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1130, 1050, 1020, 810 cm⁻¹; mass spectrum m/e (rel intensity) 164 (46, M+), 109 (42), 96 (100), 83 (58), 80 (50), 79 (98), 67 (96), 66 (75), 41 (50).

Anal. Calcd for C₁₁H₁₆O: C, 80.44; H, 9.83. Found: C, 80.6; H,

A mixture of 3.3 g (0.02 mol) of the combined VPC fractions of homoprotoadamantanone obtained above, 93 g (0.165 mol) of potassium hydroxide, 10 mL (0.207 mol) of 100% hydrazine hydrate, and 100 mL of diethylene glycol was heated under gentle reflux (ca. 160 °C) for 3 h. The reaction temperature was elevated gradually to 220 °C while water formed was distilled off, and the mixture was refluxed for an additional 2 h at that temperature. Combined reaction mixture and distillate were diluted with 100 mL of a saturated sodium chloride solution and extracted with three 50-mL portions of n-hexane. Combined hexane extracts were washed with two 50-mL portions of water and dried over anhydrous magnesium sulfate. Evaporation of the solvent and purification of the residue by sublimation under slightly diminished pressure gave 1.3 g (42% yield) of a pure sample of 4-homoprotoadamantane (11): mp 129-130 °C; IR (neat) 2910, 2850, 1460, 1230, 1100, 1020, 920, 800 cm $^{-1}$; ¹H NMR (CDCl₃) δ 1.0–2.4 (complex m); 13 C NMR (CDCl₃) δ_{C} 21.0 (t), 27.3 (t), 29.3 (d), 32.8 (t), 34.1 (d), 34.7 (t), 35.9 (d), 37.2 (t), 37.5 (t), 37.7 (d), 40.8 (t); mass spectrum m/e (rel intensity) 150 (86, M⁺), 135 (44), 107 (44), 94 (54), 93 (65), 81 (62), 80 (68), 79 (100), 67 (86), 55 (46), 41 (78).

Anal. Calcd for C₁₁H₁₈: C, 87.92; H, 12.08. Found: C, 88.1; H,

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Registry No.-4, 43000-53-9; 6, 700-56-1; 7, 281-46-9; 12, 27567-85-7; 13 isomer 1, 61770-30-7; 13 isomer 2, 61770-31-8; trifluoromethanesulfonic acid, 1493-13-6.

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Selective Formation of Biaryls via Interaction of Polynuclear Arylcopper Compounds with Copper(I) Trifluoromethanesulfonate [Copper(I) Triflate]1

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Selective formation of biaryls is observed upon interacting well-defined arylcopper cluster compounds (2- $Me_2NCH_2C_6H_4)_4Cu_4$, $(4-MeC_6H_4)_4Cu_4$, $(2-Me_2NC_6H_4)_4Cu_6Br_2$, and $(2-Me_2NC_6H_4)_4Cu_6OT_2$ with equimolar amounts of CuOTf in benzene. It is shown that complex formation of the arylcopper cluster with CuOTf precedes the C-C-coupling process. In some cases these complexes are sufficiently stable to be isolated, e.g., (2-Me₂NC₆H₄)₄-Cu₆OTf₂ (from the 2/1 reaction of 2-Me₂NC₆H₄Cu with CuOTf). Decomposition of the 2-MeC₆H₄Cu/CuOTf complex with NH₃/H₂O in the presence of oxygen affords, in addition to toluene and 2,2'-bitolyl, 2-H₂NC₆H₄Me and $2-HOC_6H_4Me$. The formation of the arylcopper–CuOTf complexes and hence biaryl formation can be inhibited by suitable ligands such as PPh3. In the absence of built-in ligands in the arylcopper compound, e.g., (4-MeC6H4)4Cu4, the reaction with CuOTf can be made catalytic in CuOTf. The selective C-C-coupling reaction has been explained in terms of intraaggregate electron-transfer processes occurring in the intermediate arylcopper-copper triflate complexes. A mechanism is proposed based on valence disproportionation inside the copper core induced by charge transfer from the core to the electron-accepting OTf anions.

The 1/1 reaction of polymeric 2-(dimethylamino)phenylcopper³ (I) with cuprous halides affords stable hexanuclear copper complexes which have R₄Cu₆Hal₂ stoichiometry.⁴ The interaction of 2-[(dimethylamino)methyl]phenylcopper⁵ (II) with cuprous halides gives rise to the formation of polymeric complexes with (R'Cu-CuHal)_n stoichiometry.⁶ Both types (2/1)

$Reagents^a$	Added ligands	Reaction time, h	$Products^b$		
			RR	RH	Others
$R = NMe_2$					
RCu/CuOTfc		48	0	0	$R_4Cu_6OTf_2$
$(2/1)$ $R_4Cu_6OTf_2/CuOTf$ $(1/2)$		24	91	9	80 <i>d</i>
RCu/CuOTf		48	85	15	d
$(1/1) \ { m RCu/CuOTf} \ (1/0.1)$		288	e	e	$\rm R_4Cu_6OTf_2/RCu$
(RCu	PPh ₃	24			
$\begin{cases} (1/4) \\ \text{CuOTf} \\ (2) \end{cases}$		72	0	0	$\mathbf{RCu} \cdot \mathbf{3CuOTf} \cdot \mathbf{6PPh}_{3} f$
RCu	PPh_3	48	0	0	No reaction
$(1/3)$ $R_4Cu_6Br_2/CuOTf$ $(1/4)$		48	97	Trace	$\mathrm{Cu}\hspace{.01in}\mathrm{Br}^d$
$\mathbf{R'} = \mathbf{CH_2NMe_2}$			R'R'	R'H	Others
R'Cu/CuOTf		48	100	N.D.g	
$egin{array}{l} (1/1) \ R'\mathbf{Cu}/\mathbf{Cu}\mathbf{OTf} \end{array}$		3 <i>h</i>	0	0	i

Table I. Interaction of 2-Me₂NCH₂C₆H₄Cu and 2-Me₂NC₆H₄Cu with (CuO₃SCF₃)₂·C₆H₆ in Benzene

