The chemistry of **diazonium and diazo groups** Part I

Edited by **SAUL PATAI The Hebrew University, Jerusalem**

1978 JOHN **WILEY** & **SONS** *An Interscience e Publication* **CHICHESTER -- NEW YORK - BRISBANE -- TORONTO** Copyright @ **1978** by John Wiley *R:* Sons Ltd Reprinted February 1979.

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Library **of** Congress Catalog Card No. **75-691 3**

ISBN **0 471 99492 8 (Pt. 1) ISBN 0 471 99493 6 (Pt. 2) ISBN 0 471 99415 4** (Set)

Printed in Great Britain by Page Bros (Norwich) **Ltd,** Mile Cross Lane, Norwich.

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Foreword

The present volume, 'The Chemistry of Diazonium and Diazo Groups' is, on the whole, organized and presented according to the gencral lines described in the 'Preface to the Series', printcd on the following pages.

Some difficulty arose in the prcsentation owing to the fact that while the two groups treated, i.e. the diazo group and the diazonium group, are closely related and even occur in equilibrium with each other, thcir chemical behaviour and chnracteristics differ from each other considerably. Moreover, thc material which had to be covered proved to be much more extcnsivc than originally surmised. For thesc reasons, some of the subjects had to be divided into two or more chapters; for instance, the synthetic applications of diazonium and diazo groups arc treated in two separate chapters and even so each of these turned out to be very large. Similarly, the syntheses of the different title compounds are discussed in three separate chapters.

The plan of the present volume also included a chapter on 'Biological and Pharmaceutical Effects' which, howcver, failed to materialize. It is hoped that this will appear in one of the supplementary volumes to the series.

Jerusalem, February 1977 **SAUL PATA?**

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The Chemistry of Functional Groups Preface to the series

The series 'The Chemistry of Functional Groups' is planned to cover in each volume all aspects of thc chemistry of one of the important functional groups in organic chemistry. The emphasis is laid on the functional group treated and on the effects which it exerts on the chemical and physical properties, primarily in the immediate vicinity of the group in question, and secondarily on the behaviour of the whole molecule. For instance, the volume The Chemistry of the Ether Linkage deals with reactions in which the $C-O-C$ group is involved, as well as with the effects of the $C-O-C$ group on the reactions of alkyl or aryl groups connected to the ether oxygen. **It** is the purpose of the voluine to give a complete coverage of all properties and reactions of ethcrs in as far as these depend on the presence of the ether group but the primary subject matter is not the whole molecule, but the $C-O-C$ functional group.

A further restriction in the treatment of the various functional groups in these volumes is that material included in easily and generally available secondary or tertiary sources, such as Chemical Reviews, Quarterly Reviews, Organic Reactions, various 'Advances' and 'Progress' series as well as textbooks (i.e. in books which are usually found in the chemical libraries of universitics and research institutes) should not. **as a** rule, be repeated in detail, unless it is necessary for the balanced treatment cf the subject. Thercfore each of the authors is asked *tiot* to give an encyclopaedic coverage of his subject, but to concentrate on the most important recent developments and mainly on material that has not been adequately covered by reviews or other secondary sources by the time of writing of the chapter, and to address hiniself to a reader who is assumed to be at a fairly advanced post-graduate level.

With these restrictions, it is realized that no plan can be devised for a volume that would give **a** *cotiipletc* coverage of the subject mfith *ti0* overlap between chapters, while at the same time preserving the readability of the text. The Editor set himself the goal of attaining *reasonable* coverage with *moderate* overlap, with a minimum of cross-references between the chapters of each volume. In this manner, sufficient freedom is given to each author to produce readable quasi-monographic chapters.

The general plan of each volume includes the following main sections:

(a) An introductory chapter dealing with the general and theoretical aspects of the group.

(b) One or more chapters dealing with the formation of the functional group in question, either from groups present in the molecule, or by introducing the new group directly or indirectly.

(c) Chapters describing the characterization and characteristics of the functional groups, i.e. a chapter dealing with qualitativc and quantitative methods of determination including chemical and physical methods, ultraviolet, infrared, nuclear

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magnetic resonance and mass spectra: a chapter dealing with activating and directive effects exerted by the group and/or a chapter on the basicity, acidity or complex-forming ability of the group (if applicable).

(d) Chapters on the reactions, transformations and rearrangements which the functional group can undergo, either alone or in conjunction with other reagents.

(e) Special topics which do nct fit any of the above sections, such as photochemistry, radiation chemistry, biochemical formations and reactions. Depending on the nature of each functional group treated, these special topics may include short monographs on related functional groups on which no separate volume is planned (e.g. a chapter on 'Thioketones' is included in the volume The Chemistry of *tlie* CGrbotiyl Group, and a chapter on 'Ketenes' is included in the volume The Chemistry of Alkenes). In other cases certain compounds, though containing only the functional group of the title, may have special features so as to be best treated in a separate chapter, as e.g. 'Polyethers' in The Chemistry of the Ether Linkage, or 'Tetraaminoethylenes' in The Chemistry of the Amino Group.

This plan entails that the breadth, depth and thought-provoking nature of each chapter will differ with the views and inclinations of the author and the presentation will necessarily be somewhat uneven. Moreover, a serious problem is caused by authors who deliver their manuscript late or not at all. In order to overcome this probleni at least to some extent, it was decided to publish certain volumes in several parts, without giving consideration to the originally planned logical order of the chapters. If after the appearance of the originally planned parts of a volume it is found that either owing to non-delivery of chapters, or to new developments in the subject, sufficient material has accumulated for publication of a supplementary volume, containing material on related functional groups, this will be done as soon **as** possible.

The overall plan of the volumes in the series 'The Chemistry of Functional Groups' includes the titles listed below:

The Chemistry of Alkenes (two *volumes*) The Chemistry of the Carbonyl Group (two volumes) The Chemistry of the Ether Linkage The Chemistry of the Amino Group The Chemistry of the Nitro and Nitroso Group (two parts) The Chemistry of Carboxylic Acids and Esters The Chemistry of the Carbon-Nitrogen Double Bond The Chemistry of the Cyano Group The Clieniistry *of* Amides The Chemistry of the Hydroxyl Group (two parts) *Tlie* Chemistry of the Azido Group The Chemistry of Acyl Halides The Chemistry of the Carbon-Halogen Bond (two parts) The Chemistry of Quinonoid Compounds (two parts) The Chemistry of the Thiol Group (two parts) The Chemistry of Amidines and Imidates The Chemistry of the Hydrazo, *Azo and Azoxy Groups* The Chemistry of Cyanates and their Thio Derivatives The Chemistry *of* Diozoriiicm *atid* Diazo Groups Supplement A: The Chemistry of Double-Bonded Functional Groups (two parts) Titles in press:

The qliernistry of the Carboti-Carbon Triple Bond *Supplement B: The Chemistry of Acid Derivatives*

Future volumes planned include:

The Chemistry of Cumulenes and Heterocumulenes The *Clreniisfry of Orgariomefallic Cornpoirtrds The Chemistry of Sulphur-containing Compounds Sirpplemerit C: The Chemistry of Triple-borided Ficirctiorial Grorips* **Supplement D: The Chemistry of Halides and Pseudo-halides** *Supplement E: The Chemistry of* $-NH_2$ *,* $-OH$ *, and* $-SH$ *Groups and their Derivatives*

Advice or criticisin regarding the plan and execution of this series will be welcomed by the Editor.

The publication of this series would never have started, let alone continued, without the support of many persons. First and foremost among these is Dr Arnold Weissberger, whose reassurance and trust encouraged me to tackle this task, and who continues to help and advise inc. The efficient and patient coopcration of several staff-members of thc Publisher also rcndered me invaluable aid (but unfortunately their code of ethics does not allow me to thank them by name). Many of my friends and colleagues in Israel and overseas helped mc in the solution of various major and minor matters, and my thanks are due to all of them, especially to Professor *Z.* Rappoport. Carrying out such a long-range project would bc quite impossible without the non-professional but none the less essential participation and partnership of my wife.

The Hebrew University Jerusalem, **ISRAEL.**

SAUL PATAI

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CHAPTER 1

6eneral and theoretical aspects of the diazsnium and diazo groups

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1. INTRODUCTION

In discussing the chemistry of molecules containing the NN bond and, in particular, in this first chapter, the general and theoretical aspects of such molecules, it seems reesonablc to establish the scenario by comparing and contrasting such groups with other groups of importance in organic chemistry.

One would be hard pressed to find more logical candidates for comparison with molecules containing the N_2 group than those of C_2 and CN. Indeed such a ubiquitous trio of organic chemistry involving two homonuclear and one heteronuclear bond, the latter formed from the two different clenients of the former, almost invite sociological comparison with parents and offspring. The C_2 , CN and $N₂$ groups contain 12, 13 and 14 electrons, respectively, and the hierarchy of bonding can be conceived as beginning with the so-called single bond, so that we have as the simplest example for each group:

The **first** line of Table **1** compares average bond lengths for the central bonds of each of these molecules, and, not surprisingly, in view of the extra proton in the nitrogen nucleus as compared to carbon, the bond lengths decrease as one passes from the $C-C$ to the $N-N$ bond.

TABLE 1. Average bond lengths **(A)** in molecules containing *CC,* CN and NN groups

Bond label	CC.	CN	NN
Single Double Triple	1.537 1.335 1.202 $1.312(C_2)$	1.472 1.32 1.157 1.172 (CN)	1.451 1.25 1.098(N ₂)

Removal of **a** hydrogen atom from each of the central atoms in the molecules above produces the trio of molecules with central double bonds,

Again the same trend in bond lengths (line 2, Table 1) is observed as in the case of the corresponding single-bonded molecules. However, in addition, in all cases shown, formation of the double bond results in a decrease in the bond length.

Continuation of the dehydrogenation process yields

H-C=C--H **H-C=N** N=N

with the same gradation in bond lengths being maintained from $C=C$ to $N=N$, as well as a further decrease in bond length in each case on formation of the triple bond. Extraction of thc remaining hydrogen atoms from acetylene and hydrogen cyanide produces an increase in bond length, signalling a discontinuity in electronic structures. However, for cach type of bond (single bonded, ctc.) it is clear from the table that the NN bond length is, in each case, the lowest of the three shown.

The foregoing comparison may be cxtendcd to encompass bond encrgics, and average values for these arc given in Table 2. Examination of **the** table shows that the two lowest bond energies listed are 37 and 61 kcal/mol, those for the $N-N$ and $N=N$ bonds respectively, while the highest value is 225 kcal/niol for the dissociation of the nitrogen molecule. In fact, with the exception of *CO* which **has** a dissociation energy of 256 kcal/mol, N_2 has the highest dissociation energy of any diatomic molecule.

	$C1$ and $C1$ bonds				
Bond label	CC.	CN.	NN		
Single Double	83 146	73 112	37 61		
Triple	200 144 (C_2)	213 188 (CN)	225(N ₂)		

TABLE 2. Average bond energies (kcal/mol) of CC, CN and NN bonds

Consistent with these experimental observations are the conclusions derivable from simple molecular orbital theory for the electron configurations of C_2 , CN and N₂, and their singly-ionized forms (Table 3). The difference between the number of bonding orbitals and the number of antibonding orbitals, sometimes called the bond order, can be seen to increase from C_2^+ to N_2 , thereby suggesting that the strength of the bond in N_2 should not be unexpected.

Number of electrons	Molecule	Lowest electron configuration	State	$P_{\rm b}$ $P_{\rm a}$	Differ- ence	D_n^0 (volts)
11	C_2^+	$KK(\sigma_{\mu}2S)^2(\sigma_{\mu}2S)^2(\pi_{\mu}2P)^3$		$2\frac{1}{2}$		
12	$C2$, $CN+$	$KK(\sigma_{2}2S)^{2}(\sigma_{u}2S)^{2}(\pi_{u}2P)^{4}$		3.		(3.6)
13	C_2^- , CN, N ₂ ⁺	$KK(\sigma_g 2S)^2(\sigma_u 2S)^2(\pi_u 2P)^4\sigma_g 2P$		$3\frac{1}{2}$ 1	24	6.34
14	CN^- , N_2	$KK(\sigma_{\alpha}2S)^{2}(\sigma_{\alpha}2S)^{2}(\pi_{\alpha}2P)^{3}$ - $(\sigma_{\nu}2P)^2$	$\begin{array}{l} \Sigma_{\kappa}^+ \\ \Sigma_{\kappa+}^+ \\ 1 \Sigma_{\kappa}^+ \end{array}$	$\overline{4}$	3	7.37
15	N.	$KK(\sigma_{g}2S)^{2}(\sigma_{u}2S)^{2}(\pi_{u}2P)^{4}$ - $(\sigma_{\alpha}2P)^2 \pi_{\alpha}2P$	2 π_{σ}		24	

TABLE 3. Electron configurations^a of C₂, CN and N₂ molecules and their singly-ionized forms

 $P_{\rm b}$ = Number of bonding electron pairs.

 $P_{\rm A}$ = Number of antibonding electron pairs.

 D_0^0 = Dissociation energy.

^a*G.* Herzberg, *Spectra of Diafotnic Molecules,* 2nd ed., D. Van Nostrand *Co.* Inc. Princeton, **N.J.** (1950).

Energy and geometry parameters relating to the ionization of the diatomic species **C2,** N, and CN are given in Table **4.** Again, although limited data are availablc, a consistent pattern is followed, N₂ is observed to have a high ionization energy and would be predicted to have a relatively high electron affinity.

	Ionization	Electron		Bond lengths (λ)		
Molecule	energy (kcal/mol)	affinity (kcal/mol)	XY^-	ХY	XY^+	
$\frac{\text{C}_2}{\text{CN}}$		71.5		1.24		
	348.9	$88-1$	1.07	1.18	$1 - 17$	
N,	359.3			$1 - 10$	$1 - 12$	

TABLE 4. Energy and geometry parameters^a for ionized forms of C_2 , N_2 and CN

^aK. Watanabe, T. Nakayama and **J.** Mottl, *J. Quanr. Specr. and Energy Tramfer, 2,* **369 (1962).**

The theoretical aspects, and related physicochemical properties of molecules containing the $N=N$ bond will be developed, in the following sections, by considering individual molecules and their substituted derivatives, each representative of a particular type of azo or diazo compound. The simplest relatives of N_2 will be considered first, CNN being taken as **a** hypothetical precursor of diazomethane, and $N_{2}H^{+}$ and $N_{2}H$ being considered as prototypes of the diimide bond. The discussion on CNN is followed by sections on diazomethane itself and the two of its structural isomeric forms, nitrilimine and diazirine, which contain the $N=N$ bond. Relevant information on substituted diazirines is also included.

Section I11 discusses the results of theoretical studies of diimide and its monoand di-substituted forms as well as being concerned with related experimental work on these molecules. Although not strictly within the bounds of the title of this volume, diimides have been included here both for completeness and purposes of comparison in discussions of the NN bond.

In all cases, where available, spectral data and the results of theoretical calculations are given, together with some thermochemical and kinetic data, particularly where the latter may assist in understanding the structure and properties of the molecule. In general each section is developed historically, in order to permit the appropriate assessment of the data to be made by the reader wishing to use them, and also to indicate to the researcher topics which require further study.

II. DIAZO COMPOUNDS

A. The **Prototype, CNN**

Molecules containing two nitrogen atoms joined together and with one of these nitrogen atoms attached to a single carbon atom are referred to as diazoalkanes (R_2CNN) . A number of extensive reviews of this subject^{1, 2} have been published in the previous decade, in addition to several since 1970, some as recent as 1972³⁻⁶. There has been much interest in diazoalkanes for many years, particularly because of their usefulness as intermediates, and more recently as **a** consequence of work on the reactivity and structure of carbenes (R_2C) , which are intermediates in the photolysis and thermolysis of diazoalkanes.

The parent species of the diazoalkanes can be viewed, at least from the theoretical point of view, as diazomethylene, CNN. Robinson and McCarty, Jr⁷, following the photolysis of diazomethane in a krypton matrix, observed a featurc in the spectra at 4240 **A** and tentatively suggested that this might be assigned to CNN or to HCNN. At about the same time, Goldfarb and Pimentel⁸ observed features at 4182 and 3968 **A,** following the photolysis of diazome!hane isolated in a nitrogen matrix,

1. General and theoretical aspects of the diazonium and diazo groups 5

which they too considered must be duc to a reactive species, presumably CNN, since these fcatures vanished when the sample was warmcd. In I966 thc rcsults of the first complete spectral study on CNN, prepared by the photolysis of matrix-isolated cyanogen azide with 2100-2800 Å radiation, were reported by Milligan and Jacox⁹. They first suggested the following mechanism for the production of diazomethylene from cyanogcn azidc :

$$
N_{3}CN + h\nu (\lambda < 2800 \text{ Å}) \longrightarrow N_{3}^{*} + \cdot CN \tag{1}
$$

$$
N_3^* + C N \longrightarrow N_3 NC \tag{2}
$$

$$
N_{3}NC \longrightarrow 2 N_{2}+C
$$
\n
$$
N_{3}NC \longrightarrow N_{2}+CNN
$$
\n(3a)
\n(3b)
\n(3b)

$$
N_3NC \longrightarrow N_2+CNN \tag{3b}
$$

However, observations on the systems containing ^{15}N led them to modify their conclusions to suggest that the principal mode of forniation of the CNN is by the attack of a carbon atom on a nitrogen moleculc.

Diazomethylene was deduced to possess a triplet electronic ground state with a linear structure. The carbon-nitrogen bond was found to have approximately triple-bond character and the nitrogen-nitrogen bond approximntcly double-bond character, with the length of the former being estimated as 1.15 A, and **tlint** of the latter as **1.25 A.**

More recently, CNN was prepared by trapping carbon atoms in a pure N₂ matrix¹⁰ and medium-intensity bands of ¹²CN₂ (in a N₂ matrix) were observed at 2858, **1253** and 394 cm-', which shifted *to* 2836, 1232 and **392** C11i-l when I3C **was** vaporized. Stretching force constants of 19.5 and 14.7 mdyn/ \AA were calculated for the CN and NN bonds, respectively.

What appears to be the only theoretical work done on CNN was reported by Thomson¹¹ in 1973. Using the linear molecules program $ALCHEMY¹²$ he performed restricted Hartree-Fock calculations for the CNN configuration, ${}^3\Sigma^-$, $(1\sigma)^2(2\sigma)^2$ - $(3\sigma)^2(4\sigma)^2(5\sigma)^2(6\sigma)^2(1\pi)^1(7\sigma)^2(2\pi)^2$. Calculated energies, geometries and atomic populations are reported in Table *5.*

				Total atomic populations		
Basis set	Energy	R(NN)	R(CN)			
Double zeta Double zeta + polarization	-146.59094 -146.68096	2.565 2.530	2.220 2.190	5.72 6:12	7.43 6.92	6.85 696

TABLE *5.* Calculated energies' and structure of CNNl'

⁴ Energies are in hartrees and bond lengths in bohrs: 1 hartree = 627.7 **kcal/mol; 1 bohr =** 0.529 **Å.**

In addition **a** dissociation energy (to atoms) of 0.198 hartrec and a dipolc momcnt of **1.022 D** were calculated for CNN, with the convcntion that thc right-hand nitrogen atom is positive.

The calculated bond lengths are in reasonable agreement with the estimates of Milligan and Jacox9. The atomic populations arc seen to bc particularly scnsitivc to the size and nature of the basis set. Thomson also calculated the dissociation energies for NCN, NCC and CNC to **be** 0.212, 0.270 and 0.273 hartrecs, respectively. **As** expected the calculations suggest that CNN is less stable than any of thcse lattcr three species.

B. Diazornethane (H,CNN)

1. Introduction

Diazomcthane is the simplest diazoalkane and as such has served as the prototype for many studies, both experimental and theoretical. From a classical point of view it may be considered as a resonance hybrid of **a** variety of planar forms:

$$
\bar{C}H_2 - \stackrel{+}{N} \equiv N \tag{a}
$$

$$
\bar{C}H_2 - N = N^+ \tag{b}
$$

$$
\tilde{C}H_2-N=N
$$
 (c)

$$
CH_2 = \stackrel{+}{N} = N^-
$$
 (d)

Hence it is not unreasonable to expect diazomethane to be capable of assuming the properties of a carbene source, or an acid or **a** base, or an electrophile or a nucleophile, or a 1,3-dipole⁵.

For many years there was considerable controversy concerning the structure of diazomethane until electron diffraction¹³ and microwave spectroscopic experiments¹⁴ showed the arrangement of carbon and nitrogen atoms to be linear and the overall structure to be planar. In 1960 the chemical evidence for the linear structure of diazomethane was provided by Paulsen¹⁵, and Schmitz and Ohme¹⁶ in the form of independent syntheses of the first diazirines,

which **are** isoelectronic with their corresponding diazoalkanes.

2. Theoretical studies

The first calculations¹⁷ of the electronic structure employed the simple Hückel molecular orbital method to calculate a resonance energy of 0.6β and, at least within the assumptions of the method, values of the bond lengths and dipole moment in reasonable agreement with experiment. Hoffmann¹⁸ some years later applied his extended Hückel technique to calculations on the excited states of diazomethane. Yoshida and Kobazashi have calculated the electronic spectra of diazomethane by the ASMO-LCAO-SCF-CI¹⁹ and INDO-SCF-MO-CI²⁰ methods. Brintzinger²¹ has suggested a qualitative description of the chemical bonds in diazomethane. What appears to be the first ab initio study of diazomethane was that performed by André and coworkers²². Apparently, however, an error in geometry unfortunately invalidates the results. Bastide and Henri-Rousseau²³ have done CNDO calculations to examine the transition between the linear and a bent form of diazomethane, the CNN angle in the latter was taken as 75°. A rotational barrier of 22 kcal was calculated and an interpretation²⁴ of orientation effects in $1,3$ -dipolar addition reactions was proposed. Hart²⁵ suggested that seven isomers of diazomethane are, in principle, possible, and carried out *ob irtirio* calculations using the **LW** [533; 31 Gaussian lobe basis set, on five of these structural isomers, diazomethane itself being included. Caballol and coworkers²⁶ employed the INDO technique to obtain wavefunctions for the ${}^{1}A_{1}$, ${}^{1}A''$ and ${}^{3}A''$ states of diazomethane. The ground-state

geometry was taken from Herzberg²⁷ and the excited state geometries were obtained from the ground-state geometry by bending the CNN angle in the molecular and bisector planes. The planar structures were found to be more stable and the CNN angle was found to be **158"** and **152"** for the first excited singlet and triplet state, respectively. Two minima of depth -14.9 and -13.9 kcal/mol were found in the electrostatic potentials for ground-state diazomethane, the former at the carbon end and the latter at the nitrogen end of the molecule. These authors argue that this agrees with the experimental results which suggest **a** carbon attacks. In the case of the first excited **'A"** singlet state, protonation is predicted to occur within the molecular plane with a single minimum (-23.6 kcal/mol) appearing in the neighbourhood of the terminal nitrogen. Electrostatic potential results for diazomethane in the triplet state are quite similar to those for the singlet state, the well depth being - **24-4** kcal/mol in the triplet case.

Leroy and Sana²⁸ have employed the Gaussian -70 programme of Hehre and coworkers²⁹ and a variety of basis sets to calculate the electronic structure and some thermodynamic properties of this compound. With the contracted Gaussian basis of Clementi, a total electronic energy of - **147.18569** hartree, and bond energies of 104 and 181 kcal/mol for the CN and NN bonds, respectively, were obtained. In these calculations the geometry of diazomethane determined experimentally by **Cox** and coworkers³⁰ was employed. With a 4-31G basis set, Leroy and Sana²⁸ calculated the dipole moment to be 1.52 D to be compared with an experimental value of $\mathbf{i} \cdot 40$ D³⁰, and a total electronic energy of -147.60552 hartree.

The energy of formation at $0K$ was obtained by the use of the equation

$$
\Delta E_{\rm f} (0~{\rm K}) = E_0^0 = \Delta E_{\rm f}^{\rm th} + \frac{1}{2} \sum_i h \nu_i
$$

where ΔE_f^{th} , the theoretical energy of formation was taken to be that calculated with the $4-31G$ basis set, and the v_i are the frequencies of the nine normal modes for which the values from Moore and Pimentel 31 were employed. The energy of formation at *25 "C*

$$
\Delta E_{\rm f} (298.16 \text{ K}) = E^0 = E_0^0 + (E^0 - E_0^0)
$$

was then obtained by using the previously calculated value for E_0° , together with the value of $E^0 - E_0^0$ found by Moore and Pimentel³¹. The heat of atomization (ΔH_a) at **298.16** K **was** calculated from the experimental values of the heats of atomization of the elements 32 .

In addition Leroy and Sana²⁸ calculated values for the thermodynamic functions mentioned above by another method, namely that proposed by Hehre and coworkers³³ in which use is made of reactions and their calculated energies in which the molecule of interest, in this case diazomethane, is involved either as a reactant or as a product. The two such reactions selected by Leroy and Sana are

(a)
$$
2 CH_{2}N_{2} + NH_{3} = (CH_{3})_{2}NH + 2 N_{2}
$$

\n $\Delta E^{th} = -145,260$ kcal
\n(b) $CH_{2}N_{2} + NH_{3} = CH_{3}NH_{2} + N_{2}$
\n $\Delta E^{th} = -72,064$ kcal

So that correlation effects could be considered to cancel out, both the **C-H** and $N-H$ bonds had to be assumed equivalent, as well as the $C-N$ bonds in all molecules in which they were present. The energy of formation of diazomethane can then be obtained from

$$
\Delta E^{\text{th}} = \sum k_i N_i \Delta E_{\text{fi}}
$$

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or, more specifically for each of the reactions (a) and (b)

(a)
\n
$$
\Delta E^{\text{th}} = 2\Delta E_f^{\text{exp}}(N_2) + \Delta E_f^{\text{exp}}[(CH_3)_2NH] - \Delta E_f^{\text{exp}}[(CH_2N_2) - 2\Delta E_f^{\text{th}}(CH_2N_2)]
$$
\n(b)
\n
$$
\Delta E^{\text{th}} = \Delta E_f^{\text{exp}}(N_2) + \Delta E_f^{\text{exp}}[(CH_3)NH_2] - \Delta E_f^{\text{th}}(CH_2N_2)
$$

where ΔE_f^{exp} is an energy of formation referenced to 0 K and corrected for the zero point energy. Table **6** reports the values obtained by Leroy and Sana for the thermodynamic functions of diazomethane. **A** value of 78 kcal/niol was also found for the **CH** bond energy. Thc calculated value for the heat of formation of diazomethane at **298.1 6 K (63.5** kcal/niol) may be compared with that obtained experimentally $(51-60 \text{ kcal/mol})^{34}$ (see also Table 8).

TABLE 6. Calculated thermodynamic properties of diazomethane (kcal/mol)²⁸. Reproduced **by** permission of Springer-Verlag

	$\Delta E^{\rm th}$	ΔE_f (0 K)	ΔE_f (298.16 K)	$\Delta H_{\rm a}$ (298.16 K)
Method 1 Method 2	55.26	58.85	61.74	440.56
Equation (a)	57.45	$61-03$	63.91	438.39
Equation (b)	60.26	62.83	$65 - 72$	436.58
Average value	57.66	60.90	63.79	438.51

Leroy and Sana haw also performed geometry optimizations within the **STO-3G** basis. Thcir results togcther with those for the force constants are compared with the experimental values of Moore and Pimentel in Table 7.

Leroy and Sana, in discussing the significancc of their *ab inifio* calculations in respect of the structure and bonding of diazomethane, note that the $C-N$ and $N-N$ bonds in diazomethane have the characteristics expected of a double and a triple bond respectively, and hence the molecule behaves as one possessing six π electrons. Four of these six electrons form a π bond extending over three centres, while the remaining two arc essentially localized on the two nitrogen atoms. These authors concludc that the most important structural forms are those labelled (a) and (d) in Section II.B.1.

Walch and Goddard³⁵ have very recently reported on the results of *ab initio* generalized \fnlrnce bond **(GVB)** and configuration interaction **(CI)** calculations (using **n** double-zeta basis) on the ground and low-lying excited states of diazomethane. **All** calculations employed thc experimental geometry for the ground state

as found by Cox, Thomas and Sheridan³⁰: $R_{CH} = 1.077 \text{ Å}$, $R_{CM} = 1.300 \text{ Å}$, $R_{NN} = 1.139$ Å and $\angle HCH = 126.1^\circ$. In contrast to the conclusion of Leroy and Sana, Walch and Goddard conclude that the calculated wavefunction **is** basically that of a singlet biradical with strong bonding between the radical *x* orbitals on the *C* and terminal N resulting from the interaction with the π pair on the central N:

$$
\sum_{n=0}^{n} (-\ddot{y}) = \dot{y}
$$

Further, Walch and Goddard suggest that the GVB model of diazomethane would support a mechanism based on **a** biradical attack in the 1,3-addition of diazomethane to olefins, analogous to that in ozone rather than the usual mechanism dependent on the dipolar nature of diazomethane.

Results *of* theoretical studies, in which both diazomethane and diazirine are considered, are discussed in the section on diazirine.

3. Thermochemical properties

Foster and Beauchamp³⁶ have concluded that the proton affinity (PA) of diazomethane can be ordered with that of ammonia, azomethane and methylamine in the following way:

$$
\mathsf{PA(NH_3)}\!<\!\mathsf{PA(CH_2N_2)}\!<\!\mathsf{PA(CH_3N=NCH_3)}\!<\!\mathsf{PA(CH_3NH_2)}
$$

Consequently they place the heat of formation of diazomethane as greater than **64** kcal/mol but less than 73 kcal/mol.

Table **8** summarizes the various values suggested for the heat of formation of diazomethane.

Method	$\Delta H_{\rm f}$ (25 °C) (kcal/mol)	Reference	
Kinetics	$67+$	a	
Thermochemistry	$64 - 77$	h	
Group additivity	71	c	
Appearance potential	49	d	
Photodissociation	$51 +$	е	
Proton affinity	$64 - 73$		
Theoretical	64		

TABLE 8. Heat of formation of diazornethane

D. **W.** Setser and **B.** *S.* Rabinovitch, *Can. J. Chwt., 40,* **1425** (1962).

 \circ J. C. Hassler and D. W. Setser, *J. Amer. Chem. Soc., 87,* 3793 (1965).

S. W. Benson, F. R. Cruickshank, D. M. Golden, G. R. Haugen, H. **E.** O'Neal, **A. S.** Rodgers, R. Shaw and R. Walsh. *Cizern. Rev., 69.* **279** (1969).

^{*d*} G. S. Paulett and R. Ettinger, *J. Chem Phys.*, 39, 825, 3534 (1963); **41. 2557** (1964).

 \bullet A. H. Laufer and H. Okabe, *J. Amer. Chem. Soc.,* **93.** 4137 **(1971).**

1 Referencc **36.**

0 Reference **28.**

C. Isomers of Diazomethane

having six known or suspected structural isomers. As Hart²⁵ has pointed out, diazomethane is unique among small molecules in

Of these isomers, diazirine and cyanamide have been prepared, and are stable molecules at room temperature. Derivatives of isodiazomethane, carbodiimide and isocyananiide are known and there is some evidence that isocyanamide has been prepared37* **38.** Carbodiimide and isodiazirinc have apparently not yet been prepared. Isodiazomethane, although not stable at room temperature, has been prepared³⁹⁻¹¹. Only the diazo isomers, namely isodiazomethane, diazirine and isocyanamide, will be discussed here.

1. Nitrilimine

Investigations of the reaction of triphenylmethyl sodium on diazoniethane by Miiller and **coworkers** in **193430** led to the discovery of this isomer of diazomcthane. Anselme⁴² has summarized the properties of isodiazomethane as they were known in 1966. The preparation involves the acidification at -80 °C of an ether suspension of diazomethyl anion ($[CH-N=N]^m$ +), with an aqueous solution of potassium dihydrogen phosphate or ammonium chloride, as in

where M⁺ may be Li⁺, Na⁺ (C₆H_b)₃PCH₃. Isodiazomethane, a pale yellow liquid, is then obtained by evaporation of the solvent at -50 °C *in vacuo*. Decomposition

begins to occur at **15** "C and the compound explodes at **35-40** "C **'O. A** maximum occurs at **247** nm in the ultraviglet spectrum, compared to absorption at **412** rim and **321** nm with diazomethane and diazirine, respectively. No infrared spectra have yet been reported, but it appears from niolecular weight determinations in benzene that isodiazomethane is monomeric¹¹. The formation of isodiazomethane by acidic hydrolysis of $[CH-N=N]$ ⁺ may be explained through the assumption of a kinetically controlled reaction, by which protonation occurs rapidly at the more nucleophilic nitrogen (presumably the terminal nitrogen) to give the thermodynamically less stable isomer.

Apparently the only theoretical work which has been done on isodiazomethane is that by Hart²⁵ mentioned previously. Results of calculations on isocyanamide were also reported at that time. The total electronic energies were obtained as -147.2425 , also reported at that time. The total electronic energies were obtained as -147.2425 , -147.2868 and -147.3386 hartree for isodiazomethane, diazomethane and isocyanamide, respectively. This would imply that isocyanamide is more stable than isodiazomethane or diazomethane by approximately **63** and **34** kcal/mol, respectively. Indeed, Müller, Beutler and Zeeh⁴³ now suggest that the compound previously labelled isodiazomethane may in fact be isocyanamide. If this be so it is of interest to seek out any evidence for the existcnce of nitrilimine. Mills and Thompson", in a report in **1954** on the vibration-rotation bands of diazomethane, note that a peculiarity in the results for the band near 2100 cm^{-1} leads to the suggestion that this may be due to nitrilimine. On the other hand, Moore and Pimentel⁴⁵, 10 years later, find no evidence for any tautomeric form of diazomethane. More recently however, Ogilvie⁴⁶, again on the basis of an unexplained infrared band observed at **2169** cm-' in the photolysis of diazomethane (gas phase and an argon matrix), has postulated the existence of nitrilimine.

Hart²⁵ has calculated the total electronic energy of the anions $[HCNN]$ ⁻ and [CNNHI- **a:** - **146.6485** and - **146.6389** hartree, respectively, making the former [HCNN]⁻ more stable than [CNNH]⁻ by approximately 6 kcal/mol. Calculations, assuming the entropies of these anions are equal, yield a value of **2.3** x **10'** for the equilibrium constant *K* for the equilibration of the two anions. Hart has calculated electrostatic potential maps for the two anions. For [CNNH]⁻ there are minima of **-225.0** and - **187.0** kcal/mol in the region of the nitrogen and carbon lone pairs, respectively, while for [HCNN]⁻ the corresponding values are -191.0 and - **229.0** kcal/mol. These results suggest that protonation of [CNNH-1 is more likely to occur at the nitrogen atom to give isocyanamide as the product while protonation of [HCNN]⁻ should occur at the carbon atom to give diazomethane. These values indicate that the dilference in proton affinity for nitrogen over carbon in [CNNH]⁻ is exactly reversed in [HCNN]⁻. Protonation at the 'secondary' nucleophilic site in each anion generates a common intermediate, nitrilimine. Hence it is conceivable that a feasible tautomeric pathway could involve a quickly attained equilibrium (involving proton gain and loss) through nitrilimine as an intermediate: FILENTY THE COTESPONDING VARIATE RESERVING THE CONTROLLING THE CONTROLLING THE CONTROLLING THE CONTROLLINGT THAT THAT A THAND THE CONTROLLING THAND T

$$
\text{CN\~M}\xrightarrow{\text{H}\,^+} \text{HCNNH}\xrightarrow{\text{H}\,^+} \text{H\~CNN}
$$

A hypothetical reaction mechanism for the production of isocyanamide from diazomethane can then be constructed as shown in Figure **1.** The anion tautomerism could occur through nitrilimine or directly as a single step (broken line). Unfortunately there is very little experimental evidence to support any mechanism. Of course a variety of different possibilities for such a mechanism exist. The formation of a dianion (CNN2-) intermediate or the formation of a cyclic transition state involving **a** water molecule cannot be excluded. With the suggestion from Walch and Goddard³⁵ of a diradical structure for diazomethane, various mechanisms involving radicals become more reasonable. As Hart²⁵ points out, although his calculations suggest that isocyanamide is more stable than diazomethane, the experimental evidence suggests that diazomethane is thermodynamically the more stable. Since geometry optimization was performed with isocyanamide but not diazomethane, it is probable that the energy of the latter is not **at** the minimum value.

FIGURE **1. Total** electronic energy changes in a hypothetical reaction mechanism for **the** production *of* isocyanarnide from diazomethane. Reproduced by permission of the *Australian Journal of Chemistry.*

2. Diazirine and derivatives

a. *Spectra, structure and thermochemistry*. The cyclic isomers of diazoalkanes (diazirines) were first prepared in 1960 independently by Paulsen⁴⁷ and Schmitz and Ohme⁴⁸. Studzinskii and Korobitsyna¹⁹ have briefly considered the subject of diazirines as part of a review on aliphatic diazo compounds. Shortly thereafter the molecular structure, dipole moment and quadrupole coupling constants of diazirine were determined⁵⁰ from the rotational spectrum in the frequency region 7.8-41 GHz. Measurements on H_2 ¹³CN₂ and H_2C ¹⁴N¹⁵N permitted interatomic distances of 1.228, 1.482 and 1.09 Å to be calculated for NN, CN and CH, respectively. The HCH, NCN and NCH angles were given as **117",** 48'57' and **118'24'** and the dipole moment as 1.59 debye.

Schmitz and Ohme⁵¹ and Graham⁵² report the syntheses, by two different methods, of diazirinc itself. Diazirinc is a colourless **gas** at room tempcratiire and has **a** boiling point of $-14 \degree C$. Graham⁵² reports a number of sharp, regularly spaced peaks between 282 and **324** nm in the gas-phase ultraviolet spectrum and remarks on the striking similarity in shape of the spectrum with that of **2,3-diazabicyclo[2.2.1]-2** heptene⁵³. The proton n.m.r. spectrum in carbon tetrachloride solution at 0 °C is a singlet at + **241** Hz (relative to external benzene), a relatively high-field absorption for methylcne protons, which is consistent with a strained three-membercd ring structure. In the infrared spectrum of diazirine, Ettinger⁵⁴ found strong absorption in the 1600 cm^{-1} region, which he assigned to double-bond NN stretching. In comparison, Le Fevre and coworkers^{55, 56} concluded that the $-N=N-$ linkage, in a simple cnvironment found in a series of azo compounds, has a charactcristic frequency about 1580 cm^{-1} . Values of the thermodynamic functions for diazirine (ideal gas, **1** atni) as calculated by EttingerS4 are given in Table **9.** In a brief prcliminary note, Merritt⁵⁷ has reported on the ultraviolet spectrum of diazirine under

high dispersion, and finds the *0-0* band at **30,964 cm-I. A** number of frequencies in the ground and upper state are also found and tentatively identified.

The mass spectra and appearance potcntials of diazirinc and diazomethane have been determined⁵⁸ and the heat of formation of diazirine estimated to be 79.3 kcal/mol as compared with 49.3 kcal/mol for diazomethane. By the use of bond energies the strain energy associated with the diazirine ring structure was estimated to be about 22 kcal/mol. The heats of formation of the ions $CH_2N_2^+$ and CHN_2^+ were also estimated to be 314 and 355 kcal/mol, respectively.

T(K)	$-(G^0 - H_0^0)/T$	C_{p}^0	S^0	$(H^0 - H_0^0)/T$
$298 - 15$	48.44	10.19	56.87	S.43
400	51.01	12.39	60.17	9.16
600	55.05	15.94	65.91	10.87
800	58.39	18.35	70.85	12.45
1000	61.32	20 04	75.14	13.81

TABLE 9. Thermodynamic functions (cal mol⁻¹ deg⁻¹) for diazirine⁵¹ (ideal gas, **1 atm)**

That the cyclic isomer is more stable than the linear isomer is readily seen from the dilference in the estimated heats of formation, approxiniatcly 30 kcal/mol. The difference is also evident in tlie reaction chemistry, the diazoalkancs bcing some of the most reactive substances known in organic chemistry while diazirines arc much more stable. Graham⁵² notes that diazirine is decomposed relatively slowly by sulphuric acid with liberation of a molecule of nitrogen. It is stable to *t*-butoxide ion in t-butyl alcohol and, although storable in glass, is decomposed by ultraviolet radiation. Apparently it can be handlcd safcly at room tempcraturc as a liquid or a gas although, as a consequence of the lack of agrccnient on this point, it is advisable to take suitable precautions.

Bellss has suggested that **a** different set of assumptions concerning thc processes occurring in the mass spectrometer ion source will yield values of ΔH_f differing from those of Paulctt and Ettinger⁵⁸ but more in accord with the results of other experiments. The production of CH_2^+ from CH_2N_2 (either diazirine or diazonicthane) can bc expressed by

$$
A(CH_2^+) = \Delta H_f(CH_2^+) - \Delta H_f(CH_2N_2)
$$

where $A(CH₂⁺)$ is the appearance potential of $CH₂⁺$ from the particular isomer under consideration. Hence ΔH_f (CH₂N₂) can be obtained from the experimental appearance potential if a value for $\Delta H_i(\text{CH}_2^+)$ is available. Bell argues that the value of 333 kcal/mol⁶⁰ used by Paulett and Ettinger for $\Delta H_1(\text{CH}_2^+)$ is an older one, and in a previous application⁶¹ has led to tile conclusion that the major product in an ionmolecule reaction is formed via an endothermic process, whereas if the value suggested by Bell is uscd the process bcconics exothermic. Thc ionization potcntial of CH₂(${}^{3}\Sigma_{\rm g}$) to give CH₂⁺(${}^{2}\pi$), probably the ion ground state, is 240 kcal/mol⁶². Bell suggests that a very probable value for ΔH_f of CH₃(¹A₁) is 85 kcal/mol, on the basis of kinetic and thermochemical experiments⁶³⁻⁶⁵. Jordan and Longuet-Higgins⁶⁶ have calculated a value of 10 kcal/mol for the separation of the $CH_2(^1A_1)$ state from the ground state CH₂(${}^{3}\Sigma_{g}^{-}$). Hence $\Delta H_{1}[\text{CH}_{2}({}^{3}\Sigma_{g}^{-})] = 75$ kcal/mol and $\Delta H_{1}[\text{CH}_{2}^{+}({}^{2}\pi)] = 75$ 315 kcal/mol. **If** a correction of 40 kcal/inol is applied to allow for thc possibility that CH⁺ will not be formed in the 2π state in electron-impact experiments⁶⁷ but rather in the higher energy ${}^{4}\Sigma_{a}^{-}$ state⁶², then $\Delta H_{1}[CH_{2}^{+}({}^{4}\Sigma_{a}^{-})]$ is 355 kcal/mol. Table 10

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summarizes Bell's values, with two heats of formation for each of CH_2N_2 , $CH_2N_2^+$ and CHN_2^+ from each isomer, calculated on the assumption (i) that $A(CH_2^+)$ corresponds to a process yielding $CH_2^+(P_{\alpha})$, or (ii) that $A(CH_2^+)$ corresponds to a process giving $CH₃⁺(² π).$

		Diazomethane	Diazirine
$A(CH_2^+)$ ⁵⁸		284	254
$A(CH_2N_2^+)$ ⁵⁸		208	235
$A(CHN2+)$ ⁵⁸		341	327
ΔH_I (CH _a N _a)	(i)	71	101
	(ii)	31	61
ΔH_f (CH ₂ N ₂)	(i)	279	336
	(ii)	239	296
ΔH_f (CHN ⁺)	(i)	360	276
	(i)	320	336

TABLE 10. Appearance potcntinls and calculated thcrrnochemical values⁵⁰ (kcal/mol)

Differentiation between values labelled (i) and (ii) **is** riven in the text.

Bell favours the value of 71 kcal/mol for ΔH_f (diazomethane), since kinetic studies⁶³ suggest that the value should be greater than 67 kcal/mol, and the value of 61 kcal/mol for ΔH_f (diazirine) on the basis that the value of 101 kcal/mol would cause the reaction to yield $CH₂$ and $N₂$ to be 16 kcal/mol exothermic, and this would seem to conflict with the experimentally known relative thermal stability of diazirine compared to diazornethane.

Paulett and Ettinger⁶⁸, in their reply to Bell, argue that his estimate of 75 kcal/mol for the hcat of formation of the ground state of methylene is too low and present evidence to support a value of 95 ± 5 kcal/mol. They present values of 104 and 134 kcal/mol for the heats of formation of diazomethanc and diazirine. respectively, in the case where the $A(CH_2^+)$ corresponds to processes in which CH_2^+ is generated in an excited state.

Lau⁶⁹ has reported an absorption maximum of $30,989$ cm⁻¹ for diazirine, and notcs a red shift whcn the spectra of derivatives of diazirine were obtained in solution, which leads him to suggest that the electronic transition is a π^* - π transition.

Following up on his earlier note, Merritt⁷⁰ provides further details of his work on the electronic absorption spectra of *n*-diazirine, N_1^{15} -diazirine, d_1 -diazirine and d_2 -diazirine. Merritt notes that, as in diimide, the diazirine molecule contains two adjacent nitrogen atoms, each with a lone pair of electrons. Hence a splitting of the degeneracy in the electronic spectrum resulting from the perturbation of the lonepair electron orbitais wouia be expected. Mcrritt reports two systems. separated by approximately 220 cm⁻¹, at 30,970 cm⁻¹ and 31,187 cm⁻¹, both of which are present at **14K.**

b. *Thorctictll studies.* What appears to be the first quantum chcmical study of diazirine was done by Hoffman⁷¹ using his extended Hückel calculations (EXH). The experimentally determined geometry of Pierce and Dobyns⁵⁰ was employed to calculate that diazomethane is 70 kcal/mol more stable than diazirine and that the ionization potentials of diazomethane and diazirine arc 11.96 and 12.78 eV, respectively, compared with the Paulett and Ettinger⁵⁸ experimental values of 9.03 and **10.18** eV, respectively. The iciativc simplicity of **the EXH** nicthod must, of

course, be kept in mind. Frey and Stevens⁷² and Amrich and Bell⁷³ have shown that the major primary photochemical reaction in the **3200** *A* photolysis **of** diazirines is probably direct elimination of nitrogen to form **a** carbene, while a minor pathway is rearrangement to form the diazo compound.

The b_2 orbital is $C-N_1$ and $C-N_2$ bonding but N_1-N_2 antibonding. The antibonding a_2 orbital is only $N_1 - N_2$ antibonding. Hence the $a_2 \leftarrow b_2$ excitation (to which the 3200 Å transition is assigned) weakens the $C-N_1$ and $C-N_2$ bonding more than that of $N-N$, and the breaking of one or both of these bonds is consistent with the known photochemistry of the molecule.

In a study of the photolysis of diazirine in a nitrogen matrix, Moore and Pimentel⁷⁴ showed that the product diazomethane was formed from the reaction of methylene (from diazirine) and matrix N_2 . No diazomethane was found in the gas phase photolysis of diazirine. Hoffmann employed calculations on two simple models to obtain information on the path of the isomerization. Starting with diazirine geometry, a simple C-N₂ bond lengthening, keeping C-N₁ and N₁-N₂ constant was studied. In addition, using the diazomethane structure, and holding all bond lengths fixed, combinations of changes in CN_1N_2 and HCN_1 bond angles were considered, but with N_1 and N_2 constrained to remain in the original plane of the ring. A number of features were observed from the results of the calculations. The A_2 excited state of diazomethane prefers to be planar, and correlates with the B_1 excited state of diazirine. The A₁ excited state of diazomethane has a lower relative energy with a bent CNN skeleton. The results of photochemical experiments⁷³ appear to require the involvement of more than one excited state of diazomethane in the rearrangement. Hoffmann suggests that one possible pathway then involves the conversion of some of the excited B_1 diazirine into the second excited state, A_1 , of diazomethane, and return from that excited state to the ground state without internal conversion to the A_2 excited state. The application of Woodward–Hoffmann rules to the concerted decomposition of diazomethane and diazirine to methylene and N_2 leads to the conclusion that the first excited state of diazirine (B_1) and that of diazomethane (A_2) may decompose only to an A_1 or B_1 methylene and an electronically excited nitrogen molecule.

Snyder and Basch²⁵, as part of a paper in which heats of reaction are calculated from self-consistent field energies of closed-shell molecules, have included diazomethane and diazirine among the many molecules considered. Table 11 summarizes

	Diazirine		Diazomethane		
E_{DZ} (hartree) $-V/2T$ (hartree) $E_{\rm vib}$ (hartree) $E_{\rm T}$ (hartree) $E_{\rm corr-intra}$ (hartree) ΔH_f^0 (298 K) (kcal/mol)	-147.7287 0.99960 0.0321^{54} -147.6938 -0.7014 $[+101]$ ⁵⁹	-147.7702 1.00006 0.0306^{27} -147.7368 -0.7071 $[+71]^{59}$			
Reactions	Theory	Experi- mental	Theory	Experi- mental	
$\Delta H_{\text{208}}^0 \text{ (H}_2 + \text{CH}_2\text{N}_2 = \text{CH}_4 + \text{N}_2 \text{) (kcal)}$ ΔH_{298}^0 (4 H ₂ + CH ₂ N ₂ = CH ₄ + 2 NH ₃) (kcal) ΔH_{298}^0 (2 CH ₄ + CH ₂ N ₂ = C ₂ H ₆ + CH ₄ + N ₂)	-119.6 -162.5 -142.9 -97.2 (kcal)	$[-118.8]$ $[-141.0]$ $[-103.3]$	-92.2 -135.11 -102.3 -69.8	$[-88.8]$ $[-110.0]$ $[-73.3]$	

TABLE 11. Calculated and experimental data for diazirine and diazomethane⁷⁵

their various results for these two molecules, with experimental geometries employed throughout. To facilitate the calculations it was assumed that the sum of the electronic correlation energy and the difference of the electronic energy in the Hartree-Fock and the double-zcta basis did not change in a reaction having closedshell reactants and products. The second set of values given for the complete hydrogenation reaction in Table 11 results from the inclusion of estimates for interatomic and intraatomic correlation energy.

Kochanski and Lehn²⁶ employ the IB MOL programme for *ab initio* SCF-LCAO-**MO** calculations using a basis set of gaussian functions **on** diazirinc in its experimental configuration⁵⁰. Table 12, column 1, contains some of their calculated

TABLE 12. Calculated and experimental data^a for diazirine

^a All energies in hartrees.
^b Estimated from heat of formation.⁵⁸

results. The MO's $2b_2$, 6a₁ and $3b_2$ which are strongly localized on the nitrogen atoms contribute at the same time weakly to the $C-N$ bonds and strongly to the nitrogen lone pairs. Both $2b_2$ and $6a_1$ contribute strongly to the description of the lone pairs.

The C and N atoms have the following charge distributions:

 $1s^{1.98}$ $2s^{1.22}$ $2p\sigma^{1.73}$ $2p\pi^{1.25}$ N: $1s^{1.98}$ $2s^{1.89}$ $2p\sigma^{2.32}$ $2p\pi^{0.97}$

From this it may be concluded that the N atoms are σ acceptors and very weak π donors, while C is a π acceptor. Overlap populations show that the C-N bonds in diazirine are much weaker than the C-C single bonds in cyclopropene. The N=N bond is weakly σ antibonding and π bonding (MO's 1b₁ and 2b₁) so that the N=N bond is weakly σ antibonding and π bonding (MO's 1 double bond is weak. The low-lying $2b₂$ MO results from the antisymmetric mixing of two nitrogen lone pairs mostly N(2s) in character. $6a_1$ is weakly N-N bonding which contradicts the result from overlap population which implied that $6a_1$ was of weakly N-N antibonding character. $3b_2$ contributes to the nitrogen lone pairs as well as to the $C-N$ bonds. There is no pure lone pair MO. All MO's contributing to the lone pairs also contribute to some extent to the ring bonds.

An experimental study of the lower $n \rightarrow \pi^*$ excitation in difluorodiazirine, F_2CN_2 , has been performed by Lombardi and coworkers⁷⁷ and a supporting calculation on both diazirine and F_2CN_2 has been done⁷⁸. Calculations were performed using a gaussian-type orbital (GTO) basis set, using a total of 125 primitive GTO's which were contracted to 50 gaussian-type functions. This basis is equivalent to **a** socalled Double Zeta basis, in which each **A0** is represented by two Slater-type orbitals. Hencher and Bauer's structural data⁸⁷ were employed.

The calculated results⁷⁸ for F_2CN_2 are given in Table 13. The two MO's 9a₁ and 4b, represent the two 'non-bonding' orbitals conventionally assigned to the azo group, but mutual interaction separates them by 3.8 eV . The $n \rightarrow \pi^*$ transition originates at $4b_1$, and terminates at the π^* MO, 2a₂. The computed singlet-singlet and singlet-triplet excitations are **3.99** and 3.07 **eV,** rcspectively, compared to the observation of the excited singlet at **3.7** eV. Analysis of the MO's involved in the transition shows that *n* is nearly equally distributed among the $C-N-N$ atoms of the ring, but that π^* is completely localized on the N atoms. Consequently the $n-\pi^*$ excitation involves the transfer of about $\frac{1}{2}$ electron from the CF₂ group to the N=N group of difluorodiazirine.

The results of the *ab initio* calculation by Robin and coworkers⁷⁸ on diazirine are included in Table **12** for convenience in comparison with the earlier work of Kochanski and Lehn⁷⁶. In addition, Tables 12 and 13 also include some of the results obtained in comparison with photoelectron spectral data of both diazirine and F,CN,. The agreement between their observed and calculated (using 92% of the Koopmans' Theorem value) ionization potentials for both these molecules was found to be quite acceptable.

Bloor and Maksic⁸⁰ calculated diamagnetic susceptibilities and quadrupole moments of diazirine using the CNDO/ZD and *SCC* (self-consistent charge) semiempirical methods. The molecule was oriented as shown :

TABLE 13. Calculated and experimental data for difluorodiazirine⁷⁸

Total energy (a.u.) $-V/2T$		-345.3999 0.99961					
Orbital energies							
la _i 1b ₂ 2a, $1b_1$ $3a_1$ 4a, 2b ₂	-27.3726 -27.3726 -15.7494 -15.7483 -11.5465 -1.7715 -1.6915					Overlap and type ⁷⁹	
5a,	-1.5702						
2b ₁ 6a,	-0.9790 -0.9763		-0.581 $+0.158$	$N-N\sigma$ $N-N\sigma$			
3b ₂	-0.8569		$+0.100$	$C-F\sigma$			
$7a_1$ 3b ₁	-0.8447 -0.7741		$+0.195$ -0.207	$N-N\sigma$			
$1a_2$	-0.7306		-0.042	$N\!-\!N\sigma$ $F-F$			
$8a_1$	-0.7107		$+0.081$	$N - N\sigma$			
4b ₂	-0.7017		-0.145	$F-F$			
9a,	-0.6251						+0.039 N-N l.p. and -0.071 C- $F_{1,2}$ σ
$5b_2$	$-0.5800(\pi)$		$+0.354$	$N - N\pi$			
$4b_1$ 2a ₂ 56 $10a_1$ 6b,	-0.4851 $+0.0221 (\pi^*)$ $+0.2568$ $+0.2705$ $+0.2970$		-0.255		$N-N$ l.p.		
Dipole moment 0.082 D							
Quadrupole coupling constants ^a $x_{\alpha\alpha}$ (N) $x_{\beta\beta}$ (N) $x_{vv}(N)$ n	-0.614 -2.452 $+3.066$ 0.599						
Net charges							
С N F	$+0.544$ -0.018 -0.255						
Second moments of the electronic charge $\langle x^2 \rangle$ $\langle \nu^2 \rangle$ $\langle z^2 \rangle$	41.4323 ^b 97.2931 136.2211						

Values in Mc/sec computed in the principal axes system using a nuclear quadrupole moment of $Q(^{14}N) = 1.60 \times 10^{-26}$ cm².

 $\frac{b}{z}$ is the two-fold rotation axis, x is perpendicular to z in the CN₂ plane and y is perpendicular to the CN_2 plane.

Tables **14** and **15** list the electronic second moments and the molecular quadrupole moments calculated by the two methods and compare these with the *a6 initio* resultssl. Unfortunately there do not appear to be any experimental data **for** diazirine with which to compare the calculated values. However, with other molecules where such experimental data are available, agreement between such data and calculated values **is** very good **for** the second moments, but not as satisfactory with molecular

quadrupolc momcnts, the former agreement being ascribed by the authors more to the insensitivity of the quantities mentioned than to the high quality of the calculated wavefunctions.

_ . _ . _ . _ _ .					
CNDO/2D	SCCMO	ab initio ^{si}			
15.81	15.60	15.48			
9.27	9.47	9.62			
4.94	4.89	5.08			
30.02	29.96	30.22			

TABLE 14. Second moments of the electronic-charge distribution $(10^{-16}$ *e* cm²) of diazirine⁸⁰

TABLE 15. Molecular quadrupole moments $(10^{-26} \text{ Fr cm}^2)$ $(Fr = c.s.u.$ of charge) of diazirine⁸⁰

	CNDO/2D	SCCMO	ab initio ⁸¹
Q_{xx}	-0.81	0.54	2.01
$Q_{\nu\nu}$	-2.44	-4.02	-4.76
$\varrho_{\scriptscriptstyle{2z}}$	3.25	3.48	2.75

The results of MINDO/2 and EXH calculations^{82, 83} on diazirine and its dimethyl derivative are shown in Table 16. Haselbach and coworkers⁸² note that the orbital sequence for diazirine calculated by the MINDO/2 technique differs from that obtained by Robin and coworkers⁷⁸ in the order of the orbitals $a_1(n_+)$ and $a_1(c)$, and point out that it is known that for unsaturated systems MINDO/2 yields *o* orbitals with somewhat high energies. Hence the positioning of $a_1(\sigma)$ between $b_1(\pi)$ and $a_1(n_+)$ in Table 16 should be considered a consequence of the use of the MINDO/2 method.

TABLE 16. MINDO/2 calculated orbital energies (hartrees) for diazirine and 3,3-dimethyldiazirine

Diazirine ⁸²	3.3-Dimethyldiazirine ⁸³
$0.5751 b_1(\pi)$	$0.3572 b_2(n_$
0.4943 $a_1(n_+)$	$0.3862 b_1(\pi)$
$0.4322 a_1(\sigma)$	$0.4212 a_1(\sigma)$
$0.4105 b_1(\pi)$	0.4392 $a_1(n_+)$
$0.3848 b_2(n_+)$	$0.4502 b_1(\pi)$

The vertical ionization potentials $I_Y(J)$ of dimethyldiazirine as determined by photoelectron spectroscopy⁸² are given as 9.76 , 12.11 and 13.31 eV, or 0.3587 , 0.4450 and 0.4891 hartrees.

3. Substituted diazirines

a. *Spectra mid sfriictiive.* Although earlier **workersa7- 518 bs** had proposed methods for the preparation of diazirine and substituted diazirines, it appears that Mitsch⁸⁴ was the first to prepare perfluorodiazirine (F_2CN_2), a precursor of CF_2 . He

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and his coworkers subsequently measured⁸⁵ the infrared spectrum of F_2CN_2 from 4000 to 250cni-l **and** the Ranian spectrum from 2700 to 100cm-I in order to cstablish its structure. The frequencies of the infrared and Ranian bands for the fundamental modes expected for the diazirine and diazomethanc structures werc calculated and these were compared with the measured fundamental vibration frequencies of F_2CN_2 , shown in Table 17. The match between those values predicted for the diazirine structure and those observed is quite convincing.

TABLE 17. Summary of fundamental vibration frequencies of $F₂CN₂⁸⁵$. Reprinted, with permission, from Hencher and Bauer, *J. Afncr. Chcni. SOC.,~~,* **1186(1965).** Copyright by the American Chemical Society

Symmetry	Frequency (cm^{-1})	
Α,	1563 ν_{1} 1282 ν_2 805 v_3 502 $\nu_{\mathbf{A}}$	
Α,	451 ν.	
B_{1}	1248 ν_{κ} 481 ν,	
В.,	1091 $v_{\rm a}$ 544 Vq	

Simmons and coworkers⁸⁶ studied the ultraviolet absorption spectrum of CF_2N_2 in the 3750-3000 *8,* region under high resolution and confirmed the conclusions from the infrared and Raman work⁸⁵, that the molecule did indeed possess a cyclic C_{2V} structure. The results of the two studies agree quite \veil, as can be seen from Table **18,** except for v_3 , v_1 and v_4 may be described as the symmetric NN stretch and symmetric **CF,** defamztion, respectively. Hence it appears that the most significant change in geometry between the two electronic states is a stretching of the nitrogen-nitrogen bond which expands the ring and alters the FCF angle.

Vibration	Infrared (vapour) (cm^{-1})	Raman (liquid) (m^{-1})	Ultraviolet (vapour)	
				ν'' (cm ⁻¹) ν' (cm ⁻¹)
ν_{1}	1563	1560	1564.0	1436.9
ν_{2}	1282	1280	1282.9	
v_{2}	805	804	775.0	643.3
ν_a	502	500	$500-2$	516.6

TABLE 18. Comparison of the vibrational assignment for the A_1 species in CF_2N_2 as found from the infrared⁸⁵, Raman⁸⁵ and ultraviolet⁸⁶ spectra

Hencher and Bauer⁸⁷ completed the puzzle by providing interatomic distances and bond angles from electron diffraction experiments on the vapour. They confirmed the C_{2V} symmetry in which the planes of CF_2 and CN_2 are mutually perpendicular with $C-F = 1.315 \text{ Å}$, $C-N = 1.426 \text{ Å}$, $N=N = 1.293 \text{ Å}$, $\angle FCF = 111.84^{\circ}$, $\angle NCN = 53.95^\circ$. Comparison of these values for perfluorodiazirine with those for diazirine shows that the C-N distance is shorter (1.426 vs 1.482 Å) and the N=N

distance is longer $(1.293 \text{ vs } 1.228 \text{ Å})$ in the former than in the latter by about 0.06 Å. These arc summarized in Table 19. Hencher and Bauer compare the $C-F$ distance and the \angle FCF for F₂CN₂, 1.315 Å and 111.84°, respectively, with those found for perfluorocarbene (F_2C :) **, 1.300 Å and 104.9^o, respectively, and rationalize the differenccs **on** the basis of the greater extent of tielocalization of the carbon valence electrons in thc radical. In addition, the compression of the chargc density around the carbon atom and hence the shortening of adjacent bond Icngths when F is substitutcd for H on a carbon atom has been observed with many other molecules. The introduction of F atoms affects not only the adjacent $C-F$ bond lengths, but also adjacent bonds are shortened, as is observed in the present examples, where the $C-N$ distance in F₂CN₂ is smaller than in H₂CN₂. The longer N=N bond in F₂CN₂ than in H₂CN₂ cannot be due primarily to the presence of a three-atom ring since the $C-C$ distance in cyclopropene is 1.525 Å⁸⁹, close to that found in cyclopropane⁹⁰, while C=C is 1.286 Å, less than the $C = C$ distance in ethylene. To be consistent with these changes, it would be anticipated that the $N=N$ distance in F_2CN_2 (1.293 Å) would be less than that in N_2F_2 (1.224 Å)⁹¹, which it is not.

	pernuorogiazirine	
Parameter	F_2CN_2 ⁸⁷	H_2CN_2 50
$C-F$	1.315 Å	
$C-H$		1.09 Å
$C-N$	1.426 Å	1.482 Å
\angle NCN	53.95°	48.9°
\angle FCF	111.84°	
∠нсн		117° (± 2)
$N = N$	1·293 Å	1.228 Å
$F \cdot \cdot \cdot F$	2.178 Å	
$N \cdots F$	2.374 Å	

TABLE 19. Nuclear configuration of diazirine and perfluorodinzirine

Robertson and Merritt⁹², continuing the work on diazirines already begun in their laboratory, obtained thc ultraviolet abscrption spcctra, at high resolution, for 3-bromo- and 3-chloro-3-methyldiazirine. These compounds were prepared by the method of Graham⁵². Two electronic systems are found for these two substituted diazirincs, separatcd by 63 cni-' in 3-bromo-3-niethyldiazirine (BMD) and **137** cm-l in 3-chloro-3-methyldiazirine (CMD) , again providing evidence for the splitting of the degeneracy of the lone-pair nitrogen electrons as observed with diazirine⁷⁰. The strong progression is the symmetrical NN stretching frequency at 1412 cm^{-1} in BMD and 1454 cm⁻¹ in CMD. Spectra of solutions of BMD and CMD in benzenc, ethanol and chlorobenzene showed a blue shift of approximately 30 cm^{-1} as the solvent was changed from less polar to more polar, which indicates that the transition is $\pi^* \leftarrow n$.

The vapour-phase infrared spectra of BMD and CMD were measured in the 250-4000 cm⁻¹ region by Mitchell and Merritt⁹³. The N=N stretching vibrations in BMD and CMD are 1571.3 and 1573.4 cm⁻¹, respectively, little changed from 1626 cm⁻¹ in diazirine⁵¹ and 1563 cm⁻¹ in difluorodiazirine⁸⁵. The symmetric and antisymmetric C-N stretching vibrations were assigned as 991 and 807 cm⁻¹ in diazirine⁵⁴, and 1282 and 1091 cm⁻¹ in difluorodiazirine^{s5}, a considerable shift. Mitchell and Merritt assign the symmetric $C-N$ stretching vibration to 1071.6 cm⁻¹ in CMD and 1090.9 cm^{-1} in BMD and the antisymmetric vibration to 858 and 868 cm⁻¹ in BMD and CMD, respectively.

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Since Hencher and Bauer⁸⁷ had found that the C-N and N=N bond lengths of F_2CN_2 differ by an unexpectedly large amount (0.06 Å) from their corresponding values in H_2CN_2 , then it might be expected that other substituents would alter the ring geometry but to a lesser degree. Hence Wollrab and coworkers⁹⁴ observed the rotational spectra for dimethyldiazirine to test this hypothesis. Their measured structural parameters are given in Table 20. The $N=N$ and $C-N$ distances in diazirine and \overline{DMD} are quite similar, indicating that methyl substitution has little, if any, effect on the ring structure. A dipole moment of 2.19 D (± 0.07) was obtained from the Stark shift. The internal rotation splittings for $(CH₃)₂CN₂$ yield a barrier to internal rotation of 1129 ± 20 cal/mol and an angle of 120.4 ± 0.4 ° between the top axes. The quadrupole coupling constants (MHz) are $\chi_{aa} = -0.94 \pm 0.8$, $x_{bb} = 3.20 \pm 0.4$, $x_{ce} = -2.27 \pm 0.4$, with the *a* axis being that passing through the ring carbon atom and bisecting the $N=N$ bond, the C axis passing through the ring carbon atom and running parallel to the $N=N$ bond.

TABLE 20. Structure parameters of dimethyldiazirinc (DMD) **'I,** difluorodiazirine (DFD) *', diazirine (D) *tio* methyldiazirine (MD) **Oi** and **3-chloro-3-methyldiazirinc** (CMD) **8G**

Bond	DMD	DFD	D	MD	CMD
			Bond length (Å)		
$C-C$	1.499			1.501	1.498
$C-N$	1.490	1.426	1.482	1.481	(1.462)
$N = N$	1.235	1.293	1.228	1.235	1.241
$C-H_a$	1.080			1.098	
$C-H_n$	$1 - 100$		$1 - 09$	1.096	(1.09)
			Bond angle		
\angle NCN	$(48.9°)$ ^a	53.95°	48.9°	49.3°	50.2°
$\angle NNC$	$(65.5^\circ)^a$				
\angle CCC	119.7°				
\angle FCF		111.8°			
\angle HCH			117°		

H₈, H_a: symmetric (in-plane) and asymmetric (out-of-plane) hydrogen positions, respectively.

Continuing their examination of the effect of substituents on the diazirine ring, Scharpen and coworkers⁸⁵ obtained the rotational spectra of CH_3CHN_2 and $CD₃CHN₂$ (methyldiazirine). Their results are shown in Table 20 for convenience in comparison of the values of the various parameters. The $N=N$ bond length is the same as that found in dimethyldiazirine although the CN bonds may be slightly shorter. The internal rotation splittings gave a methyl group rotation barrier of 773.9 cal/mol. Quadrupole coupling constants (MHz) of $\chi_{aa} = +0.48 \pm 0.8$, $\chi_{bb} = -2.53 \pm 0.4$ and $\chi_{cc} = +2.05 \pm 0.4$ were obtained with the *a* axis along the CN bond and the *b* axis parallel to the N=N bond. A dipole moment of 2.03 ± 0.07 D was calculated.

A third paper, this time on CMD, was published by Wollrab and Scharpen^{9e}, and the structural parameters are addxi to Table **20** for convenient comparison. The N=N bond length is only slightly larger than the same parameter in the methylsubstituted diazirines^{94, 95}. The barrier to internal rotation of the methyl group is

 1689 ± 10 cal/mol. The chlorine quadrupole coupling constants (MHz) are $\chi_{aa} = -66.51$, $\chi_{bb} = 32.91$ and $\chi_{ce} = 33.60$ with the *ab* plane being the molecular symmetry plane containing the two carbon atoms, the chlorine atom and $H₂$. The **L:** axis is normal to this plane.

Mitchell and Merritt⁹⁷ measured the vapour and solid-phase infrared spectra for 3,3-dimethyldiazirine and 3,3-dimethyl- d_6 -diazirine in the 250-4000 cm⁻¹ region. The NN stretching frequency (solid phase) was 1586 cm^{-1} , while the symmetric and antisymmetric C-N stretching vibrations were assigned to 1136 and 788 \cdot 1 cm⁻¹, respectively.

Lombardi, Klemperer, Robin, Basch and Kuebler (LKRBK)⁷⁷ carried out both an experimental and theoretical study of the lower $n \rightarrow \pi^*$ excitation in difluorodiazirine, F_2CN_2 (DFD). They note that no microwave spectrum has been observed for DFD which may be due to **a** vanishingly small ground-state dipole moment. Using the results of Hencher and Bauer⁸⁷ they are able to establish the changes in the $N-N$ and $F-F$ distances on excitation, and hence the molecular structure of F_2CN_2 in the $n \rightarrow \pi^*$ excited state. Such changes are given as $+0.060 \pm 0.005$ and -0.034 ± 0.005 Å for NN and FF, respectively, yielding 1.343 Å and 2.144 Å, respectively, for the $n \rightarrow \pi^*$ excited state bond distances. These authors point out that since the spectrum is sensitive only to the change in the $N=N$ distance on excitation, their results cannot confirm or deny the large value previously found⁸⁷ by electron diffraction for the $N=N$ bond length in DFD. LKRBK estimate a value of $\mu' = 1.5 \pm 0.2$ D for the $n \rightarrow \pi^*$ excited-state dipole moment of DFD and a value of cnly *0.082* D for that in the ground state. Discussion of the tl:eoretical studies of LKRBK on DFD has been given in the appropriate section of this chapter. However, it may be desirable to summarize those portions of their theoretical results which relate particularly to their experimental work. The orbital energies are summarized in Table 13 of the theoretical studies section (II.C.2.b) to which reference may be made. Orbitals $4b_1$ and $9a_1$, the highest occupied and the third highest occupied, have energies of -0.4851 and -0.6251 hartrees, respectively, and represent the two 'non-bonding' orbitals associated with the $-N=N-$ group. As has been observed repeatedly experimentally in the diazirines, their mutual interaction separates them, in this case by 0.14 hartrees or 3.8 eV. The $n \rightarrow \pi^*$ transition involves the passage of an electron from $4b_1$ to the π^* molecular orbital, $2a_2$. Using the virtual orbital approximation yields a calculated singlet-singlet $n \rightarrow \pi^*$ excitation energy of 3-99 eV compared to their observcd value of 3.7 eV.

LKRBK note that the electronic spectrum of a polyatomic molecule is normally severely blue-shifted on substituting the hydrogens by fluorine, and hcnce it is surprising to find the $n \rightarrow \pi^*$ band of F₂CN₂ (3.7 eV) to the red of that of H₂CN₂ (3.95 eV). However their calculations predict the correct relative ordering of the $n \rightarrow \pi^*$ bands of diazirine (4.17 eV) and difluorodiazirine (3.99 eV).

The bond length changes on excitation may be rationalized by examining thc nature of the molecular orbitals (MO). The $9a₁$ MO tends to draw the two nitrogen atoms together while the 4b, MO acts in an opposing fashion. Hence, in an $n \rightarrow \pi^*$, the promotion of an electron from the $4b_1$ MO would be expected to cause the N=N distance to shorten, but promotion occurs to the $2a_2\pi^*$ MO which is more N=N antibonding than $4b_1$. Consequently the increased N=N distance in the upper state can be rationalized.

LKRBK also argue that the sccond band systcm obscrved in the vibronic spectra of diazirines by themselves and others⁹² could not be the second $n \rightarrow \pi^*$ transition $(9a_1 \rightarrow 2a_2)$ for two reasons. One, their MO energies would have the two $n \rightarrow \pi^*$ transitions separated by approximately $20,000$ cm⁻¹, a value of the order of 100times **larger** than that observcd experimentally. Secondly, symmetry considerations would suggest that the second $n \to \pi^*$ transition is forbidden. LKRKB tentatively assign the second system in the diazirines to the $\pi \to \pi^*$ triplet state. Behaviors and Mariti⁹⁸ pressured the high resolution electro

Robertson and Merritt⁹⁸ measured the high-resolution electronic absorption spectrum of 3,3'-dimcthyldiazirine (DMD), and two superimposed electronic systems were again observed, the weak second origin is 126 cm^{-1} to the red of the strong origin of the allowed singlet-singlet $\pi^* \sim n$ system. In addition, a third electronic system was observed in the 3900–4200 Å region, which these authors take as the triplet-singlet $\pi^* \leftarrow n$ transition. The N=N stretching vibration is again the **most** prominent fcatiire in all systcnis. Thc high-resolution electronic absorption spectrum of 3-methyldiazirine⁹⁹ was also obtained. Results somewhat similar to those for DMD were obtained, the two electronic systems again being observed, this time separated by 92 cm⁻¹. The N=N stretching vibration was again prominent near 1499 cm⁻¹.

Correlations of bond strctching frequencies, bond lengths and dissociation energies, as advanced by Bernstein¹⁰⁰, have been pursued by McKean and coworkers¹⁰¹ for a large variety of molecules among which are the diazirines. CH dissociation energies (D_{298}^0) of 110.8 and 98.4 kcal were predicted for diazirine and dimethyldiazirine, respectively.

The 3920 **A** absorptioa spectrum of difluorodiazirine \vas reinvestigatcd by Hepburn and Hollas¹⁰² and changes in structural parameters for the ground to the excited state were calculated as 0.036 and -0.038 Å for the NN and FF bond lengths, respectively, which may be compared to $+0.06$ and -0.034 Å for NN and FF, respectively, **as** found by LKRBK". Hepburn anc! Hollas also conclude that there is no positivc evidence for a second electronic system in thc **3520** *8,* region.

Accurate values of the ¹⁴N nuclear quadrupole coupling constants (MHz) have been measured by Pochan and Flygare¹⁰³, as $\chi_{aa} = +0.196$, $\chi_{bb} = -3.010$, χ_{ce} = +2.814, all \pm 0.150, where the *a* axis is coincident with the molecular symmetry axis and *is in the plane of the molecule. These values are in fair agreement with the* calculated results of LKRBK⁷⁷, $\chi_{aa} = -0.8 \text{ MHz}, \chi_{bb} = -1.88 \text{ MHz}, \chi_{ce} = +2.69$ MHz. Walsli orbitals having **sp** orbitals in the plane of **thc** ring (one pointing **away** from the N and containing a lone pair of electrons; the other entering into a $C-N$ σ bond) were used with unhybridized p_x and p_y orbitals to describe the ¹⁴N field gradients.

b. *Kinetics of thermal decomposition*. Relatively little work had bcen donc on the thermal decomposition of aliphatic diazo coinpounds, presumably because of their instability, until the discovery of convenient preparative procedures for various substitutcd diazirincs. Since thesc cornpounds appeared to be more stable than the corresponding diazo cornpounds, there followed a substantial amount of work on their decomposition. This section will deal primarily with the physicochemical aspccts of such dccornpositions.

Frey and Stevens^{tor} have investigated the thermal decomposition of dimethyldiazirine **(DhlD)** in **the** teniperatitrc rangc from **123** to 173 *"C,* and havc found this reaction, producing propene and nitrogen, to be homogeneous and first order. Below 100 torr, the first-order rate colistant decreases with pressure, but **at** 100 torr it has essentially attained its high-pressure limit. At an initial pressure of 4 torr, the measured first-order rate constants yield an activation energy of 33.2 kcal/mol. At this pressure and 145.1 "C, $k = 3.47 \times 10^{-4}$ sec⁻¹. At sufficiently high pressures the rate constant of a unimolecular reaction should be independent of pressure, and consequently the authors propose that this reaction is probably unimolecular. Further, no departure from first-order kinctics was observed even in runs carried to more than 80% decomposition. Since propene is known to be an inhibitor of

radical-chain reactions, and since the rate of decomposition of dimethyldiazirine is not affected by the accumulation of propene in the system, it was concluded that it was unlikely that any chain reaction is involved. Hence it was presumed that the decomposition is a true unimolecular reaction. These authors considered two possible mechanisms, first, a 1,2-hydrogen shift and a $C-N$ scission to form a diradical as an intermediate, followed by a breaking of the remaining CN bond and, second, a concerted mechanism in which N_2 is released and dimethylcarbene is formed as an intermediate, followed by the rearrangement to propenc. The authors argue that thc first mechanism is unlikely on energetic grounds.

Bottomley and Nyberg^{ios} have also investigated the gas-phase thermal decomposition of DMD, but in the temperature range 60-70 *"C* and at pressures of 200 torr, at very slow rates using precision gas-volumetric techniques. Their results and those of Frey and Stevens'o', aftcr adjustmcnt for pressurc differcnces, **fit a** single Arrhenius activation energy equation. They also conclude that the reaction is homogeneous. Bottomley and Nyberg¹⁰⁵ argue for a mechanism in which dimethyldiazomethane is formed as an intermediate, since the high value of the pre-exponential factor (10^{14}) is taken to indicate a transition state considerably 'looser' (i.e. highcr entropy) than the initial configuration, espccially whcn a rigid cyclic structure is broken.

The thermal decomposition of difluorodiazirine (DFD) yields primarily nitrogen. tetrafluoroethylenc, hexafluorocyclopropnnc and **perfliioro-2,3-diaza-l,3-butadienc,** with traces of other fluorine-containing compounds^{94, 106}.

The reactions may be represented as a serics of competitive consecutive steps:

$$
\begin{array}{ccc}\nF & C & \uparrow & \\
F & C & \uparrow & \\
(1) & & & & \\
CF_{2} = N - N = CF_{2} & & & \\
 & & & & \\
F_{2}C = CF_{2} & & \xrightarrow{+CF_{2}} & \searrow F_{2}C - CF_{2} \\
 & & & & \\
 & & & & \\
F_{2}C = CF_{2} & & \xrightarrow{+CF_{2}} & \searrow F_{2}C - CF_{2} \\
 & & & & \\
 & & & & \\
F_{2}C = CF_{2} & & \xrightarrow{+CF_{2}} & \searrow F_{2}C - CF_{2} \\
\end{array}
$$
\n(1)

By decomposing at **a** partial pressure of 50 torr in the presence of a 3-molar cxcess of C_2F_4 , the side reactions were minimized and decomposition occurred primarily to form difluorocarbene and nitrogen. Such primary decomposition appeared to be unimolecular and homogeneous in the temperature range of $123-170$ °C and total pressure of 100-800 torr. The rate constant was pressure dependent in this range. At a total initial pressure of 200 torr and 141.0 ^oC the rate constant was 1.27×10^{-4} sec⁻¹. At the same pressure the pre-exponential factor was 10^{13} ⁻¹ and the activation energy 32.2 kcal/mol. Neuvar and Mitsch^{lor} also prefer a mechanism involving difluorodiazomethane as an intermediate prior to loss of nitrogen.

In order to determine the effect of the electronegative chlorine atom on the decomposition of diazirines and to confirm that the decomposition was unimolecular, by following the reaction in the fall-off region, the thermal decomposition of 3-chloro-3-methyldiazirine (CMD) was investigated¹⁰⁸. In the temperature region of 106-170 °C, CMD undergoes a first-order decomposition, which is probably unimolecular, to form vinyl chloride and nitrogcn. No other products **wcrc** detected. Lo\v-prcssiirc experiments were performed at an initial reactant pressure of 4 torr and at nine temperatures ranging from 107 to 170 °C. At 147 °C the rate constant was 3.50×10^{-3} scc⁻¹, the pre-exponential factor $10^{11.7}$, and the activation energy 27.2 kcal/mol. In ordcr lo study thc effcct ol' pressurc pcrfluoropropane tvas added as an inert gas. For a reaction mixture which consisted initially of *50* torr of CMD and 300 ton of perfluoropropanc, and temperatures between :I0 and 148 *"C,* an Arrhenius expression was obtained for k_{∞} which contained 10^{14-1} as a pre-exponential factor and 31.1 kcal/mol as an activation energy. These authors retain their belief that the decompositions proceed via the formation of a carbenc. Substitution in the 3-position of a chlorine atom should lead to some stabilization of the resulting carbene as compared with its hydrogen or alkyl analogue. The energy of activation for DMD was 33.2 kcal/mol(4 torr) which probably could be adjusted to 34 kcal/mol to correspond to k_{∞} . The difference between this value and that of 31 kcal/mol obtained from CMD would suggest a significant stabilizing effect of the chlorine atom. Bridge and coworkers¹⁰⁸ take the pre-exponential factor of 10^{14} as indicating that the decomposition has a relatively small entropy of activation, and hence that in the transition state no new free rotations have developed. They consequently favour a cyclic intermediate in which both C-N bonds are weakened and $N=N$ strengthened.

The thermal decomposition of **3,3-tetramcthylenediazirine,** 3,3-pentamethylenediazirine and 3,3-diethyldiazirine was also investigated by Frey and Scapiehorn¹⁰⁹. The decompositions may be represented as follows :

3,3-Tetramethylenediazirine

 $k = 10^{13 \cdot 40}$ exp ($- 30,500/RT$) sec⁻¹ $P_{i, \text{reactant}} = 7 \text{ torr}$

3,3-Pentamethylenediazirine

$$
\bigotimes \begin{matrix} N \\ N \end{matrix} \longrightarrow \bigotimes + N_2 \qquad (101-137 \, \degree C)
$$

 $k = 10^{13.34}$ exp ($- 30,870/RT$) sec⁻¹ $P_{i.}$ reactant = 5 torr

3,3-Diethyldiazirine

 $k = 10^{13.73}$ exp (-31,890/*RT*) sec⁻¹ $P_{i.}$ reactant = 10 torr

In all cascs the reactions \vcre homogeneous and **first** order, and probably unimolecular.

1. Gcneral and theoreiical aspects of **the diazonium** and **diazo groups 27**

Although **it** does not properly fall under the heading of kinetics of thermal decomposition, nevertheless it seems appropriate to take note of **a** study of the polymerization of difluorodiazirine (DFD)¹¹⁰ because of the information which such work may provide concerning possible reaction intermediates. Pyrolysis or ultraviolet pyrolysis of difluorodiazirine in the presence of boron trifluoride produced polydifluoromethylene. Unlike Frey, Ogden and Mitsch¹¹⁰ favour the formation of a linear intermediate rather than the concerted clirnination of nitrogen in the decomposition of DFD. Under certain conditions **tetrafluoro-2,3-diazabuta-l,3-dienc,** tetrafluorethylcne and perfluorocyclopropanc arc formed from the decomposition of **DFD:** sition of DFD. Under
norethylene and perfluce
D:
 $F_2C\begin{pmatrix} N & \Delta \\ \parallel & \Delta \end{pmatrix}$

$$
F_zC\begin{matrix}N&\Delta\\ \parallel&\end{matrix}\begin{matrix}F\end{matrix}\begin{matrix}C=C\begin{matrix}F&\vdots\\ F&F\end{matrix}\begin{matrix}CF_{2}&\vdots\\ F_{2}C-CF_{2}&\end{matrix}\begin{matrix}F\end{matrix}\end{matrix}
$$

Tetrafluoro-2,3-diazabuta-l,3-diene could be derived from reaction of difluorocarbene with the linear intermediate or by insertion of difluorocarbcne into the difluorodiazirine double bond, followed by rearrangement:

$$
F_2C\begin{matrix} N\\ \parallel\\ N\end{matrix} \quad + \quad CF_2\colon \; \xrightarrow{\hspace*{1.5cm}} F_2C\begin{matrix} N\\ \parallel\\ N\end{matrix} \begin{matrix} CF_2 \end{matrix} \; \longrightarrow \; \; F_2C = N-N = CF_2
$$

However, Ogden and Mitsch¹¹⁰ could find no examples of insertion of difluorocarbene into an $N=N$ bond, and consequently favoured the linear intermediate.

Continuing their studies of the thermal decomposition of diazirines, Frey and Liu¹¹¹ investigated 3-chloro-3-ethyldiazirine, 3-chloro-3-n-propyldiazirine, 3-chloro-3-isopropyldiazirine and **3-chloro-3-r-butyldiazirine.** The overall decomposition reactions are given below:

3-Chloro-3-ethyldiazirine

$$
\sum_{i=1}^{E_1} C \left(\bigvee_{i=1}^{N} A_i\right) \longrightarrow \bigvee_{i=1}^{M} C = C \left(\bigvee_{i=1}^{C_1} A_i\right) \longrightarrow \bigvee_{i=1}^{M} C = C \left(\bigvee_{i=1}^{M} A_i\right) \longrightarrow \bigvee
$$

 $k = 10^{13.985}$ exp ($-30,450/RT$) sec⁻¹ (20 torr of diazirine and *60* torr of n-butane; 103-145 "C)

3-Chloro-3-m-propyldiazirine

Propyldiazirine

\n
$$
\begin{array}{ccc}\n\begin{array}{ccc}\n\text{propyldiazirine} \\
\text{or} \\
\begin{array}{ccc}\n\text{Ft} \\
\text{C1}\n\end{array} & \longrightarrow & \begin{array}{ccc}\n\text{Et} \\
\text{Ft} \\
\text{H}\n\end{array} & \text{Ft} \\
\begin{array}{ccc}\n\text{C} & & \\
\text{C1} & & \\
\end{array} & \longrightarrow & \begin{array}{ccc}\n\text{Et} \\
\text{H} & & \\
\end{array} & \text{Ft} \\
\begin{array}{ccc}\n\text{C} & & \\
\end{array} & \text{Ft} \\
\end{array}
$$

 $k = 10^{13.98}$ exp ($-30,980/RT$) sec⁻¹ (10 torr diazirine and 20 torr butane; 105-149 *"C)*
3-Chloro-3-isopropyldiazirine

 $k = 10^{14 \cdot 01}$ exp (-30,590/RT) sec⁻¹ **(10** torr diazirine and **20** torr butane; **107-145** "C)

3-Chloro-3-t-butyldiazirine

 $k = 10^{13 \cdot 36}$ exp (-29,500/RT) sec⁻¹ **(4** torr diazirine; **105-148** "C)

The values of the free energy of activation (ΔG^*) for decomposition of all the alkylchlorodiazirines fall within the range 28.75 ± 0.3 kcal/mol and the authors again interpret this as supporting the intermediate formation of a carbene by a concerted elimination of nitrogen.

The thermal decomposition of 3-chloro- and 3-bromo-3-phenyldiazirines (BPP) has been investigated in several solvents over the 60-110 °C temperature range¹¹². In this temperature range, **3-chloro-3-phenyldiazirine** (CPD) decomposes unimolecularly to yield nitrogen and the corresponding carbene. This carbene can either rcact with diazirine to give dimeric product or with the solvent to yield the appropriate compounds. The average activation energies for CPD and BPD are **27.8** and **27.4** kcal/mol respectively, the former being averaged over that obtained in three different solvents. The former value represents a decrease in activation energy of about 3 kcal/mol from that found for 3-chloro-3-methyldiazirine¹⁰⁸ and this is suggested to reflect the resonance stabilization effects of the phenyl group on the transition state of diazirine decomposition. The pre-exponential factors in both cases are approximately 10¹⁴, suggesting a typically unimolecular reaction with a relatively tight transition state. The results of Liu and Toriyama¹¹² also show that the rate of decomposition changes very little **as** the solvent is varied, which appears to indicate that the transition state of the rate-determining step **is** likely to bc more radical than ionic. Further, the substituent effects are interpreted as supporting a radical intermediate such as

In a second paper Liu and Toriyama¹¹³ have examined the effect of *para* substituents on the thermal decomposition of a series of **3-chloro-3-aryldiazirines** in various solvents. **A** summary of their results, together with those of previous workers, is given in Tablc **21.** They find, in agrcement with the data previously discussed in this section, that thc thermal decomposition of diazirines is uniniolecular with a preexponential factor of approximately $10^{13.7}$. Each of the solvents used (cyclohexene, diethyleneglycol monoethyl ether and dirncthyl sulphoxide) gave similar organic products and all the free energies of activation are approximately 26.0 ± 0.3 kcal/mol.

TABLE 21. Summary of kinetic data for the thermal decomposition of substituted diazirines¹¹³

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Although the substituent effects on the rates of reaction are small, nevcrtheless the following order can be discerned: p -CH₃O > p -CH₃ > p -Cl > H $\cong p$ -NO₂. Liu and Toriyama employ these results in an attempt to select one of

as the most likely transition state. The order of substituent effects would be expected to be the reverse of that observed if the charge separated transition state **I1** is invoked. Further, the lack of solvent effects renders **I1** unlikely as a transition state in the thermal decomposition of diazirines. In view of the suggestion of Schmitz¹¹⁴ that a mechanism passing through transition **statc I** would require thc carbon atom in that state to tend towards a sextet, and consequently that such **a** transition cannot be stabilized by withdrawal of electrons, Liu and Toriyama conclude that the order p -CH₃O > p -CH₃ > H > p -Cl > p -NO₂ would be expected, thus ruling against state **I** in the present case. Liu and Toriyama as a result conclude that transition state **111** is most probable and consider three possible structures for that state:

$$
\begin{array}{ccc}\n\bigcirc - N = N & \qquad & \searrow & -N = N \\
\text{(IIIA)} & & \qquad & \searrow & -N = N \\
\text{(IIIB)} & & & \text{(IIIb)}\n\end{array}
$$

State **III**c is believed to be unlikely in view of the known relative electronegativities of nitrogen and carbon atoms. The free radical form **may** be classified as ail S radical¹¹⁵ in which the observed properties are shifted in the same direction with donor 2nd acceptor substituents. The likely order of contribution of different

substituents to structure **Inb** would, however, be identical to the observed order. Consequently these authors believe that the transition state of 3-chloro-3-pmethoxyphenyldiazirine would be stabilized through both structures **IIIa** and **b,** and hence might be expected to exhibit the most rapid decomposition. Since 3 **chloro-3-p-tolyl-phenyldiazirine** has less resonance ability, both transition states **IIIa and b** will be involved to a lesser extent. With 3-chloro-3-p-chlorophenyldiazirinc, **LlIa** would then be the expected transition state, and its rate would be expected to be lower. Although structure **IIIa** is relatively equally stabilized by the p -NO₂ as by the **p-CH30** substituent, since p-NO, is strongly clectronegative **ILlb** is not likely, and the stabilizing effect from **IlIa** may be largely erased so that the obscrvcd rate is similar to that of the unsubstituted compound. Liu and Toriyama conclude that the most likely mechanism is one involving the breaking of one of the $C-N$ bonds leading to a diradical **(IIIa)** and a diazomethane **(IIlb)** intermediate, the contribution of the latter probably being small due to the lack of solvent effects. **As** has been noted earlier in this chapter (Section II.B.2) Walch and Goddard³⁵ obtained wavefunctions for diazomethane which lead them to suggest a singlet biradical structure for the ground state of diazomethane.

Although the recent calculations by Tschuikow-Roux and Jung¹¹⁶ on a threecentre decomposition reaction for perfluorodiazirine might be included in the section on theoretical studies of diazirines, they are included here because results of such calculations do provide information on **a** possible decomposition mechanism. These authors begin with the stable configuration for perfluorodiazirine, that is, with the NCN plane perpendicular to the FCF plane. The molecule is then perturbed within one of three models where the $\angle NCN$ bending mode is used as the reaction coordinate. In model I, the $C-N$ bonds and the $N-N$ bond are taken to have half-bond orders and bond order 2.5, respectively. The C-F bond distance and the LFCF remain unchanged from the initial state. In model **11,** the total bond order is taken as fixed, the $C-N$ bonds and $N-N$ bond then have half-bond orders and bond order **3,** respectively. The *L* FCF is chosen as the average of that in perfluorodiazirine and the $CF₂$ radical. The $CF₂$ bond distance is unchanged from the initial value. Model **I11** retains the assumption of conservation of total bond order but postulates that the fractional decrease in bond order of a bond bcing partially broken is equal to $\Delta n = 1/3$, the fraction of the number of bonds undergoing changes in the ring opening. Consequently, 2/3 and 8/3 bond orders are assigned to the $C-N$ and $N-N$ bonds, respectively. The CF bond lengths and \angle FCF are used as in model **11.** Basically what is then required is the minimum energy corresponding to each of these sets of constraints, and these are calculated as 60.3, 75.6 and 32.7 kcal/mol, for models **I, 11,** and **111,** respectively, compared to the cxpcrimental activation energy of 32.2 kcal/mol¹⁰⁷. Model III apparently is the most appropriate and in support yields a calculated pre-exponential factor in reasonable agreement with the experimental value. It should be emphasized, however, that however interesting such results may be, they do not provide sufficient information to permit a discrimination between a thermal decomposition mechanism involving a linear intermediate and one involving a concerted elimination of nitrogen. However, the weight of evidence prescnted in this section appcars to favour a linear intermediate in the decomposition of diazirines.

