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1,3-Dipolar Cycloadditions to Coordinated Azide in Cobalt Chelate Complexes of the Type LCo(chelate)N3

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Several electron-poor dipolarophiles (alkynes, alkenes, and nitriles) react with azido cobalt chelate complexes of the type LCo(chelate)N₃ under mild conditions. Coordinated five-membered-ring heterocycles are the initial products of these 1,3-dipolar cycloadditions. Nonterminal alkynes yield triazoles, alkenes yield triazolines, and nitriles yield tetrazoles. The reactivity of the dipolarophile increases with an increase in the electron-withdrawing power of its substituents. The reactivity of the cobalt complexes is influenced by the nature of the neutral trans-coordinated ligand (L, trans effect) and by the nature of the anionic chelating system (cis effect). Other reagents also react with these azido complexes, but the products do not result from 1,3-dipolar cycloadditions. Terminal alkynes behave as C-H acids and produce alkynyl complexes, HC1 produces LCo(chelate)Cl, HClO₄ produces [LCo(chelate)]⁺ClO₄, carbon monoxide produces isocyanato complexes, LCo(chelate)NCO, and heterocumulenes generally give ill-defined products but CH₃SCN undergoes ligand substitution and thiocyanate decomposition to produce CH,SCNCo(chelate)CN. Each of the products of these reactions was characterized by elemental analysis and infrared and 'H NMR spectroscopy.

Introduction

Organic azides are synthetically very useful reagents.' Among the many and varied transformations that they undergo, perhaps the most important are their 1,3-dipolar cycloaddition reactions to produce heterocycles.^{2.3} Thus, alkynes react with organic,⁴⁻⁶ silyl,^{7,8} tin,⁹ and lead¹⁰ azides to produce triazoles. But only a few 1,2,3-triazoles with free NH groups have been prepared from explosive HN,.

Organic azides also react with electron-poor alkenes (enamines and enol ethers^{4,11,12}) and strained alkenes^{1,4,13} to produce triazolines, but simple alkenes either do not react or react very slowly.

Organic azides,^{14,15} binary metal azides,¹⁶⁻¹⁹ silyl azides,^{14,20} and complex metal a zides a^{21-36} react with electron-poor nitriles under relatively mild conditions to produce tetrazoles.

In a search for convenient and reactive sources of azides for 1,3-dipolar cycloadditions, we have probed the reactivity of complexes of the type $LCo(chelate)N_3$.

Complexes of the type LCo^{III} (chelate)X have been widely studied because of their resemblance²⁷⁻³⁰ to vitamin B₁₂, and a wide variety of these complexes has been synthesized.^{31,32} Because the nature of both the anionic chelating ligands and the neutral trans ligand can be widely varied, this affords **us** the opportunity of simultaneously investigating both the cis and trans effects upon the reactivity of the coordinated azide. We report herein the reactions of a number of LCo (chelate) $N₃$ complexes **I-IV** with several representative electron-poor alkynes, alkenes, nitriles, and heterocumulenes.

Results and Discussion

I. Reactions with Alkynes. Complexes **I-IV** were all found to react with electron-deficient internal alkynes (see Table **I)** under mild conditions (\sim 20 °C) to produce triazolato complexes similar to what had previously been observed for the Ag), $Rh_2(C_5H_5)_2(N_3)_4$ ²⁶ and CpFe(CO)₂N₃³³ complexes. The reactions of the cobalt complexes all occur at lower temperatures and usually with shorter reaction times than the analogous reactions of NaN_3^{36} and HN_3 .³⁵⁻³⁹ $(R_3P)_2M(N_3)_2^{24}$ (M = Pd,Pt), $[(R_3P)_2MN_3]_2^{23,25}$ (M = Cu,

(a) Triazolato Complexes. As was found for the reaction of silyl azides' with dimethyl acetylenedicarboxylate, the cobalt complexes react to produce $N(2)$ -bound 4,5-bis(methoxycarbonyl)- 1,2,3-triazolates. The structure of the complexes **is** clearly established as the N(2) isomer from the appearance

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of their ¹H NMR spectra which show a singlet resonance at $\delta \approx 3.7$ for the six methoxycarbonyl protons. The ¹H NMR

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Table I. Triazolato Complexes Formed from the Reaction of Alkynes with Cobalt Azido Chelate Complexes

