As mentioned above the yield of CH_4 is ~0.56 per cluster when pure HSO₃CF₃ is employed, and it is increased to 0.82 per cluster when the reaction medium is 1:1 HSO₃CF₃- $O(SO_2CF_3)_2$. Neither concentrated HCl nor pure HSO₃F produces volatile hydrocarbons, but concentrated H₂SO₄ yields a trace of CH₄ along with SO₂, CO₂, CO, COS, and H₂.

Two experiments were performed to test whether the methane originates from the coordinated carbon monoxide, or from an adventitious source. In the first of these, DSO₃CF₃ was employed as reactant with the Fe₄ cluster, and infrared spectra of the methane-carbon monoxide fraction showed the presence of CD₄ and no indication of ¹H-containing methane. In the second experiment, the $[Fe_4(CO)_{13}^{2-}]$ was enriched such that 26% of the carbon monoxide contained the ¹³C isotope. The reaction of this enriched metal carbonyl with HSO₃CF₃ according to the procedure indicated above produced methane which was 25% ¹³CH₄ as judged by quantitative infrared spectroscopy. These experiments convincingly show that the methane originates from carbon monoxide in the metal cluster, and therefore confirm eq 1.

Since H_2 is a product of the reaction, it was conceivable that methane might arise from a H_2 reduction rather than direct attack of H⁺ on the metal cluster. To check this possibility the reaction was run under the uniform conditions outlined above but with the addition of 457 Torr of H_2 , which is 58 times greater than the H_2 produced in a typical run. The yield of CH_4 was 0.53 per cluster, which, within experimental error, is equal to the yield in the absence of added H₂. Two isotopic experiments also demonstrate the lack of incorporation of dihydrogen into the product. When H_2 (344 Torr) was placed over the reaction mixture containing DSO₃CF₃ in place of HSO₃CF₃, the only methane produced was CD4 as judged by infrared spectroscopy (eq 2) and, when the reaction was run with D_2 over a reaction mixture containing HSO₃CF₃, only CH₄ was produced (eq 3). From these experiments it is clear that the hydrogen in the methane originates from H^+ and not from H_2 intermediate.

$$[Fe_4(CO)_{13}^{2-}] + DSO_3CF_2 \xrightarrow{H_2} CD_4$$
(2)

$$[Fe_4(CO)_{13}^{2-}] + HSO_3CF_3 \xrightarrow{D_2} CH_4$$
(3)

One possible mechanism is attack on the CO oxygen by protons to split off water, leaving a carbide which subsequently is attacked by H^+ to produce methane (eq 4). Any carbide

$$Fe_{4} \xrightarrow{CO} \xrightarrow{+H^{+}}_{-H_{2}O} Fe_{4} \xrightarrow{C} \xrightarrow{H^{+}}_{-H_{4}} CH_{4} + Fe^{2+} + iron carbonyls$$
(4)

intermediate would have to display a different reactivity pattern from known carbides, because we have observed the conversion of [Fe₆(CO)₁₆C²⁻] into Fe(CO)₁₅C in HSO₃CF₃¹⁰ without the production of CH₄.

The work of Chatt and co-workers¹¹ on the reduction of coordinated dinitrogen (eq 5) indicates that the necessary

$$L_n \stackrel{6e^-}{MNN} _{0} \stackrel{6H^+}{\longrightarrow} 2NH_3 + M^{VI} + nL \qquad (5)$$

$$M = Mo \text{ or } W$$

equivalents for CO reduction may be supplied by a mononuclear complex of a metal such as Mo or W, which can easily reach a high oxidation state. In the iron series, however, it appears that a cluster is necessary to supply the six electrons required for the complete reduction of CO. Thus the mononuclear carbonyl Fe(CO)5 in HSO3CF3 produces the known ion $[HFe(CO)_5^+]^{12}$ The three-iron cluster $[Fe_3(CO)_{11}^{2-}]$ yields traces of CH4 when treated with HSO3CF3, and as described above a good yield of methane was obtained from the four-iron cluster.

It is probable that the reaction of metal carbonyl clusters with the surface of hydrated metal oxides is analogous to the homogenous reaction reported here.¹³ A closer comparison of the two reactions will be possible when the role of dihydrogen is delineated in the heterogenous reaction.

