

## Communication

## First Pauson–Khand reaction on sugar acetylenes

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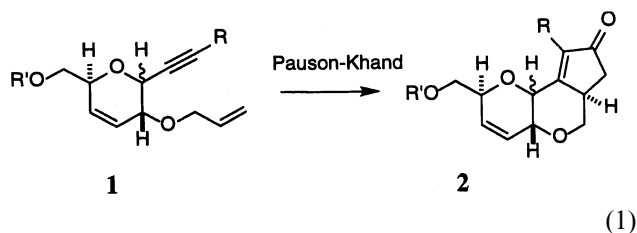
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## Abstract

Pauson–Khand reaction was achieved on the sugar acetylenes having an allylic ether substituent at the two-position of the tetrahydropyran ring to provide tri-cyclic products with high stereospecificity. This is the first example of Pauson–Khand reactions with compounds on carbohydrate or tetrahydropyranose ring, and the scope is described. © 1999 Elsevier Science S.A. All rights reserved.

**Keywords:** Pauson–Khand reaction; Sugar acetylene; biscobaltoctacarbonyl

Among synthetic reactions involving organometals as catalysts, the Pauson–Khand reaction has occupied a leading position for synthesis of cyclopentanone or cyclopentenone derivatives. Most of those examples are found in the precursor compounds having no stereogenic centers. We became interested in performing Pauson–Khand reaction, a  $[2 + 2 + 1]$ cyclization with an acetylene and an olefin attached to a pyranose ring at the adjacent position, in order to obtain optically active products that are potentially useful for natural product synthesis. We have recently established a new method to synthesize the sugar acetylenes for use as reagents [1]. The current study of the Pauson–Khand [2] reaction on the sugar acetylene **1** should give the tricyclic products **2** as shown in Eq. (1). These products would also provide some additional aspects for the sugar acetylenes as synthetic intermediates.



The general precursor sugar acetylenes (**1**) are readily available by *C*-glycosidation of glycals with silylacetylenes in acidic media [1,3]. When 2-acetoxy-glucal **3**, for example, is treated with bis(trimethylsilyl)acetylene in the presence of tin tetrachloride, and then with sodium borohydride in the presence of cerium(III) chloride, the alkynylated product **4** is obtained [4]. This acetylene alcohol (**4**) was then converted into the corresponding allyl ether (**6**) in two steps via the corresponding carbonate (**5**) in the presence of palladium as catalyst [5]. In this case about 23% of the allylic alcohol (**4**) was recovered. The product **6** was obtained with retention of configuration, which suggested the  $\pi$ -allyl complex of palladium and the carbonate **5** took place at the non-cyclic allyl ether group rather than the cyclic one. The 1,2-*cis*-en-yne compound (**6**) was first converted to the corresponding biscobalthexacarbonyl complex (**7**), and was then subjected to one of several Pauson–Khand conditions. Treatment of **7** with six equivalents of *N*-methylmorpholine *N*-oxide (NMO) at room temperature under nitrogen [6] yielded the tri-cyclic product **8** as crystals (m.p. 86.3°,  $[\alpha]_D = +195^\circ$ ) in 98% yield. To the best of our knowledge this is the first example of successful Pauson–Khand reaction related to a sugar-based acetylene [7] (Scheme 1).

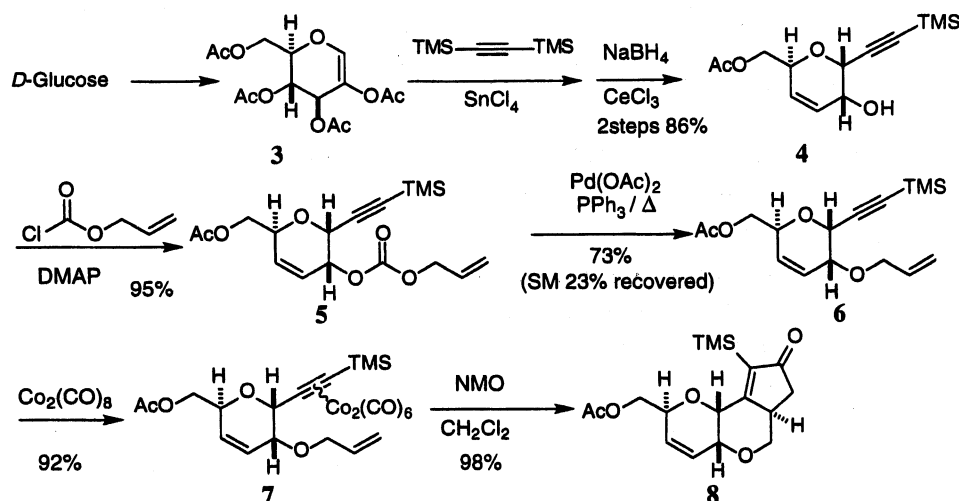
Such a successful Pauson–Khand reaction with the 1,2-*cis*-en-yne precursor **7** prompted us to examine a 1,2-*trans*-ene-yne precursor **9**, which was obtainable

