SYNLETT Spotlight 294

This feature focuses on a reagent chosen by a postgraduate, highlighting the uses and preparation of the reagent in current research

Bestmann–Ohira Reagent: A Versatile Reagent in Organic Synthesis

Compiled by Umesh Daga Patil

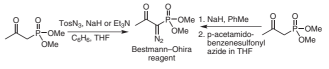
Umesh D. Patil was born in Pimpri (Mondhale), Jalgaon, India, in 1984. He pursued his B.Sc. in chemistry (2005) from Kisan Mahavidyalaya Parola, affiliated to North Maharashtra University (NMU), Jalgaon, India, and his M.Sc. in Organic Chemistry (2007) at the Department of Organic Chemistry, NMU. At present he is working towards his Ph.D. at the same department under the guidance of Dr. Sidhanath V. Bhosale. His research interests include the synthesis of biologically active heterocyclic compounds using ring transformations.

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Introduction

Bestmann–Ohira reagent [(1-diazo-2-oxopropyl)phosphonate] can be prepared by the reaction of dimethyl-2-oxopropylphosphonate, $TosN_3^{1a}$ or *p*-acetamidobenzene-sulfonyl azide,^{1b} NaH, *t*-BuOK or Et₃N in benzene and THF. An alternative is the preparation using polymer-supported sulfonyl azide and *t*-BuOK in methylenchloride.^{1c} The Bestmann–Ohiro reagent is widely used in the con-

version of primary alcohols, aldehydes, ketones, and amides into alkynes. Recently, it was employed in the synthesis of pyrazoles as well as 1,3-oxazoles.



Scheme 1

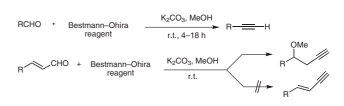
Abstracts

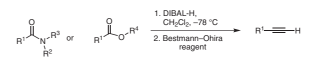
(A) The use of the Bestmann–Ohira reagent is an alternative to the Fritsch–Buttenberg–Wiechell-type rearrangement and the Corey–Fuchs procedure that allows the addition of the reagent to an aldehyde under mild reaction conditions, thus avoiding the use of a strong base under low-temperature conditions. The reaction works with alkyl and aryl as well as hindered aldehydes. In case of α , β -unsaturated aldehydes the main products were isolated as homopropargylic methyl ethers. The reaction can be easily performed under one-pot conditions.²

(B) Hamilton D. Dickson and co-workers reported that esters and amides undergo reduction to the corresponding aldehydes using DIBAL-H followed by in situ conversion into terminal alkynes utilizing the Bestmann–Ohira reagent in good to excellent yields.³ Additionally, chiral nonracemic substrates undergo this transformation with complete preservation of stereochemical integrity.³

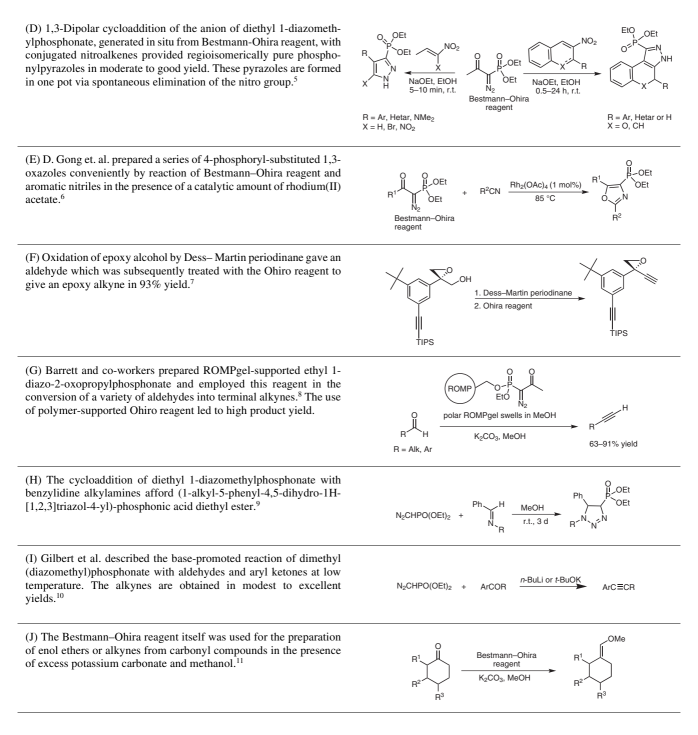
(C) E. Quesada et al. described the direct conversion of activated primary alcohols into terminal alkynes through a sequential one-pot, two-step process involving oxidation with manganese dioxide and then treatment with the Bestmann–Ohira reagent. This transformation proceeds efficiently (59–99% yield) under mild reaction conditions with a range of benzylic, heterocyclic, and propargylic alcohols.⁴

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