

Contents lists available at ScienceDirect

Tetrahedron Letters

journal homepage: www.elsevier.com/locate/tetlet



A simple and efficient hydrogenation of benzyl alcohols to methylene compounds using triethylsilane and a palladium catalyst

Maryam Mirza-Aghayan a,*, Rabah Boukherroub b, Mahshid Rahimifard a

ARTICLE INFO

Article history: Received 24 May 2009 Revised 22 July 2009 Accepted 14 August 2009 Available online 20 August 2009

Keywords:
Hydrogenolysis
Benzyl alcohols
Palladium(II) chloride
Triethylsilane
Reduction
Methylene compounds

ABSTRACT

Hydrogenolysis of benzyl alcohols using triethylsilane (Et_3SiH) and a catalytic amount of palladium(II) chloride ($PdCl_2$) is described. The reaction takes place under mild conditions affording high yields of the corresponding methylene compounds in short reaction times.

© 2009 Elsevier Ltd. All rights reserved.

Heterogeneous and homogeneous catalytic hydrogenation reactions have found widespread applications in the reduction of a large variety of functional groups.^{1–3}

Benzyl alcohols are often used in organic synthesis, either as starting materials or as protecting groups.^{4–7} Hydrogenolysis of benzyl alcohols has been described in several reports. For example, benzyl alcohols are readily converted into tetrazolyl and benzisothiazolyl ethers which can be catalytically hydrogenolyzed into toluenes over palladium-on-charcoal in good yields using hydrogen donors.8 Hydrogen iodide, generated (and regenerated) in situ by the action of iodine upon hypophosphorus acid, was found to reduce a variety of aryl carbinols to the corresponding diphenylmethanes under mild conditions.⁹ The reduction of primary, secondary, and tertiary benzylic alcohols to their corresponding hydrocarbons was performed in a two-phase system [apolar organic solvent and 50% KOH(aq)] at 50 °C, using 5% Pd/C and H₂ at atmospheric pressure. The reaction requires the presence of aromatic halides. In the absence of the halide promoter, the reaction takes place only if a halogen is present on the aromatic ring of the alcohol. 10 The use of trialkylborons with trifluoromethanesulfonic acid was found to be an effective reducing system for the conversion of a variety of tertiary, secondary, and benzylic alcohols into the corresponding alkanes, as well as alkylated alkanes.¹¹ The Raney Ni–sulfuric acid system was successfully applied for the reduction of several aryl bromides, benzyl alcohols, benzyl ethers, and benzylamines under mild conditions. Although details of the reaction mechanism for this reduction are not clear, reductive cleavage of the substrate is believed to proceed by utilizing both the adsorbed hydrogen on the Raney Ni and the hydrogen generated by the reaction between Raney Ni and sulfuric acid.¹²

Furthermore, organosilicon reagents in the presence of small amounts of a catalyst are used for the reduction of functional groups. The combination of chlorodimethylsilane and a catalytic amount of an indium compound is effective for the deoxygenation of aryl ketones and *sec*-benzylic alcohols to give the corresponding hydrocarbons. ^{13,14} Finally, ionic reduction of alcohols and ketones with triethylsilane/trifluoroacetic acid (Et₃SiH/CF₃CO₂H) was described by Mayr. ¹⁵

We have investigated the efficiency of the $Et_3SiH/PdCl_2$ system for the hydrogenation of 1-alkenes under mild conditions, 16,17 for the transformation of alcohols into their corresponding silyl ethers and for the cleavage of triethylsilyl ethers to the parent alcohols. 18 Furthermore, the versatility of the $Et_3SiH/PdCl_2$ system was demonstrated for the selective hydrogenation of the carbon–carbon double bond of α , β -unsaturated ketones to afford the corresponding saturated ketones under mild conditions. 19 More recently, this system was used for the reduction of carbonyl groups of aromatic aldehydes and ketones into a methylene group. 20 The results indicated the formation of the corresponding alcohol as an intermedi-

^a Chemistry and Chemical Engineering Research Center of Iran (CCERCI), PO BOX 14335-186, Tehran, Iran

b Institut de Recherche Interdisciplinaire (IRI, USR 3078), et Institut d'Electronique, de Microélectronique et de Nanotechnologie (IEMN, UMR 8520), Cité Scientifique, Avenue Poincaré-B.P 60069, 59652 Villeneuve d'Asca. France

^{*} Corresponding author.

