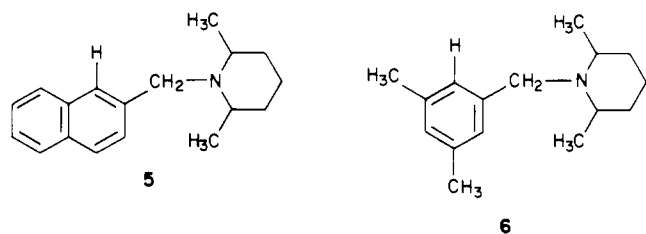


4 gave **3h** and **4h** but also varying amounts of tele substitution products **5** and **6**, respectively. Data from some representative experiments are set forth in Table I.



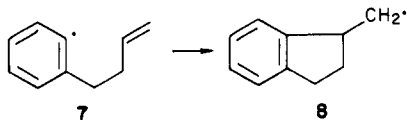
Concerning the first four entries, we note that the presence of di-*tert*-butyl peroxide causes the yield of **3h** from **3i** to increase from 37% to 63% while the presence of 1,1-diphenylethene or azobenzene lowers it respectively to 18% or 10%. Inasmuch as di-*tert*-butyl peroxide is a radical source and the other two added substances are radical and/or electron traps, these data fulfill expectations for a radical chain mechanism.

The four entries for substrate **3c** in Table I show some of the same features, but in muted form: a small augmentation of dehalogenation yield by di-*tert*-butyl peroxide, and a modest diminution by 1,1-diphenylethene. Tetraphenylhydrazine causes an increase in dehalogenation yield; conceivably it acts as a source of radicals (by homolysis to Ph₂N·). It is noteworthy that the yield of substitution product **5** wanes as the yield of **3h** waxes.

The four experiments in Table I concerning **4i** show it to be less reactive than **3i** in dehalogenation. Again acceleration by di-*tert*-butyl peroxide is evident. The crown ether¹² 12-crown-4, which chelates Li⁺ well, strongly inhibits deiodination.

The four experiments on **4b** manifest qualitatively the same effects as just discussed for **4i**, except that some tele substitution to form **6** now occurs.

Thanks to Beckwith and co-workers,^{13,14} we have available another tool to probe radical character, namely, the propensity of the *o*-(3-butenyl)phenyl radical (**7**), to cyclize to 1-indanyl-methyl radical (**8**). Reaction of *o*-(3-butenyl)iodobenzene¹⁵ with



2 in 2,6-dimethylpiperidine solution (24 h at room temperature) afforded 28% of 1-methylindan as well as 27% of 3-butenylbenzene, affirming the intermediacy of **7**. But the chlorine analogue, *o*-(3-butenyl)chlorobenzene, under the same conditions gave 3-butenylbenzene (10%) free of cyclization product. Thus a nonradical mechanism is indicated for it.

Dehalogenations of **3i** and of **3c**, through the action of **2** in *N*-deuterio-2,6-dimethylpiperidine solution, afforded **3h** free of deuterium at C-1 but deuterated in the methyl group (owing to base-catalyzed hydron exchange). These results inveigh against any mechanism that would involve 1-lithio-2-methylnaphthalene or the 2-methyl-1-naphthyl anion as an intermediate.

Our experimental findings are concordant with expectations from the mechanism of Scheme I insofar as aryl iodides are concerned. One aryl chloride result is contraindicative of that mechanism, while others are weakly in accord with it. We think that some nonradical mechanism of hydrodechlorination may experience an overlay of reaction according to Scheme I when a good source of radicals is present.

The remarkable retardation of radical chain dehalogenation by 12-crown-4 suggests that tight ion pairing if not actual covalent bonding with lithium is essential if some unidentified step(s) is to occur.

(12) Formally, 12-crown-4 is 1,4,7,10-tetraoxacyclododecane.

(13) Beckwith, A. L. J.; Gara, W. B. *J. Chem. Soc., Perkin Trans. 2*, 1975, 795.

(14) Meijs, G. F. Ph.D. Thesis, University of Adelaide, Australia, 1981.

(15) The *o*-(3-butenyl)iodobenzene was prepared by Dr. G. F. Meijs, to whom we are grateful.

Our observations furnish only meager indications of how tele substitution to form **5** or **6** occurs: In debromination of **3b** in *N*-deuterio-2,6-dimethylpiperidine at reflux (16 h), besides **3h** (48%) free of deuterium at C-1 we obtained 29% of **5** which carried deuterium at C-1. Data in Table I suggest the mechanism to be of nonradical character.

Metal Ion Complexation by Rhodium and Iridium Metallomacrocycles. The Preparation and X-ray Crystal Structures of

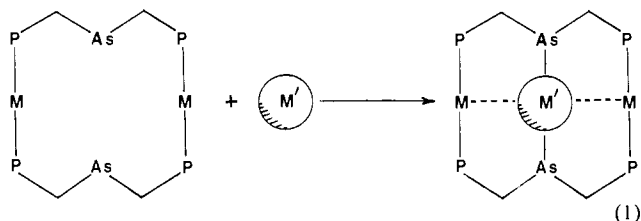
Rh₂(CO)₂Cl₂(μ-Ph₂PCH₂As(Ph)CH₂PPh₂)₂ and [Rh₂Pd(CO)₂Cl₃(μ-Ph₂PCH₂As(Ph)CH₂PPh₂)₂][BPh₄]

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The diverse reaction chemistry that has developed around diphosphine-bridged, binuclear complexes has expanded our knowledge of metal-metal bonding, given insight into the interaction of small molecules with two metal centers, and provided models for species bound to catalytically active metal surfaces.^{1,2} Polyfunctional phosphine ligands can be designed to produce more complex arrays of reactive metal centers which are expected to display multicenter metal-metal interactions and allow for multisite metal/small molecule bonding. The tripod ligand tris(diphenylphosphino)methane can stabilize or create new triangular arrays of metal centers^{3,4} while bis[(diphenylphosphino)methyl]phenylphosphine⁵⁻⁹ and 2,6-bis(diphenylphosphino)pyridine¹⁰ can be used to place three or four rhodium ions into nearly linear arrays. With metal ions other than rhodium dpmp forms six-membered chelate rings which cannot, as yet, be opened up to give linear chain complexes with metal ions in close proximity.¹¹ Thus, to date, this class of ligands has not yielded complexes that contain linear, trinuclear arrays with any transition metals except rhodium. Here we describe the creation of metallomacrocycles **1** using the tridentate ligand bis(diphenylphosphinomethyl)phenylarsine (dpma) and the subsequent complexation, as shown in eq 1, of a third, different metal ion to form



new trinuclear species **2** in which the central metal differs from the outer two. This procedure extends the range of trinuclear chains by allowing different metal ions to be placed in the center of the chain. In order to form **1** we take advantage of the

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