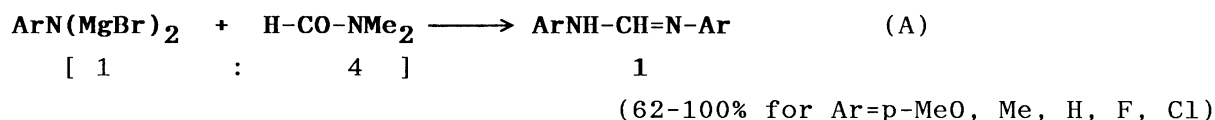


Reactions of Aryliminodimagnesium with Some N,N-Dimethylcarboxamides and Benzonitriles Affording Various Types of Amidines. Correction of Previous Results on Formamidine Formation from N,N-Dimethylformamide ¹⁾

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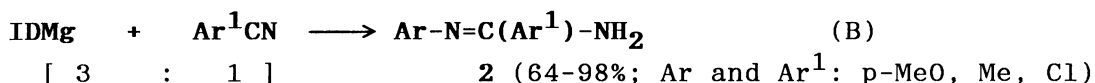
Some symmetrical and unsymmetrical form- and benzamidi-
nes were prepared by the reaction of ArN(MgBr)_2 with Ar'CN ,
 HCONMe_2 and related compounds in tetrahydrofurn.

Usefulness of aryliminodimagnesium (ArN(MgBr)_2 , IDMg), derived from anilines in tetrahydrofuran (THF), has been established as shown by the condensation ability with aromatic carbonyl and nitro compounds to afford $>\text{C}=\text{N}-\text{Ar}$ and $-\text{N}(\text{O})=\text{N}-\text{Ar}$ type products. Introduction of nitrogen functionality using IDMg is extended to formation of various type of amidines in the reactions with carboxamides and benzonitriles (Schemes A-E), and is described in this communication.

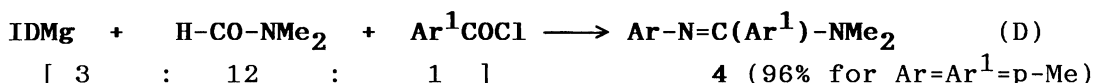
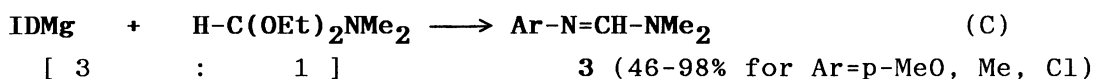


For reaction A, three (M. O., M. T., K. M.) of the present authors reported that addition of nitrobenzenes (Ar'NO_2) is required to mediate single electron transfer (SET) from IDMg to N,N-dimethylformamide (DMF: two molecules per Mg atom being used).²⁾ This description²⁾ has to be revised since reaction A was proved to proceed in later experiments without addition of Ar'NO_2 ; the erroneous description arose from failure in the detection of formamidine **1**. In the presence of p-substituted Ar'NO_2 , formation of **1** competes with that of unsymmetrical azoxy and azo compounds (see Ref. 1), whereas crowded 2,4,6- $\text{Me}_3\text{C}_6\text{H}_2\text{NO}_2$ (MesNO_2) is completely recovered and **1** is solely formed. When Ar'NO_2 is absent, the yield of **1** in the reaction with p- $\text{MeOC}_6\text{H}_4\text{-IDMg}$ (for 1 h) is affected by temperature in an interesting manner: [yield /%, temp /°C] = [58, 55], [59, 20], [100, 10], [72, 0], [29, -40], suggesting participation of equilibrium between the reactants or

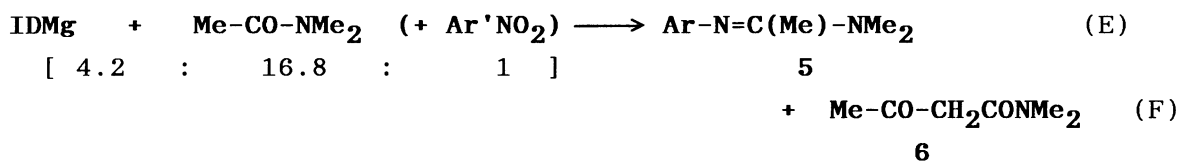
between reactant and the precursor of 1. This feature concerns that of D, and comparison of mode of A with that of E (and/or F) is given later.



Also for reaction B, an erroneous description was previously given.²⁾ The fact that IDMg reacts with p-nitrobenzonitrile only at its nitro group to afford p-cyanoazoxy- and p-cyanoazobenzenes⁵⁾ was considered to arise from general inertness of CN group. However, p-R-benzonitriles (R= MeO, Me, Cl) without strongly electron-accepting nitro group³⁾ are converted into N-monosubstituted benzamidines 2 in yields higher than that obtained by means of the conventional procedure using AlCl₃.⁶⁾



Reaction C with DMF-diethylacetal (esterification reagent^{7,8)}) leads smoothly to unsymmetrical N,N-dimethyl-N'-arylformamidine (3) in good yields, whereas triethyl orthoformate is easily converted into N-arylformamide which is hardly converted further into the symmetrical formamidine 1. Reaction A is modified into reaction D by means of stepwise addition of DMF and aroyl chloride into IDMg solution in THF: Unsymmetrical benzamidine 4 is formed in fair yield. The formyl-benzoyl exchange in D could be elucidated by the equilibrium suggested in A.