				IR data (KBr) , cm ⁻¹					
com-		triazolato				¹ H NMR data, CDCl, vs. Me ₄ Si			
plex	starting azide complex	alkyne	ring vib	$\nu(C=O)$	$\nu(C-O)$	δ	rel intens		
la	$PPh_3Co(DH), N_3$	$CH_3CO_2C \equiv CCO_2CH_3$ 827, 798, 773		1737	1296	7.42, 3.79, b 2.42 ^{<i>a</i>}	15:6:12		
1 _b	MePPh, Co(DH), N,	$CH_3CO_2C \equiv CCO_2CH_3$ 829, 804, 780		1720	1243	7.55, 3.70, $^{\circ}$ 1.99, $^{\circ}$ 1.95 ^a	10:6:3:12		
1 _c	Me, PPhCo(DH), N,	$CH_3CO_2 \cong CCO_2CH_3$	823, 801, 791, 1738, 1717 778		1238	7.35, 3.71, b 2.09, a 5:6:12:6 1.53 ^c			
1d	$PBu3Co(DH)$, N ₃	CH_3CO , $C \equiv CCO$, CH_3	828, 801, 770 1730		1290	3.68^{b} 2.26^{a} 1.1	6:12:27		
1e	$P(Bz1)$ ₃ Co(DH), N ₃	CH,CO,C≡CCO,CH,	826, 794, 773	1748, 1720	1295	6.8.3.65, b 2.95, c 2.13^{a}	15:6:6:12		
1f	$P(OCH_3)$ ₃ Co(DH) ₂ N ₃	CH_3CO , $C = CO$, CH_3	827, 792, 780	1732	1285	$3.74b$ 3.65, 2.31 ^a	6:9:12		
1g	$P(OPh)$, $Co(DH)$, N_3	$CH, CO, C=CCO, CH,$	828, 798, 775	1737	1295	7.08, 3.74, b 2.08 ^{a}	15:6:12		
1h	PhNH, Co(DH), N,	CH , CO , $C = CCO$, CH ,	838, 823, 803	1727, 1692	1298				
1i	$pyCo(DH), N_2$	CH_3CO , $C \equiv CCO$, CH_3	825, 805, 779	1725	1245	3.78^{b} 2.37 ^a	6:12		
1 _j	$NH3Co(DH)2N3$	$CH3CO2C \equiv CO2CH3$	832, 805, 776	1729	1243				
1k	$1-MelmCo(DH), N_3$	CH_3CO , $C \equiv CCO$, CH_3	834, 800, 779	1720	1295	7.1, 3.74, $\overset{b}{}$ 3.58,	3:6:3:12		
11	AsPh ₄ [Co(DH) ₂ (N ₃) ₂]	$CH, CO, C=CCO, CH,$	828, 799, 792, 1709, 1734 771		1295	2.30^{a}			
1 _m	PBu, Co(DPH), N,	$CH, CO, C=CCO, CH,$	826, 801, 779	1743, 1722	1288	7.25, 3.72, b 1.45	20:6:27		
1n	$[Co[(DO)(DOH)pn](N3)$,	CH_3CO , $C = CO$, CH_3	829, 798, 772	1740	1293				
1 _o	PPh, Co(Salen)N,	$CH, CO, C = CO, CH,$	828, 798, 775	1730		7.28.3.69	8:6		
1 _p	$PPh_3Co(Salphen)N_3$	$CH, CO, C=CCO, CH,$	822, 798, 775	1712, 1732		7.37, 3.57	26.9:6		
1q	cis -PPh ₃ Co(acac) ₂ N ₃	$CH_3CO_2C \equiv CCO_2CH_3$	830, 804, 780	1720, 1750	1295	7.37, 4.58, 3.70, b 1.65	15:2:6:11.65		
2a	PhNH, Co(DH), N,	$HC = CCO, Et$	850, 779	1712	1241, 1091	7.18, 4.16, 3.44, 2.22 ^a 1.26	5:2:2:12:3		
2 _b	pyCo(DH), N,	$HC = CCO, Et$	830, 775	1730	1236, 1090	7.25, 4.25, 2.22 ^a 1.30	5:2:12:3		
2c	$NH_3Co(DH), N_3$	$HC = CCO, Et$	829, 745	1705	1242, 1092				
2d	$AsPh4[Co(DH), (N3)$	$HC = CO, Et$	842, 831, 777	1723 sh. 1709	1242, 1231, 1092, 1078				
2e	PPh , Co(Salphen)N,	$HC = CCO, Et$	808,778	1720	1227, 1090				
2f	$pyCo(Salphen)N$,	$HC = CO, Et$	808,778	1720	1227, 1090	7.50, 4.17, 1.16	17.8:2.08:3		

^{*a*} Oxime CH₃, doublet, $J_{PH} = 1$ Hz, ^{*b*} CO₂CH₃, ^{*c*} P-CH₃ or P-CH₂, doublet, $J_{PH} = 11$ Hz.

spectrum of an $N(1)$ -bound isomer would exhibit two proton resonances for its anisochronous methoxycarbonyl groups. Each of the triazolato complexes also possesses very characteristic infrared spectra that clearly show the absence of $\nu(N_3)$ and the presence of $\nu(C=O)$, $\nu(C-O)$, and triazolato ring frequencies (see Table I).

Changing the electron-donor ability of the other ligands

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coordinated to the cobalt (III) center brings about a change in the rate constant for the cycloaddition reaction by 2 orders of magnitude.⁴⁰

$$
\begin{array}{ccc}\n\sum_{i=1}^{N_3} & & & \sum_{i=1}^{N_3} & & \sum_{i=1}^{N_3} & & \\
\sum_{i=1}^{N_3} & & & \sum_{i=1}^{N_3} & & \\
\sum_{i=1}^{N_3} & & & \sum_{i=1}^{N_3} & \\
\vdots & & & \vdots & \\
\sum_{i=1}^{N_3} & & & \sum_{i=1}^{N_3} & \\
\vdots & & & & \vdots \\
\sum_{i=1}^{N_3} & & & \sum_{i=1}^{N_3} & \\
\vdots & & & & \vdots \\
\sum_{i=1}^{N_3} & & & & & \vdots
$$

For example, the pseudo-first-order rate constants for the reactions of $LCo(DH)₂N₃$ with dimethyl acetylenedicarboxylate are $\sim 10^{-3}$ s⁻¹ when L = PPh₃ and $\sim 10^{-5}$ s⁻¹ when $L = BzI_3P$ at 27 °C in CHCl₃. Though the N(1) isomer (V) (reaction 1) is surely the kinetic product of these reactions, the isolated thermodynamically stable product is the $N(2)$ isomer (VI). Isomerization from $N(1)$ - to $N(2)$ -bound triazole is most likely sterically promoted as has been found for the analogous tetrazolato complexes.⁴¹

Following the cycloaddition reaction, the base $B = PPh_3$, is partially liberated and produces an equilibrium mixture of five-and-six coordinate complexes as depicted in reaction 2.