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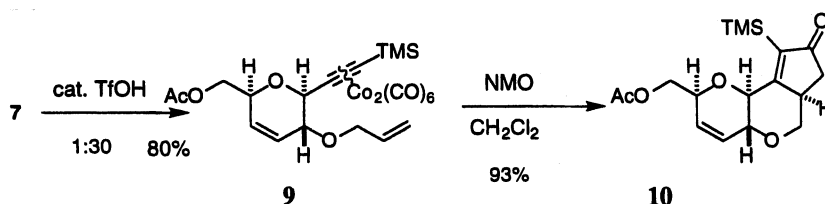
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from **7** via acidic epimerization. Treatment with trifluoromethanesulfonic acid (0.1 M solution of TfOH) at room temperature afforded **9** (containing a small amount of **7** as a mixture in a ratio of 30:1). This product was subjected to NMO in dichloromethane under the same conditions as above to provide **10** (m.p. 119°,  $[\alpha]_D = +222.1^\circ$ ) (Scheme 2).

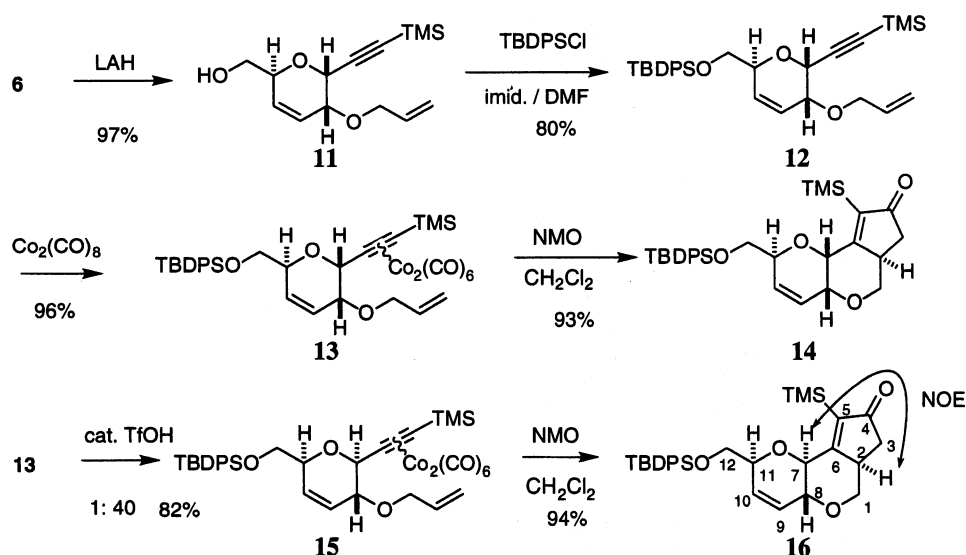
Similar examples were achieved with 1,2-*cis*- and 1,2-*trans*-en-yne precursors of acetylene bis-cobalthexa-carbonyl complex having *tert*-butyldiphenylsilyl protection at the six-position as **13** and **15**, which were prepared from the acetate **6**, respectively as shown in Scheme 3. In this case the  $\alpha$ - $\beta$  isomerization took place in 40:1 ratio under the same acidic conditions as the above



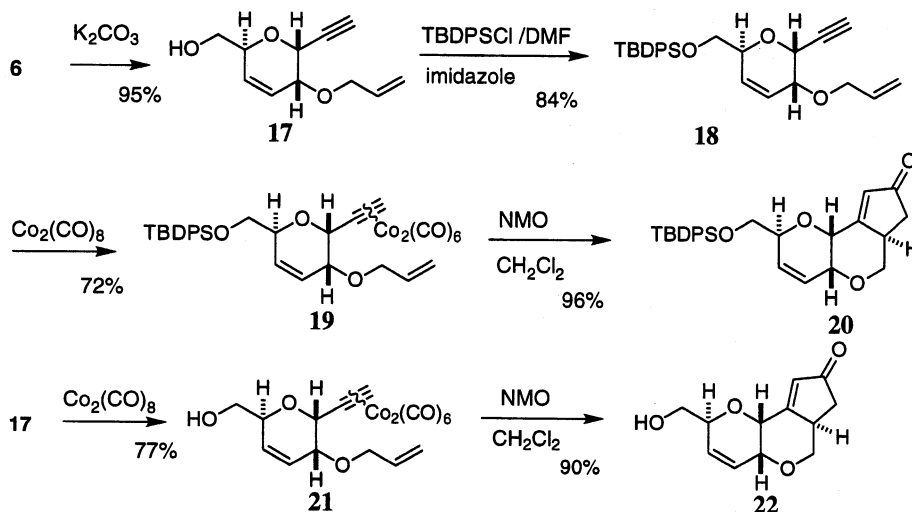
Scheme 1.



