Substituent Effect in Photochemistry of Carbonyl Compounds: *a*-Halovalerophenones

Sungsu Cho and Bong Ser Park*

Department of Chemistry, Dongguk University, Seoul 100-715, Korea Received September 18, 2003

Valerophenones containing a substituent at alpha position to the carbonyl group show the remarkable substituent effects on their photochemical reactions. α -Bromovalerophenone gives only the C-Br bond cleavage products, but the α -chlorovalerophenone follows the classical Norrish/Yang reaction pathway predominantly.

Key Words : Substituent effect, Photochemistry, Halo ketone

Photochemical reactions of alpha-halo ketones have been known for a long time in organic photochemistry.¹ The usual reaction routes of these ketones involve photoinduced carbon halogen bond cleavage reactions either from singlet or triplet excited states. The efficient cleavage reactions have been used as photoinitiators for polymerization of several vinyl monomers.² The reaction has recently found itself an useful application into a photoremovable protecting group.³ If alpha-halo ketones contain a hydrogen atom to be abstracted nearby, the carbon halogen bond cleavage reaction can occur from an intermediate of photoinduced hydrogen abstraction reactions as exemplified by photorelease of HCl from 2,5-dimethylphenacyl chloride.⁴ Most recently Wessig and coworkers, have reported a new type of reaction from irradiation of some butyrophenones containing a good leaving group such as mesyloxy or trifluoroacetoxy group at alpha to the carbonyl (Scheme 1).⁵ These papers did not mention whether halogens as a leaving group had been tested in their studies. It would be very interesting to see which reaction sequence the a-halo substituted ketones will follow among several competing reaction routes. In the course of our studies on the substituent effect in the Yang photocyclization,6 we had chances to look into photochemical properties of α -halovalerophenones and found an interesting substituent effect7 of these ketones. Here we would like to report our results on this system.

 α -Bromovalerophenone (1) and α -chlorovalerophenone



Scheme 1. Photochemical reaction of α -substituted butyrophenone.

(2) were prepared by α -bromination and chlorination of valerophenone using CuBr₂ or Br₂ and SO₂Cl₂, respectively. Photochemical reactions of both 1 and 2 were monitored by TLC and turned out to be time dependent. Some of the initially formed products were replaced by other products upon prolonged irradiation. The secondary photoreaction was confirmed later by NMR scale photolysis, vide infra. Each product was isolated by column chromatography using hexane and ethyl acetate in 10 to 1 ratio as eluents. When the reaction stopped at early stage just before the secondary photoreaction started, two major products from 1 were identified to be valerophenone (VP) and β -bromovalerophenone (1P) according to usual structural analysis. Compound 2 gave four major products, which were two isomeric 2-chloro-4-methyl-1-phenylcyclobutanols (CB),8 α -chloroacetophenone (2P) and valerophenone (VP). The starting ketones staved intact in refluxing benzene for several hours in dark, which confirmed that the process was not thermal but photochemical.



In order to monitor the reaction more closely, a sample in an NMR tube containing 1 or 2 in benzene-d₆ (0.02 M) was prepared and irradiated under the same condition as above. For 2 hours irradiation of 1, the only products detected were **VP** and **1P**. As the irradiation continued, the ratio of **VP** and **1P** decreased in compensation for appearance of acetophenone, a photoproduct of **VP**. Longer irradiation made the reaction mixture more complicated presumably due to formation of more secondary photoproducts not only from **VP** but also from **1P**. In case of **2**, the reaction mixture

Substituent Effect in Photochemistry of α -Halovalerophenones

 Table 1. Product Distribution of Photolysis of 1 and 2 in Benzene

Compounds	VP	1P	2P	СВ
I	35%	65%		
2	10%		13%	77%

showed formation of four products, whose structures were already identified in large scale photolysis. The secondary photoreaction from **VP** was also detected upon prolonged irradiation of **2** similarly to the case of **1**. Table 1 summarizes product distribution of photolysis of **1** and **2**.