^a Molar ratio of the reagents are given in parentheses. ^b Yield (%) calculated on the total amount of R (or R') in the starting organocopper. ^c Pure benzene complex of CuOTf, $(\text{CuOTf})_2 \cdot \text{C}_6 H_6$, has been used. ^d Quantitative amounts of Cu^o were formed. After the reaction the total amount of CuOTf was present in the form of a complex with RR and RH. ^e Small amount of RH (<10%) was present in solution. Heating of the reaction mixture at 80 °C for 12 h afforded 75% R as RH and 25% R as RR (for comparison: the thermal decomposition of pure I in DMF gives 60% R as RH and 40% R as RR^{se}). ^f Composition of the solid isolated (18% yield) from the reaction mixture; see Experimental Section. ^g Not detectable by NMR or by GC/MS. ^h CuOTf was added to II at −20 °C. ⁱ Yellow solid with 3R Cu·2CuOTf stoichiometry (elemental analysis; NMR in pyridine confirmed 3/2 molar ratio) was isolated.

$$4 \longrightarrow NMe_{2} + 4CuHal \longrightarrow \left[\longrightarrow NMe_{2} \right]_{4}Cu_{6}Hal_{2}$$

$$+ 2CuHal \quad (1)$$

$$- CH_{2}NMe_{2} + nCuHal$$

$$- Cu$$

$$- II$$

$$- CH_{2}NMe_{2} - Cu_{2n}Hal_{n} \quad (2)$$

of complexes are stable in the presence of an excess of cuprous halide.

During a study of the interaction of I with copper triflate $(CuOTf)^7$ we noted that depending on the I/CuOTf molar ratio either stable complexes of the type $R_4Cu_6OTf_2$ were formed (2/1 molar ratio) or decomposition of the arylcopper with formation of metallic copper occurred (1/1 molar ratio). The results of a study of the 2/1 reaction have been published elsewhere. The high stability of $R_4Cu_6OTf_2$ has been ascribed to the geometry of the 2-(dimethylamino)phenyl ligands which each are capable of spanning three copper atoms of the octahedral copper core thus stabilizing the $R_4Cu_6^{2+}$ unit. 2

A study of the products formed in the 1/1 reaction of 2-Me₂NC₆H₄Cu with CuOTf revealed that a selective coupling of the 2-(dimethylamino)phenyl ligands had taken place. While studies concerning carbon—carbon bond formation via interaction of organocopper intermediates with *divalent* copper salts have recently been reported,^{6,8} to our knowledge

$$\begin{array}{c|c} & & Me_2N \\ \hline NMe_2 & \frac{1. \ CuOTf}{2. \ NH_3/H_2O} & & \\ \hline & NMe_2 \\ 85\% & & \\ \end{array} \tag{3}$$

this is the first example of selective coupling of organic groups achieved by interaction of organocopper compounds with a monovalent copper salt. In view of the scant knowledge about the relation between reactivity and structure of organocopper intermediates we have studied this reaction in greater detail. Three other, structurally different, arylcopper compounds, $2\text{-Me}_2\text{NCH}_2\text{C}_6\text{H}_4\text{Cu}$ (II), $4\text{-Me}\text{C}_6\text{H}_4\text{Cu}^9$ (III), and $2\text{-Me}\text{-C}_6\text{H}_4\text{Cu}^9$ (IV), have been included in this study.

Results

The results of the reaction of 2-Me₂NC₆H₄Cu (I) and 2-Me₂NCH₂C₆H₄Cu (II) with CuOTf are presented in Table I. The reaction of I with CuOTf afforded well-defined R₄Cu₆OTf₂ (V)² in 80% yield. At I/CuOTf molar ratios between 2/1 and 1/1 metallic copper was formed during the reaction. NMR spectra of a 1/2 reaction mixture of R₄Cu₆OTf₂ with CuOTf in benzene-d₆ revealed that in addition to metallic copper 2,2'-bis(dimethylamino)biphenyl (VI) and N,N-dimethylaniline (VII) had formed. The chemical shift data as well as the broadening of the NMe2 proton resonances of VI and VII indicated that these products were present in solution as their complexes with CuOTf. 10 Decomposition of these complexes by workup procedures involving extraction of the reaction mixture with NH₃/H₂O solution afforded a mixture of uncomplexed VI and VII. GC/MS analysis revealed that the dimer VI was formed in 91% and the arene VII in 9%

< 1

Products^b x, x'-Bitolyl Toluene Other Reagents Workup conditions x = x' = para95d0 0 p-TolCu/CuOTfc p-TolCu/CuOTf 0 0 100d (1/0.1)x = x' = orthoo-H2NTol o-HOTol o-TolCu/CuOTf NH₃/H₂O/O₂e 20 40 10 17f (1/1)

Table II. Interaction of o- and p-Tolylcopper with CuOTf in Benzene

^a Molar ratio of the reagents given in parentheses. ^b Yield (%) calculated on the total amount of tolyl in the starting tolyl-copper. ^c (CuOTf)₂·C₆H₆ has been used. ^d Quantitative amount of Cu^o is formed. ^e Reaction time 5 h; workup in the presence of air oxygen. ^f Minor amounts of ditolyl ether (most probably the 2,2' isomer) were detected by GC/MS. ^g NH₃/H₂O solution added under N₂ atmosphere; reaction mixture stirred for 1.5 h before the final workup procedure in air was carried out.