111. DIIMIDES (DIAZENES)

As indicated in the introduction to this chapter, although not strictly speaking within the purview of this chapter, diimides have bccn included for purposes of comparison and completeness. Rcfcrencc should also bc madc to the chapter by Dr M. B. Robin

on 'Electronic Structiircs' ivhich appcared in the volume on *Tlrc Clrcmis/ry of /he* Hydrazo, Azo and Azoxy Groups in the present series. It is hoped that the present discussion will supplcment, rather than duplicate, the information given in the chapter mentioned above.

A. The **Prototypes,** *N,H+* **and** *N,H*

The molecular ion, $N₂H⁺$, may quite reasonably be considered as representing the simplest azo or diazo compound. In 1974 Turner 117 reported the discovery of three closely spaced new interstellar lines near 93.174 GHz. Green and coworkers¹¹⁸ suggested the molecular ion N_2H^+ to be an excellent candidate for the carrier of Turner's lines and, assuming the ion to be linear, have performed ab initio calculations, using a double-zeta basis set, as a function of $r(NN)$ and $r(NH)$, the nitrogennitrogen and nitrogen-hydrogen bond lengths. The minimum energy was found when $r(NN) = 1.099 \text{ Å}$ and $r(NH) = 1.041 \text{ Å}$, and the frequency of the $J = 1 \rightarrow 0$ rotational transition was calculated to be 92.2 ± 1.8 GHz, in good agreement with the observed value given previously. Similarly the quadrupole coupling constant for the outer nitrogen, deduced from observations as -5.7 MHz, was calculated to be -5.3 ± 0.5 MHz, while that for the inner nitrogen was calculated as -1.2 ± 0.5 MHz, a contribution to thc hyperfine structure too small to be resolved in the sourccs so far observed. However, the agreement between the aforementioned experimental and calculated values seems to provide rather good evidence for the association of Turner's lines with N,H+.

Even more recently, Hillier and Kendrick¹¹⁹ have extended the previous calculations on N_2H^+ . Restricted Hartree-Fock (RHF) calculations of the electronic ground state of N_2H^+ were performed with inclusions of nitrogen d and hydrogen p polarization functions.

Large-scale configuration interaction calculations were then performed including single and double excitations from the RHF configuration and again assuming the molecule to be linear. Values of 93.1 GHz , -4.70 MHz and -1.00 MHz were obtained from the calculations for the frequency of the $J = 1 \rightarrow 0$ rotational transition (actually B_e rather than B_0 was calculated), the quadrupole constant for the outer nitrogen and that for the inner nitrogen, respectively. Quite obviously the agreement of the first value with experiment is considerably improved, while that of the second suffers some degradation. The inclusion of correlation alters other calculated values, of course, and -109.414325 hartrees, 1.097 Å and 1.031 Å are found for the total electronic energy, $r(NN)$ and $r(NH)$, respectively, as compared with - 109.163339 hartree, **1.065 A** and **1.023 A,** respectively, obtained with the RHF calculations.

Vasudevan and coworkers¹²⁰ have carried out extensive calculations on N_2H^+ . Their calculated equilibrium values compared with those mentioned previously and with those of Forsen and Roos¹²¹ are provided in Table 22. Vasudevan and coworkers performed calculations as a function of NH and NN distances **as** well as the HNN angle, and found N_2H^+ to be linear in its ground state.

A6 irritio molecular orbital calculations have also been performed'?? on **:!iC** free radical HNN, using the restricted open-shell method of Roothaan¹²³ with the extended $4-31$ G basis set of Pople and coworkers¹²⁴. Standard molecular exponents were used for HNN \cdot and N₂, but the energy used for the H atom was that obtained with optimum atomic exponents. The internuclear separations of NN and NH were calculated as 1.18 and 1.020 Å, respectively, and the $H - N - N$ angle was obtained as 118°. The HNN· radical was calculated to be less stable than an H atom and an **N2** molecule in thcir ground states, the radical having **a** dissociation energy of

r(NH) (Å)	r(NN) (Å)	NH Stretching force constant (mdyn/A)	Total electronic energy (hartree)	Dissociation energy (eV)	Proton affinity (kcal) mol)	Reference
1.041	1.099					118
1.031	1.097		-109.414325 (CI)			119
1.023	1.065		-109.163339 (RHF)			119
1.026	1.093	7-1	-109.0748 (SCF)	5.48	126.5	120
1.027	1.093°	4.9	-109.2443 (CI)	5.46	126.0	120
1.024	1.093 ^a		-108967		$127 - 7$	121
	1.077		-109.157 ^b		123	121

TABLE 22. Calculated equilibrium parameters for N_2H^+

^a Assumed.

With d-type polarization functions.

 -9.0 kcal/mol. The activation energy for the decomposition of HNN \cdot to N₂ and H was found as 22.6 kcal/mol.

Very recent *ab initio* SCF and CI calculations¹²⁵ on the ground and excited states of the HNN radical, using a bond length for $N-H$ of 1.058 Å and either 1.093 Å or 1.199 Å for N-N, predicted the ground state to be non-linear with a $H-N-N$ angle of 124° and a binding energy of 2.2 kcal/mol.

B. Diirnide (Diazene)

1. Introduction

Although the molecular ion, N_2H^+ , may be taken as the simplest representative of the class of molecules containing the NN bond, the molecule $H-N=N-H$, diiniide (or diazene) is perhaps a more appropriate candidate for this position, since **it** is the simplest uncharged molccule of this type. Like diazomethane, as discussed earlier, diimide has a number of possible isomers, the *cis* and *trans* form, and 1,1dihydrodiazine $(H_2 \bar{N} = \bar{N})$. Diimide is the parent molecule of the azo compounds. The unsymmetrically substituted diazenes, of which 1,1-dihydrodiazine is the parent compound, are both important and interesting molecules. Lemal¹²⁶ has pointed out that these unstable isomers of azo compounds might be expected to lose molecular nitrogen with ease and hence scrve as an important method for producing carboncarbon bonds, as azo compounds do. However, as a consequence of their instability, other reactions appear to occur preferentially. Nevertheless, the properties ana reactions of 1,1-disubstituted diazenes remain of importance in chemistry. Unfortunately, discussion of these compounds would lead us too far afield of the intention of this volume. Hence, attention will be limited to $1,1$ -dihydrodiazine itsclf for comparison with the *cis* and *fratis* forms of diiniide. **A** comprehensive review of nitrcncs of which Reference 126 is a part has been published previously.

The nomcnclature unfortunately presents some difficulty. Lema1126 has pointed out that N_2H_2 is an aminonitrene, but prefers to label R_2N_2 , where the two R atoms or groups arc attached to the same nitrogen, as I,]-disubstituted diazene, or simply diazenes. Hünig and coworkers¹²⁷ suggest that the systematic name diazene would be appropriate if NH was analogously named 'azene'. They employ diimine in preference to diiniide on the basis of less ambiguity and continued association with the older literature. In the present article, diimide and diazene will be employed interchangeably where no confusion can arise.

hy where no contusion can arise.
Although a number of workers believed that they had direct or indirect chemical evidence for the existence of diimide, it was not until 1958 that Foner and Hudson¹²⁸ first detected diimide by mass spectrometry as an *mle* 30 peak during the decomposition of hydrazoic acid or hydrazine in an electric discharge under reduced pressure. A fast-flow system was used and the diimide was detected downstream from the discharge. Foner and Hudson also showed that N_2H_2 could be trapped at -196 °C as a yellow solid consisting of ammonia and N_2H_2 , and regenerated in the gas phase on warming. Thermal decomposition of anthracene-9,10-biimine at 100 °C *in vacuo*

at the inlet of a mass spectrometer also apparently produces diimide¹²⁹, thereby suggesting a relatively high thermal stability for diimide. In contrast, diimide cannot be detected during the thermal decomposition of hydrazine in argon at **1470-** 2770 °C¹³⁰. Photodecomposition of hydrazine¹³¹ with vacuum ultraviolet radiation and the combustion of hydrazine at low pressure¹³² has also led to the observation of diimide.

2. Spectroscopy and structure

shown below: The various nuclear configurations which are, in principle, possible for N_2H_2 are

One of the prime tasks of the spectroscopists was not only to determine in which of these forms diimide exists in its most stable form, but also to elucidate the nature of the electronic states, and in particular the electronic ground state. The next section attempts to trace the development of the answers for these and other questions, and to indicate those which remain unanswered at this time.

Walsh¹³³, in his interesting and far-reaching semi-quantitative predictions of shapes, spectra, and the electronic orbitals of polyatomic molecules, suggested that diimide would possess a planar non-linear structure. Wheland and Chen¹³⁴ assumed NH and NN distances to be **1-014** *8,* and **1.23 A,** respectively, and calculated, using simple, semi-empirical, LCAO-MO theory with overlap integrals neglected, a value of 100" for the most stable NNH angle. With the assumption that the *cis-lrnns* isonierization passes through a planar intermediate in which one NNH angle is 180°, whereas the other is allowed to take on whatevcr value leads to the most stable configuration, an activation energy of 33 kcal/mol was calculated for an activated complex in which the NNH angle that is not equal to 180° has the value of 76° .

Blau and coworkers^{135, 136} obtained the infrared spectrum of solid diimide between 350 and 4900 cm⁻¹ and that of the gas in the 3200 to 1300 cm⁻¹ regions by decomposition of hydrazine with a microwave discharge. Infrared bands were observed at 3205, 3095, 2898, 1821, 1495, 1406 and 1362 cm⁻¹. They deduced that the solid at liquid nitrogen tcmperatlire has a planar structure and that the molecule appears to be primarily in the cis form, with molecular parameters of NH, NN and \angle HNN, being assumed as 1.014 **A,** 1.230 *8,* and IOO", respectively. Unfortunately, the structure of the molecule in the gaseous state could not be unambiguously determined from the spectrum.

Dows and coworkers¹³⁷ investigated spectra of glow discharge decomposition products of $HN₃$ after condensation at 90 K and during slow warming. Absorptions at 860 and 3230 cm⁻¹ were attributed to N_2H_2 . Three prominent absorptions at 1290, 1325 and 3080 cm⁻¹ observed in the photolysis of $NH₃$ in solid nitrogen were attributed13* to intermediates D, **E** and F, respectively, possibly corresponding to NH_3 , N_2H_2 and NH, respectively. It was established¹³⁹ that species D contained two hydrogen atonis but it was not possible to distinguish between the amine radical, *cis-* and *trans*-diimide, or NH_2N_3 . Milligan and Jacox¹⁴⁰ propose that D and E may be N_aH_2 or $HN=N-N=NH$. Emission features near 3000 cm⁻¹ from an ammoniaoxygen flame¹⁴¹ were interpreted as due to NH_2 , but with N_2H_2 as an alternate possibility. Rosengren and Pimentel14? produced diimide by photolysing hydrazoic acid, both unlabelled and labelled with deuterium and ^{15}N , in solid nitrogen at 20 K. To trans-diimide the following features were positively assigned: 1285.8 cm⁻¹ (HNNH), 1480.5 and 1058 cm⁻¹ (HNND), 946.2 cm⁻¹ (DNND). The only other absorptions which displayed isotopic frequency patterns appropriate to diimide were bands at 1279 and 3074 cm⁻¹ which they assigned to *cis*-HNNH.

Trombetti¹⁴³ examined the infrared spectrum of N_2H_2 vapour in the 3100 cm⁻¹ region and the vacuum ultraviolet spectra of both N_2H_2 and N_2D_2 , and concluded from an analysis of the former spectrum that N_2H_2 in the ground state has a planar *trans* conformation with $r_{N-N} = 1.238$ and $\angle N-N-H = 109^{\circ}$, assuming r_{N-H} to be between 1.05 and 1.08 Å. Trombetti attributes five bands to diimide in the discharge products (77 K) of N₂H₄ (3095, 2950, 2900, 1404 and 1359 cm⁻¹) and N₂D₄ (2521, 2291, 2199, 1032 and 999 cm⁻¹). Trombetti interpreted his results in terms of a *trans* structure only, assigning the three infrared active fundamentals of trans- N_2H_2 at 3095, 1404 and 1359 cm⁻¹ (2291, 1032 and 999 cm⁻¹ for N_2D_2). The remaining bands were tentatively assigned as combinations involving unobserved Raman-active fundamentals which were deduced to be 1552 and 1496 cm⁻¹ (1498 and 1167 cm⁻¹ for N_2D_2). Trombetti also observed two electronic band systems near 3500 and 1730 *8,* and, from intcnsity variations in the 1730 *8,* system, decided that the ground state is a triplet $({}^{3}B_{g})$ and that the first excited state may have a B_{u} symmetry. Trombetti¹⁴⁴ has also measured the infrared spectra of solid N_2H_2 and N_2D_2 from 600 to 4000 cm^{-1} at liquid nitrogen temperature. For both compounds five bands were observed and tentatively assigned to the three allowed fundamentals $(N_2H_2$: 1359, 1404, 3095; N_2D_2 : 999, 1032, 2291 cm⁻¹) and to two combination bands $(N_2H_2: 1496, 1552; N_2D_2: 1167, 1498 cm^{-1})$ of *trans*-diimide.

Bondybey and Nibler¹¹⁵ have prepared N_2H_2 as a discharge product of N_2H_4 and have studied the infrared and Raman spectra for both solid and matrix isolated

HNNH, HNND and DNND. The five in-plane vibrations of isolated *trans*-HNNH have all been determined: Raman A_g modes, 3128, 1583 and 1529 cm⁻¹; infrared B_u modes, 3131 and 1286 cm⁻¹. Isotopic substitution established the characteristic NN stretch as the 1529 cm⁻¹ band. The A_u out-of-plane torsion was not definitely assigned but could be a weak infrared feature observed at 832 cm^{-1} . No evidence of cis-diimide was dcteclcd. Dilution studies of the solid discharge products showed that a number of hydrogen-bonded complexes form between HNNH and NH₃, and Bondybey and Nibler suggest that their work indicates that the carlier interpretations13G, **143** of the solid spcctra in ternis of 'isolated' *tram-* and cis-diiniide must be discountcd. Thesc authors also concludc that thcir results are not inconsistent with Trombetti's conclusion that the electronic ground state of N_2H_2 is a triplet.

The infrared absorption spectrum of diimide in the $3.2 \mu m$ region has been reexamined at higher resolution than previously employed¹⁴⁶, and that of dideuterodiimide (N_2D_2) at 4.3 μ m was also obtained for the first time. The following geomctrical paranictcrs werc obtained for the planar *trnm* conformation of diimidc: r_{N-H} = 1.028 Å, r_{N-N} = 1.252 Å, $\angle N-N-H = 106^{\circ}51'$. It is interesting to note that this N-N bond distance is the same as in azomethane, while $\angle N-N-H$ is lower than $\angle N-N-C$ in the same molecule by approximately 5 degrees^{147, 148}. Carlotti and coworkers¹⁴⁶ argue that the evidence provided on line intensities agrees with the intensity alternation which would be expected for an A_g or A_u ground state. This disagrees with the earlier conclusions of Trombetti¹⁴³, whose spectral data were recently confirmed by Willis and Back¹¹⁹. Hence Carlotti and coworkers offer the possibility that the ultraviolet and the infrared spectra may not have the same lower state. They speculate that it may be possible that planar trans-diimide converts slowly to a less stable *cis* form which absorbs in the ultraviolet, so that an explanation for the apparently long lifetime of the ultraviolet absorption could be provided.

Back and coworkers¹⁵⁰ have examined the absorption spectra of N_2H_2 and N_2D_2 in the gas phase in the region 3000-4300 Å. The spectra were attributed to the ${}^{1}B_{g} \leftarrow {}^{1}A_{g} (\pi^{*} \leftarrow n_{+})$ transition of *trans*-diimide, which the authors suggest may be allowed as a consequence of vibronic interaction. From Franck-Condon calculations the $H-N=N$ angle in the upper state was calculated to be 132°, or 25 degrees larger than the ground-state value and the increase in the $N=N$ bond length was given as approximately 0.05 **A.** As these authors point out, it might be expected that diimide becomes non-planar in its first excited singlet state. However, if the excited state of diimide were non-planar, progressions in the spectra corresponding to excitation of the ν_4' torsional vibration should be observed, but there are no unassigned bands of significant intensity.

The near-ultraviolet absorption spectrum of diimide in liquid ammonia at -50 °C has been measured¹⁵¹. At this temperature the diimide and ammonia should exist as a pale yellow liquid. Compared with the gas-phase spectrum, the absorption of diimide in aminonia is shifted to the red by approximately 500 **A,** broadcned considerably, and the sharp vibrational structure found for the gas phase essentially disappears. These effects are attributed to a strong interaction between ammonia and diimide, possibly through hydrogen bonding. A complementary experiment with diazomethane, in which a much smaller shift to the red was observed, leads Back and Willis to conclude that the bonding of the diimide hydrogens to the $NH₃$ lone pairs is much more important than the bonding of the $NH₃$ hydrogens to the lone pairs of azomethane or diimide, which might be expected from the strong intramolecular interaction between the tivo lone pairs in azo compounds which make them less available for intermolecular interactions.

A brief surnrnary of this section may be both appropriate and useful. The evidence would appear to favour the planar *trans* configuration of diiniide as its most stable structure with values for the geometrical parameters as shown:

The electronic ground state appears to be a singlet and the first excited state apparently possesses a planar structure with an $N=N-H$ angle widened to 132°.

3. Theoretical studies

The first theoretical studies of diimide were those of Walsh¹³³ and Wheland and Che n^{134} . The results and conclusions of these authors have been briefly described in the previous section and only brief additional references will be made to them here. In 1953, Walsh¹³³ predicted from the number of electrons in the molecule for the then unknown N_2H_2 , a planar bent ground state, existing in either a *cis* or a *trans* form. Wheland¹³⁴ could not distinguish between the cis and *trans* forms because of the assumptions made in his calculations, but he did find the most stable NNH angle to be about 100° and an activation energy of 33 kcal/mol for the cis-trans isomerization.

The spectroscopic studies in the mid-60's on the structure of diimide prompted Alster and Burnell¹⁵² to carry out extended Hückel (EXH) calculations on the cis-trans thermal isomerization of N_2H_2 . They considered three possible activated complexes: (i) an asymmetric bent structure in which one NNH angle was held at 180" while the other NNH angle was varied to produce the most stable configuration (lowest electronic energy), (ii) a 90" twisted structure in which one NNH bond is held at 90° to the second NNH bond and both angles are varied to minimize the energy, (iii) the linear N_2H_2 structure. The NN and NH bond distances were held fixed at 1.22 and 1.03 **A,** respectively. Values of 128" in the cis, trans and asymmetric bent geometries and 143[°] in the 90[°] twisted structure for the NNH angle were found to minimize the energy. Energy differences from the cis form, which these calculations predicted to be of lowest energy, are 8.5, 11.6, 16.6 and 20.6 kcal/mol for the trans, asymmetric bent, 90" twist and linear forms, respectively. Hence the 8.5 kcal/ mol corresponds to the *cis-trans* transition energy and the remaining values to the predicted activation energies for the isomerization of the cis form.

Very shortly after, Robin and coworkers¹⁵³ published the results of their work on the electronic states of azoalkanes. They performed ab inirio calculations in a Gaussian-type orbital (GTO) basis with limited configuration interaction, and, in addition, measured the electronic spectra of a series of *trans*-azoalkanes, $R-N=N-R$, where R was CH_3 , C_2H_3 , $(CH_3)_2CH$ and $(CH_3)_3C$, also *trans*difluorodiazirine and one cis-azoalkane, 3-methyl-l-pyrazoline,

In their calculations, *cis-* and *trans-HN=NH* were taken to have planar structures with $\angle NNH = 112^{\circ}39'$, $D(N-H) = 1.021$ Å and $D(N=N) = 1.240$ Å. Two basis sets wcre employed, a conventional 'energy-minimized' gaussian basis and an augmented GTO basis. The authors note that it appears that the energy-minimized basis yields excited states of unusually high energies due to a neglect of the tails of the wavefunctions. It is predicted that in the case of $trans\text{-dipnide}$, the lowest excited triplet state will lie below the singlet ground state when calculatcd in the energyminimized basis, but **2** eV abovc it when calculated in thc augmcntcd basis. Consequently their calculations primarily employed the augmented basis. Ionization potentials of 10.97 and 10.46 eV were calculated for *cis-* and *trans-diimide*, respectively, compared to the Foner and Hudson¹²⁸ value of 9.85 eV.

Gordon and Fisher¹⁵⁴, as part of a study of the isomerization mechanism of diazacumulenes using the INDO mcthod, examincd diiniide and difluorodiimide. In both cases the *cis* form was found to be thermodynamically more stable than the *tram* form by **4.5** and **4.1** kcal/mol for diimide and difluorodiimidc, rcspectivcly. Activation energies of 46 and 72 kcal/niol arc predicted for inversion and rotation on one nitrogen, respcctively, if only singlet states are involved \vith diimide. **A** triplet isomerization proceeding by a rotation around the double bond has a calculated activation energy of 40 kcal/mol for diiniide. For difluorodiimidc the calculated singlet and triplet activation energies are 68 and 52 kcal/mol, respectively.

SCF-LCAO-MO calculations using Gaussian functions and cmploying the IBMOL program with a basis set consisting of 9s and **5p** for N and C and 4s for Fl were reported by Lehn and Munsch¹⁵⁵. The bond lengths of diimide were taken from the results of Blau and Hochheimer¹³⁶ and the NNH bond angles were optimized. The results are given in Table 23. The *trans* form is found to be more stable than the *cis* by 10.5 kcal/mol. Thermal *trans*-to-cis isomerization by inversion at one nitrogen (activation energy of 50 kcal/rnol) is predictcd, thc activation encrgy for rotation around the $N=N$ bond being substantially higher at 84.1 kcal/mol. Veillard. Lehn and Munsch¹⁵⁶ had pointed out earlier that nitrogen inversion is accompanied by a $2s \rightarrow 2p$ electronic population transfer at the inverting nitrogen atom. With diimide a decrease of approximately **0.3** and 0.1 elcctron was observed for tlic nitrogen 2s and hydrogen (on the inverting nitrogen) 1s populations, respectively. Simultaneously the population of the nitrogen **2p** orbital lying in the direction of the **lone** pair increases by 0.47 electron. From a comparison of population changes on inversion in other molccules the authors conclude that **a** high inversion barrier corresponds to a large population transfer.

Symmetry groups	Conformation	Optimized NNH angle	Total cnergy (hartree)	' Energy barriers (kcal/mol)
$C_{\rm 2h}$	trans	70°	-109.924240	
C_{2V}	cis	63°	-109.907558	cis -trans = 10.5
$C_{\mathbf{h}}$	bent (b)	68°	-109.844374	b -trans = 50.1
$D_{\tt{wh}}$	$linear$ (1)	0°	-109.737979	1- <i>trans</i> = 116.9
C_{2}	perpendicular (perp)	56° a	-109.790334	perp- <i>trans</i> $= 84$ ·1

TABLE 23. Computed angles and energies for diimide¹⁵⁵

^aAngle not optimized.

Scliaad and Kinser **(SK)Is7** point out that diiniidc gives *cis* addition of hydrogcn to a double bond, which is usually interpreted in ternis of a cyclic transition *state,* which seems to imply a *cis* structure for diimide or perhaps an easy interconversion between the *cis* and *trnris* forins. With this in mind, calculations were done employing a minimal basis of Slater-type orbitals (Is for each hydrogen and Is, 2s, 2p,, **2p,,** 2pz for each nitrogen and carbon atom) expressed in terms of normalized gaussian functions. In their calculations on the various forps of diimide, **SK** optimized bond angles and bond lengths. The same conformations of diimide as those studied by Lehn and Munsch¹⁵⁵ were also considered by Schaad and Kinser, except that in place of the perpendicular form, a twisted form with a variable out-of-plane angle was employed. Only the closed-shell singlet states of the electronic configurations were considered.

Table **24** shows the results of Schaad and Kinser for the various configurations. When the configurations yielding the lowest energies had been found, the bond lengths and angles were varied to minimize the energy of that configuration. The lowest energy configurations found were:

$$
cis: \qquad (1a_1)^2 (1b_2)^2 (2a_1)^2 (2b_2)^2 (3a_1)^2 (4a_1)^2 (1b_1)^2 (3b_2)^2
$$

 $trans:$ $(1a_g)^2(1b_u)^2(2a_g)^2(2b_u)^2(3a_g)^2(3b_u)^2(1a_u)^2(4a_g)^2$

Table **24** summarizes the values of the total electronic energy and bond angles and lengths for the optimum geometries. The table indicates that the *trans* structure is **7.6** kcal/mol lower in energy than that of the cis and the lowest energy path for the trans-cis isomerization involves the bent conformation with an activation energy of **72.4** kcal/mol. These authors employ group frequencies to calculate statistical mechanical values of various thermodynamic functions. It turns out that the ΔG_{298} value for cis-trans isomerization is 7.5 kcal/mol yielding a value of 2.7×10^5 (25 °C) for the equilibrium constant, $K = [trans]/[cis]$, which implies that approximately 2.2×10^{18} molecules of *cis* isomer are present per mole. Hence, double bond reduction could take place only with the cis isomer which is replenished through equilibrium with the trans form.

	Electronic		Bond lengths (A)	Bond angles	
Conformation	energy (kcal/mol)	NN	NH	NNH	ϕ^*
cis	$-66,844.1$	1.27	$1 - 13$	112°	0 ⁵
trans	$-66,851-7$	1.28	$1 - 13$	104°	180°
linear	$-66,654.9$	2.28	2.77	180°	
twisted	$-66,767.6$	2.53	2.24	105°	90°
bent	$-66,779.1$	2.42	2.42 2.08	115° 180°	

TABLE 24. Total electronic energy **and** geometrical parameters for

Wong and coworkers¹⁵⁸ performed LCAO-MO-SCF calculations employing a basis set of Gaussian lobe atomic wavefunctions on trans-diimide, cis-diimide and 1, I-dihydrodiazine in the expected closed-shell, ground-state, electronic configuration with full geometry optimization. In addition they made additional calculations for an open-shell configuration using one of Trombetti's experimentally measured geometries for trans-diimide. **A** summary of their results is given in Table **25.** The *in situ* adjusted Gaussian lobe function calculations referred to in the table involve the determination of the long-range components of the **2s** and 2p groups of nitrogen by the SCF procedure. The extended Gaussian lobe function caiculation makes use of a basis in which **a** one-component s orbital and three one-component **p** orbitals centred at the middle of the NN axis have been added as supplementary functions.

Examination of the wavefunctions and coefficients of the molecular orbitals **(MO)** shows that the first two **MO's** are nitrogen **1s** orbitals, and the remainder could approximately represent one σ_{NN} bond, two σ_{NH} sp² bonds, two sp² lone pairs, and one pure P_x NN bond.

			Bond lengths		
Basis set	Structure	NΝ	NH	Bond angle HNN	$E_{\rm tot.}$ (hartree)
SCF-LCAO-GLO in situ adjusted	trans ^a cis^a $H_2N^+ = N^{-b}$	2.370 2.353 2.403	1.90 1.91 1.89	102.5° 115.5° 122°	-109.93017 -109.92913 -109.92233
Extended GLO	trans cis $H_2N^+ = N^-$	2.260 2.353 2.423	1.90 1.91 1.90	102.5° 115.5° 122°	-109.95274 -109.94423 -109.94145
Open shell $^{1}B_{c}$ ${}^{3}B_{\alpha}$	trans	2.340	2.01	109°	-109.79297 -109.81556
Closed shell $A_{\mathbf{g}}$	trans	2.340	2.01	109°	-109.9340

TABLE 25. Results of Wong, Fink and Allen¹⁵⁸ for diimide

^a Same electronic configurations as given by Schaad and Kinser¹⁵⁷.

^b $(IA_1)^2(2A_1)^2(3A_1)^2(4A_1)^2(1B_2)^2(5A_1)^2(1B_1)^2(2B_2)^2$.

It is instructive to examine the ground-state molecular-orbital symmetries and order of occupancy. The 1B_1 of $H_2N^+=N^-$ and *cis*-N₂H₂. 1A_u of *trans-N₂H₂* are all $2p_x \pi$ bonded molecular orbitals. The $2B_2$ in $H_2N^+=N^-$ is a pure $2p_z$ bonded with hydrogen atoms. In $3B_2$ of *cis*-N₂H₂, $4A_g$ of *trans*-N₂H₂, the molecular orbitals are hybrid orbitals mixing 2p,, 2p,, **2s,** Is *of* nitrogen and bonded to hydrogen atomic orbitals.

The theoretically predicted ionization potentials for trans-diimide, cis-diimide and 1 ,I-dihydrodiazine are 10.87, 10.84 and 9.95 eV. respectively, which may be compared with the experimental value of 9.85 eV¹²⁸. Use of the relation

$$
E(\text{binding}) = E(\text{total}) - \sum_{i} E_i \text{(atom)}
$$

where $\sum_i E_i$ (atom) = -109.77432 hartree for N_2H_2 yields binding energies of 4.85, 4.62 and 4.55 eV for *trans*-diimide, *cis*-diimide and $H_2N^+ = N^-$, respectively. Hence all these species are predicted by Wong and coworkers to be possible, although diimide is thermodynamically unstable with respect to $N_2 + H_2$, as shown by Schaad and Kinser¹⁵⁷. From Table 25 it is readily seen that the calculated order of stability places *trans*-diimide first, followed by cis-diimide, then $H_2N^+=N^-$ followed by the *frams* form in the triplet state. Hence a planar, *trams* structure for a ¹Ag electronic ground **state** is favoured.

Tinland¹⁵⁹ applied the CNDO/S method¹⁶⁰ to the cis and *trans* forms of diimide with the geometries of Robin and coworkers¹⁵³. The computed ionization potentials of *cis-* and trorrs-diimide are 12.61 and 11.96 eV, respectively. and it is predicted that the *fratis* form is more stable than thc *cis* forin by 0.47 **eV.** The general features of the shape of diimide, molecular cnergy levels aad simplc molecular orbital pictures as determined by extended Hückel calculations are presented by Gimarc¹⁶¹. Gimarc also makes use of symmetry arguments in discussing probable *cis-trans* isomerization mechanisms. In the *trans* isomer of N_2H_2 , the highest occupied orbital is $3a'_8$ while in the *cis* isomer it is $2b_8$. On rotation from *trans* to *cis* through the non-planar transition state, the *trans* $3a_k$ orbital moves much higher in energy to become a₂ in the *cis* form. On the other hand, the lowest unoccupied orbital in the *trans* form is b''_g which falls in energy to become $2b_2$ in the *cis* form on rotation from */rms* to *cis.* Hencc the orbitals cross, and consequently the electrons in the highest occupied orbital $(3a'_g)$ of *trans*-diimide, on rotation from *trans* to *cis* through the non-planar transition statc, would have to transfer to the b orbital, which falls during rotation. to avoid moving to the higher energy. **As** Gimarc points out, this would be a violation of the principles of conservation of orbital symmetry¹⁶². In the C_s pathway the a, b symmetry is lost and $3a'_g$ can be converted directly to $2b_2$. Hence the planar C_s transition state for the N_2H_2 isomerization is strongly favoured since thc linear state can be excludcd on cncrgetic grounds and the non-planar rotation by symmetry conditions.

The $n \rightarrow \pi^*$ transition energies and one-electron orbital energies for the *cis* and *Irnm* forms of diiniide, as obtained from ab *itiitio* minimal basis **SCF-CI** calculations, are reported by Ditchfield and coworkers¹⁶³. The minimal basis set of Slater-type atomic orbitals (STO) is replaced by a sum of four gaussian-type functions, one such sum for each STO. Wavefunctions, ψ_i^l , describing singly excited configurations are formed from the ground-state wavefunction by promoting electrons from doubly occupied orbitals ψ_i to virtual orbitals ψ_i . Mixing of configurations representing different singly excited virtual states is then performed. These authors calculate that the *trans* isomer should be more stable than the *cis* for the ground states.

Since the azo group contains two pairs of non-bonding electrons, there are two possible $n \rightarrow \pi^*$ transitions, although only one low-energy u.v. band is generally observed. **If** diirnidc exists in the *fratis* form, thcn **a** relatively strong interaction between the lone pairs will occur, and consequently a large splitting in the energies of the two *n* orbitals, and finally two $n \rightarrow \pi^*$ transitions of quite different energies. Ditchfield and coworkers find a splitting between the two *n* orbitals of approximately 5-6 eV and 2.5-3.5 eV in the *trans* and *cis* compounds, respectively. The two $n \rightarrow \pi^*$ transitions in the *trans* compound are separated by about 6 eV, but only 2 eV in the *cis* compound. Values reported for the transition energies for diimide arc **4.37** and 3.37 eV for *cis* and */rotis* forms, respectively, to be compared with Trombetti's experimental value of 3.55 eV for the *trans* structure.

Genson and Christoffersen¹⁶⁴, continuing their work on *ab initio* calculations by a technique employing molecular fragments and employing the floating spherical gaussian orbitals of Chu and Frost¹⁶⁵ as basis functions, applied this method to a number of nitrogen-containing molecules, including diimide. Bond lengths of 2.30 and 1.911 bohr for $N-N$ and $N-H$ bond distances, respectively, were used. 2.30 and 1.911 bohr for N-N and N-H bond distances, respectively, were used.
They too find that the *trans* form is the more stable with energy of -93.56754 hartree They too find that the *trans* form is the more stable with energy of -93.56754 hartree
compared to -93.56048 hartree for the *cis* form. The value of *E(cis) - E(trans*) was calculated as 4.52 keal/mol.

Ab initio calculations were performed with the IBMOL-5 program, using a gaussian-type basis, on I ,I -dihydrodiazine and on *cis-* and /ram-diiniides by Wagniere¹⁶⁶. The geometry of Robin and coworkers¹⁵³ was employed, namely $I(N) = 1.240 \text{ Å}$, $I(NH) = 1.021 \text{ Å}$, $\angle(NNH) = 112^{\circ}39'$. Calculations involved two basis sets (I and II), one with 11s and 7p on nitrogen and 6s on hydrogen contracted to a $(5, 3/2)$ set, the other (11) with the same contracted basis, augmented by polarization functions in the plane of the molecule, p_x and p_y , on each hydrogen atom, and by diffuse s, p_x , p_y , p_z functions on each nitrogen atom. The lowest

1. General and theoretical aspects of the diazonium and diazo groups 41

energies were, of course, obtained with basis set **I1** and are listed below for the lowest closed-shell state:

Calculations by Wagniere¹⁶⁶ on the excited states of the three molecules showed that the **3A,** state of 1,l-dihydrodiazine lies 0.090 eV in energy below that of the closedshell state ${}^{1}A_1$ (basis set II).

Hariharan and Pople¹⁶⁷ have carried out *ab initio* calculations intended to examine the influence of polarization functions on molecular orbital hydrogenation energies, and in so doing have used a **4-31 G** basis set, a **6-3 1G** basis set and a **6-31G** basis set with polarization functions added. The **4-31G** set has an s inner-shell function which is a linear combination of four s-type gaussian functions and a valence shell (s and p) represented by inner (three-gaussian) and outer (one-gaussian) parts. For hydrogen, similar **(31)** s-type functions are used. The **6-31G** basis set differs from the **4-31G** set in improvement of the inner-shell function. Then a set of six single (uncontracted) d-gaussian functions is added to the **6-31G** set for heavy atoms, the hydrogen functions being unmodified. This set is labelled **6-3 1G*.** The basis set called **6-31G**** is arrived at from the **6-31G*** set by adding three p-polarization functions on each hydrogen atom. Bond lengths of **1-25** *8,* and **0.99 A** for NN and NH, respectively, **109.47"** for the NNH bond angle, and the *rrms* conformation, were employed for diimide. Total energies as follows were calculated with the various basis sets (hartree):

These values may be compared with -109.81102 and -109.79320 hartree for *trans*and *cis*-diimide obtained earlier with the 4-31G set¹⁶⁸. For the complete hydrogenation reaction

$$
HN = NH + 2 H_2 \longrightarrow 2 NH_3
$$

they calculated the following energies (kcal/mol) :

Hence these authors suggest that the addition of d-functions $(6-31G \rightarrow 6-31G^*)$ produces an energy lowering per N atom (in N_2H_2) or 19.9 kcal/mol while the inclusion of p-functions on hydrogen $(6-31G^* \rightarrow 6-31G^{**})$ lowers the energy per hydrogen atom by 2.0 kcal/mol (for N_2H_2).

Baird and Swenson160 have used the closed-shell **SCF-LCAO-MO** method with a minimal basis of Slater-type orbitals each represented by a linear Combination of three gaussian-type orbitals **(STO-3G)** to calculate various properties of diirnide in the ground, $^{1}(n, \pi^{*})$, $^{3}(n, \pi^{*})$ and $^{3}(\pi, \pi^{*})$ states. In their work all N-H bond lengths were taken as 1.01 Å and the two H-N-N angles were assumed to be equal, while the $N-N$ distance, the $H-N-N$ bond angle, and the dihedral angle between the two N-H bonds were taken as variables. An absolute minimum is claimed for the *trans* conformation, with the *cis* isomer 5 kcal/mol higher in energy. Optimized geometries were obtained for the trans, cis, bent and twisted forms and these are

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shown in Table 26. A value of 98° was found for the dihedral angle of the twisted form at the maximum in their potential energy curves. Activation energies of 60 and 84 kcal/mol were found for the *trans* to *cis* isomerization through the planar and rotational transition states, respectively. These authors note also that the $N-N$ bond lengths in the planar excited states, $\frac{1}{n}$, π ^{*}) and $\frac{3}{n}$, π ^{*}), do not differ greatly from those in the ground state, and attribute this to the fact that the highest occupied molecular orbital (HOMO) in both *cis* and *trans* diimide is $N-N$ antibonding, so that removal of an electron from the n-MO of *cis* diimide increases the $N-N$ overlap population by 0.21 , which almost compensates for the decrease of 0.29 due to single occupancy of the π^* MO. As they also point out, ionization of the least tightly bound electron would be expected to produce a decrease in the central bond distance for diimide and azo compounds, an effect not often observed in polyatomic molecules.

It is interesting to note that Back, Willis and Ramsay¹⁵⁰, from their absorption spectral data and Franck–Condon calculations, predicted that the $\iota(n, \pi^*)$ state of diimide would have an $N-N$ bond length increased by 0.05 Å and an HNN angle 25° larger than the ground-state value, for the *trans* form. Baird and Swenson¹⁰⁹ calculated 0.05 Å and 12° for the corresponding increases, providing reasonably satisfactory agreement.

Calculations show that *trans* to *cis* isomerization through a rotational mechanism is preferred for both the $(0, \pi^*)$ and $(0, \pi^*)$ states of diimide with the activation energies being zero for both states, compared to 25 and 23 kcal/mol, respectively. for passage through the planar transition state. Rotational isomerization for the $\pi(\pi, \pi^*)$ state should be symmetry-forbidden, and the *trans* to *cis* barriers are calculated to be 71 and 65 kcal/mol for the rotational and planar transition states, respectively.

In the course of analysing the photoelectron spectra of a number of azo compounds, Brogli and coworkers¹⁷⁰ have performed some extended Hückel and MINDO/2type calculations on *cis*- and *trans*-diimide. From the latter type of calculations they have shown that the splitting for the nitrogen lone-pair orbital energies passes through a maximum at about 120° for the $N-M-H$ angle in the *trans* form but a minimum at about 100° for the cis form. This appears to be supported, in part, by their spectral data. Gimarc¹⁶¹ argued that such behaviour in cis-azo groups is due to a 'throughbond' interaction of the n_+ -orbital with a lower lying N-N σ -orbital.

Merényi and coworkers¹⁷¹ used the program REFLECT, which employs a basis set of gaussian orbitals, in this case four s-type orbitals on each hydrogen atom and seven s-type and three p-type on each nitrogen atom, to calculate barriers to rotation and inversion for diimide and its protonated form. The geometry of Lehn and Munsch¹⁵⁵ was used. Their results are summarized in Table 27. With both diimide

and protonated diimide the inversional barrier is lower than the rotational barrier for isomerization, but the corresponding barriers for the protonated form are lower than those for thc unprotonated form.

Form	Total encrev (hartree)	Barrier (keal/mol)
$H - N = N - H$		
trans	-1098175	
cis	-109.8033	$8.9(c-t)$
planar bent	-109.7348	51.9 ^a
twisted	-109.7044	71.0
$H_2N^+=NH$	$-110-1360$	
planar bent	-1100880	30·1
twisted	-110.0325	64.9

TABLE 27. Effect **of** protonation on *the* encrgics and barriers to rotation and inversion of diimide 171

^a Relative to the energy of the *trans* isomer.

An extensive study¹⁷² by means of *ab initio* SCF and CI methods has been carried out on both the ground and excited states of diirnide for the purpose of investigating the ultraviolet spectrum of this nioleciile. The NN and NH distances were held fixed at 1.238 Å and 1.065 Å, respectively¹⁵⁸, and the NNH bond angle was varied from *90"* to 170" for a variety of electronic configurations. An absolute minimum was obtained for the *tram* form with an energy of - 109.9345 hartree, first ionization potential of 10.78 eV and *cis-trans* energy difference of 12.6 kcal/mol. A barrier to isomerization of 109 kcal/mol was calculated for the ground state by employing a linear configuration for the transition state.

The most recent reports of theoretical studies on diimide appear to be that of Vasudevan and coworkers¹⁷², mentioned previously, and that of Winter and Pitzer¹⁷³. In the latter work, both NH and NN bond distances and the NNH bond angle were varied to achieve a geometry optimization to 0.01 bohr and 0.3° for the bond distances and angle, respcctivcly. Hartree-Fock calculations employed the **9SSP/4S** gaussian basis set of Huzinaga".' contracted to a **[4s3p/Zs]** basis by Dunning¹⁷⁵. A summary of geometries, energies and energy differences for all calculations of these and previous workers is shown in Table 28.

Ahlrichs and Staemmler¹⁷⁶ have published the results of *ab initio* calculations on the structure and geometry of the three isomers of N_2H_2 , including 1,1-dihydrodiazine, and using gaussian basis sets with polarization functions. Thesc authors refer to the work of Carlotti and coworkers¹⁴⁶ and Frost and coworkers¹⁷⁷ as establishing experimentally that diiniide possesses a singlei ground state in a planar *tmns* configuration. The latter authors have studied the photoelectron spectrum of **N2H,** and conclude that they are probably observing the molecule in the singlet ground statc.

The calculations of Ahlrichs and Staeinmer, the results of which are summarized in Table 28, account for electron correlation by means of a configuration interaction **(CI)** computation bascd on the pair natural orbitals (PNO) of the respective pair correlation functions (PNO CI). This trcatmcnt is stated by these authors to bc essentially cquivalcnt **10 ;i** configuration interaction calculation **in** \vhich all doubly substituted configurations have been included with the Hartree-Fock function. In

TABLE 28. Geometries and energies for cis- and trans-diimide

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addition, thc coupled electron-pair approximation was ernploycd to approximate the effect of unlinked cluster terms arising from the doubles (CEPA). The energy values given in Table 28 are those arising through the use of CEPA. Not included in the table arc their values for H,NN which are 1.221 **A,** 1.012 **A,** and **122.3"** for the NN and NH distances and NNH angle, respectively. Their best energy for the triplet (ground) state of H_2NN is -110.3341 hartree (CEPA). for these arising from the domestion of the two states of H₂NN which are 1.221 and NNH angle, respectively and NNH angle, respectively μ NN is $-110-3341$ hartree (C ind the *trans* isomer to be to I approximately 5.8

Ahlrichs and Staemmler find the *trans* isomer to be the most stable isomer with the *cis* isomer and H₂NN approximately 5.8 and 26.0 kcal/mol above the *trans* forni. In addition thcir CEPA calculations yield a value for *AE* of *59.6* kcal/mol for the reaction

$$
N_2H_2
$$
 (trans) \longrightarrow H+N₂+H

so that the isomerization apparently competes with this process. Further, their value of 55 kcal/mol for the barrier to the *trans-cis* isomerization suggests that such isomerization is unlikely to be the first step of the decomposition of N_2H_2 pproximately 5.8 and 2
 2 calculations yield a value of a value of a value
 $\frac{1}{2}$ (*trans*) —————> $H + N_2$

pparently competes with

parrier to the *trans-cis* is

e the first step of the dec
 $2 N_2H_2$ ————> $N_2H_$

$$
2 N_2 H_2 \longrightarrow N_2 H_4 + N_2
$$

Unpublishcd work from this laboratory*78, in which **a** variety of molcculcs containing the NN bond have been compared using an STO-3G basis, will be referred to in more detail later. While not yielding cnergy values of Hartrec-Fock quality, such calculations do have the advantage of being relatively inexpensively extended to larger molecules, thus permitting a comparison of a larger numhcr of molecules possessing the NN bond. With this method, an energy of **7-3** kcal/niol was calculated for the *trans* to *cis* isomerization, with the *trans* possessing the lower energy of the two forms.

C. Disubstituted Diimides

1. N_2F_2

a. *Theoretical studies*. Next to diimide itself, probably the most studied diimide, from the point of view of theoretical calculations, has been difluorodiimide.

Extended Hückel calculations were employed by Kaufman and coworkers¹⁷⁹ to examine some properties of molecules containing nitrogen and fluorine, among them being cis- and $trans\text{-}N₂F₂$. The values given by Sanborn¹⁸⁸ for the structural parameters \yere used. Calciilatcd N-F overlap populations of *0.33* and 0.32 for thc *trans* and *cis* forms, respectively, and 1.09 and 1.16 for that of the N=N bond in the *trans* and *cis* forms, respectively, indicated that the N- F distance in *trans-N*₂F₂ should be shorter than that in cis-N₂F₂, while the N=N distance in cis-N₂F₂ should be shorter than that in trans- N_2F_2 . In addition, their calculated total electronic energies suggested that the *cis* form should be more stable than the *trans*.

Ray and Narasimhan¹⁸⁰ have utilized extended Hückel theory to study the transition state in the isomerization of trans- N_2F_2 to cis- N_2F_2 . The structural data of Bohn and Bauer²⁰¹ were employed in the calculations. Configurations generated by rotating one NF bond around the NN axis were considered. Two types of *trans* configurations were examined, one (T) in which the experimentally observed bond lengths and angles of the *trans* form were used, and the other (T') in which those of the *cis* form (C) were employed. Likewise C' refers to a hypothetical *cis* form in which the *trans* bond lengths and angles are used. Linear configurations. LT and LC, were also considered, in the former the $N-N$ and $N-F$ bond lengths of *trans-* N_2F_2 were used, while in the latter those of the *cis* form were employed.

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Table 29 summarizes the calculated values of electronic energy. The calculated binding energy for *trans*-N₂F₂, 255.4 kcal/mol is in good agreement with the experimental value *07* **244.3** kcal/mol **193.** The cis *(C)* configuration is predicted to be more stable than the *rratls(T)* by approximately 15 kcal/mol, compared with **3** kcal/mol found experimentally (Armstrong and Marantz). The T' and C' configurations have

		Binding energy (kcal/mol)			
Configuration	Total energy (eV)	Calculated	Observed ¹⁹³ , 207		
т	-539.4168	255.41	244.3		
C	-540.0626	270.33	247.3		
T'	-5391941				
\mathbf{C}'	-539.6203				
LT	-531.7076				
LC	-531.8146				

TABLE 29. Total electronic energies for N₂F₂ as found by EXH met hod **IHa**

higher energies than the respective **T** and C forms. Barrier heights in the isomerization of *rrans-* to *cis-N2F,* are shown in Table **30.** According to the results of extended Hückel calculations, of the three transition-state configurations considered, that achieved by a rotation around the $N=N$ bond possesses the lowest energy.

$12a$ lion of N_2F_2 ¹⁵⁰					
Transition state	Barrier height (kcal/mol)				
rotation	47				
LT	178				
LC	176				

TABLE 30. Barrier heights for *trans* \rightarrow *cis* isomer-
ization of N_2F_2 ¹⁹⁰

Gordon and Fisher¹⁵⁴ have utilized the CNDO and INDO methods of Pople and coworkers¹⁸¹ to calculate energies and structural parameters for N_2F_2 , the latter being obtained by minimizing the energy. As can be seen from Table 31, thc *iratis* form is found to be **4.1** kcal/rnol higher in energy than that of the cis form, in good agreement with the experimentally derived value of 3.0 kcal/mol found by Armstrong and Marantz¹⁹³. If only singlet states are involved in the isomerization of *cis-* to *trans-* N_2F_2 , then activation cnergies of 68 and 85 kcal/mol for inversion and rotation on onc nitrogen, respectively, are found. If the transition form esists in a triplct statc, then it is prcdictcd that thc lowest activation energy, 52 kcal/mol, will be achieved through rotational isonierization. In **the** singlet and triplet rotational transition states, dihedral angles of 100° and 70°, respectively, were calculated.

CNDO/2 calculations have also been performed on N_2F_2 by Herndon and co\vorkers'". Geometry optimization **was** also employed and gave 1.26 **A** for the N-F distance in both the cis and *trans* forms (experimental: cis, 1.41 \AA ; *trans*, 1.40 \hat{A}^{201}); and 112^o and 109^o for the FNN angle in the *cis* and *trans* molecules respectively (experimental: *cis*, 114°; *trans*, 106°²⁰¹). In view of these results, these

		Geometrical parameters				
		Bond lengths (\AA)		Bond angles (degrees)	Relative	
State	Form	$N-F$	$N = N$	\angle FNN	Dihedral	cnergy (kcal/mol)
Singlet	linear	$1 - 25$	1.20			143.8
Triplet	linear	1.25	1.20			128.6
Triplet	cis	1.23	1.30	111	0	98.5
Triplet	bent	$1 - 25$ $(1.23)^a$	1.26	115		94.0
Singlet	trans	$1 - 24$	1.24	132	100	85.0
Singlet	bent	1.24 $(1.23)^a$	1.22	114	--	$67 - 7$
Triplet	trans	1.24	1.27	119	180	66.7
Triplet	trans	1.24	1.30	113	70	520
Singlet	trans	1.25	1.25	109	180	$4-1$
Singlet	cis	1.25	$1 - 24$	116	0	0.0

TABLE 31. Total relative electronic energies and energy-optimized structural parameters for N₂F₂ as found with the CNDO and INDO methods¹⁵⁴

^a Bond lengths in brackets refer to the $N-F$ lengths at the $N-F$ which is bending.

workers varied the value of the β constant for fluorine in the CNDO method to bring the experimental and theoretical $N-F$ distances for the *trans* into coincidence. Table 32 summarizes their results, which do not differ greatly from those of Gordon and Fisher¹⁵⁴ except in predicting an activation energy of 81.1 kcal/mol for the inversion mechanism.

Gimarc¹⁶¹ applied the extended Hückel method to N_2F_2 and discussed the results primarily from the point of view of the molecular orbitals involved and the shapes of the molecule. For molecules like N_2F_2 which possess 24 valence electrons, the highest occupied orbitals for linear geometry would be the doubly degenerate upper

		Geometrical parameters					
	Bond lengths (A)			Bond angles (degrees)		Relative	
Form	$N-F$ $N = N$		Dihedral \angle FNN		Symmetry	energy (kcal/mol)	
trans	1.40	1.19	144	180	C_{2h}	155.2 (max)	
linear	1-45	1.15	180		$D_{\infty h}$	$106 - 8$	
trans	1.40	1.23	110	95	C_{2}	85.70 (max)	
trans	1.393	1.19	112, 185	180	$C_{\rm s}$	$81 \cdot 10$ (max)	
trans	1.40	1.23	110	90	C_{2}	79.65	
bent	1.393 (1.393)	1.19	112	---	C_{s}	75.40	
1,1-Difluoro	1.40	1.19	133		C_{2v}	39.0 (min)	
trans	1.395	1.22	105	180	$C_{\rm 2h}$	3.14 (min)	
cis	$1 - 40$	1.21	114	0	C_{2v}	0.00 (min)	

TABLE 32. CNDO/2 results for N_2F_2 ¹⁸² using adjusted β_F value of 14.07

 π_{μ} orbitals, $\pi_{g}(x)$ and $\pi_{g}(y)$. These are completely antibonding, being composed of p atomic orbitals antiparallel to each other, and each would contain **a** single electroil only. The degeneracy of these $\pi_g(x)$ and $\pi_g(y)$ orbitals is removed when the molecule bends, the two levels becoming a_2 and b_3 for cis, and b''_6 and a'_6 for trans bending, and these four orbitals can now contain p_z and s atomic orbitals as well as p_z or p_w . When cis bending occurs, the energy of the a₂ level changes very little from that of the $\pi_{\mathfrak{g}}(y)$, but the energy of the b₂ level decreases substantially below that of the linear $\pi_{\mathfrak{g}}(x)$, because the lower lobes of the F atom p_x atomic orbitals move from antibonding positions into favourable bonding positions with respect to the neighbouring N atom p_x orbitals. On the other hand, when *trans* bending occurs, b_{μ}^{\prime} remains similar in energy to $\pi_{g}(y)$, but the a_{g}^{\prime} level is lower in energy than $\pi_{g}(x)$, since the p_x orbitals on the F atoms move into bonding positions with respect to their neighbouring N atoms. As Gimarc points out, it is then the b₂(cis) and a_g(trans) encrgy levels which give *cis* and *trans* shapes to N_2F_2 .

In discussing the isomerization of N_2F_2 , Gimarc notes that the rotational mechanism involves the crossing of highest filled and lowest unoccupied a and b orbitals. The principle of conservation of orbital symmetry consequently forbids the rotational mechanism. In the inversional mechanism the highest-filled $a_{\rm g}$ orbital can convert into **b**₂ because a,b symmetry is lost.

Turner¹⁶³ has used the CNDO theory to calculate the total electronic energies of the *cis* and *trans* forms of N_2F_2 as a function of bond angle between 90° and 125°. Minimum energies of -78.6953 and -78.6958 hartree were found at 110[°] and 105" for the *cis* and *trotis* forms, respectively. Within the limits of error of the calculation the energies are the same. The calculated rotational barrier (*trans* \rightarrow *cis*) is S5.l kcal/mol. Turner suggests, since no significant change occurs in the population analysis in passing from the *frnns* to the cis form, that the origin **of** the barrier is probably not due to any alteration in chemical bonding but is simply an atom-atom $repulsion.$

The method and results of *nb itzitiu* minimal basis SCF-CI calculations performed by Ditchfield and coworkers¹⁶³ on diimide have been discussed in a previous section. The same paper reported -11.4 and 5.67 eV for the energies of the *n* and π^* orbitals of cis-N₂F₂, and -10.05 and 5.87 eV for the corresponding orbitals of *trans-N₂F₂*, yielding vertical singlet $n \rightarrow \pi^*$ transition energies of 5.75 and 4.22 eV for the *cis* and *trans* forms, respectively.

Brundle and coworkers¹⁸⁴ have employed photoelectron spectroscopy and ab *initio* gaussian orbital calculations to study the effect of the substitution of fluorine for hydrogen in **a** planar molecule, which has been shown, at least in some cases, to have a much larger stabilizing effect on the σ MO's than on the π MO's. There is some difficulty in studying diimide and difluorodiimide for this purpose, as these authors point out, since there are no photoelectron data for *trans*-diimide and the photoelectron spectrum of trans- N_2F_2 may contain bands due to an isomeric impurity. However, Table 33 summarizes their calculated and experimental ionization potentials for these two molecules. The highest MO in N_2F_2 , 7a_g, contains some $N-F$ c-antibonding character in addition to the $N-N$ antibonding lone-pair component. The agreement between the calculated energy of 13.92 eV and the experimental, 13.4 cV, is quite good. What the authors refcr to as the perfluoro shift, or simply the change in orbital energies on substitution of fluorine for hydrogen, is $+3.6$ eV for the $4a_g-7a_g$ orbital pair. The $1a_u \pi$ MO of N_2H_2 is calculated to possess an energy of 13.07 eV, which correlates with the $2a_u$ level of $N₂F₂$ at 14.24 eV (calc.). The $1a_u-2a_u$ shift then is approximately $+1.1$ eV.

Companion and Hsia¹⁸⁵ have employed a bicentrically rescaled, energypartitioned, version of the CNDO/2 method in a study of atomization of a variety of nitrogen fluoride molecules, including N_2F_2 . The geometrical data of Bohn and Bauer²⁰¹ were employed. Calculated values for the heats of atomization were 217.1 and 216.0 kcal/mol for the *trans* and *cis* forms respectively, which may be compared with the experimental values¹⁸⁶ of 251.6 and 254.9 kcal/mol for the *trans* and *cis* forms, respectively. The electron affinity and first ionization energy were calculated as 2.00 and 14.62 cV for N_2F_2 .

N_2H_2		N_2F_2					
Orbital	$0.92 K.T.^a$	Orbital	$0.92 K.T.^a$	Adiabatic $_{\rm IP}$	Vertical IP		
$4a_g$	9.84	7a _g	13.92	12.8	13.4		
1a _u	13.07	$2a_n$	14.24	13.65	$14-1$		
3b _u	15.97	$6b_u$	16.60				
$3a_{\rm g}$	17.42	$6a_{\alpha}$	17.50				
$2b_u$	23.93	$1b_{\rm r}$	18.57				
$2a_{\rm g}$	34.80	la _u	19.80	$18-0$	18.70		
$1b_u$	$391 - 4$	$5b_{11}$	20.20		19.80		
1a _g	391.5	$5a_{\alpha}$	20.39		21.0		
		4b _u	24.06	22.3	$22 - 7$		
		4a _v	35.8				
		3b _u	$42 - 4$				
		$3a_{\alpha}$	43.5				
		$2b_u$	396				
		$2a_{\rm g}$	396				
		1b _u	661				
		la _g	661				

TABLE 33. Observed and calculated ionization potentials (eV) for N_2H_2 and trans- N_2F_2 ¹⁸⁴. Reprinted, with permission, from Brundle and coworkers, J. Amer. Chem. Soc., 94, 1451 (1972). Copyright by the American Chemical Society

^a 0.92 K.T. refers to 92% of the value as given by Koopmans' theorem.

b. Spectra and structure. The electron diffraction pattern of a sample of N_eF_e was determined in 1947 by Bauer¹⁸⁷, who assumed a 50-50% mixture of cis and trans isomers, and fitted the data with $N-F$ and $N=N$ bond lengths of 1.44 and 1.25 Å, respectively, and FNN bond angle 115°. Sanborn¹⁸⁸ prepared large amounts of N_zF_z by maintaining an electric discharge in a stream of nitrogen trifluoride in the presence of mercury vapour. Two fractions were obtained by gas chromatographic separation on a silica-gel column held at a low temperature, each with a molecular weight of 66, but with quite different chemical reactivity. One fraction reacted with mercury and explosively with diborane, while the other did not, and the infrared spectra of the two fractions were different. Sanborn, from his infrared data, decided that the inactive form was *trans*-1,2-difluorodiazine $(\tau$ -1,2-DFD) and the active form was suggested to be $1,1$ -difluorodiazine (1,1-DFD). For the fraction suggested to be τ -1,2-DFD, three fundamental vibrations (cm⁻¹) of symmetry A_g (1010; $N-F$ symmetric stretch; 1636, $N=N$ stretch; 592, $N=N-F$ symmetric bend), one of A_u (360, torsion) and two of B_u (989, N-F asymmetric stretch; 421, N=N-F asymmetric bend) were identified. On the other hand, 1,1-DFD gave three A₁ fundamentals (896, N-F symmetric stretch; 1524, N=N stretch; 552, F-N-F symmetric bend), one B_2 (737, out-of-plane bend) and two B_1 (952, N-F asymmetric stretch; 573, $F-N-F$ symmetric bend). Sanborn provided chemical,

mass spectral, dipole nionient and infrared spcctral evidence for thc identification of his 'active' fraction of N₂F₂ as 1,1-DFD and not *cis*-1,2-difluorodiazine.

Ettinger and coworkers¹⁸⁹ agreed with Sanborn that the less active (lower boiling, $-111.4 \degree C$) isomer was the *trans* configuration, but argued that the structure of the -111.4 °C) isomer was the *trans* configuration, but argued that the structure of the more active (higher boiling, -105.7 °C) isomer was much more in doubt, for a variety of reasons. They provided alternative interpretations for Sanborn's experimental data and, in addition, noted that n.m.r. spcctra of both the high- and low-boiling isomers showed only one nitrogen resonance, while two nitrogen resonances might be expected for 1,1-DFD. They noted also that the dipole moment change for the $N=N$ stretching mode is not along the axis of the least moment of inertia, but is along an axis of intermediate moment. Consequently, a *cis* isomer as well as **a ^I**,I-isomer could show only PR structure. Sanborn190 while conccding that the shapes of the infrared bands did not preclude a *cis* isomer, insisted that the infrared band at 737 cm⁻¹ in 'active' N_2F_2 must be a fundamental vibration, and can only be a bending fundamental, which would be, he argued, out of position for a *cis* isomer, but fitting perfectly for 1.1-DFD.

Herron and Dibeler¹⁹¹, in their mass spectrometric study of N_2F_2 , measured the appearance potentials of N₂F₂⁺, N₂F⁺ and NF⁺ from the *trans* isomer as 13.1, 13.9 and 17.0 eV, respectively, while those from what they considered to be the cis isomer were, for N_2F^+ and NF⁺, 14.0 and 16.9 eV, respectively. The *cis* isomer apparently does not produce a parent ion. Hence the appearance potentiais of the normal fragment ions N_2F^+ and NF^+ are identical within experimental uncertainty $(\pm 0.2 \text{ eV})$ for both *cis-* and *trans-N₂F₂*. Herron and Dibeler reasoned that this information suggested a similarity in molecular structure and hence favoured the identification of the 'active' isomer as cis-l,2-DFD rather than 1,l-DFD. Heats of formation for the *cis* and *trans* isomers were reported as 16.4 and 19.4 kcal/mol, respectively with associated bond dissociation energies $D(FN=NF)$ as 106 and 103 kcal/mol, respectively.

Measurements of the ¹⁰F chemical shifts and splittings for the two isomers of N_2F_2 were reported by Noggle and coworkers¹⁹². An analysis of the ¹⁹F- ${^{14}}N_1^{13}$ double-resonance spectra for the reactive isomer indicated that both nitrogens in the molecule have the same chemical shift, thus providing strong evidence in favour of the *cis* structure for the 'active' isomer. Calculated and observed spectra agreed quite well when the *cis* and *trans* structures were assumed.

The heats of formation of the two isomeric forms of N_2F_2 were measured calorimetrically by Armstrong and Marantz¹⁹³, by reacting the N_2F_2 with ammonia.

$$
N_2F_2(g) + 8/3 NH_3(g) = 2 NH_4F(C) + 4/3 N_2(g)
$$

Mean heats of reaction of 211.7 and 209.2 kcal/mol were measured for the trans and 'active' isomers, respectively. The difference in the heats of reaction of the two samples, adjusted for the purity of the samples, yields a heat of isomerization (active \neq trans), $\Delta H_{f298}^0 = 3.0$ kcal/mol, indicating that the 'active' form is more (active \rightleftarrows *trans*), $\Delta H_{2.98}^0 = 3.0$ kcal/mol, indicating that the 'active' form is more stable thermodynamically than the *trans* form. Use of ΔH_{2208}^0 [NH₃(g)] = -11.04 kcal/ stable thermodynamically than the *trans* form. Use of $\Delta H_{2.08}^{9}[\text{NH}_3(\text{g})] = -11.04 \text{ kcal/mol}^{194}$, and $\Delta H_{2.08}^{9}[\text{NH}_4\text{F}(\text{c})] = -111.0 \text{ kcal/mol}^{195}$, produces values of 19.4 and 16.4 kcal/mol for $\Delta H_{2.298}^0$ of the *trans* and 'active' isomers of N₂F₂, respectively. Armistrong and Marantz calculate the total binding energy of trans- N_2F_2 to be 244.3 kcal/mol, which, with the use of 107.9 kcal/mol for E(N=N), yields **a** value of 68.2 kcal/mol for $E(N-F)$.

The microwave spectrum of the 'active' isomer of N_2F_2 was measured by Kuczkowski and Wilson¹⁹⁶, who concluded that the structure was planar and *cis* with NN and NF bond lengths of 1.214 and 1.384 **A,** respectively, and FNN angle

 $0.114.5$ ². A dipole moment of 0.16 D was obtained from the rotational Stark effect. Thus, by 1963, the evidencc in favour of the *cis* structurc for the 'active' isomer seemcd strongly to outweigh any other previously hcld conclusion.

The gas-phase infrared and liquid-phase Raman spectra of $trans\text{-}N_{2}F_{2}$ has been measured by King and Overend¹⁹⁷, with the following vibrational assignments: $v_{NN} = 1522$, $s_{VNF} = 1010$, $s_{N} = 600$, $a_{VNF} = 990$, $a_{N} = 423$ and torsion = 363.5 cm⁻¹. These authors note that their assignment of the infrared spectrum is essentially the same as that of Sanborn¹⁸⁸. In a subsequent paper¹⁹⁸ these same authors restudied some of the bands at higher resolution to obtain estimates of the rotational constants, which values are consistent with the values of Bauer¹⁸⁷ from electron diffraction measurements.

The Raman spectrum of gaseous trans-1,2-difluorodiazine has been reported by Shamir and Hyman¹⁹⁹. The three symmetric vibrations were obtained and assigned **as 101s** cm-', N-F symmetric stretch; **1523** cm-l, N=N stretch, and 603 cm-', $N=N-F$ symmetric bend, which agree quite well with those found by King and Overend¹⁹⁷ for the liquid, where in all cases the liquid-phase values are lower than those in the gas phase, as might be cxpected.

The Raman spectrum of cis -N₂F₂ was also measured by King and Overend²⁰⁰.

	$cis-N2F2$ (gas) ²⁰⁰			$trans-N.F2$					
						Frequency (cm^{-1})			
		Frequency (cm^{-1})	Description			Gas ¹⁹⁹	Liquid ¹⁹⁸	Calc ¹⁸³	Description
А,	ν_1 v_{2} v_3	1525 896 341	$\nu_{\rm NN}$ $\nu_{\rm NF}$ $\delta_{\rm NF}$	$A_{\rm g}$	ν_{1} ν., $\nu_{\mathcal{R}}$	1523 1018 603	1522 1010 600	1636 1010 592	$\nu_{\rm NN}$ $\nu_{\rm NP}$ $\delta_{\rm NF}$
A ₂ B ₁	$\nu_{\rm A}$ ν_5 ν_6	(550) 952 737	Torsion $\nu_{\rm N,F}$ $\delta_{\rm NF}$	$A_{\rm II}$ $B_{\rm u}$	ν_{1} v_{5} $\nu_{\mathbf{a}}$	362 990 420	363.5 990 423	360 989 421	Torsion $\nu_{\rm NF}$ $\delta_{\rm NF}$

TABLE 34. Vibrational assignment of *cis-* **and** *trans-N₂F₂*

Bohn and Bauer201 have determined the structures of *cis-* and irons-I ,2-difluorodiazine from electron diffraction measurements, using sectored-microdensitometer data. The gcometric configurations and interatomic distances werc found to be essentially the same **as** those previously estimated on the basis of the visual tecliniquc. Table 35 records the structural parameters from the work of Bohn and Bauer²⁰¹ and Kuczkowski and Wilson¹⁹⁶. The N=N and N-F distances as obtained by Bohn and Bauer²⁰¹ are similar to those expected for other molecules with $N=N$ and $N-F$ bonds. However, in the *traits* isomer, $\angle NNF = 105.5^{\circ}$ is surprisingly small. With the cis compound there is agreement between the two sets of data shown in Table 35 for the N=N bond distance and $\angle NNF$ but not for the N-F bond distance. But the major structural difference between the *cis* and *trails* isomers is the larger \angle NNF in the former compound. This can be rationalized on the basis of a repulsion between the two F atoms in the *cis* isomer which are about 1 angström closer together than in the *trans* form. The $N=N$ bond length is also somewhat shorter in the *cis* form, while the N-F distance is slightly longer in the *cis* form.

Calculations of vibrational coupling effects and potential functions of both the *cis* and *trans* isomers have been made by Pearce and Levinzo2 who cniployed the **3**

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structural parameters of Bohn and Bauer²⁰¹, and compared their calculated results with the experimental data^{198, 199, 200}. Calculated and observed frequencies agreed quite well in all cases.

	$trans-N2F2$	cis -N ₃ F_2			
	Bohn and Bauer ²⁰¹		Bohn and Bauer ²⁰¹ Kuczkowski and Wilson ¹⁹⁶		
N=N(Å) N-F(Å) N · · · F(Å) F · · · F(Å)	1.230	1.214	1.214		
	1.396	1.410	1.384		
	2.093	2.209			
	3.339	2.381			
\angle NNF (deg)	105.5	114.4	114.5		

TABLE 35. Structural parameters for cis- and trans- N_2F_2

c. Thermochemical properties. Colburn and coworkers²⁰³ were the first to report on the properties of N_pF_p . Their values for a variety of physical properties are given in Table 36. For comparison purposes the table is augmented by the data given in the review paper of Pankratov²⁰⁴. Colburn and coworkers²⁰³ considered the cis isomer to be chemically more reactive, yet Lustig²⁰⁵ found that the isomerization of the *trans* to the *cis* isomer took place below 100° C nearly quantitatively in a static system. Lustig²⁰⁵ points out that this isomerization is consistent with the heats of formation of Armstrong and Marantz¹⁹³ (see Table 36), which indicates that the cis isomer is thermodynamically more stable.

	trans-Difluorodiimide	cis-Difluorodiimide
Vapour pressure	$\log P_{\text{torr}} = (-742.3/T) + 7.470$	$\log P_{\text{torr}} = (-803.0/T) + 7.675$
Boiling point $(^{\circ}C)$	$-111-4$	$-105-7$
Heat of vaporization (cal/mol)	3400	3670
Critical temperature $(^{\circ}C)$	-13	-1
Critical pressure, estimated (atm)	55	70
Melting point $(^{\circ}C)$	-172	below -195
Heat of isomerization (kcal/mol)		$27.5 + 5.0$ (est) ²⁰³
Heat of formation ΔH_t^0 (kcal/mol)	19.4193	16.4^{193} 25.3204 20.5 (-105 °C) ²⁰⁴

TABLE 36. Physical properties of cis and trans difluorodiimide²⁰³

Pankratov²⁰¹ has determined the heat of formation of the 'active' *cis* isomer of N_2F_2 from reaction with acid potassium iodide solution. He obtained 25.3 ± 2.0 kcal/mol at 25° C and 20.5 ± 2.0 kcal/mol at -105.7° C (for the liquid isomer). Pankratov calculates a dissociation energy of 97.5 kcal/mol for the $N=N$ bond in the cis isomer by using 61.14 kcal/mol for the heat of formation of NF and his value for N_2F_2 .

d. *Kinetics of isomerization*. Binenboym and coworkers²⁰⁶ have carried out experiments designed to provide information on the *cis-tram* isomerization of difiuorodiiniide under homogeneous conditions. **As** has been mentioned, the isomerization is catalysed by heated metal reaction vessels, and difluorodiimide also attacks glass at elevated temperatures. In the study by Binenboym and coworkers, **N2F2,** highly diluted in argon was heated in a single-pulse shock tube. Since the tube walls remain cold, the reaction can be studied under homogeneous conditions. The isomerization was studied between *297* and 342 "C and at pressures up to 1.6 atm. Vapoti; phase chromatography was employed to calculate *cis* and *trans* concentrations. The isornerization was identified as first-order with rate constants given by

$$
k = 10^{14} \exp(-32{,}200/RT) \sec^{-1}
$$

These authors point out that the pre-exponential factor is high and similar to that found with the carbon-carbon double-bond system²⁰⁷. However, their calculations of the energy required to decouple the π -bond overlap yielded 84 kcal/mol, much higher than their experimental value of 32 kcaljmol. Hence they conclude that the isomerization cannot take place by rotation and a decoupling of the π -bond overlap so that the transition state presumably retains its double-bond character. They tentatively suggest that a linear configuration in the transition state may be consistent with their results.

Apparently no further **work** on the *fratis-cis* isomerization was done until, in 1974, Vedeneev and collaborators²⁰⁸ reported on experiments done in a Teflon reactor from 100 to 150 °C and at initial *trans*-N₂F₂ pressures of 15, 25 and 32 torr. They report that an autoaccelerated reaction begins when the first trace of cis- N_2F_2 is formed. Values of $\Delta H = -2070 \pm 145$ cal/mol and $\Delta S = -1.55 \pm 0.37$ e.u. were obtained.

2. Tetrazene

Wiberg and coworkers have recently reported on the preparation³⁰⁹ of tetrazene, $N₄H₄$, and its photoclectron spectrum²¹⁰ resulting from their earlier work on azo compounds.²¹¹ Discussion of the work contained in the latter reference would lead us too far outside the domain of this volume.

These authors believe that N_4H_4 is $H_2N-N=N-NH_2$, probably, at least with their methods of preparation, in the *trnns* form. Thcrmal decomposition produces $N₂$ and hydrazine by one route or alternative rearrangement to ammonium azide NH_4^+ N₃, the former in about 75% yield and the latter 25%. Gaseous tetrazene is unstable at room temperature, whereas decomposition of solid tetrazene begins at approximately $0^{\circ}C$. Their assignments of the bands of the photoclectron spectrum, plus CNDO/S calculations lead to the following orbitals and energies: $\pi_3(8.51 \text{ eV})$, $n_{+}(10.13 \text{ eV}), \pi_{2}(11.67 \text{ eV}), n_{-}(13.92 \text{ eV})$ and $\pi_{1}(14.44 \text{ eV}).$

D. Monosubstituted Diimides

1. Introduction

Relatively little work, of either theoretical or experimental nature, had been reported on the physicocheniical properties of the interesting molecules, $R-N=N-H$, until quite recently. A review²¹² has been published in 1971 by Kosower. He states that, up to 1965, there were no reports of the direct observation of *a* monosubstituted diiniidc, although there were many descriptions of reactions in which $R-N=N-H$ had been invoked as an intermediate. Monosubstituted diimides (i) are oxygen sensitive, (ii) disappear via bimolecular reaction and (iii) react (often) with base. It is claimed that methyldiimide is the only monosubstituted diimide that can be readily isolated in pure form.

2. Spectra and structure

Absorption data are reported by Kosower for the monosubstituted diimides. The $n \rightarrow \pi^*$ bands fall between 3490 and 3750 Å for unconjugated systems and between 4030 and 4175 *8,* for conjugated compounds. Kosower notes that the absorption coefficients are all very low, between 18 and 100, thus implying a trans structure for all diimides listed. The $n \rightarrow \pi^*$ transition for trans CH₃N=NCH₃ appears at 3430 Å²¹², while that for $CH_3N=NH$ is given as 3610 Å, thus suggesting that alkyl substitution for hydrogen may decrease the n^+-n^- splitting.

Steinmetz²¹³ has measured the microwave spectrum of $CH₃NND$, the deuterated species being used because **of** its higher stability than rhe hydrogen species. The deuterated species can be stored at liquid nitrogen temperature without decomposition, whereas the hydrogen species will still decompose at this temperature. To obtain a structure thc HCN angle and CH bond length were assumed to be 109.5" and 1.09 A, respectively and values of 1.24, 1.47 and 1.014 *8,* wcrc taken for the NN, CN and NH bond lengths, after consideration of other similar compounds. This pcrniitted angles CNN and NNH to be calculated as 112 and 1 lo", rcspcctively. **A** methyl group rotational barrier of 1665 ± 40 cal/mol was also calculated.

Ackerman and coworkers²¹⁴⁻²¹⁵ have measured the infrared spectra (4000- 200 cm^{-1}) of methyldiimide and its deuterated forms, in the gas phase, condensed phase (77 **K),** and in nitrogen matrices (20 K), and have made thc following assignments (deuterated species in brackets): $N-H$ stretch at 3130 (2320), $N=N$ stretch at 1575 (1550) and the N-H out-of-plane bend at 844 cm⁻¹ (662). The N-H stretch agrees with that found for *trans* diimide¹⁴³. It is interesting to note, however, that the $N-H$ out-of-plane bends in diimide itself have been assigned at 1359 cm⁻¹ (N_2H_2) and 999 cm⁻¹ $(N_2D_2)^{111}$, much higher than those for the monosubstituted molecules.

Tentative assumptions²¹⁴ of geometrical parameters for methyldiimide were:

By comparing their results with those for trans-diazene, Ackerman and coworkers214-215 concludcd that tlie *frotis* form can be cxpccted to be the most stable structure of methyldiimide.

3. Theoretical studies

Very few calculations of any type have been reported in the literature for monosubstituted diimides. This is somewhat surprising since some relatively simple monosubstituted diimides, whosc existcncc has not yet been verified experimentally, could be studicd with advantage from a thcorctical point of view. Of course the experimentally known monosubstituted diimides may still be too large for inexpensive theoretical **work.**

Ditchfield and coworkers 163 , in their work discussed in an earlier section, have calculated the *n* and π^* orbital energies and singlet $n \rightarrow \pi^*$ transition energies for a number of monosubstituted diirnides. Their results are summarized in Table **37.** Using thcir standard geometry, Ditchficld and coworkcrs found that the *trans* isomer should be more stable than the cis in the ground state, with the energy difference between the two forms decreasing in the order $CH_3 > H > NH_2 > OH > F$.

			$\Delta E(n \rightarrow \pi^{*})$		
R	ε_n	ε_{-}	Calculated	Observed (λ_{\max})	
CH ₃ (cis)	-9.30	6.68	4.63		
$NI2$ (cis)	-9.92	6.83	5.72		
HO (cis)	-10.05	6.42	5.54		
F (cis)	-11.14	5.67	5.75		
CH ₃ (trans)	-8.64	6.88	3.58	$3.45 - 3.55$ ²¹²	
$NH2$ (trans)	-9.98	6.32	4.37		
F (<i>trans</i>)	-10.05	5.87	4.22		
HO (<i>trans</i>)	-9.98	6.32	4.37		

TABLE 37. *n*, π^* Orbital energies and singlet $n \rightarrow \pi^*$ transition encrgies (eV) for $RN=NH$ molecules¹⁶³

In both the *cis* and *trans* monosubstituted diimides, the $n \rightarrow \pi^*$ transition (the lowest of two is reported) is shifted to higher energy in order $CH_3 < HO < NH_2 < F$ for the *cis*, and $CH_3 < F < HO < NH_2$ for the *trans*, although the differences for the highest three encrgies in the *trans* series is small. As Ditchfield and coworkers point out the changes of the *n* and π^* orbital energies resulting from σ withdrawal are more important than the π -donating effects.

Again, **as** with diimide itself, the cxistence on thc azo group of two pairs of nonbonding electrons gives rise to two possible $n \rightarrow \pi^*$ transitions. However, only one low-cnergy u.v. band is usually observed. If the molecules have a *trans* structure, a strong interaction **will** exist betwecn thc lone pairs, which gives risc to a large splitting in the energies of the two *n* orbitals, and consequently two $n \rightarrow \pi^*$ transitions of very different energies. Ditchfield and collaborators find the splitting between the two *ti* orbitals to be approxiinatcly 5-6 and **2.5-3.5** eV in thc *[ratis* and *cis* isomers, respectively, thus supporting this explanation. The two $n \rightarrow \pi^*$ transitions are separated by approxiniatcly 6 and 2 **CV** in the *trans* and cis compounds. Consequently, it would be expected that the second $n \rightarrow \pi^*$ transition in the *trans* isomers will be in a much dimerent region of the **U.V.** spcctruni.

Mcrényi and coworkers¹⁷¹, in their work described in a previous section, have performed *ab initio* calculations on $CH_3N=NH$ and its protonated form for the purpose of obtaining barriers of rotation and of inversion. Their results are reproduced **in** Table **38.** In addition, a proton aflinity of 212.5 kcal/mol and *n* z-orbital energy of -0.468 hartree were calculated for the *trans* form. The high values for the barrier of inversion with $CH_3N=NH$ are attributed to the orbital associated with the lonc pair of electrons.

In this laboratory²¹⁶, we have employed the Gaussian 70 programme²⁹ and a minimal STO-3G basis set to calculate the total electronic encrgies of a number of niono- and di-substituted Jiiniidcs in thcir cis and *frmiy* configurations. In all cases the nuclear configurations were optimized to ± 0.001 Å for bond lengths and ± 0.1 for bond angles. Tablc **39** summarizes **[lie** optimized values for the various bond

Molecule	Configuration	Total energy (hartree)	Barrier (kcal/mol)
$CH3N = NH$	trans	-148.8032	7.8
$CH3N = NH$	cis	-148.7907	
$CH_3N = NH$	Invertomer (H inverts)	-148.7193	52.6 (from <i>trans</i>)
$CH_2N = NH$	Invertomer $(CH3$ inverts)	-148.7297	46.1 (from <i>trans</i>)
$CH3N=NH2+$		-149.1419	
$CH3N=NH2+$	Invertomer	-149.1030	$24 - 4$
$CH3N = NH2$	Rotamer	-149.0371	65.8

TABLE 38. Total electronic energies of CH₃N=NH and CH₃N=NH₂⁺ in various configurations¹⁷¹

TABLE 39. Calculated structural parameters^a of mono- and di-substituted diimides

		Bond lengths		Bond angles			
Molecule	NN	NX	XH	NNX		NCH HCH	
HNNH (trans)	1.267	1.061		105.3			
$HNNH$ (cis)	1.263	1.064		$111 - 4$			
HNNH (invert)	1.246	1.082		$108 - 0$			
		1.006 ^b		(182.0)			
$H_1C-N=N-CH_3(t)$ 1.268		1.514	1.089	109.6	109.8	109.3	
$H_3C-N=N-CH_3(c)$ 1.264		1.513	1.088	119.2	109.8	109.0	
$H_2C - N = N - CH_3$ (i) 1.247		1.549	1.088	$111 - 4$	1100	109.0	
		1.432^{b}	1.096	(182.0)	$110-2$	109.1	
$CH3-N=N-H(t)$	1.268	1.060(NH)		104.8 (NNH)			
		1.515(NC)	1.089 (CH)	109.9(NNC)		109.6 109.6	
$CHr - N = N - H(c)$	1.263	1.066(NH)		111.0(NNH)			
		1.517(NC)	1.089 (CH)	115.6 (NNC)	109.5	109.5	
$CH3 - N = N - H$ (i)	1.250	1.082 (NH)	1.096 (CH)	107.6 (NNH)	109.5	109.5	
on $CH3$		$1.436(NC)^{5}$		(183.0) (NNC)			
$F-N=N-F(t)$	1.282	1.371(NF)		$106 \cdot 1$ (NNF)			
$F-N=N-F(c)$	1.277	1.373(NF)		111.5 (NNF)			
$F - N = N - F$ (i)	1.250	1.398(NF)		109 2 (NNF)			
		$1.351(NF)^{b}$		(182.0) (NNF)			
$NC-N=N-H(t)$	1.277	1.062 (NH)		104.2 (NNH)			
		1.446 (NC)		109.5 (NNC)			
		1.159 (C=N)		109.5 (NNC)			
$NC-N=N-H(c)$	1.273	1.064 (NH)		111.9(NNH)			
		1.447(NC)		115.3 (NNC)			
		1.159 (C≡N)					
$F-N=N-H(t)$	1.270	1.056 (NH)		104.0(NNH)			
		1.370(NF)		107.4 (NNF)			
$F-M=N-H(c)$	1.264	1.065 (NH)		108.7(NNH)			
		1.372 (NF)		110.6 (NNF)			
$NC-N=N-CN(t)$	1.281	$1.441(N - C)$		109.1(NNC)			
		1.160 (C=N)					
$NC-N=N-CN(c)$	1.281	$1.440(N - C)$		117.4 (NNC)			
		1.160 (C== N)					

 α Bond lengths in \hat{A} and bond angles in degrees.
 α Bond length of inverting NX.

lengths and angles. **As** can be seen froni the table, in the cases of diimide and dimethyldiimide calculations were also performed on the invertomers. By bending one of the NX bonds where X is either H or $CH₃$, the bend angle corresponding to the top of the encrgy barrier was located. Optimization of the nuclcar geometry was then done with this bend angle (normally approximately 180") held fixed.

The total clectronic energies (including nuclear repulsion) and the transition energies for the *cis* to *trans* isomerizations are given in Table 40. It can be seen that these calculations predict that the *trans* (singlet) form will be the more stable of the two, *cis* and *trans*, for all the molecules examined except N_2F_2 . In the case of N_2F_2 , although other data are not entirely conclusive, nevertheless the evidence available points to the *cis* isomer as the more stable form (see Section III.C.l.a–c). However the calculated isomerization energy given in Table 40 is suspiciously small. We used the optimized N_2F_2 geometry from Table 39 and calculated, using an extended basis (6-31G), for the *cis* and *trans* forms of N_2F_2 , energies of -307.441691 and - 307.44078 1 hartree, respectively, corresponding to an isomerization energy of about 0.6 kcal/mol. This is still much smaller than the 3.0 kcal/mol value of Armstrong and Marantz¹⁹³ or the 2.1 kcal/mol obtained by Vedeneev and coworkers²⁰⁸. The calculated inversion energy of 62.1 kcal/mol is also much higher than the experimental value of 32.2 kcal/mol²⁰⁶.

Molecule	Energy (hartree)	Transition energies (kcal/mol)		
HNNH (trans)	-108.55696	(trans to cis) $+7.3$		
$HNNH$ (cis)	-108.54525			
HNNH (invert)	-108.45100	$(66.5)_{\ell-1}$		
CH ₃ NNCH ₃ (t)	-185.72658	$+6.8$		
CH ₃ NNCH ₃ (c)	-185.71578			
$CH3NNCH3$ (i)	-185.62762	$(62.1)_{t-1}$		
$CH2-N=N-H(t)$	-147.14214	$+5.2$		
$CH2-N=N-H(c)$	-147.13388			
$CH3-N=N-H$ (invert on $CH3$)	-147.04272	$(62.4)_{\ell=1}$		
$F-N=N-F(t)$	-303.44088	-0.1		
$F-N=N-F(c)$	-303.44102			
$F-N=N-F(i)$	-303.30685	$(S4.2)_{c-1}$		
$NC-N=N-H(t)$	-199.09309	$+6.1$		
$NC-N=N-H(c)$	-199.08341			
$F-N=N-H(t)$	-206.00450	$+1.1$		
$F - N = N - H(c)$	-206.00281			
$NC-N=N-CH(t)$	-289.62259	$+6.6$		
$NC-N=N-CH(c)$	-289.61203			

TABLE 40. Calculated electronic energies for mono- and di-substituted diimides

The results of our calculations for mcthyldiimidc predict that the *frrrns* isomer is more stable than the *cis* by approximately 5.2 kcal/mol which agrees reasonably well with the conclusions of Merényi and coworkers although their total electronic energies reflect the higher quality of their \\zvefunctions. However, their **CH3** inversion energy of 46.1 kcal/mol is much lower than our 62.4 kcal/mol.

The results of wry similar calculations **have** just bcen publishcd by HOWCII and Kirschenbaum²¹⁷ and, as expected, our results agree quite well with the results of these authors.

58 **J. B.** Moffat

IV. DlAZONlUM COMPOUNDS

A. Introduction

A number of books concerned, at least partially, with diazoniuni compounds have appeared in the last 25 years. Of these, two, one by Saunders²¹⁸, and the other by Zollinger', were published prior to 1965. More rcccntly, Overberger and colleagues?, have discussed organic compounds with nitrogen-nitrogen bonds and have included a section on diazoniuni salts. **A** chapter on carbonium ion formation from diazonium ions, written by Friedman²¹⁹, is contained in the series on carbonium ions by Olah and Schleyer.

The reactivity and stability of arencdiazonium ions have been reviewcd recently by Zollinger²²⁰, and the diazotization of heterocyclic primary amines has been discussed by Butler?21. Porai-Koshits' summarized the current state of the problem of the structure and reactivity of aromatic diazo compounds.

Most textbooks refer to diazoniuin salts as ionic compounds in which the diazo group is bonded to an aromatic residue, so that the diazonium ion is the appropriate cation. It was originally believed¹ that aliphatic diazonium salts are incapable of existence. There are, however, evidences that stable non-aromatic diazoniuni salts can be prepared²²² and olefin diazonium compounds are also reported²²³.

For the purposcs of the present discussion **we** will consider that any cation of the form RN; **may** be properly considcrcd as a diazonium ion. Consequently, the ion $N₂H₂⁺$, discussed previously in section III.C, qualifies as the simplest diazonium ion. However most of the subsequent discussion will be concerned with benzenediazonium salts, and cation and monosubstituted forms, since most of the theoretical work has dealt with these compounds.

B. Crystal Structure and Spectroscopic Data

What appears to be the first crystal structure information on diazonium salts was reported in **a** preliminary short note by Ramming in *1959'?'.* This was followed by more-complete data in 1963²²⁵. A single crystal of benzenediazonium chloride $(C₆H₅N₂Cl)$ was examined by X-ray methods. The crystals are orthorhombic, space group C222, with unit cell dimensions: $a = 15.152 \text{ Å}$; $b = 4.928 \text{ Å}$; $c = 9.044 \text{ Å}$. The shortest Cl⁻ to N distance is 3.225 Å . The space group requires the $N_{(2)}-N_{(1)}-C_{(1)}\cdots C_{(1)}$ atoms to be situated on a two-fold axis of symmetry so the distances between these atoms could **be** found from one projection normal to this axis. Planes containing chloride ions bisect the $N-N$ bond of the diazonium ions, and every $(N=N-)$ ^r group is surrounded by four chloride ions in a planar arrangement normal to the $N-N$ axis, two of them are at a distance of 3.184 Å and two at a distance of 3.511 Å from this axis. The distances from the former chloride ions, the connecting line of which runs nearly normal to the planc of the benzcnc ring, to the N₍₂₎ and N₍₁₎ are 3.225 Å and 3.237 Å, respectively, where N₍₁₎ refers to the nitrogen atom attached to $C_{(1)}$. The corresponding distances from the other set of chloride ions to the N₍₂₎ and N₍₁₎ atoms are 3.548 Å and 3.559 Å. The latter chloride atoms arc approximately in thc plnnc of the benzcnc ring and arc scparatcd from the $H_{(2)}$ atoms by only 2.5 Å. As Romming points out, this is half an angstrom less than the sum of the van der Waals radii, so that a strong interaction is presumed to **exist** bctwcen thesc atoms.

Classical valence concepts suggest that the $N_{(1)}$ atom is equivalent to the nitrogen atom of an ammonium ion and should carry the positive charge. In the structurc studied by Rømming, the distances from the $N_{(2)}$ atom to the chloride ions are nearly equal to, but slightly smaller than, those from the $N_{(1)}$ atom to the chloride ions.

Further, in the case of benzenediazonium chloride, it might be expected that some shift of electron density from $N_{(2)}$ to $N_{(1)}$ might occur. If the shift is substantial enough, the two nitrogen atoms might then share the positive charge, thereby resulting in the approximately equal separations of the two nitrogen atoms from a chloride ion.

Table **41** summarizes the internuclear distances and angles found by Kamniing for the benzenediazoniuni ion. The NN bond length is approximately that found for gaseous nitrogen. The $N_{(1)}-C_{(1)}$ bond is 0.085 Å shorter than the aliphatic N-C bond. The carbon-carbon bonds are **all** found to be shorter than those in benzene **(1.397 A).** The perturbation of the CCC angles is not unexpected as a consequence of the inductive effect of the N_2^+ group. Interestingly, however, the distortion in the diazonium case is more substantial than that found with benzonitrile²²⁶, thereby demonstrating the stronger inductive effect of the N₂⁺ group when compared with the nitrile group. Of course, it is well to note that the bcnzonitrile data to which reference is made were measured in the gas phase.

TABLE 41. Internuclear scparations and **angles** in the benzencdiazoniirni ion²²⁵

Bond	Length (\hat{A})	Bond	Angle (degrees)
$N_{(1)} - N_{(2)}$	1.097	$N_{\Omega} - N_{\Omega} - C_{\Omega}$	180.
$N_{\rm th} - C_{\rm th}$	1.385	$C_{(1)} - C_{(2)} - C_{(3)}$	117.6
$C_{(1)} - C_{(2)}$	1.374	$C_{(2)} - C_{(3)} - C_{(4)}$	119.8
$C_{\text{eq}}-C_{\text{eq}}$	1.383	$C_{(3)} - C_{(1)} - C_{(5)}$	$121 - 7$
$C_{(3)} - C_{(4)}$	1.376	$C_{10} - C_{11} - C_{22}$	124.8

At about the same time, Andresen and Rømming 227 reported on the crystal structure of benzenediazonium tribromide. The compound crystallizes in the monoclinic system with space group C2/c. The tribromide ions are situated in centres of symmetry and the benzcnediazoniurn ions on two-fold axes. **The** Br-Rr bond length is 2.543 Å, and the $N-N$ and $N-C$ bond lengths in the diazonium ion are **1.11** and **1.41 A,** respcctively, the latter *two* agreeing reasonably well with the results found for the chloride. The tribromide ion is centrosymmetric. The brominebenzene plane separation is **3.68 A** which is approximatcly 0.10 *8,* less than thc sum of the van der Waals radius for bromine and the half-thickness of the benzene molecule. The shortest distance between a central bromine and a nitrogen atom is **4.34 A.** However, the shortest distance from any broniinc atom to any nitrogen atom is 3.31 Å , where the nitrogen atom is $N_{(2)}$, that is, an outer nitrogen, and the particular Br here is believed to carry the negative charge. This suggests that $N_{(2)}$ carries a substantial positive charge, again in agreement with Romming's conclusions for benzenediazonium chloride.

