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Chemoselective Hydrogenation of Aromatic Nitro Compounds Using Diammonium Hydrogen Phosphite and Commercial Zinc Dust

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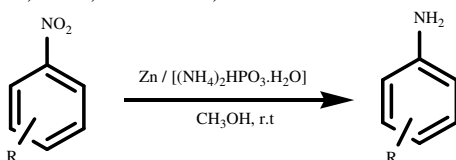
Abstract: The aromatic nitro compounds are reduced to their corresponding amines at room temperature in good yields by employing diammonium hydrogen phosphite as hydrogen donor and zinc as catalyst. The hydrogenation is fast and selective in the presence of the other sensitive functionalities such as halogens, -OH, -NH₂, -OCH₃, -CN, -COCH₃, -COOH, -COOR *etc.* It was observed that, this system is equally competitive with existing methods.

Keywords: Commercial zinc dust, Diammonium hydrogen phosphite, Aromatic nitro compounds, Catalytic transfer hydrogenation.

Introduction

Aromatic amines are an important class of compounds frequently used as key intermediates in the synthesis of pharmaceutical products, dyestuff and polymers¹. Various methods have been developed for the preparation of aromatic amines from the corresponding aromatic nitro compounds²⁻⁴. The methods employed generally are metal/acid reduction⁵, catalytic hydrogenation⁶, electrolytic reduction⁷, homogeneous catalytic transfer hydrogenation⁸, heterogeneous catalytic transfer hydrogenation⁹ *etc.*, are in practice. However, these methods have one or more limitations. In recent years, metal mediated reactions have wide scope in organic synthesis because of their simple work-up and selectivity. Several methods have been developed for the reduction of nitro compounds based on the use of metals from our laboratory as well as from other laboratories¹⁰⁻²⁰. Diammonium hydrogen phosphite was introduced by Rose *et al.*²¹ as a reducing agent in a number of reactions and also as a corrosion inhibitor for lubricating grease.

In this context, we wish to report a rapid, selective and simple reduction of substituted nitro compounds to corresponding amino derivatives by using low cost commercial zinc dust and diammonium hydrogen phosphite²¹ at room temperature in methanol medium (Scheme 1). This new system reduced with ease a wide variety of nitro compounds to corresponding amines. Many primary and secondary functional groups like halogens, carboxylic acid, phenol, ester, nitrile *etc.*, are tolerated.



R= -OH, -NH₂, -CH₃, -OCH₃, -CN, -COOH, -COCH₃, -COOR and halogens

Scheme 1

Experimental

Materials

All the nitro compounds were purchased from Aldrich Chemical Company (USA) and zinc dust from SISCO Research Laboratories Pvt. Ltd., Bombay (India). Diammonium hydrogen phosphite was prepared according to the published procedure²¹.

All the solvents were of analytical grade or were purified according to standard procedures. TLC was performed on silica gel plates obtained from Whatman Inc with the eluting systems chloroform:methanol(90:10) and chloroform:methanol(95:05). For preparative TLC, the plates were prepared from Kieselgel 60GF254 Merck, Darmstadt, and for column chromatography 60-120 mesh silica gel was used. The melting points were determined by using a Thomas-Hoover melting point apparatus and are uncorrected. IR spectra were recorded on SHIMADZU-FTIR-8300 spectrometer.

General procedure for the preparation of aromatic amines

A mixture of aromatic nitro compound (5 mmol) and commercial Zn dust (6 mmol) in methanol (5 mL) was stirred with diammonium hydrogen phosphite (10 mmol) at room temperature. The reaction was exothermic. After completion of the reaction monitored by (T.L.C), the mixture was filtered off. The organic layer was evaporated and residue was dissolved in CHCl₃ or ether (2x30 mL). The extract was washed with saturated 50% sodium chloride solution (30 mL). The organic layer was dried over anhydrous Na₂SO₄ and the solvent was evaporated under reduced pressure to obtain the desired amino compound.

