm 0.04$

<sup>&</sup>lt;sup>a</sup> From ref 11. <sup>b</sup> Value of  $k_q$  from ref 6a. <sup>c</sup> From ref 13b.



**Figure 1.** Logarithm of  $k_q$  for the reaction of  ${}^3BT^*$  with phenol in various solvents (numbered as in Table 1) versus the solvents'  $\beta_2^H$  values. Solvent 2 was not included in constructing the solid correlation line<sup>14</sup> (slope -3.9). The dashed line has the predicted (eq II) slope of -4.9; see text.

line)]. That is, the magnitudes of the HBA solvents' suppression of the rate are less than those for all free radical + phenol H-atom abstraction<sup>7</sup> reactions examined to date. This would be the case if the  ${}^3BT^* + PhOH$  reaction had a significant SPLET component. Roughless However, this appears improbable for two reasons: (i) even for reactions where a SPLET mechanism is strongly favored, it does not occur in dioxane (8); and (ii) the addition of 10 mM acetic acid to THF and triethyl phosphate produced a very small increase in  $k_q$  (Supporting Information) rather than the decrease expected if the SPLET mechanism occurred.

We propose that the slope of the solid line in Figure 1 (-3.9) is lower than predicted (-4.9) because, in HBA solvents, the <sup>3</sup>BT\* + PhOH reaction occurs by two mechanisms, A and B. A involves Leigh's<sup>3</sup> exciplex process with HB formation between <sup>3</sup>BT\* and only those PhOH molecules that are not H-bonded to a solvent molecule and for which a plot of log  $k_A$  versus  $\beta_2^H$  would fit eq II (slope -4.9). The contribution of mechanism B increases as the HBA activity of the solvent increases. It must involve PhOH molecules that are H-bonded to solvent molecules and does not occur in the non-HBA solvent, octane. This mechanism could involve either all, or some subset, of PhOH···S species. We suggest that **B** occurs by electron transfer to <sup>3</sup>BT\* from (reactive) PhOH···S followed by, or concerted with, proton transfer from the resulting highly acidic phenol radical cation, (PhOH)\*+, to the S to which it is already H-bonded. The stronger this H-bond, the more facile the B mechanism will be. The overall KSE is described by eq III. We predict that KSEs for quenching of 3BT\* by other phenols will also be described by this equation.

$$\log(k^{S}_{A} + k^{S}_{B}) = \log k^{O}_{A} - 8.3 \times \{1 - (1 - 3.9/4.9)\} \alpha_{2}^{H} \beta_{2}^{H}$$
$$= \log k^{O}_{A} - 6.6 \alpha_{2}^{H} \beta_{2}^{H} = \log k^{S}_{O}$$
(III)

In our review of KSEs on H-atom abstractions from phenols,<sup>8</sup> the (formally trimolecular) ETPT mechanism, B, in which the electron and proton go to *different acceptors*, was described as an

unidentified fourth possible mechanism for intermolecular H-abstractions from phenols by strongly oxidizing radicals and nonradicals. Unimolecular (i.e., intramolecular) examples of this mechanism are known in chemistry and biology,<sup>8</sup> but the only intermolecular example of phenol oxidation by this mechanism that we are aware of involved an added solute as the proton acceptor<sup>16</sup> and hence is not, strictly speaking, a KSE.

In conclusion, a fourth mechanism, B, for H-atom abstraction from phenol in PhOH···S HB complexes has been identified and quantified. Although the mechanism B rate increased with the solvents' HBA strength ( $\beta_2^{\text{H}}$ ), the overall rate decreased as  $\beta_2^{\text{H}}$  increased. Nevertheless, the exciting possibility remains of "inverse" KSEs where the rates of H-atom abstraction from a HBD actually increase as the solvents become stronger HBAs.

**Acknowledgment.** R.E.G. thanks the Ministerio de Educación y Ciencia for the Ramón y Cajal contract. We also thank Dr. Mario Foti for some extremely helpful comments.

**Supporting Information Available:** Transient absorption spectra and kinetic data for quenching of <sup>3</sup>BT\* by pnenol in different solvents. This material is available free of charge via Internet at http://pubs.acs.org.

