New Preparative Method for 2-Arylpropanoic Acids by Oxidative Aryl Migration in Aryl α -Seleno- and Aryl α -Telluro-ethyl Ketones

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Oxidation with m-chloroperbenzoic acid of the ethylene acetals of aryl α -phenylseleno- or aryl α -phenyltelluro-ethyl ketones prepared by treating the corresponding α -bromo compounds with diphenyl diselenide-sodium or diphenyl ditelluride-sodium, respectively, affords hydroxyethyl 2-arylpropanoates in moderate to good yields via aryl group migration.

It has recently been reported that oxidation of alkyl phenyl selenides¹ or tellurides² with *m*-chloroperbenzoic acid (MCPBA) in methanol gave dialkyl ethers. The reaction is accompanied by phenyl migration in selenides and tellurides having a phenyl group vicinal to the phenylselenium or phenyltellurium moieties. Using this reaction we have succeeded in developing a new method for the synthesis of 2-arylpropanoic acids which are pharmaceutically important

compounds exhibiting anti-inflammatory and analgesic activities ³

Aryl ethyl ketone (1) was brominated at the α -position to give (2) which was then converted into acetal (3) with ethylene glycol in the presence of toluene-p-sulphonic acid [overall yield of (3) from (1) was 88—98%]. When (3) was added to a tetrahydrofuran-dimethylformamide solution of diphenyl diselenide or diphenyl ditelluride and sodium wire under a N_2

ArC—Et
$$\xrightarrow{Br_2}$$
 ArC—CHMe $\xrightarrow{H^+}$ HOCH₂CH₂OH ArC—CHMe O Br

(1)

(2)

(3)

PhSeCI

ArC—CHMe (PhM)₂-Na (PhM)₂-Na (THF - DMF)

ArC—CHMe (II I O O MPh

(5)

M = Se, Te

(4)

Ar — C—CHMe (4)

Ar — C—CHMe (4)

Ar — C—CHMe (6)

atmosphere and the resulting mixture was stirred under reflux for 6—20 h, the bromine atom was substituted by the PhSe or PhTe group to afford (4) in 50—80% (for M = Se) and in 35—60% (for M = Te) yield.†‡ This substitution did not proceed by using the $(PhM)_2$ -NaBH₄-EtOH (M = Se, Te) system which is known as a source of PhM⁻ anion and the starting compound (3) was recovered. Attempts to prepare (4;

Table 1. 2-Arylpropanoic acids from aryl ethyl ketones (1).

	Yield (%)	
M X in Ar	(4) ^a	(6) ^b
Н	75	80
Me	83	86
$\mathbf{B}\mathbf{u}^{\mathrm{i}}$	70	82
Ph	81	56
Br	74	85
Н	62	80
Me	34	62
$\mathbf{B}\mathbf{u^i}$	50	85
	H Me Bu ⁱ Ph Br H Me	X in Ar (4)a H 75 Me 83 Bui 70 Ph 81 Br 74 H 62 Me 34

^a Isolated yield based on (3). ^b Isolated yield based on (4).

M=Se) by acetalation of aryl α -phenylselenoethyl ketone (5), prepared by selenation of (1) with phenylselenenyl chloride, resulted in decomposition to (1) and diphenyl diselenide.

Compound (4) was then treated with 5 equiv. MCPBA in methanol at room temperature for 10 min to 1 h. After normal work-up (addition of aq. NaCl, washing with aq. Na₂S₂O₃ and aq. NaHCO₃, and diethyl ether extraction), evaporation of the solvent left an oily compound which was subjected to column chromatography [SiO₂, hexane–EtOAc (10/1 to 1/1) as eluent] to give the hydroxyethyl ester of 2-arylpropanoic acid (6).† The isolated yield of (6) was 55—58% based on (4). Alkaline hydrolysis of (6) readily afforded the corresponding acid (7). Typical results are summarized in Table 1. We assume that the reaction proceeds via a selenone or tellurone intermediate (A) in which aryl group migration occurs. Direct oxidation of (5) with MCPBA in methanol did not produce any of (6) or the analogous ester.

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References

- 1 S. Uemura and S. Fukuzawa, Tetrahedron Lett., 1983, 24, 4347.
- 2 S. Uemura, S. Fukuzawa, and A. Toshimitsu, J. Chem. Soc., Chem. Commun., 1983, 1501.
- 3 For example, K. Fujii, K. Nakao, and T. Yamauchi, *Synthesis*, 1983, 444, and references therein.

[†] All new compounds showed satisfactory combustion analytical and spectroscopic data (¹H and ¹³C n.m.r. and i.r.).

[‡] All compounds except (4; X = Me, M = Te) (pale yellow crystals, m.p. 105—106 °C) are yellow oily substances.