Free Radical Reactions of Organomercurials[†]

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I. Introduction

Organomercury compounds¹⁻⁸ have been known since the middle of the past century, and they were one of the first types of organometallic compounds studied. However, due to the low reactivity of the mercurycarbon bond, their application in synthetic organic processes was not important, and their utility was centered on the synthesis of other more reactive organometallics. This low importance was even more diminished when the Grignard reagents were discovered at the beginning of this century. However, in the past **20** years, organomercury compounds have again acquired interest in organic synthesis, in spite of their toxicity, in relation mainly to the solvomercuration reaction, which permits the preparation of functionalized organomercurials with high selectivity; these organometallics are adequate precursors **for** functionaliied organic compounds.

The numerous general methods for obtaining organomercury compounds are described in detail in *Hou* $ben-Weyl³$ and include, basically, substitution or addition reactions; they are shown as follows:

(a) Substitution reactions

$$
RH + HgX_2 \rightarrow RHgX + HX
$$

 $(R = alkvl, alkvnvl, aryl)$

 $RMet + HgX_2 \rightarrow RHgX + MetX$

 $RMet + RHgX \rightarrow R_2Hg + MetX$

 $(Met = Li, \frac{1}{2}Mg, \frac{1}{3}B, \frac{1}{3}Al, \frac{1}{3}Tl, \frac{1}{3}Si, \frac{1}{2}Zn, \frac{1}{4}Pb, ...)$

 $RX + Hg \rightarrow RHgX$

 $(X = Br, I, N_2^+, NHNH_2)$

(b) Addition reactions **⁵⁰⁶**

 $RCH=CH_2 + HgX_2 + HY \rightarrow RCHY-CH_2HgX$

 $(Y = OH, O₂H, OR', O₂R', OAc, NR'₂, NHCOR',$ N_3 , NO_2)

$$
N_3, N
$$

RC=CR + HgX₂ → RCX=CR(HgX)
(X = F, Cl, OAc, SCN)

$$
R \longrightarrow + HgX_2 + HY \longrightarrow R
$$

The reactivity³ of the prepared organomercury compounds can be summarized in the following way:

(a) Mercury-metal interchange (transmetalation)
\n
$$
RHgX + Met \rightarrow RMet + Hg + X^{-}
$$
\n
$$
R_2Hg + 2Met \rightarrow 2RMet + Hg
$$
\n(Met = Li, Na, K, Mg, Al, ...)

This reaction is particularly interesting for obtaining β -functionalized organometallic intermediates derived from alkaline-earth metals.⁹

(b) Mercury-hydrogen interchange (hydrogenolysis) by means of mineral acids or reducing agents, especially sodium borohydride m hydride
RHgX + HX \rightarrow RH + HgX₂

$$
RHgX + HX \to RH + HgX_2
$$

 $RHgX + H^{-} \rightarrow RH + Hg + X^{-}$

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[†]This review is dedicated to the memory of Professor Vicente Gômez-Aranda.

.IC& **Barhmga** was **born** In Tardiema, Spain, **h 1940.** He **obtained** his **Ph.D.** degree (soivomercuration of dlenes) at **the** University of Zaragoza in 1966 under the direction of Prof. V. Gomez-Aranda. Following this, he spent 3.5 years as a postdoctoral research fellow of the Max Planck Gesellschaft at the Max Planck Institut für Kohlenforschung, Mülheim a.d. Ruhr, Germany, in the group of Prof. H. Hoberg studying aluminum chemistry. In **1970** he took a position as a Research Associate at the University of Zaragoza, where he was promoted to Associate Professor in **1972.** In **1975** he moved to the University of Oviedo as Professor in Organic Chemistry, in the Department of Organometallic Chemistry, where he is now head of the group of organic synthesis. His major research interest is focused on the development of new synthetic methods in the area of heterocyclic chemistry and functionalized systems. He has published a number of publications in this area involving mercury salts and also organoalkaline and organomagnesium compounds.

Miguel Yus is currently Professor in Organic Chemistry at the University of Oviedo, Spain. He received his **B.S.** degree in **I969** and his Ph.D. degree in 1973 from the University of Zaragoza (under the direction of **V. G5nmz-Aranda** and **J.** Barluenga). Atler **2** years of postdoctorai study (Max Planck Institut fur Kohlenforschung, Mülheim a.d. Ruhr, Germany), he joined the Faculty of Chemistry of the University of Ovledo. **He** was a visiting scientist at ETH-Zirrich. CH **(1983).** at the University of Oxford, UK **(1984).** and at Harvard University (1985). His research interests are focused on new methodology in organic synthesis by means of organometallic reagents.

(c) Mercury-carbon interchange in the presence of a catalyst

on new methodology in organic synthesis by mean
etalli^c reagents.
Mercury–carbon interchange in the prese
alyst

$$
RHgX + CO + R'OH \xrightarrow{Pd^{II} cat.} RCO2R'
$$

$$
RHgX + CO \xrightarrow{AlBr_3} RCOR'
$$

$$
R_2Hg + R'COCl \xrightarrow{AlBr_3} RCOR'
$$

$$
RHgX + R'Cl \xrightarrow{Pd^{II} cat.} RR'
$$

$$
RHgX + CH2=CHR' \xrightarrow{Pd^{II}} RCH=CHR'
$$

(d) Mercury-halogen interchange (halode-
ercuration)
 $RHgX + Hal_2 \rightarrow RHal + XHgHal$ mercuration)

$$
RHgX + Hal2 \rightarrow RHal + XHgHal
$$

$$
(Hal = Br, I)
$$

More recently, different methods for the generation of carbenes and radicals from organomercury compounds have been developed. In the first case, the starting materials are haloalkylmercury compounds:8,10 ore recently, different methods for the generation
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ids have been developed. In the first case, the
ing materials are haloalkylmercury compounds:^{8,10}
RHgCH₂HaI - $(CH_2:)+RHgH$

$$
\mathsf{RHgCH}_{2}[\mathsf{Hal} \longrightarrow (\mathsf{CH}_{2}:) + \mathsf{RHgHaI} \xrightarrow{\qquad} \qquad \qquad \longrightarrow
$$

In the generation of radicals $11,12$ the starting materials are typical organomercury compounds; the thermal, photochemical, or chemical-mainly using a hydridetreatment of these organometallics yields the corresponding radicals: (Hal = CI, Br, I)
the generation of radicals^{11,12} the
pical organomercury compour
chemical, or chemical—mainly
nent of these organometallics
ing radicals:
 $RHgX \xrightarrow[or H^+]{heat or A\nu} R^* \frac{(1) \n $\left.\times \text{CN}\right.^2 \cdot \text{N} + \text{atom donor}}{(2)H-\text$$

$$
RHgX \xrightarrow{\text{heat or } h\nu \atop \text{or } H} R^* \xrightarrow{(1) \text{ or } CN} R
$$

From the above-described reactivity, it can be deduced that organomercury compounds are adequate precursors for carbanions (transmetalation), carbocations (halodemercuration), carbenes, or radicals. From these possibilities, the transformations of organomercury compounds, which involve intermediate radi*cals,* have acquired great interest in the past decade due to the importance in organic synthesis of radical reactions, which have been developed in the past years, above **all** in the important field of carbon-carbon bond formation. $11-17$

The present review considers the reactions of organomercury compounds, which occur through a radical mechanism, paying special attention to their potentiality in organic synthesis, that is, pointing out the applicability of these processes.

II. Radicals Generated Thermally or Photochemlcally

A. General Theoretical Comments

The thermal generation in solution of substituted benzyl radicals from substituted dibenzylmercury¹⁸⁻²³ or benzylmercury iodides²⁴ has extensively been studied from a theoretical and spectroscopic point of view. Thus, the kinetica of the decomposition of these mercurials has been investigated, concluding that they are processes of first order, $18-21$ the corresponding values of σ have been deduced (although these might be flawed because benzyl mercurials decompose by a chain mechanism), 22 and the resulting radicals have been studied by spin resonance spectroscopy (ESR).²⁵ This theoretical and spectroscopic study has been extended to diarylmercury²⁴ or dialkylmercury compounds, 26 fluoroalkylmercury derivatives,²⁷ and β -substituted organomercurials.28

B. Thermally Generated Radlcals

The most studied generation of radicals by heating²⁹ has been carried out starting from dibenzylmercury.

The reaction has no synthetic utility because a mixture of reaction products is usually obtained, 1,2-diphenylethane always being present. Thus, while dibenzylmercury reacts with acetic acid at 130 "C, yielding benzylmercury benzoate as the main product,³⁰ its treatment with nitrobenzene³¹ or nitrosobenzene³² leads to a mixture of different compounds (Scheme 1). When decomposition is performedin the presence of maleic anhydride, the mixture indicated in the Scheme 1 is obtained.33 The corresponding radical produced by reaction of the benzyl radical with the anhydride is postulated as an intermediate.

Perhaps the most studied reaction with the benzyl radical thermally generated from dibenzylmercury is the process with aromatic systems such as anthracene, $^{34-36}$ **9,1O-dihydroanthra~ene,~~-~~** pyridine,39 quinoline,39 isoquinoline, 39 and indene. 40 In all these cases the reaction is initiated by generation of the benzyl radical, which attacks the aromatic system, yielding a new radical; the final stabilization of this intermediate leads to the corresponding mixture of products. An indirect route for generating benzyl radicals is to thermally decompose dimethyl- 41 or diethylmercury⁴² in the presence of toluene.