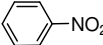
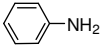

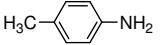
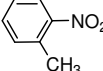
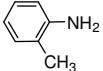
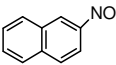
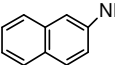
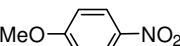
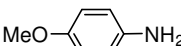

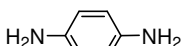
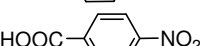
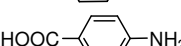
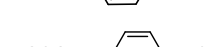
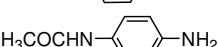
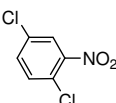
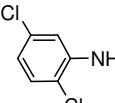
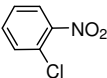
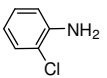
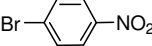
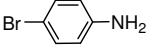

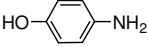
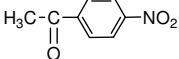
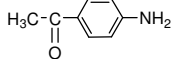
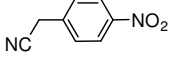
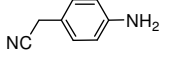
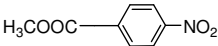
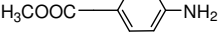
Results and Discussion

The result of this reduction of various nitroarenes was shown in Table 1. In most cases the reaction was over within 5-20 min. The usual side products of nitro reduction such as azoxy, azo, and hydrazo compounds were not observed in the final product. At the same time, it was also noteworthy that the present method was highly chemoselective and some sensitive functional groups such as -Cl, -Br and -COOC₂H₅ did not undergo any change under the reaction conditions. Moreover, many other substituted groups, such as -CH₃, -OH, -OCH₃, were intact during the reaction.

In order to test the selectivity, reduction was attempted with 4-nitroacetophenone, 4-nitrophenylacetonitrile and 4-nitrobenzoic acid gave the corresponding anilines without

affecting the other reducible groups. Even the reduction of chloronitrobenzene showed high selectivity for chloroanilines without any dehalogenation of chloronitrobenzene. It is worth to note that; all the substituted nitro compounds reduced by this system were obtained in good yields.

Table1. Reduction of nitroarenes to anilines with diammonium hydrogen phosphite and commercial zinc dust in methanol.

Entry	Nitroarenes	Time min	Amines	Yield ^a %	m.p. ^{°C} Found (Lit).
1		15		85	185(184-186) ^b
2		08		90	45(44-45) ²²
3		08		90 ^c	200(200-202) ^c
4		20		87	111-13(113) ²²
5		08		88	56-57(57) ²²
6		04		90	139-141(141) ²²
7		10		60	184-186(186) ^c
8		15		89	162-164(163) ²²
9		05		90	70(71-72) ²³
10		07		90	99-100(99) ²⁴
11		07		90	65-66(66) ²³
12		06		90	187-189(189) ²³
13		12		85	101(103-107) ²⁴
14		15		80	46(45-48) ²⁴
15		12		80	108(110-111) ²⁴

^a Isolated yields are based on single experiment and the yields were not optimized;

^b Boiling point; ^c Isolated as benzoyl derivative.

And also a controlled experiment was carried using substituted nitro compounds with diammonium hydrogen phosphite but without zinc dust, did not yield the desired product. Furthermore, an attempted reduction of a substituted nitro compounds using zinc dust in the

absence of diammonium hydrogen phosphite, did not yield the desired product, even if the reaction mixture was stirred for more than 24 hours. This clearly confirms that methanol serves as solvent and not as hydrogen source.

Conclusion

In conclusion, we report here a novel approach for the preparation of aromatic amines from the corresponding aromatic nitro compounds using diammonium hydrogen phosphite in the presence of zinc dust. This method is mild, exceedingly efficient and highly selective. The obvious advantages of proposed method over earlier methods are: (i) selective reduction of nitro compounds, in the presence of other reducible or hydrogenolysable groups, (ii) ready availability and ease of operation, (iii) rapid reduction, (iv) high yields of substituted amines, (v) avoidance of strong acid media, (vi) no equipment of pressure apparatus, (vii) cost effectivity and (viii) prevention of unwanted by products such as, hydroxylamines, nitroso, hydrazo, and azo compounds.

The catalyst is non-pyrophoric in nature and other interesting behaviors of zinc dust lies in fact that it can be recycled after simple washing with ether and dilute HCl, rendering thus process is more economic. The present method offers an economical, safe, and environmentally benign alternative to available procedures.

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