XCo(chelate)PPh₃
$$
\frac{-PPh_3}{+PPh_3}
$$
 XCo(chelate)
X = triazolate (2)

By addition of triphenylphosphine to the reaction solution, one

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can isolate the corresponding complex which loses triphenylphosphine upon washing with diethyl ether. The tendency for formation of the five-coordinate complex is a function of the chelating ligand system and increases in the order²⁸ (DH)₂ \sim Salphen \lt (PH)₂ \lt (NH₂) \lt Salen \lt BAE (cis effect). These equilibria also occur for the bases $L =$ AsPh₃, SbPh₃, P(OPh₃)₃, and P(OEt)₃ but not with the more tightly bound base pyridine. These observations lead to the conclusion that the triazolate has a greater trans-labilizing ability than azide. When the chelate is Salen or BAE, both the $PPh_3Co(chelate) triazolate$ and $Co(chelate) triazolate$ complexes may be isolated. Loss of triphenylphosphine from the former upon washing with ether is visually evident by the formation of crystalline triphenylphosphine in the ether wash solutions.

Similar to these reactions, the diazido cobalt complexes react with dimethyl acetylenedicarboxylate to produce bis(triazolate) complexes (compounds **11** and **In** of Table I).

(b) Alkynyl Complexes. Terminal alkynes react as CH acids and undergo a ligand substitution reaction rather than a 1,3-dipolar cycloaddition with the cobalt azide complexes to produce alkynyl complexes with liberation of $HN₃$ according to reaction 3 (see Table II). The more electron withdrawing
LCo(chelate)N₃ + RC=CH \rightarrow

$$
RC \equiv CCo(chelate)L + HN3 (3)
$$

the R group, the more stable is the alkynyl complex. Propynal and (more slowly) propynal diacetate produced only the alkynyl substitution product independent of the nature of the cobalt azide complex, whereas ethyl propiolate produced predominantly the triazolate complex. But ethyl propiolate reacted with $PPh_3Co(Salen)N_3$ to produce the alkynyl complex, suggesting that there is little difference in the stability of the triazolate and alkynyl complexes. The alkynyl complexes have also been obtained by reaction of the corresponding chlorocomplexes with the sodium or silver acetylide.^{42,43} The velocity⁴⁰ of the cycloaddition or ligand substitution as well as the nature of the product formed (alkynyl or triazolate complex) is related to the acidity of the alkyne. By comparison, trimethylsilyl azide gave no cycloaddition products in these reactions, only $HN₃$ liberation occurred.⁴

(c) Cleavage of the Coordinated Triazole. We have recently shown⁴¹ that the complexes $PBu_3Co(DH)_2(5-R-tetrazolate)$ react with alkyl halides such as benzyl bromide and methyl iodide to alkylate the tetrazole ring regiospecifically at the $N(1)$ nitrogen. Since all the triazolate complexes contain $N(2)$ -bound triazolate, it was of interest to determine if they would react similarly to produce $N(1)$ -alkylated triazoles. Accordingly, seven of the LCo(DH),(triazolate) complexes were reacted with benzyl bromide in CDC1, at room temperature in a 5-mm NMR tube, and the progress of the reaction was followed by 'H NMR spectroscopy for 3 months. The complexes with $L = PBu_3$, CH_3PPh_2 , and $(CH_3)_2PPh$ were all found to react whereas when $L = 1$ -methylimidazole, pyridine, $P(OCH₃)₃$, or PPh₃, either no reaction or decom-

position occurred. The results were determined by comparing the 'H NMR spectra of the reaction solutions to that of an independently prepared sample of 1 **-benzyl-4,5-bis(methoxy**carbonyl)-1,2,3-triazole. The latter shows singlets at δ 7.22, 5.15, 3.90, and 3.83 in the ratio 5:2:3:3. The 2-benzyl-1,2,3-triazole isomer was not available, but it should be readily distinguishable from the 1-benzyl isomer by having a benzyl CH₂ resonance at a value different from δ 5.75 and having isochronous methoxycarbonyl methyl resonances. In each of the three cases where alkylation occurred, it occurred regiospecifically at the $N(1)$ nitrogen and at approximately the same rate as for the alkylation of coordinated tetrazole⁴⁵ (compare reactions 4 and *5).* Both reactions were run under

pseudo-first-order conditions at 36 $^{\circ}$ C in CDCl₃. The complex $PPh_3Co(DH)_2$ (triazolate) also reacted with benzyl bromide, but alkylation of PPh₃⁴⁶ to form [BzlPPh₃]⁺Br⁻ occurred rather than alkylation of triazolate; this is consistent with the previous conclusions that $PPh₃$ dissociation occurs for this complex.