The pyrolytic decomposition of diphenylmercury36~40~43 or phenylmercury acetate44 as a source of phenyl radicals and their coupling reaction with aromatic systems such as anthracene, 43 9,10-dihydroanthracene, 36 or indene⁴⁰ have also been described.

The pyrolysis of different dialkylmercury compounds in the presence of 2-propanol leads to the radical derived from the alcohol, which suffers disproportionation or oxidation, giving, in the case of diisopropylmercury, the products showed in Scheme **2.** The process, which can also be activated photochemically, has been studied by using 0-deuteriated 2-propanol; thus, deuterium incorporation has been found to take place in both radicals.45

In regard to cyclopentadienyl derivatives, the reaction of ethyl radicals-generated by heating diethylmercury-with ferrocene⁴⁶ and the formation of cyclopentadienyl radicals from substituted dicyclopentadienylmercury⁴⁷ have been studied. On the other hand, benzyl radicals—obtained by pyrolysis or photolysis of dibenzylmercury-have been used for trapping of nitrogen-^{48,49} and phosphorus-containing⁵⁰ radicals.

SCHEME 351

(R=Me, Et, L-Pr)

SCHEME $4^{53,54}$

SCHEME $5^{57,58}$

The addition of dialkylmercury compounds to tetracyanoethylene (TCNE) through a charge-transfer mechanism, which involves two radicals has been described (Scheme 3).⁵¹

When the thermal or photochemical decomposition of the mercurial is performed in the presence of carbon tetrachloride as a solvent, olefins and alkyl chlorides are obtained as reaction products through a chain radical process.⁵²⁻⁵⁴ Scheme 4 shows two examples as well as the proposed mechanism; when the alkyl substituents contain deuterium atoms at the β -position with respect to the mercury atom, an isotopic effect of 4.9 ± 0.1 (K_H/K_D) is observed.⁵³

The above-described reaction has also been performed in the presence of perchloroethane;⁵⁵ in both cases an in-depth mechanistic study relating the reaction conditions, structure of reagents, initiators, etc. has been carried out, concluding that both processes are mechanistically similar. Finally, the spontaneous decomposition of di-tert-butylmercury in carbon tetrachloride has been demonstrated to occur via a radical $mechanism.⁵⁶$

C. Photochemically Generated Radicals

The photochemically induced reaction of different primary or secondary alkylmercury chlorides or bromides 1 with several anions derived from nitro derivatives **2** leads to the corresponding products **3** through an $S_{RN}1$ type mechanism (Scheme 5 and Table 1).^{57,58}

The process is inhibited with di-tert-butyl nitroxide, it does not occur in the dark, and the reaction works only when the solvents are completely deoxygenated. Additional evidence for a radical mechanism is the

TABLE 1. Nitro Derivatives 3 from Organomercurials 1 and Nitronate Salts $2^{57,58}$

organomercurial 1		nitronate salt 2	nitro derivative 3
$\rm R^1$	x	\mathbf{R}^2	yield, %
$\rm (CH_2)_4 COCH$	Cl	Me ₂	56
	Br	Me,	68
	Cl	(CH ₂) ₅	60
PhCH ₂	Cl	$(CH_2)_5$	87
	Cl	Me,	100
$n\text{-}C_6H_{13}$	Сl	Me,	90
	Br	Me,	50
$c - C_6H_{11}$	Cl	Me,	76
	Cl	(CH ₂) ₅	84
i -Pr	Cl	Me,	63
сн,=снсн,	Сl	Me ₂	50
$^{\prime\prime}$ OMe	Cl	Me ₂	14.5
c -C ₅ H ₉ CH ₂	Cl	Me,	35
Me	Cl	Me,	$\mathbf{2}$
t-Bu	Cl	Me ₂	0
Ph	C1	Me ₂	0
(E) -Me ₃ CCH= CH	Cl	Me,	0
$2\text{-CH}_2\equiv$ CHCH ₂ OC ₆ H ₄	C1	Me,	0

SCHEME 65s*5g

SCHEME 757,58

SCHEME $8^{60,61}$

$$
\underline{t} - \text{BulkgCl} + \text{Nu} \xrightarrow{\text{hV}} \underline{t} - \text{Bulk} \xrightarrow{\text{hV}}
$$

above-described reaction (Scheme 5) starting from **(5** hexeny1)mercury chloride: the result of this process is a mixture of the corresponding cyclization products through the initially generated radical (Scheme 6).⁵⁹ The proposed mechanism is shown in Scheme 7.^{57,58}

As shown in Table 1, the reaction does not work when a tertiary organomercurial (i.e., tert-butyl) or arylmercury chloride is used. However, this problem can be overcome in the case of the tert-butyl mercurial by carrying out the reaction in the presence of a stoichiometric amount of 18-crown-6 (Scheme 8 and Table 2).60,61

The photochemically generated radicals from organomercurials 1 have been used with success to produce coupling reactions with heteroatoms, yielding carbonheteroatom bond formation.^{62,63} The products 5-8 obtained in this way are listed in Scheme 9 and Table 3.

^aIn all cases the potassium salt is used.

SCHEME $9^{62,63}$

As shown in Table **3,** when (5-hexeny1)mercury chloride is used as starting material, a mixture of products is obtained arising from the corresponding 5-hexenyl and the equilibrated cyclopentylmethyl radical intermediates. The proposed mechanism for products *5* and **6** is included in Scheme 10.

Radicals photochemically generated from organomercurials **9** can be added to olefins **10,** yielding the corresponding coupling products **11** through a nonstereospecific chain radical process (Scheme 11 and Table **4) *64**

TABLE 3. Coupling Products 5-8 from Mercurials 1 and Reagents $(R^2Y)_2$, R^3SH , R^4H al, and Y⁻M^{+62,63}

organomercurial 1				reagent		products 5-8	
R ¹	$\mathbf X$	R^2Y	R^3	$R4$ Hal	${\bf Y}^-{\bf M}^+$	yield, %	ref
(E) -Me ₃ CCH=CH	Cl	n -BuS				100	62
		PhS				100	62
		PhSe				95	62
		PhTe				29	62
				PhSO ₂ Cl		84	62
					(EtO) ₂ POK	76	62
					PhP(OBu)OK	84	62
					p -Me $PhCO2Na$ n -PrSO ₂ Na	81 75	62 62
			Ph			99	62
			$t - Bu$			100	62
			PhCH ₂			64	62
(Z) -HOCH ₂ C(Cl)=CH	C ₁		Ph			61	62
$CH2=CH$	Cl		Ph			100	62
		PhSe				91	62
(E) -EtC(OAc)=CEt	$_{\rm Cl}$		Ph			92	62
(E) -Me ₃ CCH=CH	AcO		Ph			100	62
			PhCH ₂			97	62
$Ph_2C = CH$	Br	MeS				100	62
					p-MePhSO ₂ Na	61	62
				Me ₂ CHI		50	62
(E) -Me ₃ CCH=CH	PhS					99 ^a	62
$Ph_2C = CH$	n -BuS PhS					100 ^a 100 ^a	62
$Ph_2C = CMe$	PhS					100 ^a	62 62
(E) -PhCH=CH	PhS					100 ^a	62
(E) -n-PrCH=CH	PhS					100 ^a	62
	Cl				(EtO) ₂ POK	56	62
					p -Me $PhSO2Na$	71	62
(E) -PhCH=CH	C1				p -Me $PhSO_2$ Na	21	62
			$n-Bu$			100	62
$CH_2=CHCH_2CH_2$	Cl	PhS				92	63
		PhSe				85	63
		PhTe				92	63
					p-MePhSO ₂ SePh	87 ^a	63
$CH_3CH_2)_4CH_2$	Cl	PhS				78	63
		PhSe				82	63
		PhTe				83	63
					p-MePhSO ₂ SePh	82 ^a	63
				PhSO ₂ Cl		46	63
t -BuCH ₂	$_{\rm Cl}$	PhS		CCl_3Br		56	63
		PhSe				74 86	63
		PhTe				78	63 63
					p-MePhSO ₂ SePh	75 ^a	63
i -Pr	Cl	PhS				100	63
		PhSe				$100\,$	63
c- C_6H_{11}	C1	PhS				65	63
		PhSe				72	63
$\mathrm{c}\text{-}\mathrm{C}_{5}\mathrm{H}_{9}\mathrm{CH}_{2}$	\rm{Cl}	PhS				86	63
		PhSe				84	63
	Br	${\tt PhS}$				43	63
		PhSe				53	63
		PhTe				45	63
					p-MePhSO ₂ SePh	48 ^a	63
$CH_2=CH(CH_2)_3CH_2$	$\rm Cl$	PhS				88 ^b	63
		PhSe				93 ^b	63
		PhTe				85 ^b	63
					p-MePhSO ₂ SePh	$81^{a,b}$	63
				PhSO ₂ Cl		54^b	63
			${\bf Ph}$			58^b	63
PhCH ₂	$\rm Cl$	PhS				15	63
		PhSe				72	63
		PhTe				80	63
	PhCH ₂	PhS			p-MePhSO ₂ SePh	68^a	63
		PhTe				8 100	63 63
n -Bu	$_{\rm Cl}$	PhS				85	63
	n -Bu	PhS				100	63

'Product of the type **5 is** isolated. bMixture of the cyclopentylmethyl and As-hexenyl derivatives in different ratios depending on the reaction conditions.