Rømming and Waerstad²²⁸ isolated the crystalline, orange-red complex of a diazonium salt and cuprous halide obtained from a Sandmeyer reaction at low temperature and in concentrated sollition, and found it to consist of normal benzenediazonium ions and $(Cu_2 Br_3)$ ions. Each of the 'inner' nitrogen atoms, N₍₁₎, is surrounded by four bromine atoms lying in a plane approximately perpendicular to the C-N-N direction **with** a nitrogcn to bromine distance of **3.65 A.**

Although extension of the present discussion to so-called double diazonium salts $(Ar N₂X+MXn)$ may lic outside the intended scope of this chapter, some mention of these compounds should be made for the sake of completeness. Porai-Koshits and coworkers have published the structure of *thc* double salt of ferric

chloride and o-methoxybenzenediazonium chloride²²⁹ and of p -N,N-dimethylaminobenzenediazonium chlorozincate^{230}. In the former case the ionic structure **[o-CH30CGHIN,]+.[FeCI,]-** was shown to apply and it was suggested that the positive charge lies primarily on the terminal nitrogen atom, $N_{(2)}$. The length of the $C_{(1)}-N_{(1)}$ bond was measured as 1.48 Å, substantially larger than that found by Rømming for the chloride²²⁵ or the bromide²²⁷. However the $N_{(1)}-N_{(2)}$ length of 1-11 *8,* was in agreement with the earlier values. The average distance between chloride ion and nitrogen atom was found to be 3.36 and 3.53 Å, for $N_{(2)}$ and $N_{(1)}$ respectively, again suggesting a concentration of positive charge on the terminal nitrogen atom, $N_{(2)}$.

In the case of p-N,N-dimethylaminobenzenediazonium chlorozincate²³⁰, the structure was found to consist of planar cations $[(CH₃)₂NC₆H₄N₂]+$ and tetrahedral $[ZnCl₁]$ ²⁻ anions. In this case no evidence was found for the shortening of the $C_{(2)}-C_{(3)}$ or $C_{(5)}-C_{(6)}$ bonds in the benzene ring, although the $C_{(6)}-C_{(1)}-C_{(2)}$ angle is reported as 130". Again, although **a** division of electron density was not possible, the results were interpreted as suggesting that $N_{(2)}$ possesses a positive charge.

C. Theoretical Studies

What appears to be the first calculations done on any diazonium ion wcre reported by Schuster and Polansky²³¹ in 1965. π -Bond orders and electron densities as calculated by the Huckel method were given for a number of diazonium ions including that of methyl and phenyl. In all cases for RN_2^+ , the nitrogen atom adjacent to the R group had a calculated π density of approximately $+0.8$ to $+0.9$ while that on the neighbouring nitrogen atom was approximately $+0.1$ to $+0.2$. Klasinc and Schulte-Frohlinde²³² performed the same type of calculation on a variety of substituted phenyldiazonium ions and established correlations between the rate constants for thermal decomposition of the diazonium ions and the $C-N$ bond orders, and a quantity labelled $(\Delta H)_n$ which is related to the difference between the π -electron energies of the diazonium and corresponding carbonium ion.

Russian workers^{233, 234} were also among the first to calculate properties of diazoniurn ions using Hiickel theory. Correlations between absorption energies and the differences in energies between the highest occupied and the lowest unoccupied orbitals wcre developed.

Evleth and Cox²³⁵ measured the visible-ultraviolet absorption spectra of benzenediazonium, 2-, **3-, 4-methoxybenzcncdiazoniurn,** 2,4-, 25, 2,6-, 3,4-dimethoxybenzenediazonium. and 2,3,5-, 2,4,5-, 2,4,6-, **3,4,5-trinicthoxybenzenediazonium** cation in the 200-450 nm region. Hiickel theory was employed to calculate transition energies which correlated well with those measured experimentally. The so-called Coulomb and resonance parameters were adjusted to provide the best fit with the first two electronic transitions in the benzencdiazonium cation. π -Electron densities of **1.34** and 0.71 were calculated for the adjacent and terminal nitrogen atoms ot' this cation.

The results of experiments and calculations on the electronic absorption spectra of the benzenediazonium cation and its p- and m-substituted forms were reported by Sukigara and Kikuchi²³⁶. The near-ultraviolet absorption spectrum of benzenediazonium fluoroboratc in cthanc! solution produced two peaks **at** wavclengths 259 and *395* nm, and molar extinction coefficicnts 5640 and 1160, respcctivcly, \vhilc in qucous solution the peaks appear at 263 and 298 nm. From their observed fluorescence cniission and cxcitation spectra, thesc authors concludcd that the radiative transition occurs from the same excited state, when the system was excited to the lowest excited singlet state. No emission was observed when the diazonium salt was excited by 259 nm light. The phosphorcscence emission spectrum **(77** K) was observed only when the diazonium salt solution was excited by a 295 nm light. On the other hand, an e.s.r. signal **(77** K, methanol solution) was obtained only when the diazoniurn salt was irradiated by a **259** nm light. The observed c.s.r. spectrum resembled that obtained for the phenyl o-radical.

Sukigara and Kikuchi²³⁶ used the earlier X-ray structural data of Rømming²²⁵ in their Pariser and Parr²³⁷ calculations on the benzenediazonium cations. Values of -2.39 and -2.42 eV were taken for β_{CC} and β_{CN} , respectively, where

$$
\beta_{AB} \equiv \int X_A^*(1) H_{\text{core}} X_B(1) dv
$$

 β_{NN} and ω (the π_{z} electron density on the nitrogen atom attached to the carbon atom and x is the direction in the plane of the ring and perpendicular to the NN **axis)** were taken as adjustable parameters whose best values were **-3.10** eV and **1-05,** respectively. Sukigara and Kikuchi ascribe the 295 nm and 259 nm absorption bands to the ${}^{1}A_{1}{}^{-1}B_{1}$ and ${}^{1}A_{1}{}^{-1}A_{1}$ transitions respectively, the calculated transition energies (experimental in brackets) being **4.1** 8 **(4.20)** and **4.75 (4.79)** cV, respectively. They also deduce from thcir results that the nitrogen atom adjacent to the ring bears almost all of the positive charge of the benzenediazonium cation and that the NN bond order is approximately three.

Sukigara and Kikuchi conclude that two processes occur from the excited **IB,** state, namely, radiative transition to the ground state and intersystem crossing to a phosphorescent state. No detectable photochemical decomposition occurs after the ¹A₁-¹B₁ transition. Since neither fluorescence nor phosphorescence was detected after the ${}^{1}A_{1}$ - ${}^{1}A_{1}$ transition, no radiative process occurs from the excited ${}^{1}A_{1}$ state and no internal conversion or intcrsystem crossing to **a** radiative state occurs. From this excited state the molecule presumably decomposes rapidly to givc phenyl c-radical.

In their second paper, Sukigara and Kikuchi²³⁶ measured the visible and nearultraviolet absorption spectra of p -fluoro, p -chloro, p -bromo-, p -iodo, p -methoxy and **p-dimethylamino-benzenediazonium** cations and applicd the PP method in their calculations. Table **42** summarizes their cxpcriniental results.

In their PP calculations on the p -substituted cations, Sukigara and Kikuchi use the electron affinity of the electron-accepting group $-C_6H_4N_2^+$ and the core resonance integrals, β_{CX} , where $X = F$, Cl, Br, I, O or N, as adjustable parameters. These authors conclude that thc strong absorption in the visible and nearultraviolet regions can be assigned to **the lA,-'A,** transition, and that the weak

absorptions of the ${}^{1}A_{1}{}^{-1}B_1$ transition are hidden in the large absorption bands of the ${}^{1}A_{1}{}^{-1}A_{1}$ transition. The calculations also show the large stabilizing effect that an electron-donating group at the para position exerts on the lowest excited ${}^{1}A_1$ electronic state. Evidence is also presented to support the contention that the strong absorption bands of the p -iodo-, p -methoxy- and p -dimethylamino-benzenediazoniuni cations have the character of intramolecular charge-transfer bands, while the strong bands of the p -fluoro-, p -chloro- and p -bromo-benzenediazonium cations are associated with local excitation within the benzenediazoniuni franiework.

In addition, it was shown that the vibrational frequency of the NN group increased linearly with the $N-N$ bond order. Further the $N-N$ bond order decreases and the contribution of charge transfer to the ground statc increases with an increase in the electron-donating capability of the substituent group. This suggests that the contribution of a resonance structure
 $x = \sqrt{\frac{1}{N}} \times \sqrt{\frac{1}{N}}$ suggests that the contribution of a resonance structure

$$
X = \left\langle \begin{array}{c} \overline{} \\ \overline{} \end{array} \right\rangle = N = \overline{N}
$$

becomes larger in the ground state as the electron-donating capability **of** thc para substituent increases. This had previously been predicted by Zollinger¹.

Finally, Sukigara and Kikuchi examined the electronic structures and electronic absorption spectra of m -fluoro-, m -chloro-, m -bromo- and m -methoxy-benzenediazoniurn cations. **A** summary of their cxperimcntal rcsults is given in Table 43.

TABLE 43. The observed peak wavelengths and molar extinction coefficients for meta-substituted benzencdiazoniiim cations?". Rcproduced **by** permission of the Chemical Society of Japan

Substituent	λ_{\max} (nm)	$\log \varepsilon_{\rm max}$
F	262 308	4.160 3.457
СI	266 317	4.000 3.299
Br	268.5 323	3.625 2.972
OCH ₂	273.5 348	3.982 3.475

Treiger and coworkers²³⁸ also applied the PP method to the benzenediazonium cation. Again β_{NN} was taken as an adjustable parameter whose best value was found to be -4.50 eV. The nitrogen atom adjacent to the carbon atom **was** calculated to possess a charge of $+0.58$, while that on the remaining nitrogen atom was $+0.27$ in semi-quantitative agreement with the results of Hückel calculations 2^{33} . These authors notc that, in the transition to the cxcitcd state with an cncrgy of 4.04 eV $(A_1 \rightarrow B_1)$, electron density is transferred from the *ortho* and the *meta* positions of the benzene ring to the diazonium group, resulting in a decrease in bond order of the $N-N$ bond and the positive charge on the diazonium group, while in the case of the transition with an energy of 4.77 eV $(A_1 \rightarrow A_1)$ the transfer occurs from the para position of the benzene ring.

Treiger and Bagal^{2:59} applied the Pariser-Parr method²³⁷ to a series of substituted benzencdiazonium cations in their ground and lowest excited states. **As** in the work of Sukigara and Kikuchi²³⁶, values of adjustable parameters were obtained by fitting to experimental data for the bcnzcnediazoniuni cation. **A** linear correlation was established betwcen the frequcncics **(2250-2305** cm-l) of the strctching vibrations of the N-N bonds and the calculated values of the π -electronic orders of these bonds for the various substituted benzencdiazonium ions. However, the largest deviation from this linear correlation was displayed by those molecules containing elcctron-donating substituents in the *orrho* and *pnm* positions, particularly in the case of the methoxy group. The authors suggest that the method of calculation employed is unsuitable when the resonance structure contributes substantially to charge transfer.
Treiger and Bagal²³⁹ also calculated values for some of the electronic transitions

and found reasonably good agreement $(\pm 0.04 \text{ eV})$ with the experimental values in most cases. These authors conclude that the absorption due to charge transfer in the para derivatives proceeds from the 4.71 eV band of the unsubstituted compound, while for the *rm/n* and *orrho* derivatives it can be referred to the **4-18 cV** band.

Quite recently²⁴⁰ low- and high-energy photoelectron spectroscopy and *ab initio* molecular orbital calculations have been applied to diazocyclopentadiene $(C_5H_4N_2)$. **It** appears that this is the first report either of such experiments or of such calculations on any diazo compound of the form RNN where R is larger than $CH₂$. Single determinant, all-electron, SCFMO calculations employing a basis of gaussian functions fitted to three Slater-type orbitals were performed on diazocyclopentadiene and its singly ionized states. Bond lengths of **1.130, 1.340, 1.431** and **1.104** *8,* were taken for $N-N$, $C-N$, $C-C$ and $C-H$, respectively.

Table **44** summarizes the calculated charge distributions of diazocyclopentadiene where $C_{(1)}$ is attached to $N_{(1)}$. The diazo group in this molecule is essentially neutral. However, the nitrogen atom attached to the carbon atom of the ring shows a net increase in its π population compared to that of the nitrogen atom in the nitrogen molecule. This increase is balanced by a decrease in **2s** and 2po density from that found in N_2 . The four remaining carbon atoms not attached to nitiogen each have approximately the same negative charge which is approximately balanced by the positive charge on the attached hydrogen atom. The carbon atom, $C_{(1)}$, attached to nitrogen, has a small net positive charge, and more π but less σ density than the

	$N_{(1)}$	$N_{(2)}$	$C_{(1)}$	$C_{(2)}$	$C_{(3)}$	$H_{(2)}$	$H_{(3)}$	N (in $N2$)
1s	1.995	1.997	1.993	1.994	1.994	0.710	0.714	1.997
2s	1.481	1.818	1.161	1.287	1.299			1.766
$2p\sigma$	1.093	1.139	1.547	1.909	1.966			1.236
$2p\pi$	1.192	1.261	1.255	1.093	1.053			2.000
	$(1.289)^{a}$	(0.740)						
Atomic								
charge	-0.05	0.05	0.05	-0.28	-0.31	0.29	0.29	0
					π -bond overlap populations			
	$N_{(1)} - N_{(2)}$ 0.32	$C_{(1)}-N_{(1)}$ 0.21	0.13	0.40	$C_{(1)}-C_{(2)}$ $C_{(2)}-C_{(3)}$ $C_{(3)}-C_{(4)}$	0.18	$N-N$ (in $N2$) 0.50 ^b	

TAULE 44. Calculated charge distribution **of diazocyclopentadiene240.** Reproduced, with permission, from Aarons and coworkers, *J. Chon. Soc., Furaduy Tratrs. 11, 6,* 1106 (1974).

Valucs in brackets are for the nitrogen **2p** orbital in thc molecular plane and are not included in the π overlap populations.
^b For one π orbital only.
other four carbon atoms. The N-N π -bond overlap population is much smaller in this diazo molecule than in N₂, and the authors interpret the C-C π -bond overlap populations as suggesting that a dicnc-type structure may be more appropriate than a delocalizcd one. Finally they regard the bonding as resulting from an initial $\sigma \rightarrow \pi_{\kappa}$ electron promotion in N₂ followed by π bonding with the five-membered cyclopentadienylidene ring.

Some calculations have been performed in this laboratory²⁴¹, using the Gaussian 70 programme²⁹ with an STO-3G basis, to obtain more information on the diazonium cation. The calculations were done on the benzenediazonium cation and on the same cation with a fluorine substituted at the *ortho, meta* or *para* positions of the ring. Rømming's²²⁵ geometry was employed in all cases. The energy decreased as the fluorine was removed from the *orrho* to the *mefa* to the *para* position, although as total decrease was only **3** kcal/mol this may not be significant, given the approximate nature of the calculations. What is more interesting, however, is the observation that the charge on the nitrogen atom $N_{(1)}$ attached to the carbon atom is $+0.09$, while that on the remaining nitrogen atom, $N_{(2)}$, is $+0.28$. The carbon atom C₍₁₎ itself carries a charge of $+0.10$. No significant change in this distribution of charge is observed when fluorine is attached at any one of the *ortho, meta* or *para* positions. In addition, the calculations show that the σ electrons yield contributions to the total charges of $N_{(1)}$ and $N_{(2)}$ of $+0.43$ and $+0.10$, yield contributions to the total charges of $N_{(1)}$ and $N_{(2)}$ of $+0.43$ and $+0.10$, respectively while the π electrons produce -0.34 and $+0.18$, respectively, suggesting that $N_{(1)}$ has a deficiency of σ ele cases greater than 50% of the positive charge is distributed around the benzene ring, most of it rcsiding on the hydrogen atoms.

For comparison purposes similar calculations were performed on several diazo compounds, some of which may be hypothetical in the sense that they have not yet been synthesized. Atomic charges obtained are **as** follows:

In these cases the charge on the nitrogen atom $N_{(1)}$ attached to the carbon atom is much more dependent on the groups attached to the carbon atom.

V. ACKNOWLEDGEMENT

The financial support of the National Research Council of Canada is gratefully acknowledged.

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CHAPTER 2

Diazonium-diazo equilibrium

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I. INTRODUCTION

Diazonium ions $RN_{\alpha}^{+}(R =$ substituted arene, alkene, heterocyclic compound, alkane, etc.) are strong Lewis acids. They form covalent bonds with a number of bases with formation of diazo compounds, e.g. benzenediazonium ion with OH- gives benzenediazohydroxide C_6H_5N = NOH. Diazo compounds differ from azo compounds, such as azomethane $CH_3N=NCH_3$, in that the base is bound to nitrogen by an atom other than carbon (diazocyanides, $ArN = NCN$, are an exception). In principle there is of course no difference between the diazo compound, for example a diazohydroxide or a diazocyanide, and the azo compounds, such as azobenzene¹.

The diazonium ions which have a hydrogen atom at the α -carbon (or an acidic hydrogen at some other position, see Section IV) may also react with bases as Brönsted acids, vielding diazo compounds, e.g.

$$
C_2H_3O_2CCH_2N_2^+ + B \xrightarrow{\bullet} C_2H_3O_2CCH = N = N + BH^+
$$
 (1)

The present chapter discusses the kinetics and mechanism of reversible formation of diazo compounds from diazonium ions and nucleophiles. The main emphasis is on the reaction of arenediazonium ions with hydroxide ions.

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II. DIAZONIUM-DIAZO EQUILIBRIUM OF ALIPHATIC DIAZO COMPOUNDS

The equilibrium system diazoalkane \Rightarrow alkanediazonium ion \Rightarrow alkanediazotate (equation 2) cannot be studicd, because alkanediazonium ions are not stable and usually decompose² immediately with formation of nitrogen and compounds derived from the cation RCH⁺: **DIAZO COMPOUNDS**
allibrium system diazoalkane \Rightarrow alkanediazonium ion \Rightarrow alkanediazotate
n 2) cannot be studied, because alkanediazonium ions are not stable and
decompose² immediately with formation of nitrogen and

RCH₂N₂ + OH⁻
$$
\xrightarrow{\longrightarrow}
$$
 RCH₂N = NOH $\xrightarrow{\longrightarrow}$ RCH₂N = NO⁻ + H⁺ (2)
\n
$$
\left|\begin{array}{ccc}\n-\text{II}^+ \\
-\text{II}^+ \\
\text{RCH} = N_2\n\end{array}\right|
$$

Existence of the diazonium ion can be infcrred, in some cases, from the structure of products and the presumed reaction mcchanism. When R is strongly electron withdrawing, a diazo compound is formed on nitrosation of the amine under suitable conditions^{2, 3}: Existence of the diazonium ion can be inferred, in some cases, fro

products and the presumed reaction mechanism. When R is stron

drawing, a diazo compound is formed on nitrosation of the ami

conditions^{2, 3}:

C₂H₃ **e** inferred, in some cases, from the structure of
mechanism. When R is strongly electron with-
ed on nitrosation of the amine under suitable
 $\,_{\rm N}N = NOH \xrightarrow{-OH^-} C_2H$, $O_2CCH_zN_z^* \xrightarrow{-H^+} C_2H$, $O_2CCH = N_z$ (3)
alkanediazot

$$
C_2H_1O_2CCH_2NH_2 \xrightarrow{HONO} C_2H_1O_2CCH_2N=NOH \xrightarrow{-OH^-}
$$

$$
C_2H_1O_2CCH_2N_2^+ \xrightarrow{-H^+} C_2H_1O_2CCH=N_2
$$
 (3)

In strongly basic media, the primary alkanediazotates $RCH_aN₂O⁻$ are transformed into a mixture of diazoalkane and the products derived from $RCH₂⁺$ (Reference 4). Obviously the diazoalkane is formed by action of base on $RCH_2N_2^+$ which is present in a very slight amount, according to the equilibrium shown in equation **(2).**

The instability of alkanediazonium ions is further demonstrated by acid-catalysed decomposition of diazoalkanes. The decomposition of diazodiphenylmethane is a general acid-catalysed reaction⁵, which means that the decomposition of the diazonium ion is faster than **the** splitting off of the proton.

$$
(C_{6}H_{5})_{2}CN_{2} \xrightarrow{H^{+}} (C_{6}H_{5})_{2}CHN_{2}^{+} \xrightarrow{X^{-}} (C_{6}H_{5})_{2}CHX+N_{2}
$$
 (4)

Some substituted alkenediazonium ions can exist in the form of salts with anions of strong acids as SbF_{σ} (Reference 6) for example. The increased stability is due to resonance of the diazonium group with p and π electrons. Nucleophiles react with the activated double bond which results either in substitution (if halogens are present instead of RO), or in addition of H₂O or ROH giving an alkanediazonium ion which is immediately decomposed⁶.

111. ARENEDIAZONIUM ION \rightleftharpoons **DIAZOHYDROXIDE** \rightleftharpoons **DIAZOTATE EQUILIBRIA**

In contrast to alkancdiazonium ions, arenediazonium ions are usually quite stable in aqueous solutions in the form of salts with anions of strong acids. This stability is duc to resonance interaction of the diazonium group with ring π electrons (and elcctrons of substitucnts in certain cases) and to the fact that the bond between nitrogen and **a** sp'-hybridized C atom is stronger than that with the sp3-hybridized C atom.

Substituted arenediazonium ions can react with nucleophiles in three ways' which are schematically represented below for the reaction of 4-hitrobenzenediazonium ion with hydroxyl ion (the first reaction proceeds by the S_N1 mechanism, and the water molecule is the real nucleophile^s). The first and second reactions are practically irreversible, the third one is reversible and much faster except for diazonium ions having strongly activated rings such as 2,4-dinitrobenzenediazonium ion⁹ or 1,4benezenetetrazonium ion¹⁰, $N_2C_6H_4N_2$, for which the rates of the second and third reactions are comparable.

A. Development of Ideas Regarding the Reactions ArN₂⁺+OH⁻

In **IS58** Griess described the preparation of a diazonium salt by action of nitrous gases on picramidel'. When studying the reactions of further similarly prepared diazonium salts he found¹² that benzenediazonium salt is transformed by action of potassium hydroxide into a crystalline potassium salt, the reaction being reversible. Development of azo chemistry and its application in thc dyestuff industry was influenced in an important way by investigations of Schraube and Schmidt¹³ published in **1594.** The sodium salt prepared by action of sodium hydroxide on p-nitrobenzenediazonium compounds did not couple with aromatic hydroxy compounds, and on acidification it slowly gave the original diazonium salt. Similarly, while the potassium salt prepared from benzenediazonium salt coupled readily with alkaline 2-naphtholate, it rearranged on heating for several hours with concentrated potassium hydroxide into a stable, non-coupling isomer. Since the sodium salt prepared from the 4-nitrobenzenediazonium salt gave **4-nitro-N-methylnitrosan1inc** by reaction with methyl iodide, the authors¹³ presumed that the isomers not capable of coupling were salts of derivatives of nitrosamine, and that the reactive isomcr was a diazotate. Hantzsch, who studied the stereochemistry of aromatic oximes several years earlier, came to the conclusion that on the basis of analogy between both types of compounds the matter was stereoisomerism in this case, *too''.* **Is.** Hc ascribed the syn structure to the reactive form and the *anti* structure to the non-reacting one.

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The existence of two stereoisomers was proved later in the case of the structurally similar azobenzene^{16, 17} and substituted benzenediazocyanides^{18, 19}; the i.r. spectra of substituted *syn*- and *anti*-benzenediazotates were also interpreted²⁰ on the basis of the structures suggested by Hantzsch.

Two equivalents of hydroxyl ions are consumed in the transformation of a diazoniuni ion into the respective diazotate. Hnntzsch follo\ved **this** reaction conductometrically²¹, and assumed that the reaction proceeds in two separate steps, as with other dibasic acids, and he even described thc isolation of the presumed diazohydroxide and the measurement of its dissociation constant.

Wittwer and Zollinger²² measured the diazonium \Rightarrow diazotate equilibrium with three substituted benzenediazonium salts and found that the neutralization curve showed but one step, even though two equivalcnts of base were consumcd for one equivalent of diazonium ion. This case was similar to that described before by Schwarzenbach²³. The diazonium ion reacts with the hydroxyl ion as a Lewis acid. The equilibrium constant K_1 of this reaction is smaller than the equilibrium constant *K₂* of the reaction of the formed diazolydroxide (which is a Brönsted acid) with a further OH⁻ ion. Therefore, the diazohydroxide is transformed quickly into the diazotate ion, so that the concentration of thc former **is** slight throughout the measurement. This result was confirmed also by spcctral dctcrniination of the diazonium ion \Rightarrow diazotate equilibrium in a series of substituted benzenediazonium ions?'. Only the absorption bands corrcsponding to the starting diazoniuni ion and final diazotate ion were found in the spectra. The slopes of the dependence of $log[ArN^{\dagger}_{\sigma}]/[ArN_{\sigma}O^{-}]$ (calculated from the absorbances) on pH had the value -2 . This means that the diazoniiim ion reacts practically **at** oncc with two equivalents of OH- ion, and the diazohydroxidc concentration is negligible. I. Therefore, the diazohydroxide is transformed quickly into
aat the concentration of the former is slight throughout the
sult was confirmed also by spectral determination of the
tate equilibrium in a series of substitute were found in the spectra. The slopes of the dependence of

(calculated from the absorbances) on pH had the value -2 .

Iliazonium ion reacts practically at once with two equivalents of

azohydroxide concentration is neg

$$
ArN_{2}^{+}+OH^{-} \xrightarrow{\longrightarrow} ArN_{2}OH \qquad K_{1}
$$
 (6)

$$
ArN_{2}OH + OH^{-} \xrightarrow{\cdots} ArN_{2}O^{-} + H_{2}O \quad K_{2}
$$
 (7)

6. *Effect of Benzene Ring Substituents on the Diazonium /on* ..+ *syn-Diazotate Equilibrium*

In alkaline aqueous medium the following rcversiblc rcactions of diazonium ions take place^{25, 26}: syn-ArN₂OH syn-ArN₂OH syn-ArN₂OH syn-ArN₂OH syn-ArN₂OH syn-ArN₂OH syn-ArN₂OH syn-ArN₂O+ (7) *R₁* syn-ArN₂O⁻+H₂O *K*₂ (7) *kyn-ArN₂OH* syn-ArN₂O+ (7) *kyn-ArN₂O+ (7) kyn-ArN₂O+ (7) k*

$$
ArN_{2}^{+} + OH^{-} \xrightarrow[k_{1}]{}^{k_{1}} \text{syn-ArN}_{2}OH \qquad K_{1}
$$
 (6)

$$
syn-ArN2OH+OH- \xrightarrow{\longrightarrow} syn-ArN2O-+H2O K2
$$
 (7)

$$
syn-ArN_{2}OH \xrightarrow[k_{-8}]{k_{3}} anti-ArN_{2}OH \t K_{3}
$$
 (8)

$$
syn-ArN_{2}OH \xrightarrow[k_{-s}]{k_{s}} anti-ArN_{2}OH \tK_{3}
$$
 (8)
\n
$$
syn-ArN_{2}O^{-} \xrightarrow[k_{-s}]{k_{s}} anti-ArN_{2}O^{-} \tK_{4}
$$
 (9)
\nanti-ArN_{2}OH+OH^{-} \xrightarrow[anti-ArN_{2}O^{-}+H_{2}O K_{5}] (10)

$$
anti-ArN_2OH+OH^- \xrightarrow{\longrightarrow} anti-ArN_2O^- + H_2O \quad K_s \tag{10}
$$

Benzenediazonium ions which have a hydroxyl group in the *ortho* or *para* position split off the proton even in mildly acidic medium; the pK_a value of 4-hydroxybenzenediazonium ion is 3.40²⁷. The diazo compounds formed are only slightly reactive and do not form diazotates evcn in a strongly alkaline medium.

$$
HO - \left(\bigcup - N_2^+ \right) \rightleftharpoons \left[-O - \left(\bigcup - N_2^+ \longleftrightarrow O = \left(\bigcup - N_2\right) + H^+ \right] \tag{11}
$$

Besides these reactions, irreversiblc substitutions of the nitrogen or substituents in the *orrho* or para positions by **OH-** group (equation 5) and decomposition rcactions of diazohydroxides (leading to formation of hydroxy compounds and othcr, often polymeric, substances) may be significant. These side-reactions make determination of equilibrium constants difficult. For example, *m*-nitrobenzenediazohydroxide decomposes so rapidly that at **a pH** of about *9* (the region of its maximum concentration) practically all the diazonium salt is transformed into the dccomposition products within less than a minute.

With most substituted benzenediazonium ions the rate of $syn-anti$ isomerization is many orders of magnitude slower than that of the reaction of the diazoriium ion Is many orders of magnitude slower than that of the reaction of the diazonium ion
with OH-, so that it does not interfere with the determination of the ArN:
 $\frac{1}{2} \approx syn-$ ArN:
 $\frac{1}{2} \approx syn-$ ArN:
 $\frac{1}{2} \approx syn-$ ArM:
 $\frac{1}{2} \approx$ $ArN₂O⁻$ equilibrium. Diazonium ions which have strong $-M$ type substituents (e.g. nitro groups) in *ortlio* or *para* positions form an exception. **s**
traction than that of the rate of syn-anti isomerization
agnitude slower than that of the reaction of the diazonium ion
t does not interfere with the determination of the ArN⁺₂ = syn-
n. Diazonium ions which have

The transformation of diazonium ions into syn-diazotates can also be expressed by the following equations:

$$
ArN_2^+ + H_2O \longrightarrow \text{syn-ArN}_2OH + H^+ \quad K_{A_1}
$$
 (12)

$$
syn-ArN2OH \xrightarrow{2} syn-ArN2O^- + H^+ K\Lambda2
$$
 (13)

Equilibrium constants of the overall reaction of the cases described by equations $(6)+(7)$ and $(12)+(13)$ respectively are given by the relations:

$$
K = K_1.K_2 = [ArN_2O^-]/[ArN_2^+][OH^-]^2
$$
 (14)

$$
K_{\Lambda} = K_{\Lambda 1}.K_{\Lambda 2} = [\Lambda r N_2 O^{-1}][H^{+}]^{2}/[\Lambda r N_2^{+}] = K.K_{w}^{2}
$$
 (15)

where $K_w = [H^+] [OH^-]$.

As the determination of an equilibrium does not dcpend on the path by which it **has** been reached, both methods are correct. In fact practically all arenediazoniuin ions are transformed into diazohydroxides by reaction with hydroxyl ions. The reaction of diazonium ion with water in alkaline medium is slower by many orders of magnitudc. Also diazohydroxide rcacts far quicker with hydroxyl ion than with water in the pH region used for measurements of equilibrium constants of this reaction (except perhaps for the most reactive diazonium ions). On the contrary, the use of equations such as (13) and **(14)** is more usual and more convenient for acidbase equilibria; on introducing logarithms into equation (15) we get the relation:

$$
\frac{1}{2}(pK_{\Lambda_1} + pK_{\Lambda_2}) + \log ([ArN_2O^-]/[ArN_2^+]) = pH \tag{16}
$$

This means that $(pK_{A1} + pK_{A2})/2$ equals the pH of the medium in which $[ArN_2^+]$ = $[ArN₂O⁻]$ (as in the case of pK_A of the Brönsted acids). This pH was called pH_m by Lc\\ is and Suhr?' (Table **1).**

The ratio **[ArN:]/[ArN,O-]** changes a hundrcd times on changing **pH** by unity, hence at $pH = (pH_m + 1)$ the reaction mixture contains 1% diazonium ion and 99% diazotate ion. Concentration of diazohydroxide is negligibly low (except for very reactive diazonium salts) in the whole pH range, being maximum at $pH = pH_m$. The equilibrium constants K depend on ionic strength and they increase about ten times on changing the ionic strength from **1 M** to **0.1 M ?I.**

x	K ª (l^2/mol^2)	$K^{\mathfrak{d}}$	
		(l^2/mol^2)	pH_m ^a
$4-NO2$	1.3×10^{9}	2.0×10^{10}	9.44
4 -CN	2.9×10^8	3.9×10^{9}	9.77
$3-C1$	4.0×10^6	1.9×10^{7}	10.70
4-Ci	3.8×10^{5}	6.7×10^{5}	$11 - 21$
н	1.6×10^{4}	2.8×10^{4}	11.90
4 -CH ₃	6.6×10^{2}	2.2×10^{3}	12.59

TABLE 1. Equilibrium constants K **and** pH_m **values of** the reaction $XC_6H_4N_2^+ + 2OH^- \rightleftharpoons XC_6H_4N_2O^- + H_2O$

Refcrence **24:** room temperature, ionic strength 1.0 **M.**

Reference **28: 23** *"C,* ionic strength **0.004-0.03 M.**

Dependence of log K on the Hammett σ constants is linear with the slope $\rho = 6.3^{24}$, $6.58²⁸$ or $6.66²⁹$. The *p* values found are unusually high. One of the reasons for this is that K is a product of two equilibrium constants, so that the final value is a sum of ρ_1 and ρ_2 characterizing the formation and dissociation²¹ of diazohydroxide, respectively. Similarly it is also possible to correlate pH_m with the σ constants; in this case the ρ value is negative and half that from equation (17).

$$
\log (K/K_0) = \rho \sigma = \log (K_1/K_1^0) + \log (K_2/K_2^0) = \rho_1 \sigma + \rho_2 \sigma = (\rho_1 + \rho_2) \sigma
$$
 (17)

It is impossible to determine the values of K_1 and K_2 or of ρ_1 and ρ_2 separately by measuring the concentrations or the pH of the equilibrium system; since $K_2 \gg K_1$ only the overall constant *K* is accessible. For dctcrniination of the constants of both reaction steps it is necessary to determine the rates of the reactions considered; in this case only the rcaction of diazoniuni ion with hydroxyl ion and the reverse decomposition of diazohydroxide are suitable for measurement.

C. Kinetics and Mechanism of the Reactions: Diazonium Ion \Rightarrow syn-Diazo*hydroxide* \rightleftharpoons *syn-Diazotate*

The rate of thc reaction of diazohydroxide with hydroxyl ion (equation **7)** is far greater than that of the dccornposition of diazohydroxide into diazoniuni ion and hydroxyl ion (in neutral and alkaline media). Therefore, the rate-limiting step of the transformation of diazonium ion into syn-diazotate consists in the reaction of diazonium ion with hydroxyl ion (equation 6), whereas the rate of the reverse reaction (transformation of diazotate into diazonium ion) is limited by the splitting of the diazohydroside.

$$
\vec{v} = k_1 [ArN_2^+] [OH^-]
$$
 (18)

$$
\overline{v} = k_{-1} [\text{ArN}_2 \text{OH}] \tag{19}
$$

At a pH comparable with pH_m or higher (the conditions suitable for determination of k_1) $[ArN_2OH] \ll [ArN_2O^{-}]$, and equation (20) can be written for the rate of the revcrse reaction *^c*

$$
\overline{v} = k_{-1}[ArN_2O^-]/K_2[OH^-] = k_1[ArN_2O^-]/K_1K_2[OH^-]
$$
\n(20)

The rate of formation of diazotate ion \vec{v} increases linearly with concentration of hydroxyl ion, and, on the contrary, the rate of the reverse reaction decreases linearly with $[OH^-]$. The observed rate constant k_{obs} is a sum of the constants in both directions (equation 21) and has its minimum at $pH = pH_m$ (Figure 1, curve a)

$$
k_{\text{obs}} = k_1[\text{OH}^-] + k_1/K_1 K_2[\text{OH}^-] = k_1([\text{OH}^-] + 1/K[\text{OH}^-])
$$
 (21)

when the concentrations of diazonium and diazotate ions are, after reaching the equilibrium, the same. Logarithms of the rate constants measured **by** spectrophotometric stopped-flow method^{28, 29} correlate with σ constants (equations 22, 23).

$$
\log k_1 = 2.61\sigma + 3.72 \qquad \text{(Reference 28)} \tag{22}
$$

$$
\log k_1 = 2.06 \sum \sigma + 3.97 \quad \text{(Reference 29)} \tag{23}
$$

FIGURE 1. The **pM** dependence of log *kobs* of the reaction **3-nitro-4-chlorobcnzcnediazoniurn** ion[≥]syn-diazohydroxide=syn-diazotate at 20 °C. (a) Transformation of diazonium ion into the diazotate; (b) transformation of diazohydroxidc (in equilibrium with diazotatc) into diazoniurn ion.

In spite of OH- ion being a much stronger base than the anions of C-acids (acetoacetanilide, acetylacetone) and naphtholate ions, the rate constants found arc generally lower than those of coupling reactions with the anions mentioned by several orders of magnitude³⁰. At the same time the respective ρ value (2.06) is lower than those of the coupling reactions $(3-4)$, too. This fact is inconsistent with the rule31 (which often is not fulfilled) that the more selcctive a reagent is, the more slowly it reacts. The reason is probably that the OH^- ion is strongly solvated in water, and partial desolvation must take place before the $N-O$ bond can be formed. This process necessitates considerable cncrgy and is obviously thc main cause of the relatively lower reactivity of the OH^- ion. The $N-O$ bond formation proceeds very quickly, and this step is little selective. Since the substituents in the benzene nucleus of diazonium ions have small effect on the desolvation of the OH- ion, the overall influence of substituents is relatively small which makes itself felt in the low value of the ρ constant. On the contrary, the naphtholate ions and the anions of C-acids are far less solvated, and the carbon atoms which are involved in the reaction do not form any hydrogen bonds, so that the predoniinant part of the activation energy is consumed in the coupling reaction (formation of $C-N$ bond).

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Under the conditions suitable for determination of k_1 it is possible to obtain only the complex constant k_{-1}/K_2 (equation 20) for the reverse reaction. The determination of k_{-1} (and K_2) necessitates starting with syn-diazotate and following the reaction at lower pH values when a considerable part of the diazotate is transformed into diazohydroxide in a rapid pre-equilibrium. The reaction proceeds practically only in onc direction, and *kobs* is defined by equation **(24).**

$$
k_{\text{obs}} = k_{-1}/(1 + K_2[\text{OH}^-]) = k_{-1}[\text{H}^+]/(K_A + [\text{H}^+])
$$
 (24)

With increasing proton concentration $\log k_{\text{obs}}$ increases gradually more and more slowly eventually becoming pH independent (Figure 1, curve b) when almost all the diazotate has been transformed into diazohydroxide in a rapid pre-equilibrium³².

In diluted hydrochloric acid **/robs** again increases with increasing proton concentration, and the same is true in chloroacelatc buffer for increasing concentration of chloroacetic acid; this means that the transformation of diazohydroxide into diazonium ion is a general acid-catalysed reaction³²; the value of the Brönsted coeficient is about **0.2:**

$$
k_{\text{obs}} = k_{-1} + k_{\text{H}+} [\text{H}^+] + k_{\text{HA}} [\text{HA}]
$$
\n(25)

Logarithms of the rate constants k_{-1} and dissociation constants K_{Λ_2} (Table 2) correlate with σ constants:

$$
\log k_{-1} = -3.4 \sum \sigma + 5.4 \tag{26}
$$

$$
\log K_{\Lambda 2} = 1.3 \sum \sigma - 8.1 \tag{27}
$$

TABLE 2. Rate and equilibrium constants of the rcaction TABLE 2. Rate and equilibrium constants of the reaction
arenediazonium ion + OH⁻ $\frac{k_1}{k_{-1}}$ *syn*-diazohydroxide $\frac{k_{A_1}}{\sqrt{K_{-1}}}}$ *syn*-diazotate + H⁺

Reference 29.

$$
b \text{ At } pH = pH_{in}.
$$

The equations (23). *(26)* and (27) describe quantitatively the reaction of substituted benzenediazonium ions with OH⁻ ion and the subsequent dissociation of *syn*diazohydroxide to *syn*-diazotate. From the values K_1 and K_2 it is also possible to calculate the maximum concentration of syn-diazohydroxide (at $pH = pH_m$). The concentrations are slight, in accordance with the predictions of Zollinger²², and they increase with increasing value of σ constants, because K_1 increases with increasing σ value much faster than K_2 does $(\rho_1 > \rho_2)$. The constant ρ_1 of the reaction (6) can be calculated either from the difference of the ρ constants derived from the equilibrium constants *K* and K_{Λ_2} or from the rate constants k_1 and k_{-1} . The average value of p_1 is **5.4.**

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The reaction of diazonium ion with hydroxyl ion produces first the diazohydroxide which then partially isomerizes to nitrosamine ArNHNO. The proton transfer between nitrogen and oxygen (be it inter- or intra-molecular) represents a reaction several orders of magnitude faster than thc splitting off of hydroxyl ion from diazohydroxide, so that the reaction produces an equilibrium mixture of both tautomers. The extent of this isomerization is not known. The ρ constant of log K_A of substituted phenols³³ is 2.23; the inserted N=N group diminishes the ρ constant by a factor of about 0.4 so that for diazohydroxides we obtain $\rho \approx 1$. For dissociation constants of substituted formanilides³¹ (which are structurally similar to nitrosamines) the experimental ρ value was 1.53. The difference is relatively small, and the value **1.3** (equation 27) is almost in thc middle, so that, from this point of view, it is impossible to decide whether the compound is a diazohydroxide, a nitrosamine or a mixture of both.

Splitting of diazohydroxides is subject to general acid catalysis, hence the reverse reaction of diazonium ion with water must be subject to general base catalysis,

$$
ArN_{2}^{+} + H_{2}O + X^{-} \longrightarrow \left[ArN_{2} \cdots O_{N}\right]^{H} \longrightarrow ArN_{2}OH + HX
$$
 (28)

where X stands for H_2O , RCO_2^+ or even OH^- , although in the last case there is a possibility that OH^- reacts directly with the nitrogen atom of the diazonium ion.

Wittwer and Zollinger²² found that the dependence of log k on pH for the reaction of diazotized nietanilic acid with the dianion of **2-hydroxynaphthalenc-6-sulphonic** acid at $pH > 12$ is linear with a slope of -2 . The rate-limiting step is the reaction of the dianion \vith the diazonium ion which is present in a rapid pre-equilibrium with diazotate **in-** \t thc given pH, niore than 99% of !he clectrophile is present in the .. **1\120-,** and concentration of ArN: depends linearly on **[H+]?.** In the case _. ,'cry reactive diazonium ions and with thc use of sufficient conccntration of coupling component (e.g. substituted naphthol) the coupling rate becomes greater than the rate of reaction of diazonium ion with hydroxyl ion²⁹. The splitting off of OH- ion from diazohydroxide is rate limiting. Thc reaction rate is directly proportional to the first power of [H+] and independent of the concentration of the coupling component (naphtholate). *r.*

D. Kinetics and Mechanism of **syn-anti** *lsomerizations*

syn-Arenediazotate ions formed by reaction of diazonium ion and hydroxyl ion isomerize into anti-diazotate ions. Isomerization of syn-diazohydroxides, which in alkaline solution arc present in a substantially smaller concentration, takes place to a very small extent. New equilibria are established, e.g. **netics and Mechanism of syn-anti Isomerizations**
Arenediazotate ions formed by reaction of diazonium ion and hydroxyl is
ize into *anti*-diazotate ions. Isomerization of *syn*-diazohydroxides, which
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ions formed by reaction of diazonium is

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present in a substantially smaller concent

lew equilibria are established, e.g.

anti-ArN₂OH syn-

and the acid-base equilibrium

and hence also **a** new overall equilibrium

anti-ArN₂OH syn-ArN₂O⁻ anti-A

uilibrium

anti-ArN₂OH anti-ArN₂O⁻ + H⁺,

v overall equilibrium

ArN₂⁺ + 2 OH⁻ anti-ArN₂O + H₂O⁻

ubstituted benzenediazonium ions this equilibrium In the case of the substituted benzenediazonium ions this equilibrium is usually established within a fraction of a second. The syn-anti isomerization proceeds slowly (except for diazotate ions with $-M$ type substituents in the *para* or *ortho*

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position), e.g. the isomerization half-life of benzenediazotate ion is 38 min at 80 °C²⁴. In all the cases known up to now the rate constant k_1 of the reaction of diazonium ion with hydroxyl ion is greater than $k₄$ of isomerization (equation 9) at least by several orders of magnitude, so that the two reactions can be followed separately. The effect of substituents on the isomerization rate was determined with **4-** and 5-substituted 2-nitrobenzenediazotate ions³⁵ whose isomerizations are, due to activation by the o -nitro group, rapid enough even at room temperature. The respective logarithms of the constants k_4 correlate with σ and σ ⁻ constants³⁵ according to equation (29)³⁶. *syn*-2-Nitrobenzenediazotate ion is the basic member of the series.

$$
\log k_4 = 2.1[\sigma + 2.4(\sigma^2 - \sigma)] - 2.83 \tag{29}
$$

In contrast to thc syn diazotates \vhich split off hydroxyl ion within **a** half-life of several msecond or less, the anti-diazohydroxides are much morc stable. Their dissociation constants $K_{\Lambda5}$ (equation 10 but with H_2O as the reagent) can be determined by usual methods, e.g. potentiometric titration of anti-diazotates with acids^{26,37} or from the pH dependence of the transformation of *anti*-diazotate into diazonium ion39. **37s 38** (Table **3).**

Thc other rate and equilibrium constants (Tables **3** and **4)** were determined by the reverse **37-30.** . anti-diazotate was injected into a solution of acid or

TABLE 3. Rate constants **of** isomerization of substituted benzenediazotate ions (k_4) and diazohydroxides (k_{-3}) and dissociation constants of *anti*-diazohydroxides at 20 °C³⁵

Substituent	k_4 (sec ⁻¹)	k_{-3} (sec ⁻¹)	$\mathfrak{p}K_{\mathsf{As}}$
н	2×10^{-6} ⁴	1.5×10^{-2} b.c	7.29 b
$4-NO2$	5.4×10^{-2} a	4.8×10^{-3}	6.13 ^b
$2-NO2$	1.5×10^{-3}	1.9×10^{-2}	6.15
$2-NO2-5-Cl$	1.0×10^{-2}	4.0×10^{-2}	5.60
$2,4$ -diNO ₂	20.9		5∙0

Reference **40, 25** "C.

Reference **37,** 25 *"C.*

^c Rate constant of splitting of anti-diazohydroxide.

TABLE 4. Rate and equilibrium constants of *syn-anti* isomerization of substituted benzenediazotate ions at 20 *"C* **³⁴**

Substituents	k_4 (sec ⁻¹)	k_{-4} (sec ⁻¹)	K_{Λ}
$4-NO2$	5.4×10^{-2} a		600 ^b
$2-NO1 - 4-CH2$	7.8×10^{-4}	4.2×10^{-5}	18.5
$2-NO2$	1.5×10^{-3}	8.3×10^{-5}	$18 - 0$
$2-NO2-4-Cl$	4.3×10^{-3}	3.7×10^{-4}	11.5
$2-NO - 5-CI$	1.0×10^{-2}	8.3×10^{-4}	12.0
2 -Cl-4-NO ₂	3.2×10^{-2}	9.2×10^{-5}	350
2,6-Cl ₂ -4-NO ₃ ^c	5.5×10^{-3}	1.8×10^{-5}	3.0
$2-NO2-4,6-CI2$			0.70

*^a*Reference **40,** 25 *"C.*

Reference **26.**

Reference **35.**

 \dagger The generalized equation is: $\log k = \rho[\sigma + r(\sigma^+ - \sigma) + \log k_0]$. The constants ρ , σ and $\sigma^$ have their usual meaning; thc reaction constant *r* has been interpreted **as** the ratio **of** conjugation in a particular reaction series to that in the limiting cases requiring σ^- .

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buffer containing a sufficient amount of reactive coupling component. The diazonium ion thus formed reacted rapidly with the coupling component to form the rcspective azo compound, so that the overall reaction **was** pseudo first order. The reaction course is represented in Scheme 1, and the dependence of $log k_{obs}$ on pH is given in Figure 2 for benzenediazotate ion and its 2-nitro-4-chloro and 2.4-dinitro derivatives.

Anti-syn isomerization of bcnzenediazotate is very slow24, the main reaction path being the splitting of hydroxyl ion from *anti*-diazohydroxide (Scheme 2[†]); from the

FIGURE 2. pH-Dependence of log *kobe* of the conversion of unsubstituted **(U),** anti-2-nitro-4-chloro- (**A)** and **anri-2,4-dinitrobcnzenediazotate** (*CB)* into diazonium ion.

t **As** with syti-diazohydroxides, in this case also, two tautomeric forms **arc** possible: diazohydroxide and nitrosamine. The diazohydroxide is, without doubt, the form which is split into diazonium ion and hydroxyl ion. Neithcr from the splitting ratc nor from the **pH** dependence of log k_{obs} is it possible to determine the population of the both tautomers.
The constant k_{-1}^{anti} is a product of the rate constant of the splitting of *anti*-diazohydroxide and of the molar fraction of the diazohydroxide in the mixture of the two isomers. The same is true also of the constant k_{-1} of the splitting of syn-diazohydroxides.

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dependence of $\log k_{\text{obs}}$ on pH at pH > 6 and from the pH-independent region the constants $K_{\frac{1}{2}}$ and k_{-1}^{anti} (the rate constant of splitting of *anti*-diazohydroxide) were determined, respectively (Table 3). At $pH < 3$ the acid-catalysed splitting of antidiazohydroxide begins to be significant. Logarithms of $k_{\perp 1}^{n_1}$ of benzenediazotate ion and its 4-methyl, 4-chloro and 3-chloro derivatives correlate with the σ constants according to equation (30) :

$$
\log k_{-1}^{anti} = -2.6\sigma - 1.89\tag{30}
$$

The ρ value -2.6 is comparable with that (-3.4) found for splitting of syn-diazohydroxides³².

$$
H^+ + \text{anti-C}_{6}H_{5}N_{2}O^- \xrightarrow{\longrightarrow} \text{anti-C}_{6}H_{5}N_{2}OH \xrightarrow{\overset{R_{21}^{3}H^{\prime}}{\longrightarrow}} C_{6}H_{5}N_{2}^+
$$

With increasing values of the substituent constant σ the splitting rate of diazohydroxides decreases and simultaneously the isomerization rate increases. With the 2-nitro-4-chloro derivative³⁵ the splitting of the *anti*-diazohydroxide is slower than isomerization of the diazotate and diazohydroxide, and the subsequent splitting of syn-diazohydroxide; these become the main reaction paths. At the highest pH values the reaction proceeds according to Scheme 3. Splitting of syn-diazohydroxide is the

$$
anti-ArN2O- \xrightarrow{\longrightarrow} syn-ArN2O- \xrightarrow{+H+}{syn-ArN2OH \xrightarrow{k_{-1}} ArN2}
$$

rate-limiting step, and the dependence $\log k_{\text{obs}}$ *vs* pH has a slope of -1 (Figure 2) With increasing proton concentration the transformation of syn-diazotate into diazonium ion becomes more and more rapid. The *anti-syn* diazotate isomerization becomes the rate-limiting step, and the reaction rate becomes independent of pH₁. Further increase in the proton concentration results in the increase of concentration of anti-diazohydroxide whose isomerization becomes the main reaction path and, at the same time, the rate-limiting step (Scheme 4). The reaction rate, which at first increases again, finally becomes pH independent when almost all diazotate ion has changed into *anti*-diazohydroxide. The acid-catalysed splitting of *anti*-diazohydroxide becomes significant at pH about zero.

$$
H^{+} + \text{anti-ArN}_{2}O^{-} \xrightarrow{\text{non-} \text{anti-ArN}_{2}OH} - \xrightarrow{k-3} \text{syn-ArN}_{2}OH \xrightarrow{\text{non-} \text{ArN}_{2}OH}
$$

Further increase in the electron-withdrawing ability of substituents results in an even greater decrease in the syn-diazohydroxide splitting rate and an increase in the isomerization rate. The splitting of syn-diazohydroxide becomes rate limiting in a broader pH range and the reverse is true of the rate-limiting transformation of antiinto syn-diazotate. In the case of the 2,4-dinitro derivative³⁸ (Figure 2) the noncatalysed splitting of syn-diazohydroxide (k_{-1}) is rate limiting up to pH about 4.5, and the reaction rate is directly proportional to the *anti*-diazohydroxide concentration. Further lowering of pH causes the proton-catalysed splitting $(k_{\text{H+}})$ to become more and more significant. At $pH < 1$ the rate of the proton-catalysed splitting of syn-diazohydroxide is substantially greater than that of the reverse isomerization to *anti*-diazohydroxide. The transformation of *anti*- to *syn*-diazohydroxide becomes rate limiting, and $\log k$ is pH independent.

t The reaction rate does not become completely pH independent, because the individual reaction paths overlap.

E. *Some Conclusions from the Results on* **syn-anti** *lsomerizotion*

Isomerization rates of syn-diazotates to anti-diazotates depend strongly on polar effects of the substituents, especially on their $-M$ effect. The value $r = 2.4$ in equation (29), which describes the relative importance of the mesomeric effect, is probably the highest given in the literature up to now. The *anti*: *syn* diazotate concentration ratio is practically independent of the polar effects of substituents²⁶, ³⁵ but it rapidly increases wiih increasing stcric rcquirements of substituents in *ortlio* positions³⁵ (Table 4). This means either that the *syn* and *anti* derivatives have (in contrast to all other published interpretations) thc configuration *trans* and *cis,* respectively, or that the */rms* configuration is stabilized by substituents in *ortho* positions. A similar case is described in connection with the study of *syn-anti* configuration of substituted *N*-alkyl-*N*-nitrosoanilines⁴¹ where the *ortho* substitution shifts the equilibrium in favour of isomer **1** :

The isomerization rate of *anti-* to *syn-*diazohydroxides (Table 3) depends generally much less on polar effects and especially on the $-M$ effect (equation 31) than the isomerization of diazotate ions, and it is usually far quicker. **It** can be presumed that

$$
\log k_4 = 1.0[\sigma + 0.65(\sigma^2 - \sigma)] - 1.8
$$
 (31)

the isomerization mechanism is different for the two cases. Two mechanisms can be considered for diazotate ions²⁶, either rotation about the $N-N$ bond in 2 or inversion at the N atom positively charged in the transition state **3.** The rotation about the $N-N$ bond of the nitrosamine tautomer is the most probable mechanism for diazoanhydrides.

A reaction mechanism, according to which an anti-diazotate would be formed directly by reaction of hydroxyl ion with the diazoniuni ion which is in equilibrium with syn-diazotate, is untenable, since in alkaline solution the isomerization rate is independent of hydroxyl ion concentration³⁹. The mentioned mechanism would necessitate $v = [ArN_2^+][OH^-]$, and the isomerization rate would have to be inversely proportional to [OH⁻], because $[ArN_2^+]$ decreases with the square of [OH⁻].

Substituted **anti-benzenediazohydroxides** are by about one order of magnitude stronger Brönsted acids than syn-diazohydroxides. Direct comparison is possible only in the case of the *m*-nitro derivatives^{26, 32}, but it obviously holds also for the other *n7cfa-* and para-substituted diazohydroxides, because the constants of the acid-base reactions are approximately the same: 1.3 for *syn3'* and **1.17-1.45** for $anti^{26}$, 37 . Hence it follows that the ratio *anti*: *syn* will be by about one order of magnitude smaller for diazohydroxides than that for diazotate ions. These conclusions do not hold for *ortho*-substituted diazohydroxides.

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For the majority of substituted benzenediazonium ions it is possible to determine the equilibrium constant K of the reaction $ArN_2^+ \approx syn-ArN_2O^-$ by direct measurements on solutions in the pH range near pH_m . If the absorbances and the pH values of the reaction mixtures are measured after establishing the equilibrium of the syn-anti isomerization, it is possible to determine the equilibrium constant of the reaction $ArN_2^+ \rightleftharpoons anti-ArN_2O^-$. As irreversible side reactions take place simultaneously, the values obtained are usually very inaccurate, or not accessible at all (e.g., with the m -nitro derivative). In the case of the diazonium ions having strong $-M$ type substituents in *ortho* and *para* positions (e.g. 2,4-dinitrobenzenediazonium ion) the rate of syn-anti isomerization at pH close to pH_m is greater than the rate of the reverse transformation of syn-diazohydroxide to the diazonium ion, so that the equilibrium constant of the reaction $ArN_2^+ \Rightarrow syn-ArN_2O^-$ cannot be determined by measurements³⁸.

IV. DIAZONIUM-DIAZO EQUILIBRIA OF HETEROCYCLIC **DIAZONIUM IONS**

Diazotization of heterocyclic primary amines can be expressed by the following reaction sequence⁴²:

$$
HArNH2 \xrightarrow{HONO} HArNH2NO \xrightarrow{HX} HArNHNO \xrightarrow{HX} HArN = NOH
$$

\n
$$
HArN2 X - \xrightarrow{-HX} ArN2 XrN2 \xrightarrow{KrN2} ArN2 \xrightarrow{KrN2} Ar=N2
$$

In this Scheme (contrary to general usage in this series) the symbol Ar represents a heteroaromatic ring minus two hydrogen atoms and the symbol HAr represents a heteroaryl group.

If the aromatic ring contains an acidic hydrogen atom, the diazo compounds $Ar = N₂$ along with the corresponding diazonium ions are usually the main reaction products. The equilibrium $HArN_2^+ \rightleftharpoons Ar=N_2+H^+$ was measured⁴³ in a series of azols in aqueous solutions at 0° C. The acidity of the diazonium ions increased with the number of ring N atoms. In principle, the reaction is similar to that in the aliphatic series (alkanediazonium ion \Rightarrow diazoalkane + H⁺).

If the five-membered ring does not contain any acidic hydrogen atom, the product of nitrosation of the amine in diluted mineral acid is usually a mixture of the nitrosamine, the diazohydroxide and the diazonium cation. The tendency to form nitrosamines and diazohydroxides usually increases with increasing number of ring heteroatoms. High yields of nitrosamines were obtained with substituted triazoles and tetrazoles^{41, $\overline{15}$, thiadiazoles^{41, 46, ⁴⁷ and oxadiazoles⁴⁴. The situation is not quite}} clear, e.g. substituted 5-amino-I ,3,4- and -1,2,4-thiadiazoles **(4,** *5)* give high yields of nitrosamines^{14, 16}, however, the 5-substituted-3-amino-1,2,4-thiadiazole (6) forms only a relatively unstable diazonium salt^{18, 19} Commonly, the structure and yield of the isolated product depend on various factors such as the solubility of products, the rates of the substitution reactions \vith the nucleophilcs present and of decomposition reactions.

From i.r. (in solid state) and n,m,r . spectra it follows¹¹ that the products currently denoted as nitrosamines are, in fact, mixtures of the nitrosamine and the tautomeric diazohydroxide. In alkaline medium these are transformed into *anti*-diazotates⁴⁶. Dissociation constants were measured¹⁴ in the case of triazole derivatives (7); the pK_A values were between 3.54 and 3.13 for X varying from 4-CH₃ to 4-NO₂, i.e. far lower than those of substituted bcnzencdiazohydroxides (Tables **2** and 3).

Increasing proton conccntrations causes gradual transformation of the nitrosamine into the diazonium ion; the position of no such equilibrium was as yet measured. In one case the extent of transforination **was** followed1G by measuring the coupling rate of the formed **3-1iiethyl-l,2,4-thiadiazo-5-diazoniuni** ion with *2* naphthol in dilute sulphuric acid. Under thcse conditions 2-naphthol rcacts in its undissociated forni, so that the change of the coupling rate expresses thc change of the equilibrium concentration of the diazoniuni ion. The reaction rate increased steeply up to the highest sulphuric acid concentration used (1.65 M) , which means that the nitrosamine predominated all the time in the reaction mixtures.

The extraordinary stability of five-membered heterocyclic nitrosamincs (and diazohydroxides) compared with the carbocyclic aromatic diazohydroxides is due, first of all, to the strong electron-withdrawing effect of heterocyclic rings. This effect increases the reactivity of the diazonium ion and the stability of the nitrosamine in the same way as in substituted benzcnediazoniuni ions (Table 2), but to a greater extent. 1,2,4- and **1,3,4-Thiadiazole-5-diazonium** ions are far more reactive than 2,4-dinitrobenzenediazonium ion⁵⁰ (towards 2-naphthol). The isolated nitrosamines are obviously present in the form of the more stable (*anti*) isomer. Although so far no data have been published about *syn-anti* isomerization of heterocyclic nitrosamines, it can be presumed from analogy with substituted benzenediazohydroxides that the isomcrization half-lives will be shortcr than 1 sec in most cascs. Another factor given by Butler¹² is the stabilization by intramolecular hydrogen bonding in the case of the diazohydroxidc **8.** However, this form is, according to spcctral analyses¹⁴, far less populated than the nitrosamine and the isomeric diazohydroxide 9.

All these effects should be present in six-membered heterocyclic diazonium ions, too. However, up to now no stable nitrosarnine derived from a six-membered

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heterocyclic amine has been isolated. By acidification of pyridine-2- and -4-diazotates"' and **9-alkylpurinc-G-diazotate52** *(10)* solutions were prepared @robably containing the respective *anti*-diazohydroxides nitrosamines. The pH dependence of log *k* of their transformation into the corresponding diazonium ion (or the products derived from the latter) has a similar character as that of the substituted benzencdiazoliydroxides, and thc reaction is subject to general acid catalysis, too. **A** rather substantial dilf'erence is encountered with **pyridine-4-diazohydroxide,** which at $pH < 4$ becomes protonated to give the less reactive conjugated acid (11) .

V. REACTIONS OF DIAZONIUM IONS WITH OTHER N UCLEOPHILES

Arenediazonium ions react with a number of nucleophiles to give diazo compounds'. **G3-i5.** The free electron pair for the bond formation can be supplied by a carbon, oxygen, nitrogen, sulphur or phosphorus atom of the nucleophile. In the first case azo compounds are formed which are usually very stable, although the reaction with CN- ions (forming arenediazocyanides) is reversible. The other adducts are called diazo compounds, however some of them (e.g. those containing $N-P$ bonds) are denoted as azo compounds.

Some diazo compounds are very unstable, e.g. diazoethers formed by reaction of diazonium ions with substituted phenolate ions; here it is questionable whether the nucleophile and the azo group arc connected by a covalent or an ionic bond. In other cases, the primarily formed diazo compound is rapidly transformed to other more stable compounds. Reaction of diazonium ions with azide ion N_3^- produces unstable diazoazides which, in a subsequent (as a rule rate-limiting) step, either decompose to give nitrogen and an aryl azide or isomerize to arylpentazoles which are also unstable⁵⁶. Some diazo compounds are formed in other ways than from the reaction of arenediazonium ion with nucleophile.

The following discussion is limited to diazo compounds for which it was possible to determine the equilibrium constant of the reaction of at least some derivative of arenediazonium ion with the nucleophile.

A. Arenediazoethers

Reaction of diazonium ions with alkoxide ions is analogous with the first reaction step of that with hydroxyl ion. Very reactive heterocyclic diazonium salts, e.g. 3-phenyl-I **,2,4-thiadiazol-S-diazoniuni** fluoroborate, give high yields of diazoethers even on reaction with pure methanol⁵⁷. For preparation of substituted benzenediazoethers it is more convenient to alkylate silver diazotates with methyl iodide⁵⁸⁻⁶⁰; alkylation of a sodium or potassium salt produces the aryl derivative of N-methylnitrosamine¹³.

Reaction of 4-nitrobenzenediazonium ion with methoxide ion in methanol produces, in **a** rapid reversible step, the diazoether probably having the *cis* configuration⁶¹; it is transformed to the more stable *trans* isomer and to nitrobenzene in about equal amounts⁶² (Scheme 6). The isomerization rate constant is 7.23×10^{-3} sec⁻¹ at 0°C.

Rate constants of formation of the *syri* isomers were measured at 23 *"C* by the stopped-flow method⁶¹ for 4-nitro- and 4-cyano-benzenediazonium ions $(3.0 \times 10^8$ and 2.0×10^8 l mol⁻¹ scc⁻¹, respectively), and equilibrium constants were determined for 4-nitro- $(5.6 \times 10^7 \text{ J/mol})$, 4-cyano- $(8.6 \times 10^6 \text{ J/mol})$ and 3-chloro $(4.2 \times 10^5 \text{ J/mol})$ derivatives⁶¹. Both k_1 and K are by several orders of magnitude greater than the corresponding values of the rcaction with hydroxyl ion in watcr. The half-life of the decomposition of 4-nitrobenzencdiazoether to diazonium ion and methoxide ion is 130 niscc, i.e. two orders of magnitude greater than that for 4-nitrobcnzenediazohydroxide in water.

SCHEXII: *6*

6. *Arenediazoarnino Compounds (Triazenes)*

Diazonium ions give a number of compounds by reactions with nucleophiles containing an HN ζ group^{53, 55}. The most important and most studied are arenediazoamino compounds (triazenes) which are formed by reaction of diazonium ions with primary and secondary aliphatic and aromatic (or heterocyclic) amines HNR^iR^2 (R^1 is hydrogen or alkyl, R^2 is alkyl or aryl):

$$
ArN_{2}^{+} + HNR^{1}R^{2} \xrightarrow{k_{1}} ArN = N\bar{N}HR^{1}R^{2} \xrightarrow{\longrightarrow} ArN = NNR^{1}R^{2} + H^{+}
$$
 (32)

So far it has not been possible to prove *cis-trans* isomerism of these reaction products such as, for example, that of diazotates^{63, 61} Measurements of dipole moments indicate⁶³ that the substances measured are *trans* isomers.

The rcaction with aromatic aniines can result eithcr in a reversible reaction at the amino nitrogen to give a triazene or in an irrcversiblc coupling reaction at a ring carbon atom to give an azo compound. Coupling at carbon is negligible in thc case of aniline in neutral and slightly acidic media⁶⁵; it is of minor importance with alkylanilines⁶⁶, but it represents the main reaction in the case of more reactive aromatic amines such as substitutcd naphthylamincs. In more strongly acidic media the rcvcrse reaction of the triazcne becomes incrcasingly important and thc content of azo compounds in reaction products increases.

The reaction with primary amines involves a further complication because two tautomcric forms are produced, the second being split into compounds other than the original diazonium salt and amine (Scheme 7). If R is aryl, the triazenes are split predominantly into the less reactive diazonium salt and less basic amine 67 . If R is alkyl, the splitting produces the aromatic amine and alcohol⁶⁸, the amount of the diazonium salt being negligible⁶⁹.

The equilibrium constant for reaction (32) has not yet been determined in any case. An attcnipt 10 dctcrrnine the cquilibriurn concentration in **thc** rcaction with dicthylamine (where the reaction is not complicated by C-coupling or by decomposition to

products different from the starting substances) was made impossible by simultaneous decomposition of the diazonium salt⁷⁰.

Thc equilibrium constant can be calculated from the rate constant of formation of triazene and from that of the reverse reaction in those cases where two tautomeric forms of triazene are not formed.

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\nts different from the starting substances) was made impossible by simult
\nposition of the diazonium salt⁷⁰.
\nequilibrium constant can be calculated from the rate constant of forma
\nz and from that of the reverse reaction in those cases where two taut
\nof triazene are not formed.
\nArN=NNHR
$$
\overbrace{\hspace{2cm}}^{\text{H+-}} \text{ArNH}_2 + \text{RN}^+_{\text{H}} \text{ (R = Ary!)}
$$

\n
$$
\begin{bmatrix}\n\overbrace{\hspace{2cm}}^{\text{H+}} & \text{H+N}^+\\
\overbrace{\hspace{2cm}}^{\text{H+}} & \text{H+N
$$

The kinetic equations (33) and (34) are prcsumed for formation and decomposition of triazenes^{65, 66, ⁷¹; $[R^1R^2NH]$ represents the concentration of the non-protonated} amine. *KA* is the dissociation constant of thc protonated triazcne; its value has not yet been determined in any case.

$$
\vec{v} = k_1 [ArN_2^+] [R^i R^i NH]
$$
 (33)

$$
\overline{v} = k_{-1}[ArN = N\overline{N}HR'R^2] = k_{-1}[H^+] / K_A[ArN = N\overline{N}R^2]
$$

= $k_{\text{B}+}[H^+] [ArN = N\overline{R}^2]$ (34)

$$
k_{\text{H}+}[\text{H}^+][\text{ArN}=\text{N}\text{H}^!\text{R}^{\prime}]
$$
 (34)

$$
K = [ArN = NNR'R^2][H^+]/[ArN_r^+][R'R^2NH]
$$
 (35)

The rate constants of formation⁷² and reverse reaction⁷³ of triazenes were measured for the reaction of *meta-* and *para-* substituted benzencdiazonium ions with dimethylamine in water at 25 and 20 °C.

The observed ρ constants of the formation and the reverse reaction of triazenes are 3.47 and -3.8 , respectively. The resulting $\rho = 7.28$ for the equilibrium constant (equation 35) is only slightly higher than that for the reaction of diazonium ions with hydroxyl ion^{28, 29}. The calculated rate constant k_1 of the reaction of benzenediazonium ion with dimethylamine is 450 I mol⁻¹ sec⁻¹; the $k_{\text{H}_{+}}$ constant of the reverse reaction can only be roughly estimated to be $10-20$ mol⁻¹ sec⁻¹. The slope of the dependence of $\log k$ on pH in the measured pH range 1-3 was (instead of the expected -1 , according to equation (34), provided that only minute amounts of the triazene are protonated) only about -0.75 . The calculated *K* value is about 30. For the sake of comparison with the equilibrium constant of the reaction of benzenediazonium ion with hydroxyl ion, which is equal to 2.0×10^{12} /mol² (Reference 26) the above value must be divided by the ionic product of water K_w (which corresponds to a change of equation (32) in thc scnse that **thc** protonated triazenc loses its proton by reaction with hydroxyl ion). The resulting value, about 3×10^{15} $\frac{12}{\text{mol}^2}$ is about 11 orders of magnitude greater than that for the formation of diazotate. This difference is due, first of all, to the far greater acidity of the protonated triazene as compared with benzenediazohydroxide. Obviously here K_A is close to unity whereas it is 10^{-8} for the diazohydroxide³².

The above-mentioned **valucs,** cspecially those for tlic reverse rcaction and for the equilibrium constant, must be accepted with rescrve. It cannot be excluded that isomerizationt takes place side by side with the reaction and that the rate-limiting step changes with pH.

t Ritchie⁷⁰ studied the reaction kinetics of 4-chlorobenzenediazonium ion with diethylamine and observcd **a** slowcr subsequcnt reaction independent of diethylamine concentration.

C. Arenediazosulphonates

The formation of diazosulphonates was described in 1869⁷⁴ and has been much studied because of its technical importance. The rate of formation of diazosulphonate is directly proportional to the concentration of arenediazonium ion and SO_3^2 ; the diazonium ion does not react with HSO_3^{-40} , ⁷⁵. In the first, very rapid, step a product is formed which was called $syn\text{-}sub$ honate by Hantzsch⁷⁶, and it is transformed slowly into the other isomer (anti):

$$
ArN_{2}^{+} + SO_{3}^{2-} \xrightarrow{\begin{array}{c} k, \\ k-1 \end{array}} ArN_{\otimes} \xrightarrow[\begin{array}{c} 1 \\ SO_{3}^{-} \end{array}} Ar-N_{\otimes} N-SO_{3}^{-} \tag{36}
$$

The log K values of formation of syn-sulphonates (Table 5) correlate with the σ constants⁴⁰; the ρ value is 5.5. The syn-sulphonates are very reactive and are rapidly decomposed into the starting substances. Half-life of the reverse reaction of 4-chloro derivative⁷⁰ is 15 msec. On the contrary, the *anti* isomers are extraordinarily stable. The values given for the 4-methoxy derivative are 10⁷ for $K_3 = [anti]/[syn]$ and 10^{-1} sec⁻¹ for the rate constant of the *anti* to *syn* transformation⁷⁷.

TABLE 5. Rate and equilibrium constants of the reaction of substituted benzenediazonium ions with sulphite ions

Substituent	K^a (l/mol)	k_1 ^b (1 mol ⁻¹ sec ⁻¹) k_3 ^{<i>a</i>, <i>c</i>} (sec ⁻¹)	
$4-NO2$	3.2×10^8	4.5×10^8	1.8×10^{-3}
4 -CN	1.1×10^8	4.3×10^{8}	
4-CI	2.7×10^5	2.4×10^{7}	0.5×10^{-3}
н	3.6×10^{4}		1.9×10^{-3}

^a Reference 40, ionic strength 1 M.

 b Reference 70, ionic strength $10⁻³$ M.</sup>

 c At 0 $°C$.

D. Arenediazosulphones

Arenediazosulphones are formed by reaction of diazonium salts with anions of sulphinic acids⁷⁸; usually the reaction is carried out in slightly acidic aqueous medium⁷⁹. Only one isomer was proved in the reaction products^{79, 80}. The dipole moments found⁸¹ suggest *trans* structure of the products. The presence of the single isomer is explained⁸² by lowering of the rotation barrier about the $N=N$ bond by contribution of expanded octet structures such as 12b, c:

Since the water-solubility of the products is very low, the reaction kinetics of substituted benzenediazonium ions with benzenesulphinic acid were measured in

 m ethanol⁸². In contrast to the reactions of diazonium salis with most other nucleophiles, the diaznsulphone is formed so slowly that the reaction can be followed by usual spectrophotometric methods. The dependence of the logarithms of the rate, and equilibrium constants on thc *u* constants is expressed by equations **(38)** to (40):

$$
\log K = 3.76\sigma + 5.26\tag{38}
$$

$$
\log k_1 = 2.40\sigma + 2.52\tag{39}
$$

$$
\log k_{-1} = -1.36\sigma - 2.74
$$
 (40)

In spite of anions of sulphinic acids being far weakcr nucleophiles than the sulphite anion, the *k-,* values are more than four orders of magnitude lower than those for the diazosulphonates. Obviously this is caused by thc reverse reactions of diazosulphonates involving the more stable *trans* isomer. This fact also influences the values of the equilibrium constants. The small ρ observed for the diazosulphone equilibrium is explained⁸² by the contribution of structures such as 12b where a partial charge remains localized on the nitrogen.

E. Arenediazothioethers

Arenediazothiocthers are formed very easily and rapidly by reaction of diazonium salts with thiols. They are very unstable and decompose, evcn explosively in some cases, to give nitrogen and the corresponding sulphide⁸³. In contrast to aromatic hydroxy compounds the reactions with thiophcnols, thionaphthols and even dithioresorcinol do not lead to coupling in the ring⁸⁴.

TABLE 6. Rate and cquilibrium constants for the rcaction

x', XC,H,N: + C,H,S- . syn-XC,H,N=NSC,H, anti-XC,H,N=NSC,H, in nicthanol at **23** *"C*

The reaction of bcnzenediazoniuni ions with thiophcnoxidc ion was followcd by **the** stopped-flow mcthod in methanolic buffers"'. At relatively high concentrations of thiophenoxide ion the reaction of diazonium salts having a positive σ value for the substituent (4-chloro- to 4-nitro-) proceeds in two clearly separated steps. Thc initial rapid reaction is probably the formation of a syn-diazothioether, and the slower reaction is *syn-anti* isomerization. With the unsubstituted, 4-methyl and 4-methoxy derivatives the reactions followed first-ordcr kinetics over the entire course of the reaction. The formation of syn-diazothioether was probably the rate-limiting step followed by a rapid $syn-anti$ isomerization. Due to the isomerization, reliable values for the equilibrium constant could be obtained for the 4-nitro and 4-cyano derivativcs only (Table 6):

$$
ArN_{2}^{+}+C_{6}H_{5}S^{-} \xrightarrow[k_{-1}]{k_{1}} \text{syn-ArN=NSC}_{6}H_{5} \xrightarrow[k_{-1}]{k_{1}} \text{anti-ArN=NSC}_{6}H_{5} \qquad (41)
$$

The k_1 values approach the values for diffusion-controlled reactions; the influence of substitucnts is far smaller than in the other reactions studicd and decreases with increasing value of σ constant. Equilibrium constants are about three orders of magnitude higher than those for reactions with CH₃O⁻ ion.

F. Arenediazocyanides

Substituted benzenediazonium ions rcact vcry rapidly and reversibly with cyanide ions to form syn-diazocyanides⁸⁵ which are slowly transformed (half-life of one to several hours at room temperature) into *anti*-diazocyanides⁸⁶. Hantzsch⁸⁵ suggested cis and *trnns* configuration for tlie **sj'n** and *otiti* isomers, rcspcctively, which agrees

with the measured dipole moments and i.r. spectra of the two forms^{18, 19}:
\n
$$
ArN_2^+ + CN^ \xrightarrow[k]{k_1 \atop k_2}
$$
 $Ar-N_{N} \xrightarrow[N]{Ar-N_{N} - CN}$ (42)
\n $\xrightarrow[\text{CN}]{}$

Equilibrium constants **of** formation of tlie *syti* isonicr wcre measured by the tise of conventional spectrophotometric methods⁸⁶ and by the stopped-flow technique²⁸. The latter method affords more accurate results, because the determination of the equilibrium constant is not disturbcd by slowcr subscquent reactions". The effect of the ring substituents on the equilibrium and rate constants, determined also by the stopped-flow method, was quantitatively expressed by the Hanimett equation:

$$
\log K = 3.53\sigma + 1.82\tag{43}
$$

$$
\log k_1 = 2.31\sigma + 2.32\tag{44}
$$

The relatively small ρ value 3.53 is (analogously to diazosulphonates) due to the partial positive charge on the nitrogen atom of the diazo group caused by the strong $- M$ effect of the CN group (mesomeric structure $Ar - N - N = C = N$). The strong $- M$ $-M$ effect of the CN group is obviously also responsible for the fact that electronwithdrawing substituents slow down the isomerization⁸⁷.

Substituent	K (l/mol)	k_1 (1 mol ⁻¹ sec ⁻¹)	k_{-1} (sec ⁻¹)
4-NO.	6.0×10^{4}	1.7×10^{4}	0.28
$4 - CN$	1.4×10^{4}	7.4×10^3	0.53
	(2.2×10^7) ^a	(2.2×10^6) ^a	(0.1) ^a
$4-Br$	3.7×10^{2}	6.5×10^{2}	1.75
	$(3.2 \times 10^9)^{6}$	$(1.6 \times 10^{8})^{b}$	$(0.05)^{b}$
$4-C1$	4.5×10^{2}	6.8×10^{2}	1.5
	$(2.5 \times 10^9)^{b}$	$(1.5 \times 10^8)^{b}$	(0.06) ^b
4 -CH ₂	1.5×10	9.0×10	$6-0$
	$(1.4 \times 10^8)^{b}$	$(3.8 \times 10^7)^{h}$	(0.27) ^b

TABLE 7. Rate and equilibrium constants for the reaction of substituted benzenediazonium ions with cyanide ion in water at 23 °C²⁸

^a In methanol⁶¹.

^b In dimethyl sulphoxide⁵⁸.

The reaction of some substituted benzenediazonium ions with cyanide ion was measured in water, in methanol⁶¹ and dimethyl sulphoxide^{ss}. The equilibrium constants arc three aiid sxcn orders of mngnitudc grcater **in** iiictlianol **and** dimethyl sulphoxide, rcspectivcly (Table 7). The influence of tlie medium is dccisivc in the case of *k,,* whereas the reverse reaction showed a relatively lower decrease. **A** large increase in the equilibrium constant accompanying a change from a protic to polar aprotic solvent is typical for reactions with small, negatively-charged, nucleophiles⁸⁹.

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CHAPTER **3**

Structural chemistry

S. **SORRISO**

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1. INTRODUCTION

A. General

It is normal in a chapter on stiuctural chemistry to discuss bond distances and angles in molecules containing the characteristic group, and to explain the modifications undergone, or induced, by the latter. In the case of diazonium and diazo **groups** this is not sufficient because the structural data are often lacking. This makes necessary recourse to other kinds of results to discuss at least the possible configuration.

The configuration of diazo compounds has been the object of much work since their discovery about a hundred years ago'. Nevertheless, many problems still remain not completely solvcd. For example, due to thc instability of the molecules examined, the type of isomerism present in some covalent diazo compounds and the reasons for its existence are still objects of discussion.

Thc book written by Zollingcr' in 1961 covers the configurational aspects of the present groups. Therefore, with very few exceptions, work carried out before this year will not be covered since it has been revicwed at length in that book. **A** historical introduction to the configurational problems in the diazonium and diazo groups may be found in the book by Saunders".

Strictly structural results are relatively recent and they have not been reviewed previously. These, which have been obtained exclusively from X-ray measurements, **will** be treated cxtensivcly in the present chapter.

B. Usefulness and Errors in an X-Ray Examination

X-Ray data in chemistry give information on the molecular configuration and thus they are very helpful in indicating uncquivocally the type of isomerism present. Besides, the refinement of the structure by means of modern computational methods often allows information to be obtained also on the hybridization of the atoms and on the bond order.

The reliability of an X-ray crystallographic determination is usually indicated by the R factor, the residual error index or discrepancy index, and by σ , the standard deviation 3.4 .

The R factor is a function of the observed (F_0) and calculated (F_c) structure factors:

 $R = \sum (|F_0| - |F_c|)/\sum |F_0|$

H does not have **a** prccisc theorctical meaning and depends greatly on temperature factors, the complexity of the structure and the amount of data collected. The lower the value of R, the lower is the discrepancy between observed and calculated structure factors. Consequently, it is also useful to the crystallographer for following the progress of structure refinement. Initial *R* values of 0.50 and final ones of 0.10 are normal1. At present, with niodcrn computer techniques, *R* values between *0.08* and 0.03 are being reported for the most reliably determined structures⁴. The *R* factor is a function of the intensities. Therefore, when heavy atoms are present, they may dominate the structure factor calculation. Consequently, the bond lengths involving hcavy atoms will be more precisely determined than those involving light ones.

The standard deviation in bond lengths and angles is generally 0.002 **A** and $0.2-1.5^\circ$, respectively, in more recent work. However for the former, given the possibility of systematic errors, it is reasonable to assign a value of 0.01 Å , even if account is takcn of thermal motion (corrected value). For lighter elements, the standard deviations in bond distances are usually higher. In fact, since X-rays are diffracted by electrons, they give statistically the position of the centroid of negative charge rather than that of the nucleus. This means that while for hcavy atoms the centroid of electron cnargc and that of the nucleus are statistically coincident, in light atoms there is a certain difference. For example, for hydrogen the difference⁵ between the two is ~ 0.1 Å, which for lithium decreases to ~ 0.01 Å.

Caution is necessary in interpreting the crystallographic data in ternis of the electronic structure and bond order. **In** fact, the standard deviation in bond length niay be subject to different errors, such as the inadequacy of the atom-form factors, of thc crystal model and of the temperature factors. Tlicse may become so important as sometimes to render the experimental data suspect, even if σ values are excellent.

II. BlAZONlUM SALTS

A. Outer Diazoniurn Salts

I. Structural data

a. *Benzenediazonium chloride*, PhN₂Cl (1). This molecule has been examined in two studies. In the first study^{$6, 7$} the measurements were performed at room temperature. The compound is completely ionized; $R = 0.06$; standard deviations 0.006-0.010 Å and $0.6-0.7$ °. The second study⁸ was performed at -160 °C; R not specified; standard deviations 0.002 *8,* and *0.2".* Each diazoniuni group' is surrounded by four CI- ions situated in a plane perpendicular to the $N-N$ axis. Two of the CIions lie at **3,184** and the other two at **3.51** 1 **8,** to this axis. The fornicr chlorine ions lie 3.225 and 3.237 Å away from $N_{(2)}$ and $N_{(1)}$, respectively, and the others at 3.548 and 3.559 Å. The separation between the latter and the $H_{(2)}$ atom (see Figure 1) is only 2.5 *8,.* Because this distance is half an Angstrom less than the sum of the van der Waals' radii, a strong interaction must be present between these two atonis'.

b. Benzenediazonium tribromide, PhN₂Br₃</sub> (2). Completely ionized; $R_{ok1} = 0.08$ *,* $R_{hko} = 0.073$ and $R_{hot} = 0.073$ (where *h*, *k* and *l* are used to indicate the plane referred to); estimated standard deviations 0.04-0.05 **8,** and **3-4".** The spatial arrangement of tribromide ions about the diazo cation is shown in Figure **2.** The $C_{(1)}-N-N$ arrangement is linear. The shortest distances between the bromine atoms and $N_{(1)}$ and $N_{(2)}$ are 3.46 and 3.31 Å, respectively.

FIGURE 1. The layer of chloride and benzenediazonium ions. Reprinted, with permission, from Romming, *Acrn Client. Scad.,* **17, 1444 (1963).**

	Reference	Reference 8		Reference	Reference 8
$N_{(1)} - N_{(2)}$ $N_{(1)}$ – $C_{(1)}$ $C_{(1)} - C_{(2)}$ $C_{(2)} - C_{(3)}$ $C_{(3)}$ -- $C_{(4)}$	1.097 Å 1-385 Å 1.374 Å 1.383 Å 1.376 Å	1.093 Å 1.410 Å 1.396 Å 1.392 Å 1.395 Å	$C_{(1)} - N_{(1)} - N_{(2)}$ $C_{(1)} - C_{(2)} - C_{(3)}$ $C_{(2)} - C_{(3)} - C_{(4)}$ $C_{(3)} - C_{(4)} - C_{(5)}$ $C_{(6)} - C_{(1)} - C_{(2)}$	180° 117.6° 119.8° 121.7° 124.8°	115.9° 120.6° $121 \cdot 1^{\circ}$ 126·1°
		ш п Br.	(1) $\mathfrak{tBr}_{(n)}$ B٢ B٢ شام Br. Br. π		

Uncorrected bond lengths and angles

FIGURE 2.Arrangement of tribrornide ions about a benzenediazonium ion. The lines $Br_{(1)}^{\text{I}}-Br_{(1)}^{\text{II}}$ and $Br_{(3)}^{\text{III}}-Br_{(3)}^{\text{IV}}$ form angles of about 85° and 35° with the plane of the benzene ring, respectively. Reprinted, with permission, from Andresen and Rømming *Ac~n Cheni. Scad.,* **16, 1882 (1962).**

Uncorrected values		
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c. p-N,N-Dimethylaminobenzenediazonium tetrachlorozinc(ii) $(p\text{-Me}_2\text{NC}_6\text{H}_4\text{N}_2)_2$. $ZnCl₁$ ^{10,11} (3). Ionic structure; $R = 0.141$; only the atomic coordinates have been reported; uncertainty in atomic coordinates has not been specified. From these data we have extracted the following bond distances and angles:

The chlorine ions form a tetrahedron about the Zn atom (Cl- $2n$ -Cl 106-110² and Zn-Cl 2.28-2.35 Å). The shortest distances between $Cl_{(1)}$ and $N_{(1)}$ and $N_{(2)}$ are 3.672 and 3.435 Å, respectively, and those between $Cl_{(2)}$ and $N_{(1)}$ and $N_{(2)}$ 3.701 and 3.420 Å.

d. o-Methoxybenzenediazonium tetrachloroiron(III), o-MeOC₆H₄N₂·FeCl₄¹² (4). Ionic structure. $R = 0.13$; standard deviation 0.01–0.02 Å. The chlorine ions form a tetrahedron around the Fe atom (Cl-Fe-Cl 107.5-113.2° and Fe-Cl 2.165- 2.230 Å). All the atoms C, N and O are coplanar. The shortest distances between the chlorine and the two nitrogen atoms are Cl- $\mathbb{N}_{(1)}$ 3.51 and Cl- $\mathbb{N}_{(2)}$ 3.36 Å.

e. p-Benzenebisdiazonium tetrachlorozinc(II), $p-\text{N}_2\text{C}_6\text{H}_4\text{N}_2$ ·ZnCl₄¹³ (5). Ionic structure; $R = 0.065$; standard deviations 0.012-0.017 Å and 0.8-1.2°. The

p-benzenebisdiazonium ion is, within the accuracy of the measurements, planar and the $N-N-C \cdots C-N-N$ arrangement is linear. The diazonium group is surrounded by four chloride ions situated in a plane perpendicular to the $N-N$ axis. The shortest distances between the chlorine and nitrogen are $Cl-N_{(1)}$ 3.26 and $Cl-N_{(2)}$ 3.43 Å. These values are almost equal to the van der Waals' separation between these two atoms $(3.3 A)$. Thus it is not possible to draw any conclusion as to the charge distribution in the diazonium group from the geometrical parameters.

f. 1:1 Complex benzenediazonium chloride-acetic acid, PhN_2Cl MeCO₂H¹⁴ (6). $R = 0.075$; standard deviations 0.006-0.016 Å and 0.5-1.2°. The crystals are built up of benzenediazonium ions, chloride ions and acetic acid molecules. The distance between the oxygen and chloride ions (3.01 Å) may indicate the presence of a hydrogen bond^{14, 15}, O₍₂₎ ... H₍₁₎ ... Cl⁻.

2. Discussion

The results reported in Section II.A.1 leave no doubt as to the structure of the diazonium group. It may be seen that, with the exception of the compound 5, in all the compounds examined the $N-N$ distance lies in the range 1.09–1.11 Å, virtually the same as that observed in dinitrogen $(1.097 \text{ Å})^{16}$, which indicates that in the ground state the extreme form 7 clearly predominates over 8 and 9. The presence of

an essentially triple $N-N$ bond has also been arrived at from the value of the frequency of the N-N stretching vibration band (\sim 2290 cm⁻¹)¹⁷⁻²¹ which is between the Raman frequency of dinitrogen $(2330 \text{ cm}^{-1})^{22}$ and the stretching vibration frequency of the C=N group (\sim 2255 cm⁻¹ for PhCN)²³. Also in agreement with this

i See Note Added in Proof, p. 135.

suggcstion are thc theoretical calculations carried out, using different methods, by Shuster and Polansky²⁴ and, more extensively, by Bochvar and coworkers²⁵ and by Sukigara and Kikuchi²⁶⁻²⁸.

'The fact that the N-N bond **is** virtually a triple bond suggests that the positive charge of the benzenediazonium ion, according to the valence theory, is localized mainly on the nitrogen atom which is bonded to the phenyl ring. Several authors^{11, 20} do not agree with this conclusion. They base their opinions on the fact that for most of the diazonium salts above, the shortest distance $Hal - N_{(2)}$ is always shorter than, or of the same order as, the distance $\text{Ha} - \text{N}_{(1)}$, which suggests that the positive charge of the diazo cation is slightly more localized on the terminal nitrogen or, at least, equally distributed over both nitrogen atoms. We feel that this shorter $\text{Hal}-\text{N}_{(2)}$ distance probably does not depend cxclusively on the electrostatic interaction between the nitrogen and the halogen ion, but also on that between the latter and the hydrogen *ortho* to the phenyl ring²⁹. This suggestion might be supported by the fact that in benzenediazonium chloride⁷ (Figure 1) the distance between the $H_{(2)}$ atom and CI- ion is *2.5* A, about *0.5* **8,** smaller than the sum of the van der Waals' radii.

X-ray data for the complexes o -MeOC₆H₄N₂.FeCl₄ and $[p$ -Me₂NC₆H₄N₂]₂.ZnCl₄, together with results of **MO** calculations, havc been used by some authors?5 to suggest the presence of a 'bcnzenoid zone' and of a 'quinoid' one in niethoxy- and amino-diazo cations. **We** shall now scc how far this suggestion is supported by the experimental results.

 o -MeOC₆H₄N₂·FeCl₄: The bond lengths in the diazonium group do not indicate the presence of conjugation with the rest of the molecule. In fact, the $N-N$ bond $(1.11 \text{ Å})^{13}$ is virtually the same as that in benzenediazonium chloride ($\sim 1.09-$ **1.10 Å). In addition, the C₍₁₎-N₍₁₎** bond (1.48 Å) is noticeably greater compared to the same bond in **1** (1.385-1.410A), whercas it should, of course, be shorter according to the canonical forms **7–9**. In agreement with these results, the methoxy group does not support the presence of conjugation with the diazoniuni group. In fact, the O-C_{sp}, distance (1.36 Å) is of the same order as that in similar molecules, e.g. 1,4-dimethoxybenzene (1.36 Å)³⁰ and p, p' -dimethoxybenzophenone (1.37 Å)³¹.

 $[p-Me_2NC_6H_4N_2]_2$. ZnCl₄: For this molecule the fact that the N-N distance is sensibly longer (1.18 Å) than in benzenediazonium chloride may be an indication that there is an increase in the contribution of the extreme forms **8** compared to that in the parent cornpound **1,** i.c. strong conjugation between the substituent and the diazonium group. In agreement with this, the $N_{(3)}-C_{\rm sp}$: distance (1.35 Å) is shorter than that observed in 2,5-dichloroaniline $(1.40 \text{ Å})^{32}$ and p-toluidine $(1.43 \text{ Å})^{33}$, and is similar to that found in p-nitroaniline $(1.37 \text{ Å})^{34}$ and N,N-dimethyl-p-nitroaniline (1.35 **A)3s,** for which conjugation between the nitro and aniino groups is accepted. These points, important for defining the structure of this compound, unfortunately are doubtful because the $C_{(1)}-C_{(2)}$ and $C_{(6)}-C_{(1)}$ distances are very different (1.53 and 1.36 A, respectively) whereas they should be cqual (within experimental error, which is less than 0.1 Å)^{10, 11}. Furthermore, the present results are uncertain since the authors^{10, 11} do not report the exact standard deviations in the atomic coordinates. To conclude, the above discussion on the coniplexcs **3** and **4** shows that the present structural parameters cannot be used to prove thc existencc of a 'quinoid zone' about the arnino group and a 'benzenoid zone' involving the diazonium group²⁵. On the other hand, in this case, the theoretical calculations²⁵ cannot be invoked to support this suggestion since for the benzenediazonium cation they are very sensitive to the input parameters used. It seems obvious that, given the utility of X-ray measurements, amine complexes with different anions should be examincd before definite conclusions can be drawn.
Further information on thc electronic structure of diazo cations substituted in the phenyl ring with electron-donor groups has been obtained using other techniques.

In different solvents, good correlations have been found^{17, 19, 21, 36-38} between the frequency of the stretching vibration band (ν_{NN}) , its integrated band intensity, the N-N bond order and the Hammett σ_p values³⁹ for diazonium cations p -XC₆H₄N₂⁺ (X = H, Me, Cl, Br, NO₂, SO₂NH₂, I, F, OMe, CO₂Et, OH, O⁻ or SO₃). For p-OH-, p-OMe- and p-Me₂N-derivatives σ^+ values have been used instead of σ_p ones. This procedure led to the result that the points for the first two derivatives were on a straight line but that for the amine compound was not¹⁷. This indicates further polar conjugation between the substitucnt and the diazonium group, which is much greater in the amine derivative.

