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ARTICLE TYPE

Iron-catalysed cross-coupling of halohydrins with aryl aluminium reagents: A protecting-group-free strategy attaining remarkable rate enhancement and diastereoinduction

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Non-protected halohydrins are cross-coupled with aryl aluminium reagents to produce aryl alkanols in the presence of the iron-bisphosphine catalysts. Remarkable reaction rate 10 enhancement and diastereoinduction are realized by the *in situ* generated aluminium allegiidae offering a new offering a set of the set of

- *situ* generated aluminium alkoxides, offering a new method for the reactivity and selectivity control of the iron-catalysed cross-coupling reaction.
- The directing effect of non-protected hydroxyl groups (called *neighbouring group participation* when the directing group is near the reaction centre) is recognised as a classical, yet powerful chemical tool in organic synthesis for controlling stereo-, regio-, and chemoselectivities as well as reaction rate.¹ This synthetic control may also be expected to be operative in cutting-edge
- ²⁰ cross-coupling technology,² but there have been very few reports on attempts to actively implement it in designing such reactions.³ In addition to the paucity of systematic research efforts, the increasing interest in protecting-group-free syntheses⁴ prompted us to investigate the cross-coupling reactions of protected/non-
- ²⁵ protected halohydrins to eventually find the novel reactivity of organoaluminium reagents for iron-catalysed cross-coupling reactions. Herein, we report a new cross-coupling reaction of *non-activated* alkyl chlorides and aryl aluminium reagents, in which the free hydroxyl group, or more precisely, *in situ* ³⁰ generated aluminium-alkoxide, facilitated the reaction and
- enhanced the diastereoselectivity.

We have reported previously that the cross-coupling reactions of alkyl halides with various organometallic reagents proceed efficiently in the presence of the iron-bisphosphine catalysts ³⁵ (FeCl₂-SciOPPs, Fig. 1). ⁵,^{‡a} However, we did not observe

sufficient reactivity of the catalyst when a free hydroxyl group was present in the coupling substrates or when a free alcohol substrate was added to the reaction mixture. We assumed that catalyst poisoning resulted from the formation of inert iron-⁴⁰ alkoxide species.^{5a}



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In contrast to the previous observations, we found unexpectedly the iron-catalysed cross-coupling reaction of a non-⁴⁵ protected chlorohydrin proceeded readily when using aryl aluminate as the nucleophile. It should be noted that primary alkyl chlorides usually show low reactivity in iron-catalysed cross-coupling reactions (regardless of the presence of a free hydroxyl group), and require certain elaborate catalysts.⁶ Thus, ⁵⁰ we first reinvestigated the iron-bisphosphine-catalysed crosscoupling of various phenyl metal nucleophiles by using 6-chloro-1-hexanol **3** as a model substrate, in order to confirm the unexpected unique reactivity of aryl aluminates (Scheme 1 and Table 1).

	phenylmetal reagent (Ph_nM ; ≥1 equiv)	cat. 2 (5 mol %)	
CI (~)4 OR	THE	THE 80 °C 12 h	PII (*/4 UR
3a: R = H	0 °C to rt. 1 h	then H ₃ O ⁺	4a : R = H
3b: R = Sit-BuMe ₂	,	5 - 5 -	4b : R = Si <i>t</i> -BuMe ₂

Scheme 1. Iron-catalysed cross coupling reaction of protected or nonprotected halohydrins with various phenyl metal reagents

As shown in entry 1, PhMgBr gave the desired cross-coupling product 4a in only 4% yield, along with the formation of 5-60 hexen-1-ol and 1-hexanol in ca. 30% combined yield. The reaction with diphenylzinc, Ph₂Zn·2MgCl₂ (5b),⁷ was sluggish, giving 4a in 6% yield (entry 2). Triphenylzincate (Ph₃Zn·MgBr, $(5c)^7$ gave the product in a higher yield than the neutral diphenylzinc; however, the yield and selectivity of the reaction 65 were both low (entry 3). When diphenylborate (Ph₂B(pin)Li, 5d)^{5a} was used, no desired product was obtained (entry 4). While neutral phenyl aluminium $(Ph_3Al \cdot 3MgCl_2, 5e)^8$ resulted in almost complete recovery of halohydrin 3a, the reaction with phenyl aluminate (Ph₄Al·MgCl, **5f**)⁸ proceeded smoothly and selectively 70 to give the desired 4a in 91% yield (entries 5 and 6).^{‡b,c} We suspected that the efficient reaction might be explained by the formation of a heteroleptic aluminate species (e.g., Ph₃AlOR·MgCl), which generates an anionic iron (ferrate) or Fe/Al mixed cluster species that exhibits the unprecedented 75 reactivity (see mechanistic discussion).⁹ The reaction of the protected chlorohydrin 3b with phenyl aluminate, thus, gave a 1:1 mixture of the coupling product and the alkane by-product in only 40% combined yield (entry 7), and the results of the reactions using the other phenyl metal reagents were almost the same as ⁸⁰ those reported previously.^{6, ‡d}