SCHEME 1164

SCHEME 12⁶⁴

The proposed mechanism for the reaction described in Scheme 11 is indicated in Scheme 12.64

The corresponding photostimulated addition to diarylethenes 12 has been studied when in the starting organomercurial 9 ($R^1 = t$ -Bu),⁶⁵ taking into account the influence of the substituents in the aryl groups. Thus, for $X = H a 1:1$ mixture of the products 13 and 14 is obtained, the mechanism being not a chain radical one. However, for $X = OMe$ the major product is 13, and for $X = NO₂$ the major product is 14; in these cases a chain radical mechanism has been proposed (Scheme 13).⁶⁵

Recently, the photochemically induced addition of cyclohexylmercury chloride (9 with $R^1 = c \cdot C_6 H_{11}$) to different substituted olefins 15 has been described.⁶⁶ The process, which leads to the products 16, takes place with partial retention in the configuration, and the highest stereospecificity has been found for $R^2 = I$ in the olefin 15 (Scheme 14 and Table 5).

^aThe compounds 10 with $R^2 = H$ have the *E* configuration. ^bThe products 11 with R² = H appear as an E/Z mixture. ^cThe only reaction product is the cyclopentylmethyl derivative. ^dThe organomercurial R^1_2Hg is used.

TABLE 5. Olefins 16 from Cyclohexylmercury Chloride and Alkenes 1566

	reactant 15			
stereo-				product 16
chemistry	х	\mathbf{R}^2	yield, %	E/Z ratio
E	CO ₃ Me	n -Bu ₃ Sn	20	36
Z	CO ₂ Me	n -Bu ₃ Sn	34	2.5
E	CO ₃ Me	n -Bu ₃ Sn	66	23
Z	CO ₂ Me	n -Bu ₃ Sn	70	2.1
E	CO ₂ Me		34	20
Z	CO ₂ Me		45	0.9
E	HgCl	Cl	28	4.3
z	HgCl	Cl	39	0.7
Z	Сl	Cl	70	0.8
E	Сl	Cl	63	0.7

The former results can be explained admitting that, in the case of a secondary organomercurial such as cyclohexylmercury chloride, the elimination of group X is faster than the establishment of equilibrium between I and II (Scheme 15).⁶⁶ When instead of the cyclohexyl group a more bulky one, such as *tert*-butyl, is present in the mercurial 1, the Ib configuration is very disfavored and the elimination step takes place through the configuration IIb (Scheme 15).⁶⁶

The addition of radicals-photochemically generated from organomercurials 1-to pyridine constitutes an adequate method for radical alkylation of systems of this type.^{67,68} The reaction can be carried out either by starting from the isolated organomercurial 1 or in situ starting from olefins and performing previous meth-

SCHEME 1566

	organomercurial 1				
R	X	yield, %	o/p ratio		
Et	Cl	64	2.0		
n-Bu	Cl	66	2.4		
$n - C_6H_{13}$	C1	83	a		
t-BuCH ₂ CH ₂	Cl	64	2.5		
t -BuCH ₂	Cl	54	1.9		
c -C ₅ H ₉ CH ₂	Cl	77	1.9		
i -Pr	Cl	72	1.6		
$c - C_6H_{11}$	Cl	69	3.1		
	Cl	90	4.1		
t -Bu	Cl	94	1.4		
$MeOCH_2CH_2$	CF ₃ CO ₂	73	2.0		
MeOCH(Me)CH ₂	CF_3CO_2	76	2.5		
MeOCH(Et)CH ₂	CF_3CO_2	78	$2.5\,$		
$MeOCH(n-Bu)CH2$	CF ₃ CO ₂	80	2.7		
$MeOCH(t-Bu)CH2$	CF_3CO_2	67	2.8		
MeOCH(Me)CH(Me)	CF ₃ CO ₂	81	1.9		
OMe	CF ₃ CO ₂	86	2.2		
NHCOMe	CF_3CO_2	69	2.3		
OMe	$\rm CF_3CO_2$	78	3.0		
^a Not given.					

TABLE 7. Acetylenes 19 from Organomercurials 9 and Substituted Phenylacetylenes 1869

SCHEME $16^{67,68}$

1 17 SCHEME 1769 $RHgCl$ + PhCaCX \longrightarrow PhC=CR **9 18 19**

The relative reactivity of acetylenes 18 with cyclohexylmercury chloride is as follows: $X = PhSO₂(65)$ $>$ I (19) $>$ n-Bu₃Sn (1.0), the mechanism being similar to that described for the radical reaction with substituted olefins.

Recently, the photostimulated radical addition of tert-butylmercury chloride to acetylenes **20,** which bear electron-withdrawing groups, has been described,

oxymercuration with mercury trifluoroacetate in methanol (Scheme 16 and Table 6).

The above-described reaction (Scheme **16)** has been extended to other systems such **as** substituted pyridines or N, N, N', N' -tetramethyl-p-phenylendiamines.^{67,68}

The reaction of photochemically generated radicals-from organomercury compounds-with olefins or aromatic compounds has been applied to phenylacetylenes **18** bearing another substituent such as iodine, phenylmercapto, phenylsulfonyl, or a mercury or a tin atom; therefore, products **19** are prepared in this way (Scheme 17 and Table **7).69**

TABLE 8. Olefins 22 and 23 from tert-Butylmercury Chloride and the Acetylenes 2070

	acetylene 20		product, ^a yield, %	
\mathbf{R}^1	\mathbf{R}^2	22	23	
н	Ph	63	41	
н	COMe	90	85	
CO ₂ Et	Ph	72	94	
CO ₃ Et	CO ₂ Et	97	96	
^a Mixture of Z/E isomers.				

SCHEME 18'O

yielding vinyl mercurials 21 ;⁷⁰ these organomercurials are transformed in situ into the corresponding reduced or iodinated systems **22** and **23,** respectively (Scheme **18** and Table

Finally, the photochemical decomposition of organomercurials bearing carboxymethyl groups leads to carboxymethyl radicals; the results, depending on the solvent used (benzene⁷¹ or tetrahydrof ur^2), yield different reaction products.

D. Heteroradicals

By thermal or photochemical decomposition of mercury carboxylates—either isolated or generated in situ by reaction of the corresponding alkaline carboxylates with mercury acetate-can be transformed, depending on the reaction conditions and the groups R, into mercurials arising from a mono- or didecarboxylation. The proposed mechanism includes the formation of an oxygenated radical of the type $RCO₂$ as is shown in Scheme **19.73** The same reaction is also known for sulfinic acid derivatives.⁷⁴ However, a similar radical mechanism in this case has not been postulated.

A convenient method for generating bis(trifluor0 methyl) nitroxide radicals is to heat mercury bis(trifluoromethy1)nitroxide at 85 **0C;75,76** this radical has been used in the synthesis of new organic and inorganic compounds, as well in the trapping of other radicals.⁷⁷

SCHEME 20

SCHEME 2184,85

Silyl radicals have been generated from bis(trimethylsilyl)mercury, and they have been used, for instance, in the silylation of aromatic systems such as substituted pyridines.^{78,79} The reaction of bis(trimethylsily1)mercury or the ethyl derivative with pyridines bearing a metal like silicon, germanium, tin, or lead,⁸⁰ arenes like benzene or toluene,⁸¹ and alkyl or acyl chlorides82 has also been studied. In all cases a trimethylsilyl radical **has** been proposed as a reaction intermediate. In a similar way, the reaction of bis(trialkylgermany1)mercury with aromatic compounds such as naphthalene, anthracene, or anisole,⁸¹ pyridine,⁸⁰ and alkyl or aryl chlorides 82 through the corresponding germanyl radicals has been studied.

Finally, compounds containing a mercury-tin bond have been used **for** generating stannyl radicals, which are adequate precursors for compounds with tin-tin bonds.⁸³

I I I. Radicals Generated by Means of Hydrldes

A. Radical Mechanism in the Reduction of Organomercurials with Hydrides

The reduction of organomercury compounds by means of metal hydrides, especially sodium borohydrides, has widely been studied from a mechanistic point of view, the mechanism being of a chain radical type (Scheme **20).**

The intermediates have been trapped **by** carrying out the reaction in the presence of air;^{84,85} in these cases, together with the expected oxygenated products, others, arising from a transposition process of the initially generated radical, are obtained (Scheme **21).**

Furthermore, the reduction of (5-hexenyl)mercury bromide with sodium borohydride leads to the expected products coming from the initially formed radical; on the other hand, the use of sodium borodeuteride instead of the corresponding hydride indicates an isotopic effect, which is based on the influence of winning a hydrogen or a deuterium atom for the radical intermediate from the species RHgX $(X = H \text{ or } D)$ (Scheme 22).^{86,87}

SCHEME 2286,87

SCHEME 23#'

SCHEME 2493

However, the most important contribution to the radical mechanism in the hydride-promoted reduction of organomercurials is in relation to stereochemical studies. $88-92$ Thus, for instance, in the reduction with sodium borodeuteride or in the reduction/oxidation of both isomers of norbornylmercury bromide, the same mixture of products was obtained, with loss in the initial stereochemistry (Scheme **23).**

When the generated radical is especially stable, for instance the benzyl radical, the obtained products depend on the hydride used, the reduction competing with the symmetrization reaction. In Scheme **24** the proposed mechanism for these processes is included.93

A similar radical mechanism to that described above has been proposed for the reduction with other reducing agents such as tin hydrides,⁹³⁻⁹⁵ lithium aluminum hydride,⁹⁶ or lithium naphthalenide.⁹⁷ In regard to the starting organomercurials, they can either lack functionality or be β -substituted.^{98,99}

In subsections C of this section, the synthetic potential of radicals from the hydride reduction of organomercurials is discussed.