Many workers have shown interest in the electronic structure of the latter compounds. In the spectral zone characteristic of quinones some similarity has been observed⁴⁰ between the u.v. spectra of p -amino-, p -methylamino- and p -dimethylamino-benzenediazonium cations and that of diphcnylquinoniethane, which has a quinoid structure **(10).** This led to the suggestion that these molecules have an

analogous structure **(1 1).** There are many possible criticisms of this suggestion and it certainly appears that a completely quinoid structure for all three of the above derivatives cannot explain the other experimental results. In fact, if this were indeed the structure, there should be no appreciable variation in the electronic configuration of the diazo cation on introducing two methyl groups in place of the amine hydrogens. The opposite is the case, because on passing from the amino to the dimethylamino derivative, v_{NN} goes from 2183 to 2166 cm⁻¹ and the band at 357 goes to 382 nm⁴¹, indicating an increase in quinoid character. Also, the N-N integrated band intensities (12.6 and $13.5 \text{ mol}^{-1} \text{ cm}^{-2} \times 10^4$ respectively, for $p\text{-}NH_2\text{-}G_6\text{H}_4\text{N}_2\text{Cl}$ and $p\text{-}Me_2\text{N}C_6\text{H}_4\text{N}_2\text{Cl}$ ¹⁸ show the same difference between these two compounds. Very probably the present amino derivatives also arise from a resonance hybrid between the canonical forms **7** and **8.** However, there is a noticeable contribution from **8** which, instead, in these compounds becomes **11.** This resonance form would account not only for the frequency and integrated intensity of the $N-N$ stretching vibration band but also for the fact that for dimethylamino derivative even the σ^+ value does not lie on the ν_{NN} versus σ_p plot. Form 11, further, would explain the much higher stability and the colour of these compounds as compared with the other diazonium salts of simple anions, the great majority of which are colourless⁴⁰.

3. Effect of the anion on the structure of diazonium cations

Much information on this niatter has been obtained from infrared measurements. These have been confined to the study of the $N-N$ stretching vibration band in complex molecules and to comparison with spectra of diazonium salts of simple anions. It is not appropriate to underline the various contributions, which are very numerous⁴²⁻⁵⁹, in a chapter on structural chemistry. However, a concise discussion of the results available may be useful.

Independently of the physical state, all the diazonium salts studied to date show one absorption band in the N=N region. Some authors⁵⁰ report two N-N bands in this region only in p-dialkylamino derivatives. Of these bands the more intense, at \sim 2170 cm⁻¹, is assigned to the N--N stretching vibration and the other, at \sim 2250 cm⁻¹, appears to be the first overtone of the band at \sim 1100 cm⁻¹, assigned to the C-N vibrations of the alkylamino groups. Instead, other authors^{45-48, 56} have also observed a complex band, usually with two peaks, for many diazonium salts, indcpendently *of* substitution at the phenyl ring. The rcason for the appearance of a complex N-N band is not clear. In most cases it is probably due to solidstate lattice effects, since there is only a single peak on regrinding the mull⁵⁰. To interpret the presence of complex $N-N$ bands as caused by weak bonding between the diazoniuni group and the anion secms hazardous, although it cannot entirely be excluded. On the other hand, X-ray data on the complex diazonium salts examined above when compared to those for PhN,CI do not indicate the presence of this bond.

Nuttall and coworkerS"O observed that the salts *of* the very strong complex fluoro acids show a v_{NN} value higher than that in other complex diazonium salts. For these authors this fact indicates that the former are completely ionic, while the latter have **il** weak covalent bond between the empty orbital *of* the diazonium group and the halogen lone pair, or between the π orbital of the diazonium group and the metal atom of the anion. The first suggestion might be preferred⁵⁰ for the following reasons. **In** complex diazoniuni salts with p-dimethylarnino substituted cations, in which the diazonium group is less positive than in other derivatives, the v_{NN} value does not show the clcar diffcrcnce between the fluoro derivatives and thc others.

To conclude. the extensive spectroscopic results together with X-ray data suggest that the electronic structure of the diazo cation in diazoniuni salts of complex anions is substantially the same **as** that in the analogous simple salts.

5. Inner Diazoniurn **Salts**

1. 2-Diazonium-4-phenolsulphonate monohydrate

 $N_2C_6H_3SO_3$ $OH \cdot H_2O$ ⁶⁰ (12): $R = 0.128$; standard deviation 0.01-0.02 Å. The compound has **a** zwitterionic structurc; the water molecules connect t hcse zwittcrions by means of hydrogen bonding.

Lincorracted

2. p-Benzenediazonium sulphonate

 $N_2C_6H_4SO_3$ (13): The structure of this molecule has been determined twice. In the first study⁶¹, $R = 0.13$, standard deviations were not reported. According to the authors, the dimensions of the phenyl ring indicate that the molecule is quinoid in character. The second study²⁹ has, $R = 0.052$, standard deviations 0.002-0.004 Å and $0.1-0.3^{\circ}$. The C-H bond length is $0.91-0.99$ Å²⁹.

There has been discussion as to whether p -benzenediazonium sulphonate has a quinoid structure (14) or a benzenoid one, of the predominant extreme form (15).

The data available suggest that this molecule, and also 12, has a zwitterionic structure although, according to some authors⁴¹, a small contribution from canonical form 14 cannot be entirely excluded from the spectroscopic results. In fact, the $N-N$ stretching band frequency lies at 2296 cm⁻¹ (in petroleum ether)²⁰ in benzenediazonium chloride and at 2284 cm^{-1} in p-benzenediazonium sulphonate. Instead, in p-diazo oxide, $OC_6H_1N_2$, for which a prevalent quinoid structure is now believed to exist, ν_{NN} (in CHCl₃) lies at 2088 cm⁻¹. On the other hand, the structural data do not appear useful in this case since a small participation of canonical form 14 could cause a variation in the bond lengths of the same order as that of the experimental errors.

3. Structural chemistry **105**

In agreement with a zwitterionic structure, the $N_{(1)}-N_{(2)}$ distance (1.08 and 1.09 **A,** respectively, in **2-diazonium-4-phenolsulphonate** monohydrate and in p -benzenediazonium sulphonate²⁹) is of the same order as that found in most diazonium salts $(1.10-1.11 \text{ Å})^{7-9}$, ¹¹. In turn, the $C_{(1)}-N_{(1)}$ bond length is 1.40-**1.41 A** and compares very favourably with the values observed for unsubstitutcd diazonium salts (1.39-1.42 **A).** Finally, these results are confirmed by the value of the $C-S$ bond length, which is very near to that found in similar molecules^{62, 63}.

111. D!AZOTATES

A. isomerism

Addition of a strong base to an aqueous solution of a diazonium salt leads to rather complex transpositions, which have been of interest for a long time'. The reaction scheme which is presently agreed on^{1, 64-72} is the following: trong base to an aqueous solution of a diazo

inspositions, which have been of interest for

hich is presently agreed on^{1, 64-72} is the followin
 $ArN_x + OH = \frac{slow}{\sqrt{300}}$ $syn-ArN_xOH$
 $syn-ArN_xOH + OH = \frac{fast}{\sqrt{300}}$ $syn-ArN_xO = +H_xO$
 $syn-A$

$$
ArN_{2}^{+} + OH^{-} \xrightarrow{\text{slow}} syn-ArN_{2}OH
$$
 (1)
\n
$$
rN_{2}OH + OH^{-} \xrightarrow{\text{fast}} syn-ArN_{2}O^{-} + H_{2}O
$$
 (2)
\n
$$
syn-ArN_{2}O^{-} \xrightarrow{\text{slow}} anti-ArN_{2}O^{-}
$$
 (3)
\n
$$
ArN_{2}^{+} + OH^{-} \xrightarrow{\text{anti-ArN}_{2}OH}
$$
 (4)
\n
$$
anti-ArN_{2}OH \xrightarrow{\text{anti-ArN}_{2}O^{-} + H^{+}}
$$
 (5)

$$
syn-ArN_{2}OH+OH^{-} \xrightarrow{\text{fast}} syn-ArN_{2}O^{-} + H_{2}O
$$
 (2)
\n
$$
syn-ArN_{2}O^{-} \xrightarrow{\text{slow}} anti-ArN_{2}O^{-}
$$
 (3)
\n
$$
ArN_{2}^{+} + OH^{-} \xrightarrow{\text{anti-ArN}_{2}OH}
$$
 (4)
\n
$$
col(Ar)N_{2}OH \xrightarrow{\text{outi}ArN_{2}OH}
$$
 (5)

$$
syn-ArN_2O^- \xrightarrow{\text{slow}} anti-ArN_2O^-
$$
 (3)

$$
ArN_2^+ + OH^- \xrightarrow{\longrightarrow} anti-ArN_2OH \tag{4}
$$

$$
anti-ArN2OH \xrightarrow{2} anti-ArN2O^+ + H^+
$$
 (5)

There is kinetic evidence only for the participation of syn-diazohydroxide in the reaction^{1, 64, 71, while *anti*-diazohydroxide has been identified spectroscopically but} not isolated in the pure state^{1, 64}.

The existence of two diazotate ions in aqueous solution at appropriate pH, which is now accepted by most workers¹, was in doubt until very recently^{72, 73}. Their existence now seems definitely established even for the *syn*-isomer of p -nitrophenyldiazotate, which could not be detected previously because of its high lability⁷⁴.

The problem of the type of isomerism present in diazotate ions has been widely studied'. However, since the techniques available were unsuitable, the results obtained were never conclusive. Most workers^{1, 20} agreed that the isomerism present in diazotates is geometrical isomerism (equation 6). However, the evidence put forward to support this claim was essentially based on the analogy with similar compounds or on kinetic results. Direct confirmation has been obtained only recently, via the three-dimensional X-ray structural analysis of potassium-syn-
methyl-diazotate⁷⁵. The results obtained, which are reported in Section III.B, may
be extended, so far as this type of isomerism is concern methyl-diazotate⁷⁵. The results obtained, which are reported in Section III.B, may be extended, so far as this type of isomerism is concerned, to the corresponding phenyl derivative (PhN₂OK), since Müller and coworkers⁷⁶ found that both com-

pounds are prepared in the same way and have similar properties.\n
$$
0^-
$$
\n
$$
N=N
$$
\n
$$
A_r
$$
\n
$$
C_i
$$
\n
$$
(6)
$$
\n
$$
A_r
$$
\n
$$
C_i
$$
\n
$$
(7)
$$
\n
$$
C_i
$$
\n
$$
(8)
$$
\n
$$
2
$$
\n
$$
2
$$
\n
$$
C_i
$$
\n
$$
(9)
$$
\n
$$
2
$$
\n
$$
2
$$
\n
$$
(10)
$$
\n
$$
2
$$
\n<math display="block</p>

The problem of the configuration of the *syn* and *anti* isomers is now largely clarified and will not, therefore, be treated at length.

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B. Structure of Potassium-syn-methyl-diazotate

 $MeN₂O-K$ (16): This is the only diazotate for which structural parameters are available; $R = 0.084$; standard deviations 0.008-0.01 Å and 0.5°. The C-N and N-N bond lengths in **16,** (1.477 and 1.246 **A,** respectively) arc of the same order as those observed in azomethane $(1.474$ and 1.244 Å, respectively)⁷⁷. By contrast, the N-O distance (1.306 Å) is shorter than that observed usually $(1.39-1.41)^{5}$.

These results may be explained by invoking the participation of the extreme forins **17a** and **17b** to the resonance hybrid, with the prcdorninance of the former. The data reported for **16** exclude the altcrnative theory on the configuration of diazotates^{1, 78-s1}, according to which the stable isomer *(anti)* has the same configuration as that put forward by Hantzsch, whilst the labile one *(syn)* has thc structurc **18.**

IV. DIAZOALKANES

A. General

The structures of only a few diazoalkanes not containing other characteristic groups (diazomethane, 2-bromodiazofluorene and phenyl(triphenylsilyl)diazomethane) have been determined. The structural parameters for diazomethane have been reviewed previously^{1, 82, 83} and will not be reported here.

In the diazoalkanes examined to date, the C-N bond length $(1.28-1.32 \text{ Å})$ is greater than that observed for a $C_{sp} = N$ bond $(1.23-1.26 \text{ Å})^{34}$. At the same time, the $N-N$ distance (1.12-1.13 Å) is greater than that found for dinitrogen and for benzenediazonium chloride (1.09)^{8, 16}. In agreement with most workers⁸⁵⁻⁹², this evidence suggests that the $=CN_2$ group is a resonance hybrid between the canonical forms **19** and **21.** Form **21** is commonly not reported sincc its contribution is very small compared to that of **19** and **20,** as it possesses fewcr covalent bonds and there is a largcr charge separation.

The participation of the two extreme forms 19 and 20 to the resonance hybrid is confirmed by the infrared spectra. In fact, the frequency of the $N-N$ stretching vibration band **(2088** cm⁻¹ in ether and in CCl₁ for the molecule CH_2N_2 ^{98, 93} is

smaller than that observed for the **p-dimcthylarninobenzenediazoniuin** cation (2166 cm⁻¹, in petroleum ether)²⁰, for which a relevant contribution of a quinoid form **is** now proved.

There has been some **work** on the relative contributions of forms **19** and **20** by means of n.m.r. and electric dipole moment measurements.

Interpretation *of* the n.m.r. spectra was based on the observation that in form **19** the electron density on the carbon atoms α and β to the diazo group is different from that in *20.* Consequently, the chemical shift of the protons bonded to these carbon atoms should give some indication as to the participation of the **two** canonical forms^{94, 95}. To this purpose, Ledwith and Friedrich⁸⁹ determined the nuclear magnetic resonance shielding parameters for the following compounds : H_2CN_2 , MeCHN₂, CH₂CHCHN₂, EtOCOCHN₂, MeOCOCHN₂, PhCHN₂, Ph_2CN_2 , (p-MeOC₆H₄)₂CN₂, diazocyclopentadiene and 9-diazofluorene. For these molecules a greater shielding than that for similar compounds^{96, 97} was found. According to the authors⁸⁹ this may be due to a considerable participation of form *20* and/or to the presence of diamagnetic anisotropy in the diazo group analogous to that observed for acetylenes and nitriles^{98, 99}.

It now appears that the high shielding found in n,m,r , spectra is due in the main to the presence of diamagnetic anisotropy. In fact, from the observed moments for the molecules Ph₂CN₂ (1.27-1.42 D)¹⁰⁰, ¹⁰¹, (p-ClC₆H₁)₂CN₂ (0.65 D)^{100, 101} and using $\mu(\overline{C_{\text{spr}}-C}) = 1.59 \text{ D}^{102}$, a value of $\mu(\overline{\text{CN}_2})$ of *c.* 1.0 D, having the direction indicated¹⁰³, is obtained. Taking into consideration the fact that the theoretical moment of form **19** is smaller than that of **20** (5.46 and 6.24 D, respectively, for diazomethane) $%$, we deduce that the first predominates over the second. In this context it is worthy of note that for diazomethane **19** and **20** are present in the ratio 66/34⁹⁰, in agreement with an independent estimate of 70/30 obtained from microwave quadrupole measurements¹⁰⁴. rs that the high shielding found in n.r.
of diamagnetic anisotropy. In fact, fr
 Ph_2CN_2 (1.27–1.42 D)¹⁰⁰, ¹⁰¹, (p-CIC
CI) = 1.59 D¹⁰², a value of μ (CN₂) of

B. X-Ray Data

I. 2-Bromodiazofluorene

 $C_{13}H_7N_2Br$ ¹⁰⁵ (22): $R = 0.09$; standard deviations 0.031-0.040 Å and 1.8-2.8°. All the atoms lie in the same plane apart from Br, $C_{(7)}$ and $N_{(2)}$, which are slightly shifted from this plane.

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Uncorrected

2. Phenyl(triphenylsily1) diazomethane

 $Ph(Ph_3Si)CN_2^{106}$ (23): $R = 0.117$; standard deviations 0.016-0.020 Å and **0-5-1.4".** The **C-H** bond length is 1.05-1.08 **A.** The molecule **is** monomeric. The structural parameters of the CN_2 group are very close to those found for diazomethane^{1, 82, 83}. The Si-C bond distance (1.865 Å) is similar to that observed for tetraphenylsilane (1.872 Å)¹⁰⁷. The present data do not suggest the participation of a form $\equiv \overline{S}i = C - \overline{N} = N$, involving 3d orbitals, to the resonance hybrid.

Uncorrected

V. AROMATIC DIAZOCYANIDES

A. Synthesis

The following isomers of these compounds have been isolated: the addition product PhN,CN.HCN and the two isomeric forms, *sytz* and *mri,* of the diazocyanide. The former is labile and the latter stable. The addition product is prepared in acid medium and in excess of cyanide108-111. The other two compounds **are** also prepared in acid medium, but the cyanide is added slowly and is always in less than molar ratio with respect to the diazonium salt and the $acid¹¹²⁻¹¹⁵$. If the reaction is carried out at -5 \degree C and in sufficiently concentrated solution, the labile form precipitates as a crystalline compound. At room temperature, the labile isomer slowly isomerizes to the stablc one. Finally, if the reaction is carried out by adding a neutral solution of the diazonium salt to **a** concentrated solution of potassium a neutral solution of the diazonium salt to a concentrated solution of potassium
cyanide in a water-alcohol mixture at -20 °C, the compound obtained is
completely different from those mcntioned above^{108, 116}. On X-ray compound has been assigned the formula $ArN_2C(NH)CN$.

B. Structures

The structural parameters of the stable isomer of the diazocyanides have been determined in threc works. Unfortunately, because of their low stability, no structural

study has been carried out on the labile *syn* isomers. Even qualitative data on the latter would have allowed definite conclusions to be arrived at regarding the type of isomerism present in diazocyanides. This has bcen much discussed (see Section V.C).

The bond distances in the *anti*-diazocyanides fall in the following ranges: C_{sp} $-N$, 1.29-1.42 Å; N-N, 1.25-1.32 Å; N-C_{sp}, 1.36-1.41 Å and C=N 1.14-**I**.15 Å. Comparison of these distances with those for *trans-azobenzene* and its symmetrical derivatives gives useful information on the electronic structures of diazocyanides. In *trans-azobenzenc¹¹⁷, trans-4,4'-dichloroazobenzenc¹¹⁸, <i>trans-*4,4'-azodiphenetole¹¹⁹, *trans*-4-phenylazobenzene¹²⁰ the bond distances are: $C_{\rm sp}$:- N, $1.43-1.46$ Å, and N-N, $1.23-1.25$ Å. The presence of some delocalization in the azobenzenes (bond order 1.9 for N-N and 1.1 for $C_{\rm sp2}$ -N)⁹⁴ has been suggested by comparison with the aliphatic analogues. For the diazocyanides, which have no symmetry elements, this delocalization increases since the $C_{\text{sp}}-N$ bond length is shorter, and the $N-N$ greater, than those of *trans*-azobenzenes.

I. p-Chlorobenzene-anti-diazocyanide

 $0.7-1.9^\circ$. The molecule is planar. CIC₆H₄N₂CN¹²¹ (24): $R = 0.082$; standard deviations 0.007–0.030 Å and

2. o-Bromobenzene-anti-diazocyanide

and 0.7". The molecule is almost planar. BrC₆H₄N₂CN</sub>¹²² (25): $R = 0.059$; the standard deviations are 0.006-0.011 Å

3. o,p-Dibromobenzene-anti-diazocyanide

 $Br_2C_6H_3N_2CN$ ¹²³ (26): $R = 0.168$; almost planar.

 (26)

 $N_{(1)}$ $C_{(1)}$ - $C_{(1)}$ -

 $C_{(5)}$ - $C_{(6)}$ $C_{(1)}^{-}$ ${\rm C_{(6)}}$ -

4. p-Chlorobenrene-anti-diazoimidoglyoxynitrile

 $CIC_6H_4N_2C(NH)CN$ ¹¹⁶ (27): $R = 0.09$; standard deviations 0.007-0.014 Å and *0.5-0.8".* Other bond lengths: C-H, *0.90-1.05 8,* (**f.** 0.1), and N-H, **1-05** (**L-** 0.1 I). The molecule is essentially planar. The structure of the $C_6H_4N_2$ group is very similar to that observed in the other anti-diazocyanides.

Uncorrected

C. Isomerism

Since there are no literature data on the structure of the syn isomers of diazocyanides, it may be useful to coniment on the type of isomerism present in these .pounds.

For the basis of the structural results obtained for the behiche-*anti*-diazocyanides
seen above there are two possibilities for the isomerism of diazocyanides:
 $A rN = NNC$ (7) (7) On the basis of the structural results obtained for the benzone-anti-diazocyanides

$ArN = NNC$	$ArN = NCN$	(7)
$N = N$	CN	
Ar	CN	$Mr = N$
Ar	Cr	Ar
$syn (cis)$	$anti (trans)$	

The disagrcemcnt as to the significance of the chemical evidence, which has occupied several workers¹²⁴⁻¹³⁰, shows that it cannot constitute conclusive proof for either of the possibilities **(7)** and (8). The only physical results currently available have been obtained from dipole moment and infrared spectra measurements.

The electric dipole moments of some diazocyanides¹¹⁴, measured in benzene, are shown in Table 1. What conclusions can be drawn from these data? Since the group moments for nitrile and *iso*-nitrile are very similar¹³¹, information on the configuration may be obtained¹³² from an examination of several members of the series rather than by treating each singly. The situation **(7)** leads virtually to the same difference in observed dipole moments in each pair of $syn/anti$ isomers. Since, as seen from Table **1,** this is not the case, this possibility may be excluded. The possibility remains of a geometrical isomerism (8). We can try to verify this as follows. The group moment $\mu(2,4,6\text{-}Br_2C_6H_2N)$ may be assumed equal, to a good approximation, to μ (PhN), of a geometry
moment μ (2,
 $\overrightarrow{(\text{PhN})}$, which which is c. **2.0** D I14. Using this value, from the moment observed for 2,4,6-tribromobenzene-anti-diazocyanide, $\mu(N-C=N)$ is estimated to be *c.* 2.0 D¹¹⁴. With this value and the usual bond and group moments¹⁰², the following theoretical moments (μ_{calc}) for the *anti* isomers of $\text{XC}_6H_4N_2CN$ molecules are then deduced $(X, \mu_{\text{calc}}, \mu_{\text{obs}})$: 4-Cl, 2.4, 2.73; 4-Br, 2.4, 3.78; 4-NO₂, 2.0, 1.47. The discrepancy between observed and calculated moments do not allow conclusions to be drawn as to the presence of geometrical isomerism.

-0.000 , -0.000				
Compound	μ_{syn}	μ_{anti}	$\mu_{syn}-\mu_{anti}$	
p -CIC ₆ H ₄ N ₂ CN	2.93	3.73	-0.80	
p -BrC ₆ H ₄ N ₂ CN	2.91	3.78	-0.87	
$o\text{-}BrC_6H_4N_2CN$	3.79	5.32	-1.53	
p -NO ₂ C ₆ H ₁ N ₂ CN	2.04	1.47	$+0.57$	
$o, m, p - Br3C6H2N2CN$	2.5	4.0	$+1.5$	

TABLE 1. Observed dipole moments (D) for some aromatic diazocyanides¹¹⁴

Interpretation of the i.r. spcctra is based on the assumption that two geometrical isomers should have **C=N** stretching bands very close to one another in frequency $(v_{CN})^{115}$, 133. This is observed, for example, in 1-cyano-1-propene¹³⁴, which gives v_{CN} at 2221 cn1-l for the *cis* isomer and at 2223 for the trans isomer. By contrast, the difference between the v_{CN} values for a pair of structural isomers is of the order of 100 cm^{-1 115, 135, 136}. Several authors have used these observations to obtain information on the type of isomerism present in diazocyanides. The results obtained are shown in Table *2.* The data in this table rcveal some disagreement among the various workers. A difference in the v_{CN} values between the labile and the stable isomers has been found only by some authors¹¹⁵. It seems hazardous to use this evidence alone as suficient to indicate the presence of structural isomerism. In fact,

the integrated intensity of the $C \equiv N$ stretching vibration is greater in the stable isomer (nitrilic) than in the labile one (isonitrilic) while an isonitrile usually shows a more intense $C \equiv N$ band than the corresponding nitrile^{115, 135}. 136. In addition, the difference in the v_{CN} values is much smaller in a pair of diazocyanides than between nitrile-isonitrile pairs. Some workers explain this fact¹¹⁵ by invoking a strong

 $Ar-N=\underline{N}$ ₇ $C \equiv N$ conjugation, which would not only lower the frequency for this isomer but would also lead to an almost linear $-NNCN$ group. This explanation is discounted by the X-ray structural parameters obtained for the benzene-antidiazocyanides (see Section V.B). In fact, these molecules have a non-linear structure with a C=N bond equal to 1.14 Å, which is of the same order, if not lower, as that found in most nitriles studied to date¹³⁷.

Compound	Isomer	ν (cm ⁻¹)	$A \times 10^{-4}$ (cm ⁻² mol ⁻¹ l)
p -MeOC ₆ H ₄ N ₂ CN	syn anti	2150^a 2190	0.11 0.47
p -CIC ₆ H ₁ N ₂ CN	syn anti	2187° 2187	
$p-\text{Br}C_6H_4N_2CN$	syn	2154^a 2187 ^b 2185c	0.026
	anti	2192^u 2187 ^b 2185c	0.17
$p\text{-}NO_2C_6H_4N_2CN$	syn	2162 ^a 2187^b	0.037
	anti	2193 ^a 2187°	0.064
$o,m,p-\text{Br}_3\text{C}_6\text{H}_2\text{N}_2\text{CN}$	syn anti	2187c 2185	

TABLE 2. Frequency (v_{CN}) and intensity of absorption (A) of the C=N stretching vibration band for some aromatic diazocyanides

^a In CCl₄ 115. b In CCI_s 132.

 ϵ The tribromo compounds were examined in CHCl₃ and CS₂ solutions and the monobromo one in nujol mull¹³³.

To conclude, the results of Kazitsyna and coworkers¹¹⁵ also do not constitute proof that situation (7) is present. On the basis of indirect evidence, i.e. by analogy with similar compounds, we tend, like most workers¹, toward isomerism of geo- \therefore rical type (8) .

VI. DIAZOKETONES

A. Isomerism

I. General

Different diazoketones may be obtained, depending on the position of the carbonyl group with respect to the diazo group. Most work has been centred on the a-derivatives because they are the easiest to prepare and have also numerous applications in organic syntheses^{1, 138-141}.

The configurations adopted by α -diazoketones depend on the conjugation between the diazo group, which in this case acts with $a + M$ effect, and the carbonyl group, which exerts $a - M$ effect. The atomic orbital scheme involved in this delocalization is shown in Figure 3. For the only molecular structure of α -diazoketones solved to α date, 1,4-bisdiazo-2,3-butanedione¹¹², the following values were obtained: CNN aate, 1,4-bisdiazo-2,3-butanedione¹², the following values were obtained: UNN
176^{.8°}, CCN 116^{.5°}. These results indicate that in the *α*-diazoketones the carbon atom $C_{(2)}$ (Figure 3) is essentially sp² hybridized and both nitiogen atoms have sp

FIGURE 3. Atomic orbital scheme involved in conjugation between diazo and carbonyl groups.

hybridization. The π_v orbitals are localized on the latter two atoms, which each furnish an electron to the bond formed between them. Instead, the $\pi_{\mathbf{z}}$ orbitals are delocalized over several atoms (e.g. five in the case of diazoacetone, $MeCOCHN₂$) to different extents depending on the nature of the substituents **R'** and **R2.** The nitrogen atom $N_{(1)}$ donates two electrons to this molecular orbital and all the other atoms donate one each. The delocalization of the π_x electrons on the atoms of the molecular skeleton causes the central $C_{(1)}-C_{(2)}$ bond to become partially double bonded. It follows that two rotational isomers may exist, each having the $COCN₂$ group more or less planar (Figure **3)** depending on the electronic, steric and electrostatic effects exerted by **R'** and **R2.**

Apart from this geometrical isomerism, α -diazoketones may also, in theory, show structural isomerism due to the transposition of the hydrogen atoms α to the carbonyl group. Depending on whether the hydrogen migrates to the carbonyl oxygen or to the terminal nitrogcn there will be keto-enol or diazo-isodiazo isomerisin, respectively.

Identification of the type of isomerism present becomes more difficult since diazoketones may form an intcrrnolecular hydrogen bond with another diazoketone molecule or with a hydroxylic compound¹⁴³⁻¹⁴⁵. This explains why early results, although experimentally valid, often led to incorrect conclusions.

We now examine the theorctical and experimental evidence for the type of isomerism present in α -diazoketones in different media.

2. Ultraviolet/visible and infrared spectra

The ultraviolet/visible spectra of α -diazoketones in mixtures of non-polar and hydroxylic solvents show a characteristic behaviour. The $-COCN₂$ group has an intense band **at** *c.* 245-250 nm, which becomes weaker with increase in the hydroxylic/non-polar solvent ratio, and another band at 270-290 nm which at the same time increases in intensity¹⁴⁶⁻¹⁵⁰. There are no shifts in the bands on going from one solvent to the othcr. In almost all the cases studied to date the curves show an isosbestic point which seems less clear the lower is the availability of the carbonyl oxygen lone pair, i.e. the higher is the frequency of the $C=O$ stretching mode¹⁴⁸. The presence of an isosbestic point suggests that two species exist in solution in equilibrium with one another¹⁵¹. There are four possibilities for our molecules:

the keto-enol equilibrium

the diazo-isodiazo equilibrium

the conformational equilibrium

$$
cis \xrightarrow{(30)} trans
$$

and, finally, the presence of a hydrogen bond **(31).**

The existence of an isosbestic point does not necessarily mean that situations *28* or *29* are present since it is also observed for molecules which do not contain transposable hydrogen atoms^{148, 149}. The same may be said of the conformational *5*

equilibrium, 30, since a similar dependence on solvent and an isosbestic point are also found in the spectra of compounds for which this isomerism is not possible, e.g. 2-diazocyclohexanone¹⁵², 3-diazonorcamphor and 3-diazocamphor¹⁵³.

For a variety of reasons, we favour the possibility (31). In agreement with Fahr¹⁴⁸ it is our opinion that in mixed solvents there is formation of a hydrogen bond between the diazo compound and the hydroxylic solvent. The evidence for this conclusion is derived from the fact that the isosbestic point becomes more evident the greater is the availability of the oxygen lone pair to form this type of bond. The presence of the latter is proved by the behaviour of the $O-H$ stretching vibration band of phenol in the presence of various α -diazoketones¹⁴⁹. This interpretation explains why the same behaviour in mixed solvents is observed for molecules for which two conformers are possible and those for which they are not^{149, 152, 153}. In agreement with this, for diazoacetone, MeCOCHN₂, and diazoacetophenone, PhCOCHN₂, it was observed that the band due to the $\pi \rightarrow \pi^*$ transition in the *cis* isomer overlaps that in the *trans* isomer.¹⁵³

As to the infrared spectra of a-diazoketones in mixed solvents, Foffani and coworkers¹⁴⁹ observed that integrated intensities and frequencies of the $N-N$ stretching vibration band are almost invariant for diazoacetone, diazoacetophenone and benzoyl(diazo)phenylmethane on varying the ratio between non-polar and hydroxylic solvents. Morcover the $C=O$ stretching band at 1662 cm⁻¹ in diazoacetone decreases in intensity, in C_2Cl_4 as solvent, with increasing quantity of phenol. At the same time the peaks at 1643 and 1633 cm^{-1} , which are very weak in intensity in non-polar solvents, became more intense with increasing phenol concentration. These phenomena may be due to the presence of hydrogen bonding between the carbonylic compound and the hydroxylic solvent, which stabilizes one conformer more than the other. According to this, for molecules having two conformers, one peak in the $N-N$ stretching band disappears in the presence of hydroxylic solvents¹⁵⁴. This interpretation, which reconciles the u.v. and i.r. results, appears plausible but requires further experimental justification.

The atoms involved in the hydrogen bond may, in theory, be the terminal nitrogen, the carbon of the diazo group and the carbonyl oxygen. Involvement of the nitrogen or of the carbon atom may be excluded since the i.r. spectra of phenol in the presence of diphenyldiazomethane **(0.004** M) in CCI, show no evidence for its presence145. The only atom available *is* thus the carbonyl oxygen.

The formation of an intermolecular hydrogen bond between two diazoketone molecules has recently been established^{144, 145} for RCOCHN₂ compounds. N.m.r. measurements were carried out in weakly accepting solvents such as CCI_4 and d_{θ} -benzene at a temperature at which interconversion between possible rotamers is very fast and it was found that the chemical shift varies regularly with the concentration of the diazo compound¹⁴⁵.

The possibility that two rotamers exist in equilibrium in apolar solvents, foreseen by Foffani and coworkers¹⁴⁹, has recently been established by Sorriso and coworkers^{155, 156} by examining the N-N stretching vibration band in CCI₄. This band is suitable for our purpose because it falls in a frequency range in which other groups present in the molecule do not absorb. The molecules $CH_2CICOCHN₂$, $CHCl₂COCHN₂$, t-BuCOCt-BuN₂, PhCOCHN₂ and PhCOCMeN₂ show a single $N-N$ stretching band, while MeCOCHN₂, MeCOCMeN₂ and PhCOCPhN₂ give split N-N bands (Figure 4). Since rotamers stabilized by a sufficiently high-energy barrier to interconversion have different i.r. spectra, the splitting of the diazo band indicates the presence of two conformers.

Insight into the reasons for the existence of two rotamers in α -diazoketones comes also from an examination of the frequencies of the carbonyl and diazo stretching vibrations, which fall around 1650 and 2080 cm⁻¹, respectively^{88, 93, 155-158}. These frequencies are clearly shifted compared to those in the corresponding ketones²³ and diazohydrocarbons^{88, 93}; v_{CO} moves to lower frequency and v_{NN} to higher frequency. This indicates that in these molecules the carbonyl group has lcss doublebond character than it does in ketones, while the $N-N$ bond is of higher order than

FIGURE 4. Profile of N-N stretching vibration band observed in CCl₄ for: (a) diazoacetone McCOCHN,; (b) **I-acetyl-I-diazoetliane,** McCOCMeN,; (c) benzoyl(diazo)phenylmethane, PhCOCPhN₂; (d) diazoacetophenone, PhCOCHN₂.

in diazohydrocarbons of the same order of substitution. In other words, a third limiting form, **34,** which occurs at the expense of form **33,** is in addition to the canonical forms **32** and **33** also present in diazohydrocarbons. The participation of the form **34** gives the central C-C bond a partial double bond character. The contribution of forms of type **33** and **34** is higher the higher the frequency of the diazo band and the lower its intensity. The oppcsilc: holds for contribution of forms of the type **32 93. 160.**

When R¹ is a phenyl group, forms of the type 35 may also contribute^{88, 149}.

3. Nuclear magnetic resonance

One of the most important applications of n.m.r. spectroscopy is in structural studies by means of the chemical shift (δ) and coupling constant (J) parameters^{96, 97, 161, 162. The chemical shift and coupling constant, which give comple-} menlary structural information, are directly connected to the electron distribution about the nucleus and to the type of bond between the nuclei, respectively. Both parameters change with change in the molecular environment. Thus, the use of the technique in studies of the type of isomerism present and determination of thermodynamic parameters for isomer equilibrium is understandable.

 α -Diazoketones of general formula RCOCHN₂ have been widely studied using p.m.r.Id4. Some results of this study are shown in Table **3.** Temperature-dependent p.m.r. spectra have been observed for the molecules $HCOCHN₂$, MeCOCHN₂, EtCOCHN,, PhCH,COCHN,, MeOCOCHN, and EtOCOCHN,. **At** room temperature there is a single sharp peak which broadens and splits into a doublet on lowering the temperature (Figure **5).** Only a single peak is observed in the

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t-butyl derivative (*t*-BuCOCHN₂) and in aryldiazoketones (PhCOCHN₂, p -NO₂C₆- $H₄COCHN₂$ and p-MeOC₆H₁COCHN₂) the intensity of which remains constant at different temperatures. As for analogous cases¹⁶³⁻¹⁶⁵ this indicates that two forms are present in equilibrium in **the** first six compounds, whilst the aryldiazoketones give only a single peak and thus only a singlc spccics, or else two in rapid interconversion or coincidentally having the same chemical shift. Given the steric effects present, the r-butyl dcrivative almost certainly exists in a single, very distorted, form.

TABLE 3. Fractional populations *(P),* equilibrium constants *(Keg)* and standard free-energy differences (ΔF) of the *cis* and *trans* forms of diazoketones (RCOCHN₂), 40% w/w solution in deuteriochloroforni. Reprinted with permission from Kaplan and Meloy, *J. Amer. Chew.* Soc., 88, 950 (1966). Copyright by the American Chemical Society

R	P_{cis}	P_{trans}	$K_{\rm eq}$ (T, °C)	ΔF (kcal/mol) (T, °C)
H^a	$0.69 + 0.02c$	0.31 ± 0.02	$0.449(-20)$	$0.40(-20)$
Me	$0.924 \div 0.008$	$0.076 + 0.008$	$0.082(-40)$	$1.16(-40)$
Mc^b	$0.903 + 0.002$	0.097 ± 0.002	$0.180(-40)$	$0.79(-40)$
t -C ₄ H ₉	> 0.99	< 0.01 ^d		
PhCH ₂	$0.962 + 0.002$	$0.038 + 0.002$	$0.040(-40)$	$1.49(-40)$
McO	$0.538 + 0.004$	$0.462 + 0.004$	$0.859(-50)$	$0.07(-50)$
E1O	$0.54 + 0.01$	0.46 ± 0.01	$0.840(-50)$	$0.08(-50)$

Values of energy of activation (E_a) , frequency factor (A) , free energy of activation $(\Delta F')$ and temperature of coalescence (T_e) for hindered internal rotation about the C-C central bond in diazoketones ($RCOCHN₂$), 40% w/w solution in deuteriochloroform

a Concentration unknown.

^b*20%* **w/w** solution in deuteriochloroforni.

Average deviation involving at least two separate samplcs.

^d The *trans* resonance line was not observed.

Errors represent the prccision of the Arrhenius plots. The free energies of activation were calculated from the relationship $\Delta F'_T = 2.303 RT \log (\tau_x Kk)$. The transmission coefficients, *K,* are assumed to be unity.

For the compounds which give doubling of the methine proton peak at low temperatures, attempts have been made to assign structures. Kaplan and Meloy¹⁴⁴ considered the possibilities: keto-enol **(28)** and diazo-isodiazo **(29)** structural isomerisms; geometrical isomerism (30) (Figure 3). Tautomeric equilibria (28) and **(29)** may be excluded since the $^{13}C-H$ coupling constant for high-temperature timeaverage species is the same as that of the species predominant at low temperature $(J = 199 \text{ c.p.s.})$ for McCOCHN₂, EtCOCHN₂ and EtOCOCHN₂. If structural

isomers had been present, the time-averaged species at high temperature would have been expected to show coupling constants different from those of the species predominant at low temperature, since the two species have different $\overline{C}-\overline{F}$ environments.

FIGURE 5. Temperature dependence of the n.m.r. spectrum of the methine proton of methyl diazoacetate (MeOCOCHN₂). Reprinted, with permission, from Kaplan and Meloy, *J. Amev. Chem. Soc., 88,* 950 **(1966).** Copyright by the American Chemical Society.

Having assigned the two peaks to the two conformers¹⁶⁶⁻¹⁶⁹, the thermodynamic parameters for the $cis \neq trans$ equilibrium have been calculated by integrating the peaks (Table **3).** The values listed in this table show that the ratio of the two conformers does not depend only on stcric factors.

The cnergy barriers to interconversion between the two rotamers fall in the range 9-18 kcal/mol. The lower values are obtained for diazo esters, in which diazocarbonyl conjugation is lower and hence the double-bond character of the $C-C$ central bond is lower.

The ¹³C-H coupling constant ($J = 199$ c.p.s.) indicates¹⁴⁴ that the C-H bond has high s character (41%) , which suggests that the methine hydrogen may participate in hydrogen bonding.

The p.m.r. evidence leaves little doubt that the isomerism present in α -diazoketones is geometrical isomerism. Further conformational studies are thus airned niore at obtaining insight into the cffect of thc R' and **R2** groups on thc *cisltrotrs* ratio and into the theoretical aspects of this isomerism rather than proving its existence.

4. Electric dipole moments

The usefulness of electric dipole moments in conformational studies has been known for some time^{102, 170-172.} For α -diazoketones this technique has been used to determine the relative contribution of the two rotamers and to study the effect of the **R1** and R? groups. The observed dipole moments, in benzene, at *25* "C and assuming $P_a = 0$, are (compound, dipole moment in debye): MeCOCHN₂, 3.37; CH,CICOCHNz, 2.80; CHCI,COCHN,, 2.65 ; CCI,COCHN,, 2.46 **lo3;** PhCO-CHN₂, 3.66; p-MeC₆H₄COCHN₂, 4.01; p-ClC₆H₄COCHN₂, 3.08; p-BrC₆H₄CO-CHN₂, 3.11; $p\text{-MeOC}_6H_4COCHN_2$, 3.73; $PhCH_2COCHN_2$, 3.31; $Ph(CH_2)_{2}$ -3.56 **lol** ; MeCOCMcN,, **2-45;** Y-BUCOC~-BUN,, 1.98; PhCOCMeN,, **2.75;** $p\text{-ClC}_6H_4COCMeN_2$, 2.95 ; $o\text{-ClC}_6H_4COCMeN_2$, 3.18 COCHN₂, 3.39 ¹⁷³; BzCPhN₂, 2.77; m-CIC₆H₄COCHN₂, 3.34; o-CIC₆H₄COCHN₂,

The observed moments indicate that whilst MeCOCHN₂, PhCH₂COCHN₂, $Ph(CH₂)₂COCHN₂$, BzCPhN₂, o -CIC₆H₄COCHN₂, MeCOCMeN₂, p-CIC₆H₄-COCMeN₂, PhCOCMeN₂ and o -CIC₆H₄COCMeN₂ may be two rotamers, only one species, the *cis* one, is present in halogen derivatives of diazoacetone. Presumably there is a strong electrostatic effect by the halogen atom on the diazo group.

The aryldiazokctones are almost entirely present **as** the cis conformer. This result agrees with the X-ray data¹⁷⁵ and with the fact that only a single sharp peak for the methine proton is observed in the n.m.r.¹⁴⁴ and the $N-N$ stretching vibration band is single in $\text{CC}l_4$ ¹⁵⁵.

Finally, the *t*-butyl derivative gives only a single rather distorted structure with an angle of c. 140[°] between the CCO and CCN planes.

5. MO calculations

MO calculations on α -diazoketones have essentially two aims: to see if there is theoretical justification for the rotational isomerism found experimentally, and to obtain information on the molecular conformation of rather complicated conipounds, information not obtainable by other means.

 C sizmadia and coworkers¹⁵³ have performed MO calculations on diazoacetone and diazoacetophenone using Hoffmann's extended Hückel method. The scheme used and the results obtained are shown in Figures 6 and 7. From Figure 6 it may be seen that two conformers, *cis* and *trans*, both with almost the same energy and the COCHN, group planar, are possible for diazoacetone. The energy barrier height to interconversion between the two species (\sim 14 kcal/mol) is of the same order as that determined experimentally from n.m.r. spectra in CDCI₃ (\sim 16 kcal/ mol)¹⁴⁴. As expected, variation in the angle ϕ has an almost negligible effect on the relative energies of the two rotamers and on the barrier height.

By contrast, the effect on changes in angle ϕ in diazoacetophenone is considerable (Figure 7). At $\gamma = 120^{\circ}$ and $\phi = 0^{\circ}$ there is only a single isomer, the *cis* form. Instead on varying the angle ϕ , i.e. on rotating the pheny! group out of the molecular planc, diazoacetophenone gives two conformers having similar energies and a barrier height of the order of 14 kcal/mol. This result is not in agreement with the i.r. and n.m.r. spectra and the dipole moments, which suggest only one isomer, the cis one. The discrepancy between theoretical and experimental evidence may be due to solvation and/or to entropic factors¹⁵³.

Sorriso and coworkers have studied the molecules CH₂CICOCHN₂, $MeCOCMeN₂$ ¹⁷⁶, PhCOCPhN₂ and PhCOCMeN₂¹⁷⁷ by the CNDO/2 method using the structural parameters of Figures 6 and 7. The calculated energies suggest

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that two conformers were present in equilibrium in MeCOCMcN₂ and PhCOCPhN, and only one, the *cis,* in the chloro derivative. Finally, two forms in rapid interconversion exist for PhCOCMeN₂.

For PhCOCMeN₂ and PhCOCPhN₂, as for the diazoacetophenone, there is concerted rotation of the phcnyl and the methyl groups about the bond with the molecular skeleton during the interconversion between the two rotamers.

The phenyl group bonded to the carbonyl conjugates with it, in agreement with experimental results^{88, 119}. In addition, in PhCOCPhN₂ the phenyl bonded to the diazo group conjugates with it. Dcspite this, diazo-carbonyl conjugation prcdominates over the others.

6. **Conclusions**

The results obtained to date show that the isomerism present in α -diazokctones is geometrical isomerism. **As** mentioned in Section **VI.A.1,** this arises because of conjugation between diazo and carbonyl groups, which makes the central $C-C$ bond partially double-bonded. This suggcstion is supported by the frequency of the $N-N$ and $C-O$ stretching vibrations compared with those in diazohydrocarbons and ketones. The presence of two conformers is confirmed by the low-tcmperaturc doubling of the methine proton in n.m.r. specira of several diazoketones of the type $RCOCHN₂$ and the doubling of the N-N stretching vibration band. MO calculations using two different methods also show that, if hindrance of all kinds is absent, the configuration preferred by the $COCN₂$ group is a planar one.

Two factors seem to control the cis $\frac{1}{x}$ trans equilibrium in α -diazoketones $R¹COCR²N₂$. The first is diazo-carbonyl conjugation, which is responsible for the energy barrier height to interconversion between the two conformers. The latter increases if R¹ is electron attracting and decreases if it is an electron donor. By contrast, it is lowered if R² is electron attracting since such a group involves the π system of the diazo group, lowering the conjugation of the latter with the carbonyl. The second factor is the influence of steric and clectrostatic effects exerted between **R', R2,** the carbonyl group and the diazo group. This factor causes variations in both the height and relative energy of the two conformers and therefore in thcir population.

Finally, for most of the molecules of general formula RCOCHN,, the *cis* conformer is preferred to the *trans* one. This may be due to two reasons. The first is the fact that the diazo group is larger than the hydrogen and thercfore repulsion between it and R is greater than that between the hydrogen and R. The second is the through-space interaction between the carbonyl and the diazo group. The latter may be related to the presence of a fraction of negative charge on the carbonyl oxygen and of one of positive charge on $N_{(1)}$, which would attract one another, or to an interaction between the π systems of the diazo and the carbonyl groups. That **a** certain stabilizing action is present in the *cis* form is shown by the clear predominance of the *cis* species over the *trans* one in diazoacetaldehyde, HCOCHN₂. N.m.r. measurements lead to a *cis/trans* ratio of 7/3 for this molecule¹⁴⁴. In the absence of the stabilizing effect, the *trans* form would have been favoured, as shown by models constructed using van der Waals' radii.

B. Configuration and Wolff Rearrangement

Knowledge of the configuration of a-diazoketones allows suggestions to be made regarding the mechanism of their decomposition¹⁴⁴. However, in order that the products obtained may be correlated with thc structures of the starting molecules, **it** is necessary that the rate of interconversion between the two rotamers be lower than their rate of decomposition (36). This is not the case for diazoacetone¹⁴⁴, for which a first-order rate constant of 1.68×10^{-4} sec⁻¹ has been calculated for the decomposition in *t*-butanol at 80 °C, and the *cis* \rightarrow *trans* interconversion has a rate constant of 10^3 sec⁻¹.

It **is** known that the decomposition of a-diazoketones **(37)** under appropriate conditions, leads to α, β -unsaturated ketones (38) and ketenes (39). The photolytic and silver-oxide catalysed decomposition of α -diazoketones at room temperatures gives mainly **38** and at **high** temperature **39.** For PhCOCMeN, and (Me,C)COC- $(CMe₃)N₂$, the dipole moment data allow a semiquantitative analysis of the results.

If two conformers with COCN, group planar **are** present for these molecules, the following *cisllrans* ratio would be expected : PhCOCMeN,, **44/56** and (Mc,C)COC- $(CMe₃)N₂$, 7/93. These values are of the same order as the ketene/unsaturated ketone ratios obtained experimentally. In fact, for thc unsaturated ketone, for decomposition of PhCOCMeN, catalysed by Ag,Olin we obtain **a** ratio of **72%,** and for decomposition of $(Me₃C)COC(CMe₃)N₂$ (pyrolysis, irradiation or acid-catalysed rearrangement)¹⁷⁹ we obtain 80-90%. This evidence suggests that there is some dependence of the mechanism of decomposition on the conformation of α -diazoketones. This would mean that the decomposition occurs not through an intermediate carbene radical but, rather, via **a** concerted migration process of an alkyl or aryl group and loss of a nitrogen molecule¹⁴⁴. The cis form presents the more suitable situation for this mechanism, which would also explain the effect of temperature on the nature of the decomposition products. In fact, increase in temperature favours the higher energy rotamers (for the two molecules concerned, the cis form) and with this the formation of the ketene, exactly as observed experimentally.

Finally, the fact that the two molecules PhCOCMeN₂ and (Me₃C)COC(Me₃C)N₂ are not planar means that the mechanism describing the Wolff rearrangement is most facilitated in the *cis* conformer and impeded in the *frms* one and that there are intermediate situations between these two extremes.

C. Structures

The molecules of α -diazoketones for which X-ray crystal structures have been reported are not, unfortunately, the more representative ones. **It** is, therefore, not possible to deduce general conclusions for this class of compounds nor to describe their clectronic configuration from the present structural results.

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I. I, 4-Bisdiazo-2,3-butanedione

 $C_4H_2N_4O_2$ ¹⁴² (40): $R = 0.029$, standard deviation = 0.001 Å. The molecule is planar with **a** centre of symmetry and has a *cis* conformation for both **COCN,** groups. **As** pointed out by the authors'"?, these results appear to be contradictory. In fact, the C-O distance, of the same order as that commonly observed for ketones¹³⁷, and the N-N bond, virtually the same as that for diazonium salts (see Section **II.A.I),** suggest **a** structurc of :hc 'diazo oxide' type **32** and **33,** in which the triple-bonded $N-N$ type 33 predominates, as in unsubstituted diazonium salts. In contrast, the values above indicate a significant double-bond character in the bonds **C-N** and **CO-CN,** which may be rationalized by invoking **a** sensible contribution by the canonical forms **32** and **34,** respectively, to the resonance hybrid.

As mentioned previously (Section **VI.A.2),** the formation of an intraniolecular hydrogen bond between the methine hydrogen and the oxygen of the carbonyl group in solution has been reported by several workers. For the niolecule treated here, there seems to be little evidence for such **a** bond in the solid state as the minimum 0-H distance **(2.66 A),** even considering the usual **X-ray** displacement of the hydrogen atom, is slightly higher than the commonly observed value of 2.52 Å^{142} .

2. 5-Diazo-6-methoxy-6-hydrouracil

 $C_5H_6N_4O_3$ ¹⁸⁰ (41): $R = 0.056$; standard deviation ≤ 0.007 Å. Only the bond distances have been reported.

3. 2'-Deoxy-5-diazo-6-hydro-0~,5'-cyclouridine

distances have been reported. $C_9H_{10}N_4O_5$ ¹⁸⁰ (42): $R = 0.052$; standard deviation ≤ 0.007 Å. Only the bond S. Sorriso

The bond distances observed for 41 and 42 suggest that as for the other diazoketones, the $COCN₂$ group in these molecules is a resonance hybrid between the canonical forms 43–45. The other parts of the molecule do not conjugate with $COCN₂$ group.

VII. DIAZO OXIDES

A. General

When alkali is added to *ortho-* and *para-hydroxy-substituted diazonium salts* and the mixture heated, compounds of the type 46 are obtained:

The nature of the bonding present and the character of the six-membered ring are the subject of this section.

These substances are of some interest as they are used as intermediates in the laboratory and in industry and because they can be viewed, theoretically, as representing a situation intermediate between the diazonium salts and the diazoalkanes.

Various names have been suggested for these compounds: diazoquinones, diazoanhydrides, diazo oxides and quinone diazides. For the derivatives of benzene and naphthalene, the names diazophenols and diazonaphthols, respectively, have also been used. In the present chapter for most cases we will adopt the name diazo oxides.

Several suggestions have been put forward as to the structure of diazo oxides¹⁸¹⁻¹⁸⁹, 47 and 48. Some of these are now unacceptable on the basis of modern electronic theory. Others, such as the cyclic structures, may be excluded on the basis of the X-ray data (Section VII.C) and of the frequencies of the $N-N$ and $C\rightarrow O$ stretching vibration bands, some of which are shown in Table 4. As can be seen from this table, these frequencies are found in regions characteristic of absorptions by a N-N bond intermediate between a double and a triple bond and by a $C-O$ bond slightly lower than a double bond.

Substituent	Solvent or physical state	$\nu_{\rm NN}$ (cm ⁻¹)	$v_{\rm CO}$ (cm ⁻¹)	Reference
Diazophenols: <i>ortho</i> derivatives		O		
Unsubstituted $5-Mc$ $4,5 \cdot Me_2$ 4,5-Me, $5 - Bu - t$ $3, 5-(Bu-t)2$ 5-MeO $3,6-McO$ 5- $PhCH2O$ $5-C1$ $6-C1$ $4, 5 - Cl2$ $3, 5 - Br_2$ $4-NO2$ $4-NO2$ $4,6-(NO2)2$ 4-COOH	K Br KBr K Br CHCI ₃ CHCI ₃ CHCl ₃ CHCl ₃ CHCl ₃ CHCl ₃ CHCl ₃ CHCl ₃ CHCI ₃ Paraffin mulls Paraffin mulls Paraffin mulls K Br KBr	Ν, 2140 2175 2175 2130 2170 2120 2150 2100 2182 2160 2135 2124 2128 2135 2087 2150 2205 2158 2195 2158 2160 2110 2157 2183 2150 2225 2215	1620 1613 1625 1625 1618 1625 1620 1613 1620 1613 1610 1613 1582 1562	190 190 190 191 190 190 190 190 190 190 190 190 192 192 41 193 193
Diazophenols: para derivatives Unsubstituted Unsubstituted $2,6$ -Me ₂ $2,6$ -Me $_2$ $2, 5 - Me2$ $2,6$ -Cl ₂ 2-Me, 6-COOH 2 -CCOH, 6 -NO α	CHCl ₃ Paraffin mulls CHCI ₃ Dioxane CHCl ₃ KBr KBr KBr	2080 2109 2090 2060 2070 2140 2225 2225	1615 1600	191 41 191 194 191 193 193 193
Diazonaphthols 1,2-Naphthaquinone-1- diazide 4-Nitro-1,2-naphtha- quinone-1-diazide 1,2-Naphthaquinone-2- diazide 1,2-Naphthaquinone-2-	CHCl ₃ Paraffin mulls CHCl ₃ Paraffin mulls	2132 2096 2015 2155 2114 2148	1621 1642 1621 1618 1610	195 192 195 192
diazide 1,4-Naphthaquinonediazide	Paraffin mulls	2014	1618	192

TABLE 4. Values of the stretching vibration frequencies v_{NN} and v_{CO} for some diazo oxides

For the values of the stretching vibration frequencies reported for other diazo oxides see Reference 196.

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The problem of the electronic structure of diazo oxides may at present be put in these terms: should these compounds be considered quinoid **(483)** or benzenoid **(48b)** in character, or a resonancc hybrid between both extreme forms **48?** We will attempt a reply to this question in Section **VI1.B.**

B. Electronic Structure

Some time ago, Sidgwick and coworkers¹⁹⁷ suggested that the diazo oxides could better be represented by a resonance hybrid between the two forms **48.** This idea was later accepted by other authors^{198, 199}. Anderson and Roedel¹⁸⁹ found that the visible and ultraviolet spectra of some diazophenols and diazonaphthols were very similar to those of the corresponding quinones and inferred that the former must have a quinoid structure. Le Fèvre and coworkers^{192, 200}, from dipole moment measurements and **i.r.** and U.V. spectra, concluded that the diazo oxides are a resonance hybrid between the canonical forms **48,** the quinoid form clearly predominating. The electric dipole moments, determined in non-polar solvent, are reported in Table 5. Vector analysis of the moments of this table may give the

Compound	
3,5-Dibromo-1,2-benzoquinone-2-diazide	2.9
1,2,-Naphthaquinone-1-diazide	4.0
1,2-Naphthaquinone-2-diazide	3.5
4-Nitro-1,2-naphthaquinone-1-diazide	4.6
1,4-Naphthaquinonediazide	4.2
p -Benzoquinonediazide	5.0

TABLE *5.* Electric dipole moments observed for some quinone diazides^a

^{*a*} In benzcne, at 25 °C. $P_{a} = 0$.

following qualitative information. As the moment observed for phenanthraquinone

is 3.0 D and $\mu \overrightarrow{(C=N=N)}$ is c. 5.4 D²⁰⁰, theoretical moments of c. 2.4 D are for the quinoid form **48a** of **1,4-naphthaquinonediazide** and 1,4-benzoquinoncdiazidc, which are much lower than the values observed for these compounds (4.2 and 5.0 D, respectively)²⁰⁰. The calculated moments become equal to the observed ones if it is assumed that the diazo oxides are a resonance hybrid between the two forms **48,** in which, however, the contribution of the zwitterionic form must be very small as it has a rather high theoretical moment **(27.4 D** *?O0.*

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The dipolc moment evidence is supported by i.r. measurements. In fact, Le Fèvre and coworkcrs observed that the frequency of the $C-O$ stretching vibration band, which is much lower than in the quinone analogues, indicates that in the diazo oxides an extreme form is present in which the $C-O$ bond is a single bond¹⁹². Results in agreement with these have also been obtained by Whetsel and coworkers4'.

The presence of a structure intermediate between the benzenoid and quinoid types was also suggested by Vaughan and Phillips²⁰¹ from thermal stability measurements.

Unusual results on the electronic structure of diazo oxides have been obtained **by** Kazitsyna and coworkers^{25, 202}. These authors performed MO calculations, using the Hiickel approximation, on *0-* and p-diazo oxides having different substituent groups. It was found that the $C-O$ bond order is slightly lower than double and the negative charge on the oxygen atom is slightly higher than that usually calculated for a carbonyl oxygen. In the molecule 49 the bond order for the bonds $C_{(4)}-C_{(3)}$

and $C_{(4)}-C_{(5)}$ is very close to that for a quinoid structure. However, on going from the atoms $C_{(2)}$ and $C_{(6)}$ towards the diazo group the situation changes. In fact, the bond order of $C_{(1)}-C_{(2)}$ and $C_{(6)}-C_{(1)}$ is noticeably higher than that for a quinoid structure and the C-N and N-N bonds are almost entirely single and triple bonds, respectively. In addition, the fraction of positive charge is localized on the two nitrogen atoms, which rationalizes the tendency of diazo oxides to undergo diazo coupling reactions. According to Kazitsyna and coworkers, the electronic structure of the present compounds is better described if the forms *50* are present, in addition to **48**. The participation of these canonical forms to the mesomerism of diazo oxides

> $O=\overbrace{}^{+}$ $N\equiv N$ (two forms) **(50)**

might account for the above-mentioned $C-O$, $C-N$ and $N-N$ bond orders and also for the fact that the negative charge is greater on the oxygen atom and on the atoms $C_{(1)}$, $C_{(3)}$ and $C_{(5)}$. Finally, it was found that the distribution of electrons on the molecule is very sensitive to the parameters used. Although this result is further evidence that the electronic system of the diazo group is very mobile it also induces caution in the interpretation of results from such calculations.

We cannot conclude this examination of the literature data without underlining the similarity in i.r. behaviour between the diazo oxides **and** the a-diazoketones. In both types of compounds the v_{NN} and v_{CO} values are noticeably shifted compared to those in the corresponding diazohydrocarbons and ketones: ν_{NN} to higher frequency and $v_{\rm CO}$ to lower frequency. From the evidence discussed above we come to the conclusion that this behaviour has a single cause: conjugation between the π system of the diazo group and the π electrons of the C-O bond. Consequently, at the present stage of work on these compounds, the most likely situation seems that of **a** resonance hybrid having the main extreme forms **48,** with the quinoid form being predominant.

C. X-Ray Data

The two diazo oxide structures determined to date are not representative of this class of compounds, and the data reported here cannot give insight on the electronic structure of these molecules.

1. 3.6-Bisdiazocyclohexanetetraone

 $C_6N_4O_4^{203}$ (51): $R = 0.079$; standard deviations 0.010 Å and 0.7-0.9°. The molecule is planar. The N-N bond length (1.10 Å) is of the same order as that of diazonium salts (see Section II.A.1), while it is slightly shorter than that found in diazoalkanes (1.12–1.13 Å). This would indicate that the form having a triple $N-N$

bond is predominant, as in diazonium salts which are not substituted in the benzene ring. Instead, the fact that the $C-O$ bond length (1.21 Å) is practically the same as that found in 1,4-benzoquinone (1.22 Å)²⁰⁴,²⁰⁵ may in part be explained by remembering that the two diazo groups conjugate with four CO groups, which decreases the conjugation effect in the CO's. Finally, the CO-CO bond length is 1.54 Å, indicating the presence of a single bond, while the $CO-CN₂$ bond (1.43 Å) is slightly longer than for an aromatic bond (1.39 Å). The above evidence is sufficient to allow the conclusion that the electronic structure of this molecule may be described by a resonance hybrid between several extreme forms, the most important of which are those of 52. The distance between molecular planes in the crystal (not $\langle 2.90 \text{ Å} \rangle$ indicates that there are no intermolecular contacts.

2. 2,6-Dichloro-4-diazo-2,5-cyclohexadien-l-one

 $C_6H_2Cl_2N_2O^{206}$ (53): $R = 0.105$; standard deviations 0.013-0.018 Å and $0.10 - 0.14$ °.

 (53)

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Note Added in Proof

The zinc atom is nearly tetrahedrally surrounded by chlorine atoms, with the \hat{A} and \hat{B} at \hat{A} and \hat{B} at \hat{A} and \hat{B} at \hat{B} and \hat{B} at \hat{B} and \hat{B} and \hat{B} and \hat{B} at $\hat{B$

CI-Zn-CI angles of *105.8"* and **I** *17.0",* both with an estimated standard deviation of **0.2".** The Zn-CI distance is **2-238** A (e.s.d. 0.002 **A),** in agreement with the results obtained by Nesterova and coworkers¹¹ for p-N,N-dimethylaminobenzenediazonium tetrachlorozinc(**11) (3).**

In the present molecule you can observe a more accentuated shortening of the distance $C_{(1)} - C_{(4)}$ found in benzenediazonium chloride (1) and in o-methoxydistance **ClI,-C14,** found in benzencdiazonium chloride **(1)** and in o-methoxy- benzencdiazonium tetrachloroiron **(4).** This effect, qualitatively cxplanable by a diminution in the p character of orbitals $C_{(1)}$ in the links $C_{(1)}-C_{(2)}$ and $C_{(1)}-C_{(6)}$ and by an increase in the character p along $C_{(1)}-N_{(2)}$, has been attributed to the presence of the diazonium group¹³.

CHAPTER 4

Them *oc* **h e m i st ry of diazo compounds and organic azidest**

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1. INTRODUCTION

A search of the literature has revealed no information on diazonium compounds and very little in the way of experimental data on diazo compounds and azides, so the emphasis of this chapter is on estimation. The literature searched included *Chemical Abstracts,* **IUPAC's** *Biillefin of Tlie~niocliernistry and Tlierrtiodytiamirs,* and the two classic monographs^{1, 2}, Stull, Westrum and Sinke's *The Chemical Thermodynamics of Organic Compounds* and **Cox** and Pilcher's *Tlrevniochenristry of Organic arid* **Organometallic Compounds.**

The methods of estimation are based on additivity techniques developed by Benson and his coworkers³⁻⁵. It would not be appropriate to give a detailed discussion of additivity principles here because they have been adequately covered previously, in earlier volumes in this series^{6, 7}. Nor would it be appropriate in a review article of this size to estimate the thermodynamic properties of all possible

t Since the volume *The Chcnlistry of the Azido Groicp* **(1971)** did not contain **a** chapter on thermochernistry, it **was** decided to annex the material available on the azido group to this chapter.

compounds of interest. The main emphasis here is on the methyl, ethyl, vinyl, and phenyl derivativcs in **the** hope that thcy will form **a** basis for further **work. A** few other compounds are included.

The thermochemical properties discussed here are the heat of formation, the entropy and the hcat capacity for the ideal gas state at a temperature of 298.15 **K** (25 "C). For **the** sake of brevity, the terms molar, gas and *295.15* K will be omitted from the therniochemical symbols. Otherwise, the nomcnclature will be that recommended by **IUPAC8.** For example, the symbol for the hcat of formation of methyl azide in the ideal gas state at 298.15 K is denoted here by $\Delta_f H^0(\text{CH}_3\text{N}_3)$. Since the unit of cnergy recommended by **IUPAC** is the joule, all heats of formation will be in units of kJ/mol, followed by a value in the previously accepted unit, kcal/mol. The conversion factor is cal $=$ 4.18 J. The symbol for the standard molar entropy is S^{θ} (chemical compound) and that for the standard molar heat capacity is c_n^{θ} (chemical compound). The units for all entropies and heat capacities will be J/(mol K). **As** with the heats of formation, the value of the entropy and heat capacity in the previously acceptcd unit cal/(mol K) will also be givcn.

Most of the measurements reported in this review are good to $\pm 10 \text{ kJ/mol}$, 2 kcal/mol, for the heat of formation and ± 5 J/(mol K), 1 cal/(mol K), for the entropy and heat capacity. Most of the estimates are good to \pm 20 kJ/mol, 5 kcal/mol, for the heat of formation and ± 10 J/(mol K), 2 cal/(mol K), for the entropy and heat capacity.

II. DIAZO COMPOUNDS

A. Diazomethane

Diazomethanc is thc only diazo compound for which experimental results have been reported. Langer, Hipple and Stevenson⁹ obtained a value of $195 \pm$ 25 kJ/mol, 46 ± 6 kcal/mol, for the heat of formation of diazomethane from the appearance potential of CH_2^+ in the mass spectrum of diazomethane. Moore and Pimentel¹⁰ calculated the entropy to be 242.4 J/(mol K), 58.0 cal/(mol K), and the heat capacity to be 52.7 J/(mol K), 12.6 cal/(mol K), from statistical mechanics and the infrarcd spectrum of diazomethane.

B. Diazoethane

No measured thcrmochemical data have been rcportcd for diazoethanc. The heat of formation, entropy and heat capacity of diazocthane may be estimated by assuming that the difference between diazoethane and diazomethane is the same as thc differcncc bctwcen ethylacetylene and niethylacetylenc, equation (1).

CH,(CHN,) - H(CHN2) = CH,(CH,C=CH) - H(CH,C=CH) **(1** ¹

From equation (1) , it follows that

$$
CH_3(CHN_2) = H(CHN_2) + CH_3(CH_2C \equiv CH) - H(CH_2C \equiv CH)
$$
 (2)

By using equation *(2).* inserting values for diazomcthanc from the previous scction and valucs for cthyl- and methyl-acetylene from Stull, Wcstrum and Sinkel, **I** estimatcd thc hcat of formation of diazoethane to be 172 kJ/niol, 41.2 kcal/mol, the entropy as 285 J/(mol K), 68.2 cal/(mol K) , and the heat capacity as 73.5.J/(nioI **K),** 17.6 cal/(mol K). The results are summarized in Table 1.

C. **Diazopropylene**

No experiments have been reported on the thermochemical properties of diazopropylene. Values may be estimated by assuming that the difference between diazopropylene and diazomethane is the same as the difference between 3-vinyl-I-propyne and propyne, equation (3).

$$
CH2=CH(CHN2) - H(CHN2) = CH2=CH(CH2C=CH) - H(CH2C=CH) \t(3)
$$

From equation (3), it follows that

$$
CH2=CH(CHN2) = H(CHN2)+CH2=CH(CH2C=CH) - H(CH2C=CH)
$$
 (4)

Unfortunately, neither Stull, Westrum and Sinkel nor **Cox** and Pilcher2 list thermochemical properties of 3-vinyl-I-propyne. Furthermore, it is not possible to estimate values using group additivity because one of the groups, namely the $(C-C_d, C_t, H_2)$ group, has not been evaluated⁴.

Another solution is to assume that the difference between diazopropy!ene and diazomethane is the same **as** the difference between 1,4-pentadiene and propylene, equation **(5).**

$$
CH2=CH(CHN2) - H(CHN2) = CH2=CH(CH2CH=CH2) - H(CH2CH=CH2)
$$
 (5)

From equation **(9,** it follows that

CH,=CH(CHN,) = H(CHN,)+CH,=CH(CH,CH=CH,) - H(CH,CH=CH,) **(6)**

By using equation **(6),** inserting values for diazomethane from the previous section and values for 1,4-pentadiene and propylene from **Stull,** Westrum and Sinkel, I estimated the thermochemical properties of diazopropylcne. The results are given in Table **1.**

D. Diazotoluene

No experimental values have been reported for the thermochemical properties of diazotoluene. Valucs may be cstimated by assuming that the difference between diazotoluene and diazomethane is the same as the differexe between 3-phenylpropyne and propyne, equation **(7).**

$$
Ph(CHN2) - H(CHN2) = Ph(CH2C \equiv CH) - H(CH2C \equiv CH)
$$
 (7)

From equation **(7),** it follows that

$$
Ph(CHN2) = H(CHN2) + Ph(CH2C \equiv CH) - H(CH2C \equiv CH)
$$
 (8)

Unfortunately, neither **Stull,** Westrum and Sinke' nor Cox and Pilcher2 list the thermochernical properties of 3-phenylpropyne. Furthermore, it is not possible to cstimatc values using group additivity because one of the groups, namely, the $(C-C_B,C_t,H_2)$ group, has not been evaluated⁴.

Another solution is to assume that the difference between diazotoluene and diazomethane is the same as the difference between 3-phenylpropylene and propylene, equation **(9).**

$$
Ph(CHN2) - H(CHN2) = Ph(CH2CH=CH2) - H(CH2CH=CH2)
$$
 (9)

 \dagger Where C_d is a double-bonded carbon atom and C_t is a triple-bonded carbon atom.

[‡] Where C_B is a benzenoid carbon atom.

From cquation (9), it follows that

$$
Ph(CHN_2) = H(CHN_2) + Ph(CH_2CH = CH_2) - H(CH_2CH = CH_2)
$$
 (10)

Although no values for 3-phenylpropylene are known^{1,2}, it is possible to estimate its thermochemical properties using group additivity bccausc all the groups are available4. (While it is not noted in the text of Reference 4, it is probable that thc values for the group $(C - C_B, C_d, H_2)$ are assigned, that is, estimated by analogy with a similar group.) The results are given in Table **1.**

111. ORGANIC AZIDES

A. Hydrogen Azide

The thermochemical properties of hydrogen azidc have becn measured experimentally¹¹⁻¹⁵. They are listed in Table 2 along with similar data for organic azides¹⁶⁻¹⁹.

Experimental work on the heat of formation of hydrogen azide has been reviewed **by** Evans, Yoffe and GrayI3. (The reference for the heat of formation of hydrogen azide in another review⁴ is wrong.) Dows and Pimentel¹⁵ used statistical mechanics to calculate the entropy and heat capacity of hydrogen azide from its infrared spectrum.

B. Methyl, Ethyl, Vinyl and Phenyl Azides

No thermochemical properties have been measured for methyl, ethyl, vinyl and phenyl azides, but they can be estimated with reasonable accuracy using additivity $techniques³⁻⁵$, as follows.

The heats of formation of cyclopentyl azide, cyclohexyl azide and 2-hydroxyethyl azide have **all** been measured in the liquid state (see Tablc 2). Fagley and measured the surface tension of cyclopentyl and cyclohexyl azides and, from an empirical correlation of surfacc tension with heat of vaporization, estimated the heat of vaporization of cyclopentyl azide to be 41.8 kJ/mol, 10 kcal/mol, and that of cyclohexyl azide to be 46.0 kJ/mol, 1 **I** kcal/niol. Thus the heat of formation in the ideal **gas** state of cyclopentyl azide is 220.7 kJ/niol, 52.5 kcal/mol, and that of cyclohexyl azide is 154.2 kJ/mol, 36.9 kcal/mol.

Fagley, Klein and Albrecht¹⁶ measured the heat of formation of 2-hydroxycthyl azide in the liquid state (see Table 2). The heat of vaporization of ethyl azidoacetate may be estimated as follows. The boiling point of ethanol' is 352 K and that of cyclohexane' is very similar at 354 K. Therefore it may be assumed that the hcat of vaporization of 2-hydroxyethyl azide is ihe same as that for cyclohexyl azidc, namely, 46.0 kJ/mol, 11 kcal/mol. Thus the heat of formation of 2-hydroxyethyl azide in the ideal gas state is 140.0 kJ/mol , 33.5 kcal/mol .

All three compounds, namely cyclopentyl, cyclohexyl and 2-hydroxyethyl azides, contain the group $(C-C,H_2,N_3)$. Furthermore, for all three compounds, values are known⁴ for all the additional groups that make up each molecule. For example, 2-hydroxyethyl azide is made up of the groups $(O-C,H)$, $(C-C,H₂,O)$ and $(C-C,H_2,N_3)$. Thus from the known value of the heat of formation of 2-hydroxyethyl azide and from the known values of the $(O-C,H)$ and $(C-C,H₂,O)$ groups, a value is obtained for the $(C-C,H_2,N_3)$ group. Similarly, values for the $(C-C,H_2,N_3)$ group may **be** obtained from cyclopentyl and cyclohexyl azides. The results are given in Table 3. The results show that there is reasonable agreement between the two values from the cycloalkyl compounds, but that thcre is poor agreement with

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TABLE 2. The standard molar heat of formation, entropy and heat capacity of azides that have been determined experimentally at

thc valuc derived from 2-hydroxycthyl azide. In selecting the group value to be used in further calculations, **I** ignored the value from 2-hydroxycthyl azidc and obtained an average valuc from thc other two compounds.

TABLE 3. Heat of formation in the ideal gas state at 298.15 K (25 °C) for the $(C-C,H_2,N_3)$ group, determined from values for the heats of formation of 2-hydroxyethyl, cyclopentyl and cyclohexyl azidcs, and from other known' group values

Compound	$\Delta_f H^{\theta}$ (C-C,H ₂ ,N ₃)			
	kJ/mol	kcal/mol		
$HOCH_2CH_2N_3$ $cyclo$ -C ₅ H ₁₀ N ₃ $cyclo$ - $C_6H_{11}N_3$ Selected value	332.3 $277 - 1$ 257.5 267.5	79.5 66.3 61.6 64.0		

Ethyl azide is made up of two groups, namely, the $(C-C,H_3)$ group and the $(C-C,H₂,N₃)$ group. The value of the heat of formation of the $(C-C,H₃)$ group is known⁴ to be -42.2 kJ/mol, -10.1 kcal/mol, giving the heat of formation of ethyl azide as 225.3 kJ/mol, 53.9 kcal/mol, as shown in Table **4.**

The heat of formation of phenyl azide has bcen measured in the liquid statc (see Table *2).* The heat of vaporization may be estimated as follows. The boiling point of benzene' is 353 K and that of cyclohcxane is 354 K. Therefore it may be assumed that the heat of vaporization of phenyl azide is the samc as that of cyclohcxyl azide, namely, 46.0 kJ/niol, **11.0** kcal/mol, giving the heat of formation of phenyl azidc in the ideal gas state as 385.8 kJ/mol, 92.3 kcal/niol, as shown in Table **4.**

No expcrimental results have bcen reported for thc heat of formation of methyl azide. However, it may be estimated as follows. From the above discussion on ethyl and phenyl azidcs, the difference in the hcats of formation of ethyl and phenyl azide $is - 160.5$ kJ/mol, -38.4 kcal/mol. From the literature⁴, the difference between the heats of formation of ethyl and phenyl chlorides is -159.7 kJ/mol , -38.2 kcal/mol . It is therefore a good approximation to say that

$$
\Delta_{\rm f} H^{\theta}(\text{EtN}_3) - \Delta_{\rm f} H^{\theta}(\text{PhN}_3) = \Delta_{\rm f} H^{\theta}(\text{EtCl}) - \Delta_{\rm f} H^{\theta}(\text{PhCl})
$$
\n(11)

It may bc assumed then that

$$
\Delta_{\rm f} H^{\theta}(\text{MeN}_3) - \Delta_{\rm f} H^{\theta}(\text{EtN}_3) = \Delta_{\rm f} H^{\theta}(\text{MeCl}) - \Delta_{\rm f} H^{\theta}(\text{EtCl})
$$
 (12)

It follows from equation (12) that

$$
\Delta_{\mathbf{f}} H^{\theta}(\text{MeN}_3) = \Delta_{\mathbf{f}} H^{\theta}(\text{EtN}_3) + \Delta_{\mathbf{f}} H^{\theta}(\text{MeCl}) - \Delta_{\mathbf{f}} H^{\theta}(\text{EtCl})
$$
 (13)

From equation (13) and known values for the heats of formation of ethyl azide (Table 4) and methyl and ethyl chlorides', the heat of formation of methyl azide is 246.6 kJ/mol, 59.0 kcal/mol. This value is in very good agreement with an earlier estimate of 238.3 kJ/mol, 57.0 kcal/mol, by Evans, Yoffe and Gray¹³.

Similarly, the heat of formation of vinyl azide \vas estimated from equation (14)

$$
\Delta_{\rm f} H^0(\rm CH_2 = \rm CHN_3) = \Delta_{\rm f} H^0(\rm EtN_3) + \Delta_{\rm f} H^0(\rm CH_2 = \rm CHCl) - \Delta_{\rm f} H^0(\rm EtCl)
$$
 (14)

obert Shaw

TABLE 4. Estimation of the standard molar heat of formation, entropy and heat capacity of methyl, ethyl, vinyl and phenyl azides in the **TAULE 4.** Estimation of the standard molar heat of formation, entropy and heat capacity of methyl, ethyl, vinyl and phenyl azidcs in the

4. Thermochemistry of diazo compounds and organic azides 145

No measurements have been reported for the entropies or heat capacitics of methyl, cthyl, vinyl or phenyl azides. However ccrnparison of the values for hydrogen azide, **HN3,** with those for hydrogen isocyanate, HNCO (see 'Table *5),* suggest that it would be a good approximation to assume that the entropy and heat capacity of methyl azide is equal to those for methyl isocyanate, cquations **(15)** and (16).

$$
S^{\theta}(\text{MeN}_3) = S^{\theta}(\text{MeNCO}) \tag{15}
$$

$$
c_p^{\theta}(\text{MeN}_3) = c_p^{\theta}(\text{MeNCO})
$$
 (16)

Values for the entropies and heat capacities of ethyl, vinyl and phenyl azides were estimated similarly to those for methyl azide.

TABLE 5. Comparison of the standard molar entropy and heat capacity of hydrogen azide, hydrogen cyanate and hydrogen isocyanate in the ideal gas state at 298.15 K (25 °C)

	\mathbf{C}^{θ}						
Compound		$J/(mol K)$ cal/(mol K) Reference			$J/(mol K)$ cal/(mol K) Reference		
HN ₃ HOCN	238.7 230.3	$57-1$ $55-1$	15 20	43.5 44.3	$10-4$ $10-6$	15 20	
HNCO	237.8	56.9	20	45.1	$10-8$	20	

IV. SUMMARY OF THERMGCHEMICAL PROPERTIES OF DIAZO COMPOUNDS AND ORGANIC AZIDES

For convenience, the measured and estimated heats of formation, entropies and heat capacities discussed in the previous sections have been collected in Table 6.

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V. VALUES FOR ESTIMATING THERMOCHEMICAL PROPERTIES BY GROUP ADDITIVITY

The principles and use of group additivity have been dcscribed in considerable detail elsewhere^{3-7, 21}. A simple example was given earlier in the discussion of ethyl azide.

Values of the thermochemical properties of the groups are summarized in Table **7.**

TABLE 7. Group values for estimating the thermochemical properties of diazoalkanes and organic azides

Group	Source molecule	$\Delta_f H^{\theta}$		S^{θ}			
			kJ/mol kcal/mol		kJ/mol kcal/mol		kJ/mol kcal/mol
$C_d - C, H, N_2$	CH ₂ CHN ₂	$214 - 4$	51.3	158.0	37.8	47.6	$11 - 4$
$C_d - C_d$, H, N ₂	$CH3=CHCHN3$	$222 - 4$	53.2	166.8	39.9	53.5	12.8
$C_d - C_R$, H, N ₂	PhCHN,	$191 - 4$	45.8	199.8	47.8	62.3	149
$C - C$, H_2N_3	EtN ₂	267.5	64.0	195.2	$46 - 7$	$64 - 4$	$15-4$
$C_1 - H_1 N_2$	$CH3=CHN3$	340.2	81.4	180.2	$43 - 1$	54.3	13.0
$C_R - N_3$	PhN ₂	316.8	75.8	116.2	27.8	34.7	8.3

VI. ACKNOWLEDGEMENTS

^Iam pleased to thank Theodore Mill, Dale *G.* Hendry, David M. Golden and the Stanford Research Institute for the use of their libraries, Eileen Fellows for typing, and Kitta Reeds for editing the manuscript.

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CHAPTER 5

Detection and determination of diazo and diazonium groups

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This chapter covcrs diazoalkanes, diazoketones, diazoesters, diazonium salts and diazoaniino compounds (triazenes, tetrazenes), as well as related compounds of the general formula $RN = NX$, such as diazotates, diazocyanides, diazosulphonates, etc.; azo compounds, however, are excluded. The arenediazonium salts and the aromatic diazoamino compounds play a major role in the manufacture of azo dyes. Diazoalkanes and diazoketones are also important reagents and intermediates, respectively, in organic synthesis. Many of these compounds are rather unstable, and are conveniently handled in solution. Consequently spectroscopic methods as well as effective chemical methods of analysis arc of great significance in the elucidation of their structures and in their detection and determination.

1. DETECTION BY CHEMICAL METHODS

Nitrogen in diazo compounds cannot be detected by the usual method of melting the sample with potassium to yield potassium cyanide which is subsequently converted to Prussian blue, because under the reaction conditions, even at low temperatures, the diazo compounds decompose with nitrogen evolution¹.

A. **Diazoalkanes, a-Diazoketones and a-Diazoesters**

Diazoalkanes can be detected by the vigorous liberation of nitrogen when treated in the cold \vith dilute hydrochloric or sulphuric acid. Nitrogen evolution also takes place on heating with water. α -Diazoketones similarly liberate nitrogen when heated with acetic acid?. The stability of aliphatic diazo compounds towards trichloroacetic acid is of interest in this conncction and has been found to bc in the following order: $diazoalkanes < diazoketones < diacyl diazomethanes < diethyl diazonalonate³. Ali$ phatic diazo compounds can also be identified by the produck they yield with various reagents (equation I)?. Diazoketones and α -Diazoester

: detected by the vigorous liberation

hydrochloric or sulphuric acid. Not

water. α -Diazoketones similarly liberation

is tability of aliphatic diazo compous

is connection and has be be detected by the vigorous inferation
te hydrochloric or sulphuric acid. Nitr
h water. α -Diazoketones similarly liber
e stability of aliphatic diazo compound
this connection and has been found to
ketones < diacyldiazo

ing with water.
$$
\alpha
$$
-Diazoketones similarly liberate nitrogen when heated d^2 . The stability of aliphatic diazo compounds towards trichloroacetic est in this connection and has been found to be in the following order: diazoketones < diacyldiazomethanes < diethyl diazomalonate³. Aliconpopunds can also be identified by the products they yield with its (equation 1)².

\nRCHN₂+HCl — \rightarrow RCH₂Cl+N₂

\nRCHN₂+AroH — \rightarrow RCH₂OAr+N₂

\nR'CHN₂+R²CO₂H — \rightarrow R'CH₂OCOR²+N₂

\nR'COCHN₂+R²CO₂H — \rightarrow R'COCH₂OCOR²+N₂

\nR'COCHN₂+R²CO₂H — \rightarrow R'COCH₂OCOR²+N₂

Diazoalkancs (1-50 mg), as well as diazoketones and diazoesters, are detected by reaction with aqueous 4-(4-nitrobenzyl)pyridinium perchlorate, followed by dilution with alkali, which precipitates blue-black crystals of *N*-alkyl-4-(4-nitrobenzylidene)-1,4-dihydropyridines, which in various solvents have characteristic absorption maxima in the visible region⁴.

5. *Arenediazonium Salts*

Arenediazonium salts generally libcratc nitrogen on heating with water or with *a* dilute mineral acid². Some of them, such as those derived from aminoanthraquinones, are very stable under these conditions². With $AuCl₃$ or PtCl₁ they furnish complex salts, such as $[ArN_2]+AuCl_1^-$ and $[ArN_2]_2^2+PlCl_6^2$ whereas with $ZnCl_2$, $CaCl₂$ or SnCl₁ they yield slightly soluble double salts². When arenediazonium bromides are treated with brominc in a potassium bromide solution, they precipitate crystalline perbromides $[ArN₂]+Br₃$ which scrve in their detection and characterization⁵. Their most sensitive method of detection, however, involves coupling with a phenol in a $Na₂CO₃$ solution, or with an aromatic amine in dilute acetic acid² to yield azo dyes'.

When a weakly acidic solution of an arcnediazonium salt is added to an alkaline solution of resorufin **(9-hydroxy-3-isophcnoxazonc),** the fluorcscence of the latter is quenched as soon as the mixture becomes acidic. Addition of a $Na₂CO₃$ solution to the above mixture precipitates brown dyestuffs if diazonium ions are present, otherwise the red fluorescence is restored⁶. Fresorum (9-nydroxy-3-isophenoxazone), the nuorescence of the latter
d as soon as the mixture becomes acidic. Addition of a Na_2CO_3 solution
we mixture precipitates brown dyestuffs if diazonium ions are present,
he red f

C. Triazenes

Aromatic triazenes behave chemically in similar fashion to arenediazonium salts, but they are more stable and liberate nitrogen only when treated with strong mineral acids (equation **2)'. LIS**

$$
Ar'NHN = NAr^{2} \xrightarrow{HX} Ar'NH_{2} + Ar^{2}N_{2}^{+}X^{-} \xrightarrow{--} Ar^{2}X + N_{2}
$$
 (2)

In contrast to arenediazonium salts which couple with m-phenylenediamine in ethanolic solution to yield azo dyes, aromatic triazenes do so only on addition of acetic acid'.

Various 3-ligdroxytriazcnes have bccn detected by a spot test which involves treatnicnt with I-naphthylaniine in acetic acid, giving at once **a pink** or red-brown colour, which intensifies on gentle heatings.

II. DETERMINATION BY CHEMICAL METHODS

A. Elemental Analysis

The Kjcldalil method for the elemental dcterminntion of nitrogen cannot be applied to diazo compounds, in particular not to a-diazoketones, since addition of hydriodic acid results in nitrogen evolution even in the cold³. Zinneke has however modified the Kjeldahl method in such a manner that the total nitrogen evolved could be trapped and measured in a specially built nitrometer¹⁰.

B. Diazoalkanes and Related **Compounds**

1. Reaction with acids

Diazoalkanes easily alkylnte carbosylic acids and phcnols to the corresponding methyl esters and methyl ethers, respectively (equation 1). Thus as early as 1910 diazomcthane and diazoethane \\crc detcrinincd by adding thc diazonlkane in anhydrous ether at $0^{\circ}C$ to an excess of a solution of p-nitrobenzoic acid in ether, and after α short time titrating the excess acid with a standard 0.1 N-Ba(OH), solution to a phenolphthalein end-point¹¹. Later diazoalkanes (CH₂N₂, McCHN₂, EtCHN₂) were similarly estimated in anhydrous ether at 0° C, using benzoic acid, and back-titrating the exccss acid with a **0.1** N-NaOH or KOH solution and phenolphthalein¹². The method was modified to the micro scale, using excess benzoic acid, 0.01 N-NaOH solution as titrant and phenolphthalein¹³. Recently diazomethane was determined by reaction with benzoic acid and the resulting methyl benzoate was measured by vapour phase chromatography¹⁴. The deuterium content of CD_2N_2 was obtained by its reaction with PhCO._D in anhydrous ether, followed by determination of the deuterium content in the formed methyl benzoate, either by n.m.r. or by mass spcctronietry15.

2. Gravimetric determination

Diazoalkanes (CH₂N₂, MeCHN₂, EtCHN₂) were also determined by adding an aliquot of an ether solution *of* the diazoalkane to a cold solution *of* 3,s-dinitrobcnzoic acid. Ekess acid was removed by base and aftcr conventional work-up and cvaporation to dryness, the diazoalkanc was estimated by the **weight** of thc resulting methyl ester, and the latter's purity was assessed by a melting-point determination^{12b}.

3. Colorimetric determination

Colchiccinc aflords, with fcrric chloride, a dccp green colour, not given by its metliylated dcrivativc colchicine. This reaction scrvcs in thc determination of diazomethane. The solution of diazomethane is added to a known amount of colchiceine, and the excess *of* unrcacted colcliiccine is estirnatcd aftcr addition *of* FcCI, by colorimetric comparison with colchiceine-FeC l_a solutions of known concentration. The method gives results in good agreement with those obtained by the titration method with benzoic acid¹⁶.

The detection of diazoalkanes by the colour reaction with 4-(4-nitrobenzyl)pyridinium perchlorate has already been mentioned⁴. In its quantitative version diazoalkanes RCHN₂ (R = Me, Et, n-Pr, n-Bu, allyl) give with this reagent on addition of piperidine a red-violet colour. Measurement of its absorbance at 545 nm and comparison with a calibration curve yields the concentration of the diazoalkane⁴.

4. Gasomet ri c determination

Aliphatic diazo compounds on treatment with dilute sulphuric acid liberate nitrogen which is measured in a nitrometer (equation 1)¹⁷. The method uses up to 1-g samples, but has been adapted to the micro scale, making use of the hydrazino apparatus of Ma and Mattei¹⁸.

C. Diazonilsni Salts, Diazophenols and Diazotates

The most commonly used methods in the determination of diazonium salts include reductometric titrations, coupling titrations and gasomctric and colorimetric met hods.

1. Reductometric titrations

Most of these reductions must be carried out in the absence of oxygen.

a. Titanous chloride. Knecht and Thompson titrated benzenediazonium chloride with a TiCl₃ solution in cold dilute hydrochloric acid, using H acid (8-amino-1naphthol-3,6-disulphonic acid) as an external indicator (equation 3)¹⁹. The titration can also be carricd out by adding an excess of **a** TiCI, solution, and back-titrating the excess with a standard acid green solution in the presence of sodium tartrate¹⁹.

$$
2 \, PhN2+ Cl-+4 TiCl3+2 HCl
$$
 \longrightarrow $PhN=NNPhNH2+4 TiCl4$ (3)

The method **was** modified to the micro scale by adding an excess of the TiCI, solution to an acetate-buffered solution of the sample at room temperature, and back-titrating the excess TiCI, with a ferric **alum** solution, using ammonium thiocyanate as an indicator²⁰. It should be pointed out that this method is also applicable to azo groups and to nitroarylhydrazines. Siggia has also described this procedure in detail as **a** mcthod for the analysis of azo and nitro groups as well as hydrazines and diazonium salts²¹. Detection and determination of diazo and diazonium groups
carried out by adding an excess of a TiCl₃ solution, and b
vith a standard acid green solution in the presence of sodiu
2 PhN:⁺Cl⁻+4 TiCl₃+2 HCl - PhN=NNPh Arn acetate-buffered solution of the sample at room temperature, and
the excess TiCl₃ with a ferric alum solution, using ammonium thio-
indicator²⁰. It should be pointed out that this method is also applicable
i and t

b. *Cltroniorts chloride.* With this reducing agent diazonium salts are quantitatively reduced to the corresponding hydrazines (equation 4). Excess CrCI₂ solution is

$$
ArN2+ CI-+4 CrCl2+4 HCl
$$

added to **a** weakly acidic or neutral solution of the diazoniuni salt, and back-titration of the excess $CrCl₂$ is accomplished potentiometrically with a standard ferric alum solution 2° . This reduction method has certain advantages over the popular coupling (Section **II.C.2)** and gasometric (Scction II.C.3) methods becausc it is fast as well as quantitative. It is. howcver, also applicable to nitrobenzcnes and hydrazines.

c. *Vonodirrni(rr) sri/p/io/e.* Diazophcnols and diazonaphtliols containing **CI,** CO,H or SO,H groups \verc determined titrimetrically in **an** inert atmosphere by reduction of their acid solutions with excess **VSO,,** and back titration of the esccss reducing agent with a fcrric alum solution, using safraninc as indicator. The rcaction requires two equivalents of reducing agent **per** molc of didzo compound, and is accompanied by a quantitative liberation of nitrogen 23 .

d. *Sodium hyposulphite*. This reagent also served in the quantitative reduction of diazoniuni salts to arvlhydrazines in HCI solution **at** 0 "C. Esccss reagent was used and its excess was back-titrated with a standard acid green solution¹⁹.

e. *Hyrlriodic acid.* **A** sample of 20-30 mg diazoniuni salt is reduced with 57% HI in **a** sealed tubc betwcen 100 **anti** 300°C (equation *5)* and the liberated iodine is back-titrated with a sodium thiosulphate solution and a starch indicator²⁴. My expansion in HCl solution at 0 °C. Excess reagent was used
ack-titrated with a standard acid green solution¹⁹.