Most attempts at liberating and isolating the triazole from these complexes have been unsuccessful. Typically, the chelate system also reacts with the liberating reagents to produce oily decomposition products, making isolation of the free triazole difficult. However, treatment of the complexes with HCl in acetone liberates the triazole in low yield and reaction with acetyl chloride produces the $N(1)$ acetyltriazole which can be hydrolyzed to the triazole (see Experimental Section).

II. Reactions with Alkenes. Most of the alkenes investigated did not produce pure products, probably due to the general thermal instability and base sensitivity of Δ^2 -triazoline.⁴⁷ The progress of the reactions of alkenes with azido cobalt complexes can be followed by the loss of $\nu(N_3)$ in the infrared spectrum. Generally, these reactions occur over a long period of time as with the corresponding alkyne reactions.⁴⁰ A uniform, wellcharacterized product was isolated from the reaction of diethyl maleate with the azido cobalt complex $N_3Co(DH)_2NH_3$ in Me₂SO/CHCl₃ as shown in reaction 6. This product exhibits $\nu(C=O)$ at 1742 cm⁻¹, $\nu(NN)$ at 1634 cm⁻¹, and $\nu(CO)$ at 1203 and 1044 cm^{-1} . The complex contains CHCl₃ of solvation. Similar reactions with maleic anhydride, acrolein, ethyl

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acrylate, acrylonitrile, and cyclopentadiene dimer did not produce analyzable products, but in each case $v_{\rm as}(N_3)$ completely disappeared indicating that a reaction had occurred.

111. Reaction with Nitriles. These azido cobalt chelate complexes undergo 1,3-dipolar cycloadditions only with very electron-deficient nitriles. In the reaction of a coordinatively unsaturated azido complex, partial activation of the nitrile by prior coordination to the metal through formation of a σ -donor bond may occur as previously discussed.^{22,48} A similar catalytic effect by the central metal in the coordinatively saturated cobalt chelate complexes is not likely. An increase in coordination number is also prohibited on steric grounds, and substitution of the trans ligand is not anticipated as in each case the coordinated bases are all stronger donors than nitriles. $49,50$

Both $N(1)$ - and $N(2)$ -bound tetrazoles may be formed by the reaction of azido complexes with nitriles. Molecular orbital calculations $51-57$ indicate that these two bonding modes are essentially electronically and energetically equivalent, and for Pd and Pt both isomers are obtained. $24,50,52$ However, we have recently shown⁴¹ that in the $R_3PCo(DH)_2$ (tetrazolate) complexes, N(2) coordination of the tetrazole is sterically promoted. It is therefore likely that the products listed in Table III contain $N(2)$ -bound tetrazolates.⁴¹ The relative rates of these reactions are a strong function of the nature of the nitrile (o-phthalodinitrile, benzonitrile, and acetonitrile did not react), the trans base, and the chelate ring system.40 Both of the diazido complexes $[Co[(DO)(DOH)pn](N_3)_2]$ and Ph₄As- $[Co(DH)₂(N₃)₂]$ did not react, and the complex PPh₃Co- $(DH)_{2}N_{3}$ reacted much more slowly⁴⁰ than the more electron-rich Schiff-base complexes $PPh_3Co(Salen)N_3$ and $PPh_3Co(Salphen)N_3.$

As was found for triazolates, the tetrazolates are also better trans-labilizing ligands than azide and the tetrazolate complexes are isolated as base free complexes.

IV. Reactions with Carbon Monoxide. Azide complexes react with carbon monoxide in solution to produce isocyanato complexes and the mechanism in reaction 7 has been proposed for these reactions. $21,58$

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Table **111.** Tetrazole Complexes Formed from the Reaction of Nitriles with Cobalt Azido Chelate Complexes

complex	starting azide complex	nitrile	
9а $10a^a$ 10Ь $11a^{a,b}$	$PPh_3Co(Salphen)N_3$ $PPh_3Co(Salen)N_3$ $PPh_3Co(Salphen)N_3$ $PPh_3Co(Salphen)N_3$	NCCF, NCCO, C, H NCCO, C, H NCCH, CN	

^{*a*} The product does not contain PPh₃. ^{*b*} Only one nitrile group reacts.

The high velocity of these reactions for coordinatively unsaturated complexes supports the proposal of coordination to and activation of CO by the central metal atom. In contrast the reaction of the azido cobalt oximes with CO probably occurs by direct interaction to form the isocyanato complexes $PPh_3Co(DH)_2NCO(12a)$ and $pyCO(DH)_2NCO(12b)$. More drastic conditions are necessary to cause these reactions (5 days at 50 \degree C and 40 atm of CO), and a catalytic effect of the cobalt atom is precluded by the steric bulk of the other ligands. These same complexes may be prepared by metathesis of the chloro complexes with NaNCO.

V. Reactions with Heterocumulenes. A variety of simple heterocumulenes such as carbon disulfide, isocyanates $(RN=C=O)$, isothiocyanates $(RN=C=S)$, and thiocyanates $(RSC=N)$ react with organic azides⁵⁹ as well as metal azide $\text{complexes}^{\{31,23-26,60\}}$ to produce five-membered heterocyclic rings. Upon reaction of the oxime complex $PPh_3CO(DH)_2N_3$ with methyl thiocyanate, the first thing which occurs is ligand substitution of PPh_3 by CH_3SCN . Subsequently, the azide group is replaced by cyanide so that one eventually isolates the complex $NCCo(DH)₂NCSCH₃$ as depicted in reaction 8.