SCHEME 25"

B. Theoretical Comments and Selectivity

The generation of radicals starting from organomercurials by means of metal hydrides, mainly sodium borohydride, together with their further use in the formation of carbon-carbon bonds by reaction with electron-poor alkenes has been called "the mercury method" (Scheme 25).¹¹ This methodology has extensively been studied from a theoretical point of view.¹⁴

Thus, the 5-hexenyl radical is easily generated from the corresponding mercurial, and it cyclizes to form the cyclopentylmethyl radical, the relative reaction rate of the both species with an alkylmercury hydride or an $electrophilic$ olefin-i.e., acrylonitrile or methyl acrylate-in a competition experiment has been investigated: the mentioned radicals are trapped by the hydride with a rate greater than 10^7 L \cdot M⁻¹ \cdot s⁻¹.¹⁰⁰

When the kinetics of the reaction between different primary, secondary, or tertiary radicals with several electrophilic olefins was studied, it was found that the radicals behave as nucleophiles, and the reactivity increases as follows: primary \le secondary \le tertiary.^{101,102} In the case of the cyclohexyl radical, an involved study was carried out considering the effect of the substituents of the electrophilic olefin on the reaction rate, 103-105 finding interesting correlations between the rate constant and the electronic effect of the substituent in the electron-poor alkene. This study has also been extended to α - and β -functionalized radicals; in this case the steric effects play an important role in the reaction rate. 106

Especially interesting is the kinetics in the reaction of radicals generated by "the mercury method" with substituted styrenes:^{107,108} an important correlation between the reaction rate constant and the corresponding σ values is found for the aromatic ring substituents.

Upon comparison of " σ radicals" (an unpaired electron in an sp^3 orbital) and " π radicals" (an unpaired electron in a p orbital) generated by the above-mentioned method, it was found that the former are as selective as the latter in the reaction with different electron-poor olefins. 109,110 The chemoselectivity of several σ - and π -radicals with the halogens chlorine and bromine in a mixture of carbon tetrachloride/bromotrichloromethane was explored; in general, the selectivity is drastically dependent on the temperature and steric effects: at **273** K the methyl radical is the least selective of the series, while at **403** K it appears as the most selective. $111,112$

Numerous studies have been carried out concerning the selectivity of different radicals prepared from organomercurials and sodium borohydride with electrophilic olefins. Thus, it has been shown that the trans-disubstituted olefins of the type $YCH=CHCO₂R$

SCHEME 26114

 N aB H ₄ CVHGOAC . \sim CVHaH \sim

SCHEME 27115

 $(R = Me, n - C_6H_{13}, C - C_6H_{11}, t - H)$

SCHEME 28116,117

react more rapidly than the cis isomer, this "cis effect" increasing with the size of the group Y .¹¹³. In the reaction of cyclohexyl radicals with systems of the type 24, it has been pointed out that the relative rate increases as follows: $Me_2C > MeN > NH > PhN > S >$ O (Scheme 26).¹¹⁴

The regioselectivity in the addition process of primary, secondary, or tertiary radicals to 2-methylmaleic anhydride shows that, in any case, the major reaction product is α, α' -disubstituted (Scheme 27).¹¹⁵ In relation to the stereochemistry of the reaction mentioned in Scheme 27, it has been observed that the major diastereoisomer has both substituents in the cis position (anti attack of RHgH). The selectivity is considerably increased with the bulkiness of the generated radical R. This can be easily explained by considering that the attack of the alkylmercury hydride takes place preferentially on the opposite side of where the alkyl

TABLE 9. Diastereoisomers 27 and 28 from Cyclic Olefins. Alcohols 25, and Electrophilic Olefins 26¹¹⁸

					product $27 + 28^a$	
	alcohol 25		olefin 26			27/28
cyclic olefin	R	X^1	\mathbf{X}^2	\mathbf{X}^3	yield, %	ratio
cyclopentene	Et	CN	н	CN	60	60/40
		н	Cl	CN	66	72/28
		н	н	$\mathbf{C}\mathbf{N}$	65	71/23
		н	Me	CN	46	77/23
		н	н	COMe	51	87/13
		н	н	CO ₂ Me	60	88/12
		н	н	Ph	15	90/10
	Me	н	н	CN	65	78/22
	i-Pr	н	н	$_{\rm CN}$	50	77/23
	t-Bu	н	н	$_{\rm CN}$	8	80/20
dihydrofuran	Me	CN	н	CN	20	64/36
		н	н	CN	45	86/14
		н	н	COMe	40	88/12
		н	н	CO ₂ Me	48	93/7
cyclohexene	Me	Н	н	CN	67	65/35
		н	н	CO ₂ Me	65	70/30
		н	н	Ph	12	75/25
dihydro- pyran	Me	$_{\rm CN}$	н	CN	69	58/42
		н	н	$_{\rm CN}$	76	66/34
		н	н	COMe	68	73/27
		н	н	CO2Me	64	75/25
^a Overall yield for the one-pot reaction.						

SCHEME 30120

group in the α -position of maleic anhydride is attached (Scheme 28). 116,117

Finally, the diastereoselectivity in the addition of cyclic β -substituted radicals, generated by solvomercuration-reduction of cyclic olefins such as cyclopentene, 2,3-dihydrofuran, cyclohexene, and 2,3-dihydropyran, to electrophilic olefins has been described. In all cases the trans diastereoisomer turned out to be the major one (Scheme 29 and Table 9).¹¹⁸

Recently, the mechanism of the 1,2-migration of vinyl or formyl substituents in free radicals generated via a mercuration-reduction tandem reaction was studied from a theoretical point of view.¹¹⁹

C. Applications in Organic Synthesis

Intramolecular Reaction with Olefins $\mathbf{1}$

Linalool undergoes cyclization through the tandem oxymercuration-reduction to give a complex mixture of products, where the major one (29) is probably formed by a radical mechanism via the corresponding

SCHEME 31¹²⁰

SCHEME 32lz1

SCHEME 33121

SCHEME 34¹²²

mercury hydride intermediate (Scheme **30).** However, it has not been possible to trap the corresponding radical by reaction with oxygen.¹²⁰ The same principle has been used to prepare the strobane structure **30** from

SCHEME 35^{123}

SCHEME 3612'

SCHEME 37125

$$
R^{1}HgOAc + X^{1}CH=CX^{2}X^{3} \xrightarrow{\text{NABH}_{4}} R^{1}X^{1}CH-CHX^{2}X^{3}
$$
\n34

\n26

\n35

epimanool via mercuration-reduction (Scheme 31).¹²⁰

The intramolecular formation of monocyclic spiranic systems has been described as shown in Scheme 32. Some examples of this strategy are included in Scheme 33.121

The reduction step can also be carried out by using tributyltin hydride, and the resulting radical can be trapped with molecular oxygen. Thus, this sequence has successfully been used in a synthesis of prostaglandins (Schemes 34^{122} and 35^{123}).

In relation to intramolecular reactions with amino mercurials, the cyclization of the adduct **31** is known to give the product **32** (Scheme 36). In this reaction, which constitutes a reasonable alternative to the same process promoted by palladium, no isomer **33** was detected. 124

2. Intermolecular Reaction with Olefins

In this section the intermolecular reaction of different unsubstituted and functionalized radicals-generated by reaction of organomercurials with sodium borohydride-with olefins will be considered.

(a) Unsubstituted Radicals. The reduction of different alkylmercury acetates **34** with sodium borohydride in the presence of electron-poor olefins **26** constitutes an adequate formation method for carboncarbon bonds, yielding the corresponding coupling products 35 (Scheme 37 and Table 10).¹²⁵

Starting organomercurials **34** are easily obtained from the corresponding organomagnesium compounds by reaction with mercury acetate. 3

The reaction included in Scheme 37 fails when **cro**tonic esters are used as electrophilic olefins. This problem has been overcome by employing 1,l-dicyano olefins **36,** as shown in Scheme **38.** This procedure is a convenient route to β , β -disubstituted carboxylic acids **38** by final hydrolysis of the dicyano derivatives **37** (Table 11). 126

The reaction mechanism included in Schemes 37 and **38** has already been described in Scheme 25.l' The

TABLE 10. Coupling Products 35 **from Organomercurials 34 and Olefins 26126**

organomercurial 34		olefin 26	product 35	
R^1	X^1	X^2	X^3	yield, %
t-Bu	н	н	CO ₂ Me	83
	н	н	COMe	70
	н	н	Ph	15
	н	Me	CO ₂ Me	83
	Me	н	CO ₂ Me	<1
c -C ₆ H ₁₁	н	н	CN	61
	н	н	CO ₂ Me	62
	н	Н	COMe	70
	н	Н	CHO	27
	н	н	Ph	45
	н	Me	CN	50
	н	Me	CO ₂ Me	84
	н	Cl	Сl	27
	Me	Н	CN	53
	Me	н	CO ₂ Me	24
	CO ₂ Me	н	CO ₂ Me	34
	н	н	$n\text{-}C_{6}H_{13}$	2
	н	Н	OEt	2
$n\text{-}C_6H_{13}$	н	н	CO ₂ Me	64
	н	н	COMe	51
	н	н	Ph	20
	н	Me	CO ₂ Me	31
	Me	н	CO ₂ Me	49