E-mail address: m.mirzaaghayan@ccerci.ac.ir (M. Mirza-Aghayan).

ate during the reduction of aromatic aldehydes. Based on the above observation, the present work was aimed at exploring the efficiency of molecular hydrogen, generated in situ by the reaction of $\rm Et_3SiH$ with EtOH in the presence of a catalytic amount of $\rm PdCl_2$, for the reduction of benzyl alcohols to the corresponding methylene compounds.

Reduction of various benzyl alcohols with Et_3SiH in the presence of palladium dichloride in ethanol occurs at room temperature to yield the corresponding methylene products in high yields (Scheme 1, Table 1).

The hydrogenation reaction requires the use of an inert atmosphere and anhydrous solvent. In a typical experiment, $PdCl_2$ (10 mol %) was added at room temperature to a stirred mixture of alcohol (1 equiv) and Et_3SiH (2 equiv) in dry ethanol (5 ml). An exothermic reaction occurs during the first 5 min and then the temperature decreases to room temperature. The resulting mixture was stirred for the time indicated in Table 1 prior to GC–MS analysis.

The reduction of the benzyl alcohols takes place quantitatively. For example, the reaction of 1 equiv of benzyl alcohol (Table 1, entry 1), 4-methoxybenzyl alcohol (Table 1, entry 2) and 1-phenylethanol (Table 1, entry 7) with Et₃SiH/PdCl₂ (2 equiv/10%) in ethanol (5 ml) for 10 min at room temperature led to the formation of toluene, 4-methylanisole and ethylbenzene, respectively, in 100% yields. Likewise, the reaction of 4-hydroxy-3-methoxybenzyl

OH
$$Ar$$

$$R$$

$$Et_{3}SiH$$

$$EtOH$$

$$R=H, CH_{3}, Et, Ph$$

$$R=H, CH_{3}, Et, Ph$$

Scheme 1.

alcohol (Table 1, entry 4), 4-tert-butyl-benzyl alcohol (Table 1, entry 6) and 1-phenyl-1-propanol (Table 1, entry 8) gave 2-methoxy-4-methylphenol, 4-tert-butyltoluene and propylbenzene in 96%, 90% and 95% yields, respectively. Increasing the reaction time to 30 min led to an increase in the yield to 95% for entry 6.

It should be noted that for certain compounds an excess of triethylsilane or longer reaction times were required to drive the reaction to completion. For example, the reaction of 2-biphenyl-4-ylpropan-2-ol (Table 1, entry 9) with Et₃SiH (alcohol/Et₃SiH: 1/ 2 or 1/4) produced, respectively, 72% and 82% yields of the corresponding product after 30 min. However, high conversion (96%) was obtained when the reaction was conducted in the presence of an excess of triethylsilane (alcohol/Et₃SiH: 1/6). Similarly, the reaction of the secondary alcohol benzhydrol (Table 1, entry 10) with 2 equiv of Et₃SiH in the presence of PdCl₂ led to the formation of diphenylmethane in 87% yield, after 30 min at room temperature. The yield was increased to 100% by increasing the reaction time to 1 h under the same conditions. A yield of 100% was obtained when the ratio of benzhydrol/Et₃SiH was increased to 1/3, after 30 min at room temperature. 1-Indanol was converted quantitatively into indane with 2 equiv of Et₃SiH after 10 min at room temperature. We also observed the reduction of conjugated alcohols such as cinnamyl alcohol to propylbenzene using 8 equiv of Et₃SiH in 94% yield after 10 min at room temperature. Finally, we examined the hydrogenolysis of heterocylic alcohol derivatives. A trace amount of 3-methylpyridine was obtained after 1 h at room temperature in ethanol when the reduction of 3-pyridinylmethanol was carried out using 2 equiv of Et₃SiH. Using an excess of triethylsilane (8 equiv) and performing the reaction at reflux gave 3methylpyridine in high yield (95%) after 4 h. On the other hand, thiophene-2-methanol was readily reduced to 2-methylthiophene in 70% yield using 6 equiv of Et₃SiH in the presence of PdCl₂, after 4 h at room temperature.