. A sample of 20–30 mg diazonium salt is reduced with 57% HI

tween 100 and 300 °C (equation 5) and the

2. Coupling reactions

This method is based upon the coupling of arenediazonium salts with aromatic aniines or phcnols to yield azo dyes (equation 6). Thc method is very selective

$$
ArN_{2}^{+}X^{-} + PhNH_{2} \longrightarrow ArN = N \longrightarrow NH_{2} + HX
$$
\n
$$
ArN_{2}^{+}X^{-} + PhOH \longrightarrow ArN = N \longrightarrow NH_{1} + HX
$$
\n(6)

because only few other compounds couple with these reagents. The determination is carried out below $0^{\circ}C$ by adding the sample to an excess of coupling agent, usually **I-phcnyl-3-mcthyIpyrazolin-5-onc** or 2,4-dianiinotoluene in acidic solution,

or R-salt (sodium 2-naphthol-3,6-disulphonate) in a $Na₂CO₃$ solution, and backtitrating the excess reagent with **a** standard 0.1 N-benzcnediazoniuni chloride solution or **a** 0.1 **N-/??-nitrobcnzenediazoniuni** chloride solution, using an external indicator **(H** acid or resorcinol in basic solution). The accuracy and precision of the method are lower than those of the gasomctric method (Section **II.C.3)** because of the slow rate of coupling and the dificulty in observing the end-point25.

Traces of diazonium salts were determined by coupling the salts with R-salt in aqueous solution or with phloroglucinol or naphthol in etlianolic solution. Tenths of a p.p.m. of the formed azo dye were then determined spectrophotometrically²⁶. Additional colorimetric determinations based on the coupling method were also reported²⁷.

The isomeric nitrobenzenediazoniuni salts in mixtures of azo coupling products were determined by coupling them with 2-naphthol. The products were chromatographed on alumina to separate the *orrho* isomer from the niixture of the *nwin* and *para* isomers; the *ortho* isomer was determined spectrophotometrically and the *neta* and *para* isomers were determined by differential spectrophotometry at two different wavelengths. Each isomer was determined with an accuracy of $2-5\frac{2}{3}$.

3. Gasometric methods

Gasometric methods are the most general and most specific for measuring diazonium salts and are widely used in industry. An accuracy and precision of about **1%** are obtainable, but the procedure is tedious and time consuming. **A** single analysis may take **up** to I hour.

Solutions of aliphatic and aromatic diazonium salts readily decompose on heating with mineral acids to furnish phenols and nitrogen. When heated with acid and cuprous chloride, chlorobenzenes and nitrogen arc obtained. The nitrogen is quantitatively liberated in these reactions and is determined volumetrically in a nitrometer. Thc method is very selective since few compounds liberate a gas under the above reaction conditions. Siggia has described in detail a general procedure and nitrometer for the determination of diazonium salts by decomposition in tlie presence of cuprous chloride in concentrated hydrochloric acid²⁹.

A simple apparatus, bascd on nitrogen evolution, has been described for the determination of aliphatic and aromatic diazonium salts, stabilized salts, diazotates and triazenes. Nitrogen evolution is effected by boiling the sample with 30% H_2SO_4 and 0.75% Cu₂Cl₂³⁰. Recently another apparatus was devised for the rapid volumetric determination of the nitrogen evolved by HCI-Cu₂Cl₂ from diazonium salts and diazoamino compounds. The reproducibility of the method is $\pm 0.3\%$ and the maximum error is $\pm 1.0\%$ ³¹.

Siggia and coworkers used a graphical kinetic approach to nnalyse mixtures of diazonium compounds, which is based on the first-order rates of decomposition in the presence of cuprous chloride. The nitrogen evolved is measured in successive time intervals. In the case of a mixture of two diazonium salts a plot of $log(100-x)$ versus *f* (where 100 is the total diazo nitrogen and *s* is the ?< diazo nitrogen evolved in time *I)* is **a** straight line as soon as most of the faster-reacting component has been decomposed. By extrapolating to $t = 0$, the composition of the mixture can be obtained³².

Diazo compounds were determined gasometrically on a semimicro scale by dcconiposing samples by light irradiation in a special apparatus containing a nitrometer. The method had a standard deviation of $\pm 0.07\%$ and the whole procedure required 30-35 min. It has been applied to diazonaphthols which cannot be easily determined by the coupling method³³.

4. Miscellaneous methods

Terent'ev and Tubina described three methods for the determination of arenediazonium salts. The first method involves a direct titration of a solution of a secondary aliphatic amine (hcxamcthylcnimine, piperidine, diethylaminc) containing sodium carbonate with the solution of the diazonium salt at $15-20$ °C until the colour formed with a 1% solution of sodium 4-amino-5-hydroxynaphthalene-2,7disulphonate persists for about 30 sec. The second method is based upon the reaction between the diazonium salt and an excess of a secondary aliphatic amine, making the reaction mixture basic, and distilling the excess aniine into aqueous **4%** boric acid and titrating it with 0.1 N-HCl. The third method involves reaction between the diazonium salt and excess m -toluidine, making the solution basic, and steam-distilling the excess amine, which is then brominated and the excess bromine is titrated with sodium thiosulphate and starch as indicator³⁴. mixture basic, and distilling the excess amine into aqueous 4%
ating it with 0.1 N-HCl. The third method involves reaction
um salt and excess m-toluidine, making the solution basic, and
excess amine, which is then brom

Arenediazonium salts dissolved in non-aqueous solvents (pyridine, dirnethylformamide) were determined by potentiometric titration with 0.1 N-sodium methoxide in benzene-methanol, using a saturated calomel electrode and a glass electrode. A probable reaction (equation 7) has been suggested. The error was less than $\pm 1.8\%$ ³⁵.

D. Triazenes and Tetrazenes

I. Gasometric methods

Aliphatic triazenes are readily decomposcd by mineral acids, liberating nitrogen (equation S), whereas aromatic triazenes require, in addition, heating in order to liberate nitrogen (equation 2). Aliphatic and aromatic triazenes were determined ArN₂+ Br⁻ + MeONa - ArN₂OMe + NaBr
 Tetrazenes
 ethods

se are readily decomposed by mineral acids, liberating nitrogen

as aromatic triazenes require, in addition, heating in order to

quation 2). Aliphatic and

$$
R^{\prime}N = NNHR^2 + HCl \xrightarrow{---}{R^{\prime}Cl + N_2 + R^2NH},
$$
 (8)

gasonietrically by conventional procedures after boiling samples with concentrated HCI **36** (see also Section II.C.3). Pierce and Rising described a micro-scale niodification of the above procedures and an apparatus for the determination of aromatic triazenes³⁷. Lohr has observed that the method of Picrce and Rising gave excellent results with pure aryl-substituted triazenes but was not rcadily adapted to samples which were solutions, slurries or pastes³⁸. He made the method applicable to samples in any physical state, using cuprous chloride in cmcentratcd HCI to catalyse the quantitative liberation of nitrogen. Two other similar proccdurcs \vhich are applicable to triazenes as well as to diazonium salts have already been mentioned^{30, 31} (Section II.C.3). The determination of 1-aryl-3,3-dimethyltriazenes by the above procedures ($Cu₂Cl₂–HCl$) has also been reported³⁹.

2. Miscellaneous methods

If the aromatic amine produced on hydrolysis of an aromatic triazene (equation 2) can be broniinatcd, this can serve as a method for the determination of aromatic triazenes. The triazene **is** dissolved in glacial acetic acid. HCI and KBr solutions are added, and the niixturc is titrated with **a** *0.5* N-KBr-KBrO, solution, using KI-starch paper as indicator¹⁰.

Small amounts of **I-aryl-3,3-dialkyltriazcnes** \vere dctemiincd by hydrolysis to furnish arenediazonium cations (equation 2) which were coupled with N -ethylnaphthylamine. The absorbance of the obtained dye was ineasurcd spcclropliotometrically at the absorption maximum. The accuracy of the method was within *5%* and it has been shown that the rate of hydrolysis **of** the triazenes depends on the pH and on the electronic character of the substituents on the benzene ring⁴¹. Similarly, triazenes stabilized with aliphatic amines were hydrolysed with aqueous acetic acid, and those stabilized with aromatic amines were hydrolysed with dilute **HCI,** and the resulting diazonium ions were coupled with excess **I-phenyl-3-methylpyrazolin-5-one,** and the excess of the latter was back-titrated with $0.5~\text{N-}NaNO_2$. The error was less than \pm 2% and a determination took up to 1 hour⁴².

I-Aryl-3,3-dialkyltriazenes were **also** determined by their hydrolysis with 0.1 N-H₂SO₄ to furnish a dialkylamine which reacted with the acid, and the excess acid was titrated potentiometrically with **0.1** N-NaOH. This procedure was modified in the case the compound contained functional groups, such as carboxyl groups, which interfered with the determination. In that case the reaction mixture was made basic and the dialkylamine formed was steam-distilled into an excess of standard acid and was then determined⁴³.

4-Amino-l-(5-tetrazolyl)-l-tctrazene hydrate was determined by hydrolysis with sulphuric acid to give amidinohydrazine which was estimated by oxidative titration with 0.1 N-KBrO₃, using iodine as indicator¹⁴. Tetramethyl-2-tetrazene was determined by titration with boron trifluoride etherate, perchloric acid in acetic acid, aluminium chloride in chloroform, or sulphuric acid in acetic acid, using crystal violet as indicator. The titration gave sharp, yet fading, colour changes at a molecular ratio of $1 : 1^{45}$.

E. Other Diazo **Compounds**

p-Mcthoxyphenyl diazothio(2-naphthyl) ether was determined as follows: hydrolysis in alkaline solution yielded **p-methoxybenzcnediazoniuin** ions and 2-napht halenethiol. The diazonium ion was couplcd with 2-naphthol and the dye formed was determined spectrophotometrically. The thionaphthol was determined amperometrically by reacting it with excess silver nitrate and back-titrating the excess of the latter with a standard ethanolic n -dodecyl mercaptan solution⁴⁶.

111. POLAROGRAPHY

The anodic half-wave potentials of several aryl-substituted diazoalkanes, such as $4-RC_6H_4CN_2Ph$, $4-O_2NC_6H_4CN_2R$ and $4-R^1C_6H_4CN_2C_6H_4R^2-4$, x-diazoketones, $4-RC_6H_4CN_2COPh$, and α -diazoesters, $4-RC_6H_4CN_2CO_2Et$, have been recorded at rotating platinum elcctrodes and their shifts with change in R have been correlated with substituent constants⁴⁷. Polarographically reduced α -diazoacetophenones gave three waves which have been attributed to three reactions⁴⁸. $NC_6H_4CN_2R$ and $4-R^1C_6H_4CN_2C_6H_4R^2-4$, α -diazoketones,
d α -diazoesters, $4-RC_6H_4CN_2CO_2Et$, have been recorded at
rodes and their shifts with change in R have been correlated
nts⁴⁷. Polarographically reduced

Arenediazonium salts are reduced at a dropping-mercury electrode. Two halfwaves have been observed arid tabulated, but they have only qualitative significance since the shapes of the waves are irregular. It has been suggested that the product of reduction is an arylhydrazine (equation **9).** In spite of the fact that the waves are too

$$
ArN_2^+ + 4e^- + 3H^+ \longrightarrow ArNHNH_2 \tag{9}
$$

complex, their hcights vary directly with the concentration. Thc first wave has a half-wave potential of less than -0.3 V. It is therefore possible to determine arenediazonium salts at -0.3 V versus SCE, using a dropping-mercury electrode. The diffusion current measured is then compared with a calibration curve of diffusion current against diazonium salt concentration⁴⁹.

Another observation is that the first half-wavc of thc arenediazonium group occurs at a potential well below that of the azo group. This finding has becn utilizcct in the determination of diazonium salts by a coupling titration in an amperometric titration cell equipped with a dropping-mercury electrode and operating at -0.3 V versus SCE. A known aliquot of 1-phenyl-3-methylpyrazolin-5-one in the cell is titrated with the diazonium salt solution. As long as coupling occurs no rcduction of the diazoniiim salt takes place and no current flows. As soon as the pyrazolinone has been consumcd, the end of the titration is indicated by a current flow which is directly proportional to the concentration of excess added diazonium salt. This new technique is more generally applicable than cither the gasonietric or colorimetric methods of determination of diazonium salts^{50, 49}.

Diazonium salts derived from aniline and from the three isomeric aminobenzoic acids exhibit two polarographic waves in the range of $0 \vee 0 - 1.5 \vee$ versus SCE. At a single pH the diffusion currents are proportional to the concentrations of the diazonium salts.⁵¹ Two distinct waves were obtained in a series of arenediazonium salts and the half-wave potentials of the first wave wcre tabulated at different pH values together with the diffusion current per unit of concentration⁵².

A German patent describes an apparatus containing a polarographic analyser for the continuous determination of diazonium salts in reaction mixtures, and for the automatic regulation of diazo couplings3.

Aromatic triazenes are reduced at thc dropping-mercury elcctrodc at much highcr voltages than arencdiazonium salts. **As** a result, it is possiblc to measurc the rate of formation and hydrolysis of triazenes by following the ratc of increase or dccreasc in diazonium ion concentration, using the technique described above¹⁹. 1-Aryl-3,3dialkyltriazenes have been determined by reduction at a dropping-mercury clcctrodc, giving well-defined waves at pH 5-7 $(E_1 = -1.0$ versus SCE). Results of the polarographic analysis were reliable and compared well with those of other methods¹³. A positive shift of **0.18** V has becn observed in the half-wave potentials of triazenes on their polarographic reduction in the presence of tetraalkylammonium ions, in contrast to negative shifts observed in the reduction of other conipounds".

The polarographic method has becn used to cstimate 1.3-diphcnyltriazenc in tlie atmosphere, which accumulates there during prodaction of cellular rubbcr. Particles collected from the air were extracted with ether and the residue obtained on evaporation of the solvent was nitrated. The triazene in the product **was** determined polarographically by recording the wavc starting from *0.8* V versus SCE and comparing it with a calibration curve of diffusion current against triazene concentration⁵⁵.

Measurement of polarographic half-wavc potentials of isomeric diazosulphonatcs indicates that the isomers form an interconvertible *syn-anti* pair 1 and 2, rather than

$$
Ar-N
$$

\n
$$
Ar-N
$$

\n
$$
ArN=N-OSO2K
$$

\n
$$
N-SO3K
$$

\n(3)
\n(1)
\n(2)

a diazosulphonate (2)-diazosulphite (3) pair. The more stable *anti* isomers 2 are reduced at a more negative potential than the labile *cis* forms 1^{56} .

The polarographic reduction of azodicarbonitrilc **(4)** to thc azodicarbonitrilidc radical anion (5) is a reversible one-electron process at $E_1 + 0.40$ V versus SCE⁵⁷.

NCN = NCN

\n
$$
NCN = NCN^{\perp}
$$
\n(4)

\n(5)

IV. SEPARATION AND CHROMATOGRAPHIC METHODS

A. Diazoalkanes, a-Diazoketones and a-Diazoesters

The aliphatic diazo group, in view of its instability, is not expected to occur in nature. Nevertheless several antibiotics containing this group have been isolated. Thus azaserine $(O$ -diazoacetyl-L-serine)⁵⁸ and 6 -diazo-5-oxo-L-norleucine⁵⁹ were isolated and purified by chromatography on pretreated alumina, followed by chromatography on carbon columns and recrystallization from aqueous ethanol or methanol. Azaserine was also analysed by Craig countercurrent distribution between water and phenol, and the norleucine derivative was analysed by paper chromatography and paper ionophoresis. Alazopeptin [L-alanyl-(6-diazo-5-oxo)-L-norleucyl- $(6-diazo-5-oxo)-L-norleucine]$ has been isolated from fermentation broths of *Streptomyces griseoplanus* by crystallization⁶⁰.

Bicyclic α -diazoketones were purified by chromatography on basic alumina and recrystallization⁶¹. Several unsaturated α -diazoesters were separated from their reaction mixtures by chromatography on Florisil and rccrystallization, whereas other similar esters decomposed on chromatography on the same adsorbent"?. Diethylphosphonodiazomethane was purified by distillation *(69-70* °C at 0.35 mm)⁶³.

0. Arenediazonium **Salts** *and Diazotates*

Arenediazonium salts are relatively rarely isolated and are generally handled in solution because of their instability and tendency to explode in the solid. They can, however, be isolated free of inorganic salts when diazotization is carried out with organic diazotizing agents. such as aniyl nitrite in ethanol, and are then precipitated by ether. They should be dried and handled carefully in small quantities only. Nowadays the solid diazonium salts used in industry have been superseded by the so-called stabilized diazonium salts, as **\yell** as by the diazotates, and, above all, by the diazoamino compounds. They can also be stabilized as coniplcx salts with inorganic salts⁶⁴.

Stabilized arenediazonium salts were separated by paper chromatography using as solvent a mixture of *95%* EtOH-concentrated HCI-distilled H20 (30 : 0.5 : **43.5),** and the spots were developed with Naphthol AS. R_f values and colours of spots were reported⁶⁵.

Arenediazoniuni chlorozincates and fluoroboratcs were scparaied by thin layer chromatography on silica gel G plates. The best solvent mixtures were $Me₂SO-$ MeCO₂H-H₂O (6 : 1 : 1) and Mc₂SO-88% HCO₂H-CHCl₃-dioxane (25 : 1 : 5 : 20). Spots were developed with a blue-dye-forming coupler (2,3-dihydroxynaphthalenc), followed by treatment with ammonia. The diazonium salts were also recovered preparatively from thick plates⁶⁶.

Arenediazonium salts \vere analysed by pyrolysis gas chromatography on Chroniosorb **103** at temperaturcs betueen **160** and 300 "C. The pyrolysis products \yere identified by infrared or laser Raman spectroscopy of the trapped peaks. For routine analysis thc samplcs were pyrolysed at 160 *"C* and the nitrogen from **ihe** pyrolysis was eluted on a column of Molecular Sieve **5A** and determined quantitatively by coniparison of nitrogcn **peak** arcas wiih calibration curvcs of injected nitrogen samples. Precision and accuracy of the method were comparable to those of analyses by other methods⁶⁷.

 $anti-Di$ azotates are obtained by adding solutions of diazonium salts to highly concentrated solutions of alkali hydroxides at room or higher temperatures. They are then isolated by either salting out or precipitation with ethanol^{es}.

C. Triazenes and Tetraxenes

Aromatic triazenes arc light yellow solids when pure but upon precipitation from a reaction mixture are often orange to deep red in colour⁶⁹. In earlier years mixed mcthyl- and dimethyldiazoaminobenzenes⁷⁰ and nitrodiazoaminobenzenes⁷¹ have been precipitated in pure form by cadmium hydroxide in methanol. More recently chromatography on alumina has been recommended for purification⁶⁹. Thus **bisthiadinzolyltriazcnes** have been separated quantitatively on alumina by elution with pyridine-glacial acetic acid $(92:8)^{72}$.

Various aromatic triazenes were separated by paper chromatography. Thus diazoaminobenzenes substituted in the benzene rings⁷³ and *para*-substituted 1-aryl-3,3dimethyltriazenes'J were separated using a variety of solvent systems and detecting agents. R_f values were given.

Several phenyltriazenes and nitrophenyltriazenes found as contaminants in ¹**-phenyl-3,3-diinethyltriazene** and in **l-(p-nitrophenyl)-3,3-dimethyltriazene** were separated from these two compounds by thin layer chromatography on silica gel using xylene and benzene as solvents, and ultraviolet detection¹⁵.

Mixtures of various tetraalkyl-2-tetrazenes were quantitatively separated and recovered as pure compounds by gas chromatography on polyethylene glycol on firebrick at 70 "C. **A** linear relationship was found between peak area and amount of compound injected⁷⁶.

V. INFRARED AND RAMAN SPECTROSCOPY

A. Diazoalkanes, a-Diozoketones, a-Diazoaldehydes, a-Diazoesters, Diarocyclopentadienes and Related **Compounds**

All the bands in the infrared spectra of gaseous CH_2N_2 , $CHDN_2$ and CD_2N_2 have been recorded and assigned⁷⁷. The strongest band in these spectra is the stretching vibration of the NN bond, which appears in the vapour phase at 2102 cm^{-1} in $CH₂N₂$, at 2097 cm⁻¹ in CHDN₂ and at 2096 cm⁻¹ in $CD₂N₂$ ^{77b}. In CCl₁ solution this band occurs in $CH₂N₂$ at 2075 cm^{-1 78}. The assignments of the major bands of the **above** three compounds have been tabulated7g. The infrared spectrum of $CH₂N¹⁵N$ has also been recorded^{77b}.

The spectra of several monoalkyl-, alkylaryl- and diaryldiazomethanes in CH_2Cl_2 . display their strongest band, namely the stretching vibration of the NN bond in the $2070-2010$ cm⁻¹ region⁷³. 4-Diazooctane displays the NN stretching vibration in CCI₁ at 2050 cm⁻¹⁸⁰, whereas in diazopropyne it appears at 2075 cm⁻¹⁸¹. Bis-(trifluoromcthyl)dia17.omethane and **bis(perfluoroethy1)diazornethanc** exhibit this band at 2137 and 2119 cm⁻¹, respectively⁸². Dicyanodiazomethane displays the band at 2140 cm-' which indicates considerable contribution from the diazonium methylid structure **6.** The high symmetry of this molecule is reflected in the simplicity of the

spectrum which contains only four bands in KBr at 2225, 2140, 1240 and 1215 cm⁻¹⁸³. Three 5-nitro-2-fury1 diazo compounds were shown to display the NN band in the region 21 10-2070 cm-' **81.** Mcrcury derivatives of diazoinethyl compounds *7* exhibit the NN band in the region of **2080-2037** cm-', whereas germanium, tin and lead substituted diazomethancs show it in the 2050-1950 cm⁻¹ region⁸⁵.

Yates and coworkers⁷⁸ and Fahr⁸⁶ tabulated the infrared spectra of a large number of a-diazoketones and compared them with those of related types of compound. The α -diazoketones show a mutual interaction between the carbonyl and diazo groups, which can be represented by resonance contributions from structures of

type **8**. This leads to lower carbonyl frequencics and to higher values of the NN diazo frequency, which by now possesses some triple bond character. Thus diazoketones of type RCOCHN₂ (R groups may be alkyl or aryl) display a strong band in the region of 2100–2080 cm⁻¹, whereas those of type R¹COCR²N₂ exhibit a strong band in the range of $2075-2050$ cm⁻¹. The intense carbonyl band for both types of diazoketone occurs in the rangc of **1645-1605** crn-I, with the exception of diazocainphor and diazoacenaphthenone which have this band around 1680 cm^{-1} . The α -diazoketones also exhibit a very intense band in the $1390-1380$ cm⁻¹ region which in some compounds is **a** composite band and in others is composed of two strong bands. Fahr also discussed the mutual interaction of the diazo and carbonyl groups in the infrared spectra of bisdiazo compounds, $N₂CHCO(CH₂)_nCOCHN₂$, and in analogous compounds⁸⁷. It has also been found that with increasing contribution from diazonium structures 8 the integrated intensity of the NN stretching band decreases⁸⁸. α -Diazoketones also display a strong band between 2.356 μ m and 2.444 μ m which is attributed to the first overtone of **the** asymmetric diazo stretching vibration".

The diazo stretching vibration of α -diazoaldehydes, RCN₂CHO (where R is alkyl), falls in the region of $2100-2080$ cm⁻¹, and the carbonyl band appears around **1690-1650** cn-'. Both bands are split, which may be due to the presence of *cis-trans* conformers, resulting from the partial double bond character of the bond between the carbon attached to the diazo group and the carbonyl carbon⁹⁰.

The NN stretching frequencies of many other α -diazoketones have been tabulated^{90a}, ⁸⁵, ⁹¹, as well as those of 2-diazo-1,3-dikctones⁹¹, α -diazo- β -oxophosphonyl compounds⁹¹ and mercury derivatives of diazomethyl compounds (7, when R is acyl or aroyl)⁸⁵. The NN stretching vibration of α , β -unsaturated α -diazoketones occurs in the region of 2101-2066 cm^{-1 a}?. The diazo and carbonyl bands of 2-oxo-1-diazocycloalkancs were tabulated⁹³. 16-Diazoandrostan-3 β -ol-17-one exhibits bands at **2075, 1650** and **1324** cn-I **91.** whereas the bands of diazoisofenchone occur at 2092, 1698 and 1335 cm^{-1 95}. The absorption frequencies of several cyclic 2-diazo-1,3dicarbonyl compounds were also recorded and these display two bands around **2180** and 2140 cm^{-1 96}.

Thc interaction bctwcen diazoketoncs and hydroxylic solvents was investigated by infrared spectroscopy⁹⁷. Infrared studies showed that diazoketones, $RCOCHN₂$, donate and accept protons in hydrogen bonding⁹³. The conformations of hindered α -diazoketones were derived from infrared spectra and dipole moments⁹⁹.

 α -Diazoesters exhibit the NN stretching frequency practically in the same region as *α*-diazoketones and *α*-diazoaldehydes⁷⁸, ^{90a}, ¹⁰⁰, ¹⁰¹, ⁶², ¹⁰², ⁸⁵.

The NN stretching frequency of diazomethylene sulphones occurs in the range of 2 145-2109 cm-I **91** and that of phosphonyl- and phosphono-diazonicthanes falls in the range of 2132-2075 cm^{-1 85}.

The strongest band in the infrared spectra of diazocyclopentadiene in CCI₄ lies at 2082 cm-I **Io3, lo4.** The NN stretching frequency of many substituted diazocyclopentadienes has been reported^{104, 105}. A complete vibrational analysis has been made for diazocyclopentadiene and its perdeuterated analogue¹⁰⁶. The diazo band of α -diazoanthrone lies at 2070 cm⁻¹ and that of the carbonyl group at 1647 cm⁻¹¹⁰⁷.

B. Diazonium Salts, Diazophenols, Diazotates, Diazosulphonates and Diazoethers

Arenediazonium salts in Nujol mulls exhibit the NN stretching frequency around 2260 ± 20 cm^{-1 108}, occasionally in the range of 2295-2110 cm^{-1 109}. It is observed that changes in the anion produce only negligible shifts in this band, whereas changes of substitution in the benzene ring result in a marked effect on the frequcncy of the band¹⁰⁹. The infrared spectra of benzenediazonium tetrafluoroborate, tetrachloroborate and tetrabromoborate have been rccorded1I0. **A** systcmatic relationship between the NN stretching frequency of methoxy-substituted benzenediazonium cations and *o+* constants has been observed, which is useful in the identification and classification of these compounds¹¹¹. The NN stretching frequencies of several substituted arenediazonium salts¹¹² and of several *o*-dialkylaminobenzenediazonium salts¹¹³ have been recorded. The infrared spectra of 20 para-substituted benzenediazonium cations measured as solid **[14gC14]2-** salts indicated that electron-donating *para*-substituents caused increase in the $Ar-N$ stretching frequency at 1400– 1300 cm⁻¹ and decrease in the NN stretching frequency, whereas electron-attracting substituents were less effective in causing such shifts. A linear correlation was also found between these frequencies and Hammett σ_p values¹¹⁴.

The ¹⁵N=¹⁵N stretching band of the ¹⁵N₂-benzenediazonium ion in a KBr pellet was observed at 2227.5 cm^{-1} , i.e. shifted 74.5 cm⁻¹ from the respective $^{14}N \equiv ^{11}N$ frequency. Its C_6H_5 ⁻¹⁵N stretching fundamental was at 1478 cm⁻¹ and the band due to the in-plane bending of the $C^{-15}N \equiv 15N$ moiety was found at 511 cm⁻¹. The corresponding bands in the unlabelled compound were at 1500 and 518 cm⁻¹, respectively115.

The NN stretching frequencies of fourteen substituted ethylenediazonium salts were observed¹¹⁶ to occur in the range of $2260-2110$ cm⁻¹.

Diazobenzenes with hydroxyl groups in *orflro* or *porn* position, the so-called diazophenols and diazonaphthols, exhibit their NN stretching frequency in **tlic** range of 2175-2015 cm-l, whereas their carbonyl frequency occurs in the range of 1640- 1560 cm-I, the latter being close to the carbonyl frequency of simple tropolones. This indicates that they exist as **a** resonance hybrid **9ll'.** With increasing contribution from the diazonium structure **9a** the diazo frequency is shifted to highcr

The infrared spectrum of potassium syn-methanediazotate has been recorded in a Nujol mull: **it** has strong bands **at** 1393, **1335,** 1140 and **I1** 15 cni-l, and additional bands at 1600, 1587, 1013, 938 and 777 cm^{-1 110}. Potassium syn-1-phenylethanediazotate has in a Nujol mull bands at 1600, 1490, 1375, 1160, 1150, 1057, 758 and 697 cm^{-1} , whereas the *anti*-diazotate has bands at 1600, 1490, 1410, 1300, 1166, 1079, 1015, 760, 695 and 611 cm^{-1 120}.

In earlier years Le Fevre and coworkers recorded and tabulated the infrared spectra of potassium syn- and *anti*-benzenediazotate, sodium p-nitrobenzenediazotate, and potassium p -methoxybenzenediazotate in the 1800-600 cm⁻¹ region. These authors were however unable to make definite spectral assignments for the NN stretching vibration¹²¹. The infrared spectra of two crystalline modifications of potassium syn-benzenediazotate and of two crystalline modifications of potassium syn-p-toluenediazotate were recorded and tabulated"?. **A** very thorough infrared study of *syn*- and *anti*-arenediazotates has been made by utilizing ¹⁵N-labelled compounds. It has been found in this study that the intense NN stretching vibration occurs around 1350 cm⁻¹ in the syn compounds and around 1375 cm⁻¹ in the *anti* compounds. The band is shifted by substituents on the benzene ring but the metallic cations have no effect on its location. The NO stretching vibration in these diazotates displays two close and strong bands in the region of $1200-1120$ cm⁻¹. It has been concluded that in both the syn- and *anti*-diazotates the bond order of the NN bond exceeds markedly that of the NO bond, and that the *anti*-diazotates have a planar *trans* structure with respect to the NN bond, while the *syn*-diazotates possess a nonplanar distorted *cis* structure¹²³.

The infrared spectra of several arenediazosulphonates were recorded and tabulated but no definite NN band assignments could be made¹²¹. The infrared spectrum of p-nitrobenzenediazo methyl ether, $p-\Omega_2NC_6H_1N=NOMe$, was recorded and tabulated'23: the NN stretching vibration occurs as **a** shoulder **at** 1508 cm-' and the NO stretching vibration as an intense band at 995 cm^{-1} .

C. Diazocyanides and Diazoisocyanides

The infrared spectra of several *cis-* and *trans-arenediazocyanides* were reported^{124, 121} and the band occurring around 2170 cm⁻¹ was assigned to the C=N stretching vibration¹²⁴. It has however been claimed that the assignments in Reference 124 were erroneous and that the stable and labile forms of arenediazocyanides were actually nitrile-isonitrile isomers 125 . Recently the original conclusions that the labile and stable pairs of arenediazocyanides were geometrical stereoisomers have been reaffirmed¹²⁶. Thus it has been shown that the unstable *cis* isomers exhibit the cyano stretching vibration at $2175-2150$ cm⁻¹, which shifts to $2195-$ 2190 cm-' as they isonicrize to the more stable *trnm* isomers. The diazo stretching vibration occurs in the *cis* isomers at 1480–1420 cm⁻¹ and in the *trans* isomers at 1460-1390 cm-' The above-mentioned band at 2195-2190 cm-l in the *lrans* isomers of **p-chlorobenzenediazocyanide** and in p-bromobenzcnediazocyanide, which is absent in the *cis* isomers, has bcen used to determine them in solution during their formation in the isomerization of the cis isomers¹²⁷.

Azodicarbonitrile **(4)** and its stable radical anion *5* exhibit infrared bands at 2217 and 2165 cm⁻¹, respectively³⁷.

The spectra of **p-chlorobenzencdiazoisocyanide** and p-bromobenzenediazoisocyanide have been reported. Their most intense band is split, and its two parts occur in the ranges of $2135-2130$ cm⁻¹ and $2100-2095$ cm⁻¹. This band has been assigned to the NC isocyanide stretching vibration while a band occurring at 1440 cm^{-1} has been assigned to the NN diazo stretching vibration. Several other bands have bcen tentativcly assigned and resonancc contributions to the structure of the diazoisocyanides havc been discussed'28.

D. Triazenes and Tetrazenes

Diazoaminobenzene displays its bands at 3.1, 3.3, 6.2, 6.6, 6.8, 6.9, 7.0, 8.0, 8.3, 8.5, 9.3, 11.2, 13.0, 13.3 and 14.5 um ¹²⁹. The infrared spectra of several triazenes have been reproduced and tabulated in the 1800-600 cm⁻¹ region, and tentative band assignments have been madc, thc NN diazo band presumably occurring around 1400 cm-I **and/or** 1600 cm-I **I?*.** The spectrum of the diazoaminothiadiazole **I0** has

('10)

been recorded. This compound and its analogues show a strong broad band in the 2700 cm⁻¹ region, indicating strong hydrogen bonding⁷².

Tautomeric equilibria of triazenes substituted in the benzene rings have been calculated by comparing integrated intensities of $14NH$ and $15NH$ bands in the 3300 cm⁻¹ region in specifically labelled compounds¹³⁰.

Infrared group frequencies of triazenes, **ArN=NNHAr,** have been assigned by laser Raman spectroscopy. Thus the band in the range of $3500-3300(w)$ cm⁻¹ was assigned to the NH stretch; the band at $1522(m)$ cm⁻¹ to the NH bend; and the band at 1178 (m) cm⁻¹ to the CN stretch¹³¹.

In a study of infrared spectra of compounds of high nitrogen content, the spectrum of **1 -guanyl-4-nitrosoarninoguanylisotetrazene (11)** was obtained. A band at

> $HN=CNHNHN=NC=N$ \overline{N} H₂ \overline{N} HNHNO **(11)**

1062 cm-1 was assigned to the NN diazo stretching vibration. Other assignments were also made132.

VI. ELECTRONIC SPECTROSCOPY, FLUORESCENCE AND PHOSPHORESCENCE

A. Diazoalkanes, Diazocyclopentadienes, a- Diazoaldehydes, a- Diazoketones and a-Diazoesters

The spectra of diazomethane and diazoethane were recorded in the vapour phase133. Their spectra as well as the spectrum of 1-diazopropane were also recorded in hexane and in other solvents¹³⁴. Their absorption begins around 550 nm, displaying **a** broad and diffuse maximum in the **450** nm region, which is of very low intensity $(\varepsilon$ 3-10). Alkyl substitution shifts the band slightly bathochromically, whereas aryl substitution causes **a** largc shift. This is manifested in the change in colour from the yellow diazomethane, to the yellow-red 2-diazopropane, and to the blue-red diphenyldiazomethane¹³⁵. At wavelengths shorter than 300 nm a second region of rapidly increasing absorption is found¹³³. The vacuum ultraviolet absorption spectrum of gaseous diazomethane and deuterated diazomethane were obtained in the 135-200 nm region'3G.

The concentration of diazomethanc in ether solution has been determined spectrophotometrically at 410 nm from the molecular extinction coefficient, ϵ 7.2¹⁵.

The ultraviolet spectra of three 5-nitro-2-furyl diazo compounds have been tabulated and recorded⁸⁴. The ultraviolet spectrum of dicyanodiazomcthane in acetonitrile shows absorption at 241 and 313 nm $(\varepsilon 12,400$ and 107)⁸³. The spectra of several diaryldiazomethanes were recorded¹³⁷.

Crystalline diazocyclopentadiene $(m,p. -23$ to -22 °C) is yellow but is red as a liquid. Its electronic spectrum in isooctanc displays *a* maximum at 298 nm ($\log \epsilon$ 4.17) and a long tailing absorption into the visible, the latter being responsible for its red colour¹⁰³.

The ultraviolet and visible absorption spectra *of* diazocyclopentadiene and several **of** its substitution products were tabulated. **As** with other aromatic compounds, substitution with conjugating substituents on the diazocyclopentadiene ring resulted in bathochromic shifts¹⁰⁴. The ultraviolet spectra of several polyphenyldiazocyclopentadienes were reported and reproduced^{105c, d}.

The ultraviolet spectra of seven α -diazoaldehydes have been tabulated. They exhibit two short wavelength bands at approximately 250 and 280 nm $(\varepsilon \sim 10^4$ and 5×10^{3} –10⁴), in addition to a long wavelength band of low intensity in the region of 360–400 nm $(\epsilon \sim 40)$. As with α -diazoketones there is a marked influence of solvents on the relative intensities of the short wavelength bands^{oob}.

The ultraviolet spectra of numerous α -diazoketones belonging to several classes have been tabulated13'. Conjugation *of* the diazo and carbonyl groups shifts the long wavelength **weak** band hypsochromically relative to this band in diazoalkanes, while the short wavelcngth intense band is bathochromically shifted. 1-Diazo-2-butanone absorbs at 247, 269 and 360 nm (ε 7650, 7110 and 21) and α -diazoacetophenone absorbs at 250 and 294 nm (ε 12,300 and 13,500). Substituted diazoacetophenones have two bands in the 240-270 and 270-300 nm regions with molecular extinction coefficients, ϵ 10⁴ \rightarrow 3 \times 10⁴ \cdot ¹³⁷. The ultraviolet spectra of several cyclic 2-diazo-1,3dicarbonyl compounds were reported⁹⁶. 16-Diazoandrostan-3₈-ol-17-one has in ethanol bands at 250 and 297 nni *(E* 4139 and 3592)94. a-Diazobenzyl phenyl ketone absorbs **at** 267, 317 and 417 nni *(E* 19,953, 5012 and IOO)l3".

The ultraviolet and visible spectra of numerous diazoacctophenones in several solvents have been reported. They all show maxima at approximately 210, 250 and 290 nm, and a low-intensity band at about 400 nm. The hypsochromic shift of the long wavelength band in polar solvents may result from an $n \rightarrow \pi^*$ transition, due to the diazo group?39. Thc interaction of diazocarbonyl compounds with hydroxylic solvents has been investigated, and it has been concluded that their behaviour in the ultraviolet and infrared are reasonably consistcnt with the possible intervention of conformational isomeric forms97. **140.** In fact, it has been shown quantum-niechanically that the ultraviolet spectra of α -diazokctones, such as diazoacetone and α -diazoacetophenone, can be interpreted satisfactorily if it is assumed that they exist in *cis* and *trans* conformations¹⁴¹.

Ethyl diazoacetate absorbs in ethanol at 249 and 377.5 nm *(E* 10,050 and 16)'42; its spectrum in water has been reproduced¹⁴³. Azaserine, another α -diazo ester, exhibits a maximum at 250.5 nm (ε 19,700) at pH 7 in water⁵⁸. The ultraviolet spectra of the following esters have been reported: **12** (in ethanol) has niaxima at 213 and 248 nm (E 4860 and 16,200); **13** (in ti-hexane) at 220 and 245 **nm** *(E* 7850 and 12,650), and **14**

(in *n*-hexane) at 217 and 245 nm (ε 8100 and 12,700)⁶². Several α -substituted α -diazoesters have two bands in ethanol in the regions of 262-264 and 360-415 nm. Thus ethyl benzyldiazoacetate absorbs at 263 and 415 nm^{102} . Diethyl diazomalonate absorbs in cyclohexane at 252 and 352 nm (ϵ 7400 and 23), and dimethyl diazomalonate absorbs in cyclohexane at 225(sh), 250 and 352 nm (ε 7300, 7900 and 22)¹⁰⁰.

B. Arenediazonium Salts, Diazophenols, Diazotates, Diazosulphones and Diazosulphonates

Arenediazonium chlorides in 0.1 N-HCI display maxima in the 220-300 nm region $(\epsilon 10^4 - 2 \times 10^4)$, which are strongly dependent on ring substituents and their position. The unsubstituted benzenediazonium chloride absorbs at 262 nm (ε 12,390)¹⁴⁴. Benzenediazonium fluoroborate in acetonitrile absorbs at 261 and 296 nm (log ε 4.09 and 3.28)¹⁴⁵. Its spectrum and that of several mono and polymethoxy derivatives have been recorded in the 200-450 nm region and assignments of electronic transitions were made by means of molecular orbital theory145. The electronic spectra of para- and meta-substituted benzenediazonium fluoroborates were rccorded and tabulated. The para series has a single band in the range of 260-380 nm (log $\varepsilon > 4.0$), and the *meta* series has two bands around 270 nm (log *E-4.0)* and in the range of 300-350 nm (log ε c. 3.5). As indicated before, the maxima are strongly dependent on substituents and their positions. Molecular orbital calculations were made to correlate these transitions with substituents¹⁴⁶. The electronic spectra of several **o-dialkylaniinobenzenediazoniurn** tctrafluoroborates show two bands at about 275 and 440 nm (log $\varepsilon \sim 3.75$ for both maxima)¹¹³.

Stabilized arencdiazoniuni salts **as** their zinc chlorides have been accurately determined spectrophotometrically by thcir maximum absorbance at 380 nm. The method does not require pretrcatment and compares favourably with results obtained by the gasometric method. Seer's law is followed at 380 nrn for aqueous solutions of $1-10$ p.p.m. concentrations. The method can be applied to other substituted benzenediazonium salts with maxima in the 350-400 nm region, since decomposition products and other impurities do not absorb in this region¹⁴⁷.

It has been reported that several arenediazonium salts exhibited fluorescence at room temperature and phosphorescence at liquid nitrogen temperatures¹⁴⁸.

The ultraviolet spectra of diazophenols and diazonaphthols have been reported. **It** has been argued from these spectra and from dipolc moment nieasuremcnts that these compounds are resonance hybrids with contributions from quinonoid resonance structures $9b^{149}$, 150 . A similar conclusion has been reached from the electronic spectra of p -aminobenzencdiazonium salts, which should be assigned the quinonoid structure **15151.**

The ultraviolet spectra of syn- and anti-diazotates in aqueous alkaline solution differ markedly. The *anti* isomers have absorption maxima between 270 and 280 nm (log *E c.* **4.1),** whereas the *syn* isomers show **a** very broad inflexion in that region, the centre of which has $\log \varepsilon$ c. 3¹⁵². The ultraviolet spectra of several *syn*- and *anti*diazotates obtained on alkaline equilibration of diazonium salts have been tabulated. **It** is observed that the spectral shifts on going from the diazonium salts to the diazotates depend strongly on the substituents and their positions. The ultraviolet spectra indicate that in strong alkali the predominant species in these equilibria is the iso $diazohydroxide¹⁴⁴$. The ultraviolet spectra of the components in equilibrium (equation 10) in the alkaline solution of p -nitrobenzenediazonium ions have bcen obtained

$$
p\text{-}O_2NC_6H_4N_2^+ \xrightarrow{\text{OH}^-} (p\text{-}O_2NC_6H_4N=N-OH) \xrightarrow{\text{OH}^-} p\text{-}O_2NC_6H_4N=NO^-+H_2O (10)
$$

by computation. Whereas the diazoniuni salt exhibits maxima at 260 and 31 **1** nm ($\log \epsilon$ 4.14 and 3.24), the corresponding *anti*-diazohydroxide, and the syn- and *anti*diazotates display their maxima at 315, 305 and 332 nm (log ϵ 3.70, 3.92 and 4.10), respectively¹⁵³.

The electronic spectra of several diazosulphones, $Ar^1N = NSO_2Ar^2$, have been recorded, and it appears from these spectra that the diazo group has the *trans* configuration¹⁵⁴. Benzyl benzenediazosulphone has maxima at 290 and 438 nm $(\epsilon 14,200$ and 129) in cyclohexane¹⁵⁵.

The electronic spectra of ten pairs of alkali salts of isomeric diazosulphonates have been recorded. In aqueous solution the *sytr-* and **anti-o-chlorobenzenediazo**sulphonates exhibit maxima at 238, 300 and 448 nm ($\log \varepsilon$ 3.7, 3.2 and 2.3), and at **228,** 292 and 428 nm (log *E* **3.8,** 3.9 and 2.3), respectively. Small differences only are found in the ultraviolet spectra of each pair¹⁵⁸. The electronic spectra of numerous *ortho-, meta-* and *para-substituted anti-benzenediazosulphonates have been tabu*lated. The unsubstituted isomer absorbs at 219, 255, 290 and 416 nm (log ε 3.93, 3.14, 4.07 and 2.25). The two long wavelength bands have been assigned to the $\pi \rightarrow \pi^*$ and $n \rightarrow \pi^*$ transitions, respectively. The effects of substitution on the spectra have also been discussed¹⁵⁷.

C. Diazocyanides and Diazoisocyanides

Electronic spectra of five pairs of isomeric *cis* and trans halo- and nitrobenzenediazocyanides have been obtained. They display three bands in the regions of 220- 270, 300-340 and 390-480 nm ($\log \varepsilon \sim 3$, 3-4 and 4). The difference between the spectra of each pair is small¹⁵⁸. The spectra of *cis-* and *trans-p-chlorobenzenediazo*cyanides have been reported. They are quite similar but the maxima of the *trans* isomer are bathochromically shifted and are of somewhat higher intensity as compared with those of the cis isomer. This indicates that the *cis* isomer has steric hindrance, inhibiting resonance¹⁵⁶. The electronic spectra of *cis-* and *trans-*diphenyl-4-diazocyanides have been recorded. They are very similar but not identical, and show the differences expected of geometrical isomers¹⁵⁹. Azodicarbonitrile (4) and its radical anion *5* display maxima in acetonitrile at 240 and 445 nni *(E* 7830 and IS), and at 222, 372 and 525 nm (ϵ 5270, 4560 and 120), respectively⁵⁷.

The ultraviolet spectra of **p-chlorobenzenediazoisocyanide** and its p-bromo analogue have been reported. They indicatc that the rigid. semipolar-bonded isocyanide group ($N^+ \equiv C^-$) is non-chromophoric in character¹²⁸.

D. **Triazenes** *and Tetrazenes*

The electronic spectra of 19 substituted derivatives of 1,3-diphenyltriazenes have been determined. They all display three bands, in the rcgions of 230-240, 280-300 and 340-360 nm $(\epsilon \sim 2 \times 10^3, 10^4 \text{ and } 2 \times 10^4)$. No conclusions could be drawn from this study about their structure in solution¹⁶⁰. The spectra of 1-aryl-3,3-dimethyltriazenes display the same bands as the above series, but it was concluded here that the compounds possess a *trans* configuration about the $-N=N-$ bond¹⁶¹. Electronic spectra of numerous triazenes have been recorded, such as those of $ArN=$

NN(Ph)X (Ar = p-anisyl, 1-naphthyl; $X = H$, Me, Ac)^{162a}; NO₂C₆H₄N= $NNHC_6H_4X$ (X = H, Me, Ph, Cl, MeO, NHAc, NO_2)^{162b}; $RO_2CC_6H_4N=$ $NNHC_6H_1X$ (R = H, Me; X = H, Me, Ph, Cl, MeO, NHAc, CO₂H, CO₂Me, NO₂)^{162c}; isomcric chlorophenyltriazenes^{162d}, and polysubstituted diazoamino-
benzenes^{162e}.

I-Aryl-3,3-bis(2-hydroxyethyl)triazenes, uscd in dyeing and printing, were analysed spcctrophotometrically. Their ultraviolet spectra in ethanolic or aqueous solution exhibit maxima from 220 to 245 nm and from 315 to 390 nm, occasionally also a masimuni in the 250-295 nni region. In acid solution, a band corresponding to a diazonium cation is observed. The conversion of these triazenes to the diazonium salts is accompanied by **an** isosbestic point, indicating a quantitative relationship between the triazenes and the diazonium compounds. The dissociation products of the triazenes (a diazonium salt and an amine) do not interfere in the determination when the long wavelength absorption band is used¹⁶³. 1-Aryl-3,3-dialkyltriazenes have been determined in concentrations down to 50 μ M (in 96% EtOH) by absorbance measurements at wavelengths between 217 and 380 nm 161.

The ultraviolet spectra of tetramethyl-2-tetrazene and tetraethyl-2-tetrazene in ethanol display maxima at 277 nin *(E* 5300) and at 2S5 nm *(E 7600),* respectively. 'The tetrazenes are basic in nature and can act as proton acceptors, affording in proton donor solvents **a** ncw band around **240** nnia5. The purity of tetra-alkyl-2 tetrazenes, which were separated by gas chromatography of their mixtures, was deterniined from thc ratio of their absorbance in basic aqueous solution at 248 and 377 11111 *ir,.*

VII. NUCLEAR MAGNETIC RESONANCE SPECTROSCOPY

A. Diazoalkanes, a-Diazoketones, a-Diazoesters, a-Diazosulphones and Diazocyclopentadienes

Diazomethane in CCI, displays a singlet at δ 3.20 p.p.m. ¹⁶⁵ As compared with allenes and ethylenes this is a large upfield shift which may be due to resonance structures with a negative charge on the carbon. This is also manifested in the nucleophilic character of diazomethanc¹⁶⁶. The chemical shifts of several diazoalkanes have been tabulated¹⁶⁷. The chemical shifts of the α -protons fall in the range of 3.0– 5.0 p.p.m., and it can be seen that they are shifted downfield when the negative cliargc on thc x-carbon can be stabilized by appropriate substituents, as, for example, in phenyldiazomethane (δ 4.58) or in ethyl diazoacetate (δ 4.96)¹⁶⁷.

The 'H chemical shifts of several 5-nitro-2-furql diazo compounds have been reported⁸¹, as were also the ¹⁹F chemical shifts of bis(trifluoromethyl)diazomethane and bis(perfluoroethyl)diazomethane⁸². Trifluoromethyldiazomethane cxhibits a quartet in the ¹H spectrum at 4.6 p.p.m. (CDCI₃, J_{HF} 4.0 Hz) and a doublet in the ¹⁹F spectrum at -54.45 p.p.m. (in CDCI₃ solution, with CFCI₃ as external standard $)^{168}$.

In the ${}^{1}H$ spectrum α -diazoacetone exhibits the methine singlet at 5.18 p.p.m. and the methyl singlet at 2.00 p.p.m. ¹⁶⁹ The α -methine protons of α , β -unsaturated a-diazoketoncs resonate in the range of 5.2-5.7 p.p.m. **y2** It has been shown that the methine proton of aryl *a*-diazoketones resonates in CCl₁ at about 5.8 p.p.m., in benzene at about 4.9 p.p.m. and in $(CD₃)₂SO$ at about 6.9 p.p.m. ⁹⁸

Nuclear magnetic resonance at diffcrent temperatures has demonstrated that ~~-cliazoaldeliydes, u.-diazoketones and a-diazocstcrs exist as equilibrium mixtures of *cis* and *trans* conformers. In diazoacetone the *trans*-methine proton resonates around 5.2 p.p.m., about 0.5 p.p.m. upfield from the cis-methine¹⁷⁰.

The methine protons of methyl and ethyl diazoacetate resonate at 4.96 p.p.m. 167 The same proton in several other similar α -diazoesters was found to display singlets in the region of 4.6-4.7 p.p.m. **62** The methine proton of a-diazo-N,N-diethylacetamide resonates at 2.40 p.p.m. ¹⁰¹ The methyl group on the double bond of ethyl 2-methyldiazoacetate resonates at 1.55 p.p.m. ¹⁰¹, whereas the methylene of the benzyl group of ethyl 2-benzyldiazoacetate resonates at 3-61 p.p.m. **Io2**

The methine protons **of** m-tolyldiazosulphone and of r-butyldiazosulphone resonate in CCI₄ at 5.07 and 4.79 p.p.m., respectively⁹³. In diethylphosphonodiazomethane this proton resonates at 4.23 p.p.m. $(J_{PCH} 11 Hz)$ ⁶³.

The aromatic protons of diazocyclopentadiene exhibit two multiplets centred at 6.7 (α -protons) and 5.8 p.p.m. (β -protons)^{104, 167}. It has been concluded that diazocyclopentadiene has very little contribution from structure **16,** but that it possesses almost complete carbanion character in the ring as indicated in structure **17Io7.**

The ¹H chemical shifts of 9-diazofluorene have been reported¹⁶⁷. The ¹H chemical shifts of several nitro- and iodo-substituted diazocyclopentadienes¹⁰⁴ and of polyphenyl-substituted diazocyclopentadienes^{105c, d} have also been reported.

B. Diozonium Salts and Diazotates

The first example of **IH** and 19F n.ni.r. spectra of an alkyldiazonium ion has been reported'68. **Trifluoromethyldiazoniethane** in fluorosulphonic acid exhibits at -60 °C in its ¹H spectrum a quartet at 6.3 p.p.m. (J_{HF} 6.1 Hz), which has been assigned to the methylene group of the **2,2,2-trifluoroethyldiazoniurn** ion **(18).** In

C F,C H, N

(18)

the ¹⁹F spectrum it exhibits the CF_3 signals as a triplet at -64.58 p.p.m. from CFCI₃ $(J_{HF}$ 6.1 Hz).

The ¹H n.m.r. spectra of p-methoxy-, p-methyl-, p-chloro- and p-nitro-benzenediazonium salts in **D20** were reported. The chemical shifts of the aromatic protons are in the region of 7-7-9.2 p.p.m., and the *ortho* coupling constants are about 9 Hz. The effects of the substituents on the ¹H chemical shifts in the diazonium ions and in the corresponding aniines and ammonium salts are in the same direction. The chemical shifts are also closely related to the Hammett σ constants¹⁷¹. The aromatic multiplets of several **o-dialkylaminobcnzenediazoniuni** fluoroborates fall in the region of $6.5-8.2$ p.p.m. 113

The methyl singlets of potassium *syn*- and *anti*-methanediazotates occur at 3.18 and 3.70 p.p.m., respectively^{172, 165}. The methine, methyl and aromatic protons of potassium *syri-1* -phenylethancdiazotate resonatc at 1 .I9 (doublet), *5.58* (quartet) and 7.05-7.43 (multiplet) p.p.m. $(J_{\text{CFL-CH}}\ 6.8\ \text{Hz})$, whereas those of the *anti* isomer have their signals at **1.36** (doublet), 4.79 (qiiarlet) and 7.09-7.44 (broadened singlet) p.p.ni. $(J_{\text{CH-CH}}$ 6.8 Hz)¹²⁰.

C. Triazenes

The triazene 19 exhibits in its ^{19}F n.m.r. spectrum a doublet centred at 4.27 p.p.m. $(J 8 Hz)$, upfield from CFCI₂CFCI₂; its ¹H n.m.r. spectrum contains a septet at

The methyl chemical shifts of several 1 **-aryl-3,3-dimethyltriazenes** occur in $CDCl₃$ in the 3.5 p.p.m. region. These triazenes exhibit hindered rotation about the $N_{(2)}-N_{(3)}$ bond because of its partial double bond character, as concluded from low temperature n.m.r. studies¹⁷³. In a series of papers Wiberg and Pracht¹⁷⁴ studied silyltriazenes 20. They have tabulated ¹H chemical shifts of numerous substituents¹⁷⁴⁸

$$
R^N = NN \begin{matrix} R^2 \\ R^3 \end{matrix}
$$

(20)

and have shown by variable temperature n.m.r. that where $R^1 = R^2 = Me$ and $R³$ = SiMe₃, SiEt₃ and SiMeEt₂, only a single methyl singlet appears and that this is due to fast fluctuations of the silyl group \mathbb{R}^3 between $N_{(1)}$ and $N_{(3)}$ ^{174a, b. The same} authors also applied low temperature n.m.r. to demonstrate that silyltriazenes, where $R^2 = R^3 = \text{SiMe}_3$ and SiMeEt_2 , exist as *cis-trains* isomers, which can be isolated at low temperatures¹⁷⁴⁰. Finally they have shown that in the last-mentioned group of triazenes hindered rotation occurs about the NN single bond, as evidenced from low temperature n.m.r. studies^{174d}. Values of free energy of activation were presented for the different processes reported in these studies.

VIII. MASS SPECTROMETRY

A. Diazoalkanes, a-Diazoketones and a-Diazoesters

The mass spectral peaks and rclative intensities of diazomethane have been tabulated. Its strongest peaks are due to $CH_2N_2^+$, N_2^+ and CH_2^+ , the relative intensitics being 96.5, 50.0 and 100.0, respectively. The spectrum is to **a** large extent similar to that of diazirine, with the exception that the $CH_2N_2^+$ and N_2^+ peaks are more intense in diazomethane¹⁷⁵.

The more abundant peaks of several α -diazoketones and their relative intensities were schematically presented. These peaks arc listed below for each of the diazoketones and the most abundant peak of each spectrum is undcriined:

From the above it is seen that all of these molecules have molecular peaks as well as N_2^+ and CHN₂⁺ peaks. Ionization potentials and bond dissociation energies of these compounds were also presented, and it was concluded that fragmentation in the mass spectrometer involved a competitive intervention of a mechanism of direct electron impact at the diazo compound, and **a** process of primary ionization of the carbonyl fragment formed on thermal cleavage¹⁷⁶.

The **mass** spectral peaks and relative intensities of several diazo-I ,3-dicarbonyl compounds were tabulated. The molecular ions are of low abundance, but the compounds give peaks at $[M-N_2]^+$ and $[M-N_2-CO]^+$, the latter being due to an electron-impact induced Wolff rearrangement. A ¹³C-labelled compound was used to investigate the migration mobility of substituents in this rearrangement¹⁷⁷. In a study of the major fragmentation patterns of diazoketophenanthrenes it was observed that the more abundant ions were M^+ , $[M-N_2]^+$, $[M-N_2-CO]^+$, $[M-N_2-CO^ H$ ⁺ and $[M-N_2$ - CO - HC = CH_]+ 178.

The mass spectral peaks of diazoesters **12-14** were reported. Molecular peaks are absent in their spectra but they exhibit $[M-N_2]^+$ and CHN_2^+ peaks⁶². The mass spectral patterns of several 2-diazo-I ,3-ketoesters wcre tabulated but, in contrast to the diazo-1,3-dicarbonyl compounds mentioned above, the fragments $[M-N_2]^+$ do not give a Wolff rearrangement'".

B. Diazophenols

The zwitterions of diazotized *o-* and p-aminophenol are volatile and give intense molecular ions and strong $[M-N_2]^+$ and $[M-N_2-CO]^+$ peaks, which are formed via the quinonoid structures of these molecules. By contrast, the *meta* isomer which cannot be stabilized through a quinonoid structure yields a complex mass spectrum. The zwitterions of diazotized anthranilic acid and diazotized sulphanilic acid did not afford molecular ion peaks178.

C. Triazenes

The mass spectra of four diaryltriazenes 21 $(R¹ = H$ or alkyl) give molecular ions of low abundance, presumably because of their rapid central cleavage to the abundant ions 22 and 23. The tautomerism in triazenes 21 $(R^1 = H)$ is also reflected in

their mass spectra, but it is not known whether the tautomers are formed before or after electron impact. Elimination of nitrogen as $(M-N_2)^+$ or $(M-R^2N)^+$ peaks is also common in these triazenes¹⁸⁰.

IX. ELECTRON SPIN RESONANCE SPECTROSCOPY AND CHEMICALLY INDUCED DYNAMIC NUCLEAR POLARIZATION

Many diazo compounds are unstable and decompose easily with evolution of nitrogen. Until recent years only chemical and indirect physical methods were available to the chemist to study which of the reactions of diazo compounds proceed by a free-radical mechanism. With the emergence of the powerful spectroscopic methods of this section, it has beconie possible to probe into this problem in a straightforward manner. This section, however, will be restricted only to those cases where the diazo compounds themselves become free radicals, leaving out the majority of cases where the primary reaction products are free radicals.

On shaking diazofluorene with sodium in ether solution it yields a deep blue, oxygen-sensitive free radical, **24,** which has been detected by e.s.r. Diphenyldiazomethane behaves similarly, yielding **a** free radical, which probabiy has a contributing structure **2S".**

In the **diazo** coupling between an arenediazoniurn fluoroborate and methanolic sodium phenoxide which yields a p -hydroxyazobenzene, it has been shown by CIDNP that the first step of the reaction involves a reversible electron transfer reaction (equation **1** I) to give two free radicals which then combine to yield the azo

$$
ArN_{2}^{+} + \bigotimes_{(26)} O^{-} \Longleftrightarrow ArN = N \cdot + \bigotimes_{(26)} H \longrightarrow O \longrightarrow ArN = N - \bigotimes_{(11)} O
$$

compound. This was proved by using a diazonium salt, ^{15}N -labelled at both positions, and detecting not only enhanced absorption of the azo product, but also enhanced absorption and ernission of the first formed aryldiimine radical **2618'.** This reaction was also studied by ¹³C and ¹⁵N CIDNP, which as above showed not only polarization of formed products, but also enhanced absorption of the ^{13}C and **I5N** resonances of the original diazonium ion. **A** mechanism was proposed for this reaction, involving diazoethers¹⁸³. In similar fashion ¹³C CIDNP of p-methoxybenzenediazonium ions (labelled with ^{13}C at $C_{(1)}$) showed, on addition of a sodium hydroxide solution, not only emission from the formed anisole but also enhanced absorption which has tentatively been attributed to $C_{(1)}$ in the original diazonium ion. This indicates participation by the above type of aryldiimine radical (26)¹⁸⁴.

A phenyldiazotate σ radical 27 has been observed by e.s.r. in the decomposition of N-nitrosoacetanilide¹⁸⁵.

¹⁵N CIDNP of the thermal decomposition of labelled diazoaminobenzene shows in addition to polarization of the formed products also enhanced absorption of *the* starting material, but only in thc central nitrogen atom. This presumably originates from recombination of free radicals **28** and **29.** Radical **28** must however be very

different from radical 26, generated in the decomposition of arenediazonium fluoroborates in alkaline solution^{182, 183}, because in the latter, atoms $C_{(1)}$, N₍₁₎ and $N_{(2)}$ are equally polarized, whereas in the diazoaminobenzene only the $N_{(2)}$ is polarized¹⁸⁶.

X. ELECTRON SPECTROSCOPY

Electron spectroscopy is used to study the sequence of molccular orbitals and their energies in molecules. It has also found analytical and structural applications, but as yet it has presumably not been used in the analysis of diazo compounds¹⁸⁷.

The ultraviolet photoelectron spectra of 2-diazopropane and diazocyclopcntadicnc have been obtained. From these spectra it has been possible to conclude that in these two compounds, as well as in diazomethane itself, the highest occupied orbital is a non-bonding $b_2(\pi)$ orbital¹⁸⁸.

The X-ray photoelectron spectrum **(ESCA)** of benzcnediazonium fluoroborate displays two nitrogen Is pcaks with a pcak separation of **1.3** cV. It has becn concluded that tlie higher binding energy pcak is associatcd \\ith the nitrogen atom directly attached to the phenyl group¹⁸⁹.

XI. DIPOLE MOMENTS

Dipole moments have contributed to structural, configurational and conformational assignments of organic molecules.

Dicyanodiazomethane has a dipole moment of **3.8** D, which indicates a considerable contribution from the diazonium methylid structure 6 to the ground state of the molecule⁸³.

Thc dipolc momcnts of diazoacetonc and its three chloromethyl dcrivatives have been determined, and from these values it has been inferred that the molecules possess a rigid planar COCHN₂ structure, where the CO and $CN₂$ groups may have either the *cis* or *trans* conformation. Diazoacetone itself exists in both conformations, but in the chloro derivatives only the *cis* conformers are present. Comparison of experimental and theoretical dipole moment values for $CH₂ClCOCHN₂$ and CHCI,COCHN, Icd to angles of 150" **and 40°,** rcspcctivcly, bctwcen the niolecular plane containing the COCHN₂ grouping, and the plane defined by C(carbonyl)- $C(alkyl)-X$ (X = Cl in the former molecule, X = H in the latter molecule)¹⁹⁰. Dipole moments of *para*-substituted phenyldiazoketones showed that the *cis-trans* cquilibrium at **tlie** COCHN, grouping is almost entircly shifted to the cis conformation, in contrast to diazoacetone¹⁹¹. Dipole moment measurements of hindered diazoketones also point to occurrence of two pscudo *cis-trans* (distorted) conformations⁹⁹. Diethyl diazomalonate has a dipole moment of 2.66 D¹⁰¹.

Of the configurational pairs of arenediazocyanides, the stable forms have the larger dipole moment and therefore possess the trans configuration, whereas the labile forms have the cis configuration¹⁹², ¹²⁴², ¹⁵⁹.

The dipole moments of 1-aryl-3,3-dimethyltriazenes indicate that they presumably have a *trans* configuration about the $-N=N-$ bond^{161, 69a}.

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CHAPTER 6

Basicity, acidity and hydrogen bonding

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I. INTRODUCTION

The electron distribution in primary diazoalkanes is often represented by resonance forms $1a-1d$.

Forms 1b and 1c dictate the reactivity of diazoalkanes to acids and electrophiles. with forms 1a and 1d playing a minor role. Proton abstraction by bases is governed by form 1a whereas reactions with nucleophiles are best rationalized by consideration of forms 1b and 1c. In the equations that follow, the choice of representation will depend on the reaction in question, and does not imply a predominance of that form in the overall electron distribution.

The structure of the discussion follows largely that of the classic review of Huisgen¹. Emphasis is however placed on quantitative evaluation of basicity and acidity. When there is a lack of thermodynamic data, as is often the case for the compounds discussed, kinetic results are examined. Qualitative examples are presented as illustrations of specific points, and do not represent a complete coverage of the subject. Extensive reference is made to comprehensive^{2, 3} and more recent^{4, 5} reviews.

II. BASICITY OF ALIPHATIC DIAZO COMPOUNDS

A. Gas Phase Basicity of Diazoalkanes

In contrast to its elusive behaviour in solution, the methanediazonium ion (2) can be directly observed and induced to react in the gas phase.

The most unequivocal methods of formation are by nucleophilic displacement processes⁶, and by fragmentation of the azomethane molecular ion⁷:

$$
N_2 + CH_3FH \xrightarrow{\text{H}} \text{CH}_3N_2^+ + HF
$$
 (1)

$$
ICH.N = NCH.1^+ \xrightarrow{---} CH. - \stackrel{+}{N} = N + CH. \tag{2}
$$

Ł.

 (2)

This latter means of generation of the methanediazonium ion lends itself admirably to the study of subsequent proton transfer reactions forming diazomethanes, by ion cyclotron resonance spectroscopy⁸. Thus at 13 eV (4.2 eV above the appearance potential of the methanediazonium ion), an increase in the pressure of azomethane results in its protonation by the diazonium ion:

$$
CH3N2+ + CH3N = NCH3 - \longrightarrow CH2N2 + CH3NH = NCH3
$$
 (3)

As is often the case for the chemistry of alkyl diazonium ions in solution, the loss of a proton is accompanied by competitive loss of nitrogen. The species so formed are characteristic fragments of the trimethyldiazenium ion (3), initially produced by a nucleophilic displacement of nitrogen.

$$
CH_{3}N_{2}^{+} + CH_{3}N = NCH_{3} \longrightarrow (CH_{3})_{2}N = NCH_{3} + N_{2}
$$
\n(4)

The proton affinity (P.A.) of diazomethane, shown to be inferior to that of azomethane by observation of the reaction given by equation (3), was found to be intermediate between those of methylamine and ammonia, by the reactions observed upon introduction of each of these bases into the azomethane system, (equations $5 - 8$).

$$
CH3N2+ + CH3NH2 - \longrightarrow (CH3)2NH2 + N2
$$
 (5)

$$
CH3N2+ + CH3NH2 \longrightarrow CH2N2+CH3NH3
$$
 (6)

$$
CH3N2+ + NH3 - \longrightarrow CH3NH3 + N2
$$
 (7)

$$
CH3N2+ + NH3 \longrightarrow x \longrightarrow CH2N2 + + NH4
$$
 (8)

Hence

$$
P.A. (NH3) < P.A. (CH2N2) < P.A. (CH3N = NCH3) < P.A. (CH3NH2)
$$
 (9)

The existence of the methanediazonium ion in the gas phase, as an intermediate or part of a transition state, in the reaction of hydrogen halides with diazomethane (equation 10), is much less evident.

The estimated upper limit of 24.8 kJ/mol for the activation energy of this reaction involving hydrogen iodide was considered to be compatible with a simple polar transition state such as 4.

$$
HI+CH_{2}N_{2} \xrightarrow{\delta_{-}} I -- - H -- - \sum_{i=1}^{n} CH_{2} \cdots N \xrightarrow{\delta_{+}} CM_{3}I + N_{2} \qquad (11)
$$
\n
$$
(4)
$$

The extreme rapidity of these reactions even at -78 °C limited the measurements of rate constants in this study to ratios based on competitive experiments. Thus at $300 K:$

$$
\frac{k_{\text{H}}}{k_{\text{H}}}
$$
 = 9 ± 2 $\frac{k_{\text{H}}}{k_{\text{H}}}$ = 4 ± 2 $\frac{k_{\text{H}}}{k_{\text{H}}}$ = 50 ± 15

Rate constants for the reactions of hydrogen chloride and hydrogen bromide with diazomethane have, however, been determined¹⁰:

$$
k_{\text{HCl}} = (7.0 \pm 1.0) \times 10^5 \text{ cm}^3 \text{ mol}^{-1} \text{ sec}^{-1},
$$

$$
k_{\text{HBr}} = (3.0 \pm 1.0) \times 10^6 \text{ cm}^3 \text{ mol}^{-1} \text{ sec}^{-1}
$$

The significance of these figures is obscured by the contribution of a wall reaction, providing an experimentally-observed negative activation energy of -1.6 kJ/mol. Experiments with deuterium chloride and diazomethane provided evidence that the first associative step is reversible¹¹. All three deuterated species of the product methyl chloride were obtained. When deuterium chloride was injected with an excess of diazomethane, which was subsequently quenched with hydrogen bromide, there was 20% deuteration of the methyl bromide. This hydrogen isotopic exchange was considered to be impossible via an intermediate complex of the form 4. The alternative complexes postulated¹¹ did not include 5, which nevertheless merits consideration as it allows for both competing substitution and deprotonation reactions, with a minimum of structural reorganization.

$$
\begin{array}{c}\n\overrightarrow{H} \\
\overrightarrow{H} \\
\overrightarrow
$$

6. **Solution Basicity** *of* **Diazoalkanes**

1. Site of protonation in 'super-acid' media

Of the two potentially basic sites in simple diazoalkanes, reaction at one-the carbon in limiting structures **(lb)** and (1c)-leads to the habitually observed products of reactions with acids¹². The alternative site—the nitrogen of limiting structures **(la)** and (Id)-is not apparcntly involved. An earlier postulate of N-protonation13 was subsequently revised in the light of more extensive data^{14, 15}.

Protonation of **2,2,2-trifluorodiazoetliane** in fluorosulphonic acid-deuteriochloroform gave **a** solution of the carbon-protonated species16 indicated by the **'H** and ¹⁹F n.m.r. spectra.

$$
F_{3}C - N = N + FSO_{3}H \xrightarrow{CDCl_{3}} F_{3}C - CH_{2} - N = N + FSO_{3}^{-}
$$
 (12)
\n
$$
\delta_{H} = 4.6 \text{ p.p.m., } qJ = 4.0 \text{ Hz} \qquad \delta_{H} = 6.3 \text{ p.p.m., } J = 6.1 \text{ Hz}
$$

\n
$$
\delta_{F^{10}} = +54.45 \text{ d} \qquad \delta_{F^{10}} = +64.58, \quad tJ = 6.1 \text{ Hz}
$$

The principal evidencc for this protonation site was the presence of a triplet in the **FI9** spectrum. Analogous C-protonated diazonium ions were formed from 1,1,1,3,3,3 hexafluoro-2-diazopropane at -70 °C in FSO₃H-CDCI₃¹⁷, and for 1-phenyl-2,2,2trifluoro diazoethane at -60°C in SO₂-HFSO₃¹⁸. From the spectral data presented for these examples, an N-protonated species cannot be so rigorously excluded.

When the diazo group is adjacent to a carbonyl group, another potential site of protonation is introduced. Thus on admixture of diazoacetone with HF-SbF, in $SO₂$ at -80 °C, protonation occurs on the carbonyl oxygen, to give a mixture of *Z* and *E* diazonium ions, *(6)* and **(7),** in the ratio 4 : 1 **19.**

For *6:*

For $7:$

 $\delta_{\text{CH}_{4}}$ = 2.05 p.p.m., δ_{CH} = 5.9, J = 2 Hz, δ_{OH} = 9.47, J = 2 Hz

 δ_{CH} = 2.19, δ_{CH} = 6.1, *J* = 1 Hz, δ_{OH} = 9.29 p.p.m.

The observation of geometrical isomers renders improbable protonation on carbon or nitrogen as in **8** and **9** respectively.

The difference in coupling constants between the methine proton and that derived from the acid in *6* snd **7** tends 10 confirm 0-protonation. In FSO,H-SbF,-SO, at -60 *"C* neither this **H-H** coupling nor, indeed, the resonance of thc OH proton could be observed due to its rapid exchange with the solvent protons.

Protonation on oxygen was also observed for **a** series of aryl primary diazoketones of general formula $ArCOCHN₂$ and for alkyl diazoketones, primary²⁰ and secondary¹⁹. The p.m.r. spectra of 2-diazo cholestan-3-one derivatives in FSO₃H-CDCI₃-SO₂ at -70 °C were interpreted as indicating protonation on carbon²¹. The principal evidcnce cited **as** argument against oxygen protonation, i.e. the dissimilarity of the chemical shift of the acidic proton at $\delta = 9.3$ p.p.m. to that for protonated ketones, is, in the light of subsequent values for oxygen-protonated diazoketones¹⁹, of $\delta = 8.2 - 9.3$ p.p.m., probably invalid. The rapid proton exchange with the solvent, on warming the solution, is also more compatible with O rather than C protonation.

2. Thermodynamic basicity of diazoalkanes

For direct observation of alkyl diazonium ions in solution, they must possess features protecting them against spontaneous decomposition into molecular nitrogen and carbocations, and must bc able to cxist in weakly nucleophilic media. In the examples cited abovc, the first criterion was fulfilled by strongly electron-withdrawing groups diminishing the stability of possible carbocationic products. The second was assured by the extrcmcly feeble nucleophilicity of the counterions SbF_6 and FSO_3 , extrapolated from their extremely weak basicity²². These conditions have so far prevented thc establishment of **a** dircctly observable equilibrium. In the highly acidic media mentioned above, the diazo compounds studied were completely protonated; in more basic solutions thc cqiiilibriuni concentrations of the diazonium ions are too low. Thus the protonation equilibrium preceding the ratedetermining loss of nitrogen, in the hydrolysis of ethyl diazoacetate²³⁻²⁵ (10) (equation 15), cannot be directly observed, even in 2.5 -molar aqueous perchloric acid²⁶.

O N₂, O N₁⁺ O OH
\n
$$
\parallel \qquad \qquad \parallel \qquad \qquad \parallel
$$
\nROC–CH+H₃O $\xrightarrow{k_1}$ ROC–CH₂ $\xrightarrow{k_2}$ ROC–CH₂
\n(10, R = C₂H₃) (15)

The overall rate constant for this reaction is given by equation (16):

$$
k_{\text{obs}} = \frac{k_2 k_1 [\text{H}_3 \text{O}^+]}{k_{-1} + k_2} \tag{16}
$$

Furthermore, as in this case $k_2 \ll k_{-1}$

$$
k_{\rm obs} = \frac{k_2 k_1 [\rm H_3O^+]}{k_{-1}} = K k_2 [\rm H_3O^+] \tag{17}
$$

where $K = k_1/k_{-1}$.

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Thus the observed rate constant contains the protonation equilibrium constant K , multiplied by k_2 , the rate constant for the displacement of nitrogen, which has been shown to be nucleophile assisted for this substrate²⁷. Although the value of k_2 is inaccessible directly, its sensitivity s to the strength of the participating nucleophile n_i , as expressed by the Swain-Scott equation²⁸, has been determined²⁷. Identical *s* values for two nucleophilic substitutions indicate a common sensitivity of the transition states to reactant properties. This is equated by the Leffler postulate^{29, 30}, **to** a common position of the transition states along the reaction coordinate, the position being governed by thc difference in free energy between reactants and products. The activation frce energy is determined by this energy difference and **a** nunibcr of factors, including stcric, solvational and entropy changes dependent on the transition-state structure. **It** has bccn proposed that sutstituent effects on structurally related diazonium ions are small¹². If one accepts this proposition, it follows that thc reactions of water with structurally related diazonium ions, displaying similar sensitivity to substitution by other nucleophiles, should have similar activation cncrgies. This rescmblancc in transition state structures (with idcntical entering and lcaving groups) implies similar free cnergics of reaction dominated by the frce energy of formation of molecular nitrogen, and common interactions unique to the transition state. Hence thc **s** values for thc hydrolysis of **10** and diazoacetonc (11) of 0.3 and 0.4²⁷ imply approximately equal values of k_2 for nitrogen displacemcnt. Consequently, comparison of the overall second-order hydrolysis rate constants of **10** and **11** givc a mensurc of the relative acidity of their corresponding diazonium ions.

$$
\log k_{\rm obs}(11) - \log k_{\rm obs}(10) = -pK(10) + pK(11) \simeq 1.3 \tag{18}
$$

This type of comparison cannot bc justifiably cstendcd to the hydrolysis of primary aryldiazosulphones Arg_2CN_2H , which although following the same mechanism have a somewhat higher s value³¹.

An estimate of the acid dissociation constants of alkancdiazoniuni ions can be made in some cascs, wherc the deprotonation is lcss rapid than the compctitivc loss of nitrogen. This situation is characterized by rate-determining protonation. When $k_2 \ge k_{-1}$, equation (16) becomes:

$$
k_{\rm obs} = k_1 [\mathrm{H}_3 \mathrm{O}^+] \tag{19}
$$

An increasing concentration of additional base may, however, sufficiently accelerate the deprotonation, so as to make the second step partially rate determining, i.e. the dependence of thc observed rate constant on acid concentrations may change. This situation was rcalized in the hydrolysis of diazoacctatc anion **(12)** in thc presence of hydroxide ion³², and in the hydrolysis of 2-diazo butan-3-one (14) and ethyl diazopropionate³³ (13) in the presence of acetic acid/acetate buffers.

Analysis of the kinetic data^{32, 31} provides values for k_{-1}/k_2 , whicl, with certain approximations, allow estimates of k_{-1} . Combination with the known k_1 value gives limiting values of $K < 10^5$ mol/l, $K < 5 \times 10^3$ mol/l and $K < 1$ mol/l for the protonated forms of 14, 13 and 12, respectively, i.e. $pK < -5$, $pK < -3.7$ and $pK < 0$.

A more involved analysis of kinetic data for general acid-catalysed hydrolyses of diazo compounds can lead, in principle, to an estimate of the pK_a 's of the conjugate acids. The significance and limitations of this approach will be discussed in Section II.B.6.

3. Acid-catalysed hydrogen-deuterium exchange

Primary deactivated diazoalkanes of the form RCHN₂ where $R = -CO_2R^{23}$, 24 , $-COR³⁵⁻³⁸$, $-SO₃R^{39, 40}$, $-CF₃⁴¹$, establish an acid-base equilibrium with aqueous acid, prior to nitrogen loss. This equilibrium is principally demonstrated by a kinetic isotope effect $k_{\text{H},0}/k_{\text{D},0} \approx 0.4^{+2}$, or by hydrogen-deuterium exchange in a deuterated solution, either in the diazo compounds or products derived therefrom. For simple diazoalkanes, precise kinetic measurement is difficult due to their extreme acid lability, and so the second probe is more often used.

In the case of diazomethane, the results obtained appear somewhat confusing. Whereas it undergoes exchange in a two-phase system of alkaline D_2O and dibutyl ether or tetrahydrofuran^{13, 41}, in a system of chloroform- D_2O there is no exchange unless catalytic quantities of a weak acid, such as phenol, ammonium chloride or benzoic acid, are added¹⁵. In dioxane- D_2O , reaction of diazomethane with an acid or phenol gives the trideuterated ester or ether respectively¹⁵. Monodeuteration on carbon is observed in deuterated acetic acid⁴⁶, while a pre-equilibrium is reported to be established with benzoic acid in toluene¹⁷. In strong aqueous acid solution, only one deuterium is incorporated^{11, 48}.

That the exchange occurs via the conjugate base of diazomethane (see Section IV) cannot be excluded *a priori*. However, the implication of acidic catalysis by weak acids, but not by D_4O^+ , suggests that the determining factor is the strength of the conjugate base of the catalyst. Certain diazocarbonyl compounds, 12³², 14¹⁹ and benzoyl phenyldiazomethane (15)⁵⁰, ⁵¹ have been shown to undergo protonation by water and, at least in the first two cases, subsequent regeneration by the hydroxide ion. For bases stronger than water, deprotonation is favoured over nucleophilic substitution. In support of this argument, the methanol isolated after the hydrolysis of diazomethane in dioxane- D_2O is monodeuterated on carbon when the hydrolysis is carried out at pH 3, trideuterated for pH 13, and contains a mixture of mono-, di- and trideuterated species when the reaction is performed between pH 6 and 7^{32} .

The same phenomenon has been more precisely defined by a study of the methanolysis of diazobutane⁵³. In neutral or acidic deuterated solution there is no exchange, however in strongly basic solutions (up to 1.8 M-NaOMe) exchange is observed to the extent of 87%. The approximate inverse proportionality of the observed overall rate to methoxide ion concentration (except at concentrations 1.0 m) suggests establishment of a partial pre-equilibrium dependent on base concentration. The alternative interpretation of base-catalysed exchange may be supported by the deviation from linearity of the ratio CD₂HOMe : CDH₂CMe against [OMe⁻] at base concentrations of approximately 1.5 M. At these concentrations, however, it is difficult to decide what confidence can be placed in kinetic equations, not corrected for variations in the activity of the substrates⁵⁴.

4. Base-catalysed decomposition of N-nitrosourethanes and ureas

The alkanediazonium ion-diazoalkane equilibrium in basic solution can be established by preliminary formation of the acidic species by another reaction sequence. This situation is realized, in principle, by the decomposition in basic solution of N -nitrosourethanes⁵⁵ (16) and ureas⁵⁶ (17).

The implication of diazotates **(18) as** intermediates in this scheme follows from their isolation in the reactions of differently substituted N-nitrosourethanes in ethereal solution^{53, 57}, and their subsequent conversion into previously obtained products in basic prctic solvents. The exact status of the diazohydroxide **(19)** and

the diazonium ion, either frcc or in an ion pair, as precursors **of** the diazoalkane will be discussed later. The reactions of the diazohydroxide leading to **the** formation of the diazoalkane compete with its decomposition to nitrogen, solvated hydroxide ions and carbonium ions. Thus **the** yields of substitutcd diazoalkancs will be dependent not only on the substituent effect on their stability, but also on that of the

alternative carbonium ions. For example, in the formation of pyrazoles derived from diazoalkenes⁵⁸, the observed effect of electron-withdrawing groups of increasing the pyrazole yield is compatible with destabilizing the incipient carbonium ion, or in facilitating development of the negative charge in the transition state for diazoalkane formation. The effects of alkyl substituents, complementary in stabilizing carbonium ions and inductively destabilizing the diazo group, are evident for examples 20, 21, 22, 26 and 27 of Table 1.

$RR^{\prime}CN_{2}$	R	R'	$[OH^-]$	Yield $(\%)$	Reference
20	H	н	unknown	100	57
21	$n-Pr$	H	3 _M	60	53
22	n -Octyl	H	3 _M	31	53
23	Cyclopropyl	н	3 _M	0	59
24	Ph	H	unknown	100	57
25	Vinyl	н	3 _M	100	53
26	n -Hexyl	CH ₃	3 M	3 ^a	60
27	Cyclohexyl		3 M	$\mathbf 0$	53
28			3 _M	100 ^a	53

TABLE 1. Yields of diazoalkanes from basic hydrolysis of the corresponding diazotates

^a From deuterium-incorporation experiments.

Separation of the two effects can apparently be made for examples 23, 24, 25 and 28. The stability of the cyclopropyl carbinyl cation⁶¹ accounts for its preferential formation, and, on the contrary, the relative instability of the cyclopropyl cation⁶² allows the establishment of the diazo compound 28 as an intermediate in the reaction. The stability conferred to alkyl carbonium ions by substituent phenyl or alkyl groups does not, however, determine the direction of reaction of the diazohydroxides. The quantity of phenyl- and vinyl-diazomethane formed may rather be due to a dominant stabilization of the transition state in the base-catalysed elimination, reflecting a conjugative stabilization of the product diazo group. The above argument⁵³ is entirely dependent on the assumption that the benzyl and allyl cations are as readily formed in water as is the 2-octyl cation which is generated in preference to 2-diazooctane. Solvolysis rates in nucleophilic solvents, where the bimolecular contribution to the rate-determining step is unknown⁶³, offer debatable support to this thesis. Gas-phase heats of formation from the corresponding alkanes⁶⁴ (ΔH_f Ph-CH₂ = 854 kJ, H₂C=CH-CH₂ = 940 kJ, C₃H₇-C-CH₃ = 906 kJ) in the absence of

solvational energies offer a support only for the benzyl cation. For this particular example the alternative diazoalkane product was obtained at unspecified base concentration ('five- to six-fold excess'); at a lower, equally unspecified base concentration, there was minor formation of benzyl alcohol.

That the diazoalkane formation is dependent on base concentration and strength is supported by the observation that when cyclohexane diazotate is quenched in

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methanol rather than water, providing a more basic medium, diazocyclohexane is observed⁵³. It is worth noting, in this respect, that cyclopropyldiazomethane (23), completely by-passed in the aqueous diazotate hydrolysis, is prepared by reaction of the corresponding N-nitrosourethane with a concentrated solution (2 **M)** of sodium triethylene glycolate in triethylene glycol at -25 °C⁵⁹. For the octane-2-diazotate system, the inferred yields of diazooctane increased from **3%** to **15%** with increasing base concentration⁶⁰. The effect of base on the partition between the two reactions in methanol was examined by Kirmse for a series of cyclopropyl^{65, 66, 67} and alkenyl⁶⁸ substituted diazoalkanes.

The yield of allene **30,** derived from diazoalkane *29,* increased from 0.52% to 80% for both *cis* and *trans* starting ureas (28) with increasing base strength. The importancc of the diazonium ion, either free or in an ion pair as the immediate precursor of the diazoalkane, as depicted above, is controversial. **As** regards secondary alkyl diazotates, it has been established from results with optically pure and labelled subfrom a solvated diazotate species⁶⁹

Thc influcnce of diazo group and carbonium ion stability on the reaction course implies that both **sets** of products are formed from a common intermediate-the diazotate species favoured by Moss⁶⁹ or free diazonium ion preferred by Kirmse^{56, 70}.

An alternative explanation is that an equilibrium between the two intermediates is established rapidly relative to the two competing product-forming reactions

18-Oxygen labelling experiments showed, among other things, that this equilibrium is not established in secondary diazotate hydrolysis⁶⁹. As the benzyl and alkyl cation-forming processes are presumed to have activation cnergies lower than, or equivalent to, those for secondary cations, the equilibrium must also be excluded in these cases (assuming equivalent diazonium ion stability). This conclusion is **in** part supported by the observed base catalysis. The equilibrium will be displaced to the diazohydroxide form by increasing base concentration. This, combined with the expected first-order dependence of the proton abstraction reaction on base, would give an ovcrall zero-order dependence for diazoalkanc formation, which is not observed. The irrefutable evidence, from trapping expcrimcnts7" for frce diazoriium ions in the systems, is restricted to cyclopropyl diazonium ions, which may arise from diazocyclopropane rather than diazohydroxide prccursors.

The partition of the two reaction sequences appears to occur at the diazotatediazohydroxide stage. The mechanism for thc basc-cntalysed cliniination of watcr has been formulated as a two-step process⁵⁸.

$$
\begin{array}{ccc}\nR & H \\
\searrow & H \\
\se
$$

This however would appear to present no advantngcs over *n* concertcd scheme similar, in essence, to that proposed for the base-catalyscd cleavagc of **15** ".

It should be noted that whcn diazoalkancs are decomposed in basic alcohol ~~l~itionsS3 diazoniuni ions, and not dinzotic estcrs analogous to **19,** arc tlic carbonium ion precursors. This follows from the inverse-order dependence of reaction rate on base concentration (rather than zero order).

In conclusion the partition of diazotatcs in basic solution, between diazoalkane and carbonium ion products, appears to be dominated by the base strength, and substituent effects on the carbonium ions, rather than by diazonium ion acidity.

5. Acidic species alkylated by diazoalkanes

The acid strength and reaction conditions necessary to protonatc diazoalkanes vary largely with their substituents. Whereas diazomethane methylates readily and smoothly carboxylic acids and phcnols, their reactions with diazoesters and diazoketones require gentle warming. Likewise, dialkyloxonium salts are efficiently methylated in dichloromethane^{72} according to:

$$
(CH3), \tilde{O}H[SbCl6]+CH2N2 \xrightarrow{\qquad} (CH3), \tilde{O}[SbCl6]+N2
$$
\n(27)

In contrast, the efficiency of quaternization of protonated amines, by diazoacetophenone **(31)** in alcohol, varics apparently as a function of the acid strength as indicated in Table **2.**

$$
\begin{array}{ccc}\n & 0 & N_2 \\
\hline\nN-H + Ph-C-C-H & \xrightarrow{\text{II}} \quad Ph-C-CH_2-N \\
\hline\n(31)\n\end{array}
$$

TABLE 2. Efficiency of quaternization of pyridiniurn **salts** by **31** (equation 28)

Acid	$pK_{\rm a}$ ⁷⁴	Yield ⁷³ (%)
-H	5 ₂	88
\overline{H}	$5-4$	68
CН, Ņ7 н	595	27
ÑŦ Ĥ	4.90	21 ^a

^a*See* discussion in Scction II.B.6.

Diazomethane reacts at room temperature with alcohols⁷⁵, the efficiency and rapidity of reaction increasing with substitution of electron-withdrawing groups in the latter. Reduction of the already feeble acidity of the hydroxyl group by solvation may prevent reaction; thus trichloroethanol is methylated in heptane but not in ether⁷⁶. On the other hand, incrcasc of the proton acidity by prior complexation of the hydroxyl oxygen with an electrophilic species dramatically increases the efficiency of the process^{75, 77}.

$$
R-\overset{\circ}{O}-\overset{\circ}{A I}(\overset{\circ}{O}R')_{3} \xrightarrow{\hspace{2cm} \text{CH}_{3}N_{1}} \hspace{0.1cm} R-\overset{\circ}{O}-\overset{\circ}{A}I(\overset{\circ}{O}R')_{3} \xrightarrow{\hspace{2cm} \text{N}} R\overset{\circ}{O}CH_{3}+\overset{\circ}{A}I(\overset{\circ}{O}R')_{3} \hspace{0.1cm} (29)
$$
\n
$$
\overset{\underset{\circ}{\downarrow}}{\overset{\circ}{H}} \overset{\circ}{CH}_{3}
$$

Similar catalysis, preferably by boron trifluoride, renders aniines sufficiently acidic to undergo this reaction^{78, 77}.

Whcreas certain carbon acids such as acetylacctonc react with diazoalkanes through their enolic forms⁷⁹ (equation 30), sufficiently acidifying but non-enolizable substituents such as sulphone^{50, 81} and nitro⁸² groups allow direct methylation on carbon.

6. Kinetic basicity of aliphatic diazo compounds

In contrast to the paucity of thermodynamic data related to diazoalkane basicity, thcre exists a wealth of relevant kinetic data. This results from the facility of rate mcasurements by following the evolution of nitrogen, or currently by monitoring thc diazo ultraviolet absorptions. **A** comparative analysis of the data is not, however, as straightforward as thcir measurement. The second-order rate constant may be simply a measure of the protonation rate (equation 19) as for an AS_E2 reaction.

Alternativcly, for **A-2** and **A-1** mechanisms, it may be expressed by equation (17) where k_2 is respectively dependent on or independent of the nature and concentrations of nucleophiles present. **Examplcs** of all thrce kinetic fornis and intermediate situations³², ³³, ⁸³ have been found for acid-catalysed diazoalkane hydrolysis¹². Consequently in the ignorance of the exact mechanism, caution should be exercised in drawing conclusions as to structure/reactivity relationships. For example, **in** the reaction of protonatcd pyridines with **31** (Table *2),* the reduced yield for the 2-picolinium ion relative to the pyridinium homologue may represent not only its inferior acidity, but also a diminished nucleophilicity of its conjugate base due to steric hinderance. Independent evidence suggests that this diazo compound hydrolyses by an A-2 mechanism³⁸, although a conflicting opinion has been expressed⁸⁴.

The increasing reactivity in the order **diazomethane-diazoethane-diazopropane** towards phenol in toluene⁸⁵ may not only be a reflection of increased basicity, but also of more rapid nitrogen loss from the intermediate diazonium cation. The inertness of a diazo compound to acid, for example diazomalonitrile to sulphuric acid in anhydrous acetonitrile⁸⁶ allows no extrapolation to its behaviour in a protic nucleophilic medium.

Similarly, the acidity scale developed on the basis of reaction rates of a series of acids with weakly basic diazoalkanes⁸⁷⁻⁸⁹, such as 32 and 33, in chlorinated hydrocarbon solvents, is not necessarily **a** function uniquely of acid strength. Such a situation would be realized only if the protonation of the diazoalkanes in question were rate determining, or if the decomposition of the diazonium ion formed were unassisted. In the **first** case the observed rate constants would indeed be measures *of*

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kinetic acidities; in the second, in that they would depend on the equilibrium conccntration of the diazonium ion, they would be measures of thermodynamic acidity. In view of thc quantitative convcrsion of **1 ,I** ,I **,3,3,3-hcxafluorodiazo-2-propane (34)** into its conjugate acid in highly acidic media¹⁷, and the subsequent S_N2 type displacement of nitrogen from the latter by the fcebly nuclecphilic fluorosulphonate anion, both these possibilities scem to be unlikely. It should be noted that the observed linear $\log k_{\text{obs}} - pK_a$ plots³⁷⁻⁸⁹ are consistent also with the remaining A-2 type mechanistic possibility. Both the thermodynamic strength of the acid, and the kinetic nuclcophilicity of its conjugate basc which are included in the rate constant, can be related to the aqucous acid strength by linear free energy relationships.

The data in Table 3, and the subsequcnt discussion, refer to hydrolysis of diazoalkanes where protonation by the hydrated proton has been shown to be rate detcrmining. For several substrates, data for various sets of reaction conditions are included as minor differences in the latter can significantly affect the former.