± 20

yield (calculated on the amount of R in I).

o-TolCu/CuOTf

(1/1)

$$\begin{array}{c} 4 \ 2\text{-Me}_2 N C_6 H_4 C u + 2 C u O T f \xrightarrow{C_6 H_6} (2\text{-Me}_2 N C_6 H_4)_4 C u_6 O T f_2 \\ I & V \\ V + 2 C u O T f \xrightarrow{C_6 H_6} 4 C u^0 \end{array} \tag{4}$$

$$+ 4 \text{CuOTf} \cdot [(2-\text{Me}_2\text{NC}_6\text{H}_4)_2 \text{ and } \text{Me}_2\text{NC}_6\text{H}_5]$$
 (5)

 $NH_3/H_2O/N_2g$

The different reactivity of CuOTf toward arylcoppercopper halide compounds is clearly demonstrated by the formation of the dimer VI in 97% yield in the 1/4 reaction of (2-Me₂NC₆H₄)₄Cu₆Br₂ with CuOTf in benzene [cf. eq 1, which shows that $(2-Me_2NC_6H_4)_4Cu_6Br_2$ is stable toward CuBr⁴]. Most probably this reaction involves ligand displacement with retention of the hexanuclear cluster structure 11 followed by the irreversible interaction of (2-Me₂NC₆H₄)₄Cu₆OTf₂ with CuOTf.¹²

$$(2-Me_2NC_6H_4)_4Cu_6Br_2 + 2CuOTf \\ \iff (2-Me_2NC_6H_4)_4Cu_6OTf_2 + 2CuBr \quad (6) \\ V \\ V + 2CuOTf \rightarrow 2(2-Me_2NC_6H_4)_2 \cdot CuOTf + 4Cu^0 \quad (7)$$

The interaction of I with CuOTf in a 1/1 molar ratio afforded the dimer VI in a significantly lower yield (85%) while the arene VII was formed in 15% yield.

$$2\text{-Me}_{2}\text{NC}_{6}\text{H}_{4}\text{Cu} \xrightarrow{\text{CuOTf}} (\text{VI + VII}) \cdot \text{CuOTf}$$

$$\text{polymer}$$

$$\xrightarrow{\text{NH}_{3}/\text{H}_{2}\text{O}} \text{VI (85\%) + VII (15\%)} (8)$$

$$2\text{-Me}_{2}\text{NCH}_{2}\text{C}_{6}\text{H}_{4}\text{Cu} \xrightarrow{\text{CuOTf}} \text{VIII} \cdot \text{CuOTf}$$

$$\text{tetramer}$$

$$\xrightarrow{\text{NH}_{3}/\text{H}_{2}\text{O}} \text{VIII (100\%)} (9)$$

The corresponding 1/1 reaction of tetrameric 2-[(dimethylamino)methyl]phenylcopper (II) with CuOTf resulted in the formation of a 1/2 complex of the dimer with CuOTf. Not a trace of the arene could be detected. Hydrolytic workup of the reaction mixture afforded exclusively the dimer 2,2'-bis-[(dimethylamino)methyl]biphenyl (VIII) (100% yield). Accordingly, reaction sequence 9 provides an excellent synthetic route for the synthesis of pure VIII.13 It is worthy of note that complexes with well-defined 2-Me₂NCH₂C₆H₄Cu/CuOTf ratios are not formed. Insoluble yellow solids with varying II/CuOTf ratios were obtained starting from reaction mixtures with II/CuOTf molar ratios greater than 1.

< 1

±80

As is seen from eq 8 and 9, CuOTf is not consumed in the reaction of the arylcopper compounds. Therefore, in principle reactions of this type might proceed in a catalytic fashion. However, complex formation between CuOTf and the biaryl coupling products VI and VIII (cf. ref 10) may account for the fact that this is not observed. Indeed, the reactions of arylcopper compounds I and II with CuOTf were effectively blocked by the presence of other complexing ligands such as triphenylphosphine (see Table I). In separate experiments it was established that this was not due to the formation of arylcopper triphenylphosphine complexes. Surprisingly, both I and II do not interact with triphenylphosphine. 14 In contrast, CuOTf does form a stable 2/1 complex with triphenylphosphine. 10 It is this complex formation which blocks the interaction of CuOTf with the arylcopper compounds. 15

These observations strongly suggest that in the absence of external or of built-in ligands (-NMe2 or -CH2NMe2) only catalytic amounts of CuOTf are required to effect the coupling reaction. In order to confirm this hypothesis the interaction of o- and p-tolylcopper with catalytic amounts of CuOTf was studied. The results are compiled in Table II.

The reaction of p-tolylcopper (III) with both stoichiometric and catalytic amounts of CuOTf resulted in the quantitative formation of p,p'-bitolyl (IX).

$$4-CH_{3}C_{6}H_{4}Cu + nCuOTf$$

$$\xrightarrow{C_{6}H_{6}} \frac{1}{2}(4-CH_{3}C_{6}H_{4}) + Cu^{0} + nCuOTf \quad (10)$$
IX, 100%
$$n = 1 \text{ or } 0.1$$

NMR spectroscopy unambiguously showed that this reaction involves the intermediate formation of an insoluble 4-CH₃C₆H₄Cu/CuOTf complex. Addition of CuOTf to a solution of p-tolylcopper in benzene-d₆ causes a complete disappearance of the resonances due to the methyl and the aromatic protons when the CuOTf/III ratio reaches 1/4. NMR spectroscopy further reveals that this insoluble complex decomposes in about 2 h to give p,p'-bitolyl as the only product.

Interestingly, using o- instead of p-tolylcopper different results were obtained. Insoluble purple-colored complexes (o-CH₃C₆H₄Cu)_x (CuOTf)₄ were isolated from the reactions of o-tolylcopper (IV) with CuOTf in benzene (see Table II). Attempts to purify these complexes failed. Essentially, no reaction leading to o,o'-bitolyl (X) or toluene was observed. An NMR spectrum of a 1/1 mixture of o-tolylcopper with CuOTf showed that these complexes are stable at room temperature. After 28 h only minor amounts (<5%) of o,o'-bitolyl were present in solution. It is worthy of note that decomposition of this complex with a NH₃/H₂O solution in the presence of oxygen afforded not only toluene arising from hydrolysis of the organocopper, but also o,o'-bitolyl, 2-aminotoluene (XI), and 2-hydroxytoluene (XII). ¹⁶ As shown in Table II the solvolysis products XI and XII are only formed in the presence of oxygen. ¹⁷ Addition of NH₃/H₂O to the reaction mixture in a nitrogen atmosphere afforded toluene (80%) and o,o'-bitolyl (20%), but not XI and XII.