TABLE 11. Carboxylic Acids 38 from Organomercurials 9 and Dicvano Olefins 3612s

SCHEME 38¹²⁶

38

$$
SCHEME 39129
$$

$$
R^{1}R^{2}C=CH_{2} \xrightarrow{BH_{3}} (R^{1}R^{2}CH-CH_{2})_{3}B \xrightarrow{Hg(0AC)} R^{1}R^{2}CH-CH_{2}HgOAC
$$

39

$$
R^{1}R^{2}CH-CH_{2}CH(X^{1})CHX^{2}X^{3}
$$

NabH_a

$$
\begin{array}{c}\n\text{for } x \in \mathbb{R}^n, \\
\text{corresponding reduction products } R^1H \text{ are obtained as}\n\end{array}
$$

byproducts in this process. Another possibility of carrying out the coupling reaction of organomercurials and electrophilic olefins by generating the corresponding radicals consists in the formation of the starting mercurials by a tandem hydroboration¹²⁷-mercuration¹²⁸ process as shown in Scheme 39 and Table 12.129

The method described in Scheme 39 can be considered a convenient procedure for coupling between electron-rich and electron-poor olefins.¹²⁹

(b) a-Substituted Radicals. (a-Acetoxyalky1)mercury chlorides **41** can be prepared from ketones via mercuration of the corresponding hydrazones.¹³⁰ When these organomercurials **41** are used in the coupling process with electrophilic olefins **26** in the presence of

TABLE 12. Coupling Products 40 from Olefins 39 and 26¹²⁹

	olefin 39		olefin 26		product 40
\mathbb{R}^1	\mathbb{R}^2	X^1	X^2	X^3	yield, %
н	t-Bu	н	н	CN	47
н	$4 \cdot \text{MeC}_6\text{H}_4$	н	н	CN	50
н	2-AcOC ₆ H_4CH_2	н	н	CN	65
н	$2-\text{HOC}_6\text{H}_4\text{CH}_2$	н	н	CN	48
н	$BrCH_2CH_2$	н	н	CN	53
н	AcOCH ₂	н	н	CN	51
н	$TsOCH_2CH_2$	н	Н	CN	71
н	$EtO2C(CH2)8$	н	н	CN	57
н	EtO	н	н	$_{\rm CN}$	55
н	PhOCH,	н	н	$_{\rm CN}$	55
Me	Et	н	н	$_{\rm CN}$	65
Me	n-Pr	н	н	CN	57
Me	ClCH ₂	н	н	CN	50
	β -pinene	н	н	CN	53
н	$c\text{-}C_6H_9{}^a$	н	н	CΝ	54
		н	н	CO ₂ Me	47
		н	н	COMe	44
		н	$_{\rm Cl}$	$_{\rm CN}$	52
		н	Cl	СI	28
		н	Me	Ph	13
		CN	Me	CN	37
			oc-o-co	н	53
			OC-NH-CO	н	50

 $^{\circ}$ c-C₆H₉ = 3-cyclohexenyl.

SCHEME $40^{131-133}$

$$
R^{1}R^{2}C=0 \quad \frac{N_{2}H_{4}}{R^{1}R^{2}C=NNH_{2}} \quad \frac{(1) Hg(ORC)_{2}}{(2) KCl}
$$
\n
$$
R^{1}R^{2}CH=CX^{2}X^{3} \quad (26)
$$
\n
$$
R^{1}R^{2}CH=CX^{2}X^{3} \quad (26)
$$
\n
$$
R^{1}R^{2}C(ORC)CH(X^{1})CHX^{2}X^{2}
$$
\n
$$
41
$$
\n
$$
NABH_{4}
$$
\n
$$
42
$$

SCHEME $41^{134,135}$

$$
\begin{array}{ccc}\n\text{One} & \text{One} \\
R^1 \text{CH}-\text{CHR}^2 & + & X^1 \text{CH}=CX^2X^3 & \xrightarrow{\text{NABH}_4 \text{ or } } & \text{N^1}\text{CH}-\text{CHCH}(X^1)\text{CHX}^2X^3 \\
\downarrow \text{HgX} & & \text{26} & & \text{NABH(CN)}_3 & & \text{R}^2\n\end{array}
$$

43 44

SCHEME 42^{136,137}

$$
R^{1}R^{2}C=CHR^{3}
$$
\n(1) Hg(0AC)₂/R⁴OH (46)
\n(2) x¹CH=CX²x³ (26)/NABH₄ or NaBH(0Me)₃
\nOR⁴
\n
$$
R^{1}R^{2}C_{C-CHCH(X^{1})CHX^{2}X^{3}}^{1}
$$

sodium borohydride, the corresponding products **42** are isolated (Scheme 40 and Table 13).¹³¹⁻¹³³

47

The reaction indicated in Scheme 40 can be carried out without isolating the mercurial **41,** in a one-pot process, the yields being lower in the in situ procedure due to the resulting byproducts and the necessary purification of the final products **42.'32**

The selectivity of α -acetoxyalkyl radicals has been investigated: when the groups $R¹$ and $R²$ in the mercurial **41** are bulky, the reaction rate is lower with diethyl fumarate than with methyl acrylate.¹⁰⁶ Likewise, the relative nucleophilicity of different radicals of type **41** has been studied by using competition reactions with other primary, secondary, and tertiary radicals: the acetoxyalkyl radical turned out to be less nucleophilic than cyclohexyl radical, a logical consequence of the ability of the acetoxy group for attracting electrons. 133

TABLE 13. Coupling Products 42 from Organomercurials 41 and Olefins 26131-133

	organo-					
	mercurial 41		olefin 26		product 42	
\mathbf{R}^1	\mathbf{R}^2	$\overline{X^1}$	\mathbf{X}^2	$\overline{\mathbf{X}^3}$	yield, %	ref
Me Me	Me Et	н н	н н	CN $_{\rm CN}$	70 65	131, 132 131, 132
Me	n-Pr	н	н	CN	66	132
Me	i-Pr	н	н	$_{\rm CN}$	62	131
Me	t-Bu	н	н	CN	46	132
Et i-Pr	Et i-Pr	н н	н н	CN CN	71 33	131 131
t-Bu	t-Bu	н	н	CN	<2	132
	$-(CH2)4$ -	н	н	CN	72	131
	$-(CH2)5$ -	н	н	$_{\rm CN}$	72	131
		н	н	CN	77	132
Me	COMe	н	н	$_{\rm CN}$	<2	132
Me	Ph	н	н	$_{\rm CN}$	<2	132
Me	Me	н	н	CO ₂ Me	68	132
	$-(CH2)5$ -	н	н	CO ₂ Me	55	132
		н	н	CO ₂ Me	75	132
	$-(CH_2)_5$ -	н	н	COMe	43	132
		н	н	COMe	58	132
		$\ddot{}$				
Me	Me $-(CH2)5$ -	н н	н н	Ph Ph	13 12	.132 132
		н	$\mathbf H$	Ph	17	132
	$-(CH2)5$ -	н	Me	CN	49	132
		н	Me	CN	58	132
Me	Me	н	Me	CO ₂ Me	23	133
		н	Me	CO ₂ Me	51	132
Me	Me	н	C1	$_{\rm CN}$	72	132
		н	C1	CN	70	132
	$-(CH2)5$ -	н	Cl	Cl	27	132
		H	C ₁	$_{\text{Cl}}$	31	132
Me	Me	$_{\rm CN}$	н	CN	76	132
	$-(CH2)5$ -	$\mathbf{C}\mathbf{N}$	н	CN	69	132
		CN	н	CN	52	132
Me	Me $-(CH2)5$ -	CO ₂ Et CO_2Et	н н	CO ₂ Et CO ₂ Et	54 40	132 132
		CO ₂ Et	н	CO ₂ Et	61	132
		Me	н	CN	9	132
	$-(CH2)5$ -	oc-o-co		н	75	132
		$OC-O-CO$		н	57	132
		CO ₂ Me	Me	$\mathrm{CO_{2}Me}$	16	132
Me	Me	н	Cl	CO ₂ Me	54	133
		н н	MeO $_{\rm Eto}$	$\mathrm{CO}_2\mathrm{Me}$ $\mathrm{CO_{2}Et}$	5 52	133 133

(c) β-Substituted Radicals. β-Oxygenated and β -nitrogenated organomercurials, which are the most important β -substituted mercurials, are precursors of the corresponding radicals and will be considered separately.