Thc rcactivity of differently substituted stabilized diazo compounds within the same class towards strong acids in non-nucleophilic solvents has been correlated with the $N=N$ stretching frequency in the i.r. spectrum¹⁰⁷. The hydrolysis rates for series of primary diazoketoncs and sulphones have been similarly correlated108. Generally, the higher thc absorption frequency, the more the electron distribution described by the form $\tilde{C}-\tilde{N} \equiv N$ is predominant in the ground state¹⁰⁹. This is favoured by clcctron-withdrawing groups attached to the carbon; ips0 *facta,* these decrease the acid lability. For reactions that proceed by a rate-dctermining protonation step there is a general trend in this sense¹⁰⁸, but extrapolation from one class of compounds to another allows only qualitative comparison¹⁰⁵ of ground-state effects by this means.

In the series PhCN₂R the protonation rate decreases in the order $R = H$, Ph, CN,

0 0

CF₃, Si(Ph)₃, (EtO)₂PO-, (Ph)₂PO-, R'O-C-, −R'C-. This variation represents well thc expected different degrees of stabilization of the ground state, by clectron withdrawal (except for the triphenglsilyl substituent which may also provide a steric retardation of the reaction).

It is interesting to note that for the unique series whcre the effcct on the protonation rate of an alkyl group relative to a hydrogen substituent has been directly measurcd, the protonation of **1 -p-nitrophenyldiazoethane** is slower by a factor of almost 10 than that of **p-nitrophenyldiazoinethane.** The substitution of a methyl group has apparently less effect for diazoketones. Whereas protonation rates of primary diazoketones are normally inaccessible, that of diazoacetone has been measured by hydrogen isotopic exchange rates¹⁰¹. Furthermore, in the hydrolysis of **7-(2'-diazo-)acetyl-bicyclo[2.2.l]hept-2-ene (37,** the protonation is rate determining due to acceleration of the subsequent nucleophilic displacement of nitrogen by participation of the neighbouring double bond¹⁰⁰. In both cases, protonation is

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Solvent: A, water; B, dioxane-water 60-40 v/v; C, dioxane-water 2-1 v/v; D, dimethylsulphoxide-water 4-1 w/w; E, water 2-2 M in acetonitrile; F, ethanol-water 93.8% w/w; G, dioxane-water 40-60 v/v; H, toluene.

^a lonic strength mol/l.

⁶ Extrapolated from Arrhenius plot.

· Measured with one acid concentration only.

⁴ Reaction followed by nitrogen evolution.
• Value extrapolated from data for differing salt concentrations.

If the acid added, p-toluene sulphonic acid may not be completely dissociated in this system.

" Measured by hydrogen isotopic exchange.

three- to four-times more rapid than for methyl-substituted diazoketones measured under the same conditions. This behaviour contrasts with the accelerating influence of alkyl substitution on the overall reaction of diazoalkanes with phenol in toluene⁸⁵.

The stabilizing effect of benzoyl relative to carboethoxy groups is dependent on the nature of other substituents present. Thus the effcct of their different electronwithdrawing capacity is diminished by introduction of an additional stabilizinggroup:

$$
\frac{k_{\text{CH}_4\text{CN}_3\text{CO}_2\text{CH}_3}}{k_{\text{CH}_4\text{CN}_4\text{CO}_4\text{H}_4}} \simeq 570 \qquad \frac{k_{\text{C}_4\text{H}_4\text{CN}_3\text{CO}_4\text{CH}_4}}{k_{\text{C}_4\text{H}_4\text{CN}_4\text{CO}_4\text{H}_4}} \simeq 1.0 \tag{31}
$$

The more rapid proton-deuterium exchange of primary arylsulphonyldiazomethanes relative to the corresponding diazokctones¹⁰, though not a direct measure of protonation rate, indicates a superior stabilization provided by the carbonyl group.

The variation of protonation rate with nuclear substituents in several series of aryldiazoalkanes give normally good correlations with Hammett σ values (Table 4). The rho (ρ) values derived therefrom are generally in the range -1.5 to -1.8 , The rho (ρ) values derived therefrom are generally in the range -1.5 to -1.8 , indicating a large sensitivity to electron availability. The correlation in the diazofluorene¹¹¹ (36) examples with Hammett parameters modified as by Taft¹¹² is indicative of the non-benzenoid nature of the aromatic substituent site rather than any peculiarity of the proton transfer. On the other hand, the correlation for diaryldiazoketones⁸³ with the σ^+ parameters of Brown¹¹³ is inconsistent with the simply visualized C-protonalion mechanism. It should be noted however that as the rate constants for the hydrolysis of these compounds arc successfully correlated with those for aryldiazophosphonate esters¹⁰⁵, which are themselves correlated with ordinary σ values, there appears to be a lack of sensitivity to the constants used. Likewise while the one electron oxidation potentials of several series of arylsubstituted diazo compounds correlate with $\sigma^{+ (114, 115)}$, for diaryldiazoalkanes they are also proportional to the logarithms of the protonation rates, themselves dependent on σ .

Other traditional indices of transition-state structure in relation to reactants and products are not immediately definitive. It is a commonly held thesis that primary hydrogen isotope elfects for proton transfer attain their maximum value for approximately 'symmetric' transition states, with a continuous decline for increasingly exoor endo-energetic reactions116. One might then expect decreasing overall solvent isotope effects for increasingly 'uphill', hence slow, reactions for a series of related compounds. This is not apparent from the data presented, although within a series of substituted aryldiazoketones there is some relation⁸³. With specific reference to 3-diazobutan-2-one **(14)** and ethyl diazopropionate **(13),** despite a difference in reaction rate of 28, the measured solvent isotope effects are *2.5* and 2.46 respectively⁹⁸.

Furthermore, an analysis of this global isotope effect, from experiments in $H₂O-D₂O$ mixtures¹¹⁷, for transfer to the two substrates respectively, produces almost identical fractionation factors¹¹⁸, $\phi_1 = 2.1 \pm 0.03 \times 10^{-1}$ and 2.08×10^{-1} , and $\phi_2 = 7.78 \pm 0.04 \times 10^{-1}$ and $7.85 \pm 0.12 \times 10^{-1}$, for the proton 'in flight' and residual protons of the lyonium species. The latter results give values of α_i , a parameter describing the position of the transition state along the reaction coordinate, of 0.27 and 0.29 , surprisingly close to that of 0.30 for the much more reactive diazotate anion¹¹⁷ (12). This similarity is extended to the values of the exponent α_B in the Brönsted relation

$$
\log k_{\text{HA}} = \alpha_{\text{B}} \log K_{\text{HA}} + C \tag{32}
$$

Here K_{HA} is the dissociation constant of an acid, whose reaction with a substrate has a rate constant k_{H_A} , and α_B , describing the sensitivity of the latter to the former,

• Solvent index given in Table 3.
• Data for three series of substituents in positions 2, 3 and 4.

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has been interpreted in the same fashion as α_i above. The values in question, for reaction with carboxylic acids, are 0.61 ± 0.03 , 0.59 ± 0.04 and 0.51 ± 0.03 for 14, 13 and 12 respectively. The identity of these values, within experimental error, for the diazoketone and ester is surprising in vicw of their reactivity difference and the valucs. reflecting the degree of proton transfer in the transition state, appear incompatible with such a therinodynamically unfavourable reaction. (The acid dissociation constants of the diazonium ions formed are $\lt 10^5$, $\lt 5 \times 10^3$, $\lt 1$, whereas K_{IIA} of the acid catalysts 6×10^{-4} .) The disparity between the values of α_B and α_i is a result of the greater acid strength of the hydronium ion relative to carboxylic acids. For proton transfer from the stronger acid, the transition state is expected to be niorc 'rcactant like', in the oft-quoted terms of the Hammond postulate¹¹⁹ or its more quantitative formulation^{29, 30}, and to be characterized by a lower value of α .

A more refined free energy of activation-free energy of reaction relationship derived by Marcus¹²⁰, equations (34) and (35), has met with some success in the analysis of diazoalkane protonation data. For the reaction

S+HA
$$
\xrightarrow{\text{wr}}
$$
 S ... H-A $\xrightarrow{\text{wr}}$ S⁺-H ... A⁻ $\xrightarrow{\text{wr}}$ S⁺-H+A⁻ (33)
\n(I) (II) (IV)

$$
\Delta F_{\text{HA}}^{\text{+}} = W^{\text{r}} + \Delta F^{0\text{+}} \left(1 + \frac{\Delta F_{\text{R}}^{0}}{4\Delta F^{0\text{+}}} \right)^{2} \quad \text{for } 1 > \frac{\Delta F_{\text{R}}^{0}}{4\Delta F^{0\text{+}}} > -1 \tag{34}
$$

$$
\Delta F_{\text{HA}}^* = W^r \qquad \qquad \text{for } -1 > \frac{\Delta F_{\text{R}}^*}{4\Delta F^{0+}} \tag{35}
$$

$$
\Delta F_{\text{IIA}}^{\text{+}} = W^{\text{r}} + \Delta F_{\text{R}}^{\text{o}} \qquad \qquad \text{for } \frac{\Delta F_{\text{R}}^{\text{o}}}{4\Delta F^{\text{o}+}} > 1 \tag{36}
$$

- $\Delta F_{\mathrm{H}\Lambda}^{\ddag}$ is the free encrgy of activation observed for the proton transfer from **HA** to **S,** i.e. from state (I) to state **(IV).**
- *Wr* is the energy that must be supplied to form the 'reactant complex', i.c. state **(11)** from state **(I),** and involves diffusion, desolvational energy and distortion of bonds, if ncccssary to allow the proton to be transferrcd.
- ΔF_R° is the standard free-energy difference within the reaction complex, i.e. ΔF° state (III) – ΔF° state (II) and is ostensibly equal to $\Delta F_{\text{HA}}^{\circ}$ – $\Delta F_{\text{SH}}^{\circ}$.
- ΔF^{0*} is the activation free-energy within the reaction complex for the case where $\Delta F_{\rm R}^0=0.$

From this equation, the rate of change of α with differing ΔF_R^{α} is inversely proportional to ΔF^{0+} which measures the inherent reactivity of the system. The data for the hydrolysis of 12 catalysed by phenols for which $\alpha = 0.74$, carboxylic acids where $\alpha = 0.51^{121}$, and a series of protonated amines displayed in Figure 1¹²², allowed calculation of ΔF^{0+} , and hence the other parameters of equation (35).

The corresponding parameters could be calculated from the data for hydrolysis of **13** and **14**, with inclusion, however, of the data for catalysis by H_3O^{+34} , and for hydrolysis of diphenyldiazomcthane (35), catalysed by carboxylic acids and phenols⁹³. The collected parameters are listed in Table *5.*

The first points of note are the values of $\Delta F_{\rm SH+}^0$, i.e. the free energies of the reactions:

$$
SH^+ + H_2O \xrightarrow{} SH + H_3O^+ \tag{37}
$$

In all cases they are unacceptably large: for 14 and 13 they correspond to dissociation constants 10¹⁰ mol/l and 10^s mol/l, respectively, compared to independently estimated values of $\lt 10^5$ mol/l and $\lt 5 \times 10^3$ mol/l (Section B.2). Furthermore, the variation of the values derived for protonated diazoacetate anion indicates that these values do not have an absolute thermodynamic significance. Their significance

FIGURE 1. Predicted and experimental relations between log k_{HA} for the hydrolysis of 12, and log K_{HA} of the catalysing acid. The \bigcirc are points for oxygen acids, the \times for protonated amines. The solid lines are predicted from equation (35) by insertion of the parameters listed in Table 5. Reprinted with permission from M. M. Kreevoy and S. Oh, J. Amer. *Chem. Soc.*, 95, 4805 (1973). Copyright by the American Chemical Society.

is rather in the context of the reaction complex, in that differently organized reactant complexes, in solvational and hydrogen-bonded terms (state II), may be formed with different types of acids. Such a variation with acid type should dictate a variation in ΔF^{0*} and, more obviously, in W^r. Although the values of ΔF^{0*} obtained by the

Acid HA	Base			ΔF^{0*} (kJ/mol) W^r (kJ/mol) $\Delta F_{\rm SH}^0$ (kJ/mol)
$R^1R^2R^3NH^+$	12	5.9	58	-28.5
$RCO2H$, ArOH	12	17.0	34	-14.7
$RCO2H, H3O+$	14	80	71	-24.8
$RCO2H, H3O+$	13	8·0	60	-19.8
RCO ₃ H, ArOH	35	5.5	72.3	$-25+2$

TABLE 5. Parameters of the Marcus formalism for diazoalkane hydrolysis

above analyses are probably underestimated due to simplifications made in the algebraic derivation^{123, 124, 125}, their significance, relative to the term W^r for related compounds, is nevertheless a valid point of comparison.

The magnitude and variation of the values of W^r are the most significant features of the data in Table 5. They are markedly superior to the free energies estimated for the diffusion together of substrate and acid, and desolvation of the latter, i.e. between 37.2 kJ/mol¹²¹ and 42 kJ/mol¹²⁵. The difference in hydrolysis rate for 13 and 14 is seen thus to stem primarily from the different work terms W^r . It is also this term that represents the stabilizing effect of the aryl groups in 35. The major

kinetic consequence of these large work terms is that no matter how thermodynamically favoured be the proton transfer, depending on the acid strength, the rate can never approach the diffusion limit¹²². Differing interpretations of the high values of W^r in terms of hydrogen bond formation in the reaction complex^{93, 122}, and nonequilibrium solvated species³⁴ appear to be complementary. Evidence for the former is derived from an analysis of the Hammett ρ values for solvolysis of 35. This value can be separated by use of certain approximations into ρW^r and ρc terms⁹³, the latter referring to the actual proton transfer, state $(II) \rightarrow$ state (III) . It was found that the ρW^r term was by far the more important, indicating that the aryl substituent effects are primarily felt in the H-bond formation. The differences in hydrolysis rates of 12, 13 and 14 can be attributed to differing degrees of electron delocalization from the diazo group into the adjacent carbonyl substituents. The reflection of these differences in the work terms W^r and not ΔF^{0+} , for the same acid types, implies that this electron delocalization is reversed before the proton transfer occurs34.

This interpretation demands an intermediate with negative charge located on the diazo carbon, but available for hydrogen bonding, and with a consequently rearranged solvation sphere. The energy changes along the reaction profile **are** illustrated schematically in Figure **2.**

A corresponding energy diagram for nitroalkane deprotonation is included. This represents the same electron reorganization phenomenon, but in a reversed sense in that the tetrahedral carbanionic species (state **111)** formed by proton transfer, relaxes into the π -delocalized anion¹²⁶. In this case, kinetic evidence for the schemc includes substituent effects of different magnitude on the transition state, leading to the carbanionic structure **(111)** and on the final statc (IV) of the nitronale anion. This behaviour leads to values of $\alpha_B > 1$ and α_B , inexplicable conventionally^{126, 124} in terms of 'degree of proton transfer'. Similar results of anomalous α values for diazonium ion deprotonation would be expected, but are as yet unobtainable.

The above diagram shows how differing diazo compounds, despite differences in overall rcactivity towards acids, can have identical indices of transition-state position along the reaction coordinate within the 'reaction complex'.

111. LEWIS BASICITY OF ALIPHATIC DIAZO COMPOUNDS

A. Kinetics *of* **Electrophilic Addition**

The problems involved in defining a basicity scale for diazoalkanes towards Lewis acids ate niore complex than those conccrning proton basicity. **As** before, instability of the adducts formcd preclude direct determination of their equilibrium concentrations. Kinetic measurements may be subject to specific solvational effects. Uncertainty may exist as to the nature of the rate-determining step, or even of the electrophilic species. Whereas the effective catalytic species in the $Cu^H (CF₃CO₂)₂$ catalysed decomposition of diazoalkanes has been claimed¹²⁷ to be Cu^T , a contrary conclusion has been drawn concerning catalysis by $Cu^{I}[P(OR)_{3}]_{n}$ complexes¹²⁸. HgCl₂ reacts with diphenyldiazomethane (35) more rapidly than HgI₂ in an etherhexane solution¹²⁹, but less rapidly than HgBr₂ in tetrahydrofuran, and does not react at all in ether12g. **A** qualitative order of reactivity of a series of elcctrophiles towards 35 in ether-hexane or acetonitrile has been established and is reproduced in Table *6.*

The comparative reactivity of differently substituted diazoalkanes towards a single clectrophile appears to follow their reactivity tonards proton sources. Introduction of a *p*-methoxy substituent increases the reaction rate of diphenyldiazomcthane (35) with HgCl₂ in tetrahydrofuran, and, whereas 35 is unaffected by the less electrophilic alkylmercury chlorides, diazomethane reacts readily in their presence130. Decomposition **of** p-toluyldiazomethane by zinc(r1) halides is more rapid than that of phenyldiazomethane 131 .

TABLE 6. Relative rates of catalysed diphenyldiazomethane decomposition in ether-hexane or acetonitrile solution. Reprinted with permission from **D.** S. Crumrine, T. **J.** Haberkamp and D. J. Suther, *J. Org Chem.*, 40, 2274 (1975). Copyright by the American Chemical Society

Rate ^a	Catalyst
Very fast Fast Moderate Slow No reaction	AlCl ₃ , TiCl ₄ , SnCl ₄ , BF ₃ , O(Et) ₂ $HgCl_2$, $Hg(NO_3)_2$, $CuBr_2$, $FeCl_3$, ZnI_2 , $CuIP(OPr-i)_3$, $CuCN[MeCN]^b$ Hgl_2 , ZnCl ₂ , CuCl[ether], CuCl ₂ .2H ₂ O, SnCl ₂ .H ₂ O, Hg ₂ Cl ₂ [MeCN] CuCN, CrCl ₃ .6H ₂ O, AlCl ₃ .6H ₂ O $Hg_2Cl_2[$ ether], CdCl ₂ , Zn(CN) ₂ , Cr(OAc) ₃ .6H ₂ O, NiCl ₂ .6H ₂ O, $Co(NO_3)_2.6H_2O$, $CoCl_2.6H_2O$, $MgCl_2.6H_2O$, $CaCl_2$, LiCl, LiBr, NH ₄ Cl

^a Very fast, reacts completely in less than 1 min. Fast, reacts in less than 15 min. Moderate, reacts in **less** than **4** h. **Slow,** reacts while standing for **24** h.

Brackets indicate solvents in which the reactivity order differed. Taken from Reference 129.

The related zinc(n) halide-catalysed decomposition of diphenyldiazomethane and diazofluorene in acetonitrile, proceeds by a rate-determining association¹³² (equation 38) ated zinc(n) halide-catalysed decomposition of diphenyldiazomethane and
rene in acetonitrile, proceeds by a rate-determining association¹³² (equa-
 $Ph_2CN_2+ZnX_2 \xrightarrow{\text{slow}} Ph_2CN_2^+ Zn-X_2 \xrightarrow{-N_1} [Ph_2C^+ Zn-X_2]$ (38)
trast, the $Co(NO₃)₂.6H₂O, CoCl₂.6H₂O, MgCl₂.6H₂O
st, reacts completely in less than 1 min. Fast,
eaches in less than 4 h. Slow, reacts while standing
indicate solvents in which the reactivity order di
the effective effective effective energy of
the effective energy of the inter-
et al. (a) had been identified in the
equilibrium with the reectants before$

$$
Ph_2CN_2+ZnX_2 \xrightarrow{\text{slow}} Ph_2CN_2^*Zn-X_2 \xrightarrow{\text{rapid}} [Ph_2C^*Zn-X_2]
$$
 (38)

In contrast, the adducts formed by each of these two substrates with copper(1i) bromide are in equilibrium with the reactants before decomposition¹³³.

$$
Ph_2CN_2+CuBr_2 \xrightarrow{\longrightarrow} Ph_2CN_2^+Cu-Br_2 \xrightarrow{\text{slow}} [Ph_2C^+Cu-Br_2]
$$
 (39)

For the former reaction (equation 38 where $X = Cl$), it is interesting to note that the Hammett *p* values for diazofluorene and diphenyldiazomethane substituents are quite dissimilar $(-2.45 \text{ and approximately } -1.0)$, despite the minor $(7.60d)$ difference in rate. **Also** striking is the observation that the order of kinetic catalytic efficiency, $ZnCl_2 < ZnBr_3 \ll ZnI_2$, does not follow the equilibrium Lewis acidity (towards dibenzacridine 132).

A situation somewhat less complicated kinetically was encountered in the reaction of diphenyldiazomethane with peroxybenzoic acid¹³⁴.

The effect of substituents in the aryl groups of the diazoalkane on the reaction rate, as reflected by theis ρ value of -0.96 , is less marked than on the corresponding proton transfer reaction. That the electrophilic attack was indeed by oxygen and not by a proton, was supported by the similarity of the kinetic isotope effect $k_{\text{RO3H}}/$ $k_{\text{R03D}} = 1.33$ and of the solvent dependence of reaction rate to those of other peroxide oxidations. **A** similar mechanism is probably operative for the reaction of peroxybenzoic acid with diazoketones¹³⁵. Other examples of electrophilic oxygen attack on diazoalkanes are provided by the reactions of ozone136 and singlet oxygen137.

6. Reactions with Electrophilic Carbon

1. Reactions with esters

The reactions of diazoalkanes with electron-deficient carbon centres in carbonium ions³, acid derivatives^{138, 5} and carbonyl groups^{3, 139, 140} have been reviewed frequently. In the last-mentioned category, there exist few examples of reaction with the carbonyl group of esters. Whenever electron-withdrawing substituent groups such as trihalomethyl¹⁴¹ or nitro¹⁴² are attached, the reactivity of the carboxyl group *is* sufficiently enhanced to allow addition and subsequent cpoxide formation **as** for ketones.

Addition of diazo groups to esters may be favoured by other circumstances, for example the close proximity of the two groups in the same molecule¹⁴³⁻¹⁴⁵.

As these reactions werc llowevcr carried out **in** the presence of base, necessary for **the** initial nitrosolactam dccomposition, the possibility of nucleophilic catalysis, by attack on the terminal diazonitrogen146, cannot be cxcluded (see Section **V.C).**

2. Reaction with aliphatic diazonium ions

zonium ion with an unprotonated diazo group^{147, 148}. Similar intramolccular proximity can lead to the rcaction of **311** aliphatic dia-

Another diazoalkane-diazonium ion reaction has been proposed to explain azine formation from dipotassium diazomethane disulphonate in water 113 . There are, however, significant differences between this example and the above. Here the alkyl diazonium ion is presumed to react intermolecularly with the diazo group (in water), and through nitrogen rather than carbon as the electrophilic site. There exists ample precedent, howcver, for the formation of azines by dircct rcaction of two diazo compounds (Section V.C).

C. Site of Electrophilic Attack

1. Reaction with aromatic diazonium ions

Aromatic diazonium ions are **well** cstablishcd *iis* clectrophilic partncrs in dinzoalkane reactions¹⁵⁰. Whereas, as is usually observed, the majority of products implicate carbon as the nucleophilic site (equation 47), in the case of diazomethane, the formation of a tetrazole by-product indicatcd attack at nitrogen (equation 45).

+P- + f+ Ar-N=N CHz-N, Ar-N=N-CH,-N, + + CI- % CI c1- **H** \ 1 Ar- NH -N=CHCI **(46)** + CI- **(47)** C=N

It has, however, been pointed out¹⁵¹ that this product could be generated by a 1,3-dipolar cycloaddition of diazomethane *to* the arenediazochloride, in equilibrium with the separated ions.

2. Reactions with carbenes

The only unequivocal examples of electrophilic attack at the terminal nitrogen of diazoalkanes involve carbenes. The formation of azincs in thc thermolysis of diazoalkanes does not necessarily involve the prior formation of a carbene. Indeed, the singlet carbene from phenyldiazocthane forms styrene rather than the azine observed as the major thermolysis product¹⁵². The dimerization could result from an excited state-ground state interaction¹⁵³, or from an electrophilic attack of one diazo group

on another (Section V.C.) Kinetic evidence exists, however, that the formation of benzophenone azine from diphenyldiazoniethane is in part due to **a** carbene m echanism¹⁵⁴, and in part to dimerization¹⁵⁵.

Carbenes with dihalo-¹⁵⁶ and a variety of other¹⁵⁷ substituents react preferentially at the nucleophilic carbon site of diazoalkanes, including diphenyldiazomethane and diazofluorene. An increase in the bulk of the substituents at this carbon, however, **may** direct reaction to the nitrogen terminus.

R^2	40 $(\frac{9}{6})$	41 $(\%)$
H	85.0	
α -Naphthyl	52.0	7.0
α -Naphthyl	6·1	14.5

TABLE 7. Variation of product distribution, with increasing bulk of aryl substitucnts, in reaction *(SO)*

This is illustrated by the results in Table 7, for the influence of α -napthyl substituents on the distribution, bctwecn olefin **(40)** and azine **(41)** products, for the reactions of diaryldiazoalkanes with dihalocarbenes (equation 50).

Accordingly, reaction of phenyl-x-naphthyl-methylene with its diazo parent yields azine rather than olefin¹⁵⁸. Thus steric hindrance to reaction at carbon could well explain the preferential formation of azine in the reaction of diphenyldiazomethane with its corresponding carbene, and especially carbenoid, species¹³². Compilation of reactions of diazoalkanes with electrophiles with subsequent retention⁵, or loss¹⁵⁹ of nitrogen exist, and will not be repeated hcre. Thc rapidly developing area of substitution at a metallated diazo carbon⁵, where the mechanism is uncertain^{160, 161}, will **also** be left untreated. Discussion of these rcactions and of interactions of diazo compounds with metals in general will be found in Chapter 7 on 'Complex Forniation' in this volume. H,C=N-+(Ph),C- Na⁺ ——→ HC=N-N- Na⁺+(Ph),CH (51)
H₂C=N++(Ph),C- Na⁺ ——→ HC=N-N- Na⁺+(Ph),CH (51)
H₂C=N=N-+(Ph),C- Na⁺ ——→ HC=N- Na⁺+(Ph),CH (51)

IV. PROTON ACIDITY OF DIAZOALKANES

Diazomethane is deprotonated by triphenylmethylsodium¹⁶², methyllithium or phenyllithium¹⁶³ in ether at $0 °C$ to form the corresponding alkali metal salts.

$$
H_2C = \ddot{N} = N^{-} + (Ph)_1C^{-} Na^{+} \longrightarrow H\ddot{C} = N = N^{-} Na^{+} + (Ph)_2CH
$$
 (51)

Whereas hydrolysis of these salts with alcohol-water at room temperature produces whereas hydrolysis of these salts with alcohol-water at room temperature produces

a 35% yield of diazomethane, hydrolysis at -15° C with a concentrated, weakly

acidic buffer solution (of ammonium chloride or potassi acidic buffer solution (of ammonium chloride or potassium dihydrogen phosphate) yields a colourless diazomethane isomer. This 'isodiazomethane', originally considered to possess a nitrilimine^{164, 165} structure (42), is now identified as an N-aminoisocyanide (43) on the basis of spectroscopic¹⁶⁶ and chemical¹⁶⁷, ¹⁶⁸ evidence.

$$
H_{2}C=\overset{\dagger}{N}=N^{-}\xrightarrow{\qquad -11^{+}}\qquad H-\overset{\dagger}{C}=\overset{\dagger}{N}=N^{-}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}=\overset{\dagger}{N}=N-H \iff H-C=\overset{\dagger}{N}-\overset{\dagger}{N}-H
$$
\n
$$
+C=\overset{\dagger}{N}-\overset{\dagger}{N}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{N}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}=\overset{\dagger}{N}-\overset{\dagger}{N}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}=\overset{\dagger}{N}-\overset{\dagger}{N}-H
$$
\n
$$
+C=\overset{\dagger}{N}-\overset{\dagger}{N}-\overset{\dagger}{N}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{N}-\overset{\dagger}{N}-H
$$
\n
$$
+C=\overset{\dagger}{N}-\overset{\dagger}{N}-\overset{\dagger}{N}-\overset{\dagger}{N}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{C}\xrightarrow{\qquad \qquad}H-\overset{\dagger}{N}-\overset{\dagger}{N}-H
$$
\n
$$
+C=\overset{\dagger}{N}-\overset{\
$$

Diazomethane is re-formed immediately and quantitatively by treatment of 'isodiazomethane' with solid potassium hydroxide in ether163. Reaction with a buffered aqueous neutral solution, however, generates forniylhydrazine and hydrazine¹⁶³. Thus 'isodiazomethane' is the kinetically controlled product of protonation of the diazomethane anion, while under equilibrating conditions diazomethane is favoured. The possibility that the experimental conditions of the isomerization, involving the presence of a heterogeneous basic system, may upset the equilibrium in favour of diazomethane **(20)** is unlikely as *20* is in part reformed from **43** in a basic aqueous buffered solution¹⁶³. SCF-MO calculations using α LW $[533; 3]$ gaussian lobe basis set do not reproduce the energetics of the above situation¹⁶⁹. Thus **43** is calculated to be **136 kJ** more stable than **20.**

Substituted diazomethanes do not form isolable salts, but evidence of their deprotonation by organometallics is to be found in the reaction products. Diazoethane on treatment with methyllithium yields acetaldehyde N-methylhydrazone **(44),** and **4,5-dimethyl-l,2,3-triazole (45)170.** The corresponding triazole is the unique isolable product of the reaction of phcnyldiazornethane with biphcnylylphenyl-ketone disodium¹⁶².

The hydrazone product is readily visualized as arising from nucleophilic attack of the base on the terminal nitrogen of the diazo system. This process, also observed to the extent of 15% for the diazomethane-methyllithium system¹⁷¹, will be discussed in Section **V.C.** Thc triazole product has been postulated to bc the result of the nucleophilic attack of the diazoethyl anion on its conjugate acid¹⁷⁰, however a 1,3-dipolar cycloaddition of diazoethane onto the hydrazone formed concurrently is **a** more appealing possibility.

As would be expected, ethyl diazoacetate is more acidic than simple diazoalkancs in that it can be lithiated with butyllithium at $-110^{\circ}C^{172}$. The salt formed is, however, thermally unstable at temperatures above $-50^{\circ}C^{172}$ as witnessed by previous unsuccessful attempts at preparation¹⁷³. Below this temperature it is sufficiently reactive to undergo metal exchange with trimethylsilyl and tributyltin chloride and to condense with ketone carbonyl groups.

$$
N_{2} O
$$
\n
$$
H-C-C-OEt
$$
\n
$$
M_{2} O
$$
\n
$$
LIC-C-OEt
$$
\n
$$
LIC-C-OEt
$$
\n
$$
CH_{3})_{3}Si-C-C-OEt
$$
\n
$$
(CH_{3})_{3}Si-C-C-OEt
$$

This latter reaction can be conveniently effected by treatment of a mixture of diazoester and ketone with butyllithium at -65 °C. Similar types of condensations occur under the influence of weaker bases, but with more electrophilic carbonyl bonds. Thus ethyl diazoacetate condenses with aldehydes in ethanolic potassium hydroxidc solution at room temperature17J. These conditions **are** unsuitable for the corresponding reactions with ketones, due to the unfavourable equilibrium established between reactants and product adduct. Reaction does proceed, however, with small ring or *a*-carbonyl ketones¹⁷⁴. Metallation of ethyl diazoacetate with Grignard reagents occurs at $-70^{\circ}C^{172}$ whereas at $0^{\circ}C$, only nucleophilic addition to the terminal nitrogcn is observed.

Ethyl diazoacetate has long been known to dimerize in ethereal potassium alkoxide solution¹⁷⁵. The most reasonable mechanism proposed¹ involves its reaction as Brönsted and Lewis acids at carbon and nitrogen centres respectively.

t The formation of thermally stable, metallated diazoesters, where the metal is **Ag** or **Hg** ctc., which do not proceed by initial proton abstraction, is discussed in Chapter 7 on Complex Formation'.

Primary α -diazoketones display a similar balance of reactivity as acids and clcctrophilcs towards bases. **2,4,6-Trimethyl-w-diazoacetophcnone (47)** undcrgoes base-catalysed proton deuterium exchange rapidly in alcoholic solution¹⁷⁸, as do primary diazosulphones⁴⁰. The reaction product with methoxide in methanol is the tetrazine **(48)"',** analogous to **46,** however with r-butoxide in r-butanol the tetrazole dimer (49) is formed¹⁷⁸. Both types of dimer are also produced by diazoacetophenone, but with potassium hydroxide in dimethylsulphoxide and with potassium /-butoxide in t -butanol, respectively¹⁷³. The influence of different bases and conditions on the reaction course has not bcen clarified as both of **the** above dimers cm be considered to arise from initial proton abstraction or nucleophilic addition by the base.

Dimerization subsequent to proton abstraction is also observed in the reaction of diazoketones with lithium diisopropylamidc at - ⁷⁰*"C,* however **if** a ketone is also present, condensation occurs to form the α -hydroxy diazoketone¹⁷². For this reaction butyllithium is an unsuitable metallation reagent as **it** competitively adds to the terminal diazo nitrogen¹⁷⁴.

Base-catalysed condensation of diazoketones, diazophosphonate esters and diazophosphineoxides¹⁷⁹ with carbonyl groups is observed under mild conditions if the latter are sufficiently activated, e.g. in isatin derivatives¹⁸⁰, or held in the immediate vicinity of the diazo grouplsl. This condition allows even attack **at** a carboxyl function.

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It should finally be noted that the presence of an α -diazo group does not exclude condensation¹⁸³

V. LEWIS ACIDITY OF DIAZO AND DlAZONlUM GROUPS

A. Coupling of Aromatic Diazonium Ions

I. Introduction

+

The reactions of the aromatic diazonium group with nucleophiles can be considered as substitutions or additions. The former class does not provide an index of the Lewis acidity of the group and, **as** it has recently been reviewed181, will not be considered further here. General considerations of terminal addition to the diazonium group by nucleophiles, such as hydroxide, alkoxide, phenoxide, cyanide, azide, thiolate and bisulphite anions, phosphites, phosphines, ammonia and substituted amines, have been extensively discussed¹⁸⁵. Quantitative thermodynamic aspects of these reactions are presented elsewhere in this volume. The coverage of the literature on azo-coupling to aromatic¹⁸⁶ and aliphatic carbon nucleophiles¹⁸⁷⁻¹⁸⁹ will not be repeated. The following discussion will be limited to more recent results, which may define the relative 'kinetic acidity' of several classes of nucleophiles. re on azo-coupling to aromatic¹⁸⁶ and aliphatic carbon nucleophiles¹⁸⁷⁻¹⁸⁹
be repeated. The following discussion will be limited to more recent results,
nay define the relative 'kinetic acidity' of several classes of

2. Reactions with nucleophilic carbon

Generally the principal factor influencing the ease of coupling with aliphatic carbon bases is the nature of the substituents in the base, rather than in the diazonium salt.

$$
ArN = N + \text{CHR}_2 \longrightarrow ArN = N - \text{CHR}_2 \longrightarrow Ar - NH - N = \text{CR}_2 \quad (62)
$$

The rate constants for hydrazone formation for the anions of nitroethane¹⁹⁰, acetoacetanilide¹⁹¹, acetylacetone¹⁹² and acetone¹⁹³ with a series of diazonium salts have been determined. Furthermore, good correlations of reaction rates for each anion with Hammett's σ parameters were obtained. As was noticed for coupling to aromatic nuclci, the most reactive diazoniuni ions have negative deviations from this relation as the diffusion rate limit is approached. This phenomenon was ascribed to lack of additivity of substituent constants rather than to incursion of diffusion control (with one exception), or to a change of ρ values with decreasing activation free energy¹⁹⁴. Such a change in ρ with the overall reactivity of each anion type was however observed. Coupling to the highly reactive anion of acetone $(pK_n = 19 - p)$ **20)** was associated with a value of 1.89, compared to a value of 3.45 for acetylacetone. In fact the ρ values were observed to decrease regularly with increasing rate constant, even when data for coupling to aromatic nucleophiles were included, although the correlation was better when restricted to substrates of similar structural type¹⁹⁵. Such an analysis of the Hammett ρ values in terms of selectivityreactivity relationships is, however, subject to severe limitations and has recently been criticized¹⁹⁸. The uncertainty in extrapolating this sort of comparison to substrates of different charge type is typified by the similar *p* values of 4.0 and 4-15, for the coupling reactions to 1-naphthol and its anion respectively, despite the greater reactivity of the latter by a factor of 10⁷ ¹⁹⁷. Such a difference in reactivity between a nucleophile in its protonated and unprotonated forms is common to enols, naphthols, pyrazolones, naphthalene sulphonic acids¹⁹², pyrroles¹⁹⁸ and 3-methylindole¹⁹⁹.

3. Reactions with heteronucleophiles

8

The rate constants for coupling of aromatic diazonium ions with nitrogen in amines appear to decrease in the series N -methylaniline²⁰⁰, dimethylamine²⁰¹, aniline²⁰². This decrease is not reflected by a systematic change in the values of ρ 3.94, **3.47** and **3.90,** respectively. The effect of variation of aromatic substituents on the coupling rate with methylamine is difficult to assess, since electron-donating substituents encourage further reaction of the initially-formed triazene with the diazonium ion to generate a pentazadiene 203 . **Example 19.19** The system have so far defied analysis²⁰¹,
tic results for this system and the series N-methylaniline²⁰⁰, dimethylamine²⁰¹,
This decrease is not reflected by a systematic change in the values of ρ The rate constants for coupling of aromatic diazonium ions with nitrogen in
amines appear to decrease in the series N -methylaniline²⁰⁰, dimethylamine²⁰¹,
3.94, 3.47 and 3.90, respectively. The effect of variation of

The kinetic results for this system have so far defied analysis²⁰¹.

Coupling with hydroxyl, cyanide, bisulphite and azide ions leads initially to the cis-azo compounds, despite the fact that the *from* isomers are thermodynamically favoured²⁰⁵. The syn diazohydroxide formed when the nucleophile is hydroxyl ion is extremely rapidly deprotonated to the *syn* diazoate^{206, 207} (see Section VII.A). The aryldiazoazide formed with azide ion decomposes into aryl azide and nitrogen, or cyclizes to an arylpentazole, prior to nitrogen evolution²⁰⁸.

I ⁺ Ar-N=N+ + - N=N=N- -. **Ar-N=N-N=N=N-** (--t **Ar-pJ=N-N-N=N-***J*
For several solvents the kinetic nucleophilicity of anions is in the order $C_6H_5S^ N_3 < CN^{-209}$. An exception to this sequence is provided by the reaction of azide with diazonium ions (other than with *p*-nitrobenzenediazonium) in water²¹⁰. In these particular cases the rate-determining step is assumed to be decomposition of the $aryldiazoazide²¹¹$. The above-mentioned reactivity order is maintained towards triarylmethyl and tropylium ions and is regarded as following the relative ease of desolvation of the nucleophiles 2^{209} . Consequently, a considerable part of the activation cncrgy for thcse rapid rcactions is iinaffcctcd by **the** reactivity of the diazoniuni ions. Hcnce there does not exist even an approximate relation between Harnmett *p* values and ovcrall reactivity as found for azo-coupling rcactions. The *p* values in values and overall reactivity as found for a co-coupling reactions. The p values in question for $\text{COH}_{(H_2O)} = 2.16^{212}$, 2.1^{213} , $\text{~COH}_{(H_2O)} = 2.31^{211}$, $\text{N}_{3 \text{~(MeOH)}} = 2.83^{214}$ are considerably smaller than thosc for azo-coupling to carbon nucleophiles whcre $p = 4.0$, despitc the fact that these latter reactions are much more rapid. The coupling rate of 2,6-dichloro-4-nitrobenzenediazonium ion with 1-naphtholate anion is IO:'-tinics that **with** hydrosidc ion, despitc the greatcr basicity of the lattcr by a factor of 10^{6 215}. The similarity of the ρ values of azo-coupling to carbon to those of diazonium-diazo equilibria with the above anionic nucleophiles, i.e. $3.2 <$ *^p*< 5.2. may indicatc that **the** transition states for thcse azo-couplings are 'product like' l ³⁴. On the other hand, the smaller ρ values for reaction with anions (after their desolvation) imply a less sclectivc reaction, with 'rcactant likc' transition state. This reasoning¹³¹ is consistent with formation of *cis* products for 'early transition state' reactions, and direct formation of *trans* products with amines, phenols and aromatic tertiary amines.

B. *Coupling of Aliphatic Diazonium Ions*

1. Stable aliphatic diazonium ions

Stablc, isolable aliphatic diazonium salts (i.e. those froni which spontaneous nitrogen evolution would lead to very unstable vinyl cations) do not undergo reactions analogous to azo-coupling. This is due to preferred reactions with nuclcophilcs at alternative sites. Thus olefin diazonium salts undergo substitution at the electrophilic olefin positions rather than addition at the terminal nitrogen²¹⁶.

The *y*-pyridinyl-substituted diazonium ion (50) is deprotonated by triethylamine²¹⁷. The diazonium-substituted acetyl-acetone complex (51) normally

experiences substitution of thc diazonium group by nucleophilcs, but does show colouration, indicative of coupling with naphthol²¹⁴.

2. Trapping of aliphatic diazonium ion intermediates

Coupling products have been isolated from reactions involving aliphatic diazonium ion internicdiates. **In** all cases the ions could be trapped due to nitrogen loss becoming unfavourable, assisted or unassisted, by position either on a bridgehead²¹⁹ or on a cyclopropyl ring^{70, 220}.

Undcr the same conditions as the cyclopropyl diazoniiim ion **(52)** expericnccs terminal addition, the isobutyl and methyl analogues prefer nitrogen substitution²²⁰. **A** bridgehead position is not in i1sclf suficient to pcrmit diazonium ion trapping. The polychlorinated liomocubane diazonium ion **(53)** gave products typical of radical decomposition of an intcrinediatc addition specics **(54),** and also couplcd with β -naphthol²²¹. In contrast, its hydrogenated analogue yielded only normal hydrolysis products in the same reaction conditions²²². polychlorinated homocubane diazonium ion (53) gave products typi
al decomposition of an intermediate addition species (54), and also co
 β -naphthol²²¹. In contrast, its hydrogenated analogue yielded only r
olysis prod

(54) 1 **(71) (53)** R-+N,+.ONO R=

their counter-ions co-generated in ion pairs. The less stabilized methane²²³⁻²²⁶ and ethane diazonium²²⁷ ions can be trapped by

C. Coupling of Aliphatic Diazo Compounds

Generally the reactivity of diazoalkanes towards nucleophiles increases with increasing contribution of limiting forms **(lb)** and **(Ic)** to their overall electron distribution. Diazoalkanes lacking electron-withdrawing substituents require more reactive nucleophiles.

Trialkyl- or triaryl-phosphines, though more feeble nucleophiles than amines toward many metal ions²²⁸, form, unlike amines, stable addition complexes with most diazoalkanes²²⁹, even diazomethane²³⁰. This unique reactivity results from the ability of phosphorus to stabilize the betaine thus formed (55) by π -acceptance into its empty **d** orbitals. Generally the reactivity of diazoalkanes towards nucleophiles increases with
increasing contribution of limiting forms (1b) and (1c) to their overall electron
distribution. Diazoalkanes lacking electron-withdrawing substi

$$
R_2\overline{C}-N=N^+ + P(Ph), \longrightarrow R_2\overline{C}-N=N-\overline{P}(Ph), \longrightarrow R_2C=N-N=P(Ph), (73)
$$
\n
$$
(55)
$$

When the delocalization is diminished by hydrogen bonding, for example in the a-hydroxyphosphazine *(56),* the thermal stability of the adduct diminishes correspondingly¹⁷². In this case dissociation into reactants occurs at 70 °C, whereas normally higher temperatures are required for pyrolysis²³¹.

Although the addition reaction is subject to acceleration by electron-releasing substituents in the phosphine aryl groups²³² ($\rho = -1.03$), the limitations of the extent of reactions seem to be primarily steric²³³. Thus diazocyclopentadiencs substituted with alkyl or aryl groups in both the 2- and 5-positions fail to react with triphenylphosphine. Reaction does proceed with the less hindered and more nucleophilic tri-n-butylphosphine, or if the 5-substituent is replaced by a hydrogen, chlorine or nitro group²³³. In the latter two cases the electron-withdrawing substituents would be expected to stabilize both the diazonium reactant and also the

product-contributing form **55.** The increased importance of this form **has** been invoked to explain the more ready hydrolyses of the phosphazines containing these substituents, by nucleophilic attack at phosphorus²³³. Trialkylphosphites also form phosphazine adducts, but examples are scarce234. **235.**

Diazoalkanes disubstituted with strongly electron-withdrawing groups, such as -CN *86,* -CF, **23G, -S02R 237,** react with phosphines or Grignard reagents at or below room temperature. In certain cases such compounds, e.g. a-nitro ethyldiazo acetate (57) can also serve as 'diazo-transfer' reagents²³⁸.

Bisarylsulphonyldiazomethanes also react readily with carbanions²³⁹ (equation 75), and bismethylsulphonyldiazomethanes can experience terminal nucleophilic attack by enamines^{240 , 241 (equation 76)}

Products

diazofluorene²⁴³ to form azines. Electron-rich olefins also react with electrophilic α -dicarbonyl diazoalkanes²⁴² and

An alternative route to azine products from diazofluorene and **58,** via prior dissociation of the latter to nucleophilic carbenes, has been proposed^{244, 215}.

The diazo carbon of diphenyldiazomethane is sufficiently nucleophilic to form azines with stabilized diazoalkanes without passing through a carbcne intermediate⁸⁶, 246

The more nucleophilic diazoethane carbon reacts similarly with p -nitro- ω -diazoacetophenone and less readily with *a*-diazopropiophenone, whereas diazomethane does not react under the same conditions^{217, 248}.

6. Basicity, acidity and hydrogen bonding **217**

The normal course of reaction of diazoketones¹⁷⁴, diazoesters¹⁷³ or aryldiazoalkanes²⁴⁹⁻²⁵¹ with carbanions from organometallic reagents^{173, 249} leads to a hydrazone after hydrolysis. 6. Basicity, acidity and hydrogen bonding

The normal course of reaction of diazoketones¹⁷⁴, diazoesters¹⁷³ of

kanes²⁴⁹⁻²⁵¹ with carbanions from organometallic reagents¹⁷³, ²⁴³ leads

Dh₂C-N=N+ +PhMgBr - Ph₂ n bonding

¹⁷⁴, diazoesters¹⁷³ or an

reagents^{173, 213} leads to

————> Ph₂C=N-NH

diazodiketones and eno

stituted pyrazoles²⁵².

$$
Ph_2\overline{C}-N=N^+ + PhMgBr \xrightarrow{\sim} Ph_2\overline{C}-N=N-Ph \xrightarrow{\sim} Ph_2C=N-\overline{N}-Ph
$$
\n(80)

The initially-formed, azo-couplcd products from diazodiketoncs and enolate ions may be isolated as such^{242, 251} or may cyclize to substituted pyrazoles²⁵².

The addition products of diazoalkanes and triphenylphosphoranes fragment to form the mixed azine 253 .

Less-nucleophilic phosphoranes where $R = Ac$, Bz or CO.Me failed to react.

 N -Terminal attack on diazokctones by cyanide and bisulphite anions, and by amino groups, is evident from the isolation of the corresponding addition products¹. These reactions involving amine and sulphide nucleophiles are usually restricted to intramolecular cyclizations¹, although pipcridine forms a linear triazene with the sufficiently reactive 1,1,1,3,3,3-hexafluorodiazo-2-propane²³⁶.

For other nucleophiles the evidence of reaction is less obvious. The extent of hydrogen-deuterium exchange in $2,4,6$ -trimethyl- ω -diazoacetophenone in deuterated methanol, catalysed by halide salts, depends on *the* nucleopliilicity rather than on the basicity of the catalyst¹⁷⁶. The proposed exchange mechanism is given in equation (84).

$$
RCO\overline{C}H-N=N-B \xrightarrow{\text{ROD}} \text{RCOCHD}-N=N-B \xrightarrow{\text{R.O}-} \text{RCOCD}-N=N-B \text{ etc.}
$$
\n(83)

The products of rcactions of diazokctones in **the** prcsencc of hydrosidc or alkoside ions are consistcnt \vith intcrmcdiatcs of *the* abovc typc. Similarly, nuclcophilic

rather than basic catalysis has bcen invoked to explain the interaction of diazocarbonyl groups with adjacent electrophilic functions¹⁴⁶.

The formation of *trans* cinnamic esters from aryl-2-diazopropionic esters in basic ethanol²⁵⁴ could well involve such an intermediate, instead of the original diazoester, as an acidic spccies.

On the other hand, the reaction of diazoketones in ethanol-sulphur dioxide mixtures²⁵⁵ is not necessarily the result of a nucleophilic attack of ethanol on the diazo group. **A** sufficient increase of acidity of ethanol, due to coordination with the Lewis acid $SO₂$, to permit protonation, would be an equally acceptable explanation.

Finally, azo-coupling with aromatic substrates is limited to the very reactive 1,3,5-trihydroxybenzenc for diazoketones, ethyl diazoacetate and diazomethane²⁵⁶, although dicyanodiazomethane couples with the 4-position of N , N -dimethylaniline⁸⁶.

VI. HYDROGEN BONDING

A. Diazoalkanes as Proton Acceptors

The formation of a hydrogen bond, between an acid and a diazo carbon atom, has been shown to account for a considerable part of the activation energy of the overall proton transfer (Section II.B.6). In non-polar solvents, complexes so formed may have a lifetime sufficiently long to permit an alternative reaction with the solvent. Thus Closs²⁵⁷ has proposed a carbenoid transition state (59) for the trifluoroacetic acid-catalysed formation of cyclopropanes from phenyldiazomethane in olefinic solvents.

The evidence for such a transition state includes stereospecificity of cyclopropane formation and partial deuterium incorporation from deuterated catalysts, as opposed to monodeuteration of the carbonium-ion-derived products. $-C \rightarrow C \rightarrow C$

The evidence for such a transition state includes stereospecificity of cyclopropane

formation and partial deuterium incorporation from deuterated catalysts, as opposed

to monodeuteration of the carbonium-ion-

A similar hydrogen-bonded intermediate was originally proposed to explain the product distribution in the alkylation of tautomeric acid systems²⁵⁸.

$$
A-H+CH_2N_2 \longrightarrow A-H \cdots CH_2N_2 \longrightarrow N_2+A-H \cdots CH_2 \longrightarrow A-CH_3
$$
\n(87)

Collapse of intimate or solvent-separated ion pairs would currently be a more acceptable explanation^{79, 258}.

Hydrogen-bond formation to diazoalkanes from alcohols²⁵⁹ is indicated by the function of the former as transesterification catalysts²⁶⁰⁻²⁶². The nucleophilicity of the alcohol hydroxyl group is sufficiently raised by hydrogen bonding to the weakly basic diazomethane to allow attack on **a** carboxyl group.

It is significant in this respect that dicthyl oxalate is transesterified under conditions which leave the less electrophilic diethyl malonate unaffected²⁶³. The catalytic function of diazomethane is supported by the lack of formation of methyl acetate, in the reaction of phenyl acetate with ethanol²⁶⁰.

 $\epsilon_{\rm{max}}$ and

of alcohols to electrophilic double bonds²⁶⁴, and of isocyanates or isothiocyanates to o -hydroxybenzoate esters²⁶⁵ (equation 90). Similar mechanisms are proposed for the catalysis by diazoalkanes of the addition

Catalytic efficiency for this reaction increases in the order diazoethane, diazomethane, phenyldiazomethane, which is the inverse of the order of their rates of reaction with phenols. For the most efficient catalyst, phenyldiazomethane, **0.1-** 0.01 molar equivalents are sufficient to ensure complete reaction²⁶⁵. The intermediacy of proton-bridged rather than discrete diazoniuni ion intermediates was demonstrated by the lack of incorporation of deuterium into the products, when molar equivalent quantities of deuterated diazoalkanes were used as catalysts^{266, 267}.

B. **Diazoalkanes as Proton Donors**

In solution, primary α -diazosulphones²⁶⁸ and α -diazoketones²⁶⁹ can function as hydrogen-bond donors, either to themselves or to solvent molecules. This is revealed by a downfield progression of chemical shift of the proton bonded to the diazo group in **a** series of solvents following the basicity of the solvent. **It** should be noted that in the solid phase no evidence of hydrogen bonding was to be found in the determination of the crystal structure of 1,4-bisdiazo-2,3-butadione²⁷⁰.

On the basis of solvent dependence of chemical shift, diazosulphones are apparently better hydrogen-bond donors than are diazoketones. The latter possess, however, a better acceptor site in the carbonyl group as compared to the sulphone group268.

The difference in hydrogen-bond donor ability between the two classes of compound has been invoked to explain the alkylations of diazosulphones **(60),** but not of diazokctones by enamines²⁷¹. Formation of the hydrogen bond to the enamine apparently increases its immonium ion character *(62),* and hence its susceptibility to attack by the nucleophilic diazo carbon.

.4 close parallel to the above behaviour cxists in the chemistry of diazodiphenyl phosphine oxides **(61).** The p.m.r. spectrum displays a similar, solvent-dependent, chemical shift for the proton attached to the diazo group²⁷², and furthermore

61 adds elcctrophilically to enamines in exactly the same manner as do diazosulphones²⁷³.

VII. ACIDITY AND BASICITY OF AROMATIC D!AZO COMPOUNDS

A. Diazohydroxides

1. Introduction

The investigations of acid-base equilibria involving diazohydroxidcs are complicated by two major features. Firstly, both thc diazohydroxides and their conjugate bases display geometrical isomerism between the syn and the more stable *anti* forms, interconversion of which may be acid catalysed. Secondly the *syn* diazohydroxide is itself a component of a **Lewis** acid-base equilibrium, with arenediazonium and hydroxide ions. **As** the equilibrium constant for this reaction is usually much inferior to the dissociation constant of the *sytr* diazohydroxide, **the** latter, when formed from diazonium ions, is almost totally deprotonatcd by the hydroxide ions present²⁷⁴. This behaviour, summarized in equation (91), precludes a direct measure of the acid dissociation constant K_2 from equilibrium concentrations.

Analysis of kinetic data for conversion of diazonium ions into *syn* and *anti* diazotates **(65)** and **(66)**, and *vice versa*, allows separation of dissociation constants (K_2 and K_3) for syn and anti diazohydroxides (63) and (64). In some cases, the rates of reaction of **63** and **64** with acids have been determined.

2. Dissociation of syn diazohydroxides

2,6-Dichloro-4-nitrobenzenediazohydroxide is exceptional in that, for a certain pH range, it may be present in considerable concentrations in equilibrium with the other species in equation **(91)215.** This is in part due to the instability of thc corresponding diazonium ion, which results in an increase in $K₁$, and also to diminution of the isomerization equilibrium constant K_{iso} . The dissociation constant K_2 was calculated from that of the equilibrium mixturc of *syii* and *atiti* diazohydroxides and the isomerization constants K_{iso} and K'_{iso} , to be equal to 1.55×10^{-5} . Solutions of *syn* diazotates can be prepared by reaction of hydroxide ion with diazonium ions where substituents are not mesomerically electron withdrawing, and hence do not favour syn-anti isomerism²⁷⁵. Quenching of these solutions immediately after formation with buffer solutions reforms the benzenediazonium ion with a rate dcpendent on the concentration of the *syn* diazohydroxide present²⁷⁶. Hence, the variation of rate with the pH of the buffer solution employed allows determination of $K₂$. The Hammett ρ value for the dissociation constants of such a series of diazohydroxides was found to be **1.3.**

3. Dissociation of *anti* **diazohydroxides**

The rates of conversion of *anti* diazotates (66) to the corresponding diazonium ions in solutions of high pH (> 7) are directly proportional to the equilibrium concentrations of the anti diazohydroxides **(64).** Thus the variation of rate with pH readily gives the appropriate dissociation constants *K3 ?06.* Such an analysis has been applied to the reactions of pyridine²⁷⁷, purine²⁷⁸, mono-²⁷⁹, di-²⁸⁰ and trisubstituted²¹⁵ *anti* diazotates. The values of pK_a thus established vary from 4.65 for 2,6-dichloro-4nitrobenzenediazotate to 7.40 for 4-toluenediazotate. In the series of monosubstituted substrates, the dissociation constants give a Hammett correlation with $p = 1.45$ ²⁷⁹ (or 1.15²⁰⁷ as measured by potentiometric titration). The magnitude of these values and those for the *syti* isomers, both being much less than for phenol ionization, exclude, as expected, the possibility of localization of the chargc formed on the nitrogen or oxygen.

4. Reaction of *anti* **diazohydroxides with acids**

ions has a first-order and a second-order component^{275, 279}: The rate of conversion of mono-substituted *anti* diazohydroxides into diazonium ns has a first-order and a second-order component^{275, 279}:

$$
\frac{-d[ArN_2OH]}{dt} = k_1[ArN_2OH] + k_2[ArN_2OH][H_3O^+]
$$
 (92)

The first-order component, with rate constant k_1 , is a composite of direct dissociation into ArN₂⁺ and OH⁻ (for which the Hammett $\rho = -2.6$) and an uncatalysed (or water-catalysed) isomerization *anti-syn*²⁷⁹. The rate constant $k₂$ of the secondorder component represents an acid-catalysed isomerization of *anti-syn* diazohydroxides, and is associated with a ρ value of -2.4 . In the conditions where k_2 [ArN₂O+]^{[H₃O⁺] is significant, generally at $pH \le 4$, the scission of the *syn* diazo-} hydroxide is more rapid than thc reversc isomerization. In the pyridine 4-diazohydroxide system a water-catalysed reaction, possibly corrcsponding to an *anti-syti* isomerization, is significant only at $pH < 1$, in which region the aromatic ring nitrogen is already protonated. Acccleration of the reaction of 2,6-dichloro-4-nitrobenzenediazohydroxide and 2,4-dinitrobenzenediazohydroxide²⁹¹ in this region of **pH,** with increasing acid conccntration, Cali also be ascribed to acid catalysis of the

6. Basicity, acidity and hydrogcn bonding **223**

rate-determining *anti-syn* conversion. The existence of such an acid catalysis term does not resolve the question of whether the *anti* diazohydroxides exist as such in solution or are better represented as their N -nitrosoamine tautomers²⁸². Protonation of either tautomer could facilitate overall conversion to diazonium ion, either by reduction of the isomerization energy barrier or by improving the leaving group in the direct scission reaction.

5. Reaction of *syn* **diazohydroxides with acids**

Arenediazotates with 2.6-dichloro-4-nitro- and 2,4-dinitro- substituents generate their relatively reactive diazonium ion counterparts by a rate-controlling scission of the *syn* diazohydroxide at pH 4.5. In this pH region, the scissions are general acid catalysed²⁸¹. This was found also to be the case for *syn* diazohydroxides generated by quenching a solution of *syn* diazotates²⁷⁶. The Brönsted α values could be measured for the catalysis in both experimental approaches. Those of substrates with the electron-withdrawing substituents were greater $(0.34, 0.31)$ as opposed to 0.19).

8. Diazophenols

Arenediazonium ions with *0-* or p-hydroxy substituents react with only one equivalent of base to form the corresponding phenolate ion, often named a diazophenol:

The pK_a value of 3.19²⁸³ is inferior to that of 2,4-dinitrophenol (4.11) as expected from Hammett substituent constants for the diazonium group.

C. Triazenes

A comprehensive review of the acid-catalysed rearrangements of 1,3-diaryltriazenes to p -aminoazobenzenes has been given elsewherc²³⁴. 3-Alkyl-1-aryltriazenes react readily with acids to form the corresponding anilines, nitrogen- and carboniumion-derived products²⁸⁵. This decomposition prevents direct determination of their

basicity. Protonation rates are also inaccessible from the kinetic data accumulated to date. Either there exists a rapid acid-base equilibrium, with rate-determining scission of the conjugate acid in water²⁸⁶, or the rate-limiting protonation is considered to be coupled with simultaneous departure of a carbonium ion fragment in chloroform²⁸⁷. The acid-catalysed scission of 1,5-diphenyl-3-methyl-1,4-pentazadiene in water proceeds by a mechanism comparable to the former²⁸⁸, with its consequent limitations of cvaluation of protonation rate constant.

VIII. ACKNOWLEDGEMENTS

The author ivould like to express his gratitude to Professor H. Dahn and to **Mr T.** Smyth for helpful discussions throughout the preparation of this chapter.

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CHAPTER 7

Complex formation

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1. INTRODUCTION

The structure of diazoalkanes may be described in the valence-bond formalism in terms of the canonical structures **1,** *2* and 3, and their reactions may be classified

$$
\begin{array}{ccc}\n\sum \bar{C} - N = N: & \sum C = N = \bar{N}: & \sum \bar{C} - N = N: \\
(1) & (2) & (3)\n\end{array}
$$

on this basis. Thus diazoalkanes may react as 1,3-dipoles (3) or as nucleophiles through carbon **(1)** or through the terminal nitrogen **(2). In** addition, the unsaturated nature of these compounds renders possible the formation of π -type complexes.

Diazonium ions on the other hand may be described as **4** and **5,** and they react normally as electrophiles through the terminal nitrogen **(5).**
 $B = \frac{1}{N} - N!$

$$
R-\dot{N} = N: \qquad R-\dot{N}=\dot{N}: \qquad (4) \qquad (5)
$$

The possibility of certain metal derivatives acting as Lewis acids and bases enables them to form complexes with diazoalkanes and with diazonium ions. Few complexcs have been isolated wherc a diazoalkane molecule as such is present as a Jigand, but their presence as reactive intermediates has been invoked in numerous occasions. Consequently most of the evidence presented is of an indirect **nature.** On the other hand. diazoniuni ions have yielded numerous stable complexes with transition metal derivatives. While this manuscript was in preparation, an authoritative review appeared in this field'. Hence, the emphasis will be on the most recent developments.

II. COMPLEX FORMATION BY DIAZOALKANES

The number of stable complexes of diazoalkanes reported in the literature is severely limited. Nevertheless in the few examples described the metal has been proposed to bind the diazoalkane in three different ways. Hence, metal-carbon, metal-nitrogen and 'side-on' bonding have been suggested.

A. Carbon Coordination

There is **a** wealth of reactions in which diazoalkanes are thought to act towards metal derivatives as nucleophiles through the diazo carbon to give intermediates of type *6* (Scheme 1) but their direct observation has been limited to two reports.

SCHEhlE 1

Upon treatment of 8 with diphenyl diazomethane $at - 78$ °C the observed colour changc from red to brown was attributed to the formation of a complex whose structure has been assigned as **9** on the basis of chemical evidence?. Furthermore, the less stable diazornethane apparently forms with *8* an analogous coniplex at lower temperatures?.

It has bcen postulated that a similar complex is formed in the reaction of **AIEt,** with diazomethane, which at -78 °C yields a colourless solution without nitrogen evolution³. Spectroscopic and other methods to probe further into the nature of the bonding in these complexes have not been used.

The intermediate **6** may suffer **a** variety of reactions which lead to the formation of different types of products. Thus it is often the case that compounds of Groups IVb and Vb which possess a basic or potentially basic group react with diazoalkanes possessing an *a* proton to give a heterosubstituted diazoalkane as in path (a). Some transition metal derivatives of Groups lb and IIb have also been shown to react in this fashion but more often these dcrivatives induce the decomposition of diazoalkanes. Products of formal insertion of the diazo carbon into the bond between the metal and one of its substituents (by loss of nitrogen and 1,2-migration of a metal substituent to the former diazo carbon) are commonly observed with the normal elements and also with transition elements, the intermediacy of species **7a,** an 'invcrse' metal ylide, or **7b,** a metal-carbcne complex, often being postulated. These species have available other reaction paths such as polymerization via successive attack on *7* **by** diazoalkane and 1,2-migration of the growing alkyl chain, attack by the terminal nitrogen of another diazoalkane molecule to form an azinc, attack by the carbon of another diazoalkane molecule with formation of olefin. In addition it may react with compounds such as olefins to give products of formal carbene insertion and/or of carbonium ion-type rearrangements. **2 Me,SnNMe,+CH,N,** - **(Me,Sn),CNz+2 HNMe,**

The factors that govern the path chosen by the reaction are not well understood at present. We shall examine some evidence that bears on the question of stability of intermediates *6* and **7.**

Reaction (a) (Scheme 1) has been widely applied in the synthesis of α -heterosubstituted diazoalkanes $4-13$, as the examples given below testify.

$$
2 \text{ Me}_3 \text{SnNMe}_2 + \text{CH}_2 \text{N}_2 \longrightarrow (\text{Me}_3 \text{Sn})_2 \text{CN}_2 + 2 \text{ HNN} \text{R}_2
$$
\n
$$
\text{MeAs(NMe}_2)_2 + 2 \text{ HCCOOE1} \longrightarrow \text{MeAs(CCOOE1)}_2 + 2 \text{ HNN} \text{R}_2
$$
\n
$$
\begin{array}{ccc}\n\text{MeAs(NMe}_2)_2 + 2 \text{ HCCOOE1} \longrightarrow & \text{MeAs(CCOOE1)}_2 + 2 \text{ HNN} \text{R}_2 \\
\text{M} & \text{N}_2 \\
\text{M}[\text{N(SiMe}_3)_2]_2 + 2 \text{ HCCOOE1} \longrightarrow & \text{M(CCOOE1)}_2 + 2 \text{ HN(SiMe}_3)_2\n\end{array}
$$
\n
$$
(3)^{18}
$$

N2 ^M= **Zn, Cd, Hg Hg(OAc),+4 HCCOOEt** - **t Hg(CCOOEt),+P Nz+2 AcOEt (4)'O** II II **N2 N2 Hg0-l-2 HCCOOEt** - **Hg(CCOOEt),+H20** II **Nz** II **NZ 2 AgOAc+2 CH,N,** - **AgzCN,+2** AcOMe+N, **(6)z2**

$$
2 \text{AgOAc} + 2 \text{CH}_2N_2 \longrightarrow \text{Ag}_2\text{CN}_2 + 2 \text{ACOMe} + N_2 \tag{6}^2
$$

The mechanism **of** the reaction of metal amide derivatives such as equations (I) and (2), has been the subject of several discussions¹⁶. A recent proposal¹², appealing in its simplicity, involves a rate-determining nucleophilic attack of the diazo carbon on the metal with formation of a 4-centred transition state of structure **10,** in which interaction **also** takes place betwcen the diazo hydrogen and a basic ligand in the reactant.

$$
L_n M - NR_2
$$

\n
$$
N_2 C - H
$$

\n
$$
H
$$

\n(10)

The rcactions of diazoalkanes with transition metal derivatives, such as those shown in equations **(4),** (5) and *(6),* have not bcen studied from **a** mechanistic vicwpoint. In the formation of disilverdiazomethane (equation 6), acetate may be replaced by other anions provided that they give silver salts soluble in the solvent employed. Yiclds are better with anions deriving from weak acids. Added base helps the reaction, presumably by partially bonding an outgoing proton and/or the terminal nitrogen in the diazoalkane thus rendering thc carbon more nucleophilic.

Spectroscopic studies on α -heterosubstituted diazoalkanes provide some basis for judging factors that contribute to their stability. In the i.r. spectrum, these compounds present a band in the region 2150 -1950 cm⁻¹ attributed to an $N-N$ stretching vibration, sensitive to the naiure of the substituent.

For compounds 11, the $v_{\text{(NN)}}$ values decrease with progression down the Periodic

$$
Me3MCCOOEt \t M = C, Si, Ge, Sn, Pb
$$

\n
$$
N_2
$$

\n(11)

Table. This trend may be interpreted by invoking the increasing electron-donating effect in the series partially compensated, in particular, in the case of the silicon compound, by $p_{\pi}-d_{\pi}$ resonance interaction²².

The possibility of the metal in **a** metal-substituted diazoalkane expanding its valence shell has **also** been shown for compound **12.** In its reaction with crotyl bromide, an **85/15** mixture of **13** and **14** is obtained pointing to the intermediacy of an ion pair²³.

> N,
CH,CH=CHCH₂CCOOE **(1 3)** N,
CH2=CHCHCCOOE
CH₂=CHCHCCOOE ċн, **(14)** N,
<u>II</u> **(12)** $AgCCOOEt + CH_sCH=CHCH₂B$

The intermediacy of metal-carbene complexes^{24, 25} or 'inverse' metal ylides has been invoked in numerous mctal derivative-catalysed decompositions of diazoalkanes²⁶⁻¹². The nature of the rate-determining step of the reaction has been established by kinetic measurements in only **a** few cases. In the reaction of diphenyldiazomethane catalysed by $ZnCl₂$ or $ZnBr₂$ to yield, after work up, mixtures of benzophenone and benzophenone azine, the kinctics and product proportions arc consistent with rate-limiting formation of the complex $Ph_2CZnX_2^{36}$. When CuBr₂ is employed as catalyst, a copper complex which may have diphcnylmethylcne as **a** ligand is fornied in **a** rapid first step, the further consumption of diazoalkane occurring at a slower rate³².

Correlations have becn found betwecn the ratcs of decomposition of diazoalkanes and their basicity and the strength as Lewis acid of the metal compound. Thus, while diphenyldiazomethane shows no reaction with ZnCl₂ below - 15 °C, phenyldiazomethane reacts at -50 °C. The decreased basicity of the former compound may be attributed to inductive electron withdrawal and to the additional steric hindrance of the second phenyl ring³⁰. On the other hand, in the presence of the typically strong Lewis acids AICI₃ and SnCI₄, 0.1 mmol of diphenyldiazomethane reacts completely in air in less than **1** minute while the reaction takes hours in the presence of the weaker acids $ZnCl₂$ or $SnCl₂$ ³⁰.

In the decomposition of diphenyldiazornethane aidcd by different metal salts the observation was made that the products wcre related to the reduction potential of the salt in H₂O³⁸. Thus, when $T^[III](OAc)₃$ ($E = +1.21$ V) or Cu^{II}(OAc)₂ $(E = +0.15 \text{ V})$ was used the products resulted from a two-electron or one-electron transfer with formation of TI'OAc and Cu'OAc respectively, whereas when $Cr^{III}(OAc)₃$ ($E = -0.41$ V) was employed no reduction of the metal ion took place. The mechanism proposed involves internal acetate transfer within a metal-carbene complex38.

In a very limited number of cases stable metal-carbene complexes have been isolated from the reaction of metal derivatives and diazoalkanes (equations $7⁴¹$, **⁸31** and **9 9.**

$$
trans[lrCl(CO)(PPh2Me)2] \xrightarrow{(CF3),CN2} N2 + \n\begin{array}{c}\n & C \n\downarrow \\
 & C \n\downarrow
$$

$$
[Pd(PPh3),(CH2=CH2)] \xrightarrow{FW1} [Pd(PPh3),(FI)]+CH2=CH2
$$
 (8)

 $FI = 9 - Fluorenylidene$

Mn, *OC20* **CArZ**

On the other hand, methods different from path (b) (Scheme **1)** have been used to synthesize numerous complexes, most of which possess oxy, thio, seleno or amino substituents on the carbene carbon atom^{24, 25}. These complexes may be said to originate from nucleophilic carbenes^{14, 45} and their bonding is interpreted in terms of the limiting structures **18, 19** and **20.** complexes **15, 16** and **17** would thus constitute exceptions by being electrophilic carbene complexes. Presuniably the electronwithdrawing character of the carbene substituents induces strong π -back donation from the metal and hence produces a strong $M-C$ bond.

$$
L_n\overline{M} - C \begin{matrix} \overline{X} & & & \overline{X} \\ & \overline{X} & & \overline{X} \\ & & \overline{X} & & \overline{X} \\ & & & \overline{X} & & \overline{X} \\ & & & & \overline{X} & & \overline{X} \\ & & & & & \overline{X} & & \overline{X} \\ & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & & & \overline{X} & & \overline{X} \\ & & & & & & & & & \overline{X} & & \overline{X} & & \overline{X} \\ & & & & & & & & & & \overline{X} & & \overline{X} & & \overline{X} \\ & & & & & & & & & & \overline{X} & & \overline{X} & & \overline{X} & & \overline{X} \\ & & & & & & & & & & \overline{X} \\ & & & & & & & & & & & \overline{X} \\ & & & & & & & & & & & & \overline{X} & & \overline{X
$$

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The methylene insertion reaction has been the subject of several structurereactivity studies that have confirmed that a nucleophilic attack on the metal is involved in the rate-determining step^{46, 47}. The data available do not distinguish cither between a stepwise and a concerted process or betwecn a zwitterionic interrncdiate such as **213** and an ion pair intermediate such as **21b.**

$$
\begin{array}{ll}\n[L_n\overline{M}CH_2N_2^+] & [L_n\overline{M}CH_2N_2^+]X^-\\ \n& \downarrow\\ \n& \times & (21b)\\ \n& (21a)\n\end{array}
$$

In general it is found that organic substitution in the Group IVa halides tends to decrease the reactivity of the $M-X$ bond towards diazomethane. Thus, while SiCl, reacted rapidly with CH_2N_2 in ether even at -50 °C to give ClCH₂SiCl₃, further methylenation was found to be increasingly difficult. Et_3PbCl , on the other hand, did react to give Et_3PbCH_2Cl in high yield⁴⁶. A quantitative study of the substituent influence upon the reaction of 22 with CH_2N_2 has produced a ρ value

of **2.3** *06.* These data show that more electrophilic character in the metal favours the reaction.

An initial association of diazoalkanes to the metal has also been proposed for their reaction with Group **Illa** derivatives. The adduct rcacts further to give either methylenation products^{3, 48-52} or/and polymerization of the diazoalkane^{3, 53}.

Qualitative expcrimcnts have led to the conclusion that the rate of thc reaction depends on the Lewis acidity of the metal derivative. Thus, while $B E t_3$ reacts vigorously with CH_2N_2 at room temperature to give polymethylene, $B(OEt)_3$ requires hours under the same conditions3.

The fate of the intermediate adduct has been found to depend on the Lewis acidity of the metal dcrivative as well, weakcr acids favouring polymerization over insertion⁵⁰. Thus higher proportions of polymerization over insertion are found for boron derivatives as coniparcd with analogous aluminium ones, for derivatives substituted with OR groups rathcr than R, **I-l** or **X,** and for reactions in which additional donor molecules have been added³. A mechanistic interpretation may be givcn (cquation 10). Intcrmediatc **23** bchaves as a strong elcctrophile and can add The Lewis activity of the field derivative. Thus, while

the CH₂N₂ at room temperature to give polymethyler

is under the same conditions³.

f the intermediate adduct has been found to depend or

the intermediate ad

further diazonicthanc to form **24.** However, the stronger the intcraction bctween the metal and the diazo carbon, the more effective will be the competition of $M-X$ bond rupture and migration of X to carbon⁵⁴.

$$
\begin{array}{c}\nL_n\bar{M}CH_2\bar{C}H_2\\ \n\downarrow\\ \n\chi\\ \n\end{array}
$$
\n(24)

0. Nitrogen Coordination

The reaction of diphenyldiazomethane with dodecacarbonyltri-iron and also its irradiation in the presence of pentacarbonyliron in benzene leads to two compounds^{55, 58}, whose structures have been determined by X-ray studies to be 25⁵⁵

and **266fi.** The N atoms in both complexes show angles characteristic of **sp3** hybridization and the N -Fe bond lengths are close to those found in other complexes known to contain $N(sp^3)$ - Fe bonds.

A single case has becn reported where a diazo compound **as** such is bonded to a metal through its terminal nitrogen⁵⁷, as shown in equation (11). The product of the reaction is stable up to 80° C, diamagnetic, and it liberates the diazomalonate ester upon treatment with PPh₃ under irradiation or with CO.

The structure proposed may be compared with those determined by X-ray studies for aryldiazenato complexes (see Section III.A), the basis for the comparison being the importance of limiting structure **28** in the ground state of the diazoester.

Complexes 27 show i.r. absorptions at 2025, 1982 and 1951 cm⁻¹ (R = H) and 2022, 1979 and 1948 cm⁻¹ (R = Me). By analogy with $[M(\pi-C_5H_5)(CO)_2(N_2Ar)]$ $(M = Cr⁵⁸, Mo⁵⁹⁻⁶¹$ or W^{62, 63}), the first two bands may be assigned to $v_{(CO)}$ and the last to $v_{(NN)}$. The high value of $v_{(NN)}$ points to a ligand with an NNC angle close to 180" (see Section 1II.C.l).

C. 'Side-on Coordination*

A series of complexes of Ni and Pd have been synthesized in which **a** diazoalkane ligand is claimed to be bonded 'side-on'^{31, 64}. They have the general composition ligand is claimed to be bonded 'side-on'³¹, ⁶⁴. They have the general composition [ML¹L²(diazoalkane)] (29) with the ligands being *t*-BuNC, PPh₃ or 1,5-cyclo-octadiene and the diazoalkanes, 9-diazofluorene or d octadiene and the diazoalkanes, 9-diazofluorene or diphenyldiazomethane. These complexes have been synthesized via addition reactions to [ML¹L²]⁶⁵ or ligand substitution reactions with $[ML^1L^2(CH_2=CH_2)]$, as shown in the examples (12) and **(13).**

$$
[ML'L^{2}] \xrightarrow{R_{2}CN_{2}} [ML'L^{2}(R_{2}CN_{2})]
$$
\n(29)
\n
$$
[Ni(l-BuNC)_{2}] \xrightarrow{Pb_{2}CN_{1}} [Ni(l-BuNC)_{2}(Ph_{2}CN_{2})]
$$
\n(12)
\n
$$
[Pd(PPh_{3})_{2}(CH_{2}=CH_{2})] \xrightarrow{FIN_{1}} [Pd(PPh_{3})_{2}(FIN_{2})]
$$
\n(13)

$$
F1 = 9 - \text{Fluorenylidene}
$$

The complexes 29 present an i.r. absorption attributed to $v_{(NN)}$ between 1550 and **1450** cm-I; a linear end-on coordination involving the terminal nitrogen would presumably show in the region around 2000 cm^{-1} and a coordination through carbon presumably still higher. Furthermore, the trends observed in the i.r. spectra upon changing the metal, the ligands and the diazoalkane parallel the trends found upon similar changes in complexes containing an alkyne ligand, which is known to be bonded 'side-on'³¹.

The structure of the ketenimine complex $[Ni(t-BuNC)_2(t-BuN=C=C(CN)_2)]$ has been determined by X-ray studies to involve a 'side-on' coordination in which the Ni atom is nearly equidistant to the N and the adjacent C^{66} . A similar proposal has been made for the bonding in the diazoalkane complexes, where the orbital picture would involve the donation from an occupied π orbital localized between the nitrogen atoms to the metal and π back donation from the metal into the corresponding unoccupied π^* orbital. 'Side-on' coordination has also been suggested for analogous derivatives with dicyanodiazomethane as ligand⁶⁶.

111. COMPLEX FORMATION BY DlAZONlUM IONS

Since the first synthesis of an aryldiazenatof complex of a transition metal in 1964 59 , this area of chemistry has been continuously expanding both in the quantity and in the scope of the research carried out in it. Among the transition metals belonging to Groups VIb, VIIb and VIII, there only remain to be isolated complexes *of* Tc, Ni and Pd. On the other hand, the vast majority of the coniplexcs reported involvc the aryldiazenato ligand, only a handful of alkyldiazenato complexes having been characterized.

A. Structure

The terminal aryldiazcnato ligand has becn observed in two geometrical forms: singly bent and doubly bent. In the first instance the bonding is described as σ -donation from an sp hybrid orbital on the terminal nitrogen with π back donation from d orbitals of the metal into a p orbital on the terminal nitrogen (N_{11}) . A simple valencc bond description of this type of complex has becn given (structure **30)"'.** Evidcnce exists to show that lipon reaction of an arencdiazoniuni salt to form a

t The following nomenclature will be **uscd':** diazcne (HN=NH), nryldinzcnc **(ArN=NH),** aryldiazenato (ArN₂- \sim or ArN₂ $\left\langle$) and arenediazonium (ArN₇⁺).

complex where the aryldiazenato ligand is coordinated in this fashion, the oxidation statc of the metal suffers no change. Hence, the ligand may formally be regarded **as** an ArN_2^+ two-electron donor ligand.

$$
M = \overset{+}{N} = \overset{+}{N}.
$$

The doubly bent terminal ligand has been described as having a single bond by overlap between an sp² orbital on $N_{(1)}$ with a hybrid orbital on the metal, as shown in structure **31 87.** In this case, the evidence available indicates that when an arenediazonium ion becomes coordinated as a doubly bent ligand, the metal suffers a formal two-electron oxidation. The bonding may hence be formally described as that between a two-electron donor ArN_r ligand and a doubly oxidized metal.

$$
M^{\Pi} \leftarrow N \cdot N
$$

$$
M^{\Pi} \leftarrow N \cdot N
$$

$$
M^{\Pi} \leftarrow N \cdot N
$$

The **X-ray** studies reported'. **03- 69** for complexes containing a singly bent Iigand'O. *⁷¹* show that the NNC angles lie within the range **118-124",** the only exception being $[RuCl₃(PPh₃)₂(p-N₂C₆H₄Me)],$ with a value of 137°. It has been suggested that slight back donation from the metal into π^* orbitals on the aryldiazenato moiety leaves the N-N bond largely as a triple bond and the NNC angle close to 180 $^{\circ}$ (hypothetical linear terminal diazenato ligand)', whereas stronger back donation as described above predicts this angle to be 120° . Intermediate values of the angle might be expected depending upon the relative importance of the linear and singly bent formalisms to the electronic structure or, equivalently, upon the degree of the metal to ligand electron transfer process⁶⁷ (see Section III.C.1). Detailed discussions on the electronic structure of the isoelectronic nitrosyl ligand have been reported7?. **73.**

B. Synthesis

The most frequently used reagents for preparing aryldiazenato complexes have been diazonium salts, either as such or as compounds that generate them *in situ*. The complexes are produced via ligand substitution, oxidative addition or Lewis base association reactions⁷⁴.

A general mechanism has been proposed', which has received additions and partial confirmation from comparative electrochemical studies on nitrosyl and aryldiazenato complexes⁷⁵; it is shown in Scheme 2. The nature of the products has been observed to depend on the metal and its ligand, but with the exception mentioned above⁷⁵ no quantitative studies have yet been reported.

Complexcs between diazonium ions and transition metal derivatives havc been postulated as reactive intermediates in the Sandmeyer⁷⁶⁻⁷⁹ and related⁸⁰ reactions, but structural proofs are not available.

Several other methods have been used to prepare aryldiazenato complexes including' apparent inscrtion of diazonium ions into metal-hydrogen bonds followed by deprotonation of the aryldiazene complex formed⁸¹, reaction of coordinated nitrosyl with aromatic amines, reaction of metal complexes with arylhydrazines⁸², organodiazenes or diazoalkanes⁸³, and modification of complexes already containing **an** aryldiazenato ligandsl.

SCHEME 2

C. Spectroscopic Properties

I. Infrared spectroscopy

Infrared studies have provided a useful tool for examining the interaction of the aryldiazenato ligand with the transition metal to which it is bonded and with other ligands present in the complex. Several studies have shown that although overlap of $v_{(NN)}$ with the absorption of other groups in the complex and coupling between the $N-N$ stretching vibration with aryl vibrational modes occur^{61, 81, 85}, the isolation of $v_{\text{(NN)}}$ is possible by mathematically decoupling the interacting bands which shift upon ${}^{2}H$ or ${}^{15}N$ substitution⁸⁵.

The $N-N$ stretching vibration is found within the broad region defined between *c.* 1340 and 2100 cm-'. This wide range is a reflection of the different coordination modes of the aryldiazenato ligand, i.e. 30 and 31. High values of $v_{(NN)}$ ($>c$. 1650 cm⁻¹) can generally be attributed to the occurrence of complexes of type **30,** whereas low values of $v_{\text{(NN)}}$ (<c. 1500 cm⁻¹) can be generally identified with complexes of type **31.** With values in the intermediate ovcrlap region, the assignment is not

A method has been proposed⁷⁰ which brings the $v_{\rm N}$ values to a common scale and reduces the region where the ligand structure assignment is doubtful, the basis of it being the variations expected in $v_{(NN)}$ upon changing the nature of the metal and its ligands.

The mutual influence of the aryldiazenato ligand and other ligands present in the same complex can be studied in CO-containing complexes, since $v_{(CO)}$ is an easily identifiable absorption. An increase in $v_{(CO)}$ implies an increase in the C-O bond order, a consequence of diminished d_{π} - p_{π} back donation by the metal⁶⁰. This effect may be induced by substituents in the aryldiazenato ligand. Thus, in complexes 32,

$$
[Fe(PPh3)2(CO)2(p-N2C6H4X)]+ BF4 X = NO2, F, CI, H, Br, OMe, OH
$$

the better electron-withdrawing substituents produce the highest $v_{(CO)}$ and the lowest v_{CNN_1} of the series⁸⁶, pointing to electron flow from the CO groups to the aryldiazenato ligands.

The complexes of the heavier metals present lower v_{NNN} values (or/and lower $v_{(CO)}$ values when appropriate) indicating that they are better electron donors than the lighter ones. Thus for example, $\text{[RuBr}_3(\text{PPh}_3)_2(\rho-\text{N}_2\text{C}_6\text{H}_4\text{Me})$, $v_{\text{(NN)}}=1895 \text{ cm}^{-1}$, may be compared with $[OsBr₃(PPh₃)₂(p-N₂C₆H₄Me)], v_(NN) = 1855 cm⁻¹⁸⁷$

The electronic configuration of the metal influences its capability of back donating electron density, a comparison of the $v_{(NN)}$ values for the aryldiazenato complexes reported showing that they exhibit a general trend in which $v_{(NN)}$ decreases as the $dⁿ$ configuration of the metal increases⁸⁸. Since the data available are rather scarce, simultancous variations in the ligands, the charge and the metal are included in the comparison.

The i.r. spectra of complexes 33 $(v_{1NN} = 1723, v_{1CO} = 2030, 1978 \text{ cm}^{-1})^{89}$ and **34** $(\nu_{\rm (NN)} = 1689, \nu_{\rm (CO)} = 2010, 1960 \text{ cm}^{-1})$ ⁹⁰, show that a positive charge, which may be assumed to reside mainly on the metal⁸⁶, diminishes the electron density available on the latter to share with the ligands. $(\nu_{(NN)} = 1689, \nu_{(CO)} = 2010, 1960 \text{ cm}^{-1})^{90}$, show that a positive charge, which the assumed to reside mainly on the metal⁸⁶, diminishes the electron densi
ailable on the latter to share with the ligands.
 $[Fe(CO),(PPh_2),(Ph$

$$
[Fe(CO)_{2}(PPh_{3})_{2}(PhN_{2})]^{+}BF_{4}^{-}[Co(CO)_{2}(PPh_{3})_{2}(PhN_{2})]
$$
\n(33) (34)

The nature of the ligands influences the availability of electrons on the metal. In equation (14), replacement of CO by the relatively poorly accepting PPh₃ ligand

$$
[Mo(CO)2(\pi-CsHaPPh3)(ArN2)]+PPh3 \xrightarrow{\sim} [Mo(CO)(\pi-CsHaPPh3)(PPh3)(ArN2)]
$$
\n(14)

$$
[Mo(CO)2(OH)(ArN2)]4+Base
$$

$$
=
$$

$$
[Mo(CO)2(OH \leftarrow Base)(ArN2)]4
$$
 (15)

results in an increased electron release from the metal and $v_{\text{(NN)}}$ and v_{CO} occur at lower wave numbers in the products⁹¹. In equation (15) the lower v_{tNN} , and v_{tCO} values observed in the adducts⁹² may be related to increased electron release from the metal caused by donation of electron density from the Lewis base to the OH ligand through a hydrogen bond.

Metal	Group	Charge	Ligands	Coordination number
-50 First row Second row -30 $(Mo, -10)$ Third row 0	$+100$ VIb. VIIb $+50$ VIII. $\mathbf 0$	$+2 - 140$ -80 $+1$ $\mathbf{0}$ Ω $+80$ -1 -2 $+140$ $+200$ -3	$+20$ Third PPh ₂ $+50$ Fourth $PPh3$ -20 $HBPz^-$ ^a $+60$ π -C ₂ H ₄ PPh ₃ $+20$ H^-	$+50$ 4 0 5, 6, 7

TABLE 1. Correction parameters to $v_{(NN)}$ **(cm⁻¹)**

 $Pz = 1-Pyrazolyl.$

Table 1 collccts the suggested correction parameters which should be added (or subtracted) to v_{tNN} ⁷⁰ to obtain a modified frequency v_{tNN} . The magnitudes of the corrections have bcen chosen so **as** to give the best agreement with data collected from several hundred known complexes of the isoelectronic nitrosyl ligand.

The number of X-ray structure determinations being very limited, the $v_{(NN)}$ valucs provide a useful tool to predict confidently the type of ligand present in the great majority of aryldiazenato complcxes, **a** value above 1550 cm-l indicating a complex of type 30 and one below 1530 cm⁻¹ of type 31.

A relationship between electron back donation by the metal and i.r. spectra has been demonstrated in complexes containing the *monohapto*-dinitrogen ligand^{93, 94}, isoelectronic with NO+ and with ArN; **e5.** In a series of these complexes, X-ray photoelectron spectroscopy studies show that increasing electron density on the N_2 fragment, as measured by the binding energy of its core electrons⁹⁶, is coupled with decreasing NN stretching frequencies⁹⁷.

2. N.rn.r. spectroscopy

Although the 'H-n.m.r. spectra of many of the aryldiazenato complexes reported have been determined, the information they yield is very unspecific and hence they have been utilized as an analytical tool rather than as **a** probe into the electronic distribution in the complexes. On the other hand, $19F-n.m.r.$ studies are severely limited in number^{61, 98, 99}, but they have yielded valuable information on the interaction at the electronic level between metal, aryldiazenato ligand and other ligands in the complex.

The most recent and complete study⁶¹ has applied the technique of Taft¹⁰⁰ and coworkers where the differences in chemical shifts between a para- and a meta-F substituent are taken as a measure of the electronic effects acting on the ring that holds these substituents. Thus, the more positive the value of $\delta_p - \delta_m$ is, the less shielded the para-F is relative to the *mera-F* substituent, and hence the less electron flow into the aryl ring has taken place.

Some results obtained are collected in Table **2.** Comparisons among them lead to the same conclusions concerning the influence of the metal and its ligands on the aryldiazenato ligand, as reached on the basis of i.r. spectra.

Complex	δ_{p} - δ_{m} (p.p.m.)
$[Mo(\pi-C_6H_6)(CO)(PPh_3)(N_2Ar)]$	-5.02
$[W(HBPz3)(CO)2(N2Ar)]o$	-4.54
$[Mo(\pi-C_5H_4PPh_3)(CO)(PPh_3)(N_2Ar)]^+BF_4^-$	-3.86
$[Mo(HBPz3)(CO)2(N2Ar)]b$	-3.58
$[Mo(\pi-C_5H_5)(CO)_2(N_2Ar)]$	-2.20
$[Mo(HCPz3)(CO)2(N2Ar)]+BF4$	-2.11
$[Mo(\pi-C_5H_4PPh_3)(CO)_2(N_2Ar)]^+BF_4^-$	-0.14

TABLE 2. ¹⁹F-N.m.r. data for aryldiazenato complexes^a

^a Chemical shifts for $CH₂Cl₂$ solutions and referred to external hexafluorobenzcne.

 $Pz = 1$ -pyrazolyl.

D. Chemical Properties

No quantitative study on the reactivity of aryldiazenato complexes has yet appeared in the literature. Conclusions concerning the influence of the metal, of the substituents on the aryldiazenato ligand and of the rest of the ligands on the reactivity of thc complexes have been based on **a** reaction failing to produce the expected product or not, or on the total time requircd for a complex to disappcar under given conditions. Some of these data are discussed below.

In general, the reactivity of the complex is very dependent upon the type of bonding between the metal and the aryldiazenato ligand. Complcxes possessing structure **30,** which present an M-N bond with a substantial double bond character, are found to be more stable towards N_a loss than complexes of type 31, where the $M-N$ bond is essentially single. The $N-N$ bond in complexes 30 presents a partial triple bond character (see Section **1II.A)** and is found to be more dificult to saturate than the corresponding bond in complexes of structure **31,** where it is essentially a double bond. Furthermore, the tendency to react with acids of complexes 30, which present **a** ligand impoverished in electrons, is found to be diminished as compared with complexes **31,** where the ligand is rich in electrons.

The nitrogen evolution from complex **35** according to equation (16) is fastest when **a** $NO₂$ substituent is present¹⁰¹, pointing to decreased electron withdrawal

$$
[Pt(PPh3)3(p-N2C6H4X)]+ BF4- — \n\qquad [Pt(PPh3)3(p-C6H4X)]+ BF4- + N2 (16)
$$
\n(35)

 $X = NEt_2$, NMe₂, OMe, H, Me, F, NO₂

by Pt^{II} in 35 and hence decreased Pt-N double bond character, when electronegative substituents are present. In accord \rith this explanation, a negative slope is observed for the correlation between the U.V. absorption corresponding to the $n-\pi$ ^{*} transition and Hammett's substituent parameters, σ^{101} . Furthermore, in the complexes $[Pt(PPh₃)₂(L)(ArN₂)]+BF₄$ the stability towards the decomposition shown in equation (16) follows the order $NH_3 \approx$ pyridine > PEt₃ \simeq RNC > CO, thus paralleling the global electron-donating effect of the ligand¹⁶².

The complexes $[M(PPh_0)_2X_0(ArN_2)]$ $(M = Ru, Os; X = Cl, Br)$ exhibit i.r. spectra consistent with structure **30** and are air-stable solids, resistant to protonation or hydrogenation under mild conditions⁸⁷. On the other hand, the complexes $[Rh(PPh₃)₂X₂(ArN₂)]$, presumably of structure 31, are stable as solids, but when dissolved in organic solvents they yield unstable solutions which react by eliminating nitrogen or, when possible, by abstraction of HCI from the solvent⁸⁷.