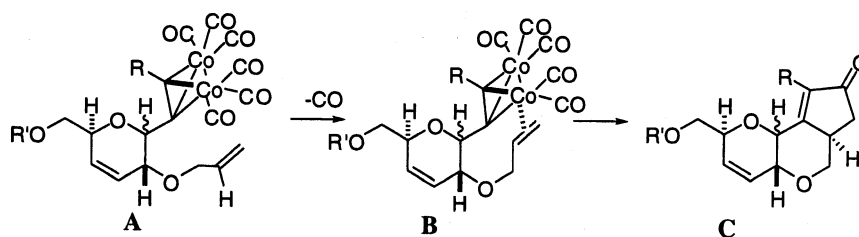
Scheme 2.



Scheme 3.



Scheme 4.



Scheme 5.

case. The P–K reaction to both **13** and **15** again proceeded with NMO to produce the corresponding tricyclic compounds **14** (colorless oil,  $[\alpha]_{\text{D}} = +78.8^\circ$ ) and **16** (m.p.  $134.4^\circ$ ,  $[\alpha]_{\text{D}} = +115.7^\circ$ ) in high yields, respectively. Among these P–K tricyclic products the last compound (**16**) from *trans*-ene-yne precursor (**15**) clearly indicated an NOE between 2-H and 7-H so that the stereochemistry of the 2-H should be  $\alpha$ . Consequently the stereochemistry of **14**, which showed no NOE, should be assigned as 2-H being  $\alpha$  as well.

The third examples are the cases of terminal acetylene (without trimethylsilyl group). Two 1,2-*cis*-precursors (**19** and **21**) were prepared from **6** by hydrolysis of the trimethylsilyl and acetyl groups with dipotassium carbonate (Scheme 4). In this case the  $\alpha$ – $\beta$  epimerization did not take place due to insufficient steric congestion without the trimethylsilyl group, so the P–K reaction was examined only with the 1,2-*cis*-en-yne precursors. Both the TBDPS derivative **19** and the free alcohol **21** afforded the tricyclic P–K products **20** (as oil,  $[\alpha]_{\text{D}} = 82.4^\circ$ ) and **22** (m.p.  $168.3^\circ$ ,  $[\alpha]_{\text{D}} = 281.1^\circ$ ) in 96 and 90% yield, respectively.

The stereochemistry of the products was assigned from NMR; thus, all of the P–K products from 1,2-*cis*-precursors showed no NOE between 2-H and 7-H. Only compound **16** indicated a clear NOE. (Compound

**10**, in fact, showed an NOE with 2-H, but the 7-H overlapped with other proton signals.) The stereochemical process might be determined by approach of the initial coordination of  $\pi$ -electrons on the side chain to exchange the carbon monoxide ligands to one of the cobalt atoms. Scheme 5 shows one possible process of the P–K reaction [8]; thus, the side chain olefin and one of the cobalt atoms in the starting material **A** approach each other, intramolecular ligand exchanges **A** to **B**. In this case, the terminal olefin is predestined to approach to the cobalt complex such that the subsequent [2+2] cycloaddition takes place through a minimum-energy surface. Fig. 1 illustrates the intermediates from 1,2-*trans* materials explaining the stereochemical course, so that 2-H directed to the  $\alpha$  orientation, where only one of the two *apical* carbonyl groups can exchange with olefin. Fig. 2 illustrates the 1,2-*cis* cases, where the tetrahydropyran ring takes an inverted conformation, but the approach between the two reaction cite should

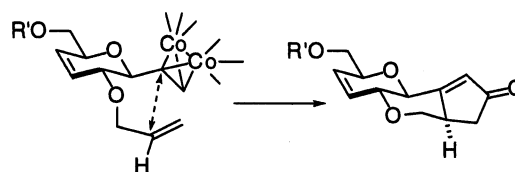


Fig. 1.

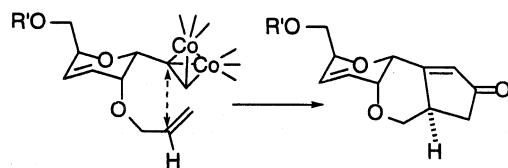


Fig. 2.

happen in a similar manner to the *trans* case. This explanation would lead to the high stereoselective product having 2-H in alpha orientation (Scheme 5).

Additional examples of this reaction including more detailed data will be described in full elsewhere.

### Acknowledgements

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