We also applied the same reaction condition (in dichloromethane containing an acid scavenger, N-methylimidazole) that Wessig used in his study mentioned above, but could not observe formation of the cyclopropyl ketones from 1 or 2. The reaction was not much different from the results in benzene except that **IP** was replaced by 1-phenylpent-2-en-1-one (**PP**) in photolysis of 1. Further experiments using other acid scavengers suggested that **IP** was the product of HBr addition to the initially formed **PP**.

The major difference of photochemical reactions of 1 and 2 is that the C-Br bond cleavage is the only reaction pathway from 1, but the Norrish/Yang reaction becomes the major reaction from 2. The resulting radical from C-Br cleavage of 1 then abstracts a hydrogen atom from any possible sources in reaction mixture including HBr to give VP. There were several reports suggesting that ion pairs could be formed from the initially formed radical pairs via electron transfer.¹ However, the extent that the electron transfer occurs can be considered negligible at least in benzene. The dehydrohalogenation product, PP, can also be formed from the radical intermediate as in the case of other α -substituted alkanones.⁹ In our studies with 1, however, the dehydrohalogenated product was not observed in benzene. Apparently the α,β unsaturated ketone is trapped by HBr via Michael addition in acidic media to give 1P as soon as it is formed.

In case of photolysis of 2, the C-Cl bond cleavage becomes only a minor reaction pathway due to the relatively faster 7-hydrogen abstraction reaction. The different photochemical behaviors of 1 and 2 can easily be attributed to bond energy difference of C-Br and C-Cl, which the former (70 kcal/mol) is ca. 14 kcal/mol weaker than the latter (84 kcal/mol).¹⁰ Considering that triplet excited state energy is ca. 70 kcal/mol for both α -bromoacetophenone and α chloroacetophenone,¹² the explanation based on the bond energies does not seem unreasonable. The better explanation can, however, be applied here with known kinetic data. Scaiano et al. have studied photoinduced C-X cleavage reactions of α -bromoacetophenone and α -chloroacetophenone by laser flash photolysis and reported that the cleavage rate of the former is over 1×10^{10} s⁻¹ while that of the latter is ca. 3×10^6 s^{-1.11} This is consistent with our results of 1 and 2. In case of 1, the C-Br bond cleavage occurs very fast with a rate which µ-hydrogen abstraction cannot compete. Typical rate constant of y-hydrogen abstraction (k_0) of α -substituted valerophenone is in the order of $10^8 \text{ s}^{-1,12}$ In case of **2**, the γ -hydrogen abstraction

reaction occurs much faster than the C-Cl bond cleavage, so the former reaction prevails. From our results, the rate constant of C-Cl bond cleavage of **2** can be estimated to be ca. $3 \times 10^7 \text{ M}^{-1}$, assuming that $k_{\rm H}$ of **2** is not much different from that of α -fluorovalerophenone.

As mentioned above, photolysis of 1 and 2 did not give the cyclopropyl ketones that Wessig et al. found from several α substituted ketones. They reported that ketones bearing a good leaving group such as sulfonate or nitrate at alpha position to the carbonyl group abstracted a µ-hydrogen with a rate of 3×10^8 s⁻¹ and formation of the cyclopropyl ketones from the resulting biradical intermediate occurred with a rate of over 4×10^7 s⁻¹. Thus the kinetic data can explain why the C-Br cleavage reaction of 1 becomes predominant. The reason why the cyclopropyl ketone is missing in photolysis of 2, however, is still puzzling. Wessig has mentioned the involvement of intramolecular H-bonded biradical intermediate shown below (BR1 and BR2) in their reactions, Even though it may explain the different reactivities between the sulfonate ester and the chloride, the more convincing evidences still await further experiments.



In summary, valerophenones containing a substituent at alpha position to the carbonyl group showed the remarkable substituent effects on their photochemical reactions. The α -bromo ketone, 1, gives only the C-Br bond cleavage products, but the α -chloro ketone, 2, follows the classical Norrish/ Yang reaction pathway predominantly. These behaviors are in a marked contrast with those of ketones bearing other substituents such as O-sulfonyl and O-nitro. Currently other ketones with these substituents are being tested in our laboratory in order to obtain the better idea of the interesting substituent effect.

