Simple Synthetic Method of Allyl- and Vinyl-Epoxides by Allylation of Carbonyl Groups with Allylic Tins Catalyzed by PbI₂-HMPA

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Allyl epoxides were prepared by the chemoseletive allylation at the carbonyl groups of α -bromo ketones with allylic tin, where PbI₂-HMPA effected as a chemoselective catalyst. Moreover, vinyl epoxides were obtained in one pot procedure by the PbI₂-HMPA catalyzed reaction of γ -chloroallyltri-n-butyltin with aldehydes. Because of the mildness of the catalyst, the formed epoxide functionalities were not decomposed.

Allylic tins are good candidates for the carbonyl-selective allylations because of its easy handling, and moderate reactivity. 1 However, allyltri-n-butyltin generally requires strong Lewis acids such as SnCl₄, and BF₃·OEt to perform effective allylations of carbonyl compounds.² Recently, milder activators such as Bu₂SnCl₂³ and InCl₃⁴ have been developed. However, these conventional activators requires stoichiometric amounts and can not be used yet for the bifunctional carbonyl compounds such as epoxy ketones because of the further transformation and decomposition of the products.⁵ Based on these background, we have reported PbI₂-HMPA as a novel catalyst for the allylation of α,β-epoxy ketones.6 The characteristic feature of PbI₂-HMPA is its mild and neutral catalytic activity. Namely, highly chemoselective reaction toward the carbonyls was established without decomposing epoxides. Beside the above competitive reaction, PbI2-HMPA would be a good catalyst to generate labile epoxides. We present here the PbI₂-HMPA catalyzed carbonylallylation in α -halo ketones without affecting reactive halides to achieve one-pot preparation of allyl epoxides. Moreover, we could found a simple and effective method to prepare vinyl

Table 1. Synthesis of allyl epoxides 3 from 1a and \mathbf{a} helesayles \mathbf{a}^a

$$\alpha$$
-halocarbonyls $\mathbf{2}^{a}$
 $Bu_{3}Sn$
 $\mathbf{1a}$
 O
 $\mathbf{2}$
 R^{2}
 Br
 R^{2}
 $R^{$

Entry	R ¹	R ²		Conditions	Product 3	Yi	eld /%
1	Ph	Н	2a	40°C, 24 h	R ¹	3a	96
2	<i>p</i> -Tol	Н	2b	n	/-\Z	3b	66
3	p-CIC ₆ H ₄	Н	2c	"	″	3с	67
4	p-NO ₂ C ₆ H	I ₄ H	2d	rt, 24 h		3d	77
5	Me	Н	2e	40°C, 24 h		3е	43
6	н	C ₅ H ₁	2f	63°C, 24 h	N R ²	2 3f	63 ^b

 $^{^{\}rm a}1$ 1 mmol, **2** 1 mmol, PbI₂0.1 mmol, HMPA 0.2 mmol, THF 1 mL, $40^{\rm o}$ C, 24 h, $^{\rm b}$ cis:trans= 19:81.

epoxides by the reaction using y-chloroallyltin.

Initially, we prepared allyl epoxides by the PbI₂-HMPA catalyzed allylation of bifunctional substrates such as α-bromo ketones 2 (Table 1). The reaction of 1a with α -bromoacetophenone 2aproceeded in good yield (entry 1).7 Allyl epoxide 3a was formed by the cyclization of the adduct I. Namely, the allylation took place chemoselectively at the carbonyl group without affecting the bromide moiety. Under the same conditions, aryl, allylsubstituted epoxides 3b-3d were obtained selectively (entries 2-Epoxides were formed in one pot reaction. bromohydrines derived from I were not isolated. Aliphatic ketone 2e was also reactive to give 3e (entry 5). In the case of α -halo aldehyde **2f**, internal epoxide **3f** was obtained (entry 6). The carbonyl-selectivity of PbI₂-HMPA catalyzed allylation is sharp contrast to the reaction under radical conditions where the allylation at halide moiety of 2 takes place. 8 As to the allylation to carbonyls, it has been carried out by using palladium catalyst. 8,9 PbI₂-HMPA was employed as an effective catalyst in place of expensive transition metals.