 Table 1. Reactivity differences between various phenyl metal reagents and effect of protection of hydroxyl group on the reactivity

3a or	3b — the	$\begin{array}{l} \mathbf{Ph}_{n}\mathbf{M} \\ \underline{\text{cat. 2}} \\ \text{en } \mathbf{H}_{3}\mathbf{O}^{+} \end{array} \qquad \mathbf{4a \text{ or } 4b } + \\ \end{array}$	alkene (R = F	OR + H or Si <i>t</i> -I	alka BuMe ₂))3 OR Ine
entry ^a substrate Ph_nM (equiv)			yield (%) ^b			- RSM (%) ^b
			4a or 4b ^c	alkene	alkane	
1	3a	PhMgBr (5a) (2.0)	4	12	17	63
2	3a	Ph ₂ Zn·2 MgCl ₂ (5b) (1.0)	6	0	0	89
3	3a	Ph ₃ Zn·MgBr (5c) (1.0)	13	23	20	30
4 ^{<i>d</i>}	3a	Ph ₂ B(pin)·Li (5d) (1.0)	0	9	0	91
5	3a	Ph ₃ Al·3 MgCl ₂ (5e) (1.0)	0	0	0	>99
6 ^e	3a	Ph₄Al·MgCl (5f) (1.0)	91	6	0	0
7	3b	Ph ₄ Al·MgCl (5f) (1.0)	20	1	20	52

^{*a*}Reactions were carried out at 80 °C for 12 h on 0.5 mmol scale. ^{*b*}The 5 yields were determined by ¹H NMR analysis using 1,1,2,2-

tetrachloroethane as an internal standard and confirmed by GLC analysis using undecane as an internal standard. ^cThe cross-coupling products **4a** and **4b** were obtained in entries 1–6 and 7, respectively. ^d20 mol% MgBr₂ was added as a co-catalyst. ^cThe reaction almost stopped for 12 h. ^fThe reaction of phenylaluminates prepared by transmetalation from AlCl₃ and PhMgBr gave the same result as that prepared from PhMgCl.

For the further study of the influence of alkoxide in the reaction, 1-hexanol was added to the iron-catalysed cross-coupling reaction of the protected halohydrin **3b** with ¹⁵ phenylaluminate **5f** (Scheme 2). Although the reactions in the absence of alcohol gave the cross-coupling product in low yield (Table 1, entry 7, and Scheme 2), stoichiometric or even 20 mol% 1-hexanol dramatically improved the reaction to give the desired product in high yield. The *in situ* generated aluminium alkoxide

²⁰ did not cause any catalyst poisoning, but enhanced the reaction.^{3e} This observation also clearly indicates that the formation of aluminium alkoxide species is a key to the observed high catalytic activity.

3b	5f (1.0 equiv), alcohol					
	THF, 0 °C, 1 h	THF, 80 °C, 12 h	4b +	alkene	+	alkane
	alcohol = none			1%		20%
	= 1-hexanol (1.0 equiv)			0		5%
	= 1-hexanol (20 mol %)			0		12%

25 Scheme 2. Rate enhancement by in situ generated aluminium alkoxide

Fig 2 shows a plausible catalytic cycle inferred from the abovementioned results and previous reports published by us and others. In the initial step, the precatalyst complex FeCl₂(TMS-SciOPP) is transformed into intermediate **A** through ³⁰ transmetalation with an aryl aluminate (Ar₃AlOR·MgX).^{‡f} The intermediate **A**, which is reminiscent of the bis(*m*-oxo)phenyl-aluminium–phenyltitanium complex ¹⁰ and iron chloride–aluminium-tert-butoxide complexes¹¹, is proposed here as a catal-



Fig 2. Plausible catalytic cycle

-ytically active species because of its expected high reactivity towards non-activated alkyl chlorides.^{6, 12} Subsequently, homolytic cleavage of the C–Cl bond proceeds to give an alkyl radical and the ferrate intermediate **B**.^{‡g} This is followed by the

⁴⁰ recombination of the resulting elusive alkyl radical with the aryl group on intermediate **B**, which occurs in a solvent cage to give the cross-coupling product and intermediate C.⁶ Finally, intermediate **A** is regenerated by transmetalation between intermediate **C** and the aryl aluminate.