Experimental Section

Synthesis of 1: A 250 mL round bottomed flask was charged with copper(II) bromide (2.75 g, 12.3 mmol), valerophenone (1.00 g, 6.2 mmol), chloroform (10 mL) and ethyl acetate (100 mL) under argon atmosphere and the reaction mixture was refluxed for 48 hours. Distilled water (20 mL) was added to the solution and insoluble solids were filtered through Buchner funnel. The filtered solution was extracted with ethyl acetate and washed with brine. The combined organic layer was dried using anhydrous MgSO₄ and concentrated at reduced pressure. The crude product was purified by column chromatography using dichloromethane and *n*-hexane in 2 to 1 to give 1.37 g of α -bromovalero44 Bull. Korean Chem. Soc. 2004, Vol. 25, No. 1

phenone (92% yield).

Spectroscopic properties of 1: ¹H NMR (CDCl₃, 200 MHz): δ 8.02 (m. 2H), 7.53 (m, 3H). 5.15 (dd, 1H, J = 7.1, 7.0 Hz). 2.15 (m. 2H), 1.51 (m. 2H). 0.98 (t. 3H, J = 7.0 Hz). ¹³C NMR (CDCl₃, 50 MHz): δ 193.4. 134.6. 133.8. 129.0. 128.9, 47.1. 35.5, 20.9, 13.7. IR (neat): 1688 (C=O) cm⁻¹, EI mass: 240 (M⁺).

Synthesis of 2: Sulfuryl chloride (8.32 g. 61.6 mmol) and *p*-toluenesulfonic acid (0.10 g) were added into 250 mL round bottomed flask containing valerophenone (1.00 g. 6.2 mmol) in dichloromethane. The reaction mixture was refluxed for 12 hours. Distilled water (20 mL) was added and the organic layer was extracted with dichloromethane. After washing with brine, the organic layer was dried using anhydrous MgSO₄ and concentrated at reduced pressure. The crude product was purified by column chromatography using dichloromethane and *n*-hexane in 2 to 1 to give 1.53 g of α -chlorovalerophenone (79% yield).

Spectroscopic properties of 2: ¹H NMR (CDCl₃, 200 MHz): δ 8.01 (m. 2H), 7.55 (m, 3H), 5.13 (dd, 1H, J = 8.0, 5.8 Hz), 2.03 (m, 2H), 1.53 (m, 2H), 0.98 (t. 3H, J = 7.3 Hz), ¹³C NMR (CDCl₃, 50 MHz): δ 193.8, 134.7, 133.8, 129.0, 128.9, 57.7, 35.7, 19.7, 13.7, IR (neat): 1693 (C=O) cm⁻¹ EI mass: 196 (M⁺).

Photolysis: The starting ketones in benzene (typically 0.01-0.02 M) were irradiated in an immersion well with argon bubbling using Pyrex (or Uranium) filtered light of a 450 W Hanovia medium pressure mercury are lamp. The reaction mixture was concentrated at reduced pressure and the resulting crude product mixture was separated by column chromatography using *n*-hexane and ethyl acetate in 7 to 1.

For NMR scale photolysis, an NMR tube containing ketones in benzene- d_6 was degassed and irradiated by attaching it to the side of an immersion well using Pyrex (or Uranium) filtered light of a 450 W Hanovia medium pressure mercury arc lamp. To control reaction temperature the sample was immersed in an temperature control bath during the irradiation.

Spectroscopic properties of **1P**: ¹H NMR (CDCl₃. 200 MHz) δ 7.97 (d. 2H. *J* = 7.5 Hz), 7.53 (m. 3H), 4.58 (m. 1H), 3.74, 3.42 (doublets of AB quartets. 2H. *J* = 17.2, 7.2 Hz), 1.97 (m. 2H), 1.11 (t, 3H, *J* = 7.2 Hz), ¹³C NMR (CDCl₃. 50 MHz) δ 196.9, 136.7, 133.5, 128.8, 128.2, 51.8, 47.4, 32.2.

12.2. IR (neat): 1686 (C=O) cm⁻¹ EI mass: 240 (M⁻).

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