TABLE 14. Coupling Products 44 from Organomercurials 43 and Olefins 2613'J3'

organomercurial							
	43			olefin 26		product 44	
\mathbf{R}^1	R ²	$\mathbf X$	$\mathbf{X}^{\bar{1}}$	$\overline{\mathbf{X}^2}$	X^3	yield, %	ref
$-(CH2)3$ -		Cl	н	н	CN	60	135
			н	н	CO ₂ Me	54	135
$-(CH2)4$ -		Cl	н	н	CN	77	134
$-(CH2)4$ -		Br	н	н	CN	66	135
$-(CH2)4$		Cl	н	н	CO ₂ Me	50	134
$-(CH2)4$ -		Br	н	н	CO ₂ Me	51	135
$-(CH_2)_4-$		Cl	н	н	Ph	22	134
			н	Me	CN	47	134
			н	Me	CO ₂ Me	38	134
			н	Cl	CN	65	134
			н	Cl	C1	25	134
			CN	н	CN ^a	61	134
			CO ₂ Et	н	CO ₂ Et ^a	53	134
			CO ₂ Et	н	CO_2Et^b	30	134
			Me	н	CN	8	134
			CO ₂ Et	Me	CO ₂ Et ^a	31	134
Ph	н	Cl	н	н	CN	$53 - 60$	134, 135
			н	н	CO ₂ Et	$42 - 44$	134, 135
			н	н	Ph	10	134
			н	Me	CN	43	134
			н	Me	CO ₂ Me	36	134
			н	Cl	CN	60	134
			CN	н	CN ^a	34	134
			CO ₂ Et	н	CO ₂ Et ^a	38	134
			CO ₂ Et	н	CO ₂ Et ^b	21	134
			Me	н	CN	6	134
			CO ₂ Et	Me	CO ₂ Et ^a	22	134
$n\text{-}C_6H_{13}$	н	Вr	н	н	CN	57	135
PhCH ₂	н	Cl	н	н	$_{\rm CN}$	55	135
			н	н	CO ₂ Et	46	135
Ph	Me	Cl	н	н	CN	46	135

^a The CN or CO₂Et groups are in a trans position. ^b The CN or $CO₂Et$ groups are in a cis position.

SCHEME 43136

$$
R^{1}CH=CHR^{3}
$$
\n(1) Hg(OAC)₂/R⁴OH (46)
\n(2) Triton X-100
\n(3) CH₂=CHX (48)/NABH₄
\n(48)/NABH₄
\n(5) CH₂=CHX (48)/NABH₄
\n(6) R¹CH₂CH₂X
\n(7) R³

(i) β **-Oxygenated Radicals.** The reaction of different $(\beta$ -alkoxyalkyl)mercury compounds 43, obtained by solvomercuration of the corresponding olefins¹⁻⁸ with electron-poor alkenes **26** in the presence of sodium borohydride or trimethoxyborohydride, gives the expected coupling products **44** (Scheme **41** and Table **14**).^{134,135}

The reaction described in Scheme **41** can be carried out in situ, without isolating the intermediate mercurials **43;** *so,* the starting materials are the corresponding olefins 45 (Scheme 42 and Table 15).^{136,137}

The yields in the coupling reaction described in Scheme 42, for β -alkoxy as well as for β -hydroxy mercurials, can be notably improved by using sodium borohydride **as** a reducing agent when a surfactant is employed **as** a phase-transfer catalyst in the reduction step (Scheme 43 and Table **16).135**

When mercuration is performed with an olefin that bears a nucleophilic group, an intramolecular addition can take place; the further coupling reaction with an electrophilic alkene in the presence of a hydride is exemplified in Scheme 44.¹³⁸

The former process has been applied to 1,3-dienic systems **50 as** starting educts **for** the mercuration step; the kinetically controlled mercuration leads to the

TABLE 15. Coupling Products 47 from Olefins 45 and 26 and Alcohols 46^{136,137}

	olefin 45		alcohol 46		olefin 26			
R ¹	\mathbb{R}^2	R^3	R ⁴	X^1	X^2	X^3	product 47 yield, %	ref
Η	H	H	Me	H	H	CO ₂ Et	50	136
H_{\rm}	Me	Me	Et	H	$\mathbf H$	CN	75	137
\overline{H}	$n-Bu$	Н	Me	H	H	CO ₂ Me	48	136
			Et	$\mathbf H$	H	CN	65	137
H_{\rm}	Ph	Н	Me	H	\overline{H}	CO ₂ Me	50	136
			Et	H_{\rm}	H	CN	48	137
Η	$-(CH2)3$ -		Me	H	$\mathbf H$	CO ₂ Me	65	136
			Et	H	H_{\rm}	CN	65	137
			Et	H	H	CO ₂ Me	60	137
			Et	H	$\rm H$	Ph	15	137
			Et	$H_{\rm 2}$	$_{\text{Cl}}$	CN	66	137
			$_{\rm Et}$	$\mathbf H$	Cl	C1	21	137
			Et	CN	$\mathbf H$	CN	66	137
			Et	CO ₂ Me	Me	CO ₂ Me	37	137
H	$-(CH2)4$ -		Me	н	H	CO ₂ Me	58	136
			Et	$\mathbf H$	$H_{\rm 2}$	CN	68	137
Me	н	Me	Me	H	H	CO ₂ Me	53	136
Me	Et	Н	Et	$\mathbf H$	H	CN	53	137
Me	$n\text{-}Pr$	H	Me	H	H	CO ₂ Me	30	136
Me	t -Bu	H	Et	H	H	CN	10	137
Me	Me	Me	Me	H	H	CO ₂ Me	32	136
			E t	H	H	CN	60	137

TABLE 16. Coupling Products 49 from Olefins 45 $(\mathbb{R}^2 = H)$ **and 48 and Alcohols 46135**

SCHEME 44^{138}

1,2-adducts, which are coupled in situ with electronpoor olefins **51,** yielding products **52** (Scheme **45** and Table 17).¹³⁹

In the field of carbohydrates the tandem mercuration-radical coupling has been used in the synthesis of branched sugars, starting from the corresponding unsaturated glycals **53** and *55* and employing the electrophilic olefin **26** (Scheme 46 and Table 18).'40

The solvomercuration-reductive coupling tandem process can be used to prepare lactones *58* and **60** starting from terminal olefins **57,** either by employing Triton X-100 as a surfactant (Scheme **47)136** or by previous isolation of the coupling products **59** (Scheme **48)141** (Table 19).

In the prior case, depending on the reaction conditions in the hydrolysis step, the corresponding γ - or

TABLE 17. Coupling Products 52 from Dienes 50, Olefins 51, and Methanol'39

		diene 50			olefin 51 product 52	
\mathbf{R}^1	\mathbf{R}^2	\mathbf{R}^3	\mathbf{R}^4	\mathbf{X}^1	\mathbf{X}^2	yield, %
н	н	н	н	н	CΝ	47
				н	CO ₂ Me	34
				н	COMe	22
				Me	CΝ	24
				Cl	CN	60
Me	н	H	Н	Cl	CN	59
н	н	Me	Me	Cl	$\mathbf{C} \mathbf{N}$	24
$-(CH2)2$ -		н	н		CN	27

SCHEME 45139

SCHEME 46¹⁴⁰

52

 $(R = CHX^1CHX^2X^3)$

SCHEME 47135

&lactones **(60** or *58)* can mainly be obtained (Scheme 48). Another possibility for obtaining lactones consists in carrying out the hydroxymercuration of the starting

TABLE 18. Branched Sugars 54 and 56 from Glycals 53 and 55 and Olefins 26140

		olefin 26			product 54 or 56	stereoselectivity
glycal	\mathbf{X}^1	\mathbf{X}^2	\mathbf{X}^3	MН	yield, %	$R_{\rm eq}/R_{\rm ax}$
53	н	Н	CN	n -Bu ₄ NBH ₄	60	67/33
	н	н	CN	n -Bu ₃ SnH	67	67/33
	н	н	CO ₃ Me	n -Bu ₃ SnH	55	71/29
	CN	н	CN	n -Bu ₃ SnH	55	90/10
	CO ₂ Me	н	CO ₃ Me	n -Bu ₄ NBH ₄	50	>97/3
	Me	$_{\rm CN}$	CN	n -Bu ₃ SnH	40	>95/ ₅
55	н	н	CN	n -Bu ₃ SnH	72	67/33
	CN	н	CN	n -Bu ₃ SnH	40	>95/ ₅

TABLE 19 Lactones 58 and 60 from Olefins 57 and Methyl Acrylate or Acrylonitrile^{135,141}

^a Based on the starting olefin 57. ^b Based on the coupling product 59. ^c Acetic acid was used in the mercuration step.

SCHEME 48141

SCHEME 49142

SCHEME 50^{142}

olefin 45 followed by the coupling reaction of the isolated mercurial 61 with the electrophilic alkene 48 and final cyclization of product 62, to give the corresponding lactone 63 (Scheme 49 and Table 20).¹⁴²

TABLE 20. Lactones 63 from Olefins 45 and 48¹⁴²

^aBased on the organomercurial 61. b Based on the coupling</sup> product 62.

SCHEME 51143

This last strategy has successfully been used in the synthesis of the antibiotic (\pm) -maiingolide 68,¹⁴² as shown in Scheme 50. Thus, starting from the allylic alcohol 64 and through the not isolated intermediate 65, the corresponding hydroxymercuration is carried out, yielding the mercurial 66 (45% overall yield), which is coupled with methacrylonitrile to afford a 1:1 mixture of diastereoisomers 67 (49% yield); the final cyclization leads to the expected 1:1 mixture of the wanted antibiotic 68 and its diastereoisomer 69, which is easily separated by chromatography.