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rings. Upon reaction of the oxime complex PPh₃CO(DH)₂N₃
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group is replaced by cyanide so that one eventually isolates
the complex NCCo(DH)₂NCSCH₃ as depicted in reaction 8.
N₃Co(DH)₂PPh₃

$$
\frac{p_{as}(N_3)}{p_{as}(N_3)} = \frac{p_{Ph_3}}{-p_{Ph_3}}
$$

N₃Co(DH)₂NCSCH₃
 $\frac{p_{as}(N_3)}{p_{as}(N_3)} = 2023 \text{ cm}^{-1}$
N₃CO(DH)₂NCSCH₃
 $\frac{p_{as}(N_3)}{p_{as}(N_3)} = 2023 \text{ cm}^{-1}$
(8)

This is not too surprising since several organic reagents are known to decompose thiocyanates to cyanides.⁶¹ One can follow the reactions of other heterocumulenes with the LCo- $(chelate)N₃$ complexes by observing the disappearance of the $v_{as}(N_3)$ stretching frequency in the infrared. Though the azide complexes completely reacted in each case no definitive well-characterizeable products could be isolated from these reactions. Surprisingly, whereas CS_2 reacted with^{21,25} plati $num, ^{24}$ copper, $62, 63$ and rhodium²⁶ azide complexes to produce thiocyanates, the azido cobalt complexes did not react with $CS₂$ either thermally or photochemically; in each case the azido complexes were recovered unchanged from these reactions.

VI. Reactions with Acids. Protonation of the azide ligand of $LCo(chelate)N_3$ complexes with acids leads to HN_3 elimination. Treatment with HCl solutions produces the analogous chloride complexes LCo(chelate)Cl. Similarly noncoordinating acids react to produce salts of the type $[Co(chelate)L]^+X^-$ (L $=$ PPh₃, py; $\bar{X} = ClO_4^-$ and HSO₄⁻).

Experimental Section

A. Reagents and Physical **Measurements.** Chemicals were reagent grade and were used as received or synthesized **as** described below.

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-
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Table V. Reaction Conditions and Physical Properties of the Alkyne Complexes Formed by Reaction of Terminal Alkynes with Cobalt Azido Chelate Complexes⁴ $\mathbb{R}^{1,1},\mathbb{C}_{2}\mathbb{C}_{2}\mathbb{C}_{2}\mathbb{N}_{3}$

 α RT = room temperature. pet. ether = petroleum ether.

Table IV. Reaction Conditions and Physical Properties of the Triazolato Products from the Reaction of Cobalt Azido Chelate Complexes with Alkynes

Table VI. Reaction Conditions and Physical Properties of the Products from the Reaction of Cobalt Azido Chelate Complexes with Nitriles^c

		%		dec		elemental anal.						
com-						$\%$ C		% H		$\%$ N		
plex	react conditions	vield	pptng solvent	pt, $^{\circ}$ C	color	calcd	found		calcd found	calcd	found	
9а	1 h, $RT,^b$ CH, Cl,	21	pet. ether	224	dark red brown	62.18	60.97		3.78 4.11	10.88	10.38	
10a	2 days, RT, CH ₂ Cl ₂		64 pet. ether with $Ph3P$		199 light brown powder		51.51 52.13^a 4.11 4.70 18.02				16.21	
	$10b$ 2 days, RT, CH, Cl,		52 ether with Ph ₂ P		213 dark brown powder	64.94	63.98		3.74 4.55 10.82		10.78	
	11a 30 days , RT, CHCl, 69 ether with Ph ₃ P				red brown		59.08 59.18		5.77 4.10 15.57		15.24	

CH₂Cl₂ adduct; see Table III for sturctures. ^b RT = room temperature. ^c Recrystallization solvent was CH₂Cl₂/petroleum ether.

All solvents were dried when necessary by standards procedures and stored over Linde 4-A molecular sieves. Melting points were determined on a Meltemp apparatus and are uncorrected. Elemental analyses were performed by Institut fur Anorganische Chemie, Munchen, and by Chemalytics, Inc. Tempe, AZ. Molecular weights have been determined with a Mechrolab osmometer. Infrared spectra were recorded on Perkin-Elmer 599 and 325 spectrometers as Nujol mulls between NaCl plates or as KBr disks. The 'H NMR spectra were recorded on CDCl₃ solutions relative to internal Me₄Si on Varian A 60 and Perkin-Elmer R24-B NMR spectrometers. The azide complexes were prepared as previously²² described.

B. Syntheses and Reactions. (1) Reactions with Alkynes. To a solution of the azide complex in a suitable solvent was added 2 molar equiv of the alkyne, and the solution was stirred magnetically at room temperature. When the azide stretching frequency could no longer be observed in the infrared spectrum of 1 aliquot taken from these solutions, a precipitating solvent was added, and the precipitate was recrystallized from a suitable solvent. The reaction conditions are given in Table **IV.**

The following complexes were obtained in a similar manner from reaction of 1.36 g (2m mol) of PPh₃Co(Salphen)N₃ with 10 mmol of alkyne in 50 mL of CHCl₃ upon addition of petroleum ether.