2-MeC₆H₄Cu + CuOTf
$$\longrightarrow$$
 [complex] $\xrightarrow{NH_3/H_2O}$ toluene
+ o , o' -bitolyl + 2-HOC₆H₄Me + 2-H₂NC₆H₄Me (11)
Discussion

Earlier we had observed that interaction of arylcopper compounds $\operatorname{Ar}_n\operatorname{Cu}_n$ with copper halides CuHal results in the formation of complexes $\operatorname{Ar}_n\operatorname{Cu}_{m+n}\operatorname{Hal}_m$ which have equal or higher stability as compared with the parent compound. However, the present study reveals that complex formation of ArCu with copper(I) salts of anions with strong electron acceptor properties gives rise to less stable complexes. This observation establishes for the first time that the nature of the counterion has a great influence on the stability of the arylcopper—copper(I) salt complex. A better understanding of this effect is important in view of the interpretation of reactions involving organocopper compounds as intermediates, e.g., the Ullmann biaryl synthesis.

The elucidation of the mode of interaction of CuOTf with arylcopper compounds²⁰ requires a discussion of the following factors: (1) the structure of the arylcopper compounds before reaction; (2) the nature of the arylcopper-copper triflate interaction including the nature of the electron-transfer processes resulting in product formation.

The Structure before Reaction. The structures of $(2-Me_2NCH_2C_6H_4)_4Cu_4^{21}$ as well as of $(2-Me_2NC_6H_4)_4Cu_6X_2$, $X = halide^4$ or triflate, in the solid and in solution are well-documented main structural features of these compounds being central tetra- or hexanuclear copper cores to which aryl groups are bound via 2e-3c bonds.

Camus and Marsich⁹ have reported that the results of molecular weight determinations of o- and p-tolylcopper were in agreement with low aggregation states (monomers or dimers), but structural details were not given.²² However, the extreme air sensitivity of o- and p-tolylcopper as well as the fact that HCCl₃²⁴ and CCl₄ had been used as solvents for the osmometric molecular weight determinations led us to redetermine the molecular weight of these compounds. Cryoscopic molecular weight determinations show that p-tolylcopper exists as a tetramer, (4-MeC₆H₄)₄Cu₄, in benzene. In contrast, o-tolylcopper exists directly after dissolution in benzene as an apparent hexanuclear species, which equilibrates in about 2 h to a tetranuclear aggregate (2-MeC₆H₄)₄Cu₄. Obviously, in solution the (2-MeC₆H₄)₄Cu₄ aggregate is thermodynamically the most stable. Other examples of tetranuclear copper cluster species are $(Me_3SiCH_2)_4Cu_4,^{25}\ (C_6F_5)_4Cu_4,^{26}$ and (2-CF₃C₆H₄)₄Cu₄.²⁶ The copper atoms in these compounds are two-coordinate by participating in two 2e-3c C-Cu interactions, whereas in arylcopper compounds containing built-in ligands, such as (2-Me₂NCH₂C₆H₄)₄Cu₄ the copper atoms become three-coordinate by an extra Cu-N coordination bond.21

2-Me₂NC₆H₄Cu is the only arylcopper compound used in this study of which the structure is not known with certainty.

The available spectroscopic data²⁷ strongly support a polymeric structure consisting of 2-(dimethylamino)phenyl groups bridging Cu atoms of one copper chain via 2e–3c Cu–C bonds and coordinating to a copper atom of a second chain via a Cu–N bond.³

The C-C Coupling. Interaction of CuOTf with organocopper compounds which have a discrete cluster structure $[(4-MeC_6H_4)_4Cu_4, (2-Me_2NC_6H_4)_4Cu_6Br_2, (2-Me_2NC_6H_4)_4Cu_6OTf_2,$ and $(2-Me_2NCH_2C_6H_4)_4Cu_4]$ results in almost quantitative formation of biaryls. The fact that arenes are formed in less than 9% yield excludes decomposition pathways involving free radicals, but instead points to the occurrence of intramolecular processes leading to pairwise release of aryl groups. It therefore would seem plausible to propose that these processes take place in an arylcoppercopper triflate precursor complex formed by extension of the copper core of the parent organocopper with one or more copper atoms of copper triflate. A representative example of such a complex is (2-Me₂NC₆H₄)₄Cu₆OTf₂. This complex, which has been isolated and characterized, has a structure consisting of an octahedral copper core to which both aryl groups and anions are bound in a well-defined way.2

$$Ar_nCu_n + mCuOTf \longrightarrow Ar_nCu_{m+n}OTf_m \longrightarrow$$

precursor complex XIII

 $\longrightarrow ArAr + 2Cu^0 + Ar_{n-2}Cu_{m+n-2}OTf_{m-2}$

The driving force in the coupling reaction is charge transfer in the precursor complex XIII from the Ar_nCu_{m+n} skeleton to the strongly electron accepting OTf groups, which reduces the electron density in the Cu_n –C region and thus the kinetic stability of the Cu_n-C bond. The occurrence of Cu_n-C bond weakening as a result of electron transfer can be concluded from the mass spectral fragmentation pattern of (2-Me₂NCH₂C₆H₄)₄Cu₄. The parent ion R₄Cu₄+· undergoes fragmentation to $R_3Cu_4^+$ (most abundant Cu-containing ion) by cleavage of a Cu₄-C bond indicating that an electron from the bridge-bond MO has been removed rather than from the Cu₄ core.²¹ This is also illustrated by the formation of biaryl and aryl halide in the reaction of (2-Me₂NCH₂C₆H₄)₄Cu₄ with CuIIX2 which involves inner-sphere-redox reaction in activated complexes of the type (2-Me₂NCH₂C₆H₄)₄Cu₄···X··· $Cu^{II}X(Cu^{II}X_2)_n.6$

In the precursor complex XIII aryl groups are at close proximity so that concerted or consecutive Cu-C bond cleavage and C-C bond formation can occur. Three different mechanisms for pairwise release of aryl groups from Cu_n clusters can be envisaged (see Scheme I): (1) two-electron