However, in the case of the mercurated lactones 70-72, the coupling reaction gives poorer yields in products 73–75 than when the corresponding iodine or selenium derivatives are used (Scheme 51 and Table $21).¹⁴³$

(ii) β -Nitrogenated Radicals. The tandem aminomercuration-reductive coupling has been studied far less than the corresponding oxymercuration process. Thus, the intra-¹³⁸ or intermolecular¹⁴⁴ amino-

TABLE 21. Coupling Products 73-75 from Organomercurials 70-72¹⁴³

organomercurial	electrophilic olefin	coupling product		
no.		no.	yield, %	
70	н	73a	6	
	Me	73b	18	
71	H	74a	20	
	Me	74b	18	
72	Н	75a		
	Me	75b		

SCHEME 52138,144

SCHEME 53145

78

SCHEME 54^{145,147,148}

mercuration is used in the first step in the generation of the amino mercurial intermediate, which is coupled in situ with an electrophilic olefin, yielding the corresponding products 76 or 77; further reduction of the products 77 is an interesting method for the synthesis of amino alcohols or diamines 78 (Scheme 52 and Table 22).¹⁴⁴

The above-described reaction can alternatively be carried out by isolating the starting amino mercurials; the coupling process with compounds 79 and electronpoor olefins 51 is shown in Scheme 53 and leads to the expected products 80 (Table 23).¹⁴⁵

The coupling reaction fails when carried out starting from β -nitro or β -azido mercurials.¹⁴⁵ However, the use of β -amido mercurials, either isolated or generated in situ, leads to the expected products. Thus, the reaction of β -acetamidomercury compounds (obtained by acetamidomercuration of olefins with acetonitrile and mercury nitrate¹⁴⁶) with electrophilic olefins 51 in the presence of a sodium borohydride leads to the coupling products 82 (Scheme 54 and Table 24).^{145,147,148}

In the case of the coupling products 83 derived from acrylonitrile, the corresponding systems have been used for the preparation of pyrrolidines 84 (Scheme 55).¹⁴⁷

TABLE 22. Functionalized Amines 77 and 78 from Olefins 45 ($\mathbb{R}^2 = H$) and 48^{144}

olefin 45		olefin 48	product 77	product 78			
R ¹	\mathbf{R}^3	x	yield, ^a %		yield, ^ª %		
H	n-Bu	CΝ	39	CH ₂ NH ₂	91		
		CO ₂ Et	31	CH ₂ OH	85		
н Ph		CN	38	CH ₂ NH ₂	85		
		CO ₃ Et	31	CH ₂ OH	80		
Н	PhCH,	CΝ	36	CH,NH,	86		
		CO ₂ Et	30	CH ₂ OH	80		
$-(CH_2)_3-$		CΝ	44 ^b	CH ₂ NH ₂	89		
		CO ₂ Et	35 ^b	CH ₂ OH	86		
$-(CH2)4$ -		CN	13(47 ^b)	CH ₂ NH ₂	87		
		CO ₂ Et	39 ^b	сн,он	85		

^aBased on the starting olefin 45. ^bTriton X-100 is used as a phase-transfer catalyst.

SCHEME 55^{147}

Other amidomercury compounds used in coupling processes are mercurated urethanes like 85 prepared by intramolecular amidomercuration¹⁴⁹ of unsaturated ure than es 86: the corresponding coupling reaction with acrylic derivatives leads to the products 87, which are adequate precursors for nitrogen-containing heterocycles 88. The tandem amidomercuration-coupling can be carried out in situ, as is shown in the case of 87b (Scheme 56).¹⁵⁰

The process described above has successfully been applied to the synthesis of piperidinic systems 91 starting from the corresponding unsaturated urethanes 89 via a tandem mercuration-reductive coupling, the mercurial 90 being the intermediate. The resulting products are also adequate precursors for alkaloid type molecules like 92 (Scheme 57).¹⁵¹

SCHEME 58^{152}

TABLE 25. Coupling Products 97 from Olefins 45 ($R^2 = H$ **)** and 48 and Amides 96155

olefin 45		amide 96	olefin 48	product 97		
\mathbf{R}^1	\mathbf{R}^3	\mathbf{R}^4	x	yield, %		
н	н	MeO	CN	32 ^a		
Me	н	Me	CN	30		
		MeO	CN	68		
n-Bu	н	Me	CΝ	77		
		Me	CO ₃ Me	50		
		MeO	CN	71		
		MeO	COMe	40 ^a		
		MeO	CO ₂ Me	61 ^a		
$-(CH_2)_4-$		н	CN	40		
		Me	CΝ	44		
		MeO	CN	63		
		NH,	CN	57		

^aA mixture of Hg(NO₃)₂ and HgO (2:1) was used in the mercuration step.

^aThe CN or CO₂Et groups are in a trans position. ^bThe CN or $CO₂Et$ groups are in a cis position.

SCHEME 59155

Recently, the in situ amidomercuration-coupling tandem reaction has been used in the key step of the preparation of the immunosuppressant tripeptide 95, starting from the lactams 93 via the intermediate 94 (Scheme 58).¹⁵²

A general method for the addition of carboxamides, ureas, and urethanes 96 to unactivated olefins 45 consists in the use of mercury nitrate.^{153,154} When this method is combined with the in situ reductive coupling reaction employing electrophilic olefins 48 as reagents, the corresponding products 97 are obtained (Scheme 59 and Table 25).¹⁵⁵

TABLE 27. Coupling Products 102 from Cyclopropanes 101, Olefins 26, and Methanol or Acetic Acid¹⁵⁷⁻¹⁵⁹

		cyclopropane 101			solvent		olefin 26		product 102	
$\overline{R^1}$	R^2	${\bf R}^3$	R ⁴	R^5	\mathbf{R}^6	$\overline{X^1}$	$\overline{X^2}$	$\overline{X^3}$	yield, %	ref.
H		$-(CH2)4$ -	$\overline{\mathbf{H}}$	$\overline{\mathbf{H}}$	${\bf Me}$	$\mathbf H$	\overline{H}	$\overline{\text{CN}}$	82	157, 158
					Me	$\, {\rm H}$	$\mathbf H$	CO ₂ Me	$74\,$	158
					Me	$\, {\rm H}$	$\mathbf H$	Ph	30	158
					Me	$\, {\rm H}$	${\bf Me}$	${\rm CN}$	70	158
					Me	Н	Me	CO ₂ Me	60	158
					Me	$\mathbf H$	Cl	CN	84	158
					${\bf Me}$	$\mathbf H$	C1	Cl	38	158
					Me	CN	$\mathbf H$	CN	87	158
					${\bf Me}$	CO ₂ Et	$\mathbf H$	CO ₂ Et	70	158
					Me	CO ₂ Et	CO ₂ Et	H	$33\,$	158
					Me	Me	н	\mbox{CN}	18	158
					Me	Me	$\mathbf H$	CO ₂ Me	$11\,$	158
					Me	CO ₂ Et	Me	CO ₂ Et	$50\,$	158
					Me	Η	Н	CN	87	157, 158
					Me	$\rm H$	Me	CO ₂ Me	$50\,$	158
					Me	CO ₂ Et	н	CO ₂ Et	68	158
					Me	CO ₂ Et	Me	CO ₂ Et	74	158
$n - C_6H_{13}$	$\, {\rm H}$	$\mathbf H$	$\mathbf H$	Н	Me	Η	н	CN	$50\,$	157, 158
					Me	н	Me	CO ₂ Me	$33\,$	158
					Me	CO ₂ Et	н	CO ₂ Et	$55\,$	158
					Me	CO ₂ Et	Me	CO ₂ Et	$40\,$	158
H_{\rm}		$-(CH2)4$ -	н	н	Me	Н	н	CN	82	158
					Me	Н	Me	CO ₂ Me	60	158
					Me	CO ₂ Et	н	CO ₂ Et	$70\,$	158
					Me	CO ₂ Et	Me	CO ₂ Et	50	158
Et	$\mathop{\mathrm{Et}}$	н	$\mathbf H$	$\mathbf H$	Me	H	н	$\rm CN$	64	157, 158
Me	Me	Me	$\, {\bf H}$	H	Me	н	Η	CN	81	157, 158
					Me	H	Me	CO ₂ Me	$50\,$	158
					Me	CO ₂ Et	Н	CO ₂ Et	$71\,$	158
					Me	CO ₂ Et	Me	CO ₂ Et	50	158
Me	н	Me	н	Me	Me	Н	н	CN	$20\,$	157, 158
${\bf Me}$	Me	Me	Me	H	Me	H	$\, {\rm H}$	CN	56	157, 158
Ph	Н	H	Н	н	MeCO	H	$\, {\rm H}$	\mbox{CN}	62	159
					MeCO	Η	Me	CN	41	159
					MeCO	н	Cl	CN	65	159
					MeCO	CO ₂ Et	$\mathbf H$	CO ₂ Et	80	159
					MeCO	CO ₂ Et	Me	CO ₂ Et	45	159
$n\text{-}C_6H_{13}$	Η	Н	Н	Н	MeCO	н	$\, {\rm H}$	CN	40	159
					MeCO	CO ₂ Et	$\rm H$	CO ₂ Et	40	159
$\, {\rm H}$		$-(CH2)4$ -	$\mathbf H$	Η	MeCO	Н	$\mathbf H$	CN	65	159
					MeCO	H	Me	CN	46	159
					MeCO	Н	$\rm Cl$	CN	81	159
					MeCO	CO ₂ Et	${\bf Me}$	CO ₂ Et	46	159
Me	${\bf Me}$	Me	H	H	MeCO	н	$\, {\bf H}$	CN	45	159
					MeCO	$\mathrm{CO_{2}Et}$	$\mathbf H$	CO ₂ Et	41	159
Me	Me	Me	Me	н	MeCO	H.	H	CN	40	159

SCHEME 60156

SCHEME 61157-159

(d) γ -Substituted Radicals. γ -Substituted organomercury compounds are easily prepared by oxymercuration of cyclopropanes 98,³ and their isolation is usually carried out as the corresponding chloromercury derivatives 99. The reductive coupling of these systems with electron-poor olefins 26 leads to the expected coupling products 100 (Scheme 60 and Table $26)$. 156

The process can be carried out in a one-pot manner without isolation of the organomercury intermediates. In this case the reaction has been studied for the methoxy-157,158 and acetoxymercuration products.¹⁵⁹ Scheme 61 shows the mentioned reaction performed with the starting cyclopropanes 101, in methanol or acetic acid as solvent, and an electrophilic olefin 26, in which the expected coupling products 102 are isolated $(Table 27)$.