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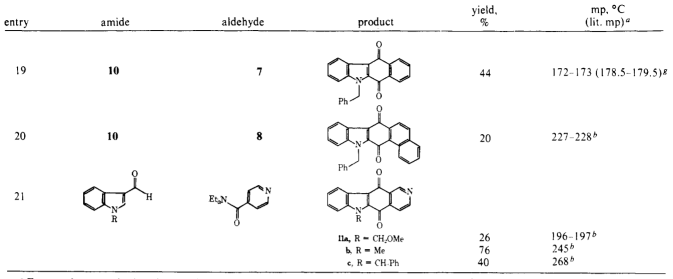
Tandem Directed Metalation Reactions. Short Syntheses of Polycyclic Aromatic Hydrocarbons and Ellipticine Alkaloids

Sir:

Directed metalation of aromatic substrates¹ is an increasingly important strategy for the regiospecific synthesis of highly substituted benzene derivatives,² heterocycles,^{2b,c,3} and natural products.⁴ The rapidly growing list of functionalities capable of promoting and stabilizing ortho metalation which includes oxazolines,^{3a,5} imidazolidine,⁶ CH=NR,^{2a,4a,7} CONR,^{31,4b,f,g,8,9} CSNR,^{3c} CONR₂,^{2b,3b,4c,d} CH(OMe)₂,⁴ⁱ OCH₂OMe,^{3d,4e,h} OCH(Me)OEt,^{3k} CH₂SMe,¹⁰ CH₂O⁻,^{3e,f} $\overline{N}COR$,^{3h} N=C,³ⁱ and halo and methoxy ArCr(CO)₃^{3j} offers attractive and, at times, highly advantageous solutions to preparative problems not readily achieved by classical (usually electrophilic) chemistry. We report on the new concept of tandem directed metalation (Scheme I) involving ortho-lithiated tertiary benzamide (1) and benzyl alkoxide (3) intermediates which provides rapid access to polycyclic anthraquinones and unusual heterocyclic benzoquinones (Table 1). We delineate the scope of this one-pot aromatic ring construction reaction and demonstrate its utility for the synthesis of carcinogenic polycyclic aromatic hydrocarbons (PAH's)¹¹ and chemotherapeutic ellipticine alkaloids (6).¹² Both classes of compounds currently constitute lively and urgent areas of study.^{13,14}

entry	amide	aldehyde	product	yield, %	mp, °C (lit. mp) ^a
	0				
1	NEt 2	н	$i \int_{h}^{2} \frac{1}{1 + 1} \int_{-\infty}^{\infty} \frac{1}{e^{-h}} h$	43	284 (287)
2	7			10	169 160 (170)
2 3	7 7	2-MeOC6H₄CHO 4-MeOC6H₄CHO	1-methoxyanthracene-9,10-dione 2-methoxyanthracene-9,10-dione	10 28	168–169 (170) 192–193 <i>^b</i>
4	7 7 7	4-MeC ₆ H ₄ CHO	2-methylanthracene-9,10-dione	15	184 (177-179)
5	7	1-naphthaldehyde	benz[a]anthracene-7,12-dione	44	165-166 (166-167)
				20	132-133 (137)
6	7	2-naphthaldehyde	benz[b]anthracene-5,12-dione	39	283 (284-285)
U		1	Benz[a]anthracene-7,12-dione	5	
_	-		3-(2-naphthyl)-phthalide	22	140-141 ^b
7	7	9-phenanthraldehyde	Dibenz[<i>a</i> , <i>c</i>]anthracene-9,14-dione 3-(5-phenanthryl)phthalide	41 24	179–180 (181–183) 162–164 ^b
8	3,4-diOMe- C ₆ H ₃ CONEt ₂	PhCHO	3-Phenyl-4,5-dimethoxyphthalide	74	81 ^b
			0 Ph		
9	\square	РһСНО		81	164-166 ^{<i>b</i>}
10	8	2-naphthaldehyde	Dibenz[a,i]anthracene-7,14-dione	10	225-226 (229)
	-	2	2naphthyl		
				70	1 48 ^b
11	8	l-naphthaldehyde	Dibenz[a,h]anthracene-7,14-dione	10	249-249.5(248-249)
				81	166(170.5-171) ^c
12		2-naphthaldehyde	dibenz[a,i]anthracene-7,14-dione	2	225-226 ^d
	O NEt ₂		dibenz[<i>b,i</i>]anthracene-6,13-dione 0	2	>300 (388-398)
13	Н	7		35	225 (227-228) ^e
	9 9		s		
				27	205 (100, 201) (
14	9	8		37	205 (199-201) ^f
		~0			
15	9			24	250*
		0			
16	9			77	252-253 (258-260) ^e
		ő	ů o		
17	9	Et ₂ N		20	192-193 <i>^b</i>
	Q	11 O	- II O		
18	Н	Et ₁ N	()	67	211-212 ^b
10	N	0			
	Ph - 10		Ph		