The complexes $[Os(CO),(PPh₃)(PhN₂)]+ PF₆$ are very reactive towards protonation, slowly extracting protons from freshly distilled methanol or ethanol, while their ruthenium analogues, although they can be protonated by acids, are stable in the presence of neutral alcohols⁸⁷. A relationship has been established between basicity of a complex and its $v_{(N,N)}$ frequency. Thus, the following order in basicities has been obtained⁸¹ for the d^6 complexes 36, 37 and 38: 36 $[\nu_{(N,N)} = 1455 \text{ cm}^{-1}] >$ **37** $[\nu_{(NN)} = 1462 \text{ cm}^{-1}] > 38 [\nu_{(NN)} = 1470 \text{ cm}^{-1}].$

> $[OsCl(CO)₂(PPh₃)₂(PhN₂)]$ $[RuCl(CO)_{2}(PPh_{3})_{2}(PhN_{2})]$ (36) (37)

> > $[IrCl₂(CO)(PPh₃)₂(PhN₂)]$

 (38)

Not only the ease of protonation but also the site **of** protonation varies with the structure of the aryldiazenato complex. Thus complexes **30** are expected to protonate on $N_{(2)}$ while complexes 31 are expected to do so at $N_{(1)}$. These *9*

expectations have been confirmed by X -ray studies in the cases of equations $(17)^{103}$ and $(18)^{104}$. **H**

$$
[Re^{I}Cl_{2}(PMe_{2}Ph)_{2}(NH_{3})(N_{2}Ph)] \xrightarrow{H^{+}} [Re^{I}Cl_{2}(PMe_{2}Ph)_{2}(NH_{3})(N=NPh)]^{+}
$$
 (17)

$$
{}^{1}Cl_{2}(PMe_{2}Ph)_{2}(NH_{3})(N_{2}Ph) \longrightarrow [Re^{1}Cl_{2}(PMe_{2}Ph)_{2}(NH_{3})(N=NPh)]^{+}
$$
 (17)
H
[Pt^{II}Cl(PEt₃)₂(ρ -N₂C₆H₄F)] \longrightarrow [Pt^{II}Cl(PEt₃)₂(ρ -N=NC₆H₄F)]⁺ (18)

H

Finally, although the discussion has been centred on aryldiazenato complexes, most of it should apply as well to **alkyl,** acyl and aroyl diazcnato complexes. Caution should be eserciscd howcver in not extcnding the gencrality too far because the stability of the aryldiazenato as ligand is greatly influenced by its capability to accept π electrons from a metal, a characteristic not shared by alkyldiazenato $groups¹⁰⁵$.

IV. ACKNOWLEDGEMENTS

The author wishes to esprcss his gratitude to Professor H. Dahn for his encouragement and advice and to Professor **J.-C.** Bunzli for helpful discussions.

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CHAPTER **8**

Synthetic applications of diazonium ions

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Di~pnrti?icrit of Clrrmistry, Uiiiwrsity of Missoitr-i- Rolln, Rollu, iMo *65401, USA,* and *Eqiiipe rlc Rcclierclre* No. *I2 rlic CiV RS, Lnbor-utoire rlc Cliiitiic, Ecole Norninle Sitpc'r-icitr-c, Paris, Fraricc*

1. INTRODUCTION

The treatments in this and the following chapter will in many cases be rather selective as a conscqucnce of thc sheer volume of material reported on an annual basis and the large numbcr of reviews which have covered various aspects during the past 20 years. (Reviews are marked with an asterisk in the References.) In the areas dealing with the alleged generation of carbcnes and carbenoids from diazoalkanes the author has refrained from employing thcsc terms, for as Kirmse has cautioned⁶⁵⁵, and the author has recently shown¹³⁰⁹, ¹³¹⁰, the assignment of reactions to thesc categories rcsts predominantly on a house-of-cards. In only **a** vcry few instances can assignments be made with any degree of rigour.

Numerous cxaniples and studies have probably been ovcrlooked, for the species are sufficiently common and their reactions so frequently employed that the processes have not been abstracted and are not detectcd by key word searches. In these instances the examples chosen reflect only a bias on the part of the writer for reading literaturc related to his synthetic and pcdagogical needs and not a feeling that a particular workcr's contribution is unworthy of enslirincment. The author has tried to cover all literature abstracted through Octobcr 1975 and all major journals which were in his hands by that date. In addition he has tried to enlist the aid of many workcrs who had published in thc area during **thc** preceding 2 years. Their responses are too numerous to acknowledge fully but results were made available to him prior to publication by Aracknl. Eistcrt, Doyle, Brokken-Zijp. Carrie, L. and Y. Vo-Quang, Hubert, Woolsey, Marchand, Moss and Rössler. Their kind aid is

gratefully acknowledged along with that of S. Julia who made his extensive notes on diazo alkane chemistry available to the author.

The origin of diazo chemistry dates from the initial generation of diazonium succinamic acid from asparagine by Piria^{991, 992} in 1849. It had only a transient existence and proceedcd to malic acid. Much chemistry has been performed on this class of compounds which furnished onc of the major bases for thc foundation of the modern chemical industry. Modern synthetic chemistry is normally assumed to bcgin with the work of Johann Peter Griess⁴¹⁶ in 1858 when he undertook to clarify the observations of Gerland12g (1853) that a red intermediate **was** formed in increasingly greater proportion at lower teniperatures whcn arninobenzoic acid was converted to hydroxybenzoic acid by the action of nitrous acid. His studies, like Gerland's,

were undertaken **at** the suggestion of Kolbe and followed the earlier observation by Hunt²²⁶ in 1849 that treatment of aniline with nitrous acid generated phenol. Griess²²⁶ modestly attributed his successcs to the fact that he operated at a lower temperature than Hunt of Gerland. He studied thc reaction of nitrous acid with picramic acid **(1)** and obtained the diazoxide **(2)** which he quickly rccognizcd as belonging to a new class of compounds. He immediately procecded to subject many othcr amino compounds to the same treatment. He prepared the first azo coupling product in **1861-624s1.** and by 1865-66 it was being prepared on a large scale by Caro??G. Griess continued his work for 3 years when he was Hofrnann's assistant in London and afterwards while working in **a** brewcry for Mcssrs Allsopp, Burton-on-Trent. He was faced with the need for CHN analyses and overcame the problem by developing a barter system with R. Schmitt in Dresden. Hempel's account is of interest: 'Regelmässig kamen von Burton an den Ufern des Trent die von Greiss dargestellen neuen Korpcr in klcinen Packeten, um in Dresden an der Elbe analysirt zu werden. Per Fracht kamen dann wohl gleichzeitig als willkommene Beilage Fässer von Allsopp's berühmten Pale Ale in ausgesuchtester Qualität.' Griess's investigations were very thorough and he discovered many of the reactions of aryl diazonium ions which had been observed up until the time of his death in 1891. Thus, by 1908 (50 years aficr his initial work) chemists had a rather thorough knowledge of the chemistry and structure of aromatic diazonium ions although work still continues in the area. The accuracy of many of the observations is all the more impressive when it is remembered that thcse progresscs were realized without i.r. and n.m.r. spectroscopy, X-ray structure determination and only very crude u.v.^{307, 1313} and visible spectra²⁰ studies.

The first important studies in the aliphatic arca after Piria's initial work occurred in 1883 when Curtius prepared diazoacetic ester (4)²⁷⁵, ²⁸⁷. Diazomethane was prepared by von Pechmann¹²⁴⁷ in 1894. The use of copper salts with aromatic diazonium salts was examined by Sandmeyer^{1091, 1346} in 1884, and Gattermann¹¹⁶ substituted finely divided copper powder for its salts in 1890.

A great resurgence in diazoalkane chemistry occurred in the 1950's when groups led by Alder¹¹ and Doering³⁰⁹ rcinvestigated the structure of the esters and acids obtained by Buchner^{169, 221, 222} from the decomposition of ethyl diazoacetate in benzene(3-9). Further impetus was gained from the production of 'evidence' that decomposition of diazoalkanes proceeds via 'carbcne' interniediatcs. Additional interest arose from the increasing demand for sypthetic pathways for strained hydrocarbons, the study of 1,3-dipolar cycloaddition reactions and studies related to examining the reliability and validity of orbital symmetry calculations.

The chemistry of alkyl diazonium ions has long been the object of a variety of mechanistic and synthetic studies with the synthetic efforts being primarily directed towards exploiting rearrangements of the Tiffeneau-Demjanov type $(10 \rightarrow 12)$ and the replacement of amino functions by various substituents. Most recently, research in this area has bcen directed towards the generation. isolation and use of diazotates as synthetic interrnediatcs and the continued development of improved means of changing amino functions to other functions.

II. REACTIONS OF ARYL DIAZONIUM IONS

The reactions of aryl diazonium ions have been systematically classified by Saunders¹¹⁰³. His classification system with minor changes is adequate and will be used here. It is summarized in thc Table of Contentst for this chapter and is used with thc permission of the publisher.

The formation of diazoniuni ions is the subject of another Chapter in this book and will not be dealt with further. Saunders¹¹⁰³ gives a very detailed discussion of the subject with appropriate references.

The aryl diazonium ions are of varying stability. Their stability is strongly influenced by the pH of the solution, temperature, the anions present, trace contaminants such as transition metal ions and the presence or absence of water¹¹⁰³.

Except for Class C, Group **3** (Reduction products) and Group *5* **(S-azo** products- diazo-thioethers) which are treated in other sections.

In the solid state some aryl diazoniurn salts exhibit appreciable stability whereas others are quite brisant explosives. There would appear to be no basic guidelines which predict stability without exceptions but it is possible to make some generalizations. It is highly probable that diazonium salts where the counter-ion is a reasonable oxidizing agent such as nitrate¹¹⁶, chromate¹¹⁰³, perchlorate¹¹⁰³, chlorate¹¹⁰³, etc. will not exhibit a high degree of stability. Conversely sulphates tend to be more stable as are the chlorides¹¹⁰³. However, the bchaviour of p-diazobenzene. sulphonate is notoriously fickle and some workers have found it quite dangerous1I0. Diazonium ions which have a potential leaving group *orrho* to the diazonium function are potentially dangerous because they can form 'benzynes' with the *o*-carboxy and o -SO_{$\frac{1}{2}$} compounds being at the extreme end of the scale⁵⁷⁰. In these cases fragmentation furnishes two neutral molecules, N_2 and CO_2 or SO_2 , all with high heats of formation. There are several diazo derivatives which have been patented as explosives. Thus 4,6-dinitro-2,1-diazoxide²⁵¹ has been used as a detonator which is less sensitive to friction than mercury fulminate or lead azide but equal or superior to both as a detonator. One should be suspicious of any

FIGURE I. Reprinted (with permission) from Saunders, *Thr Arornufic Diuzo Compouridr,* 2nd ed., Edward Arnold, London, **1949.**

compound having sufficient oxygen to convert more than half of the carbon atoms present into CO and/or a C/N ratio *c3.* All dry diazonium salts should be considered suspect until clearly demonstrated as being stable and then they should be treated with respect. In aqueous solution decomposition **of** benzene diazonium ion in strongly acidic media is independent of the anion $(Cl^-, Br^-, SO^{2-}, NO^{-}_{3})$ ¹¹²⁹, ¹¹³⁰, ³⁰⁹, ⁵⁰⁶ and proceeds according to the expression^{169, 221, 222}, $\ln k = 35.6457 - 27025/RT$.

 $Cain²²⁴$ found that the rate of decomposition fell at very high pH values. The overall trend in stability has been summarized by Saunders¹¹⁰³ in Figure 1. Snow¹¹⁶⁰ compiled a list of *32* diazonium chlorides arranged in order of their stability at *20 "C* but there appears to be no obvious correlation of structure with stability. It had been claimed that negative substituents increase stability^{1103} but the data do not support such a claim¹¹⁶⁰.

Some metals form double salts with diazonium ions and retard their decomposition. Similar effects occur upon addition of aryl sulphonic acids^{760, 810} and on the addition of sodium chloride^{506, 1130}, borates¹¹⁴³, tungstates⁷²⁵ and other neutral salts. Excess nitrite and nitrate is deleterious^{675, 1234} as is the presence of colloidal and of copper and its salts^{132, 133}.

The diazonium ion salts of strong acids are essentially **JOO%** ionized and therefore are comparable to **K** or Na salts in their degree of hydrolysis, but unlike these metal ions they do form stable covalent bonds with such nucleophiles **as** RS-, -OH and -CN. The resulting products will be discussed under Class B reactions.

A. *Class A: Derivatives Formed* **by** *Substitution or* **by** *Changes in the* **Sub***stituents of the* **Aryl** *Nucleus*

1. Group I : **Electrophilic substitution**

Although diazonium ions are of only moderate stability it is possible to perform some substitution reactions upon the **aryl** nucleus. This is facilitated by the fact that the $-N_{\tau}^+$ ion is the strongest electron-withdrawing group encountered in organic chemistry and acts both inductively and mesomerically. it is not surprising that in the benzenoid systems the diazonium compounds most prone to electrophilic substitution are the diazooxides which are formally neulral molecules. With polyannulated systems, electrophilic substitution occurs in a ring not bearing the $-N_2^+$ function¹³⁶³ (Scheme 2).

2. Group 2: Solvolysis of substituents on the aryl nucleus

When diazotization of an *ortho-* or *para*-substituted amine is carried out, there is always the possibility of the solvent, normally water, replacing a labile substituent to form an *u-* or p-diazoxidc. This can be prevented by operating in a highly acidic medium where the activity of the water is greatly suppressed by protonation and if the substituent is a chloride or bromide, by using HCl or HBr. Meldola⁷⁸² and colleagues examined these processes in great detail and found that normally two electron-withdrawing functions should be present on the initial amine and preferably in an *orrho* and/or *para* relationship to the departing group. Under such conditions

NO2 can leave as nitrite ion. Anisoles can even be solvolysed in acetic acid to furnish the diazo oxide. The failure to solvolyse in the last example **is** to be predicted on the basis that none of the required Wheland intermediates (Scheme **4)** for replacement of NO₂ places charge (PMO analysis)^{291, 292} upon more than one starred atom bearing an electron-withdrawing substituent. With the replacement of the OCH₃, the required Wheland intermediate places N_2^+ on an unstarred position and the driving force is not present. The rule appears to be that the leaving group must be on a carbon which furnishes a Wheland intermediate where the N_2^+ and one other strong electron-withdrawing substituent are present on starrcd atoms. Such a treatment is equivalent to the rules of Meldola and Hay^{784} . When NO_2 or

 $OCH₃$ can serve as a leaving group $NO₂$ is lost⁷⁸⁵. An amusing feature of the displacement of $NO₂$ is that once it begins, the amines furnish the HONO needed to carry out the diazotization.

Subsequent to formation of the diazonium ion, it is often possible to replace a labile substituent which has withstood the initial reaction conditions. This can be realized by employing a base such as an alkoxide or hydroxide. In these cases the requirements for positioning electron-withdrawing substituents are not **as** stringent. The processes are often quite rapid compared with other reactions. The attempted reduction of I **,6-dinitro-2-naphthalenediazonium** ion with alcohol under reflux failed and furnished instead the 2,1-diazoxide⁷⁸³. Sulphonic acid groups can be replaced¹¹⁴⁶ and the yield is improved by oxidizing the resulting sulphite ion to sulphate with hypochlorite⁷⁰⁶ or hydrogen peroxide⁹⁵¹⁻⁹⁵⁴ (Scheme 5). An extensive

investigation was performed by Orton⁹⁵¹⁻⁹⁵⁴ who compiled a table of the relative degrees of completion under standard conditions. He found that Br was a better activating and leaving group than CI, that the replacement of a halogen by $NO₂$ improves the reaction and the total number of substituents in the $2+4+6$ positions should be ≥ 2 . The data nicely correlate with a simple PMO analysis. The study of the resulting diazooxides from these displacement reactions has received considerable attention because they undergo many coupling reactions to furnish o-hydroxy-azo systems capable of forming metal ion complexes.

In a few cases it is possible to replace carbon substituents on an aromatic ring in the course of, or immediately subsequent to, diazotization. Elion³²⁶ found that the use of the Witt method of diazotization¹²⁹⁷ led to the replacement of a carbonyl function by **a** nitro group in a series of 3,5-dibromo-4-aminophenyl carbonyl compounds and that bromine was replaced by nitro in 3,5-dibromo-2-aminobenzoic (Scheme *6).*

8. **Synthetic applications** of diazonium ions *²⁵⁵*

3. Group 3: Exchange of diazonium anions with substituents of the aryl nucleus

Hantzsch and others^{49, 452, 455-457, 492, 498, 501, 605, 543, 1128 found that a few diazonium} salts were capable of exchanging anions between the $-N_2$ function and the aryl nucleus. The process **was** not reversible. **As** in the case of solvolysis, **Br-** was *a* better leaving group from the aryl nucleus than **CI-** and these are probably only special cases of type **A-2** reactions. Although one might write a concerted conversion of **35** to **36,** it is not possible to account for **37** going to **38** in a concerted fashion and a p-Claisen mechanism would be required **for** the transformation **39** to **40.**

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6. **Class** €3: *Derivatives in which the* **Diazo** *Group is Involved but Remains Functionally lntact*

1. Group I: Metallic diazotates and diazoanhydrides. The use of *iso***diazotates or nitrosamines as stabilized diazo compounds**

The history *of* diazonium ion chemistry is not lacking in conflicting interpretations of data and as such makes fascinating reading *for* anyone interested in the development of bonding theory and structure elucidation. Curtius²⁷⁷ noted in 1890 that an earlier structural assignment of Griess's^{11, 42} where he postulated that acidification of KON₂C₆H₅ furnished free diazobenzene, C₆H₅N₂, was incorrect. The salt was subsequently shown to be the so-called iso salt¹³ (45) which is also known as the stablc or *artti* salt. The normal salt **(43)** is also known **as** the *syrz* or labile salt. Their early history is treated by Cain²²⁴ and the modern (i.e. present) interpretation is summarized by Zollinger¹³²⁶ and by Laing⁷⁰⁹. The equilibria can be summarized as in Scheme 8. In accordance with current nomenclature the isomers should

probably be classified as *Z* for syn and E for *nnti.* The question as to whether the anti-diazohydroxide is an azohydroxide or its tautomer, the nitrosamine, has not been adequately answcred. There is reason to suspect that some assumed *iso*diazohydroxides are nitrosamines while others are anti-diazohydroxides.

The important synthetic features are: syn-diazohydroxidcs give the same typc of coupling reactions **as** diazonium ions; anti-diazohydroxides are stable to coupling but slowly cleave with acid to liberate the free diazonium ion. Clearly the position of equilibria and the structure of the *is0* compounds lie buried under **a** combination of substituent and steric effects and should be resolvable.

Iso-diazotates are normally formed by one of thc methods given in Scheme **9.**

When diazonium salts or syn-diazotate solutions are neutralized to a **pH** range **5.5-7.5** the diazoanhydrides arc formed. These are highly explosive, are converted by acids into diazonium ions and by bases into syn-diazotates.

2. Group 2: Diazoethers

The diazoethers occasionally result from failed coupling reactions³⁰⁶ but they are normally prepared by alkylation of *syn-* or *anti-*diazotates. Since the ions are ambidentate $[ArN=N-\overline{O} \leftrightarrow Ar\overline{N}=N-O]$ it is not surprising that aralkylnitrosamines are occasionally formed. The Na and K salts of the p-nitro compound furnished the nitrosamide with methyl iodide whereas the **Ag** salt yields the ether50, *62,* **182,** 485, **487, 490, 1249.**

3. Group 3: Diazocyanides, diazocarboxylic acids and related compounds

Two forms of diazocyanides arc known, E and *Z.* and the structural assignments are probably correct since they are sufficiently stable to permit a variety of transformations. A related isonitrile has been¹⁵⁹, ²²⁴, ²⁹⁹, 500, 503, ⁷²², ⁷²⁸, ⁷³⁹, ¹¹⁷⁵, ¹³²⁶ prepared by an 'unequivocal' synthesis. Thc synthesis requires the slow addition of the cyanide solution of the diazo compound in excess acid while the temperature is kept low. The *Z* isomer is formed first and this slowly rearranges to the *E* isomer⁴⁹⁰^{, 500}, ⁵⁰³. In the prcsence of excess cyanide the formation of an arcnediazoniumcyanidehydrogen cyanide complex occurs^{108, 489, 499}. In some cases the isomeric arylazo**carbonimidoylcarbonitriles** are

SCHEME 12

The *Z* isomer is normally lower melting than the *E* isomer. both are unionized, coloured and water insoluble. The isomerization to the E isomer occurs spontaneously in the melt, in solution or in sunlight. The ease and extent of isornerization are subject to steric and eixtronic effects. The *2* isomer couples and in the presence of Cu furnishes the nitrile.

An earlier proposal⁵⁵⁸ that one isomer was the isonitrile was disproven^{1191, 1192} through the synthesis of the isonitrile by generating it from **AgCN** and by dehydrating **1** -aryl-2-formyl triazencs with pyridine/SOCI, to form the isonitriles.

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The *E* nitriles undergo hydrolysis of the CN function to furnish the related acids. The CN function can also be converted into the CO_2R and CONR₂ functions⁴⁹¹.

The carbonamides react with H_2O_2 to furnish the azoxy derivatives^{\cdot}, ⁹⁸⁴⁻⁹⁸⁶ and cleave with alkali to furnish diazotates¹⁰⁶⁵.

The aniides undergo a somewhat general process when treated with cold concentrated hydrogen halide in aqueous media. The halogen enters the *pnrn* position to furnish a **p-halophenylhydrazidocarbonylamide1'z90.** The reaction occurs with other groups of similar electronic character, e.g. 2-pyridyl, 4-pyridyl, 2-pyridyl-N-oxide, benzoyl, $C(NH_2) = NOH$, $C(OH) = NOH$, $C(CH_3) = NOH$. benzoyl, $C(NH_2) = NOH$, $C(OH) = NOH$, $C(CH_3) = NOH$, does not occur with azobenzenes¹²⁹⁰. C(NH2)=NNHBz, C(NHp},)=Nph **74,** 186, **259,** *260,* **43R. 496, 617, 635, 976,** 1029, 1030, 1290 but

4. Group 4: The Diazosulphonates, their preparations, reactions and use as stabilized diazo compounds

Diazonium ions couple with $Na₂SO₃$ to furnish both labile and stable derivatives of the form $ArN₂SO₃Na$ with bonding through sulphur to the terminal diazo The reaction involves sulphite ions and not bisulphite²⁹⁰ and if the pH is too high, diazotates and other undesirable by-products are formed. If the medium is too acidic, reduction occurs^{484, 492}. Both the *Z* and *E* isomers can be reduced to the related hydrazines. The stable compounds do not couple but do react with chlorine and bromine to furnish the diazonium halides; thcy are also cleaved by lead nitrate and silver nitrate. The originally formed labile isomers isomerize extremely rapidly to the stable form but they can be isomcrized photochemically to the labile nitrogen53, **54.58,** 157, **158,** 221, **299, 392, 393,183.490,** 188,491, **487,** 499, **553,** *555, 665, 727,* **1103,** 1115, 1231, 1222 formGIC, **1337, 1344,** ¹³⁴⁷

In a chemically similar reaction, aryl-sulphinic acids couple with diazonium salts to furnish sulphones $(ArN_2^+ + Ar^sSO_2H \rightarrow ArSO_2Ar^+ + H^+)^{500}$. Diazonium salts also react with alkyl and arylsulphonamides to furnish sulphones^{317, 318}; *syn-*diazo cyanides have also been found useful for this type of coupling with sulphinic acids⁵⁰⁰.

The mixed alkyl-aryl-diazosulphones decompose by **a** homolytic path in solvents such as benzene and heterolytically in highly polar media¹⁰³. The product mixtures from the homolytic reactions were $SO₂$, biphenyl, azobenzene and phenylazobenzene **(14** : 32: 3 : 2). In polar media (acetonitrile) a 71% yield of acetanilide was obtained⁶³⁸. When the benzyl derivative was employed in benzene, **SO₂**, benzaldehyde plienylhydrazone, IV-benzyl benzaldehydephenylhydrazone and biphenyl were obtained (66 : 55 : *6* : **Z)500.** An earlier showed that phenyl benzenediazosulphone furnished primarily biphenyl, benzenesulphonic acid and only *5%* of the diphenyl sulphone.

5. Group 5: Diazoperhalides

Diazonium halides react with halogens and mixed halogens (e.g. ICI) to furnish compounds of the type $ArN_2XX'X''$ (X = halogen) ¹¹⁰³. These compounds contain active halogen and are most probably of the form $ArN^+_{z}X^-_{3}$; as such, they are reasonable halogenating reagents. Wher, boiled in alcohol they furnish the aryl halide179. Treatment of the aryl diazonium perbromides with ammonia offers **a** route to aryl azides⁴⁶³.

6. Group 6: Diazosulphides

colleagues^{157, 159}, much of the chemistry of this class of derivative is now clarified As **a** conscquencc of extensive studies by Brokken-Zijp arid her Dutch and will probably serve as a foundation and model for studies on related systems. van Beck'223 has developed **a** general synthesis for a number **of** *E* and *Z* diazosulphides. An alkanethiol was allowed to react with an equivalent amount of the diazonium fluoborate in acetone at 0 *"C* and aqueous sodium hydroxide was added over a period of **1** h. The isomers either crystallized from solution or were extracted with benzene. Afler **2 h** at 90 "C in isooctane, thermal equilibrium between *E* and *Z* isomers was established. The correctness of the structure assignments have been verified for some compounds by Brokken-Zijp¹⁵⁷, who also examined the action of light **on** both the *E* and *Z* isomers. Operating at 405 nm in methanol, pure *E* or pure *2* lead to mixtures of both. At 300 nm, E isomers and decomposition products (sulphides, disulphides) were obtained. Irradiation of the pure E isomer at 300 nm generated only the *Z* isomer. The 4-cyano-, 4-nitro- and **3,5-dichloro-4-dimethyl**amino benzene diazo-r-butyl sulphides and phcnyl diazo-r-butyl sulphides were studied.

The term 'diazosulphide' as employed by Saunders¹¹⁰³ was used by him to describe diazothioanhydrides $(ArN₂SN₂Ar')$ and $ArN₂SR$ instead of only the latter as employed in the current Iitcrature. The thioanhydrides arise from the reaction of neutral diazonium salts with hydrogen sulphide. The thioanhydrides couple with alkaline β -naphthol⁵⁶ and furnish the biaryl upon heating in refluxing benzene⁵⁶. Some of the thiodiazoanhydrides are extremely unstable and even explode under water at $0 °C$ ^{334, 561, 852}.

The diazosulphides are inore stable than their oxygen analogues and fail *to* couple or cleave with cold acids or bases. They do decompose thermally **via a** homolytic process¹²²⁵.

If **H,S** is passed into acid solutions **of** aryl diazonium ions, the main product **is** the aryl-diazoniumthiol hydrosulphide $(ArN_2SH·H_2S)$.

C. Class C: Derivatives in which the Diazo Function Disappears but the Diazo Nitrogen Atoms Remain in the New Molecule

1. Group 1: Diazo-amino and -imino compounds or triazenes^{224, 1103, 1326}

The reactions of arene diazonium ions with amines date from Griess's initial studies⁴⁴⁸ where he obtained 1,3-diphenyltriazene by reacting nitrous vapours with an excess of aniline. The process is favoured by neutral conditions and is suitable

SCHEME 14

for reactions with aromatic or aliphatic amines. With aromatic amines in which either aryl unit has a vacant *para* position, treatment with acid leads to coupling with the aromatic nucleus. In addition, equilibration is rapidly reached between both isomers and with acid, cleavage to all possible diazonium and ammonium ions results. Such a situation is simplified by blocking a *para* position.

This sensitivity to substituents is sufficiently great that *meta*-toluidine undergoes C substitution rather than N coupling⁸³.

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With primary aliphatic amines, *bis* coupling frequently occurs and 1,5-diaryl-**3-alkyl-pentaza-l,4-dienes** result. **It** is possible to generate the aryl alkyl triazenes by adding a cold neutral solution of the diazonium salt to a cold solution of the amine in the presence of sodium carbonate. The resulting triazenes are moderately stable. In the case of the methyl and ethyl p-tolyl systems, useful reagents for esterification of carboxylic acids result³⁰¹, ⁴³⁵, ⁷⁶⁴, ⁹⁹³⁻⁹⁹⁵, ¹²⁸¹, ¹²⁸³

This method of esterification offers some advantages over diazoalkanes. Most notable is the ability to generate tertiary esters in modest yields. These are not available from diazoalkanes. With secondary amines, only a **very** small degree **of** rearrangement is observed.

In the course of decomposing the secondary derivatives, extensive alkylation of the aromatic nucleus occurs. The process proceeds with a high degree of stereospecificity and retention of configuration even when the alkyl group is α -phenethyl.

8. Synthetic applications *of* diazonium **ions 263**

The simple monoaryl triazene, $C_6H_5N_2NH_2$, does not arise by the reaction of the benzene diazonium ion with ammonia but is generated by reduction of phenyl azide with stannous chloride in ether³⁰². Griess^{458, 462} obtained an explosive substance from ammonia and diazobenzene. Von Pechmann¹²⁵¹ ran a similar reaction with p-toluene diazonium ion and ammonia and obtained $p - C_7H_7N_2NHN_2C_7H_7-p$. The structure **was** verified by methylation which gave the same compound obtained from methylamine. In Griess's original attempts he obtained $C_6H_5N_2C_6H_4NH_2$.

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The reaction of the *bis* aryl triazenes with acids furnishes a phenol, nitrogen and an amine. The reaction gives two phenols and two anilines when asymmetric starting materials are employed. In the presence of copper(1) salts and halogen acids, the aryl halide is formed in place or partially in place of the phenol. The amino nitrogen is sufficiently electron poor in *bis* aryl triazenes to render the hydrogens acidic and the sodium salts are known⁷⁵⁴. The systems also are sufficiently good ligands to permit formation of platinum⁴⁶⁵, cupric³¹⁹, nickel³²⁰, mercury¹²⁷¹ and palladium⁷⁵⁵ complexes.

The *bis* aryl triazenes and alkyl-aryl triazenes decompose thermally by a homolytic proce&O7, **638, 642, 820, 848, 851, 1024-10?Ue**

The coupling of diazonium salts with hydrazines and hydrazones normally leads to formation of tetrazenes¹⁹⁹, ³⁷⁷, ³⁷⁸, ⁵¹³, ⁵⁵⁸, ⁷⁶¹, ⁷⁶², ¹¹¹³, ¹²⁸⁷, ¹³⁰⁴. A variety of other triazene reactions is summarized by Smith¹¹⁵⁹.

The diazo tetrazol **(97)** is reported to couple with hydrazine in a manner not observed with the benzene diazonium ion to form the hexazadiene *(98)572.* 577. Tetraryl hexazadienes can be prepared by oxidation of diary1 triazenes. The products undergo slow decomposition to azobenzenes $(99 \rightarrow 100 \rightarrow 101)$.

When 1,3-diaryl tetrazenes are oxidized with permanganate ion, explosive compounds believed to be octazatrienes are obtained $(102 \rightarrow 103)^{1305}$, 1306 . The chemistry of nitrogen chains of three or more N atoms is reviewed in Sidgwick¹¹¹³ and in $Smith¹¹⁵⁹$. Lieber and Smith⁷³⁰ have discussed poly-aza systems derived from aminoguanidines.

When diazonium ions condense with aryl or alkyl hydroxylamines, triazene oxides or their tautomers the hydroxy triazenes result. These compounds undergo both N and O alkylation.

If 4-substituted aryl diazonium salts are treated with methyl hydroxylamine hydrochloride and sodium carbonate at 10 *"C,* **l-aryl-3-hydroxy-3-methyl** triazenes result³⁹³ (104 \rightarrow 105). When ethyl anthranilate is diazotized and treated with

alkylamines, the related triazenes are generated. These in turn are converted smoothly in refluxing ethanol or over basic alumina into 4-oxo-3-alkyl triazenes⁷¹⁵ $(106 \rightarrow 107 \rightarrow 108)$.

This is in striking contrast with the results with diazotized anthranilic acid $(89 \rightarrow 90)$.

The oxides also result from peracid oxidation of triazencs. This latter approach leads to both possible isomers (1 and **3)** when asymmetric triazenes are employed although the presence of an electron-rich aromatic system favours the oxygen ending on the N adjacent to the electron-ric11 systcmSS, *60.* **61. 68, 69, 79. 131. 136, 31.1. 327. 805.** *806, '028.*

2. Group 2: Oxidation products

In addition to those oxidation processcs mentioned in earlier sections the diazonium function can be oxidized. Both *Z* and E diazotatcs undergo oxidation to furnish aryl nitroamines. Ferricyanide, hydrogen peroxide and permanganate have all been used under alkaline conditions⁷⁵, ¹²⁷, ³⁷³, ⁴¹³, ¹⁷², ¹¹⁸⁶, ¹²⁰³.

3. Group 4: Reactions with nitrogen bases. The formation of aryl azides

Azides have been the subject of a monograph in this series 2^{74} . Diazonium perbromides⁴⁵⁹, plumbochlorides²³⁹ and iodochlorides²³⁹ react with ammonia to furnish azides and diazonium salts react with azidc ion to furnish aryl azides, presumably via intermediate pentazoles and pcntazatrienes.

A route to aryl azides known as the 'Dutt-M'armall Reaction' involves the action of an aryl diazonium ion upon a sulphonamide such as p -toluene sulphonamide. The method is theoretically only 50% eficient in aryl diazonium ion since a second molecule couples with the liberated sulphinic acid¹⁵⁴, ³¹⁷, ³¹⁸. The reaction of azide ion with diazonium ions is treated as a Type E process.

4. Group 6: Azo compounds formed by coupling reactions (C coupling)

The coupling of aryl diazonium ions with phenols and amines to furnish azo dyes is not a majcr subject of this review but has bcen treated with a fair degree of thoroughness in texts by Fieser³⁷⁰ and by Noller⁸⁹⁵⁸.

We shall therefore limit the treatment here to some of the less well-known coupling processes and a brief treatment of the common coupling reactions.

Coupling to form C-azo derivatives received extensive attention from chemists almost from thc time that diazonium salts were discovered and lead to the development of the chemical industry in Germany with a heavy emphasis on thc synthcsis of dyes. Almost all azo dyes are prcpared by the coupling process. The dyc-forming reactions are normally performed in aqueous media and the reactions are pH dependent. Many compounds will couple in acidic media as well. The general pH limits are *3.5-7.0* for aryl amine substrate and *5.0-9.0* for phenols. An cxamination of the stability graph for diazoniuni ions reveals that this is thc same pH range where the stability is at a minimum. In general coupling occurs with phenols, aryl amines and active methylene compounds. Included in this group are such non-classical aroniatic phenols as the tropolones and proteins.

The aromatic amines couple in the *para* position when an aniline is involved but otherwise couple in the position of maximum electron density and minimum ΔE_{π} as predicted by a simple PMO analysis²⁹². Thus β -naphthyl amine couples in the 1-position and not the 3-position. Using the PMO analysis (Scheme 22) quickly leads to an explanation of why 1,3-diamino benzenes are more reactive than the 1.2-isomers or the aniltnes. The same analyses hold for phenols and polyhydric phenols.

8. Synthetic applications of diazonium ions *261*

The reaction is obviously facilitated by net electron-donating groups (negative Hammett σ values) and hindered by those groups which are net electron withdrawing. Since the anilines couple in the para position exclusively, the presence of substituents at that position stops the reaction from occurring unless strong activation of an *ortho* position is operating. With hetcrocyclic aniines such as aminopyridines, coupling is not **a** very facile process because of the inherent inertness of pyridines to electrophilic attack. The presence of α and α' amino substituents increases the activity of the γ position towards attack. A PMO analysis would seem to indicate that β substituents should be more effective.

SCHEME 22

The coupling of aminoquinolines has been examined in some detail¹⁰⁵⁴ with benzenediazonium ion as the electrophilc (Table 1).

TABLE **¹**

Coupling at Triazene ^a C-4 N.R. ⁴ C-6, C-8 C-5 C-8 C-5 Amino function at C-2 C-3 C-4 C-5 C-6 C-7 C-8				

aTriazcne = coupling with the amino function *of* thc aminoquinoline. N.R. = no react ion.

With aminophenols coupling can, in principle, be controlled by either functional group since the pH dependence of the coupling reaction is different for phenols and anilines. This can often be realized and a number of examples exist^{170, 367, 369, 420.} **424,** 1061.

The coupling with proteins primarily involves the three amino acid units froni tyrosine **(117),** histidine **(118)** and tryptophan **(119).** This phenomenon was behind

a process employed for dycing silk. Thc obvious biochemical applications of the reactions have not been overlooked. Since sevcral amino acids have functionality capable of forming ArN_2-X-R units (where X is an amino or thiol function) coupling is not limited to proteins containing tyrosine histidine, tryptophan or related units.

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Unlike the anilines, phcnols can couple in the **orrho** position **as** well as the normally more reactive *para* position. Occasionally the position of coupling in naphthols is dependent upon the diazonium ion employed. Thus 1,5-dihydroxynaphthalene is attacked at the 2-position (counter to the PMO analysis) by diazotizcd o -aminophenolsulphonic acid but at the 4-position by diazotized anthranilic acid³⁸².

a. *Corcplirrg* at *actiue CH, and* CH *groicps.* Therc is a large number of compounds which possess active methylene groups and, not surprisingly, a large proportion have been subjected to diazonium-coupling conditions. Some examples include acetoacetic ester⁸⁰⁰, nitroethane⁷⁹⁷, malonic ester⁷⁹⁶, acetone dicarboxylic acid¹²⁴⁴, β -nitroacetophenone⁹⁶⁷, glutaconic ester⁵³⁴, acetophenone- β -sulphonic acid⁹⁶⁹, many cyclic ketones²⁸⁴, ⁷³⁵, pyrazolines²⁰¹, ⁵¹⁰, ¹¹⁰⁴, ¹²³² and the nitroparaffins^{48, 71, 78} **361,** 798, 709, **801,1??1.**

The coupling products frequently exist as the hydrazones if the hydrogen replaced is borne on a $-CH_2$ - or CH_3 - unit. This point was the source of considerable controversy6?. 172- *li3,* **343, 611,** 7n6s **124J,** and one should probably treat some of thc related literature with a degree of caution. The processes can be cxceedingly complex and furnish a variety of products. Thus coupling of phcnyl diazoniuni ion with nitromethane furnishes no fewer than 12 products^{249, 335, 362, 1049, 1145, one of which} results from rearrangement while recrystallizing from refluxing ethanol³⁶², ¹¹⁴⁵.

The overall processes can be conveniently subdivided into those occurring when a hydrogen is replaced in a formal metathesis¹³⁴³ and those of the Japp-Klingmann type where a radical is split off such as an acyl or carboxylate function 1344 (equations 1 and **29n1).** The processes have been well studied and by 1958 ovcr

$$
ArN_{2}X + R - C - H \longrightarrow ArNHNCR_{2} + HX
$$
\n(1)
\n
$$
R
$$
\n
$$
H
$$
\n
$$
H
$$
\n
$$
H
$$
\n
$$
H
$$
\n
$$
ArN_{2}X + R - C - R \longrightarrow ArNHN = CR_{2} + LX
$$
\n(2)

700 pertinent references could be found^{971, 981}. The major determinant relative to whether an aryl diazonium species will replace a hydrogen is the activity of the hydrogen and not the nature of the diazonium species.

Normally the reaction will not occur with ketones lacking some additional activating group, and with attack on a methyl group there is a strong tendency to generate a formazyl ketone^{10, 47, 1246}. Similar behaviour occurs with nitromethane⁵⁹², ¹¹³², ¹¹⁹⁵.

$$
\begin{array}{ccccc}\n & & O & N_2Ph \\
 & & & \| & I \\
\text{CH}_3\text{COCH}_3 + 2 N_2^* Ph & & & \text{CH}_3C - C = N - NHPh\n\end{array}
$$
\n(3)

and with attack on a methyl group there is a strong tendency

\nrmazyl ketone^{10, 47, 1246}. Similar behaviour occurs with nitro-
\n
$$
\begin{array}{r}\n0 & N_2Ph \\
\parallel & \parallel \\
\end{array}
$$
\nH₃COCH₃+2 N₂⁺Ph

\n—A CH₃C-C=N-NHPh

\n(3)

\nCH₃NO₂+2 N₂⁺Ph

\n—A CH₃COCH₃

The formazans undergo a fairly general cyclization reaction when treated with strong acids such as sulphuric acid. The process kno\vn **as** the 'Eaniberger Triazene Synthesis' furnishes benzo-1,2,4-triazenes^{2, 42, 43, 46, 968}. The chemistry of formazans has been reviewed¹⁰²⁸.

SCHEME 24

Benzenediazonium chlorides couple with **2-benzoyl-2-cyano-thioacetanilides** in aqueous ethyl alcohol to furnish 2-arylhydrazono-3-oxo-3-phenyl propionitriles³¹².

(i) Esters, acids and their derivatives. Simple acids do not generally undergo condensation but a-keto acids (e.g. pyruvic) do react. With pyruvic acid **two** molecules of diazonium salt are consumed⁴⁷ in direct analogy with acetone and nitromethane. 9-Keto acids undergo **loss** of carbon dioxide and if an excess of diazoniuin ion is used a second molecule will react^{43, 610, 1055. Similar results accrue with acetone} dicarboxylic acid^{1244, 1248}.

Good yields of the mono hydrazones can be obtained by employing the esters and onc equivalent of the diazonium salt but if *2* moles of diazonium salt are employed with acetoacetic esters, a second diazo group enters with loss of the

acetyl group^{76, 1032, 1033, 1293. A similar phenomenon occurs with diethyl oxalo-} acetate¹⁰³², ¹⁰³³, ¹²⁹³, where the second mole of diazonium ion leads to ejection of an

ethoxyalyl group. According to Parmerter these are the only two groups which are susceptible to loss when attack is on methylene. Other leaving groups are of course encountered with the Japp-Klingemann reaction *(cide infin).* Coupling also occiirs with acetacetanilides and malonimides¹⁴⁻¹⁷, ¹⁸⁶, ²⁴⁴, ³⁶⁸, ⁴⁹², ¹⁰⁷⁴, ¹¹³⁹, ¹¹⁶², ¹²⁸⁴

It **is** not particularly surprising that malonic acid, like nitromethane, is capablc of multiple reactions involving I, *2* or 3 moles of the elcctrophile with both *tiiotzo* and *bis* products resulting from the reaction with equivalent amounts of reagents^{195, 1245}; with monosubstituted malonic acids, *bis* substitution readily occurs~os. **1259.** alyl group. According to Parmerter these are the only two groups we
tible to loss when attack is on methylene. Other leaving groups are c
ntered with the Japp-Klingemann reaction (*vide infra*). Coupling als
cetacetanilid

$$
CH2(CO2H)2 - \longrightarrow ArNHN=CHCO2H
$$
\n(133) (134)
\n
$$
--- \longrightarrow ArNHN=CM2A1NHN=CM2A1
$$
\n(135)

SCHEME 27

A limited number of reports of arylacetic acids and esters reacting with diazonium ions exists. The reaction goes moderately well with the 2,4-dinitrophenylacetic acid and its derivatives (apparently as a consequence of the highly electron-withdrawing aryl group). The acid reacts twice with loss of $CO₂$, whereas the ester consumes only 1 mole of diazonium ion.

(ii) Nitriles, nitroparaffins and sulphones. The cyano function also can serve as an activating group and a variety of dinitriles, β -cyano-carbonyl compounds, β -sulphononitriles, β -nitronitriles and β -iminonitriles undergo attack by aryl diazoniuln jonsl13, **1.15.** *250,* **273, 338, 311, 346. 384,** *705,* **738, ;:I, 1114, 1172, 1173,** *1211,* **1113. 12d0, 1273** Sulphones^{33, 248, 592, 1211, 1213} and nitroparaffins^{48, 71, 197, 434, 580, 613, 997-1001 are also} quite reactive. Hydrazones are the normal products resulting from attack of a methylene group. Subsequent attack of aldehyde hydrazones derived from activated methyl groups are responsible for thc formation of *his* aryl diazo-derived products. The reactions when carried out as two-step proccsscs are good synthetically and furnish formazans^{789-792, 1317, 1318. The aryl hydrazones of α , β -diketobutyric esters} and a-kcto-acids undergo attack by a second molecule of aryl diazonium salt in processes formally analogous to the Japp-Klingemann reaction^{44, 73, 198, 406, 713.} **1034, 1074.**

Bcnzcnediazoniuni chlorides couple \villi **2-benzoyl-2-cyano-thioacetanilides** in aqueous alcohol to furnish 2-arylhydrazono-3-oxo-3-phenylpropionitriles³¹².

(iii) Ketones. Aryl diazonium ions attack ketones bearing an electron-withdrawing **7OG, 731, i\$D, 7x7, 1057, 1'51, 1283, 1'294, I395** . The reactions are on occasion rather varied. group to the carbon α to the carbonyl group¹⁰⁰, ¹⁰¹, ¹²⁰, 111, 241, 281, 297, 342, 599, 600, 703.

Thus the oxalylindene (136) undergoes simple substitution¹²⁹⁵ while its benzo analogue (138) undergoes a Japp-Klingemann reaction⁷⁰⁶ as does the pyridine derivative (140) which furnishes a formazan¹⁰⁵⁷.

(iv) The Japp-Klingemann reaction. The Japp-Klingemann reaction⁹⁸¹ has found extensive synthctic application. The processes differ from those discussed above in that only a single active hydrogen is replaced and a functional group attached to the carbon atom being substituted is lost. The reaction strongly resembles formal attack by +NO **1208.** (This parallel occurs in transition metal chemistry as well, *vide infra.*) Fine Supp-Ringemann Feachon: The state
we synthetic application. The processes
ly a single active hydrogen is replaced a
atom being substituted is lost. The
by + NO¹²⁶⁸. (This parallel occurs in
ra.)
R₁
R₁
R₁
R₁

SCHEME 29

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The mechanism of both processes undoubtably involves initial generation of a stabilized carbanion^{510, 1104}, then attack by the diazonium ion to furnish a diazo derivative which can then cleave by nucleophilic attack upon the potential leaving group followed by ejection of the product, its tautomeric form or the stabilized anion of the product.

The enhanced stability of both of the final products (the desired product plus the final form of the leaving group) may serve as the driving force for the reaction^{117, 177},
199-202, 257, 300, 303-306, 336, 338, 343, 344, 390, 512, 513, 515, 516, 692, 620, 733, 802, 1121, 1123, 1132, 1137, **1140s 1115+ 1237. 1238, 1250. 1302.** If we extend this general mechanism to the Japp Klingemann reaction using a general refinement suggested by Dewar²⁹⁰ for ester hydrolysis, we obtain Scheme **30.**

Normally the Japp-Klingemann reaction requires two or even three electronwithdrawing substituents such as carbonyl, cyano, nitro or sulphonyl groups being present. However, electron-deficient aromatic systems such as fluorene, pyridine, quinoline and dinitrophenyl can scrvc as activating substituents as can the halogens. Halogen rarely serves as the leaving group. Perhaps this is a consequence of the fact that normally the leaving group is attacked by a nucleophile and with halogens this requires a formally positive halogen. Hence one would predict that the tendency electronegativity^{340, 341}. This interpretation accounts for the apparent anomalous behaviour of 3-bromotriacetic lactone12*8 **(148).** Acetoacetic esters undergo the

Japp-Klingemann reaction with the loss of the acetyl group and the resulting products are N-substituted derivatives of csters of dchydro amino acids. Reduction offers a ready route to the amino acid esters and treatment under Fisher-indole synthesis conditions furnishes indolesl, **.la, 128, 174-176. 180, 321, 347-330, 352-351, 356-300, 525, 58.1, 610, Gll, 683, 734. 1136, 1185, 12GO**

With 1,3-diones, similar behaviour is observed and one has a route to α -amino ketones^{298, 339. This approach is complementary to the use of p -tosyl azide in a diazo} transfer process which furnishes the α -diazo ketone which can also be reduced to the related aniin~~~, **337, I3l1.**

The Japp-Klingemann reaction of 2-carboalkoxycyclanones offers a ready source of ω -carboxy α -amino acids and of 3-(alkyl- ω -carboxy) indoles. Since the starting materials are readily available by **a** Dieckmann cyclization or by reaction of the cyclanone with dimethyl carbonate, this offers a very ready route to both systems²⁹⁵, ²⁹⁶, ²⁹⁵, ³⁵⁵, ⁵¹⁴, ⁷³³. An interesting variant on this approach employs a 2-acylbutyrolactone. Subsequcnt ring opening can be performed with a variety of nucleophiles to furnish lactones, esters or hydroxy acids^{345, 509, 1161.}

(v) Coupling with tropolones. The tropolones are prone to coupling with diazonium ions. The reaction is of sufficient reliability to permit its use as a diagnostic tool for the presence of a substituent at C-5. The processes offer ready routes to 5-amino tropolones via subsequent reduction²⁶¹, ³⁰⁸, ⁵¹⁹⁻⁵²¹, ⁶³⁹, ⁶⁷¹, ⁹¹⁰, ⁹¹²,

913, 916, 917, 921-925, 928, 930, 931, 934, 937, 939, 1135, 1313. The reaction is somewhat sensitive to steric effects and both 4-isopropyl and $4-t$ -butyl tropolone fail to couple⁹¹⁹.

The presence of halogens at positions 3 or 3 and 7 greatly decreases^{911, 926} the reactivity of the tropolone. For the test to be fully valid, it is necessary to perform the reduction step, for 5-methyl, 5-hydroxy, 5-halo and 5-isopropyl tropolones furnish coloured complexes which are cleaved by reduction without introduction of the amino function^{332, 913, 926, 930, 935, 942. Several other 5-substituted tropolones} fail to form complexes^{843, 916, 939}.

5. Group 7: Complexes of aryl diazonium and alkyl diazonium ions and diazo compounds with transition metals

Until relatively recent times onc might have dismissed out of hand the possibility of isolating and characterizing transition metal complexes of diazonium ions or of diazo compounds. This was **a** direct consequence of the fact that many transition metals catalyse the decomposition of both diazo and diazonium ions. There are of course a number of old examples \vhere aryl diazonium salts formed double salts with complex ions of such metals as platinum and mercury, but in those cases the diazonium ion was serving a rolc similar to that of an **alkali** mctal cation. During the past 11 years a sufficient number of aryl diazonium-transition metal complexes have been isolated and characterized to warrant a review with over 100 references¹¹⁹³. With diazo compounds, the nunibcr of examples is somewhat more limited, but in both cases well-documented examples now exist where the $-N_2$ function serves as a ligand with bonding to the metal occurring through the terminal nitrogen. The fact that bonding occurs through the nitrogen is probably the cause of their stability. There is reason to believe that the loss of nitrogen from diazo alkanes in the presence of metal catalysts proceeds via attack on the diazo carbon¹³¹².

The nomenclature which has been used to describe these complexes in the literature is not uniform. Thus the terms 'arylazo', 'aryldiazo'^{865, 867, 868} have been employed for the aryl complexes although the compounds can be named **as** derivatives of diimine⁵²³, diimide⁶⁰² or diazene^{385, 699}. Some workers have employed the terms 'hydrazido dianion'³²³ and 'phenyl hydrazido dianion'⁷⁶⁶ to describe the protonated complexes. It is perhaps unfortunate that ultimately the arrogance⁴⁰⁴ of *Chemical Abstracts* will probably dominate¹¹⁹³ and the *diazene* nomenclature will be employed and the **I.U.P.A.C.** proposed term, *aryl dinzennfo* will not become fixed in the literature. The use of *dinzonnto* (aryl or alkyl) would appear even more logical and consistent with the fact that the term *diazo* has been used for 111 years.

This term has been in usc for at least a hundred years before the American Chemical Society took upon themselves the rolc of final arbitrator for Chemical Nomenclature^{311, 401}, 960

The complexes appear to fall into two basic classes: those in which the $-N_2$ function formally furnishes thrce clectrons to thc mctal to satisfy the 18-electron requirement of filling the valcnce shell, and those which formally furnish a single electron to satisfy the IS-electron requirement. These groups can be identified by X-ray crystallographic analysis of the $M-N-N$ bond angle. The three-electron donor situation leads to angles in the range 170-179[°] ²⁹, ²³⁴, ³¹³, ³⁹⁶, ⁷⁴², ⁷⁸⁵ and the one-electron donor system leads to a bond angle of $125 \cdot 1^{\circ}$ ⁴²⁶. Sutton¹¹⁹³ has discussed the bonding theory which leads to predictions of **180"** and 120".

There **is** a very limited number of alkyl-diazonato complexes and structural analysis by X-ray is lacking. The spectroscopic parameters presented are totally consistent with the claimed structures.

The formation of the diazonato complexes is most commonly realized by direct reaction **of** the diazo compound or the diazoniurn salt with **a** complex (ionic **or** neutral) with or without the displacement of an existing ligand.

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The number of diazoalkane complexes with transition metals is small. For **a** summary of the aryl diazonium based complexes readers should see Sutton¹¹⁹³.

The aryl-diazonato complexes tend to be basic at the nitrogen adjacent to the metal atom¹¹⁹³ and undergo protonation on nitrogen. However, since they are prepared as ends rather than intermediates, their chemistry is not well characterized.

D. **Class** *D:* **Type 6 Ring Compounds in which the Diazo Nitrogen has been lost. The** *Pschorr Type Syntheses*

Detailed discussions of several of these classes of heterocycles have appeared in the form of monographs and as such will be described only in terms of their syntheses unless there exist additional diazo- or diazoniuni-type reactions. These reactions are treated below. The chemistry of the members of groups in which the nitrogen is lost are not of concern here and we shall only treat their generation from diazonium compounds. There are obviously other means of obtaining many of the products which do not involve diazo or diazonium chemistry and standard references and reviews cover most, if not all, alternate methods.

Y. Synthetic applications of diazonium ions *217*

In recent years the chemistry of aromatic systems has increasingly been relegated to the briefest of treatments in standard textbooks. This is perhaps a reflection of the amount of research activity in the area or the simple increase in the total amount of material covered. **As a** direct consequence the chemistries of diazonium ions normally discussed in beginning texts are the reactions in Class E and coupling with an occasional reminder that certain orrho-substituted anilines arc not suitable for diazonium reactions. The chemical reason behind this bccomes evident when one notes that the so-called 'forbidden' groups are potential nucleophiles ideally located adjacent to electrophilic diazonium groups. This proximity effect means that systems incapable of bimolecular reactions may well be capable of unimolecular cyclization. The *orflio* function need not of itself be a nucleophile or have nucleophilic character if the conditions of the reaction can cause the substituent to generate a transient intermediate capable of serving in that role. Thus an enolizable ketone can readily fulfil the role of an ethylene.

1. Subclass a: Diazo nitrogen retained

a. *Group* I: Cimolines. The presence of a vinyl group in the *orrho* position of **an** aniline or other aromatic primary amine may lead to the formation of a cinnoline378, **964, 1252. 127G** upon diazotization. The configuration around the vinyl group need not be *cis* for both *cis* and */ratis* 1,2-disubstituted ethylenes can participate¹²⁸⁵. The basic reaction shown in Scheme 37 has some limitations which

have been detailed by Simpson¹²⁸⁵. He has concluded that when $R = H$ or $CO₂H$ and $R' = \text{aryl}$, the reaction does not occur. If $R = \text{aryl}$, R' can be aryl also. When $R' = \text{aryl}$, there is a possibility that a Pschorr condensation will compete^{30, 48, 250,} **289, 603,** 621 (Scheme **38).** This process **is** discussed in detail under Group E, Class *5,*

SCHEME 38

reactions. The synthesis of cinnolines from o -vinyl diazonium ions is known as the Widman-Stoermer synthesis¹², ²⁵, ¹¹⁷, ¹⁶¹, ¹⁶², ⁷²³, ⁷²³, ⁹⁴⁴, ⁹⁶⁵, ¹¹²⁰, ¹¹²³, ¹¹⁴⁷, ¹¹⁴⁸, ¹¹⁵⁰, ¹¹⁷⁰, 1285; from o-acy] diazonium ions as the Borsch synthesis13. **28. 634-636. 683,** 724, **045,** 1117. **1122,** and from o-alkynyl diazonium ions as the Richter synthesis⁶⁰³, ¹¹¹⁶, ¹¹²¹, ¹¹⁵³, ¹²⁵². Baumgarten⁹⁰ has modified the Borsch synthesis as shown (185 \rightarrow 187 \rightarrow 188).

b. *Group 2: Indazoles*. In certain cases a methyl, methylene or methine group is rendered sufficiently acidic by substituents to make it **a** potential *C* acid. When this occurs adjacent to a potential diazonium ion, cyclization may occur to furnish 3-H-benzpyrazole which may if **a** hydrogen is at **C-3** rearrange to furnish the 1 -H-benzpyrazole (indazole) (Scheme **39).** The presence of electron-withdrawing

groups as R^1 , R^2 , R^3 , R^5 will facilitate the reaction^{65, 409, 803, 817, 818, 906, 1296, However,} there exists the possibility that the resulting indazole will be coupled to unreacted diazonium compound if there is an electron-withdrawing group at $C_6(R^5)$ or **C-4 (R3)** but not at **C-5** or C-7 (e.g. Scheme **40).**

The difference between the two substrates would appear to be a consequence of the 6-nitrobenzpyrazole having a more basic C-3 due to contributions from the anilino nitrogen. This would be strongly suppressed by electron-withdrawing groups in the *5-* or 7-positions.

c. Group 3: Benztriazoles. Bis-diazotization of ortho-diamino systems is extremely difficult, and under a wide range of diazotization conditions they afford
triazoles^{7, 148}, 330, 339, 468-470, 567, 632, 708, 814, 815, 903, 1078, 1103, 1127, 1219, 1257, 1321, 1323, 1325 The reaction was discovered by Hofmann in 1860 when he worked with nitro-ophenylenediamine⁵⁶⁷. The driving force is sufficiently great that acetylation of one nitrogen does not prevent **the** reaction and mono-alkylation has no effect. Asymmetrically substituted o -phenylene diamines may lead to two isomeric triazoles. At the time of their discovery, a debate as to structure occurred between Griess and his adherents and Kekule and his. Much research was directed towards preparing and isolating both isomers. Kekule's⁶³² structure (204 and 205) would lead to two products, Griess's to one (206) ^{469, 470}.

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Finally, Morgan and colleagues^{814, 815, 1321 prepared the two N-acetyl derivatives} of (204) and (205) . The former was known as the α -isomeride and the latter the β -isomeride. Unequivocal syntheses were employed and the argument was closed by their hydrolyses to furnish the same 'compound' (mixture) which on reacetylation furnished a mixture of the α and β acetyl derivatives with the α -isomeride being predominant. They claimed that they were able to separate the two triazoles which had slightly different absorption spectra and did not equilibrate in organic solvents. The question of which tautomer is dominant in these systems has been extensively examined using modern instrumental techniques^{625, 704, 808, 957,} **1003, llG3.**

When ortho-amino diphenylamines are submitted to diazotization conditions, they readily undergo conversion to benzotriazoles. The triazoles were shown to be thermally unstable by Graebe and Ullmann and to offer a ready route to carbazoles103. **??Q, 442.**

d. *Group 4: Benztriazines*. Normally diazonium ions will not couple with an amide to furnish **a** triazene but ihe intramolecular process occurs with great facility^{371, 601, 700, 1103, 1202, 1270 (Scheme 43) and with a wide range of substituents for}

 (216)

SCHEME 45

 (215)

8. Synthetic applications of diazonium ions **281**

R and R' to furnish 1,2,3-benzotriazines. Examples are known employing *bis* acyl hydrazides as well as diamides³⁷¹, ³⁷², ⁵³¹, ⁷⁰⁰, ⁷⁷⁰, ⁷³⁵, ⁹⁸³ (Schemes 44 and 45); diazoamine esters^{240, 770} (Scheme 46) ureas⁶⁰¹; diazonium salt esters (Scheme 46)¹³¹⁶, amino benzpyrazoles^{63, 66}, ⁶⁷, ⁴⁴⁶, ⁶⁵⁵, ¹²⁷², ¹³⁰⁹, ¹³¹⁰ (Scheme 46); amino benzamide oximes (Scheme 49)^{110, 990}; cyano hydroxylamines (Scheme 47)^{63, 66, 67, 1202; amino} oximes (Scheme 47)^{64, 72, 778, 1189; amino benzyl aryl hydrazines (Scheme 48)¹⁸⁸;} amino benzyl amides (Scheme 48)¹³⁸, ¹⁹⁰-193, ¹⁰⁴⁷, ¹²⁵³; amino thioamides¹⁰⁵⁰, 1,8-diamino naphthalenes (Scheme 49)^{7, 329, 1086, 1124, 1258, 1349, 1355; heterocyclic amino} carboxamides⁶¹⁸; amino *ortho*-2-pyrazoles and amino *ortho*-5-pyrazoles, their benzopyrazole analogues¹²⁴¹, ¹²⁴² (Scheme 49); rearrangement of a benzpyrazolodiazene⁹⁸⁷ (Scheme 50). Additional leads to 1,2,3-triazines are given in References

SCHEME 47

8. **Synthetic applications of diazonium** ions **283**

37, 138, 366, **403,** 527-530, 640, 687, 811-813, 901, 1087, 1352 and 1353 and for 1,2,4-triazenes see References 364, 842, **905,** 966, 977, 1284, 1286 and 1307. The reaction of a diazonium ion with a cyanamide (Scheme **51)987** or I-aryl diazo aryl offer a route to 1,2,4-benzotetrazines. Syntheses with aryl diazo aryl diazonium ions open routes to 1,2,3,4-benzotetrazines^{97, 533, 1320, 1322, 1324.} hydrazines42, *03,* **46, 73, 77, 268, 363, 436, 437, 779-781, 863-866, 1350, 1370. 1380** (Schemes 51 and 52)

 (262)

The products of the type **244** are not aromatic in the heterocyclic system and undergo coupling with β -naphthol and other phenols. As such they can serve as disguised and stabilized diazonium intcrmediates. In general, substituents on **N-3** render the triazine ring systems susceptible to use as diazonium ions at elevated temperatures. **A** number of syntheses depend upon the triazines and triazine N-oxides as intermediatcs.

e. Groirp 5: Bctizothindiazolcs. When p-aniino thiophennls arc subjected to diazotization conditions they couple through sulphur to furnish the $4.4'-b$ is-(aminophenyl)disulphides¹³⁸⁰. When the *ortho* isomers are subjected to the same

conditions, cyclization to furnish benzothiadiazoles results^{445, 566, 593, 605, 606}. These compounds participate in some processes charactcristic of diazonium ions. Hence the nitrogen can be replaced by treatment with carbon disulphide⁵⁹³ and as such offers a route to aryl 1,2-dithiols.

2. Subclass b: The diazo nitrogen is lost

a. *Group 6: The Pschorr type* synthesis and its heterocyclic *annlogrtes.* Although Saunders 1^{103} breaks down the various processes into subgroups, many of the reactions in the group are formally identical to the Pschorr Phenanthrene synthesis²⁸⁸. The reaction proceeds well with a variety of groups X, e.g. reactions in the group are formally identical to the Pschorr Phenanthrene synthesis²⁸⁸. The reaction proceeds well with a variety of groups X, e.g. $CH=CH_1^{1007-1023}$, $-CH_2-CH_2- 1081$, $-S- 2683$, $-O- 289$, $-NR- 5393$, $-CH_2$ ³⁸¹, $-Se-$ ¹⁰¹⁹, $-SO_2$ ²⁸⁹, $-SO_3$ ¹¹⁰⁵³. The processes are probably mechanistically very similar. **A** possible exception is the bis-diazotizaton of 2,2'-diaminobiphenyls¹⁰⁰⁷, ¹⁰¹⁷, ¹²⁷⁷.

The nature of the intermediate had been the subject of somc speculation but quite probably involves a bis(benzo)triazapine which readily loses nitrogen⁴¹². Saunders¹¹⁰³ had proposed that the reaction involved the tetra-azotized system losing hydrazoic acid from a partially reduced system but this seems unlikely on thermodynamic and electronic grounds. Chauncy and Gchlcrt have successfully replaced the copper catalyst in the Pschorr synthesis with NaI²⁴³.

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Although the Pschorr syntheses formally resemble the Gomberg-Bachmann-Hey syntheses, they differ in terms of experimental conditions. The former are normally carried out in acidic media whereas the latter are invariably performed in basic media. The Pschorr processes are homogeneous because the attacking species is part of the substrate whereas the Gomberg-Bachniann-Hey reactions are heterogeneous. Thus the Pschorr syntheses possess the inherent advantage of decreased entropy requirement.

SCHEhlE *55*

When a rigid bridge is incorporated, the configuration must be *cis* for closure to occur. DeTar has suggested that the ethylenic bridge is perhaps the best bridge avaiiable as a consequence of its rigidityz8"; with *Irons* bridging other processes $occur¹⁰⁸²$. He also noted that electronic factors seem to be of secondary importance²⁸⁸. With the presence of ϱ -hydroxy and ϱ -hydroxy substituents, difficulties occur which lead to the formation of diazooxides. The presence of ether or amine functions in the *ortho* position can lead to extensive tar formation¹²¹⁷. When substituents are present at the **3'-** and 5'-positions, two possible modes of attack operate unless one of the **2'-** (6')-positions is blocked. Since the processes exhibit only a small degree of sensitivity to electronic effects, one will invariably obtain mixtures of products when the possibility exists. However, when appreciably different $\Delta E \pi$ values are to be expected, the PMO analyses will correctly predict the product to be generated.

E. Class E: Derivatives Formed **by** *Replacement of the Diazo Group*

With the possible exception of the coupling of diazonium ions with aryl amines and phenols, Class E represents the most widely known class of aryl diazonium reactions and is very sketchily presented in nearly every basic text on organic chemistry. The emphasis is normally placed upon replacement by hydrogen and the four halogens and is usually limited to the Mai-Kornblum reduction, the Sandmeyer reactions and the Balz-Schiernann reaction. Many other replacement processes are known which are of great synthetic utility and have been the subjects of chapters in *Orgatric Reactions* and a variety of reviews. Copper and its salts are encountered with a large number of the replaceinent reactions described in this section. Each valence state has associated with it a name, $Cu(0)$ Gattermann, $Cu(1)$ Sandmeyer and CU(II) Korner-Contardi. It seems quite likely that in this case a single valence state is actually involved. The Merck Index groups all of the processes under thc name Sandmeyer reactions¹¹⁶⁸.

1. Group I: Replacement of the diazo group by hydrogen

Classically, diazonium ions were reduced with ethanol and Griess¹⁵³ made the original observation in 1564 \vhen **he** obtained benzene, dinitrophenol and acetaldehyde from the reaction of ethanol with benzenediazonium nitrate.

The processes are rclatively sensitive to conditions and reduction is favoured by the presence of water, use of the diazotate rather than diazonium ion and the presence of electron-withdrawing substituents in the *ortho* position such as nitro or halogens or hydroxy. Hodgson⁵⁴⁹ had suggested that the greater the positive character of the diazonium carbon, the greater the ease of replacement by hydrogen. The validity of such a statcment might appear qucstionable in the light of current theory since the substituents included above encompass $+E$, $-E$ and $-I$ but all are capable of stabilizing homolytic processes. The reaction is often accompanied by thc formation of appreciable quantities of thc aryl alkyl ether which is in direct parallel with the formation of phenols from watcr. When diazonium halides are employed, aryl halides can become important products. This problem was overcome by Chamberlain²³⁸ who employed alkaline conditions which generated diazohydroxides or diazotatcs which furnished no ethers. Unfortunately, even this improvement frequently leads to only moderate yields of the hydrocarbon. Parsons and Bailor⁹⁷³ found that copper bronze improved the reduction of 4-methyl**azobenzene-4'-diazoniurn** sulpliatc and other workers have successfully employed Cu(0) prepared in a variety of ways to facilitate reduction by alcohols^{545, 552, 1206}.

Under the best of conditions, reduction with alcohols is not a good reliable method and this led to a search for alternative reducing agents. Friedlander³⁹⁸, Bamberger and Meimberg¹⁵, Eibner³²⁵, and Hantzsch and Vock⁵⁰⁴ investigated the use of sodium stannite in aqueous media and found it effective. The *E* diazotates do not reduce with sodium stannites0'. **As** noted previously, acid reduction with tin furnishes the hydrazine. Surprisingly, diazotized ethylamino gallate is reduced by stannous chloride to ethyl gallate¹⁰⁰⁴. Two rarely used reduction techniques which may actually have promise are alkaline sodium pyrosulphite⁴⁴⁴ and hydroquinone⁹⁵⁶. Brewster¹⁵³ avoided the ether formation process and obtained good yields in the naphthalene series. Other methods include the use of aluminium^{597, 816, 919} in ethanol solution and cuprous oxide in alcohol or formaldehyde solutions^{548, 550,} 740, 1079

The most widely accepted reduction technique, the use of hypophosphorous acid, is due to Mai⁷¹⁶ and has been rated by Adams and Kornblum³ as the preferred method. Yields range as high as *807<* whcreas other methods rarely reach 60% and are dependent to a large extent upon the ratio of diazonium salt to hypophosphorous acid. Dificult reductions require as much as a 15-fold excess of reducing agent even though the apparent proccss involves oxidation of 1 mole of reductant to phosphorous acid. By accepted reduction technique, the use of hypophosphorous acid,
and has been rated by Adams and Kornblum³ as the preferred
range as high as 80% whereas other methods rarely reach 60%
rant to a large extent upon

$$
ArN_2^+ + H_3PO_2 + H_2O \longrightarrow \longrightarrow ArH + H_3PO_3 + H^+ + N_2
$$
 (5)

Cadogan²¹⁷ has successfully replaced the amino function on a number of anilines in a one-pot process involving treatment of the amine with excess pentylnitrite in refluxing THF or dioxan.

SCHEME 56

2. Group 2: Replacement of the diazo group by halogens including astatine

The replacement of the diazonium group by a halogen has been known since 1860⁴¹⁷ when Griess heated diazoaminobenzoic acid with concentrated hydrogen halides to obtain halobenzoic acids. Griess⁴⁵⁴ later extended his studies to prepare

$$
HO_2C-ArN_2 \xrightarrow{HX} HO_2CArX + N_2 + R_2N-ArCO_2H + R_2NH_2X
$$
 (6)

representatives of all of the halogens by adding diazoniuni sulphates to concentrated solutions of the hydrogen halide using *meta*-carboxybenzencdiazonium sulphate; he found that the yield was improved by increasing the halogen concentration 47 . This point was pursued further by others who added halide salts to furnish increased halide concentrations^{114, 559}. Cupric chloride was the additive of choice for preparing p-chloronitrobenzcnc. Although the rcactions are normally run in concentrated acidic solutions, Griess^{454, 471} did observe that thermal decomposition of solid salts such as the chloroplatinate and hexachlorostannate furnishcd the chlorides. The most common solid salt decomposition is due to Schiemann and Balz^{10, 331, 1106, 1109, 1190 which, with the improvements of Schwecten¹¹³¹, provides a} moderate scale route to aryl fluorides. The most widely known halogen and pscudohalogen replacement processes were developed by Sandmeyer¹⁰⁹¹⁻¹¹⁰⁰. Sandmeyer had been attempting to prepare phenylacetylene from copper acetylide and benzenediazoniuni chloridc and obtained some chlorobcnzene. **He** was able to show this had arisen from reaction with copper(1) chloride. Subsequently, Gattermann^{416, 419} found that freshly prepared copper powder $(CuSO_4+Zn)$ brought about the formation of chlorobenzene from benzenediazonium chloride. Ullmann¹²¹⁶ found that commercial copper bronze could also be cmployed.

The use of copper salts is not nceded for preparing aryl iodides and can be accomplished by adding an alkali iodide to a diazoniuni sulphate or by using **HI** *270,* **454, 466,** ,467, **581, 626, 763, 1110, 1212,** ¹²⁹¹

a. *Replacement by fluorine*. Although the generally preferred method for introducing fluorine is the Balz-Schiemann reaction and normally good to excellent yields are obtained, it is not free from difficulties¹¹⁹⁰. Some of these difficulties are no! surprising, **c.g.** inadvcrtent rcplacemcnt of functional groups during the diazotization step. A common problem when solubilizing substituents such as OH are present is difficulty in isolating the fluoroboratc salt. To overcome this, other pcrfluoro inorganic acids have becn examined and the hexafluorophosphates have shown some promise because they are extremely insoluble and $PF₅$ is a very weak acid¹⁰⁹³, ¹⁰³⁴. Thus the isolation of p-hydroxybenzenediazonium fluoroborate is not realized while the fluorophosphate is obtained in over 75% yield. Unfortunately, the thermal decomposition to furnish the fluoride is seldom better than the **Balz-**Schiemann approach. Attempts to improve the decomposition of diazonium fluoroborates have not becn overly successful. There is at least one report of tetramethylurea serving as the solvent of choice for decomposing diazonium hexafluorophosphates¹⁰⁸⁴.

Additives have been employed for **a** variety of reasons. Deconipositions of nitro-aryldiazoniuni fluoroborates arc often difficult to control and neutral diluents such as sand or even sodium chloride are often employed^{720, 1063, 1106.} Clearly thc use of salts with nucleophilic anions is **a** rishy approach since much of the dccomposition occurs via heterolytic mechanisms. The use of alkali fluorides

can overcome the problem of additional nucleophiles but tends simultaneously to change the mechanism from heterolytic to homolytic by interchange of anions⁹⁸.

8. Synthetic applications of diagrams in a
\nthe problem of additional nucleophiles but tends simultaneously to
\nmechanism from heterolytic to homolytic by interchange of anions⁹⁸.
\n
$$
PhN_2BF_4 + MF \xrightarrow{\text{max}} PhN_2F + MBF_4
$$
\n
$$
PhN_2F \xrightarrow{PrN_2F + MBF_4} \text{Products}
$$
\n
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\left.\begin{array}{c}\n\text{(7)} \\
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The use of transition metal catalysts has also been claimed to lead to homolytic processes. Copper(1) fluoride and chloride have been used with some success when there are no highly polar substituents such as NO₂, OH, CO₂H. These are some of the same substituents which lead to difficulties with uncatalysed reactions^{110, 111}. Apparently the use of copper(1i) fluoroborate has not been examined even though it should be free from many of the objectionable characteristics of other systems.

The **use** of various solvents has been examined and the results arc frequcntly consistent with **a** heterolytic process747-7so~ **878, 847,** e.g. halobenzenes, nitrobenzene, benzonitrile, diphenyl ether, trifluoromethyl benzene, acetophenone and methyl benzoate. When benzenediazonium fluoroborate was decomposed in pyridine solutions, the formation of *2-* and 4-phenylpyridines were the dominant arylation processes observed. These are most consistent with a homolytic process. In those cases where homolytic processes appear to be occurring, the yields of fluorides are always low.

When the Balz-Schiemann reaction is applied to diazonium ions to furnish fluorides which are highly susceptible to nucleophilic attack, problems can arise in workup. If a nucleophilic centre is already present, as in many nitrogen heterocycles, the reaction can fail totally^{98, 1064, 1065}. On occasion Pschorr-type cyclizations cause considerable difficulties³⁸⁶.

Danek, Snoble and Nouzova280 succeeded in preparing the *0-, m-* and p-fluorophenols by diazotizing the aminophenols in 40% fluoroboric acid at -10 ^oC. With the *0-* and p-diazonium salts the fluoroborates could be isolated in good yield and decomposed t hermally as the dry salts, whereas the m-diazonium ion was destroyed in solution (Scheme 57).

b. *Replacement of thc diazo group by chlorine and bromine.* The most frequently employed method used to replace a diazo group by chlorine or bromine is due to Sandmeyer and involves the diazotization of the related anilinium halide and addition to the cuprous halide. There appears to be no general trend in conditions required and it is therefore necessary to optimize temperature and concentration for the process under consideration²¹. The best results are obtained when the diazonium ion solution is run into the cuprous halide solution. This tends to minimize formation of phenol and hydrocarbon coupling reactions. The diazonium ion appears initially to form **a** complex with the copper salt which then loses nitrogen. The reaction with cuprous chloride almost invariably gives excellent results when the conditions have been optimized. **A largs** number of other metal chlorides have been investigated as

alternative sources of halide ion and catalytic activity^{554, 563} but copper appears to occupy a unique position in diazonium chemistry. Waters¹²⁶⁹ has ascribed this to the oxidation potential of copper (i) .

With bromine, the Sandmeyer processes are complemented by the use of a complex double salt such as (ArN_2) ₂HgBr₄^{884, 885, 1131}, which is thermally decomposed, by hydrogen bromide decomposition of the diazonium bromide with or without copper or its salts being present, and by the decomposition of the diazonium perbromides. Various copper (i) complexes and salts have been used when troubles have been encountered2GG- *2i1.* **272.**

Schwechten's¹¹³¹ double salt pyrolysis is an extension of Griess's earlier work and has been used to some advantage by Newman^{884, 885} who recommends employing twice as much mercury salt as employed by Schwechten.

Griess used the perbromides^{454, 464} to prepare the aryl bromides. The diazonium perbromides were used with sodium carbonate and the products were distilled from the mixtures. In some cases no diluent is needed¹⁷⁹. Newman⁸⁸⁵ has diazotized the anilinium bromide in hydrobromic acid solution by adding nitrogen trioxide until the solution remained dark coloured, and then heated until nitrogen was no longer evolved.

C. *Repplacemetit of the diazo group by iodine mid astatitie.* GriessJSr, **,06, 407** discovered that replacement of the diazonium group by iodine was extremely facile either in hydroiodic acid or upon addition of an alkali iodide. Waters^{1262, 1269} ascribes the ease of the reaction to the fact that the redox potential is near that of copper(1) and that the process involves generation of I_3^+ . Normally the reaction is mild although 2-diazodiphenyl has proceeded explosively⁵⁸¹.

The element astatine can be introduced into an aromatic nucleus via the diazonium astinide. Thus, at $-5^{\circ}C$, o-halobenzenediazonium ions form complexes with astinide ion which are subsequently decomposed at 80 *"C* to furnish, as the major radioactive product, the corresponding 1-halo-2-astinobenzene⁷⁹⁴.

3. Group 3: Replacement of the diazo group by nitrogen-containing functions through formations of a new **C-N** bond

It is possible to replace the diazonium group by a number of groups which result in new $C-N$ bonds but the only ones of wide synthetic utility are the introduction of the $NO₂$ and $N₃$ functions. It is only of academic interest to convert an arylamine into itself by generating the diazoniuni ion and then allowing it to react with hydroxylamine to furnish a mixture of azide and aniline⁷⁴⁵. In some cases this can

be accomplished by reaction with ammonia, presumably via the diazoamines which undergo prototropy to furnish the aniline plus nitrogen^{474, 1255, 1256}. It appears that neither reaction has any promise as a route for introducing labelled nitrogen. **A** possibly more promising way of introducing labelled nitrogen is via the replacement of the diazonium group by NCO in the presence of copper salts^{417, 418}. Synthetically this offers no special advantage over treating the amine with phosgene if labelling is not desired.

a. *The formation of nitro compounds*. The replacement by NO₂ can be realized by several paths including the Sandmeyer reaction, and the Korner-Contardi reaction and the Griess replacement.

A number of copper derivativcs have been employed including Cu,O, finely divided copper powder^{89, 1166, 1233}, copper(II) sulphate in conjunction with cobalt nitrite complexes⁵⁵⁵, cupric chloride⁶⁹⁴⁻⁶⁹⁸ and cuprocupric sulphite⁵⁰², all in the presence of excess nitrite. In a number of cases the diazonium fluoroborate has been decomposed with a copper salt in the presence of nitrite salts^{777, 1167}.

Double mercury salts of the type $(ArN_2)_2Hg(NO_2)_2(NO_3)_2^{502}$ have been decomposed in thc presence of copper derivatives. In line with Newman's observations on the double salts with mercury halides^{884, 885}, one might suspect that the use of the diazonium nitrite with an additional equivalent of mercury salt would prove a more useful process.

In a number of cases, the generation of diazonium ions in the presence of large excesses of nitrite ion leads to formation of the nitro compound directly. This is most readily realized if the aryl group possesses electron-attracting substituents in the *ortho* and/or *para* positions. The substituents can be electron donating in the mesomeric sense, thus bromo and nitro substituents are both effective^{167, 168, 955,} **1088, 1207.**

b. *The formafie;! gf azides.* The **aryl** azides were known as azoimino compounds well into the 20th century and as such have appeared in the literature since 1864 when Griess^{119, 450} reported that benzenediazonium perbromide reacted with ammonia *to* generate phenylazoimine. (296)

17.5% (297)

17.5% (297)

SCHEME 59

of azides. The aryl azides were known as azoimino compounds

century and as such have appeared in the literature since 1864

reported that benzenediazonium perbromide reacted wi

$$
PhN2+ Br3- + 4 NH3 \xrightarrow{---} PhN3 + 3 NH4 + 3Br-
$$
 (8)

The same compound results when benzenediazonium sulphate is treated with hydroxylamine^{374, 745}, by diazotization of phenylhydrazine^{576, 732} or reaction of hydrazine with benzenediazonium sulphate. I PhN; HSO,+NH,OH - PhN,+H,O+H,SO, D. S. Wulfman

pund results when benzenediazonium

⁷⁴⁵, by diazotization of phenylhydra

nzenediazonium sulphate.
 N_2^+ HSO₄+NH₂OH - PhN₃+H₂O-

PhNHNH₂+HONO - PhN₃+2 H₂O

PhNHNH₂+HONO - PhN₃+2 H₂O

$$
PhN2+ HSO4+NH2OH
$$

\n
$$
PhN+H2O+H2SO4
$$

\n
$$
PhN+H2CH2O
$$

\n
$$
PhN3+2H2O
$$

\n
$$
(9)
$$

A very general synthesis is known which proceeds in part via the unstable pentazole which loses nitrogen to furnish the azide. The synthesis was developed by Noelting and Michel¹⁰⁴ in 1893 but it was not until 1954 that evidence was obtained indicating the formation of an intermediate pentazole along with a diazonium azide^{254, 587, 1215 .}

The pentazoles are extremely unstable substances and their chemistry has been reviewed by Ugi^{1215a} (see Scheme 59 above).

4. Group 4: Replacement of the diazo group by oxygen. Formation of phenols, phenol ethers and phenol esters

One of the earliest reactions observed with diazonium ions was replacement by **-OR (R** = **H,** alkyl, acyl). The formation of alcohols is a common reaction of alkyl diazonium ions and is often an annoying side-reaction when aryl diazonium ions are formed.

The alkyl diazonium reaction was noted by Piria^{991, 992 in 1848 when he treated} asparagine and aspartic acid with nitrous acid and obtained malic acid. The formation of phenol by heating an aqueous solution of benzenediazonium sulphate was reported by Griess in 1866⁴⁶⁰. The reaction was employed by Fisher in 1878 to help elucidate the structure of the triphenylmethane dye, rosaniline, which has thrce amino groups which he replaced with hydroxyls³⁷⁵. The reaction is generally successful and works well with tetrazotized biphenyls. Substituents do affect the with which the reaction oc~urs171. **223, 225,** 526, **641, 806, 907,** 1329, **1335, 1348, 1365.** Care must be taken to avoid an excess of nitrous acid or nitrate ion since under the acidic conditions of the process, the resulting phenols are often prone to nitrosation or nitration⁹⁰², ¹²³⁰.

The forniation of ethers from alcohols and diazonium ions **is in** direct parallel with the reaction with water. The reaction is frequently accompanied by reduction to the hydrocarbon. The **two** reactions compete effectively and often prevent one from obtaining a good yield of the desired product unless care is taken to employ the anhydrous diazonium salt. (There is however some divergence of opinion on this matter. Thus in at least one case, **15%** water was found permissible1053.) When this is done, the ethers are often obtained in excellent yields. The reactions are usually performed in refluxing alcohol^{793, 950, 970, 1051-1053, 1141} but operating in a pressure vessel to permit an increase in temperature has been efficacious in several instances⁷⁹³, ⁹⁵⁰, ⁹⁷⁰, ¹⁰⁵⁰, ¹⁰⁵², ¹⁰⁵³, ¹⁰⁴¹. The reaction does not work well with phenols where yields of less than 20% are common and they are accompanied by large quantities of arylated phenols^{242, 571}.

5. Group 5: Replacement of the diazo group with formation of a C-C bond

a. *Formation of aryl cyanides.* In Section **II.B.3** we have discussed the formation of diazocyanides and their rcactions. Sandmeyerdo2- **Iog3** demonstrated that treatment of a diazonium salt with KCN-CuCN furnished aryl cyanides. Subsequently a large number of aryl nitriles have been prepared in this fashion and the reaction

is now recognized as a simple variant of the Sandmeycr reaction with **halides^{8, 691}, ⁹⁸⁹, ¹⁰⁷⁵, ¹⁰⁸⁰, ¹⁰⁹⁴, ¹¹²⁵, ¹²¹⁸, ¹²¹⁸, ¹³⁵¹, ¹³⁶². The yields are often only moderate** because the initially formed 'labile' diazocyanidcs rapidly isomerize to the 'stable' isorners which do not decompose to furnish the nitrilc. Unlike the Sandmcycr reaction with halogens, other metal complexes can be used, in particular tetracyanatonickel(1i). **A** claim has been made that its use is to be prefcrred over biscyanato copper(1)^{402, 691}. Potassium tetracyanatoammine copper(II) has also been employed to avoid the potential hazard of generating cyanogen¹³⁶⁹.

b. *Formation of sytnmetricol und asyimietrical* binryls. Diazoniuni hydroxides decompose in aromatic solvents to furnish biaryls. The rcaction, known **as** the Gomberg-Bachmann reaction³², proceeds by a free radical mechanism and is strongly paralleled by the related Grieve-Hey-Heil bron syntheses using aromatic nitrosamides which are decomposed in the substrate aromatic solvent^{32, 281, 328, 438,} **539, l103, 1326.** Both provide a route to unsyinnictrical biaryls (Schcrne 60) and are

mechanistically similar. The reactions have been reviewed. These proccsses offer one of the best routes to unsymmetrically substituted biaryls and furnish ready access to meta-substituted biphenyl when substituents are to be in only one ring; the related substituted aniline is employed along with the aromatic hydrocarbon. Unlike electrophilic substitution, thc products from substratcs such as nitrobenzene are *para* substituted. This is not particularly surprising if one remembers that the corresponding Wheland intermediates **(298, 299)** will be more highly delocalized with *para* or *ortho* attack than with *mefo* **(300).** With substrates such as pyridine all possible isomers result. The related unimolccular process involving the formation ot' phenanthrene and fluorcnes is known as a Pschorr synthesis and has been treated in Section D.2.b. The Gomberg-Bachmann reaction furnishes yields in thc poor to good range $(60 + \frac{9}{6})$ but infrequently higher). The Pschorr reaction (actually discovered previously by Graebe and Ullmann^{441, 412} and Fischer and Schmidt³⁸¹) being an intramolecular process can be expected to furnish good yields, 70-90% not being unusual. It should not surprise anyone to find that isomer mixtures result when the non-diazonium ring system is asymmetrically substituted. Obviously the presence of an *ot.flio* ethylene bridge causes the rnolecules *to* undergo some or even dominant closure *to* cinnolines and related heterocycles. Thus all three of these related reactions are subject to the same limitations as are other ρ -substituted diazonium ions.

A fundamental limitation of both the Gomberg-Bachmann and the Grieve-Hey-Hcilbron syntheses is the need to employ the substrate hydrocarbon in the liquid state. When this is combined with the desirability of having the substrate molecule as the most symmetrical portion of the biaryl to be synthesized, very strong limitations in scope arise. Since the reactions are normally run below **30** "C, many formal processes arc just not practicable in the laboratory. The use of solvents is not particularly encouraging since the aryl radicals generated can attack the solvent as well. Thus when chloroform or carbon tetrachloride are used, appreciable quantities of the aryl diazonium compound end up as aryl chlorides¹⁷⁴. With substrates such as pyridine, the dry diazonium salt can be added to the pyridine⁸⁰⁷ or by the reaction of the sodium diazotate with pyridine-acetic anhydride³⁸⁹. Alternatively the nitrosoacetyl derivative can be heated in pyridine solution^{517, 518}.

Cadogan²¹⁵ has smoothly converted aryl diazonium fluoborates to dihalodiazobenzenes by deconiposing the diazonium salt in **10%** acetone 90% benzene at room temperature using copper powder. If **2** mol of water are present, the biphenyls result (Scheme 58).

C. *Ary fated qiiitiancs atid other iinsaturoted species incliiditig Iietcro aroniatics atid non-classical aromatic systems.* The *Afcetwcitr reactions.* **It** is possible to arylate quinones, α , β -unsaturated carbonyl compounds, vinyl nitriles and styrene using aryl diazonium ions. In the case of quinones and quinone oximes they are frequently dissolved in acetic acid or alcohol and the diazonium salt is added at room temperature. The evolution of nitrogen is normally complete in 24 h^{22, 137, 139, 707.} **759, loil.** 1103. When olefins are uscd as substrates, copper salts are normally employed as catalysts^{767, 1330}. This process has become known as the Meerwein reaction¹⁰⁷¹. With many substrates two possible products can arise in principle, but a simple analysis on the basis of which intermediate radical will be most stable usually suffices to predict the correct product.

A large variety of olefinic substrates can be used and these include, vinyl halides269, **079, 944,** acrylonitriles163, *293,* **,195,** *670-68?.* **717, 751, 752, 1332,** phcnylacetylenes766, phenylpropiolic acid⁷⁶⁷, acetylene^{#44}, coumarin^{394, 767, 1332}, cinnamaldehydes⁷⁶⁷, **211. 210. 210. 210. 210. 21. 103.** *Pg. 103. Pg. 103. Cinnamyl esters⁶⁸⁴</sub>, ⁷⁶⁷, acrylic acids^{702, 644}, ¹⁰³⁶, butadienes¹⁵², ²⁶⁵, ⁴⁷⁶, ⁸⁴⁴, ¹⁰⁷², ¹⁰⁷³, 219. 219. 219. 219. 21* fumaric eSters767, lloc, **1235,** maleimideS'O6S. **1070** furanS1GO. 395. **396.** 717, **946** and nitrostyrenes¹⁰⁹.

Most of these reactions are straightforward but in some cases addition of aryl *767,* **844, 847, 10~8. 1070. 1072, 1073. 1196. 1235.** This is in part influenced by the reaction conditions. Cinnamic acids and maleic acids are often accompanied by decarboxylation with the aryl group formally replacing the $-CO₂H$ ¹⁰⁶⁻¹⁰⁸, 286, 407, 622, 716, 767, 771, *772,* **1035.** This is somewhat dependent upon reaction conditions, with very low pH apparently favouring retention of the carboxyl function⁶²². halogen across the double bond occurs¹⁵², ²⁶⁵, ²⁶⁹, ²⁹³, 425, 476, 679, 680, 682, 712, 751, 752,

The variety of less common unsaturated substrates also undergo the Meerwein reaction. These include oximes^{94, 95, 140, 622, 623, 980, malonic ester⁴⁸⁰, nitromethane¹²¹⁴,} ferrocene^{155, 880-882, 1275 and γ -pyridines¹⁰³¹.}

There seem to be few limitations upon the aryl diazonium compound used except that at very low pH values some lose nitrogen to form aryl halides which is facilitated by the copper catalysts normally used. At higher pH resin formation can create problems.

6. Group 6: Replacement by groups in which attachment occurs through sulphur, selenium or tellurium

Many of the aryl diazo-sulphur derivatives mentioned previously are of extremely limited stability and lose nitrogen explosively to furnish aryl-S derivatives. Griess^{440, 461} prepared diphenyl sulphide by reacting benzenediazonium salt with solutions of potassium sulphide. The same product occurred when the silver or gold salts obtained from the diazonium hydroxide were treated with hydrogen sulphide. Even in extremely small runs of these reactions, explosions occurred. The use of cupric sulphide in analogy with a Sandmeyer-type process fails and furnishes only $tars¹³²$. The reaction with elemental sulphur furnishes phenyl sulphide and, in a similar fashion, reaction with selenium furnishes the selenide¹²⁷⁰. With tellurium one obtains the diaryl dihalotelluride¹²⁷⁰.

When the hydrogen sulphide is replaced by alkyl mercaptans, the resulting aryl diazonium alkyl sulphides can be readily (often explosively) decomposed into the aryl alkyl sulphide⁷¹⁰, ¹¹⁶⁴.

Clearly this approach does not offer much attraction to cautious chemists and approaches employing the sulphur atom in modified form are more desirable.

The safe generation of thiophenols and of aryl thioethers from diazonium salts can be realized by the Leukart thiophenol reaction where the diazonium salt is allowed to react with a potassium xanthate. If the ether is desired it is pyrolysed; if the thiophenol is wanted it is subjected to alkaline hydrolysis⁷²⁶.

This process along with the somewhat similar approaches of Lustig⁷⁴⁰ and Clifford and Lichty²⁵² have fairly general applicability²⁵², ⁵⁷⁴, ⁵⁹⁴, ¹²⁷¹, ¹³⁴⁰.

When aryl diazonium salts are added to solution of calcium or sodium polysulphides, the bis(ary1) disulphides are generated and these can be subsequently converted to the thiophenols¹²⁷⁰, ¹³⁵¹, ¹³³⁸.

The reaction of aryl diazonium ions with thioureas generates aryl thioronium salts which hydrolyse to furnish the thiophenols²⁰³. When the thiourea is unsubstituted, troubles can arise. Thus a diazonium anthrone furnishes the thiol¹³³⁸ while thiophenol was not isolated from the reaction with benzene diazonium ion²⁰³. Forrester and Wardell list only the Leukart synthesis for thiophenols³⁸⁸

amongst the various syntheses given. The sulphides can also be prepared from the reaction of a dialkyl sulphide with benzyne and diaryl sulphides with differing substituents can be obtained by copper-catalysed decomposition of the aryl diazonium aryl sulphides^{540, 690}.

$$
ArN_2 + \bigvee_{\begin{subarray}{c} \parallel \\ S \end{subarray}}^{R_2N} \bigvee_{\begin{subarray}{c} \subseteq \\ S \end{subarray}}^{NR_2} \longrightarrow Ar - \overline{S} = C \bigvee_{NR_2}^{NR_2} \xrightarrow{H, O} ArSH
$$
 (12)

Aryl diazonium ions react with thiocyanate and selenocyanate ions to furnish the related aryl thiocyanate or aryl selenocyanate^{124, 355, 415, $421-423$, 888 , 846 , Subsequent} hydrolysis furnishes the corresponding phenol analogues. **A** large number of metal and metal salt catalysts wcre examined in the thiocyanate case with ferric chloride being the superior catalyst for conversion of o -nitrobenzenediazonium chloride at 15-20 *"C (76%)* and tungsten hexachloride being the best catalyst at 60-70 *"C* $(69\%)⁸⁹⁰$. With *meta*-nitrobenzenediazonium chloride, ferric ion was found to be the best catalyst (80%) ⁶⁸⁹.