Other γ -substituted organomercury compounds are the so-called homoenolates 105, which are available from aldehydes or ketones 103 by successive silylation,¹⁶⁰ cyclopropanation,¹⁶¹ and final mercuration of the isolated silylated cyclopropanols 104 .¹⁶² When the organomercury compounds 105 are allowed to react in situ with an electrophilic olefin 26 and sodium borohydride,

SCHEME 62160-164

SCHEME 63164

SCHEME 64^{165}

 R^1 HgCl + R^2 C=CR³ $\frac{NABH_4}{N}$ R^1R^2 C=CHR³ \bullet 108 45

the corresponding coupling products 106 are obtained (Scheme 62 and Table 28).^{163,164}

In the case of the aldehyde derivatives (103 with $R¹$ $=$ H), described in Scheme 62, it is necessary to treat the reaction product with potassium fluoride at the end of the reaction in order to get the final desilylation.

An alternative to the reaction described above for aldehyde derivatives consists in the hydroxymercuration of the corresponding silylated cyclopropanols (104 with $R¹$ = H) before the second reaction step. In this case a mixture of reaction products 106 and 107 is obtained,

TABLE 29. Coupling Products 106 $(R^1 = H)$ and 107 from Silylated Cyclopropanols 103 (\mathbb{R}^1 = H) and Olefins 26¹⁶⁴

cyclopropanol 103		olefin 26		product $106 + 107$	
\mathbf{R}^2	\mathbf{R}^3	\mathbf{X}^1	\mathbf{X}^2	X^3	yield, %
n-Bu	н	н	н	CN	55
		н	н	CO ₂ Et	50
		н	Cl	CN	35
		CO ₂ Et	н	CO ₂ Et	36
Me	Me	н	н	CN	47
		н	н	CO ₂ Et	45
		н	Cl	CN	45
		CO ₂ Et	н	CO ₂ Et	45

TABLE 30. Coupling Products 45 from Organomercurials 9 and Alkynes 108

SCHEME 65166

 $\begin{picture}(150,10) \put(0,0){\line(1,0){10}} \put(15,0){\line(1,0){10}} \put(15,0){\line($ $h\nu$ RHg' $RHq =$ RH \longrightarrow R $+$ H_2 RH \longrightarrow RH $+$ R' PHA ^{*} - \blacksquare Have \blacksquare

which arises from the corresponding equilibration of the radical intermediates (Scheme 63 and Table 29).¹⁶⁴

3. Reaction with Acetylenes

The reaction of radicals generated by the "mercury" method" with acetylenes has been the subject of a sole publication.¹⁶⁵ When different primary, secondary, or tertiary organomercury compounds 9 are treated with several alkynes 108 in the presence of sodium borohydride, a Z/E mixture of the corresponding coupling products 45 is obtained (Scheme 64 and Table 30).

TABLE 31. Products from the Mercury-Photosensitized Decomposition of Sulfur-Containing Compounds^{167,168}

starting material					product ratio, %				
	EtH	Me ₂ S ₂	MeSH	MeH	MeS ₃ Me	Me ₂ S	H_2S	H ₂	
Me ₂ S Me ₂ S ₂ MeSH	40.6 13.4 17.5	28.8 26.5 13.1	11.8 22.0	3.0 19 6.6	25.9	13.8	38.5	13.6	
$\rm \bf SCHEME$ 66 ¹⁶⁷					SCHEME 68^{172}				
$Me2$ \longrightarrow MeS	$+$	$Hg \xrightarrow{h\nu} Hg^* \xrightarrow{Me_2S} Me_2S^*$ Me	Hq		OMe $Ph \rightarrow$ $\ddot{}$ ACOHG	${\tt MeOH/BF}_3$	OMe $-\bullet$ Ph \rightarrow MeÒ	$Hg_2(0Ac)_2$	

SCHEME 66Is7

SCHEME 67¹⁶⁹⁻¹⁷¹

The relative reaction rate of the intermediate radical with alkenes and alkynes was studied, concluding that the last reagents react 3.0-5.2 times more slowly than the alkenes. These results contrast with the addition of nucleophiles to both unsaturated systems.

I V. Radlcals Generated by Other Methods

A. Radlcals Generated by Excited Mercury

Although of no synthetic interest, the reaction of photosensitized mercury with 3-methylpentane glass at 5-77 K leads to radicals, through the corresponding alkylmercury hydride species (Scheme 65).166

Likewise, the reaction of photosensitized mercury with different sulfur-containing compounds such as dimethyl sulfide,167 dimethyl disulfide,167 and methyl or ethyl mercaptan168 has been studied by mass spectrometry (Table 31). Scheme 66 shows the proposed mechanism for the case of dimethyl sulfide.¹⁶⁷

B. Electrochemlcally Generated Radlcals

Depending on the reaction conditions, the formation of a mixture of products has been observed in the electrochemical reduction of organomercury compounds. This process can easily be explained by considering the existence of an alkyl- or arylmercury radical intermediate. This species would be generated in the first phase of the process (first polarographic wave), and a second step (second polarographic wave) would give the reduction products. In an alternative way, the generated radical can suffer disproportionation to afford symmetrization products. However, in these processes the corresponding dimer has never been observed (Scheme 67).¹⁶⁹⁻¹⁷¹

C. Radlcals Generated by Autoxidation

The oxidation of organomercury compounds by means of mercury(I1) salts, the so-called "autoxidation",

SCHEME 69"2

constitutes a method of interest for the substitution of a mercury atom by a nucleophile (Scheme **68).172**

The proposed mechanism for this process is of a radical type, this conclusion being based on the fact that the reaction is inhibited by means of oxygen and is capable of polymerizing acrylonitrile (Scheme 69).¹⁷²

0. **Radicals Generated by Means of Metalilc Salts**

More general than the case of the "autoxidation" (section **C)** is the use of other metallic salts for the generation of radicals starting from dialkylmercury compounds, for instance, iridium(1V) **salts;** the obtained radicals have been the object of broad theoretical, spectroscopic, and chemical study.^{173,174} The proposed mechanism is shown in Scheme 70: as can be seen, in the first step a charge transfer between the iridium salt and the organomercury compound is generated. Recently, the existence of stable complexes of this type has been demonstrated, such **as** the complex prepared from the 1:1 complex mercury (II) trifluoroacetate-**EDTA** and hexamethylbenzene; its structure has been analyzed by X-ray diffraction.¹⁷⁵

On the other hand, organomercury compounds have been used **as** initiators or accelerators in radical polim-

SCHEME 711'*

SCHEME 721s0,1s1

erizations of olefins¹⁷⁶⁻¹⁷⁸ and dienes,¹⁷⁹ such as sty- $\text{rene},^{176}$ methyl acrylate, $^{176-178}$ vinyl acetate, 176 acrylonitrile,¹⁷⁸ or butadiene.¹⁷⁹ For instance, the corresponding process using copper(II) chloride **as** a reagent for the generation of the initiator, in this case, the methyl radical from dimethylmercury, is shown in Scheme **71.178**

E. Radicals Generated by Haiodemercuration

Radical intermediates have been proposed in reactions of organomercury compounds, bearing the metal atom on an sp3-hybridized carbon atom, with bromine^{180,181} or iodine,¹⁸² based on stereochemical data. The reaction products are in **all** casea the corresponding alkyl halides (Scheme **72).** In polar solvents, a competing nonradical halodemercuration can occur.183

However, when the organomercury compound bears the metal atom on an sp^2 -hybridized carbon atom, the corresponding halodemercuration gives different stereochemical results depending on the solvent used. For instance, in the bromodemercuration in pyridine, a retention in the configuration is observed, whereas in carbon disulfide the main process occurs with inversion. Taking these facts into account, a radical mechanism does not seem to be general for the mentioned pro $cess.¹⁸⁴$

V. Conclusions

From the chemistry described in this survey it can be conclused that one of the most important applications of organomercury compounds in organic synthesis is related to their potential for generating radical intermediates. The most general method for carrying out this generation, which can **also** be performed thermally,

photochemically, or by other methods, is the so-called "mercury method" by means of sodium borohydride. Primary, secondary, and tertiary unfunctionalized or α -, β -, and γ -functionalized radicals bearing an oxygenated or nitrogenated functional group have been obtained by this methodology. The further reaction of the generated radicals with different unsaturated systems through an intra- or intermolecular reaction constitutes an efficient procedure of obtaining regiospecific but not stereospecific carbon-carbon bonds and represents an adequate way to construct organic molecules.

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