3a. From 1,4-dichlorobutyne after 65 days at room temperature was obtained 1.2 (65%) of $(CICH₂)₂(C₂N₃)Co(Salphen)PPh₃·CHCl₃$ Anal. Calcd for $C_{43}H_{34}Cl_5CoN_5O_2P$: C, 56.14; H, 3.73; N, 7.61. Found: C, 55.48; H, 4.02; N, 7.45.

4a. From propargyl alcohol after 80 days at room temperature was obtained 1.0 (50%) of H(HOCH₂)(C₂N₃)Co(Salphen)PPh₃. CHCl₃. Anal. Calcd for C₄₂H₃₄Cl₃CoN₅O₃P: C, 59.13; H, 4.03; N, 8.21. Found: C, 59.88; H, 4.72; N, 7.96.

Sa. From propargyl bromide after 80 days at room temperature was obtained 0.9 **g** (47%) of $H(BrCH_2)(C_2N_3)Co(Salphen)PPh_3$. CHC₁

(2) Substitution Reactions with Terminal Alkynes. To a solution containing the azide complex in a suitable solvent was added 2 molar equiv of the alkyne, and the solution was stirred magnetically until the azide stretching frequency was no longer observable in the infrared spectrum of 1 aliquot of the reaction solution. The reaction conditions and physical properties of the products are given in Table **V.**

(3) Liberation of Coordinated Triazolate. (a) Concentrated HCl. To a solution containing 0.74 g (1 mmol) of $(CH_3CO_2)_2(C_2N_3)$ Co-(DH),PPh, in 50 mL of acetone was added 5mL of concentrated HC1, and the solution was stirred vigorously for 2 days at room temperature. The resultant brown needles of $PPh_3Co(DH)_2Cl$ (mp 220 °C)⁶⁴ were removed by filtration, washed with absolute ethanol and petroleum ether, and air-dried. Reduction of the acetone filtrate on a rotary evaporator produced a green oil. The green oil was extracted with hot petroleum ether from which 33 mg (18%) of thin yellow crystals of **4,5-bis(methoxycarbonyl)-l,2,3-triazole,** mp 129-1 30 "C (lit.34965 133-134.5 °C) separated. Similarly, reaction of the triazolate complex with HBr produced the bromide complex $PPh_3Co(DH)_2Br$ and the triazole.

(b) Acetyl Chloride. Similarly, 0.74 **g** (1 mmol) of $(\text{CH}_3\text{CO}_2)_2$ - $(C_2N_3)Co(DH)_2PPh_3$ was stirred with 1 mL of freshly distilled acetyl chloride for 10 **min** at room temperature under moisture-free conditions to produce a green oil. The oil was pumped on under high vacuum to remove the acetyl chloride and extracted with 15 mL of absolute diethyl ether, leaving behind green needles of $(Ph_3PO)_2CoCl_2$. To the filtrate was added petroleum ether, and after cooling, a mixture

of the acetyltriazole and dimethylglyoxime diacetate precipitated. The mixture was separated by fractional crystallization from diethyl ether/petroleum ether in which the dioxime diacetate is the more soluble. In this way, 48 mg (21%) of $C_8H_9N_3O_5(\nu(C=0) 1790 \text{ cm}^{-1})$ was isolated. Hydrolysis of this acetyltriazole in warm moist petroleum ether afforded 25 mg (14%) of **4,5-bis(methoxycarbonyl)-l,2,3-triazole** $(mp 130-130.5 °C)$.

(4) Cycloaddition Reactions with Alkenes. Electron-poor alkenes such as diethyl fumarate, diethyl maleate, maleic anhydride, benzoquinone, and cyclopentadiene dimer were reacted with a variety of azido cobalt chelate complexes $LCo(chelate)N₃$ with the intent of preparing triazoline complexes. In each case the intended triazoline is thermally unstable. In only one case could an analyzable wellcharacterizeable product be isolated: that being the product of reacton of diethyl maleate and $NH₃Co(DH)₂N₃$.

A solution containing 1.74 g (5 mmol) of $NH₃Co(DH)₂N₃$ and 1.72 g (10 mmol) of diethyl maleate in 70 mL of Me₂SO was heated at 65 °C for 28 days. To a petroleum ether extract of the Me₂SO solution was added 140 mL of CHCl₃ producing a black oil. Dissolution of the oil in acetone followed by addition of petroleum ether afforded a light brown precipitate which was filtered, washed with petroleum ether, and dried for 3 h under high vacuum. There was obtained 0.66 g (25%) of $(EtO_2C)_2(H_2C_2N_3)Co(DH)_2NH_3CHCl_3$. Found: C, 32.25; H, 5.52; N, 18.02. Anal. Calcd for $C_{17}H_{28}Cl_3CoN_8O_8$: C, 32.01; H, 4.43; N, 17.57.

(5) Cycloaddition Reactions with Nitriles. An excess of an electron-poor nitrile was reacted with azido cobalt chelate complexes in a suitable solvent, isolated and recrystallized as described in Table VI.