Scheme I

R
Cu
OTf
$$Cu$$
OTf
$$RR + \begin{bmatrix} Cu \\ Cu \\ Cu \end{bmatrix}$$
OTf
$$RR + \begin{bmatrix} Cu \\ Cu \\ Cu \end{bmatrix}$$
OTf
$$RR + \begin{bmatrix} Cu \\ Cu \\ Cu \end{bmatrix}$$
OTf
$$RR + \begin{bmatrix} Cu \\ Cu \\ Cu \end{bmatrix}$$
OTf

transfer from one Cu–C bond to the Cu_n cluster resulting in reduction of two Cu^I atoms in the metal core and development of a high degree of carbonium ion character at C(bridge) fol-

lowed by intraaggregate nucleophilic attack of a second Cu–C bond; (2) simultaneous one-electron transfer from two Cu–C bonds to the Cu_n cluster and concomitant coupling of the two aryl radicals or one-electron transfer followed by intraaggregate trapping of the aryl radical by a second Cu–C bond; (3) valence disproportionation inside the metal core followed by reductive elimination of R–R.

Process 3 seems to provide a rationale for the large influence of the type of the anion on the occurrence of coupling reactions because the anions attached to the copper core will affect the potentials of the various copper couples.^{28,29} The strong electron-accepting properties of the OTf anion favor the CuII oxidation state, whereas, for example, the electron-donating halide anions favor the Cu^I state. As regards the influence of the type of aryl group on the coupling process the kinetic stability of the Cu_n-C bonding will be optimal when the aryl nucleus is oriented about perpendicular to the Cu-Cu axis thus allowing maximum back-bonding to the aryl nucleus.²¹ This orientation is favored for steric reasons³⁰ in the o-methyl substituted aryl derivative whereas in the 2-Me₂N- and 2-Me₂NCH₂-substituted compounds the occurrence of Cu-N coordination further reduces the number of possible rotamers. The lower stability of the precursor complex containing the 2-Me₂NCH₂ grouping as compared with the o-methyl substituted complex can be ascribed to coordination of the hard nitrogen ligand to the copper core which favors valence disproportionation by stabilizing the Cu^{II} oxidation state.

Finally, selective biaryl formation can only be expected for precursor complexes $\operatorname{Ar}_n\operatorname{Cu}_{n+m}\operatorname{OTf}_m$ in which n equals 2, 4, 6, etc.³¹ In case of polymeric or oligomeric arylcopper compounds chains containing both odd and even numbers of aryl groups will be present. Accordingly, for these compounds the formation of a small amount of arene originating from decomposition of chains with n=3,5,7, etc., must be expected. Indeed, interaction of polymeric 2-Me₂NC₆H₄Cu with CuOTf yields in addition to 85% of the dimer 15% of the arene $\operatorname{Me}_2\operatorname{NC}_6H_5$.

Experimental Section

General. CuOTf- $\frac{1}{2}$ C₆H₆ was prepared according to the directions given by Salomon and Kochi. ⁷ o- and p-tolylcopper and 2-dimethylamino- and 2-dimethylaminomethyl-substituted phenylcopper compounds were prepared by published methods. ^{3,5,9} The reactions were carried out under dry, oxygen-free nitrogen. Solvents were carefully purified, dried, and distilled before use under nitrogen.

IR spectra were recorded on a Perkin-Elmer 577 grating IR spectrometer. ¹H NMR spectra were recorded on a Varian Associates HA-100 NMR spectrometer. Molecular weight determinations were carried out in benzene using a cryoscopic method. The spectra and the molecular weight data were obtained by Mrs. G. M. Bijlsma-Krüger and Mrs. T. van Montfort-Volp. GC/MS analyses were recorded on a Finnigan 3100D by Mrs. G. G. Versluis-De Haan. Elemental analyses were carried out under the supervision of Mr. W. J. Buis in the Analytical Department of this Institute.

Interaction of 2-(Dimethylamino) phenylcopper with CuOTf. Synthesis of (2-Me₂NC₆H₄)₄Cu₆OTf₂ (V). Solid CuOTf· $\frac{1}{2}$ C₆H₆ (2.45 mmol) was added at room temperature to a well-stirred suspension of 2-(dimethylamino) phenylcopper (I) in benzene (25 mL). The resulting brown-yellow colored reaction mixture was stirred for 48 h. The yellow precipitate was filtered off and extracted twice with benzene (20 mL) and with pentane (2 × 20 mL). The yellow residue (80%) was dried in vacuo. Anal. Calcd for C₁₇H₂₀Cu₃F₃O₃N₂S: C, 35.20; H, 3.48; N, 4.83; Cu, 32.86; F, 9.83. Found: C, 34.2; H, 3.5; N, 4.4; Cu, 32.0; F, 9.6. IR (OTf vibrations²) ν_4 , 1315 s, 1298 (sh), 1230 m, 1199 m; ν_1 1010 s; ν_5 632 s; ν_3 521 m. NMR (in toluene- d_8) δ (10 °C) 1.92 and 2.96 (2 s, br. 6 H, NMe₂, coalescence at room temperature to one singlet at 2.50), 6.50 (m, J = 8 Hz, H₃), and 8.84 (m, J \simeq 6 Hz, H₆). Decomposition (under N₂, 5 °C/min) started at 118 °C; explosion occurs at 123 °C.

Reactions of 2-Me₂NC₆H₄Cu, (2-Me₂NC₆H₄)₄Cu₆OTf₂, and (2-Me₂NC₆H₄)₄Cu₆Br₂ with CuOTf. A typical experiment involving the reaction of I with CuOTf in a 1/1 molar ratio is described. The

respective reaction conditions and results of the other reactions are in Table I.