## References

- Das, P. K.; Encinas, M. V.; Scaiano, J. C. J. Am. Chem. Soc. 1981, 103, 4154–4162.
- (2) Wagner, P. J.; Kemppainen, A. E.; Schott, H. N. J. Am. Chem. Soc. 1973, 95, 5604-5614.
- (3) (a) Leigh, W. J.; Lathioor, E. C.; St. Pierre, M. J. J. Am. Chem. Soc. 1996, 118, 12339-12348. (b) See also: Lathioor, E. C.; Leigh, W. J. Photochem. Photobiol. 2006, 82, 291-300.
- (4) The photoreduction of benzophenone by aliphatic amines had previously been suggested to proceed via amine triplet ketone charge-transfer complexes (exciplexes) which favored subsequent H-abstraction reactions. See: (a) Inbar, S.; Linschitz, H.; Cohen, S. G. J. Am. Chem. Soc. 1981, 103, 1048–1054. (b) Inbar, S.; Linschitz, H.; Cohen, S. G. J. Am. Chem. Soc. 1980, 102, 1419–1421.
- (5) (a) Pérez-Prieto, J.; Galian, R. E.; Morant-Miñana, M. C.; Miranda, M. A. Chem. Commun. 2005, 3180–3182. (b) Biczók, L.; Bérces, T.; Linschitz, H. J. Am. Chem. Soc. 1997, 119, 11071–11077.
- (6) (a) Pérez-Prieto, J.; Boscá, F.; Galian, R. E.; Lahoz, A.; Domingo, L. R.; Miranda, M. A. J. Org. Chem. 2003, 68, 5104–5113. (b) Evans, C.; Scaiano, J. C.; Ingold, K. U. J. Am. Chem. Soc. 1992, 114, 4589–4593.
- (7) H-abstraction is used to refer to two, nonexclusive, mechanisms:<sup>8</sup> (i) transfer of a proton and one of its bonding electrons (hydrogen atom transfer, HAT); (ii) transfer of a proton with an accompanying electron from a nonbonding orbital (proton-coupled electron transfer, PCET).
- (8) Litwinienko, G.; Ingold, K. U. Acc. Chem. Res. 2007, 40, 222-230.
- (9) Snelgrove, D. W.; Lusztyk, J.; Banks, J. T.; Mulder, P.; Ingold, K. U. J. Am. Chem. Soc. 2001, 123, 469–477.
- (10) Abraham, M. H.; Grellier, P. L.; Prior, D. V.; Duce, P. P.; Morris, J. J.; Taylor, P. J. J. Chem. Soc., Perkin Trans. 2 1989, 699-711.
- (11) Abraham, M. H.; Grellier, P. L.; Prior, D. V.; Morris, J. J.; Taylor, P. J. J. Chem. Soc., Perkin Trans. 2 1990, 521–529.
- (12) (a) Foti, M. C.; Sortino, S.; Ingold, K. U. Chem.—Eur. J. 2005, 11, 1942—1948. (b) Astolfi, P.; Greci, L.; Paul, T.; Ingold, K. U. J. Chem. Soc., Perkin Trans. 2 2001, 1631—1633.
- (13) (a) Litwinienko, G.; Ingold, K. U. J. Org. Chem. 2003, 68, 3433-3438.
  (b) Litwinienko, G.; Ingold, K. U. J. Org. Chem. 2004, 69, 5888-5896.
  (c) Litwinienko, G.; Ingold, K. U. J. Org. Chem. 2005, 70, 8982-8990.
- (14) The  $\beta_2^H$  value for CH<sub>2</sub>Cl<sub>2</sub> is too small. In a full paper, we will show that  $\beta_2^H$  for CH<sub>2</sub>Cl<sub>2</sub> should be revised upwards to 0.15, a value that puts log  $k_q$  in 2 essentially on the solid correlation line shown in Figure 1.
- (15) In the only previous example of acetic acid increasing the rate of an ArOH/ Y\* reaction, <sup>13c</sup> acceleration was attributed to the excellent anion solvating abilities of the acid enhancing ArOH ionization and, hence, the contribution of SPLET to the measured rate.
- (16) The quenching of triplet  $C_{60}$  by phenols is accelerated by added pyridine (Pyr) bases. <sup>17</sup> This was attributed to a trimolecular transition state ET from PhOH···Pyr to  ${}^3C_{60}$  concerted with PT to the H-bonded base. <sup>17</sup>
- (17) (a) Biczók, L.; Linschitz, H. J. Phys. Chem. 1995, 99, 1843–1845. (b) Gupta, N.; Linschitz, H.; Biczók, L. Fullerene Sci. Technol. 1997, 5, 343–353.

JA071716Y