(6) Reactions with Heterocumulenes. The azido cobalt chelate complexes reacted with a variety of heterocumulenes such as isocyanates, thiocyanates, and isothiocyanates to produce five-membered heterocycles by 1,3-dipolar cycloadditions that were usually not isolable because of the decomposition of the five-membered ring (see, for example, ref 60). From the reaction of $CH₃SC \equiv N$ with $PPh₃Co (DH)₂N₃$, there was obtained the cyano complex $CH₃SCNCo(D-1)$ H)2CN as follows: A solution containing 2.97 **g** (5.0 mmol) of $PPh_3Co(DH)_2N_3$ and 0.7 **g** (10 mmol) of CH₃SCN in 50 mL of CHCI, was refluxed for 51 days at which time the azide stretching frequency had disappeared. Addition of petroleum ether to the solution precipitated a yellow solid which was filtered, washed with petroleum ether, and air-dried; 66% CH₃SCNCo(DH)₂CN. Anal. Calcd for $C_{11}H_{17}CoN_6O_4S$: C, 34.02; H, 4.42; N, 21.65. Found: C, 33.58; H, 5.48; N, 19.90 (see reaction 8).

(7) Other Reactions. (a) Isocyanate complexes were prepared by metathesis of the corresponding chloro complex with NaNCO in $CHCl₃/CH₃OH$ or by heating a solution of the azide complex at 50 ^oC in CHCl₃ or THF under 40 atm of CO. In each case the isocyanate complex was recrystallized from CHCl₁. Anal. Calcd for $C_{27}H_{29}$ -CoN₅O₅P (12a, PPh₃Co(DH)₂NCO, dark brown): C, 54.64; H, 4.93; N, 11.80. Found: C, 52.95, 52.68; H, 4.94, 4.79; N, 12.09, 11.21. Anal. Calcd for C₁₄H₁₉CoN₆O₅ (12b, pyCo(DH₂)NCO, beige powder): C, 40.95; H, 4.66; N, 20.48. Found: C, 41.32, 41.06; H, 4.36, 5.33; N, 19.25, 18.47.

(b) Reactions with CS₂. A suspension of $\text{PPh}_3\text{Co}(\text{DH})_2\text{N}_3$ or $pyCo(BAE)N_3$ in 100 mL of CS_2 was stirred for 22 h. During this time no evolution of N_2 occurred, the azide stretching frequency was still present, and upon removal of the $CS₂$ on a rotary evaporator the complexes were recovered unchanged.

(c) **Reactions with Acids.** Treatment of the azide complexes with HCl liberated $HN₃$ and produced the chloride complexes as follows: HCI gas was bubbled through an acetone solution containing either 0.59 g (1 mmol) of $PPh_3Co(DH)_2N_3$ or $pyCo(DH)_2N_3$ to produce

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0.25 g (43%) (mp 208 °C) of dark brown crystals of PPh₃Co(DH)₂Cl or 0.25 **g** (57%) (mp 248 "C) of light brown crystals of pyCo(DH),CI.

Anal. Calcd for PPh₃Co(DH)₂Cl (C₂₆H₂₉ClCoN₄O₄P): C, 53.19; H, 4.98; N, 9.95. Found: C, 53.27; H, 5.45; N, 9.24. Molecular weight for $\text{PPh}_3\text{Co}(\text{DH})_2\text{Cl}$: calcd, 587.0; found (CHCl₃), 625.0.

Anal. Calcd for $pyCo(DH)_2Cl$ (C₁₃H₁₉ClCoN₅O₄): C, 38.68; H, 4.74; N, 17.35. Found: C, 38.64; H, 4.57; N, 16.93.

If these same complexes are left in contact with concentrated aqueous HCl for a long time, $CoCl₂·6H₂O$ precipitates.

Treatment of 0.53 g (1 mmol) of $PPh_3Co(DH)_2N_3$ or 0.41 g (1 mmol) of pyCo(DH)₂N₃ in 50 mL of acetone with 5 mL of 70% HClO₄ produces upon addition of ether $[PPh_3Co(DH)_2]^+ClO_4^-$ or $[pyCo(DH)_2]^+ClO_4$. Addition of NaBPh₄ in ethanol to the $[PPh₃Co(DH)₂]$ ⁺ClO₄ solution precipitates $[PPh₃Co(DH)₂]$ ⁺BPh₄⁻ as follows. Anal. Calcd for $C_{26}H_{29}ClCoN_4O_8P$ (0.51 g, 79%; $[PPh₃Co(DH)₂]$ ⁺ClO₄: C, 47.97; H, 4.50; N, 8.61. Found: C, 46.88; H , 4.78; N, 7.97. Anal. Calcd for C₁₃H₁₉ClCoN₅O₈ (0.33 g, 72%; $[pyCo(DH₂)]⁺CIO₄$: C, 34.41; H, 4.23; N, 12.35. Found: C, 34.07; H, 5.01; N, 11.94. Anal. Calcd for C₅₀H₄₉BCoN₄O₄ (0.54 g, 88%; $[PPh₃Co(DH)₂]+BPh₄$: C, 71.51; H, 5.89; N, 6.67. Found: C, 69.24; H, 5.04; N, 7.19.