Solid CuOTf-1/2C6H6 (2.36 mmol) was added at room temperature to a suspension of I (2.36 mmol) in benzene (25 mL). This mixture was stirred for 48 h. NMR spectroscopy of the solution showed two broad NMe resonances at δ 2.46 and 2.54 ppm [NMR in benzene- d_6 of (2-Me₂NC₆H₄+₂·2CuOTf, NCH₃, 2.52 ppm broad, ¹⁰ and of pure VI, NCH₃, δ 2.40 ppm³]. A 6 N NH₃/H₂O solution (25 mL) was added to the reaction mixture. The benzene layer was extracted with NH₃/H₂O solution (removal of copper). The benzene layer was extracted with 4 N HCl solution. The acidic water layer was made basic with NaOH solution and extracted with diethyl ether. The ether layer was dried over Na₂SO₄ and concentrated, affording a yellow oil. NMR spectroscopy showed this oil to be a mixture of 2,2'-bis(dimethylamino)biphenyl (VI) and N,N-dimethylaniline (VII). NMR (C₆D₆) VI, δ 2.40 (12 H, s, NCH₃), 7.52 (m, 2 H, H₃ or H₆), 7.15 (m, 2 H, H₆ or H₃ partly masked by $C_6D_{6-x}H_x$ resonances), and 6.94 (m, 4 H, H_{4.5}); VII, 2.50 (3, 6 H, NCH₃). Total recovery of R as VI and VII amounts to 98%, 85% as RR and 15% as RH.

Reaction of 2-(dimethylamino)phenylcopper with Triphenylphosphine. Solid triphenylphosphine (8.17 mmol) was added to a suspension of I (2.72 mmol) in benzene (20 mL). The resulting yellow colored suspension was stirred at room temperature for 48 h. The solid was filtered off and extracted with benzene (2 \times 10 mL). NMR and IR spectroscopy revealed that this solid consisted of pure I. The spectra were identical with those of an analytically pure sample of I (see ref 3). I was recovered in 96% yield.

Reaction of I with PPh₃ and CuOTf. A mixture of I (3.3 mmol) and PPh₃ (13.1 mmol) in benzene (25 mL) was stirred at room temperature for 24 h. Subsequently, solid CuOTf. $\frac{1}{2}$ C₆H₆ (5.68 mmol) was added and the reaction mixture stirred for another 72 h. The yellow precipitate was separated by centrifugation and extracted with benzene (3 × 25 mL). The benzene extract was concentrated, which afforded a yellow solid. Extraction of this solid with pentane (2 × 20 mL) and with ether (3 × 20 mL; removal of PPh₃) afforded an ochre solid which had 2-Me₂NC₆H₄Cu·3CuOTf·6PPh₃ stoichiometry (18% yield calculated on the starting amount of I). Decomposition occurred at 140–160 °C. Anal. Calcd for C₁₁₉H₁₀₀NO₉P₆S₃F₉Cu₄: C, 59.66; H, 4.2; N, 0.58; P, 7.76; F, 7.14; Cu, 10.61. Found: C, 59.2; H, 4.6; N, 0.6; P, 7.5; F, 7.1; Cu, 11.7. NMR (pyridine- d_5) δ 2.90 (br, NCH₃), 7.15–7.60 and 6.4–7.0 (m, br, complexed PPh₃); addition of H₂O afforded VII and PPh₃ in 1/5.7 molar ratio (elemental analysis, 1/6).

Reaction of I with CuO_2CCF_3 in DMF. Solid copper(I) trifluoroacetate¹⁹ (3.6 mmol) was added to a suspension of I (3.6 mmol) in DMF. This mixture was stirred for 96 h. Workup with NH_3 solution (vide supra) afforded a colorless solid which according to NMR spectroscopy was 2,2'-bis(dimethylamino)biphenyl and N,N-dimethylaniline in 9/1 molar ratio. The recovery of R in these products was 70%.

Interaction of 2-[(Dimethylamino)methyl]phenylcopper (II) with CuOTf. Reaction of II with CuOTf. Solid CuOTf. $\frac{1}{2}$ CeHe (1.62 mmol) was slowly added at room temperature to a solution of II (1.62 mmol) in benzene (20 mL). The color of the solution turned immediately to red upon the addition of the first amount of CuOTf. At 2/1 molar ratios a yellow precipitate was formed which upon continued addition of CuOTf dissolved. Finally, a green solution containing metallic copper was obtained. After 48 h the reaction mixture was worked up following the procedure described above for the I/CuOTf reactions. A yellow oil was isolated which according to NMR spectroscopy consisted of pure 2,2'-bis[(dimethylamino)methyl]biphenyl (VIII). Thus R was quantitatively recovered as the dimer VIII: NMR (CeHe) δ 2.01 (s, 2 H, NCH₃), 3.07 (d, 2 H) and 3.28 (d, 2 H, $J_{\rm gem} \simeq$ 13 Hz, NCH₂) (cf. ref 5). N,N-Dimethylbenzylamine was absent.

In a separate experiment the NMR spectrum of the reaction mixture after stirring for 48 h, but before hydrolysis, was recorded: NMR (C_6D_6) δ 2.30 and 2.14 (2 s, br, 12 H, NMe), 3.78 and 2.34 (2 d, br, 4 H, $J_{\rm gem}$ = 12 Hz, NCH₂), identical with the spectrum obtained by mixing VIII and CuOTf in an exact 1/2 molar ratio.

Synthesis of VIII. Crude 2-[(dimethylamino)methyl]phenylcopper (II), isolated by filtration of the reaction mixture of 2-[(dimethylamino)methyl]phenyllithium (59.7 mmol) with an equimolar amount of CuBr,⁵ was mixed with 120 g of naphthalene and subsequently heated at 165 °C for 6 h. Crude VIII was isolated by an acid/base workup procedure and purified by fractional distillation. The overall yield, calculated on the amount of the aryllithium, was 63%, bp 135–140 °C (0.1 mm), NMR spectrum vide supra.

Attempted Synthesis of II/CuOTf Complexes. A solution of II in toluene (3.29 mmol in 20 mL) was cooled to -20 °C. Under vigorous stirring solid CuOTf (1.65 mmol) was added. The resulting orangebrown suspension was stirred at -45 °C for 1 h and for another 1 h

at room temperature. The orange solution was filtered and concentrated affording a yellow solid. This solid was extracted with pentane (2 × 10 mL) and dried in vacuo. Elemental analysis pointed to the isolation of a complex which had 3(2-Me₂NCH₂C₆H₄Cu)-2CuOTf stoichiometry. Anal. Calcd for $C_{29}H_{36}N_2Cu_5O_6S_2F_2$: C, 34.19; H, 3.54; Cu, 31.21; N, 4.13; F, 11.20. Found: C, 35.2; H, 3.9; Cu, 28.3; N, 3.7; F, 10.8. This solid decomposed slowly at room temperature.