 Table 1 (Continued)



^a Except where noted otherwise, melting points refer to values quoted in or traced from ref 16. ^b New compound; analytical and spectral data are in full accord with the structure. ^c L. F. Fieser and G. W. Kilmer, J. Am. Chem. Soc., **61**, 862 (1939). ^d Identical (melting point, mixture melting point, UV and NMR spectra) with the sample in entry 10. ^e D. W. H. MacDowell and J. C. Wisowaty, J. Org. Chem., **37**, 1712 (1972). ^f M. S. Newman and K. G. Ihrman, J. Am. Chem. Soc., **80**, 3652 (1958). ^g V. I. Shvedov, G. N. Kurilo, and A. N. Grinev, Khim. Geterotsikl. Soedin., 1079 (1972); Chem. Abstr., **77**, 139712s (1972).

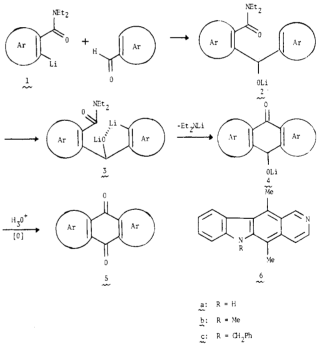
Metalation (1.0 equiv of sec-BuLi, 1.0 equiv of TMEDA, Et₂O, -78 °C, 1 h) of N,N-diethylbenzamide (7) gave the yellow-orange ortho-lithiated species 1 which, upon sequential treatment at -78 °C with benzaldehyde (1.0 equiv) and sec-BuLi (1.25 equiv) and allowing the reaction mixture to warm to room temperature overnight (12 h), afforded anthraquinone in 43% yield. Table I illustrates representative anthraquinones and heterocyclic benzoquinones obtained by this procedure. Alkoxy and alkyl groups in the benzaldehyde reactant are poorly tolerated and give low yields of anthraquinones (entries 2-4), whereas alkoxy-substituted benzamides afford (after treatment with TsOH)^{2b} only phthalide products (entry 8). The reaction of 7 with 1- and 2-naphthaldehydes and 9-phenanthraldehvde provides polycyclic anthraquinones in modest yields (entries 5-7). Entry 6 shows the preference for the formation of a linear rather than an angular anthraquinone system. 1-N,N-Diethylnaphthamide undergoes reaction with benzaldehyde and 2-naphthaldehyde to give (after TsOH) mainly phthalide derivatives (entries 9 and 10). 2-N,N-Diethylnaphthamide is a poorer substrate providing the two possible anthraquinones in low yields (entry 12).

Some of the condensed anthraquinones (entries 5, 7, 11, and 12) were reduced (aluminum tricyclohexoxide, cyclohexanol)¹⁵ to the corresponding hydrocarbons,¹⁶ thus completing short syntheses which compete favorably with reported methods for these carcinogenic PAH's.^{15,16}

Knowledge concerning the facile abstraction of the 2 proton in furan,¹⁷ thiophene,¹⁷ and N-substituted indole¹⁸ derivatives stimulated tandem directed metalation reactions on these systems. Thus treatment of ortho-lithiated benz- and 1naphthamides with thiophene-3-aldehyde gave 4,9-dihydronaphtho[2,3-b]thiophene-4,9-dione and its 7,8-benzo analogue, respectively (entries 13 and 14). Furthermore, condensation of the same thiophene aldehyde with metalated furan-3, thiophene-3, and pyridine-4¹⁹ amides afforded the easily isolable diheterocyclic benzoquinones (entries 15–17). Combination of N-benzylindole-3-aldehyde (10) with ortho-lithiated aromatic and heteroaromatic amides smoothly gave condensed indolobenzoquinone systems (entries 18–20). Most of these compounds have been accessible only by tedious and inefficient routes.