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Registry No. la, 80462-1 1-9; **Ib,** 80462-10-8; **IC,** 80462-09-5; **Id,**

80462-08-4; **le,** 80462-07-3; If, 80462-06-2; **lg,** 80462-05- 1; **lh,** 80462-04-0; **li,** 80462-03-9; **lj,** 80462-02-8; **lk,** 80462-01 -7; **11,** 80462-00-6; **Im,** 80461-98-9; **In,** 80461-97-8; **lo,** 80462-40-4; **lp,** 80462-38-0; **lq,** 80462-37-9; **2a,** 80462-36-8; **2b,** 80462-35-7; **2c,** 80462-34-6; **2d,** 80462-33-5; **2e,** 80462-3 1-3; **2f,** 80462-30-2; **3a,** 80462-39-1 ; **4a,** 80462-29-9; **5a,** 80462-28-8; **6a,** 80462-27-1; **6b,** 80462-26-6; **6c**, 80462-25-5; **7a**, 80462-24-4; **7b**, 80462-23-3; 8a, 80462-22-2; **9a,** 80462-21-1; **loa,** 80471-14-3; **lob,** 80462-20-0; **lla,** 80462-19-7; **12a,** 771 72-84-0; **12b,** 30786-24-4; 1,4-dichlorobutyne-2, 821-10-3; propargyl alcohol, 107-19-7; propargyl bromide, 106-96-7; **4,5-bis(methoxycarbonyl)-** 1,2,3-triazole, 707-94-8; diethyl maleate, 141-05-9; C₈H₉N₃O₅, 80461-78-5; (EtO₂C)₂(H₂C₂N₃)Co(DH)₂NH₃, 80462-18-6; CH₃SCNCO(DH)₂CN, 80462-53-9; PPh₃Co(DH)₂CI, 23295-34-3; pyCo(DH)₂Cl, 23295-32-1; $[PPh_3Co(DH)_2]^+ClO_4^-$, 80462-52-8; $[pyCo(DH)_2]$ ⁺ClO₄⁻, 80513-21-9; $[PPh_3Co(DH)_2]$ ⁺BPh₄⁻, 80462-51-7; PPh₃Co(DH)₂N₃, 69879-86-3; MePPh₂Co(DH)₂N₃, 80462-49-3; Me₂PPhCo(DH)₂N₃, 80462-48-2; PBu₃Co(DH)₂N₃, 51194-39-9; $P(Bzl)$ ₃Co(DH)₂N₃, 80462-47-1; $P(OCH₃)$ ₃Co(DH)₂N₃, 61024-83-7; P(OPh)₃Co(DH)₂N₃, 70106-13-7; PhNH₂Co(DH)₂N₃, 66496-40-0; pyCo(DH)₂N₃, 56589-86-7; NH₃Co(DH)₂N₃, 66496-43128-64-9; $PBu_3Co(DPH)_2N_3$, 80462-45-9; $[Co[(DO)(DOH)]$ -39-7; 1-MeImCo(DH)₂N₃, 80462-46-0; AsPh₄[Co(DH)₂(N₃)₂], pn] $(N_3)_2$], 69879-56-7; PPh₃Co(Salen)N₃, 70106-20-6; PPh₃Co- $(Salphen)N_3$, 62858-89-3; cis-PPh₃Co(acac)₂N₃, 70100-51-5; pyCo(Salphen)N₃, 69907-81-9; CH₃CO₂C \equiv CCO₂CH₃, 762-42-5; $HC=CCO_2Et$, 623-47-2; $HC=CC(O)H$, 624-67-9; $HC=CC H(O_2CCH_3)_2, 67088-67-9; NCCF_3, 353-85-5; NCCO_2C_2H_5, 623-49-4;$ NCCH₂CN, 109-77-3.

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Electron Transfer. 51. Reactions of Riboflavin Radicals with Metal-Center Oxidants'

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Riboflavin undergoes successive one-electron reductions by Eu2+ or **V2+** in aqueous HC104, giving first the radical ion, **11, and then the dehydro compound, III.** Specific rates (21 °C, 0.12 M HClO₄) for the two V^{2+} reductions are 8.0 \times 10⁴ and 3.6 **X** lo4 M-l **S-I,** both far above the substitution-controlled upper limit for inner-sphere **V(I1)** reactions. Calculated formal potentials for the two reductions in 1 M H⁺ are 0.220 and 0.150 V. Rates of reactions of the radical cation Rbwith Hg²⁺, $(NH₃)$ ₅CoBr²⁺, and the *N*,*N*-dimethylnicotinamide derivative of $(NH₃)$ ₅Co^{III} (IV) are inversely proportional to $[H^+]$ (in the range of 0.7–1.3 M H^+), indicating that the flavin radical cation is undergoing partial deprotonation to the active reducing species. Reactivity patterns for reductions of (NH_3) ₅Co¹¹¹ complexes by the radical are devoid of those substituent effects that characterize inner-sphere reductions by transition-metal centers but resemble instead the pattern for reductions by the outer-sphere reagent $Ru(MH_3)_{6}^{2+}$. This similarity, in conjunction with the characterization of the oxidation product as riboflavin (with the **ring** system apparently unaltered), indicates that reductions by Rb. of these complexes follow outer-sphere paths. Specific rates of reaction of Rb. (25 °C, 1.3 M HClO₄) with the oxidizing metal centers Fe³⁺, T^{3+} , Cu^{2+} , Hg^{2+} , VO^{2+} , and UO_2^{2+} are not related simply to oxidation potential, ionic charge, electronic structure, self-exchange rates, or the accessibility of an oxidation state lying one unit below.

Among the oxidation coenzymes, flavin derivatives rank
second to none in importance and diversity of role.² Much
of their versatility reflects the ease with which the isoalloxazine
unit, a general feature of the class, unit, a general feature of the class, can participate in both onebetween electron-transfer processes and the even-electron processes characterizing metabolic conversions.

The impetus for the present work, which deals principally

tered electron-transfer species, was the expectation that flavins, because of their dual redox functionality, might catalyze re-

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