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

Introduction

Allyl methyl carbonate is the simplest allyl alkyl carbonate. It was first synthesised by Hermann Otto Laurens Fischer in 1929. In general, a synthesis of these carbonates is possible in high yields starting from allyl alcohols, which can be converted with dialkyl dicarbonates or alkyl chloroformates under basic conditions (e.g., BuLi, pyridine) into the corresponding allyl alkyl carbonates. Allyl carbonates are highly versatile reagents, and they can be used both for nucleo- and electrophilic reactions. Furthermore, the introduction of allyl groups is of high synthetic value, because they can be easily transformed into other functional groups. Traditional allylation reagents, like allyl bromide, require the addition of a base. An advantage of allyl alkyl carbonates is that no additional base is needed because of the cleavage of the allyl carbonate moiety into carbon dioxide and an alkoxide. This is the reason for the influence of the alkyl substituent (Me, t-Bu) on the reaction.

Abstracts

(A) Allylic Alkylation of Nucleophiles; C–C Bond Formation

Tsuji and Trost pioneered the palladium-catalysed alkylation of nucleophiles. The allylic alkylation is a versatile method to construct C–C bonds, especially products with bulky quaternary carbon centres. The allyl carbonate and palladium form an η3-allylpalladium complex, which is attacked by a nucleophile. In general, C–H acidic compounds are used, but nucleophiles like diphenylmethane are also suitable.

(B) Allylic Alkylation of Nucleophiles; C–Het Bond Formation

Various heteroatoms – with aliphatic as well as aromatic substituents – can be alkylated by allyl alkyl carbonates using catalysis by palladium or iron complexes. While the reaction of 1,1-dimethyl-allyl bromide with phenol leads to the unexpected n-product due to an SN′ reaction, the analogue carbonate leads to the desired iso product.

(C) Asymmetric Alkylation of Nucleophiles; C–C Bond Formation

The Trost asymmetric allylic alkylation, often referred to as AAA, is the enantioselective version of the Tsuji–Trost reaction. The AAA is catalysed by palladium or molybdenum. The enantioselectivity can be introduced by a chiral ligand, for example by a tetradentate Trost ligand. Furthermore, branched asymmetric alkylation products can be generated by iridium catalysis. The enantioselectivity can be introduced as described above.
(D) Reductive Allylation of Alkyl Halides
The direct allylation of alkyl halides results in a $\text{C}(\text{sp}^2)-\text{C}(\text{sp}^3)$ coupling. This catalyzed reaction proceeds via an allyl alkyl cobalt intermediate. Manganese acts as reducing agent for the allyl copper complex.\(^{17}\)

(E) Barbier-Type Allylation
Allyl ethyl carbonate can be used for the allylation of aldehydes and ketones in good yield. Furthermore, crotylation, prenylation, and intramolecular allylation are also possible with the corresponding carbonate. The Barbier-type allylation is mediated by a bimetallic system of titanium/palladium and highly accelerated by manganese dust. Initially, palladium undergoes an oxidative addition with allyl carbonate. Single electron transfer (SET) of the $\eta^1$-allylpalladium complex forms a palladium(I) intermediate. This species fragments further to an allyl radical, which can form the nucleophilic $\eta^1$-allyltitanocene(IV) complex.\(^{18}\)

(F) Alder-ene Reaction
1,4-Dienes can be formed by the Alder-ene reaction of allyl carbonates and alkynes. The $E/Z$-selectivity could be increased by the use of a permethylated cyclopentadienyl ruthenium complex.\(^{19}\)

(G) Allylation of Styrenes via a Heck-Type Reaction
The iridium-catalyzed reaction of 2-vinylanilines and allyl carbonate leads to $Z,E$-dienes. This method is a cis-selective supplement to the Heck reaction, which affords the trans products. The authors discuss an amine-assisted iridium-catalysed vinyl C–H bond activation to form the reactive intermediate.\(^{20}\)

(H) Direct C–H Allylation of Arenes
The allylation of arenes is catalysed by a permethylated cyclopentadienyl ruthenium complex. The reaction proceeds via C–H activation and is directed by $N,N$-diisopropylacetamide.\(^{21}\)

References