Next, we investigated the hydrogenolysis of aliphatic alcohols such as 1-heptanol and 1-decanol using the Et₃SiH/EtOH/PdCl₂ system. The reduction failed to proceed under these conditions, even in the presence of a large excess of Et₃SiH. Indeed, no reaction oc-

 $\label{eq:table 1} \textbf{Table 1} \\ \textbf{Reduction of benzyl alcohols to the corresponding methylene compounds using } Et_3SiH/PdCl_2 \text{ in ethanol} \\ \textbf{Et_3SiH/PdCl_2 in ethanol} \\ \textbf{Ethanol} \\ \textbf{Ethanol}$

Entry	Substrate	Product	Substrate/Et ₃ SiH	Time (min)	Yield ^a (%)
1	Benzyl alcohol	Toluene	1/2	10	100
2	4-Methoxybenzyl alcohol	4-Methylanisole	1/2	10	100
3	2-Methoxybenzyl alcohol	2-Methylanisole	1/2	10	98
4	4-Hydroxy-3-methoxybenzyl alcohol	2-Methoxy-4-methylphenol ²¹	1/2	10	96
5	2-Hydroxybenzyl alcohol	o-Cresol	1/2	30	90
6	4-tert-Butyl-benzyl alcohol	4-tert-Butyltoluene	1/2	10	90
7	1-Phenylethanol	Ethylbenzene	1/2	10	100
8	1-Phenyl-1-propanol	Propylbenzene	1/2	10	95
9	2-Biphenyl-4-ylpropan-2-ol	4-Isopropylbiphenyl	1/2	30	72
			1/4	30	82
			1/6	30	96
10	Benzhydrol	Diphenylmethane	1/2	30	87
	•		1/2	60	100
			1/3	30	100
11	1-Indanol	Indane	1/2	10	100
12	Cinnamyl alcohol	Propylbenzene	1/4	10	80
	•		1/6	10	86
			1/8	10	94
13	3-Pyridinylmethanol	3-Methylpyridine	1/2	60	Trace
	•	• • •	1/6	240	34
			1/6	60	85 ^b
			1/8	240	95 ^b
14	Thiophene-2-methanol	2-Methylthiophene	1/2	60	24
	•	•	1/6	60	54
			1/6	240	70

a Determined by GC-MS

^b The reaction was conducted at reflux.

Table 2 Reduction of benzyl alcohol with $Et_3SiH(1/2)$ with different palladium catalysts in the absence of a solvent at room temperature

Entry	Catalyst	Time	PhMe/BnOSiEt ₃ /BnOH ^a (%)
1	PdCl ₂ (10%)	20 min	50/50/0
2	Pd(OAc) ₂ (10%)	20 min	8/78/14
3	PdCl ₂ (1 equiv)	5 min	100/0/0
4	Pd(OAc) ₂ (1 equiv)	15 min	16/27/57
5	$Pd(OAc)_{2}^{b} (10\%)$	19 h	84/3/2 ^c

- ^a Determined by GC-MS.
- b The reaction was conducted in the presence of 1 equiv of Cl₃C-CCl₃ as halogenating agent.
 - c 11% of benzyl chloride was also obtained.

curred after 30 min at room temperature in ethanol in the presence of 6 equiv of Et_3SiH , or when using 10 equiv of Et_3SiH in the presence of $PdCl_2$ at reflux for 4 h, and only the starting material was recovered. However, when the reaction was performed for 22 h at room temperature in the presence of excess Et_3SiH (alcohol/ Et_3 -SiH: 1/10), the corresponding triethylsilyl ethers were obtained in 62% and 56% yields along with 38% and 44% yields of the starting alcohols, respectively.