Interaction of p-Tolylcopper with CuOTf. Pure p-tolylcopper was prepared following the directions of Camus and Marsich.9 Anal. Calcd for C7H7Cu: C, 54.36; H, 4.56; Cu, 41.08. Found: C, 52.8; H, 4.6; Cu, 40.8. Mol wt (cryometry in C_6H_6) 604 (\overline{n} = 3.9) concentration independent (calcd for C₇H₇Cu, 154.7). NMR (C₆D₅N) δ 2.15 (s, 3, CH_3), 7.03 (m, 2, $J_{2,3}$ 7 Hz, H_3) and 8.09 (m, 2, H_2); (in C_6D_6) 1.96 (CH₃), 6.84 (H₃), and 7.98 (H₂).

Reaction of p-Tolylcopper with CuOTf. Solid CuOTf (3.24 mmol) was added to a solution of p-tolylcopper (3.24 mmol) in benzene (40 mL). The color of the solution turned red and a black solid precipitated. A 6 N NH₃/H₂O solution was added. The benzene layer was extracted with NH₃/H₂O solution and with H₂O and dried over MgSO₄. The NMR spectrum indicated that p,p'-bitolyl was formed in 95% yield. NMR (CCl₄) δ 2.32 (3, s, CH₃), 7.08 and 7.34 (2 d, J = 8Hz, H_{2,3}). Concentration of the benzene solution afforded white solid p,p'-bitolyl, mp 115-119 °C (lit.20 121 °C).

Interaction of CuOTf with p-Tolylcopper/2-[(Dimethylamino)methyl]phenylcopper Aggregates. p-Tolylcopper (0.8 mmol) and II (2.4 mmol) were dissolved in toluene-d₈ (3 mL). NMR (toluene-d₈) δ (100 °C) H₆ (II), 8.40 (d, br); H₂ (4-TolCu), 7.74 (d); NCH₂ (II), 3.08 (s, br); NMe (II), 1.84 (s, br); CH₃ (4-TolCu), 1.90 (s, br). Solid CuOTf (3.2 mmol) was added to this solution which was then stirred at room temperature for 2 h. NH₃/H₂O solution (10 mL, 6 N) was added. Workup as described above resulted in the isolation of a white solid. NMR spectroscopy indicated that (2-Me₂NCH₂C₆H₄)- $_2$ (VIII) (64 mol %), 2-Me₂NCH₂C₆H₄C₆H₄CH₃- $_2$ P (15 mol %), p,p'-bitolyl (15 mol %), and N,N-dimethylbenzylamine (15 mol %), p.p. -bitolyi (15 mol %), and N, 14 difficulty identify identifies identification identifies iden dimethylbenzylamine, 2.08 (NCH₃), 3.26 (NCH₂).

The corresponding reaction but now using p-tolylcopper/II/CuOTf in a 1/0.8/2 molar ratio afforded VIII (34 mol %), 2-Me₂NCH₂C₆H₄C₆H₄CH₃-p (16 mol %), and p,p'-bitolyl (51 mol %). GC/MS: VIII, m/e 268; 2-Me₂NCH₂C₆H₄C₆H₄CH₃-p, m/e 225; p,p'-bitolyl, m/e 182.

Interaction of o-Tolylcopper with CuOTF. Pure o-tolylcopper was prepared according to ref 9. NMR (C₅D₅N) δ 2.39 (s, 3, CH₃), 7.0-7.4 (m, $H_{3,4, \text{ and } 5}$), 8.42 (m, 1, H_{6}); (in $C_{6}D_{6}$) 2.62 (CH₃), 6.7-7.1 (other H_{arom}), 7.97 (d, br, J=6 Hz, H_6). Upon heating (75 °C, 1 h) of the benzene- d_6 solution in the NMR tube o,o'-bitolyl was formed exclusively [§ (CH3) 1.96 ppm]. Mol wt (cryometry in C6H6) \overline{n}_{0h} 6.1, $\overline{n}_{3/4h}$ 4.9, \overline{n}_{2h} 4.1; second run \overline{n}_{1h} 4.5, \overline{n}_{2h} 4.0, concentration inde-

Reaction of o-Tolylcopper with CuOTf. Solid CuOTf (1.2 mmol) was added to a solution of o-tolylcopper (1.2 mmol) in benzene (10 mL). A purple colored product precipitated during 5 h of stirring. Workup with NH₃/H₂O solution in air afforded a colorless oil which consisted of o,o'-bitolyl, o-hydroxytoluene, and o-aminotoluene. The recovery of tolyl group in these products amounted to 46%. These compounds were identified by NMR and GC/MS techniques. NMR (benzene- d_6) o,o'-bitolyl, δ 1.95 (CH₃); o-aminotoluene, 1.80 (CH₃) and 2.84 (br, NH₂); o-hydroxytoluene, 2.08 (CH₃). Mol % 43/37/21. Their identity was further established by GC/MS analysis. A trace amount of a compound with C₁₄H₂₄O (m/e 198) was identified to be bis(o-methylphenyl) ether (very intensive m/e 107)

Exactly the same result was obtained when after the addition of NH₃/H₂O solution O₂ gas was bubbled through the reaction mix-

Complex Formation of o-Tolylcopper with CuOTf. An equimolar mixture of CuOTf and o-tolylcopper (CuOTf added to o-tolylcopper) which was dissolved in benzene (50 mL) was stirred for 0.5 h. The purple colored precipitate was filtered off, washed with benzene (3 × 40 mL, removal of uncomplexed CuOTf) and with pentane $(2 \times 10 \text{ mL})$, and dried in vacuo. Elemental analysis of this solid pointed to the isolation of a CuOTf·1.1 o-tolylcopper complex contaminated with metallic copper. NMR (pyridine-d₅) δ 2.5 (br, NCH₃) and 8.4 (br. Hg).

