Preliminary communication

AN ORGANOMETALLIC ROUTE TO (±)-3-p-TOLUENESULFINYL-2-PYRONE

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Summary

3-Bromo-2-pyrone reacts with dimethylcopperlithium and then with p-tolyl p-toluenethiosulfonate to form 3-p-toluenethio-2-pyrone. Organocopper species are likely intermediates in this reaction. Oxidation of the sulfide produces (±)-3-p-toluenesulfinyl-2-pyrone. The overall process involves an important and useful umpolung, a reversal of reactivity at C(3) of the pyrone ring from an electrophilic center to a nucleophilic center.

As part of our research program on the use of enantiomerically pure unsaturated ketone and lactone sulfoxides for asymmetric carbon—carbon bond formation with various organometallic reagents [1], we required (+)- or (-)-3p-toluenesulfinyl-2-pyrone (3). Because even racemic pyrone sulfoxide 3 had not been made before, we chose to prepare the racemate and to investigate its chemistry before attempting the more difficult task of preparing enantiomerically pure (+)- or (-)-3. We report here a reliable and very short (3-step, 2-pot) synthesis of (±)-3 involving organocopper intermediates.

3-Bromo-2-pyrone (1a) was prepared via bromination—dehydrobromination of 2-pyrone [2] using a slightly modified (40 h instead of 24 h reflux, 1.4/1 molar ratio of pyrone/carbon tetrachloride; 59% overall yield from 2-pyrone) literature procedure [3]. Attempts at direct bromine \rightarrow lithium exchange [4], (i.e. at formation of 3-lithiopyrone 1b) using n-butyllithium at -78° C failed, giving only what appeared to be n-butyl addition products [5]. In contrast, bromine \rightarrow copper exchange was highly successful.

Treating 3-bromo-2-pyrone (1a) with 2 equivalents of dimethylcopperlithium in diethyl ether at -78° C led, after 2 h, to complete bromine \rightarrow copper exchange; hydrolytic work-up gave 2-pyrone in good yield, with less than 5% of the coupling product [6] 3-methyl-2-pyrone being formed. A mechanistic

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(1a, X = Br; 1b, X = Li; 2, X = p-ToIS;3, X = p-ToIS

rationalization of this result is shown in Scheme 1 involving transient copper(III) intermediate 4 which apparently undergoes selective reductive elimination of ethane (rather than of 3-methyl-2-pyrone) to form a 3-cuprio-2pyrone (e.g. 5); the excess dimethylcopperlithium might react with cupriopyrone 5 to form a more reactive cuprate species (e.g. 6) and possibly methylcopper (a yellow-green precipitate is initially observed) [7]. Similar bromine \rightarrow copper exchanges instead of carbon—carbon couplings are known during reaction of organocopper reagents with some α -bromoketones [8], α -bromoacrylate esters [9], and alkenyl bromides [10]; it has been possible in these cases to intercept the newly-formed organocopper species with a limited number of electrophiles either intermolecularly [8,9] or intramolecularly [10].



SCHEME 1

When 3-bromo-2-pyrone (1a) reacts with dimethylcopperlithium at -78° C for 3 h and then with *p*-tolyl *p*-toluenethiosulfonate [11] at -30° C for 2 h, a coupling reaction occurs to form 3-*p*-tolylthio-2-pyrone (2, ¹H, NMR (CDCl₃): δ 6.54 (dd, 1H, H₄)) isolated after preparative TLC in 49% yield (based on bromopyrone 1a, eq. 1). Standard oxidation of sulfide 2 produces the desired, racemic, crystalline, lactone sulfoxide 3 (m.p. 112–113°C; ¹H NMR (CDCl₃): δ 8.08 (dd, 1H, J_{4,5} 6.7 Hz, J_{4,6} 2.1 Hz, H₄). Analysis: found: C, 61.52; H, 4.30; S, 13.69; C₁₂H₁₀O₃S calcd.: C, 61.60; H, 4.31; S, 13.61%).

It is noteworthy that a C(3)-tolylthic group causes an upfield chemical shift of H(4) comparable to that caused by a C(3)-methoxy group, whereas a C(3)tolylsulfinyl group causes a downfield chemical shift of H(4) comparable to that caused by a C(3)-ethoxycarbonyl group [12].



These significant results represent an important and useful umpolung [13] or reversal of reactivity at C(3) of bromopyrone 1a from an electrophilic center to a nucleophilic center, thereby allowing regiospecific attachment of a sulfur substituent as an electrophile. Indeed C(3) attachment of other electrophiles should be possible as well; we have had some initial success with introducing an allyl group.

We are actively studying the chemistry of 3-sulfinyl-2-pyrone (3) as well as the reactivity of the intermediate organocopper species (e.g. 5, 6) toward various electrophiles, and a full report will appear in due course.

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References

- (a) G.H. Posner, T.P. Kogan, S.R. Haines, and L. Frye, Tetrahedron Lett., 25 (1984) 2627; (b) G.H. Posner, T.P. Kogan, and M. Hulce, ibid., 25 (1984) 383; (c) G.H. Posner and M. Hulce, ibid., 25 (1984) 379; (d) G.H. Posner, L.L. Frye, and M. Hulce, Tetrahedron, 40 (1984) 401.
- 2 Prepared from commercially available 5,6-dihydro-2-pyrone according to M. Nakagawa, M. Tonozuka, M. Obi, M. Kinchi, T. Hino, and Y. Ban, Synthesis, (1974) 510.
- 3 W.H. Pirkle and M. Dines, J. Org. Chem., 34 (1969) 2239.
- 4 R.G. Jones and H. Gilman, Org. Reactions, 6 (1951) 339.

- 5 For a study of addition of Grignard reagents to 2-pyrones see P. Lhoste, M. Moreau and J. Dreux, Tetrahedron, 40 (1984) 1551, and ref. therein.
- 6 G.H. Posner, Org. Reactions, 22 (1975) 258.
- 7 G.H. Posner, An Introduction to Synthesis Using Organocopper Reagents, John Wiley, New York, 1980.
- 8 (a) G.H. Posner and J.J. Sterling, J. Am. Chem. Soc., 95 (1973) 3076; see also (b) A.E. Greene, J.-P. Lansard, J.-L. Luche, and C. Petrier, J. Org. Chem., 48 (1983) 4763; (c) G.R. Clark, J. Lin and M. Nikaido, Tetrahedron Lett., 25 (1984) 2645.
- 9 (a) J.P. Marino and D.M. Floyd, J. Am. Chem. Soc., 96 (1974) 7138; (b) J.P. Marino and D.M. Floyd, Tetrahedron Lett., (1975) 3897, 3901; (c) See also J. Klein and R. Levene, J. Am. Chem. Soc., 94 (1972) 2520.
- 10 (a) E.J. Corey and I. Kuwajima, J. Am. Chem. Soc., 92 (1970) 395; (b) E.J. Corey, M. Narisada, T. Hiroaka, and R.A. Ellison, ibid., 92 (1970) 396.
- 11 Prepared by m-chloroperbenzoic acid oxidation of di-p-tolyl disulfide.
- 12 W.H. Pirkle and W.V. Turner, J. Org. Chem., 40 (1975) 1616.
- 13 D. Seebach, Angew. Chem. Int. Ed. Engl., 18 (1979) 239.