We also examined the reduction of benzyl alcohol by Et₃SiH in the presence of catalytic amounts of PdCl₂ or Pd(OAc)₂ in the absence of solvent (ethanol). The reaction of benzyl alcohol with Et₃SiH (1/2) in the presence of 10% PdCl₂ gave 50% of toluene and 50% of the corresponding triethylsilyl ether after 20 min at room temperature (Table 2, entry 1). Under the same conditions, Pd(OAc)₂ as the catalyst afforded only an 8% yield of toluene, 78% of the corresponding triethylsilyl ether, and 14% of the starting alcohol (Table 2, entry 2). On the other hand, the use of 1 equiv of PdCl₂ in the absence of ethanol yielded quantitatively, toluene, after only 5 min (Table 2, entry 3). In contrast, only partial reduction of benzyl alcohol was observed when the reaction was performed in the presence of 1 equiv of Pd(OAc)₂ (Table 2, entry 4). These results clearly indicate the influence of the chloride ions (or Et₃SiCl) on the reaction mechanism. Thus we investigated the reduction of benzyl alcohol with Pd(OAc)₂ as the catalyst in the presence of a halogenating agent such as Cl₃C-CCl₃. The reaction of benzyl alcohol with Et₃SiH (1/2) in the presence of 10% Pd(OAc)₂ as catalyst and 1 equiv of Cl₃C-CCl₃ yielded 84% of toluene, 3% of benzyloxytriethylsilane, 11% of benzyl chloride, and 2% of the starting alcohol under solvent-free conditions after 19 h at room temperature (Table 2, entry 5). We also performed the reduction of benzyl chloride using PdCl2 as the catalyst in the presence of 2 equiv of Et₃SiH and the reduction to toluene was complete after 5 min at room temperature.

The results suggest that chloride ions might play a role in the reduction mechanism. This is corroborated by previous reported work on the reduction of alcohols into their corresponding halides and alkanes using the Et₃SiH/PdCl₂ system in the presence of a halogenating agent.²² Furthermore, addition of a halogenating agent such as Cl₃C–CCl₃ to the reaction mixture led to an increase in the yield for the reduction of benzyl alcohol by Et₃SiH in the presence of Pd(OAc)₂. However, in the present work, the reduction of

benzylic alcohols does not require the presence of a halogenating agent. The only potential source of halogen is Et₃SiCl, resulting from reduction of PdCl₂ by Et₃SiH, as previously suggested.²³ Thus it is hard to draw a final conclusion regarding the exact reaction mechanism at this stage.

In conclusion, we have developed a simple and highly efficient method for the hydrogenolysis of benzyl alcohols to the corresponding methylene compounds using Et₃SiH in ethanol in the presence of a catalytic amount of PdCl₂. The reaction is easy to carry out affording high yields of the corresponding products.

General procedure for the reduction of alcohols: To a solution of alcohol (1 mmol, 1 equiv) and triethylsilane (amount indicated in Table 1) in ethanol (5 ml) was added a catalytic amount of palladium(II) chloride (10 mol %) under an argon atmosphere. The resulting mixture was stirred for the time indicated in Table 1 prior to GC–MS analysis. The pure products in entries 1 and 7 were isolated by distillation, and those for entries 2–6 and 8–10 were isolated by column chromatography using hexane/ethyl acetate (9/1) as eluent. The products²⁴ were characterized by ¹H NMR spectroscopy and mass spectrometry.