⁴⁵ We next focused on the stereoinduction by the hydroxyl group in the electrophilic coupling partner. Fu^{3a} and Yorimitsu-Oshima^{3b} reported the catalytic diastereoselective cross-coupling reactions of protected cyclic 2-halohydrins using nickel and cobalt catalysts, respectively. In addition, Knochel recently reported the ⁵⁰ iron-mediated diastereoselective cross-coupling reaction of *tert*butyldimethylsilyl-protected cyclic 2-iodohydrins.^{3c} A protected hydroxyl group near the reaction centre is thus known to induce diastereoselectivity in Ni- and Co-catalysed as well as Femediated cross-coupling reactions. However, it was unknown ⁵⁵ whether a non-protected hydroxyl group, i.e., a metal alkoxide generated *in situ*, could give rise to such stereoinduction.

We first compared the diastereoselectivities of the reactions of non-protected and acetyl-protected trans-4-cholorocyclohexanols with phenyl aluminate 5f (Table 2, entries 1 and 2). While the 60 reaction of the protected halohydrin afforded an almost 1:1 mixture of diastereomers, that of the non-protected substrate produced the trans-isomer in 94% yield with high diastereoselectivity (93/7). Although the bulkiness of the silvl protecting groups may have affected the diastereoselectivity 65 slightly, high-level diastereoinduction was not observedth (entry 3). Various aryl aluminates possessing electron-rich, electrondeficient, and sterically demanding aromatic groups could participate in the reaction, and gave the product with high diastereoselectivities (entries 4-8). The reaction of mesityl 70 aluminate gave the desired product with excellent diastereoselectivity, albeit in low yield,^{‡i} showing that the steric demand of the nucleophile also contribute to the high diastereoselectivity (entry 9). The reaction of cis-4chlorocyclohexanol also gave the trans-isomer of the cross-75 coupling product, as in the case of trans-4-cyclohexanol, suggesting that the stereochemistry at the newly formed C-C bond is controlled by that of the in situ generated alkoxide moiety in the radical recombination step (entry 10). With cis-3chlorocylohexanol, the cis-isomer was obtained as the major 80 product (entries 11 and 12). Acetylated trans-2chlorocyclopentanol gave the desired product in low yield with low diastereoselectivity, whereas trans-2-chlorocyclopentanol gave the product in good yield with high diastereoselectivity (entries 13 and 14). With trans-2-chlorocyclohexanol, the product 85 was obtained with high diastereoselectivity, but in low yield because of the side reaction that gave 13 cyclopentyl(phenyl)methanol 15). trans-4-(entry Bromocyclohexanol gave essentially the same result as that of the corresponding chloride (entry 16). Because high 90 diastereoselectivities have been observed when a bulky substituents, such as tert-butyl14 or siloxyl3a-c groups, is in the cyclic alkyl halide substrates, we consider that the observed diastereoselective induction is caused by the bulkiness of

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2 | Journal Name, [year], [vol], 00-00

35

Table 2. Diastereoselective cross-coupling of cyclic halohydrins



^aReactions were carried out at 80 °C for 12 h on 0.5 mmol scale. ^bIsolated yield. The diastereoselectivity of the product was determined by ¹H NMR ⁵ and confirmed by GLC analysis.

aluminium alkoxide: it is likely to exist in the form of aluminium alkoxide oligomer, thereby acting as a sterically demanding substituent.¹⁵

In summary, we have demonstrated the unique iron-catalysed ¹⁰ cross-coupling reaction between halohydrins and arylaluminates.

- The aluminium alkoxide generated *in situ* through deprotonation of the hydroxyl group of halohydrin by arylaluminate did not cause the expected catalyst poisoning; instead, to the contrary to the initial expectation, the reaction rate was enhanced, and high-
- ¹⁵ level diastereoselectivity was induced, thus providing a first illustration of the synthetic potential of this protective-group-free strategy in catalytic cross-coupling reactions.

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Notes and references

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³⁰ † Electronic Supplementary Information (ESI) available: [Additional data, experimental procedure, and data for new compounds]. See DOI: 10.1039/b000000x/

‡ (a) SciOPP is the abbreviation for Spin-Control-Intended *Ortho*-Phenylene bisPhosphine. (b) We interpreted that the reactivity difference

35 between 5f and 5e is derived from whether or not the reactive ferrate species A in Fig. 2 is formed with the organoaluminate species. (c) A study to find effective dummy ligands on arylaluminate is ongoing. See Supporting Information (SI). (d) The result of the reaction of phenyl metal reagents was shown in SI. (e) The yield and reaction rates were almost the ⁴⁰ same regardless of the structures of alcohols examined. See SI. (f) Neutral FeAr₂-SciOPPs, which are the reactive species in the cross-coupling of

- alkyl halides previously reported by us (ref. 5), showed poor reactivities toward primary alkyl chlorides. (g) A radical clock experiment is reported in SI. (h) The aluminum alkoxide did not improve the diastereoselectivity 45 as shown in SI. (i) The starting material was recovered (c.a. 80%).
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