A Novel Synthesis of 4-Dioxy-2-enones from 1,3-Dienes using Pyridinium Dichromate tert-Butyl Hydroperoxide

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Treatment of 1,3-dienes with the pyridinium dichromate (PDC)—ButOOH reagent system leads directly to the formation of 4-tert-butyldioxy-2-enones in good yield under mild reaction conditions.

The addition reaction of alkyl hydroperoxide and cyclopentadiene in the presence of Fe^{II} and Cu^{II} salts yields among other products, 4-alkoxycyclopent-2-ene.¹ Oxidative cleavage of endoperoxides of 1,3-dienes using acidic or basic reagents, ^{2a} Fe^{II}, Ti^{III}, Co^{II}, Rh^I and Pd⁰ salts or complexes^{2b}

gives rise to the corresponding 4-hydroxyenones as one of the products; however, the reactions generally lead to low yields and difficult separations.

The PDC-ButOOH system has been shown to be very effective in a variety of oxidative transformations.³ Interest-

Table 1 Reaction of 1,3-dienes with PDC-ButOOH

Entry	Substrate	Product ^a	Yield (%) ^b
1		Bu'00 2	76
2	\bigcirc	Bu'00	55
3	5	Bu'00 0	52
4		Bu'00 8	24°
5	9	Bu'00 OH 10	60
6	11	Bu'00 12	42°
7 (86
8 Pt	13 Ph	Ph Ph	73

^a All the products gave satisfactory spectral and analytical data. ^b Yields refer to isolated products taking into account the recovery of starting matrials (6–20%). ^c Formation of some unidentified polymeric material lowers the yield in these cases.

ingly, when 1,3-dienes were treated with a PDC and Bu t OOH mixture supported on silica gel (substrate: PDC: Bu t OOH = 1:2:3) for 8–13 h,† the only products that could be isolated in

† General procedure: Silica gel (1.505 g) and PDC (4 mmol, 1.505 g) were suspended in dichloromethane and cooled to 0 °C. This suspension was treated with ButOOH (6 mmol, 0.8 ml, 70% solution in water) to give a crimson coloured mixture. After 10 min, a 1,3-diene (2 mmol) was added and the reaction was monitored by TLC. The reaction mixture was filtered using a sintered-glass funnel, through a pad of silica gel. The solvent was removed and the product was purified by column chromatography (silica gel, eluent 2% ethyl acetate in hexane).

a reasonable yield were the 4-tert-butyldioxy-2-enones.‡ The results are shown in Table 1.

Both cyclic and acyclic dienes reacted with PDC-ButOOH and this transformation is the first report of its type in the literature. Although substrates 13 and 15 initially gave the corresponding dioxyenones, only the dienes could be isolated successfully. Surprisingly the diene moiety of the triene 11 did react enabling the isolation of compound 12. In spite of the possibility of multiple reaction pathways open to the dienes 7 and 11 only 8 and 12 respectively, were obtained. Diene 9 with two tertiary centres at the diene termini gave 10 as expected.

A set of experiments was carried out with cyclopentadiene illustrating the use of Cr^{VI} reagent in this reaction. Replacing PDC with chromium trioxide and 4 equiv. of ButOOH instead of 3 equiv., gave 2 in 74% yield. A catalytic amount of PDC (10 mol%) or chromium trioxide (10 mol%) and 6 equiv. of ButOOH (at -5°C, 24 h) in the reaction provided 2 in 65 and 70% yields respectively. Compounds 2, 4, 6 and 8 could be further converted to the corresponding 4-hydroxy-2-enones in quantitative yield following a closely related literature procedure for hydrogenolysis.⁴

The direct conversion of 1,3-dienes to the 4-dioxy-2-enones and their subsequent transformation to 4-hydroxy-2-enones could be useful in organic synthesis. *E.g.*, 4-hydroxy-cyclopent-2-enone which could be easily accessable from readily available and inexpensive cyclopentadiene, is a key intermediate in the synthesis of prostaglandins.⁵ The mechanism of this reaction is being studied in our laboratory.

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 $[\]ddagger$ 6–20% of the starting material was recovered. A small amount (5–10%) of a highly polar product formed in all these reactions has not been characterized.