Entry 9, Table 1, ¹H NMR (CDCl₃, 80 MHz): δ = 1.42 (d, J = 5.3 Hz, 6H, CH₃), 3.06 (m, 1H, CH), 7.34–7.77 (m, 9H, CH arom.). MS (70 eV), m/z (%): 196 (62) (M⁺), 181 (100).

References and notes

- 1. Brieger, G. B.; Nestrick, T. J. Chem. Rev. 1974, 74, 567.
- 2. Jonstone, R. A. W.; Wilby, A. H.; Entwistle, I. D. Chem. Rev. 1985, 85, 129.
- 3. Wang, G.-Z.; Bäckvall, J. E. J. Chem. Soc., Chem. Commun. 1992, 980.
- 4. Larock, R. C., Comprehensive Organic Transformations: A Guide to Functional Group Preparations, 2nd ed. 1999, Wiley.
- Green, P. G. M.; Wutts, T. W.; Protecting Groups in Organic Synthesis, 3rd ed.; Wiley: New York, 1999.
- 6. McCombie, S. W.. In *Comprehensive Organic Synthesis*; Trost, B. M., Fleming, I., Eds.; Pergamon: Oxford, 1991; Vol. 8, p 811. Chapter 4.2.
- 7. Entwistle, I. D.; Wood, W. W.. In *Comprehensive Organic Synthesis*; Trost, B. M., Fleming, I., Eds.; Pergamon: Oxford, 1991; Vol. 8, p 895. Chapter 4.7.
- 8. Araújo, N. C. P.; Brigas, A. F.; Cristiano, M. L. S.; Frija, L. M. T.; Guimarães, E. M. O.; Loureiro, R. M. S. *J. Mol. Catal. A: Chem.* **2004**, *215*, 113.
- 9. Gordon, P. E.; Fry, A. J.; Hicks, L. D. Arkivoc 2005, 393.
- 10. Marques, C. A.; Selva, M.; Tundo, P. J. Org. Chem. 1995, 60, 2430.
- 11. Olah, G. A.; Wu, A.-H.; Farooq, O. J. Org. Chem. 1991, 56, 2759.
- Okimoto, M.; Takahashi, Y.; Nagata, Y.; Satoh, M.; Sueda, S.; Yamashina, T. Bull. Chem. Soc. Jpn. 2004, 77, 1405.
- 13. Miyai, T.; Ueba, M.; Baba, A. Synlett 1999, 182.
- 14. Yasuda, M.; Onishi, Y.; Ueba, M.; Miyai, T.; Baba, A. J. Org. Chem. 2001, 66, 7741.
- 15. Mayr, H.; Dogan, B. Tetrahedron Lett. 1997, 38, 1013.
- Mirza-Aghayan, M.; Boukherroub, R.; Bolourtchian, M.; Hoseini, M. Tetrahedron Lett. 2003, 44, 4579.
- Mirza-Aghayan, M.; Boukherroub, R.; Bolourtchian, M. Appl. Organomet. Chem. 2006, 20, 214.
- Mirza-Aghayan, M.; Boukherroub, R.; Bolourtchian, M. J. Organomet. Chem. 2005, 690, 2372.
- Mirza-Aghayan, M.; Boukherroub, R.; Bolourtchian, M.; Rahimifard, M. J. Organomet. Chem. 2007, 692, 5113.
- Mirza-Aghayan, M.; Boukherroub, R.; Rahimifard, M. J. Organomet. Chem. 2008, 693, 3567.
- 21. Wang, Q.; Yang, Y.; Li, Y.; Yu, W.; Hou, Z. J. Tetrahedron 2006, 62, 6107.
- Ferreri, C.; Costantino, C.; Chatgilialoglu, C.; Boukherroub, R.; Manuel, G. J. Organomet. Chem. 1998, 554, 135.
- 23. Boukherroub, R.; Manuel, G.; Chatgilialoglu, C. Organometallics 1996, 15, 1508.
- 24. Pouchert, C. J., The Aldrich Library of NMR Spectra, 2nd ed. Vol. 1 and 2, 1983.