Reaction E with N,N-dimethylacetamide (DMA) leads in limited cases to unsymmetrical acetamidine 5 accompanying azoxy product, and pathway of E is greatly affected by the substituents of IDMg and Ar'NO₂ additives (see Fig. 1). When DMA and p-Cl- or p-Me-C₆H₄NO₂ were added, p-ClC₆H₄-IDMg formed 5 (25%) whereas p-MeO and p-Me reagents caused "Claisen-like" self-condensation of DMA in both the absence and presence of Ar'NO₂ to give N,N-dimethylacetoacetamide (6; reaction F).⁹⁾ The experiments of F in the absence and presence of Ar'NO₂ were carried out under conditions shown in Fig.

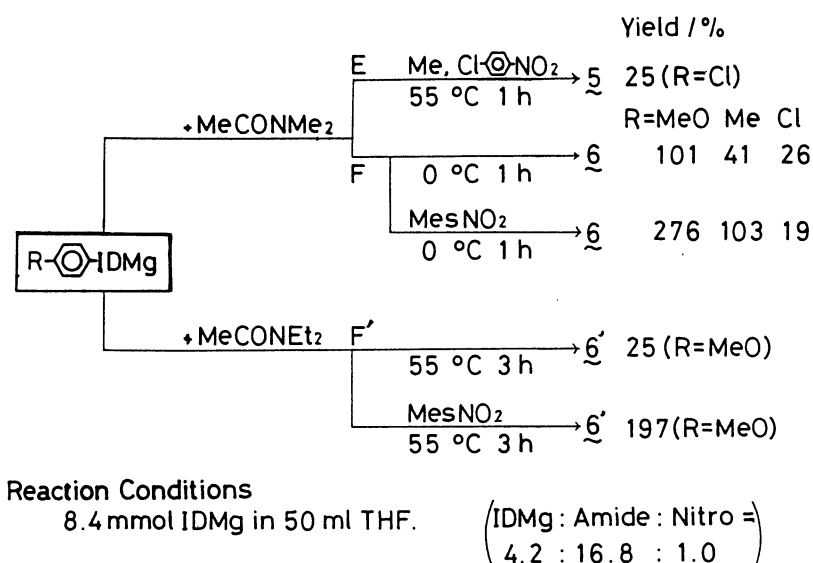


Fig.1.

1; the yields of **6**, calculated based on the amount of $\text{Ar}'\text{NO}_2$ (irrespective of its presence or absence), were compared for evaluation of its effect. Great improvement of yield of **6** (with use of p-MeO reagent) due to addition of MesNO_2 indicates that SET process, involved in F, is accelerated by the additive in a catalytic mediation manner (vide supra).

Great effect of bulkiness of not only acyl and amino moieties of carboxamides but also nitro additives on pathways of reactions A, E, and F is demonstrated by the three features. First, reaction mode of DMF having formyl moiety is distinguished from that of DMA having acetyl moiety (see Scheme A and Fig. 1). Second, the efficient reaction of DMF in reaction A is in contrast with low reactivity of N,N-diethylformamide giving no appreciable yield of **1** and also with inertness of N-phenyl- and N,N-dimethylbenzamides. Third, concerning pathway of F, N,N-diethylacetamide gave the corresponding acetoacetamide (**6'**; pathway F') in quite low yield, the yield being improved greatly by addition of crowded MesNO_2 : Ordinary p-substituted $\text{Ar}'\text{NO}_2$ undergoes the known azoxy formation (see Ref. 1).

Irrespective of comparably strong "coordinating abilities" of DMF and DMA (evaluated by Gutmann's donor numbers $(\text{DN})^{10})$ and similar conditions of A and F, the components of A are bound efficiently to give **1** while the components of F dissociate (probably after SET) to give **6**. From comparison of the processes of coordination of acyl oxygen to Mg atom of IDMg (σ -complexation) accompanied by exchange of THF ligand, small formyl and bulky acetyl moieties of A and F lead to mutual and self-condensation, respectively; the tightness of σ -complexation depends on the bulkiness of acyl group. Bulkiness effect of added $\text{Ar}'\text{NO}_2$ on E and F reflects, similarly, the degree of its access to the proximity of ligand sphere.

IDMg is a weaker donor than ArMgBr,⁴⁾ and the weak electron-accepting ability of DMF is evaluated by its negative reduction potential (E_{red} : -2.01 V)³⁾ larger than that of ordinary p-substituted Ar'NO₂ (-1.25 — -1.51 V) and comparable to that of benzophenone (-1.99 V).^{4,11)} The great structural effect of reactants on the present IDMg reactions with the weak acceptors, amides and nitriles, evokes the behavior of sterically crowded 2,3,5,6-Me₄-benzophenone: No SET takes place because σ -complexation is inhibited on the treatment with IDMg while facile SET takes place on the treatment with PhMgBr¹³⁾ (see "Less Reactive, More Selective" principle). Scope and limitations of reactions A-F, optimization of reaction conditions, and precise mechanism and characterization based on defined classification from unified structure-reactivity viewpoint (relative SET efficiency) of reactions of the magnesium reagents¹²⁾ will be reported elsewhere. At present, the fact that IDMg procedure is extended to provide novel routes to synthetically useful amidines¹⁴⁾ should be stressed.

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