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## Pd(II)Cl<sub>2</sub> Mediated Oxidative Cyclisation of Some 3-Hydroxy 4-Vinyl Furanoside Derivatives to Synthetically Valuable *Bis*-Furanosides

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Abstract: PdCl<sub>2</sub> mediated oxidation of 3-hydroxy-4-vinylfuranosides (la-k) lead to the formation of aldehydes, some of them are 'in situ' trapped as their lactols (llla,c,e,f,h) and further converted to bis-furanoside lactones lVa,c,e,f,h). Copyright © 1996 Published by Elsevier Science Ltd

Palladium (II) catalysed oxidation (Wacker process) of terminal olefins to methyl ketones is, probably one of the most well studied organic reactions. Its utility both in industrial processes and in synthetic organic chemistry is too well noted to make a mention in this short communication. Several terminal olefin derivatives possessing various functional groups have been reported to result in the formation of methyl ketones and/or aldehydes<sup>2</sup> under Wacker reaction conditions. However, till date there is no literature precedent to predict the exclusive formation of one of these products for a given substrate. Consequently there is still scope to investigate on a variety of factors to rationalise the formation of aldehyde/ketone with respect to functional groups, substitution pattern and their stereochemical disposition for a given alkene substrate, for utility in organic synthesis.

Our own interest in Palladium(II) catalysed oxidation reactions in the area of synthetic carbohydrate chemistry has earlier resulted in the finding of a new, mild method of deprotection of O-allyl and O-propenyl ethers of several alcohols, in serendipity.<sup>3</sup> Hence, we became interested in optimising reaction conditions required for Wacker oxidation of 3-hydroxy 4-vinylfuranoside derivatives to obtain exclusively aldehydes which *in situ* could be trapped as their lactols and further

oxidised to *bis*-furanoside lactones, that are valuable synthons. We describe herein our conclusive observations on the Wacker oxidation of 4-vinylfuranoside derivatives  $(1a-1)^{4a,b}$  that led to the synthesis of several valuable chirons (IV a,c,e,f,h).

Various furanoside terminal olefins (Ia-I) possessing diverse substitution pattern were oxidised under Wacker reaction conditions to obtain aldehydes (IIb,d,g,i,j,k) and lactols (IIIa,c,e,f,h) respectively (Table 1).

Thus it has been observed that when 3-hydroxy group is in *cis*-relation to the vinyl group in the furanosides (1a,c,e,f,h) the products formed were lactols (entry la,c,e,f,h) due to the trapping of

the so formed aldehydes. On the contrary if the 3-hydroxy group is in *trans* orientation (11) a mixture containing aldehyde and methyl ketone has resulted. It is worthwhile to note that there was no lactol formation in the latter case. It was also observed that when the 3-hydroxy group was protected Wacker oxidation of 1b,d,g,i,j,k invariably lead to the formation of aldehydes IIb, d,g,i,j,k irrespective of the *cis* or *trans* relation of the vinyl and 3-hydroxy group.

Lactol IIIa( $\alpha/\beta$ , 3/1) has been characterised from <sup>1</sup>H-NMR spectrum from the appearance of H-1 at  $\partial$  5.01 and H-6 ( $\alpha/\beta$ ) at  $\partial$  5.52 and  $\partial$  5.7. <sup>13</sup>C-NMR spectrum indicated C-1 at  $\partial$  100.3, C-6( $\alpha$ ) at  $\partial$  101.3 and C-6( $\beta$ ) at  $\partial$  109.1. The utility of lactols (IIIa,c,e,f,h) was shown by oxidising them further to their *bis*-furanoside synthons (IVa,c,e,f,h) (PDC/CH<sub>2</sub>CI<sub>2</sub>/4h).

Table 1: xPdCl2 mediated oxidation of alkenes

ENTRY	SOLVENTY	TIME	PRODUCT5	YIELDz (in %)
la	DMF:H <sub>2</sub> O	8 hr	IIIa	8 1
b	CH3CN:H2O	10 hr	IIb	87
С	DMF:H <sub>2</sub> O	8 hr	Illa	8 1
d	DMF:H <sub>2</sub> O	8 hr	IId	82
е	DMF:H <sub>2</sub> O	8 hr	IIIe	63
f	CH3CN:H2O	12 hr	Hif	77
g	CH3CN:H2O	14 hr	IIg	86
h	DMF:H <sub>2</sub> O	6 hr	IIIh	83
i	CH3CN:H2O	10 hr	Ш	85
j	CH3CN:H2O	13 hr	Hj	76
k	CH3CN:H2O	12 hr	IIk	89
I	DMF:H <sub>2</sub> O	18 hr	Aldehyde + methyl ketone (1:1)	68

x = PdCl<sub>2</sub> (0.2 mole equivalent), CuCl (1 mole equivalent)/O<sub>2</sub>

y = 7:1 ratio. For the convenience of isolation of product acetonitrile: water was chosen as solvent of choice in certain reactions.

z = Isolated yields

Thus, in conclusion Wacker oxidation of vinyl furanosides on resulted in the formation of aldehydes. It is perhaps due to the formation of the  $\pi$ -allyl palladium complex of the vinyl furanoside, which being hindered, facilitates the anti-markonikov hydration to give aldehydes as exclusive products. Taking advantage of the above observations various valuable chiral *bis*-furanoside lactols(IIIa,c,e,f,h) and *bis*-furanoside lactones (IVa,c,e,f,h) have been synthesised.

## Typical Experimental Procedure

To a stirred solution of PdCl<sub>2</sub> (10 mg, 0.0060 mmole) and CuCl (29 mg, 0.030 mmole) in DMF:H<sub>2</sub>O (3 ml, 7:1 ratio) under oxygen atmosphere was added **Ia** (70 mg, 0.030 mmole). The resulting black solution was stirred at room temperature for 8 hr and then extracted with ether. The organic layer was dried over Na<sub>2</sub>SO<sub>4</sub> and evaporated in vacuo. The crude product was purified by SiO<sub>2</sub> (60-120 mesh, 5 gms) column chromatography (EtOAc:hexane 1:1) to afford **IIIa** (60 mg, 81%)

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## References and Notes

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- 5. All new compounds gave satisfactory elemental analysis.

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