The reaction of γ -halo allyltin **1c** with aldehydes is suitable for the preparation of epoxides because remaining halogen functionality played a role for the cyclization. For the purpose, it is necessary that the C-C bond formation must take place at the halogen bonded carbon of **1c** to form halohydrin derivatives. Prior to use this tin reagent, we examined the basic reactivity of PbI₂-HMPA catalyzed allylation. (E)-Crotyl tributyltin **1b** afforded S_E2' allylation with PhCHO **4a** to give α -methyl homoallyl alcohol **5a** with high syn-selectivity. (eq. 1). This syn-selectivity suggests that the allylation proceeded via acyclic antiperiplanar transition state. ^{2b} In the reaction of allyltributyltin **1a** with unsaturated aldehyde **4b**, regioselective 1, 2-addition took place to give **5b** (eq. 2)

These high regio- and stereoselective reactions could be applied to the preparation of functionalized epoxides using γ -chloro allylic tin **1c** (Table 2).10 For example, the reaction with **4a** gave cis-vinyl phenyl epoxide **6a** selectively (entry 1). Similar to the case of crotyltin **1b**, the reaction proceeded by S_E2 reaction.

Subsequently, the formed chlorohydrin intermediate II resulted in the immediate cyclization to give **6a**. The formed epoxide ring was not decomposed by the catalyst.

Table 2. Synthesis of vinyl epoxides **6** from γ -chloro allyltin **1c** and aldehydes **4**^a

Entry	Entry Aldehyde 4		Product 6			Yield /%	
1 2 3 4	Ph p-CIC ₆ H ₄ p-NO ₂ C ₆ H ₄ p-CNC ₆ H ₄		R	. /	6a 6b 2 6c 6d	58 65 67 69	
5	n-C ₆ H ₁₃	4f	R	R= n-C ₆ H ₁₃	6e	27	
•	c-Hexyl PhCH=CCI	4g 4h	Ph.	R ² = CI	6g	14 67	
8	PhCH=CBr	4i	~ ~	→ Br	6h	59	

^a 1c 1 mmol, 4 1 mmol, PbI_2 0.1 mmol, HMPA 0.2 mmol, THF 1mL, $63^{\circ}C$, 24 h,

Other aromatic aldehydes, **4c-4e**, were also reactive to give aryl vinyl epoxides **6b-6d** (entries 2-4). These reactions proceeded with high stereoselectivity because epoxides obtained were *cis*-isomers. Aliphatic aldehydes, **4f** and **4g**, also gave vinyl epoxides, **6e** and **6f**, although these substrates were less reactive compared with aromatic ones (entries 5 and 6). Divinyl substituted epoxides, **6g** and **6h**, were obtained selectively by the initial 1, 2-addition of unsaturated aldehydes, **4h** and **4i** (entries 7 and 8).

From the results of Table 1 and 2, PbI₂-HMPA act as a effective catalyst for the allylation of carbonyl groups without decomposition of the generated epoxide functionalities. Probably, only carbonyl groups were activated because of the relatively low activity of the catalyst. PbI₂-HMPA was the best choice in the case to form labile functionalities such as epoxide rings. The high chemoselective reaction enabled the preparation of diepoxides, **8a** and **8b**, by the reaction between **1c** and epoxy ketones **7** (eq. 3).

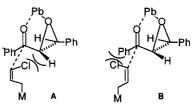
In the products **8**, the relationship between C2 and C3 was stereoselective because the chelation controlled reaction proceeded,6 however the selectivity between C3 and C4 was not so good compared with the reaction of aldehydes.¹¹

Although the correct mechanism how lead iodide complex works is not clear as yet, it seems that the catalyst acts as a mild Lewis acid or the tansmetallation occurred with allyltins.

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References and Notes

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Scheme 1