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Registry No.-I, 32691-15-9; II, 38286-29-2; III, 5588-74-9; IV, 20854-03-9; V, 61966-49-2; VI, 20627-78-5; VII, 121-69-7; VIII, 38286-37-2; IX, 613-33-2; X, 605-39-0; XI, 95-53-4; XII, 95-48-7; CuOTf, 42152-44-3; $(2-Me_2NC_6H_4)_4Cu_6Br_2$, 58616-70-9; $(2-Me_2NC_6H_4)_4Cu_6Br_2$ Me₂NC₆H₄+₂·2CuOTf, 61966-46-9; triphenylphosphine, 603-35-0; 2-Me₂NC₆H₄Cu·3CuOTf·6PPh₃, 61966-50-5; CuO₂CCF₃, 25535-55-1; $3(2\text{-}Me_2NCH_2C_6H_4Cu)\cdot 2CuOTf, \ 61966\text{-}48\text{-}1; \ bis(\textit{o}\text{-}methylphenyl)$ ether, 4731-34-4; $(2-Me_2NCH_2C_6H_4)_4Cu_4$, 37185-48-1; $(4-MeC_6-4)_4Cu_4$ H_4)₄Cu₄, 61966-47-0; 2-Me₂NCH₂C₆H₄C₆H₄CH₃-p, 61846-68-2.

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- The solubility of CuOTf in benzene (cuprous halides are insoluble) favors this equilibrium in spite of the fact that OTf is a weaker bridging ligand than bromide.²
- Thermal decomposition of 2-Me $_2$ NCH $_2$ Ce $_6$ H $_4$ Cu (II) in naphthalene at 165 °C for 6 h, which affords a 9/1 mixture of VIII and the arene (63% overall yield starting from in situ prepared II, see Experimental Section), provides an alternative route to the dimer VIII. Oxidation of II with O_2 gives rise to a mixture of VIII, the arene, and 2-hydroxybenzyldimethylamine [NMR (CCI₄)
- δ 2.27 (s, 6 H, NCH₃), 3.53 ppm (s, 2 H, NCH₂); in C₆D₆, δ 1.72 (s, 6 H, NCH₃), 3.15 (s, 2 H, NCH₂)].
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A Simple Preparation of Phenols from Diazonium Ions via the Generation and Oxidation of Aryl Radicals by Copper Salts^{1a}

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The standard method of preparation of phenols from diazonium salts consists of heating the latter in highly acidic aqueous solution; the high temperature and acidity often cause a variety of unwanted side reactions. We advocate an entirely different procedure which can be performed in a few minutes in neutral solution at room temperature, or below. The method is based on our previous observation that aryl radicals can be oxidized to phenols by cupric ion and it consists of adding cuprous oxide to a dilute solution of the diazonium salt dissolved in a solution containing a large excess of cupric nitrate. In one case the presence of silver(I) appeared to accelerate the radical oxidation. Not only is the redox procedure simpler than the thermal method, but in all cases studied to date, the yields are equivalent or superior to those obtained by the thermal procedure. In four cases in which the latter is unsatisfactory, the redox method is quite successful and it is considered the method of choice for new cases.

As indicated in all textbooks in organic chemistry the standard method for the conversion of an aromatic diazonium ion to a phenol is thermal decomposition of the diazonium ion in a highly acidic aqueous medium. The great deal of controversy concerning the mechanism of this reaction² has apparently been resolved recently in favor of a substantially free, singlet, aryl cation intermediate (eq 1).3

$$ArN_2^+ \longrightarrow N_2 + Ar^+ \xrightarrow{H_2O} ArO^+H_2 \longrightarrow ArOH + H^+$$
 (1)

The high acidity is required in order to suppress the ionization of the product phenol to phenoxide ion which couples readily with unreacted diazonium ion to produce an azo compound.^{4,5} In order to obtain high yields of phenol uncontaminated with azo compound, it is frequently necessary to add a solution of the diazonium salt to a boiling sulfuric acid solution,6 if possible with simultaneous removal of the phenol by steam distillation.7

The coupling reaction is only one of a variety of competing reactions which plague the synthesis of phenols by this route. Intramolecular nucleophiles or potential nucleophiles can also attack the diazonium group. For example, an ortho carboxamido group reacts with a diazonium function to yield a benzo-1,2,3-triazene.8 Similarly, an ortho hydroxyl group leads to the production of a diazoxazole, 9 while an ortho thiol group leads to a benzothiadiazole. 10 Intramolecular diazo coupling with a suitably placed electron-rich ring has also been observed.¹¹ Nucleophilic attack on the diazonium function by an ortho vinyl group to form an indazole (1) is also common (eq 2).12

$$\begin{array}{c}
H \\
C = C
\end{array}$$

$$\begin{array}{c}
A, H_2O, H^+ \\
N \\
H
\end{array}$$

$$\begin{array}{c}
C \\
N \\
H$$

$$C \\
A \\
H$$

$$\begin{array}{c}
C \\
N \\
H$$

$$C \\
A \\
H$$

$$C \\
A \\
H$$

$$C \\
A \\
H \\
H$$

$$C \\
A \\
H \\
H$$

$$C \\
A \\
H$$

$$C \\
C \\
A \\
H$$

$$C \\
A \\
H$$

$$C \\
A \\
H$$

$$C \\
A \\
H \\
H$$

$$C \\
A \\$$

Even a saturated carbon atom in the ortho position to a diazonium function can serve as a nucleophile, presumably after deprotonation. 13-15 For example, the diazonium ion 2

yields 85% of the indazole 3 when heated in acid solution, 13 although the phenol can be produced instead by adding the diazonium solution to a boiling sulfuric acid solution.¹⁴

Another problem sometimes encountered during thermal decomposition of a diazonium ion is replacement of an ortho or para substituent with a hydroxyl group. 16

In addition to these competing reactions in which the two nitrogen atoms are retained in the product, several reactions in which nitrogen is lost are known to occur in place of phenol