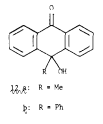
A one-step construction of the ellipticine alkaloid skeleton





was achieved by using N-substituted indole-3-aldehydes with lithiated N,N-diethylisonicotamide (entry 21).²⁰ Sequential treatment of the N-methylenemethoxyquinone **11a** with excess MeLi, 47% HI in MeOH, and SnCl₂ in HOAc-HCl-THF (5:1:5)²¹ without isolation of intermediates gave ellipticine (**6a**) (overall 40% yield).²² The same reaction sequence on the Nmethyl- (**11b**) and N-benzylquinones (**11c**) provided the corresponding ellipticine derivatives **6b** (62%) and **6c** (40%).²³

The proposed mechanism (Scheme I) for the reaction, in particular, the formation of dianion 3, is supported by independent tandem metalation of 7 with acetophenone and benzophenone under identical reaction conditions to yield the hydroxyanthracenones 12a (17%) and 12b (15%).²⁴ The generation of simple ortho-lithiated benzyl alkoxide species corresponding to 3 has been documented.^{3e,f,25} The low yields



of anthraquinones in the reaction of 7 with alkoxy and methyl benzaldehydes (entries 2-4, 8) points to complications due to competitive metalation directed by CONEt₂ and CH₂O⁻ groups and steric effects in the cyclization, $3 \rightarrow 4$. The aerial oxidation step, $4 \rightarrow 5$, is well precedented.^{26,27}

Outside of the specific application presented here, the concept of tandem directed metalation may have broader significance in organic synthesis.28

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Determination of Rotational Correlation Time from Perturbed Angular Correlations of γ Rays: **Apomyoglobin Reconstituted with** ¹¹¹Indium(III) Mesoprotoporphyrin IX¹

Sir:

Currently popular methods for determining macromolecular rotational correlation time, τ_{rot} , each possess major drawbacks. Fluorescence depolarization (FD)² and depolarized light scattering³ require transparent media. Electron paramagnetic resonance (EPR) "spin labeling" 4 and FD require introduction of an artificial reporter group, linked by one or more flexible bonds to the macromolecule. EPR line-shape analysis is complicated by inhomogeneous (nitrogen hyperfine coupling) broadening, nitrogen nuclear relaxation, and g-tensor nonaxial symmetry.⁵ NMR relaxation time analysis is complicated by competing relaxation pathways,6,7 internuclear distance uncertainty,⁸ pulse imperfections, and multiexponential relaxation.9,10

In contrast, time-resolved emission anisotropy from a perturbed $\gamma - \gamma$ angular correlation (PAC) experiment can provide direct determination of $\tau_{\rm rot}$,¹¹⁻¹⁶ in opaque media (even in vivo17), at 10⁻¹² M, without flexible and/or bulky reporter groups. Unfortunately, previous PAC determinations of τ_{rot} have been complicated by multiple $au_{\rm rot}$ processes.¹⁴⁻¹⁶ In this communication, we report the PAC results from the successful reconstitution of indium-111 mesoprotoporphyrin IX, In-MPP (see ref 18 for synthesis), into apomyoglobin by the procedure of Srivastava¹⁹ to give a singly labeled protein with unique conformation and chemical form.20

The crystal and molecular structure of the closely related tetraphenylporphinatoindium(III) chloride²¹ strongly suggests that In-MPP-myoglobin should be isostructural with the Fe(II) heme in deoxymyoglobin or deoxyhemoglobin, because the indium atom is displaced above the mean porphyrin plane by the same distance (0.6 Å) as is Fe(II) in the native proteins.