The aryl selenocyanides are convenient intermediates in aryl selenium chemistry⁵⁹⁸.

7. Group 7: Replacement of the diazo group by arsenic, antimony and bismuth

In a footnote to a paper, Koenigs⁶⁸⁶ noted in 1890 that the benzenediazonium ion gave **a** cherry-red colour with arsenite. This is a characteristic phenomenon observed when aryl arsonic acids are prepared by the Bart reaction which was described in 1910 in the German, English and French patent literature1331- **1342. 1356.**

Prior to that time arsenite had been employed as a reducing agent in strongly alkaline solution. The Bart reaction is of general utility^{482, 963} furnishing the aryl 1357-1359, 1366. It has been used to prepare heterocyclic³⁸, ¹²⁵, ²³⁰, ⁹⁷⁹, ¹¹⁵⁵, ¹³⁶⁶ and polycyclic¹⁵¹, ¹⁰⁸⁸, ¹¹⁰², ¹¹⁷⁴ arsonic acids. The reaction is strongly dependent upon pH and is catalyscd by silver-, copper- and nickel-based catalysts. **A** modification, known as tlic Scheller reaction, proceeds through the copper-catalysed decomposition of the aryl diazonium tetrachloroarsenite which furnishes an intermediate which decomposes in hot water to furnish the aryl arsonic acid¹²⁶, ¹³⁶⁴, ¹³⁶⁷. time arsenite had been employed as a reducing
 A. The Bart reaction is of general utility^{482, 963} f

fair to excellent yields<sup>87, 88, 91, 387, 604, 619, 624, 729,

has been used to prepare heterocyclic^{38, 125, 2}

¹</sup> arsenic acids in fair to excellent yields", *88.* **91, 387. 801, 619. 625.** 729. **'383, 809. BJ8. 1111. 1263.**

$$
ArN2OH + Na2HAsO3 - \longrightarrow ArAsO3Na2+N2+H2O
$$

Elemental arsenic^{1265, 1267} and elemental antimony¹²⁶⁷ are attacked by aryl diazonium salts **when** warmed in acetone and furnish the triaryl antimony dichlorides or triarylarsenic dichlorides.

ArN,CI+Sb ~ + **Ar,SbCI,**

The aryl stibonic acids arc formed by trcating a strongly alkaline solution of the diazo compound with an alkali antimonate¹¹¹², ^{1333, 1360, 1361. A modification employs} the aryl diazonium chloride-zinc chloride double salt (e.g. May's double salt). This i5 decolnposcd in thc prescnce of copper of copper(1) salts1112. **1339, 1341** or zinc dust⁸⁷⁸. Aryl bismuth compounds arc generated by decomposing the diazoniumbismutli chloi.ide tioublc salt in **ii** solvcnt such as acetone. Mono-, di- and triaryl bismuths are formed in moderate yields^{431, 432}.

8. Group 8: Replacement of the diazo group by metals

The aryl diazonium ions form a large number of double salts with a variety of heavy metals. The decompositions of some of these offer routes to aryl metal systems where a metal-carbon bond results. There is apparently only one example involving a transition metal (Pt) ¹⁰²⁴. heavy metals. The decompositions of some of these offer routes to aryl metal system
where a metal-carbon bond results. There is apparently only one example involvin
a transition metal (Pt) ¹⁰²⁴.
PhN₂BF₄ + PtHCI[P(C

$$
PhN_{2}BF_{4} + PHCI[P(C_{2}H_{3})_{3}]_{2} \longrightarrow PhN_{2}PLCI[P(C_{2}H_{3})_{3}]_{2} \longrightarrow H \times PH
$$
\n(306)\n(307)\n(308)

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This fact is rather surprising since a large number of aryl diazonium metal complexes are known¹¹⁹³ and many metals are capable of forming aryl-metal bonds. One might expect to form aryl-metal bonds in some of the cases where the decompositions of aryl diazonium ions are catalysed by transition metals and their derivatives. The absence of such products may be a consequence of oversight rather than thermodynamics.

With mercury, tin and lead it is possible to prepare the aryl derivative from aryl diazonium ions. Nesmeyanov and coworkers have prepared a number of arylmercurichlorides from the aryl diazonium chloride-mercuric chloride double salts in the presence of copper⁸⁶⁷⁻⁸⁷⁷. Alternatively, one can react the diazonium salt while strongly agitating the mercury cathode^{315, 744}. to form aryl-metal bonds in some of
tryl diazonium ions are catalysed by trans
ence of such products may be a consequer
cs.
a and lead it is possible to prepare the ary
essmeyanov and coworkers have prepare
pom the aryl d

$$
ArN_{2}Cl \cdot HgCl_{2} \xrightarrow{Cu} 2 CuCl + N_{2} + ArHgCl
$$

Stannic halides also form double salts with aryl diazonium ions and these, when reduced with zinc, copper or preferably tin in ethyl acetate solution, furnish principally the bis(aryl)stannyl dichlorides $677, 1266$.

$$
2 \text{ ArN}_2 \text{Cl} \cdot \text{SnCl}_4 + 4 \text{ Sn} \longrightarrow \text{Ar}_2 \text{SnCl}_2 + 2 \text{ N}_2 + 4 \text{ SnCl}_2
$$

In acetone solutions, tin reacts directly to furnish the bis(aryl)dichloro stannane¹²⁶⁶. Decompositions of lead double salts in the presence of copper furnish poor yield of triaryl chloro plumbane and biaryl dichloro plumbanes^{1266, 1268}.

When aryl diazonium fluoroborates are decomposed with copper powder, small amounts of the aryl copper(1) derivatives are formed and these can be stabilized by chelation with pyridine^{135, 1280}.

111. ALKYL DlAZONlUM IONS

The known reactions of alkyl diazonium ions unlike those of the arenediazonium ions are rather limited. **It** seems likely that many of the processes known for the arenes will ultimately become known when suitable reaction conditions are developed. Until fairly recently, alkyl diazonium ions have held the role of 'alleged intermediates' in deamination reactions. The basic features with numerous examples have been the **l105, 1152, 1157, 1278.** *1282.* **¹³²⁶**and we shall only conccrn ourselves here with the synthetic uses of the nitrosation of alkyl amines and with species derived from acyl nitrosamines. of several reviews83, **115, 158, 183,** *226, 258,* **477, 478,** *651, 715,* **758,** *825,* **836. 09G. 1010, 1056,**

One of the niore practical synthetic reactions of alkyl diazonium ions is the Tiffeneau-Demjanov reaction where an α -amino alcohol or an amine is treated with

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a nitrosating agent and rearrangement occurs with shortening or lengthening of the carbon chain (usually a ring is enlarged or contracted). The processes often proceed with excellent yields and in many instances provide routes which are competitive with the Wolff rearrangements and homologations with diazoalkanes, both of which are discussed in Chapter 18. An exhaustive review of both sets of processes has recently been presented by Gutsche and Redmore^{478, 1040}. The Demjanov reaction²⁸⁵ in a strict sense describes the rearrangement process of a nitrosated amine and can lead to a ring expansion or contraction.

 (315)

The Tiffeneau reaction (sometimes referred to as a semi-pinacolic rearrangement) involves the nitrosation of an α -amino alcohol^{701, 1205}. Since such compounds are readily attainable in good yield by a variety of routes from ketones, the process is of considerable synthetic utility. The reaction does not always lead to ring contraction77s and thc behaviour is a function of ring size2'5. *267,* **271, s??. 323. s~?.** 674, 743. 1194, 1209, 1210 and, as indicated in the hypothetical case illustrated, is a function of conformation.

The overall question of conformational effects on deamination in rigid systems has been summarized by Kirk and Hartshorne⁶⁴¹. The course of Tiffeneau-Demjanov reactions on a variety of aminocyclanols has been summarized according to ring size by Gutsche and Redmore¹⁰⁴⁰ and it is clear that conformational effects are of great importance. The difficulties encountered are often a direct consequence Gf the lack of conformational regularity encountered in seven or more membered rings. In the case of hydrochrysenes *(e.g.* D-homosteroids) a careful analysis has been made¹²⁷⁸. The migrating group is always anti to the leaving diazonium nitrogen^{267, 674}.

With simple amines one encounters a variety of processes which are normally of little synthetic utility. The diazotization process normally furnishes mixtures of alcohols and olefins which, on occasion, prove to be rather complex as a consequence of Wagner-Meerwein rearrangements. Several reviews dealing with various aspects of the deamination via nitrosation have appeared.

Fairly recently, work originating primarily in the laboratories of Moss⁸²¹⁻⁸⁴¹, Newman⁸⁸⁹⁻⁸⁹⁵ and of Kirmse⁶⁴⁴⁻⁶⁵⁹ has led to the development of reasonable to excellent synthetic schemes dependent upon the transient existence of alkyl diazonium ions. Thus Steinheimer¹¹⁶⁹⁻¹¹⁷¹, upon discovering that diazo transfer to 7-aminonorcarane proceeded in only moderate yield, was able to convert the same amine via the 7-N-nitrosourea or urethane into the azide in excellent yield by employing Kirmse's reaction64G with lithium azide. The reaction was followed by careful thermolysis to the 1-azetidine and represents a general route to the formal $2+2$ cycloaddition of hydrogen cyanide to an olefin¹¹⁷⁰, ¹¹⁷¹. The process had been postulated by him as proceeding through a diazonium ion and a pentazole¹¹²¹. Simultaneous work in Kirmse's laboratory revealed this to be the case^{662, 663} and that it did not involve the highly improbable cyclopropyl carboniuni ion originally claimed⁶⁴⁶.

Kirnise644~~6s4~ **G5i-6G9** has subjected a variety of A'-nitroso urethanes to basic conditions under which there appears *10* be a high probability that alkyl diazonium ions and alkyl diazotate ions are formed^{641}. The reactions involve trapping these species prior to formatior. of a diazoalkane. It seems likely that similar behaviour is occurring in the synthetic studies of Newman⁶⁸⁹⁻⁸⁹⁵, Hogan⁵⁷, Billett¹²¹⁻¹²³ and Moss^{821-831, 833-811. Kirmse has employed several bases of which formate, methoxide,} carbonate and bicarbonate are dominant. Many of his reactions have been run in methanol but in some cases ethers have been employed. The solvent and base contribute markedly to the naturc and distribution of the final products. Thus the 6-cso-bicyclo[3. I .O]hex-2-ene furnishcs *90%* + benzene with sodium carbonate/ nietlianol and **437;** bcnzene, 23% **322** and 35% **323** whcn sodium forniate/methanol is employed⁶⁵³.

When the β -phenethyl system was treated with NaBH₁, the N₂ function was removed to furnish **61%** ethyl benzene and a mixture of styrene and **u-** and 3-phenethyl methyl ethers (7 : 9 : 84) ⁶⁴⁷. The reaction to furnish cyclopropyl azidc646, *6G2,* 663 has been studied using isotopic labelling and the reaction strongly resembles the related aromatic series with a mixed path resulting from pentazole and pentazene intermediates. This fact should not be particularly surprising since

SCHEME 65

cyclopropyl carbonium ions are virtually non-existent with only one other substantiated example being known $(334 \rightarrow 345)$ ⁹⁷⁸. Thus the claim of Jacquier of having observed the conversion of 7-exo- and 7-endo-norcaranylamines to the alcohols via nitrosation has been shown to be incorrect along with their subsequent conversion to a common cyclopropyl ketone^{544, 607-609, 669.} What does occur is conversion to cyclohept-2-enyl alcohol^{514, 669}. Steinheimer¹¹⁷¹ cycled the 7-exonorcaranylamine **(336)** through the azide and back to the amine and found only the exo-azide **(338)** was formed (t.1.c. at both stages, with known endo-amine for comparison). Kirmse obtained allylic systems in the absence of azide ion^{651, 657, 659.} When Kirmse examined the behaviour of the 7-exo- and 7-endo-norcaranylamines in the absence of azide ion, he obtained 97 and 91% of the *trans-7-cycloheptenyl* methyl ether (MeOH/Na₂CO₃) and 94 and 98% norcarane with sodium borohydride⁶⁵¹.

When I, ω -diamines are subjected to the basic Kirmse sequence (amide, nitrosamide-baseaso), the nature of the final products was found to be **a** function of the chain length. With I,3-diaminopropane, cyclopropyl methyl ether was obtained in 50% yield, but with larger chains the dimethoxy ethers and ω -methoxy- α -olefins occurred (with MeOH/MeONa, the diethers were the major products) 650 .

When propargylic and allylic amines are subjected to the Kirrnse series, rearrangements occur^{654, 658, 665-667. The propargylic systems offer routes to} P-substituted propionic acids, alkoxy allenes and propargylic ethers (Scheme 68).

With allylic systems, allyl ethers (mixtures of isomers) are obtained^{654, 658}. When cyclic substrates are employed (cyclopropyl and cyclopropyl carbinyl) rearrnngements occur in good yield. In two of the examples, the diazonium species was generated by the addition of **H?C)/-OH** photochemically and the presence of rearranged products not available via **a** carbene process can be taken as evidence that reaction did involve a carbonium ion process⁶⁶⁴.

Moss821-e31. 833-811 has examined the question of when does the action of base upon N-nitrosourethanes lead to alkyl diazotates and when do diazoalkanes result⁸²¹. In the first paper of the Moss series and in a review, one finds a good summary of references to earlier work in the area by a number of workers^{821, 836}.

The reaction of α -phenethyl diazotates at -50 °C in CH₂Cl₂ with Et₃OBF₄ furnished the alkyldiazo ether which collapses to the α -phenethyl ether with 70% retention of configuration^{826, 828}. Concurrent with O-alkylation, one obtains varying amounts of *N*-alkylation to furnish azoxyalkanes^{830, 833}, 834. The yields are somewhat better in HMPA solutions than in $CH₂Cl₂$. The reaction has been employed to synthesize L-dihydroelaromycin *(359).* L-Elaroniycin **(358)** has been reported to be an antibiotic and a carcinogen 834 .

SCHEME 69

When α -phenethyl diazotates are treated with hydrazine⁸³⁹, α -phenethyl hydrazine results with 54% net inversion. When ammonia is employed⁸³⁵, inversion occurs **along** with some racemization. Similarly treatment with Grignard reagents leads to alkylation with some inversion. Maximum inversion was observed at the lowest temperature employed, with phenyl Grignard and s-butyl diazotate in ether.

Ethyldiazotates react with acyl anhydrides and halides in ether solutions to furnish intermediate diazotate estcrs which collapse with loss of nitrogen leading to a stereoselective synthesis of esters, e.g. 17-21% inversion with acetic anhydride⁸³¹. The reaction procecds with little intermediacy of the carbonium ion for the cyclopropyl carbinyl system proceeds in \sim 80% yield to the cyclopropyl carbinyl ester.

Moss⁸⁴¹ has summarized a number of potential reactions of diazotate chemistry. These include the use of allenic diazotates to generate aryl diazomethanes^{899, 900}, photolysis of epoxy diazotates to furnish oxadiazotenes^{961, 962} and the modification of D-nor-steroids via diazotate-generated carbonium ions7'4. *7iG.*

Recently, Newman's group⁸⁸⁰⁻⁸⁹⁵ (Scheme 70) has expanded upon earlier work by Huisgen⁵⁸⁵ and Müller⁸⁹⁸ involving treatment of N-nitroso lactams (in one instance, oxazolidones) with bases to furnish several interesting systems $(360 \rightarrow 362)$ and $(363 \rightarrow 365)$. Related work by Hogan⁵⁷³ (see also Reference 886) is summarized in Scheme 66 $(360 \rightarrow 362)$. Hassner has presented a detailed study of N-nitroso oxazalidone chemistry⁵¹¹.

The use of acidic nitrosation to dearninate amines has received considerable attention in the past and a number of recent reviews have appeared. The reactions far too frequently lead to vcry complex mixtures (see, for example, References **184,** *255,* 773, 1236) but on occasion are relatively

At 0° C it has been possible to prepare β , β -disubstituted vinyl diazonium salts using β-halo-aldehyde-tosylhydrazones and SbCl₅ in CCl₄¹⁵⁰.

The behaviour of 3 (or 7) amino tropoloncs towards diazotization is closer to that of alkyl amines than it is to the anilines but diazonium tropolones exhibit reasonable stabilities at other ring positions and have been used in the Pschorr synthesis with mixed success. It is therefore surprising that when the diazonium ion is in the 3-position, collapse to σ -hydroxybenzoic acids occurs with great ease even though an intermediate oxadiazole should in principle exhibit considerable stabiIity9, *262,* **521. 672, 843. 8D7, 911,** *80,* 927, **929. 932,** 933. **030** by virtue of being a 1k electron system.

The oxadiazole ring system may of course be generated but under the acidic conditions reverts to the diazonium ion **(383)** which might collapse to the hydroxybenzoic acid.

IV. BENZYNES, HETARYNES AND OTHER DIDEHYDROAROMATICS

The generation and reactions of benzynes and hetarynes (Schemes 73-77) have been the subject of a monograph⁵⁷⁰ and a chapter in The *Chemistry of Acetylene*^{570a}. This type of species is often treated concurrently with carbenes and nitrenes in texts and *Annual Reports* of the Chemical Society, as a consequence of their high

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reactivity and chemistry and their exhibiting a valence state with carbon having fewer than four formal bonds (in one resonance form one has **a** bis-1,2-carbene). **A** number of syntheses involving benzynes start formally or actually with diazonium compounds. The recent innovations using the Pbl" and **I** -chlorobenzotriazole oxidation of N-aminobenzotriazoles (Schemes **73** and **77)** both formally and

SCHEME 73

practically involves the chemistry of $N=N-X$ systems. Workers at the Robert Robinson Laboratories at the University of Liverpool have carried out a prodigious amount of work in this area. Earlier work involving the destruction of o-carboxydiazonium salts and benzothiadiazole S,S-dioxides has also furnished entry into this important synthetic area. The arynes can insert into $X-Y$ bonds, act as hot dienophiles and dirnerize to furnish biphenylenes. The ability of arynes to participate in cycloaddition reactions is very well documented (Schemes **76** and **77)** and Hoffmann⁵⁷⁰ has presented a 25-page tabulation of products.

The arynes are not limited to 1,2-benzo-type systems. Photolysis of 3-carboxybenzenediazonium systems furnish the 1,3-arynes¹¹³. The oxidation of 1-aminonaphtho[1.8a.8.de]-1,2,3-triazine furnishes the 1,8-didehydroxnaphthalcne^{113, 624a, 776a.}

The gas phase and flash photolysis of benzcnediazoniuni 4-carboxylate furnishes the **1** ,4-didehydrobenzcne1". Recs and his colleagues **have** performcd extensive studies in this field as a conscquence of syntheses dircctcd towards preparing

 $\ddot{}$

.y,y-didchydroaromatic systems such as I ,8-didehydronaphthalene **(385), 2,2'** didehydrobiphenyl and related compounds (e.g. 4,5-didehydrophenanthrene, 4,5-didchydrofluorenonc¹¹²), as well as intermediates for preparing I-azabenzocyclobutenesG and benzynes. Tetrazotization of 2,2'-diarninobiphenyI **(415)** in 2 N-HCI furnishcd six identifiable products (Schcnie 78). Oxidation of **389** furnished **384** and related products but no **392.** Work with the rclatcd fluorenonc systems was unsuccessful. When the 2,2'-diaminobiphenyl was tetrazotized and treated with

SCHEMC 78

ammonia, the triazepine (428) was obtained in high yield. Similar results were obtained employing **I** ,8-diaminonaphthnlenc which furnishes **429.** When **428** was treated with **a** variety of reagents it **was** evident that the system was a masked diazonium ion in much the same fashion as is observed with open chain triazenes.

When the tetrazotized biphenyl was allowed to react with methyl amine, **430** resulted.

The aziminc **(418)** is **a** potential 1,3-dipole and is generated thermally from the triazipene. The carboxethoxy derivative (435, $R = CO₂Et$) was found to be a good 1,3-dipolar reagent²³⁵. The N-ethyl derivative (435, $R = C₂H₅$) behaves anomalously with *N*-alkylimides and acetylenic esters¹⁰³⁸. The triazine (439) available from tetrazotized 1,s-diaminonaphthalenc is unusual in that it behaves **as** a **1,l** I-dipole towards dimethyl **acetylenedicarboxylate1044.**

The iminoazimines **(441)** are readily prepared from **418** by reaction with the appropriate imidochlorides in acetonitrile solution in the presence of potassium carbonate. They undergo thermal rearrangement with the alkyl case proceeding slowly even at room temperature⁸⁶.

Although the formation of benzotriazene derivatives has been known for many years it was only in 1971 that Rees and Storrs reported the preparation of the parent compound. The syntheses grew out of their attempts (ultimately successfuls~ *6,* to prepare benzazetes. Since these syntheses involve cyclizations of azido diazoalkanes, they will be treated in Chapter 18. An alternative route **was** through oxidation of N-aminoindazoles.

When **4-phenyl-l,2,3-benzotriazine** was pyrolysed in the vapour phase at **420- 450 0C/IO-3** Torr, the benzazete **(445)** was formed along with several other products. The product was collected at -78 °C and on warming the dimer is formed. A number of possible dimeric structures are reasonable and the structure is currently unknown.

V. CYCLOALKY NES

The compound cyclononyne is known¹²⁹ and is quite reactive. Smaller ring systems would be expected to be even more reactive and to resemble the arynes. Hence one might expect to form on a transient hasis such intcrmediates by eliniination of **HX** from a 1-halocycloalkene by treatment with strong bases¹²⁹⁹. The eliminations of the element of $M-X$ from some metallated haloalkenes are extremely slow reactions as a consequence of the strain in a potential alkyne or the high energy of the didehydro species¹³⁰¹. The formation of strained cycloalkync can however be reached via 1,2-diazocycloalkanes and their subsequent decomposition and related
systems¹³⁰, ¹³¹, ¹⁶⁵, ²⁷⁶, ²⁷⁶, ²⁷⁸, ³¹⁰, ⁴⁸¹, ⁵⁸⁵, ¹⁰⁰⁵, ¹²⁹⁸, This problem will be treated in Chapter 18 under dimerization processes in which two new C-C bonds are formed.

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CHAPTER 0

**Photochemistry of the
diazonium and diazo groups**

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1. INTRODUCTION

Light can be transmitted, refracted, scattered or absorbed by a system. It is a fundamental tenet of photochemistry that only the light absorbed can be effective in producing chemical or physical changes in a molecule.

A. **Photochemicol** *Energy'-5*

The energy of light in kcal/mol is given by thc expression:

$$
E = \frac{2.86 \times 10^5}{\lambda (\text{\AA})} \text{ kcal/mol}
$$

An energy of 1 kcal/mol corresponds to radiation of wavelength 286,000 *8,* or

1 kcal/mol =
$$
\frac{10^8 \text{ Å/cm}}{286,000 \text{ Å}}
$$
 = 353 cm⁻¹

in the infrared portion of the spectrum. When electromagnetic radiation of these frequencies is absorbed, molecules are excited to higher vibrational states. Radiation of shorter wavelength (higher frequency) contains more energy. Visible light has a wavelength from 4200 Å (violet) to 7000 Å (red), i.e. it contains 68–41 kcal/mol. Ultraviolet light of 2000 *8,* corresponds to an energy of 143 kcal/mol.

6. Photochemical Excitation

Light in the ultraviolet-visible region has energy sufficient to excite molecules to higher electronic states. Consider now the potential energy curve for the ground state (E_0) and first excited state (E_1) . According to the Franck-Condon principle transitions occur between vibrational levels for which the nuclear configurations are the same $(r' = r'')$, i.e. we have vertical excitation as shown in Figure 1. The energy of the electronic transition is measured from v_0 of the ground state to v_0 of the excited state, $E_1(\nu_0) - E_0(\nu_0) = \Delta E = h\nu$.

The major events that occur following light absorption may be summarized in terms of a Jablonski diagram. The absorption of light raises the molecule from *So* to *Sp,* as shown in Figure **2.** In solution the excess vibrational energy of **S2** will be rapidly dissipated by radiationless processes. After conversion to $S₁$ the molecule may lose its energy by four important processes: (i) fluorescence, (ii) chemical reaction, (iii) return to **a** highly vibrationally excited ground state *(S,-So)* and (iv) intersystem crossing. Intersystem crossing $(S_1 \text{ to } T_1)$ occurs between two excited states of similar energy and is very important in photochemistry, because the triplet state T_1 produced is even longer lived than S_1 . The lifetimes of T_1 are limited by: (i) phosphorescence, (ii) chemical reaction and (iii) radiationless decay to *So.* Furthermore, an excitcd moleculc may transfer its energy to another molecule; if this occurs the excited or donor molecule returns to its ground state and the acceptor molecule becomes excitcd. If the donor molecule (D) was in the triplet state at the time of its energy transfer, the acceptor rnolccule **(A)** gets promoted to its triplet state, and similarly we expect singlet-singlet energy transfer.

FIGURE 1. Potential energy curve for an clectronically excited and ground-state diatomic molecule.

FIGUKC *2.* Jablonslii diagrain. Typical **cnergy** diagram.

II. PHOTOCHEMISTRY OF AROMATIC DIAZONIUM SALTS

At the end of the last century, Andressen⁶ established that aqueous solutions of diazo compounds decompose under the influence of light. The mechanism of the photochemical decomposition is not completely understood; it appears that it is largely determined by the environment and by the chemical species present. The effect of substituents on the light sensitivity of diazo compounds has been studied by many investigators^{7, 8}.

A. Deamination through Diazonium Salts

According to Andressen⁶, diazonium salts irradiated in aqueous solution are decomposed into a phenol and nitrogen. Orton and coworkers⁹ investigated the photolysis of diazonium salts in aqueous solutions and found a noticeable increase in the rate of decomposition as compared to that of the dark reaction. When parasubstituted benzenediazonium derivatives are irradiated by ultraviolet light in methanol as the solvent, the formation of the phenol ether is rather a minor process, while the main reaction is the substitution of hydrogen for the diazo group¹⁰.

When 2,4,6-tribromobenzenediazonium sulphate was irradiated in aqueous solution, Orton and coworkers obtained 2,4,6-tribromophenol in quantitative yield⁹. When this reaction was carried out in acetic acid, 2,4,6-tribromophenyl acetate was formed, whereas in formic acid only 1,3,5-tribromobenzene was obtained.

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μ				
R	Reduction $(\%)$	Ether $(\%)$		
p -NO.	60	о		
p -Cl	60	O		
p -Mc	$40 - 50$	$20 - 30$		
p -OMe	$40 - 50$	$20 - 25$		

TADLE 1. Photolysis of compounds [RC,H,N,]+ **CI-** in niethanol solution

If the photolysis product is reactive enough, it can react under favourable conditions with the non-decomposed diazonium compound and a dye is formed in *the* exposed areas but not in the areas protected from light. The reactions taking place can be expressed as follows:

This process was first proposed by West¹¹ and further explored by Feer¹², Andressen¹³ and others.

When irradiating diazo conipounds in concentrated hydrochloric acid, Sus obtained the corresponding p-halogenodiphenylamines. **I-le** also showed that the action of light on $p₋$ and $q₋$ dialkylaminobenzenediazonium compounds results, in the case of *para* substitution, in the corresponding p-dialkylaminophenols, whercas the *ortho* isoniers produce a considerable quantity of secondary amine, along with the corresponding phenols^{14, 15}.

	R	R'	Solvent	2(%)	3(%)	4 $(\%)$
1a		$-C(H_2)_4$ -	$H_2O(20 °C, h\nu)$	51		
1b		Me Me	$H_2O(20 °C, h\nu)$	15		56
			1 M-HCl (20 °C, $h\nu$)	8	10	48
			1 M-HCl (100 °C, 90 min)		16	44
			$H2O$ (100 °C, 90 min)	20	0	57
1c	Et	Εt	1 M-HCl (20 °C, $h\nu$)			92
			$H_2O(20 °C, h\nu)$			70

TABLE 2. Photolysis of diazonium salts 1^{16}

8. Substitution of the Diazonium Group by Fluorine

In the photolysis of crystalline diazonium fluoroborate and hexafluorophosphates, the corresponding fluoroarenes were usually the only volatile product, as in pyrolysis1'. In some cases, the yield of fluoroarenes **was** higher than that obtained by pyrolysis of these salts (the Balz-Schiemann reaction), which remains the most generally used means of introducing a fluorine substituent into an aromatic ring¹⁸. Pure 4-substituted diphenylamine was isolated in **37%** yield on the first attempt when the corresponding diazonium fluoroborate was irradiated as a solid film at **30** *"C* with 3500 **A** light. Hexafluorophosphates sometimes gave better yields than tetrafluoroborates in the Balz-Schiemann reaction¹⁹:

$$
ArN_{2}^{+}BF_{4}^{-} (or PF_{6}^{-}) \xrightarrow[3500 \text{Å}]{h\nu} ArF + N_{2} + BF_{3} (PF_{5})
$$

Photolysis of aryldiazonium salts substituted with electron-withdrawing groups gave much lower yields of fluoroarenes, perhaps because with a light source available the products absorbed light about as **well** as the starting material and the competing

$\mathbf R$	X	Irradiation time (h)	Yield of RC_6H_4F (%)
$4-Et2N$	BF ₄	$4.5 - 49$	53–55
$4-Et2N$	PF_6	4	72
$4-Et2N$	PF ₆	94	74
$4-Me2N$	BF_{4}	17	55
4-MeO	BF ₁	19	69
4-PhNH	BF ₄	24	37
н	BF ₃	$\frac{2}{8}$	34
4-Cl	BF _a		10
$3-Ph$	BF ₄	$\overline{2}$	29
N_2^+	PF_6^-	12	40
N_{2}	BF_{4}^-	54	19

TABLE 3. Photolysis of solid arenediazonium fluoroborates and fluorophosphates, $RC_6H_1N_2^+X^-$

reaction destroyed them. An ionic intermediate or transition state of some **sort** appears to be implicated in the photolysis of solid arenediazonium fluoroborates and fluorophosphates. **If** aryl radicals were intermediates, they would be expected **to** abstract hydrogen atoms from other molecules of the starting material, especially when side-chains with alkyl groups are present. However, no reductive product **was** produced in the photolysis of even the dimethyl- and diethyl-amino compounds. In contrast, when crystalline **para-IV,N-dimethylaminobcnzenediazoniurn** chloride was irradiated, reduction to N,N-dimethylaniline (25%) competed strongly with conversion to **chloro-N,N-dimethylaniline (33%).** These observations are in accord

with the results of studies of isomer distribution in the phenylation products formed when solid benzenediazonium haloborates are pyrolysed or photolysed in the presence of a substituted benzene²⁰.

The lack of parallelism between fluoroarene yields for photolysis versus pyrolysis makes it unlikely that these reactions proceed exclusively through **a** common intermediate. Lewis and coworkers²¹ compared photochemical with thermal reactions of diazonium salts in aqueous solution and concluded definitely that there was no single intermediate common to both.

A wide variety of fluorinated analogues of biologically significant compounds have been synthesized and studied as potential enzyme inhibitors and as therapeutic agents²².

Kirk and Cohen studied synthetic approaches to fluoroimidazoles²³. Imidazolediazoniurn ions, prepared by diazotization of aminoimidazoles in tetrafluoroboric acid solution and irradiated *in situ*, decompose with formation of fluoroimidazoles in 30-40% yield. This procedure has been applied to 2-fluoroimidazoles, 4-fluoroirnidazole and ethyl **3-fluoroimidazole-5-carboxylate.** The ester, in turn, has served as the starting point for various transformations, including the synthesis of 4-fluorohistamine and 4-fluorohistidine. To date, fluorinated imidazoles have been obtained only by photochemical methods²⁴.

9. Photochemistry of thc diazonium and **diazo** groups **349**

Irradiation of diazotized N-acetyl-L-histidine mcthyl ester in tetrafluoroboric acid solution provides the 2-fluoroimidazole derivative in 32% yield; 2-fluorohistidine is obtaincd after ester hydrolysis and enzymic deacylation of the lattcr derivative. An analogous approach to 2-fluorohistamine failed because of decomposition during acid or alkaline hydrolysis of the N-acetyl group. The histamine analogue was obtaincd, howevcr, by use of the more labile trifluoroacetyl group for protection of the side-chain amine. In contrast to the vcry high degree of stability of 4-fluoroimidazoles, the 2-fluoro isorncrs are subject to fluorinc displacement²⁵.

C. Substitution of the Diazonium Group by Chlorine or by Hydroxy Groups

Solid anhydrous chlorides completely frce of polar influenccs were observed to bc photolysed with a quantum efficiency of well over unity. These results can be explained by considering the covalent diazo tautonier of the diazonium chloride, which decomposes symmetrically in the first step²⁶:

Free chlorine atoms and phenyl radicals so produced arc capablc of continuing the original function of the utilized photon. For example,

The observed quantum yiclds were of the order of two.

Aromatic diazonium salts react with hydrochloric or with hydrobromic acid under the influence of sunlight yielding halogenated aromatic compounds²⁷. In this way, for example, *6* is converted to **7** or *8,* respcctivcly.

The photoreaction of diazonium salts in very dilute sulphuric acid, like the thermal decomposition of diazonium compounds in aqueous acid solutions, yields hydroxy compounds.

The photoreaction proceeds very smoothly in many cases, while the corresponding thermal decomposition often leads to resin formation. **4-Anilinobenzenediazonium** sulphate *(6)* afforded **8,** and **4-dimethylaniinobenzenediazonium** chloride gave 4-dimethylaminophcno1. Huisgcn and Zahlcr have described the preparation of fluoren-9-one (11) by irradiation of 10 in aqueous sulphuric acid²³:

The photolysis of tetrazo conipounds of **1,5-di(o-aniinobenzoyInaphthalene)** takes place by means of a double Pschorr reaction to give **1,2,6,7-dibenzopyrene-3,8** quinone²⁹.

D. Spectra and Excited States **of** *Diazonium Salts*

The spectra of diazonium salts are of interest here because they identify the excited states which arise in the act of light absorption and arc potentially involved in the photolytic conditions. The simplest diazonium compounds absorb ultraviolet radiation in the range 2200-3300 **A,** which is virtually non-cxistent in the spectrum emitted by conventional light sources. When dialkylamino, arylamino, arylthio or hydroxy groups are introduced into the molecule in the *ortho* or para positions with respect to the exceptionally strong electron attracting diazonium group, then the absorption is sharply moved to longer wavelengths. Since most substituted diazonium compounds are yellow and absorb in the region 3600-4500 Å, they are sensitive to near ultraviolet and blue light, and are virtually insensitive to red and green¹⁰. Figure 3 shows the absorption spectra change of **p-CGH,NHC,;H4N:** HSO; in water **(pfi 4.6)** during U.V. irradiation3".

In aqueous solution, diazonium **snlls** stabilized by zinc chloride exhibited a major absorption band in **thc** region 350-390 nm and n minor band in the region 245- 260 nni. The introduction of progressivcly Iargcr alkyl or phcnyl groups into the *para* amino group causes a slight bathochromic shift in the major band. With every compound a minor band was observed in the region $245-260$ nm³¹.

X (nm)

FIGURE 3. Absorption spectral change of $4-C_6H_6NH_8C_6H_4N_2^+HSO_4^-$ in water (pH 4.6) during irradiation. Irradiation with a 500-\valt, high pressure **Hg** arc at about 30 cm. Irradiation time (sec); (a) 0; (b) **15;** (c) 20; (d) 30. [Reproduced, with pcrrnission, from Inoue, Kokado and Shirnada, *Nippot; Kagaku Kaislii,* **12,** 2272 (1974).]

	$R'R^iN$ N_2^+ CI ⁻¹ $_2^+$ ZnCI ₂			
R ¹	R^2	λ_{max}	Molar extinction coefficient, ε	
н	H	353	37,400	
Me	Me	375	38,900	
Eι	Et	380	40,300	
Me	н	366	38,600	
Et	H	368	41,100	
P _h	н	377	30,200	
Ph	Ph	388	24,100	

TABLE 4. Ultraviolet absorption maxima of

 $\sqrt{2}$

Although the $n-\pi^*$ transition is observable in the absorption spectra of azo compounds, it has not yet been observed with diazonium salts. The main absorption band with these salts is not greatly affected by solvents of different polarity and is probably due to π - π ^{*} electron excitation. Bathochromic shifts observed with $p-NH₂$ -substituted derivatives also confirm this conclusion. In aqueous solution no difference in spectra between p-aminobenzenediazonium chloride and its zinc chloride double salt was found, indicating that interaction between the cation and zinc chloride was negligible.

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The electronic states of the Scnzenediazoniiim cation wcre calculated by **a** method similar to that of Pariser and Parr³². On the basis of the calculations, the 259 and 295 nm absorption bands were attributed to ${}^{1}A_{1}{}^{-1}A_{1}$ and ${}^{1}A_{1}{}^{-1}B_{1}$ transitions, respectively. When the sample was irradiated in an alcoholic solution by a **295** nm light, no e.s.r. signal was detected, but fluorescencc and phosphorescence were observed; the fluorescence and the absorption spectrum of the ${}^{1}A_{1}{}^{-1}B_{1}$ transition are niirror images of each other. On the other hand, upon irradiation by **259** nm light, this sample showed an e.s.r. spectrum which was attributed to the phenyl c-radical, but no fluorescence or phosphorcscence was observed. These results

FIGURE 4. Thc near ultraviolct absorption spectrum of benzenediazoniuni fluoroborate in ethanol (* in water). [Reproduced, with permission, from Sukigahara and Kikuchi, *Bull*. *Chetn. SOC. Japan,* **40, 462 (1** *967).]*

show that thc electronically excited benzcnediazoniurn salt releases its excess energy by the following processes: (i) There is radiative transition from the **'B,** state io the ground state and intcrsystein crossing to the phosphorcscent state, although it is as yet uncertain whether ${}^{3}B_1$ or other triplet states are involved. The photodecomposition does not procced from the excited ${}^{1}B_{1}$ state. (ii) From the excited **lAl** statc, neithcr radiative transition to the ground state nor **a** non-radiative transition to any fluorescent or phosphorescent state occurs. The molecule decornposcs rapidly to give a phenyl c-radical after cxcitation *to* this electronic state.

Electronic structures and absorption spcctra of p-halogeno-, p-methoxy- and **p-dinicthylaniino-bcnzenediazoniurn** cations showed that when an electron-donating group is introduced at the *para* position, the $^1A_1^{-1}A_1$ transition shows a very large shift to longer wavelengths, while the ${}^{1}A_{1}{}^{-1}B_{1}$ transition is not so much affected³³.

Tsunoda and Yainaoka reported the relationship between the charge density and the photolytic rate3'. **It** was observcd that the benzcnediazonium cations having electron-releasing groups such as $-OMe$ and $-NMe₂$ in the para position have rclatively large quantum yields, while electron-attracting groups, such as $-NO₂$ and -Br, substituted in the *para* or *ortho* position lower the quantum yields. As

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FIGURE *5.* The fluorescence emission and excitation spectra of benzencdiazonium fluoroborate. (Reproduced, with permission. from Sukigahara and Kikuchi, Bull. *Clienr. SOC. Japan,* **40, 462 (1967).]**

FIGURE *6.* The e.s.r. spectrum of irradiated benzencdiazonium fluoroborate (at 77 K in methanol). [Reproduced, with permission, from Sukigahara and Kikuchi, *Bull. Chem. Soc. Japan,* **40, 462 (1967).]**

the conjugated system extends, the peak of the absorption band is shifted to longer wavelengths, but the quantum yield tends to be lower. The tetrazonium cations also have low quantum yields, which might be the effect of the strong electrophilicity of two diazonium groups.

FIGURE 7. The phosphorescence spectrum of benzenediazonium fluoroborate at 77 **K.** [Reproduced, with permission, from Sukignhara and Kikuchi, *Biill. Cliern. Soc.* Japan, **40, 462** (1 **967).]**

FIGURE 8. The visible and **near** ultraviolet absorption spectra **of** the para-substituted benzenediazonium fluoroborates. **(A)** *p-F;* (B) *p-CI;* (C) **p-Br;** (D) p-MeO; (E) **p-I;** (F) p-NMe,. [Reproduced, with permission, from Sukigahara and Kikuchi, *Bull. Chem. SOC.* Japan, **40,** 1078 (1967).]

Table 6 shows the transition energy and the charge density of the π -electrons calculated by the Huckel theory. The charge density **of,** for example, the p-dimethylaminobcnzenediazonium cation having a high quantum yield is much greater than that of, for example, the p-nitrobenzenediazonium cation with **its** low quantum 9. Photochemistry of the diazonium and diazo groups

TABLE 5. Wavelength maxima and molar extinction coefficients for para-substituted benzenediazonium cations. [Reproduced, with permission, from Sukigahara and Kikuchi, Bull. Chem. Soc. Japan, 40, 1078 (1967).]

λ_{max} (nm)	$\log \varepsilon_{\text{max}}$
267	4.229
281	4.195
393	4.178
327.5	4.036
315	4.283
380	4.448

yield. A correlation between the quantum yield and the charge density of the diazonium group also exists, even in the excited state. Figure 9 shows that the diazonium cation with a high charge density at the diazonium group in the excited state has a tendency to give high quantum yield. The ortho isomers do not follow this tendency, because of steric hindrance.

TABLE 6. Quantum yields and charge densities of diazonium cations,

[Reproduced, with permission, from Tsunoda and Yamaoka, J. Photographic Sci. Japan, 29, 197 (1966).]

 $G =$ ground state, $E =$ excited state.

FIGURE 9. The relationship between the quantum yield and the charge density of the **diazoniurn group.** [Reproduced, with permission, from Tsunoda and Yarnaoka, *J. Phoro*graphic *Sci. Japan, 29,* **197 (1966).]**

E. Reaction Intermediates

1. Radical species and carbonium ions

The products of the photodecomposition of p- and m-nitrobenzenediazonium salts have becn studied in various solvents. It **was** found that the dominant primary prccess in alcohol solution leads to nitrophenyl radicals, and the products are those generally expected of a reaction involving a homolytic scission of the C-N **bond.**

Product	EtOH at $O^{\circ}C$ (%)	EtOH at -10° C (%)	EtOH + I_2 at 0 °C (%)
PhNO ₂	76.6	83.5	$17-7$
p -NO ₂ C ₆ H ₄ OC ₂ H ₅	$5-4$	$5-2$	4.5
$p\text{-}NO_2C_6H_4CH(OH)CH_3$	3.6	5.3	a
p -NO ₂ C ₆ H ₄ CH ₂ CH ₂ CH ₂ OH	2.3	2.8	a
CH ₃ CHO	74.1	\boldsymbol{a}	a
$(CH_3CH(OH))_2$	1.54	a	a
$(p\text{-}NO_2C_6H_4-)_2$	$0-0$	0.0	0.0
$p\text{-}NO_2C_6H_4I$			65.5
$(\sum p\text{-NO}_2\text{C}_6\text{H}_4\text{-})$	82.5		83.2

TABLE 7. Photolysis of $p\text{-}NO_2C_6H_4N_2^+Cl^{-1}$ **SnCl₄ in ethanolic solution**

Analysis not carried out but compound presumed present.
9. Photochemistry **of** the diazonium and diazo groups **³⁵⁷**

Nitrobenzene, *a-* and (3-arylethanol and butanediol are formed but **110** dinitrobiphenyl³⁵. Addition of 2,2-diphenylpicrylhydrazyl, halogen or nitric oxide confirmed the importance of nitrophenyl radical formation in the photodecomposition. However, all of the products cannot be explained on the basis of the one primary process. The results are consistent with the occurrence of two primary processes, (1) and (2):

$$
O_{2}NC_{6}H_{4}N_{2}^{+}X^{-}+h\nu \xrightarrow{\qquad} O_{2}NC_{6}H_{4}^{+}+N_{2}+X \tag{1}
$$

$$
O_{2}NC_{6}H_{4}N_{2}^{+}X^{-}+hv \longrightarrow O_{2}NC_{6}H_{4}^{+}+N_{2}+X^{-}
$$
 (2)

where X^- is Cl^- , $SnCl_5^-$ or ion solvent dipole complexes, etc. Presumably (1) is followed by the secondary radical reactions which lead to the final products; in ethanol solution the suggested secondary reactions are:

$$
O_2NC_6H_4+CH_3CH_2OH \xrightarrow{\qquad} C_6H_3NO_2+CH_3CHOH \tag{3}
$$

$$
O_2NC_6H_4^{\bullet}+CH_3CH_2OH \xrightarrow{\qquad} C_6H_3NO_2+^\bullet CH_2CH_2OH
$$
 (4)

the secondary radical reactions which lead to the final
\non the suggested secondary reactions are:
\n
$$
O_2NC_eH_e+CH_3CH_2OH \xrightarrow{\qquad} C_eH_3NO_2+CH_3CHOH
$$

\n $O_2NC_eH_e+CH_3CH_2OH \xrightarrow{\qquad} C_eH_3NO_2+·CH_2CH_2OH$
\n $O_2NC_eH_e+CH_3CHOH \xrightarrow{\qquad} O_2NC_eH_4-CHCH_3$
\n $O_2NC_eH_4++CH_2CH_2OH \xrightarrow{\qquad} O_2NC_eH_4CH_2CH_2OH$
\n $2 CH_3CHOH \xrightarrow{\qquad} CH_3CHCHCHCH_3$
\n $CH_3CHOH \xrightarrow{\qquad} CH_3CH_2OH+CH_3CHO$
\n $O_2NC_eH_4+CH_3CHOH \xrightarrow{\qquad} C_eH_3NO_2+CH_3CHO$
\n $O_2NC_eH_4++CH_2CH_2OH \xrightarrow{\qquad} C_eH_3NO_2+CH_3=CHOH$
\n CH_3CHO

The absence of dinitrobiphenyls in the reaction products suggests that the nitrophenyl radicals are converted primarily to nitrobenzene and other products via reactions **(3)** and **(4).**

The second primary process *(2)* of lower probability was suggested to explain the ether formation in the ethanol solutions:

CH_aCHO
\nof dinitrobiphenyls in the reaction products suggests that the nitro-
\nls are converted primarily to nitrobenzene and other products via
\nand (4).
\nprimary process (2) of lower probability was suggested to explain the
\non in the ethanol solutions:
\n
$$
O_2NC_6H_4^- + CH_3CH_2OH
$$

\n $\left[O_2NC_6H_4^- - O-CH_2CH_3\right] \longrightarrow O_2NC_6H_2OCH_2CH_3 + H^+$ (5)
\n $\left[H \right]$
\nmechanisms (2) and (5), the diagram salt was photolysed in the
\ndine. The amount of *p*-nitrophenetole was not essentially decreased
\nyield of nitrobenzene, which reflects the activity of the *p*-nitrophenyl
\nreatly decreased due to the very fast reaction (6):
\n $O_2NC_6H_4^+ + I_2 \longrightarrow O_2NC_6H_4I$ (6)

To test the mechanisms **(2)** and *(5),* the diazoniurn salt was photolysed in the presence **of** iodine. The amount of p-nitrophenetole was not essentially decrcascd although the yield of nitrobenzene, which reflects the activity of the p-nitrophenyl radical, was greatly decreased due to the very fast reaction (6) :

$$
O_2NC_6H_4 \cdot + I_2 \longrightarrow O_2NC_6H_4I \tag{6}
$$

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There is a striking contrast between the products of the photodecomposition of the diazonium salt in alcohol and in aqueous solution. Nitrobenzenc, the major product in alcohol, is not detected in aqueous solution. Nitrophenol and nitrochlorobenzene dominate in water solutions. These results, coupled with product

Product	Solvent $(\%)$					
	EtOH	MeOH	H.O	$5.3 M -$ NaCl(aq)	9 _M HCl(aq)	$4.3 M -$ H ₂ SO ₄ (aq)
PhNO ₂	76.6	71.9	0.0	0.0	0 ₀	0.0
p -NO ₂ C ₆ H ₄ OR	5.3	3.5	0.0	0.0	0.0	0.0
p -NO ₂ C ₆ H ₄ Cl	0.0	0 ₀	7.4	30.2	61.6	0.0
p -NO ₂ C ₆ H ₄ OH	0 ₀	0.0	73.5	50.8	23.2	84.3
Coupled product	0.0	0.0	$4 \cdot 1$	5.2	0.0	0.0

TABLE 8. Variation of product yield in the photolysis of $p\text{-}NQ_2C_6H_4N_2^+$ **Ci⁻¹** $\frac{1}{2}$ **SnCl₄ in** various solutions

studies with nitric oxide and hydroquinone, suggest that free radical formation is unimportant in the photodecomposition of nitrobenzencdiazonium salts in aqueous solution. The results are consistent with the occurrence of a single primary photodecomposition mode, process **(2),** under these conditions. The product distribution in solutions of varied chloride content results from the competition between Cland **H,O** for the carbonium ion. 0.0 0.0 0.0 0.4 1 5.2

ide and hydroquinone, suggest the

ide and hydroquinone, suggest the

ide and hydroquinone, suggest the

are consistent with the occurrence

process (2), under these condition

chiomic c ide and hydroquinone, suggest the
hotodecomposition of nitrobenzene
are consistent with the occurrence
process (2), under these condition
d chloride content results from the
ponium ion.
 $Q_2NC_6H_4^+ + C_1^- \longrightarrow Q_2NC_6H_4C$
 Q_2

Boudreaux and Boulet³⁶ have made magnetic measurements on the irradiated aqueous solutions of **p-dirnethylaminobenzenediazonium** chloride. They have suggested that the observed magnetism is due to the C1- radical and to the $Me₂NC₈H₄$ ^{*} radical, and they have estimated that the steady-state concentration of radicals is about 1×10^{-3} m. This conclusion seems untenable in view of the work of Zandstra and Evleth³⁷. Photolytic decomposition of p-dimethylaminobenzenediazonium chloride in aqueous solution at pH above *5* leads to the formation of a paramagnetic species. The e.s.r. spectrum is the same as that obtained in air oxidation of p-dimethylaminophenol in the same pH range. Hence the structure of the photolytic radical could be established as the p-dimethylaminophenoxy radical. Similarly, **p-diethylarninobenzenediazonium** chloride was photolytically decomposed to yield the corresponding phenoxy radical.

In the presence of chloride ion, aryl chlorides as well as phenols result from the photolysis of arenediazonium salts in aqueous solution, as in the thermal reaction?'. Benzenediazonium ion in aqueous sodium chloride yields just as much chlorobenzene by photolysis as it does thermally at the same temperature, but the p -methyland **p-chloro-benzenediazoniuni** salts give a significantly higher yield of the chloride photochemically than thermally". **A** similarity, but not identity, of productdetermining steps is indicated. When $p-MeC_6H_4^{15}N+$ and $p-MeOC_6H_4^{15}N=N$

9. Photochemistry of the diazonium and diazo groups

					TABLE 9. Yield of aryl chloride from photo-		
--	--	--	--	--	---	--	--

a Thermolysis at 3 *"C.*

^aThermolysis at **49** *"C.*

Tliermolysis at **80 "C.**

are exposed to light insufficient to complete the photolysis, the residual diazonium salt is in part rearranged to $ArN^{\dagger} \equiv 15N$, significantly more than in the corresponding dark reaction. This allows us to reject the aryl cation **as** the only intermediate. **A** simple mechanism adequate to describe all results is given below:

$$
Ar-^{15}N^{+} \equiv N \xrightarrow{\lambda \nu} D^{*} \tag{7}
$$

$$
D^* \longrightarrow Ar^{-1}N^+ \equiv N \tag{8}
$$

$$
D^* \longrightarrow ArN^+ \equiv 1^sN \tag{9}
$$

$$
D^* + H_2O \longrightarrow ArOH + N_2 + H^+ \tag{10}
$$

$$
D^* + Cl^- \longrightarrow ArcI + N_2 \tag{11}
$$

Reaction (7) conceals the complex process following the absorption of light and leading to a state D^* (unestablished multiplicity) stable enough to undergo the bimolecular reactions (10) and (11). Reaction (8) is included simply because it seems unreasonable that reaction (9) \vould occur without reaction (S). **It** is attractive to identify D* with the intermediate in the photolysis of **thep-nitrobenzenediazonium** ion, which may participate **in** water reactions (10) and (11) **as** well as in ethano!, reactions (12) and **(13),** to account for all the observed products: $D^* + C^{\dagger -} \longrightarrow ACCl + N_2$ (11)

als the complex process following the absorption of light and
 D^* (unestablished multiplicity) stable enough to undergo the

ns (10) and (11). Reaction (8) is included simply because it

tha als the complex process following the absorption of light and
 D^* (unestablished multiplicity) stable enough to undergo the

ns (10) and (11). Reaction (8) is included simply because it

that reaction (9) would occur w

$$
D^* + EtOH \longrightarrow ArcOEt + N_2 + H^+ \tag{12}
$$

$$
D^* + E\{OH \longrightarrow \longrightarrow Ar + N_2 + CH_3CHOH + H^+ \tag{13}
$$

One may speculate that D* corresponds to the species observed by Lee, Calvert and Malmberg³⁵ in the low temperature irradiation of p -dimethylaminobenzenediazonium hexachlorostannate, for which the symmetric structure **(12)** was tentatively suggested by Calvert and Pitts³⁹.

2. Cis-trans isomerization

The light-induced rearrangements of diazo compounds also give some information for the reaction intermediate of diazonium salts, but only the diazosulphonates $ArN=NSO₃Na$ and the diazocyanides $ArN=NCN$ have been studied in detail⁷.

It is assumcd that these exist in two forms: the unstable *cis* form, able to take part in the reaction of azo coupling, and the stable, inactive, *trans* form which cither takcs part in such a reaction only with great difficulty or not at all.

Hartley¹⁰ found that, when irradiated in a non-ionizing solvent (benzene), the stable diazocyanide is convertcd to the unstable isomer which is in a state of equilibrium with the stable form, and can be detected by its absorption spectrum. While irradiating the stable diazocyanide in ethanol in the presence of silver nitrate, Stephenson and Waters¹¹ showed that when the irradiation lasts a sufficiently long time, silver cyanide precipitates quantitatively from the solution, and the diazocyanide is convertcd to thc diazonium nitrate. When comparing these results with the rcsults of Hartley, they arrived at thc conclusion that diazonium cyanide could be obtained only as a result of the action of light on the labile form of diazocyanide, sincc thcy wcrc ablc to show that the stable form of diazocyanide was unable to undergo this reaction in the dark and, consequently, no direct conversion of the stable diazocyanidc to diazonium cyanide took place on irradiation. In non-polar solvents the process of photostercoisomerization is confined, as **a** rule, to the formation of the unstable stereoisomer, whereas irradiation in polar solvents, e.g. ethanol or acetone, results in the complete decomposition of the diazo group:

Contrary to the above work, de Jonge and Dijkstra⁴² noticed that immediately after irradiation of thc stable diazosulphonate in solution, the anion of sulphurous acid and the diazonium cation are dctected, which slowly reacted with one another in darkness, to givc the initial stable diazosulphonatc. The elimination of one of the ions, for instance the precipitation of the $\overline{SO_3^2}$ anions with lead salts and/or the conversion of the diazo cation to an azo dye, cuts short the dark reaction, and the system does not revert to thc initial state. **.4** short duration, high intensity irradiation of neutral solutions of p-niethoxy- or **o-chloro-p-methyl-benzenediazosulphonate** results in an intermittent increase in the pH of the solution. After the light source has been turned off, the solution reverts gradually to the original state, and the pH decreases to the initial value. This phenomenon becomes understandable if we assume that, **as** a result of the action of light, there occurs at least partial dissociation of diazosulphonate according to the equation:

$$
\text{MeO} \left(\bigcircled{N} - N = \text{NSO,Na} \quad \xrightarrow{h \cdot \rightarrow} \quad \text{MeO} \left(\bigcircled{N} - N = N^+ + \text{SO}_3^{2-} + \text{Na}^+ \right)
$$

9. Photochemistry of the diazonium and diazo groups 361

Now, the solution of the initial diazosulphonate is neutral, but the product is a salt of the weak sulphurous acid, and the pH of the solution therefore increases.

During more prolonged exposure the process becomes irreversible because of the

decomposition of the diazonium cation.
 $ArN_t^+ + H_2O \longrightarrow ArOH + N$ During more prolonged exposure the process becomes irreversible because of the decomposition of the diazonium cation.

$$
ArN_2^+ + H_2O \xrightarrow{\Lambda \nu} ArOH + N_2 + H^+
$$

3. Sensitized photolysis of diazonium salts

The photolysis of diazonium salts in solution has been spectrally sensitized with dyes in the presence of activators, such as amine associates, thiourea and its derivatives, and benzenesulphinate ion⁴³. p-Anilinobenzenediazonium sulphate (13) was photolysed in the presence of methylene blue and of **1,4-diazobicyclo[2.2.2]** octane **(DABCO)** or sodium p-toluenesulphinate **(STS).** Figure 10 **shows** the

FIGURE 10. Absorption spectra of solutions of **13** containing methylene blue and activator under anaerobic **conditions.**

absorption spectra of the solution before and after irradiation. The quantum yields were also pH sensitive. Figure 11 shows the variation in the quantum yields as a function of pH under aerobic conditions. It is seen that the pH for greatest dye sensitization differs markedly between the two activators. In the **DABCO** system, the dye sensitization was restricted to an alkaline range starting at about pH 7. Lower pHs may accentuate the formation of the protonated form of **DABCO** which results in a loss of photo-initiating activity.

The data given show clearly a greater range of activity in the **STS** system. The quantum yield increased with an increase in solution in the pH range below 7, and decreased with greater alkalinity.

It may be inferred that the dye-sensitized photolysis of **(13)** does not take place via the sulphonyl radical, which is formed from the dye triplet reaction with a sulphinate ion, and which is an efficient initiator for polymerization.

The reaction related to the sensitization may be written as follows:

Sens *(G)+hu* - 5ens' 'Sens*+DABCO ___j (Sens - -- **DABCO)** - Sens.+DABCO Sens*+(ll) ___j Sens (G)+N,+Ar.

The key step of the sensitizing system may be the escape of Sens \cdot , semi-methylene blue, from the dye-activator complex, Sens - - - **DABCO,** which is formed by interaction of the excited triplet dye with unprotonated **DABCO.** The complex undergoes collisional deactivation and dissociation to the semi-methylene blue and a half-oxidized activator.

FIGURE 11. Depcndence of quantum yields on solution **pH** under aerobic conditions.

111. PHOTOLYSIS OF ALIPHATIC DlAZONlUM COMPOUNDS

Photolysis of the sodium salts **14** and **15** of camphor and fenchone arenesulphonyl hydrazone, respectively, gives both *cso-* and cndo-diazoniuni ions. In each case a mixture of pinyl and tcrpinyl products is formed rogethcr with bornyl products from **14** and fenchyl from **15,** without any cross-over of thesc series. The ratio of pinyl to terpinyl products requires that two isomeric, interconverting cations of pinyl type **16** and **17** are formed. Similar results are reported for the apocaniphor scries⁴⁴. On the other hand, the *p*-toluenesulphonylhydrazone of norbornanone protonates exclusively from the *endo* face, affording $exo-2$ -norbornanediazonium ion which gives nortricyclene and $exo-2$ -norbornyl ether¹⁵. Photolysis of the tosylhydrazone **(18)** gives the tricyclo[3.2.1 .02~']oct-3-en-6-yl cation **(19),** which rearranges reversibly to 20 at a rate competing with solvent attack⁴⁶. The tosylhydrazones, 21, **28** and **30**, were prepared from 4-hydroxy-endo-tricyclo[4.2.1.0^{2,5}]non-7-en-3-one and utilized in the deaminative pinacol rearrangement to endo-tricyclo[3.2.1.0^{2,4}]oct-6-ene derivatives¹⁷. The hydroxy compound 21 underwent retroaldol cleavage exclusively on irradiation in methanol/NaOMe; 28 and 30 afforded the desired c~ndo-tricyclo[3.2.1 .O **2~** i]oct-6-ene derivative, **29** and **32,** rcspectively. Competing processes such as double-bond participation in the case of **28.** arid hydride shift in the casc of **30,** render these pinacol rcarrangeinents lcss cficicnt than the related

9. Photochcmistry **of** thc diazonium and diazo groups 365

Favorskii reaction. The bicyclo[4.2. Ilnona-2,4,7-trien-9-yl cation has invoked interest because of its potential homoaromaticity¹⁸ and bicycloaromaticity⁴⁹. The solvolysis of syn-bicyclo^[4.2.1] nona-2,4,7-trien-9-yl p-toluenesulphonate (33) afforded exo-dihydroindenyl acetate (34)⁵⁰ and/or indene^{51, 52}, depending on the reaction conditions. Deamination, on the other hand, starting from the highly energetic diazonium ion, passes its transilion state 'early' and without significant distortion of nuclear positions (vertical ionization)". Consequently, deaniination may produce cations of undistorted geometry which are by-passed in solvolysis (leading directly to more stable bridged species). Photolysis of **bicyclo[4.2.l]nona-2,4,7-trien-9-one**

tosylhydrazone (37) in McOH-MeONa afforded syn-9-methoxybicyclo[4.2.1]nona-2,4,7-triene **(40)** and **(35)** as the major products. Minor products were *endo*dihydroindenyl methyl ether **(41),** indenyl methyl ether **(42)** and indene. Photolysis of tosylhydrazone sodium salts generates diazo compounds which are protonated by protic solvents to **give** diazoniuin ions and products derived therefrom. Various

bicyclo[2.2.l]hcptcnyl **(47)** and bicyclo[3.2.0]heptenyl **(SO)** cations have been generated by decomposition of suitable diazonium precursors **(43** and **53).** The cations **47** and **50** do not equilibrate, and their substitution pattern **is** largely independent of the specific precursor⁵⁴.

IV. PHOTOLYSIS OF DIAZO COMPOUNDS

The photolysis of diazo compounds is **an** important route to the formation of carbenes :

Many reviews and descriptions of carbenes are available⁵⁵⁻⁸⁵.

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A. The Structure of Carbenes

Two electronic configurations may occur: a singlet configuration **(A)** or a triplet configuration (B).

The electronic spectrum of methylene produced in the gas phase by flash photolysis of CH₂N₂, CHDN₂ and CD₂N₂ was recorded by Herzberg and Shoosmith⁸⁶, E.s.r. spectra of methylene trapped in a xenon matrix at 4.2 X have been one of the most recent achievements in this field^{87, 88}.

The e.s.r. signal survived warming to **20** K but disappeared upon warming to 77 K. Analysis of thc results demonstrated the roduction of **a** linear or nearlinear species with a $C-H$ bond length of 1.03 Å, which absorbed u.v. radiation at about 1414 Å, and a bent species, bond angle $102-103^\circ$, C-H bond length 1.12 Å, which absorbed radiation in the 5500-9500 **A** region. Moreover, in the presence of a large excess of nitrogen, the absorption of the short wavelength, lincar **CH2** increased at the expense of that of the long wavelength, bent $CH₂$. This experiment pointed to triplet CH₂ as ground state. The short-lived bent or singlet CH₂ was being degraded through collisions with the moderator, nitrogen, and dropping to the more stable species. Wasserman and coworkers⁸⁸ observed two species with $D = 0.6636$ cm⁻¹, $E = 0.002$ cm⁻¹ and $D = 0.6844$ cm⁻¹, $E = 0.00347$ cm⁻¹, respectively. The value of *D* is related to the average $1/r^3$ where *r* is the distance between the two spins, so that a high value of *D* implies a large spin-spin interaction and a close proximity of the two spins. The value of E indicates qualitatively the magnitude of the deviation of the spin-spin interaction from cylindrical symmetry. The site with the lower *D* and *E* values in methylene is believed to allow greater motional freedom; a bent molecule which rotates freely about the long axis will have $E = 0$. Wasserman and coworkers arrived at a $H - \ddot{C} - H$ angle of 136° in good agreement with most theoretical calculations. Angles of **150-155"** have beenestimated for phenyl- and diphenyl-carbenes. It was found that for diphenylcarbene the niagnitude of $D = 0.4055$ cm⁻¹ and $E = 0.0194$ cm⁻¹ shows a small dependence on the nature **of** the host matrix:

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The higher *D* value of diphenylcarbene shows that two unpaired electrons are largely localized on the divalent carbon atom⁸⁹.

An orbital representation comparing a triplet ground-stare molecule with a phosphorescent triplet state is given by Figure 12. Several carbenes have been assigned a triplet ground state by e.p.r. experiments⁹⁰⁻⁹⁵. Non-radical products of photolysis of diazo compounds can not be determined by the e.p.r. technique, SO that it became necessary to study the optical spectroscopy of the carbene species. Ultraviolet spectroscopy, fluorescence excitation and absorption spectroscopy have been used to study the carbene problem.

FIGURE 12. Orbital representation **of** spin state. [Reprinted, with permission, from Trozzolo, *Accounrs Chem. Res.,* **1, 329 (1968).** Copyright by the American Chemical Society.]

Closs and coworkers rcported the long-wavelength absorption band of diphenylcarbene oriented in single crystals of 1,1-diphenylethylene⁹⁶. Moritani and coworkers found two new absorption bands in photolysed solutions of diphenyldiazomethane, by correlating e.s.r. studies, fluorescence, fluorescence excitation and absorption spectroscopy⁹⁷. Gibbons and Trozzolo reported the luminescence, excitation and absorption spectra of diphenylcarbene⁹⁸. The same spectrum was obtained for all excitation wavelengths in the range 230-470 nm. The emission was identified as the fluorescence of triplet ground-state diphenylcarbene^{99, 100}.

FIGURE 13. Luminesccnce of diphcnylcarbene. [Reprinted, with permission, from Trozzolo, *Accounts Chem. Res.,* 1, 329 (1968). Copyright by the American Chemical Society.]

The photolysis of other diaryldiazomethanes gave almost exclusively the corresponding substituted diphenylniethylene which e.s.r. studies have shown to be ground-state triplet molecules in each case.

The maxima of both the absorption and emission bands were shifted to longer wavelengths from those of diphenylcarbene; the short wavelength bands were more intense than the visible range bands. The absorptions are considered to involve π ⁺ transitions (Table 10). Moritani and coworkers^{97, 101} have obtained the

in 2-methyltetrahydrofuran. [Reprinted. with permission, from **Trozzolo,** *Accoroifs Chew. Res.,* **1, 329 (1968).** Copyright by the American Chemical Society.]

electronic spectrum of 61 (λ_{max} 510 nm) by photolysis of the corresponding diazo compound **(60)** in **a** variety of rigid matrices at 77 K and also in liquid paraffin at room temperature. The final product is the free radical dinier, and the decay of **61**

was assigned to **the** doublet free-radical **62.** The e.s.r. spectra obtained by photolysis of 3-bis(diazomethyl)benzene and **1,3-bis(cc-diazobenzyl)benzenc** were much more complex than those observed with triplet states^{102, 103}. The data have been analysed in terms of the corresponding quintet ground states of the dicarbencs **63** and **64.**

In a *para* arrangement, resonance interaction permits the ground state to be a quinonoid triplet, rather than a quintet¹⁰⁴.

E.s.r. experiments also show that at 4 K or 77 K , such triplets as α -naphthylcarbene can exist in two non-interconverting isomeric forms, which are possible since the

triplet is not linear. Delocalization of one of the triplet's electrons over the π -system presumably provides enough double-bond character between the carbenic centre and the ring to provide a barrier to rotational interconversion at 77 K **Io5.**

The spectrum of 9,9'-dianthrylcarbene *(65)* reveals that it is linear, however, with two aromatic systems lying orthogonally. This structure permits extensive delocalization through allenic resonance forms such as **65a Io8.** The septet ground

state is suggested for the tris-carbene (66) on the basis of its e.s.r. spectrum¹⁰⁷.

(66)

B. Chemical Induced Dynamic Nuclear Polarization (CIDNP) Studies

In the CIDNP method, the existence of a radical intermediate in **a** reaction is implied through n.m.r. investigation of the product of that reaction as it is being formed. Polarization of the product by radical precursors may be observed experimentally as enhanced absorption or as emission in the n.m.r. spectrum of the product¹⁰⁸. The observed signal directions are determined by the initial electron spin multiplicity of the radical pair and of its precursor¹⁰⁹⁻¹¹³. A set of simple selection rules based on the radical pair model has been developed to permit qualitative prediction of the effects¹¹⁴.

Methylene has been the subject of extensive CIDNP investigation in its reactions with alkyl halides and polyhalomethanes¹¹⁵⁻¹¹⁸.

9. Photochemistry of the diazonium and diazo groups 371

Singiet methylene produced via direct photolysis of diazirine reacts with a chlorine atom of CDCI₃ to give polarized CI₂DC-CH₂Cl¹¹⁹, probably through the formation of a chloronium ylide which gives the polarized product via a radicalpair intermediate. Similar reaction of diazirine forms **a** triplet radical pair by deuterium atom abstraction from CDCl₃; recombination of the resulting radical pair then affords polarized $Cl_3C-CDCH_2^{-119, 120}$.

Polarized Cl₂DC - CH₂Cl

The fundamental difference in the mechanism of reaction of singlet and triplet methylene with toluene has been shown by an investigation of the direct and sensitized photolysis of diazomethane¹¹⁸. Ethyl benzene is formed in both cases, but direct photolysis gives methylene which inserts concertedly (no CIDNP signal being observed) whereas the sensitized photolysis gives **a** signal consistent with a mechanism involving energy transfer and triplet methylene (equations 14-17). The Polarized Cl₂DC – CH₂Cl

damental difference in the mechanism of reaction

with toluene has been shown by an investige

photolysis of diazomethane¹¹⁸. Ethyl benzene is

photolysis gives methylene which inserts conce **SENDIDE TREE CREAT SET (T,)**

Interval difference in the mechanism of reaction of singlet and triplet

toluene has been shown by an investigation of the direct and

lysis of diazomethane¹¹⁸. Ethyl benzene is formed in **RR'C=N₂(T₁) RR'C:(T₁)+SH** (16)
 RR'C:(T₁)+SH (16)
 RR'C:(T₁)+SH (16) RR'C:(T₁)+SH RR'C:(T₁)+SH REACH B is followed in both cases,

RR'C=N₂(T₃) **RDF RR'C:(T₁)**+SH **RR'C:(T₁)**+SH REACH RANGER RANGER

Sens
$$
(S_0)
$$
 $\xrightarrow{h\nu}$ Sens (S_1) $\xrightarrow{}$ Sens (T_1)
Sens $(T_1) + BP'C-N$ (S_1) $\xrightarrow{}$ Sens $(S_1) + BP'C=N$ (T_1) (14)

$$
RR'C = N_{\bullet}(T_{\bullet}) \xrightarrow{\sim} RR'C:(T_{\bullet}) + N_{\bullet}
$$
 (15)

$$
RR'C:(T_1)+SH \xrightarrow{\qquad} RR'CH+S
$$
 (16)

$$
RR'CH + S \cdot \xrightarrow{\qquad} RR'CHS \tag{17}
$$

photosensitized decomposition of methyl diazoacetate was studied by examining the effect of the diazoacetate on the CIDNP phenomena observed during the irradiation of benzaldehyde and benzoin in deuterated cyclohexancbenzene solutions. The **CIDNP** spectra generated by photolysis of benzoin were not changed significantly by the addition of !he diazoacetate. **In** contrast, the spectra observed during the irradiation of benzaldehyde were suppressed by addition of diazoacetste **and** replaced by CIDNP signals of products formed by reaction of diazoacetate with benzaldehyde or with cyclohexane- d_{12} . These results show the energy transfer from benzaldehyde to methyl diazoacetate leading to triplet **carbornethoxycarbene121.**

Photolytic decomposition of methyl diazoacetate in the presence of carbon tetrachloride and deuterochloroform affords methyl 2,3,3,3-tetrachloropropionate and methyl 3-deutero-2,3,3-trichloropropionate, respectively¹²². The slow decay of **13**

the observed emission signals due to these polarized products suggested that the products of the reaction were being formed after irradiation ceased. Primarily on the basis of this observation, a free radical chain mechanism was forwarded for the reactions of carbomethoxycarbene with carbon tetrachloride and deuterochloroform. However, this conclusion has recently been reconsidered in the light of the radicalpair theory of CIDNP which does not allow spin polarization in a radical-induced recombination $step^{123}$. The preferred mechanism to account for the formation of the methyl propionates involved carbene attack at halogen with consequent ylide formation 124 : between the reaction were being formed

educts of the reaction were being formed

cetions of carbomethoxycarbene with carbo

wever, this conclusion has recently been

ir theory of CIDNP which does not allow

combination s

The recombination of geminate pairs formed presumably by homolysis of a chloronium ylide is the more likely mechanism. The 13 C CIDNP patterns are in accord with the cage recombination of radical pairs, but inconsistent with a radical chain mechanism¹²³.

Closs compared the photolysis of diphenyldiazomethane in toluene solution¹¹¹ and the thermal or photolytic decomposition of the azo compound *67.* Both reactions afforded the same products *68-70,* but the ratio of unsymmetrical to symmetrical products **68/(69+70)** was higher in the reaction of **67** (ratio **1.5** : **1)** than in the diphenylcarbene reaction (ratio **1** : **1).** The n.m.r. spectra obtained from the **two** reactions showcd opposite polarization in confirmation of theoretical predictions for radical pairs of different multiplicity. Since there is little doubt that acyclic azo compounds decompose through the singlet state¹²⁵, this experiment assigns triplet multiplicity to the radical pairs intervening in the diphenylcarbene reaction.

C. Spin Multiplicity and Reactivity of Methylene

A principal aim in the photochemistry of diazo compounds is to correlate thc observed chemistry with the multiplicity, singlet or triplet, of the carbenes formed. **Two** typical reactions are observed: insertion into a C-H bond and addition to an unsaturated bond :

(68) (69) (70)
\n**Itiplicity and Reactivity of Methylene**
\nal aim in the photochemistry of diazo compounds is to ce
\nemistry with the multiplicity, singlet or triplet, of the carber
\nreactions are observed: insertion into a C-H bond and add
\nbond:
\n
$$
N_2CH_2 \xrightarrow{h\nu} : CH_2 \xrightarrow{\qquad \qquad}_{R-\text{C}-H} R - \text{C}-CH_2-H
$$
 (insertion)
\n $N_2CH_2 \xrightarrow{h\nu} : CH_2 \xrightarrow{\qquad \qquad}_{C-\text{C}} C$ (addition)
\n $N_2CH_2 \xrightarrow{h\nu} : CH_2 \xrightarrow{\qquad \qquad}_{C-\text{C}} C$ (addition)
\n $N_2CH_2 \xrightarrow{h\nu} : CH_2 \xrightarrow{\qquad}_{C-\text{C}} C$ (addition)
\n $M_2CH_2 \xrightarrow{h\nu} H$
\n $N_2CH_2 \xrightarrow{\qquad H\nu} H$
\n N_2

Singlet methylenc **'CH,** gencratcd by the photolysis of diazomethane inserts into the **C-H** bonds **of** alkanes with nearly random statistical preferences. An example is provided by the liquid phase photolysis of diazomethane in 2-methylbutane¹²⁶:

Only slight prefercnces for the tertiary hydrogen (1.5 **I)** and the secondary hydrogen (1.22) appear.

One may ask whether the $CH₂$ insertion is direct or whether some intermediate, such as a radical pair, is involved. Doering and Prinzbach¹²⁷ photolysed diazomethane in the presence of 2-methyl[1-¹⁴C]propenc and studied the distribution of the radioactive label within 2-methyl-1-butene, the product of allylic $C-H$ insertion.

In the liquid phase less than 2% scrambling occurred, whereas **8%** of **72** was obtained in the vapour phase.

Studying the configurational relationship between substrate and insertion product, Franzen reported that methylene reacted with the tertiary carbon-hydrogen bonds in the *trans* diacetate (73) with inversion^{128, 129}. However, generally the insertion of

carbene has been shown to proceed with retention130. Optical activity is maintained in the reaction with the methoxycarbonylcarbene¹³¹ and methyl- $(-)(S)$ -2-methoxypropionate and silicon-hydrogen bond^{131, 132}.

Ho and Noyes had shown that the reactior. of triplet methylene with propane involved abstraction¹³³, since dimeric products, hexane and 2,3-dimethylbutane, were isolated :

 \cdot ^{LH3} + \cdot \cdot \cdot + \cdot \cdot + \cdot \cdot + \cdot \cdot + \cdot \cdot \cdot + \cdot \cdot \cdot + $\$

Average values for the relative rates of abstraction by triplet carbene are: primary/ $secondary/tertiary = 1/2/7$.

Incisive information concerning the chemical characteristics of singlet and triplet carbenes comes from study of addition reactions involving carbon-carbon double $bonds¹³⁴⁻¹³⁷$.

As in C-H insertion, singlet methylene adds stereospecifically in *a* concerted step: both bonds of the cyclopropanc are formed simultaneously since such a step

may occur with spin conservation. The additions of **'CH2** (from photolysis of diazomethane in liquid phase) to cis-2-butene and trans-2-butene yield only *cis-***¹**,Zcyclopropane and *trans-* **1,2-dimethylcyclopropane,** respectively. These additions are stereospecific; the geometrical relationship of the olefinic substituents is preserved in the products:

This behaviour has been taken as evidence that the addition of ${}^{1}CH_{2}$ to an olefin is a direct reaction and no diradical intermediate intervenes. Triplet methylene generated in the gas phase by the mercury photosensitized decomposition of ketene adds to olefins with greater discrimination: *e.g.* its addition to butadiene is *6.6* times faster than to isobutene; the corresponding ratio with singlet methylene is 3.2 . The increased preference of ${}^{3}CH_{2}$ for butadiene can be attributed to the

	~………	
Olefin	1 CH ₂ ^a	3CH_2
	0.91	0.96
	1.07	0.64
	1.00 ^c	1.00c
	0.58	0.33
	0.65	0.31
	$3 - 20$	6.64
	0.87	0.56
	0.72	0.35
	0.68	0.35

TABLE 11. Relative **rates** of mcthylene addition to Oe fins¹³⁸

a From photolysis **of** ketcne with 2600 A.

^b From mercury photosensitized reaction of kctene with 2537 A.

^CStandard.

radical character of this addition, which presumably involves a diradical intermediate

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The addition of triplet methylene to *cis-* and trans-2-butenes is an indirect, twostep reaction and takes place non-stereospecifically. Both *cis-* and trans-**1,2-dimethylcyclopropanes** are formed from eithe; starting olefin. This phenomenon, which is now believed to be diagnostic for triplet carbene additions, is most readily understood in terms of the Skell-Woodworth hypothesis.

Skell argued that the addition of a triplet carbene to an olefin would generate **a** triplet trimethylene intermediate, in which rotation about the single *C-C* bonds would be at least as rapid as thc spin inversion needed before formation of the final cyclopropane *C-C* bond could occur. The addition of **a** triplet carbene to *cis*or trans-2-butene should then yield both *cis-* and trans-I **,2-dimethylcyclopropanes.** On the other hand, addition of a singlet carbene could be stereospecific, since there would be no spin restriction on the simultaneous formation of both cyclopropane *C-C* bonds.

SCHEME 3

Dilution with a 200-fold excess of inert perfluoropropane induces some **inter**system crossing in either the diazomethane, or more likely in the methylene itself,

Methylene produced by irradiation of diazomethane in the vapour phase in the presence of high pressure inert gases gives non-stereospecific addition to 2-butene¹⁴⁰, ¹⁴¹:

Benzophenone-photosensitized decomposition of diazomethane induces a limited non-stereospecificity in the addition reaction¹⁴². The sensitized reactions should

'CH, + w - - Ph,C=O + **hi*** 'Ph,C=O 'Ph,C=O 'Ph2C=0 + N,CH, Ph,C=O + 'N,CH, JN,CH, __f 'CH, + N, **(1.9) (1** 1 'CH, + \==, -> (trace) **(mainly) SCHEME** *5*

produce triplet methylene, while direct irradiation is known to produce the singlet. In the presence of an inert gas or in inert solvents, singlet methylene may decay to the triplet before reacting with the unsaturated substrate.

A number of factors can influence the stereospecificity of the addition, notably solvent cage effects or unusual rapid intersystem crossing. In spite of some difficulties, Skell's criterion appears to work well in practice in distinguishing between singlet and triplet carbene species.

Theoretical justification has been forthcoming, through Extended Huckel Theory (EHT) calculations¹⁴³⁻¹⁴⁶.

A theoretical analysis of the addition *of* **'CH,** to ethylene suggests initiation of the reaction as in **74,** a ' π -approach' in which the vacant p orbital of the carbene begins to overlap with the π -system. As reactants move along the reaction coordinate towards product geometry, the π -approach goes over to a ' σ -approach'¹⁴³. Transfer of electron density from the olefin's π -system to the carbene's p orbital occurs;

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More recent calculations suggest that π -approach 74a is energetically preferable to π -approach **74b** in the addition of singlet methylene. Addition of CF_2 is calculated to follow **a** similar course, with less charge transfer from olefin to carbene in the transition state¹⁵⁰. Related calculations for additions of CH_2 and CF_2 to isobutene

again suggest that **74a** is preferred to **74b,** but only by a small energy difference. The results of these calculations must be treated cautiously, as a guide to thought rather than as firm conclusions.

D. Photolysis of Aromatic Diazo Compounds in Olefins

I. Phenylcarbene

matrix isolation¹⁵¹, which has been extended to 4 K ¹⁵², and of its emission spectrum¹⁵³, and by extended Hückel calculation¹⁵⁴. The ground state of phenylcarbene has been designated as triplet on the basis of

The chemical behaviour of phcnylcarbene generated by photolysis of phenyldiazomethane appears to be substantially singlet rather than triplet. Gutsche reported only about **3%** non-stereospecificity in the addition of phcnylcarbene to

9. Photochemistry of the diazonium **and diazo** groups 379

trans-2-butene and about 2.5% non-stereospecificity in its addition to cis-2-butene ¹⁵⁵. CIoss reported corresponding values **of 0.5-1%** and **3-5%156.** The high stereospecificity can be taken as support for a singlet state of the reactive intermediate¹³⁷. The results could be consistent with triplet phenylcarbene addition if there was a very fast ring closure allowing only incomplete rotamer equilibration in the intermediate triniethylene diradical. Moss and Doelling have presented evidence that the properties of triplet phenylcarbene are quite different from those of the singlet¹⁵⁷. These authors find that irradiation of a frozen solution of phenyldiazomethane in *cis-* or trans-2-butene yields not only cyclopropanes but products of abstractionrecombination reactions as well. They suggest that triplet phenylcarbene reacts largely by hydrogen abstraction.

2. Diphenylcarbene

In the photolysis of diphenyfdiazomethane in the presence of olefins the major reaction path with cis- and trans-2-butene is hydrogen abstraction from the allylic positions, followed by recombination of the radical fragments. More highly substituted olefins give even greater percentages of abstraction-recombination :

$$
Ph_{2}CN_{2} + \sqrt{}
$$

\n
$$
-10^{n}C_{1}^{h_{1}}
$$

\n
$$
Ph Ph + Ph + Ph + CHPh_{2} + \sqrt{CHPh_{2}}
$$

\n
$$
2.2\% - 7.0\% + 45.4\%
$$

With less substituted olefins cyclopropanes predominate, and with isobutene and propene only cyclopropanes result. Clearly the mode of reaction *of* diphenylcarbene (Table 13) depends intimately upon the olefin¹⁵⁸:

TABLE 13. Mode of reaction of diphenylcarbene with olefins

Closs reports that cis-2-butene gives a cis : trans ratio of either 87 : 13¹⁵⁹ or 77 : 23¹⁶⁰ and that trans-2-butene gives a *trans* : cis ratio 96 : 4¹⁵⁹. Thus they must be formed mainly from the singlet state:

There is a growing conviction that singlet and triplet diphenylcarbene are in equilibrium, thus leading to reactions of both spin states occurring at the same time:

From cis-p-deuterostyrene and diphenylcarbene roughly 65% cis and *35% lrans* adducts **were** formed1SR:

Obviously, in systems in which triplet abstraction is the primary reaction, the cyclopropanes will be formed largely from the singlet. This appears to be the case with the 2-butene.

The stereospecificity increases at lower temperatures, for example, at -10 ^oC with cis-2-butene, *cis-* and *trans-J* **,2-dimethyl-3,3-diplienylcyclopropanes** are formed in a ratio of 3.2, while the corresponding ratio with *trans*-2-butene is only 0.04 ; at -66° C the product ratio obtained from *cis*-2-butene is 9.0.

The *cis* : *trms* cyclopropane ratio can be affected by addition of hexafluorobenzene, a molecule inert to diphenylcarbene158. With *80-95%* hexafluorobenzene, there can be no more than 12% singlet reacting. Presumably unreactive collisions allow diphenylcarbene to approach equilibrium more closely and the groundstate triplet is naturally favoured. The presence of oxygen docs not change the isomer ratio although the total yield of cyclic and acyclic hydrocarbons is greatly diminished and approaches zero for high oxygen concentration.

All these observations are most consistent with a mechanistic scheme as outlined in below:

It is postulated that intersystem crossing is much faster than any other reaction. Furthermore, it is assumed that the reverse crossing is also very fast so that both electronic configurations are effectively in equilibrium.

One might well expect a triplet to be more likely to undergo 1,4-addition than a singlet, since the intermediate diradical might not be restricted to 1,2-closure. Even with triplets, however, acyclic dienes yield only 1,2-adducts^{158, 161}.

Triplet diphenylcarbene reacts with alkynes by addition and abstraction, and each of the reactions is a two-step process. With monosubstituted acetylenes, indenes are the sole products as shown. Addition of diphenylcarbene to the less substituted

end of the acetylenes generates **76,** which may or may not be initially the triplet carbene shown, but in any event it should quickly relax to yield **77.** In the reaction of diphcnylcarbene with dimethylacetylene, the main products are the cyclopropene **(79)** and the allene **(80).** Compound *80* is probably the product of hydrogen

abstraction reflecting the unavailability of an unsubstituted end of the acetylene for attack by diphenylcarbene. Howcvcr, both **78** and *79* are formed through addition to give **81.** The added methyl group causes a new effect: the most obvious diflerence between **81** and **76** involves the methyl-aromatic ortho-hydrogen interaction indicated in **81.**

Competition between cis - β -deuterostyrene and methylacetylene for the irradiation product of diphenyldiazomethane gave mainly addition to styrene and only traces of indene. This is reasonable, since styrene is a much better triplet trap than propene. Phenylacetylene competes more successfully and *25%* of the products in **a** competition reaction with styrene are derived from addition to the acetylene.

(Dimethoxyphosphory1)phenylcarbene *(84),* produced photolytically from 83, reacts with alkynes **10** give satisfactory yields of dimethyl I **-phenyl-2-cyclopropenyl**phosphonates (85)¹⁶². With monosubstituted alkynes, ring closure to dimethyl-3-indenylphosphonates (88) competes with the formation of **85.** The carbene **84**

apparently attacks the unsubstituted alkyne carbon atom with formation of the diradical **86** which yields first **87** and then **88.**

Moritani and coworkers¹⁶³ found that cyclopropanes were not formed by attack of **89** on 2-butene but that some products of abstraction-recombination could be found. The similarity between carbene **89** and diphenylcarbene may seem odd since **89** appears flat, at least on paper, and diphenylcarbene has been shown¹⁶⁴ to be bent and twisted. Carbenes **90** and **91** are even less flexible and both give products

specific fashion. Carbene **90** gives a spectrum of products which seems to implicate **a** mixture of spin states. Irradiation of diazoanthrone in cyclohexane again led to products of hydrogen abstraction, with no cyclopropane being formed167. Olefins not containing allylic hydrogen did form adducts, however¹⁶⁸.

3. Fluorenylidene

Fluorenylidene, generated by photolysis from 9-diazofluorene. unlike diphenvlcarbene, adds to π -systems without the complication of large amounts of hydrogen abstraction. Fluorenylidene adds easily to a variety of olefins¹⁶⁹. With cis-2-butene as the solvent, **cis-dimethylspirocyclopropane-9-fluorene** and its trans isomer are formed in a ratio of 1.95, while irradiation in *trans*-2-butene gives the same products in **a** ratio of 0.06. This memory effect becomes much smaller when the reaction mixture is diluted with hexafluorobenzene **as** an inert solvent. **At** the highest dilution,

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Olefin	$90\% C_{6}F_{6}$	Pure olefin
	0.37	0.66
	1.00	1.00
	$1 - 70$	0.47
	0.86	0.38
	0.40	0.43
	0.69	0.56
	9.00	3.5

TAOLE *15.* **Relative rates** of **fluorenylidene addition** *to* **olefins**

when the solution **was** approximately 0.1 M in olefin, the corrcsponding ratios were 0.25 and 0.14. These observations fit a mechanism involving stereospecific addition of **a** metastable singlet fluorenylidene and a competing addition from thc triplet ground state occurring with complete loss of stereospecificity.

It is possible selcctivcly to filter out the triplet state by allowing it to rcact with **a** triplet trap such **as** butadiene, with which the triplet seems to react particularly fast. **As** can be seen from Table 16, the more butadiene is added, the more stereospecific the addition becomes. Of course at high butadiene concentrations very little product is observed from the reaction with the 2-butcne, and precisc numbers become very difficult to obtain.

Butadiene (mmol)	cis-2-Butene	Cis/trans ratio of cyclopropane
	$10-0$	$2 \cdot 1$
3.0	60	$10-1$
$11-5$	3.0	49

TABLE 16. Reaction of fluorenylidene with cis-2-butene in **thc** presence of **I** ,3-butadiene

Fluorenylidene was found to add to cyclopcntadiene to give only **92** and to add to cyclohexadiene to give **95.** No trace of the hypothetical 1,4-adducts **93** and **96** could be found. Diphenylcarbene also gives $1,2$ -adducts with cyclopentadiene¹⁷⁰. The

reactions of fluorenylidene and cyclopentadienylidene^{171, 172} with cyclooctatetraene are normal and no 1,4-addition products are found. The further rearrangements of the initially formed adducts *97* and **98** have been studied.

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4. Cyclopropenylidene, cycloheptatrienylidene and cyclopentadienylidene

For cyclopropcnylidene *(99)* and cycloheptatrienylidene **(loo),** it was hoped that the singlet carbcne would derive important stabilization from the aromaticity expressed by the resonance hybrids which are shown below :

Important contributions by these forms would be shown by nucleophilic behaviour and, indeed, only olefins with electron-poor π -bonds have proved to be suitable substrates for these systems¹⁷³.

When tropylidcne is generated at lower temperatures, heptafulvalene is the only product. It is not captured by simple alkenes, but it does add stereospecifically to electron-poor olefins such as dimethyl fumarate or maleate, and to mizture of *para*-substituted styrenes in such a way that ρ can be deduced to be +1.05. These findings suggest that tropylidene is a nucleophilic singlet, the electron-donating character being imparted by resonance contribution form 100.

9. Photochemistry of the diazonium and diazo groups 389

The ground state of cyclopentadienylidene is known to be a triplet in which one unpaired electron is localized in plane $sp²$ orbital, while the other electron is in the π -system¹⁷⁴. The singlet state of the carbene may be expressed as the resonance hybrid 101. Recent calculations indicate that the configurations **(101a)** and **(101b)** are within 0.2 eV **175;** the resulting extensive configuration interaction will lead to mixing and the singlet is therefore best represented as the entire resonance hybrid 101.

Photolysis of diazocyclopentadiene in *trans-* and cis-4-methyl-2-pentene led to spiro[2.4]heptadienes **102** and **103,** respectively, in good yields. The addition is nearly, but not completely, stereospecific. Addition of either hexafluorobenzene or

perfluorobutane has little effect on the stereochemistry. The reacting state seems to be the singlet, although the ground state is known to be the triplet.

A variety of phenyl-substituted cyclopentadienylidenes has been examined. The properties do not seem much changed by phenyl substitution¹⁷⁶. Norbornadiene gives an initial adduct which rearranges further on photolysis¹⁷⁷. Alkynes give cyclopropenes which though isolable rearrange further¹⁷⁸.¹⁷⁹.

The photolytic addition *of* **105** to the double bond of cis-4-methyl-2-pentene proceeded **non-stereospecificallyl*o.** The properties of **106** can best be described **as** arising from the triplet state. The steric requirement of **106** is quite large and hence

might allow many non-reactive collisions to occur, and thus give the carbene time to undergo intersystem crossing. In addition, heavy atoms are known to enhance the probability of spin-forbidden transitions through coupling **of** spin and orbital angular momenta.

4,4-Dimethylcyclohexadienylidene undergoes additions similar to that of the five-membered ring compound¹⁸¹ (Table 17).

E. Photoiysis of Diazocarbonyi Compounds in Olefins

1. Intermolecular addition reactions

Diazo esters react with olefins **by** two distinct paths to produce cyclopropanes. Diazoacetic ester might lose nitrogen under the influence **of** heat or light to produce a carboalkoxycarbene which adds *to* the olefin or to the acetylene giving a cyclopropane or cyclopropene carboxylate.

PATH A

Alternatively, the diazoacetic ester might react with the olefin to form a pyrazoline which loses nitrogen to produce the cyclopropane carboxylate. In non-photochemical reactions of double or triple bonds conjugated with a carbonyl, imine or nitrile group, 1,3-dipolar addition of the diazo group usually occurs to give a pyrazoline or pyrazole which is stable under the reaction conditions.

PATH B

Carboalkoxycarbenes are involved in reactions above 100 "C and in photochemical reactions. Carboalkoxycarbenes behave as electrophilic species. With *cis-* and rrans-2-butene, photochemically produced carboalkoxycarbene adds in a stercospecifically **cis** manner. Thus, Doering and Mole1*? showed that methyl diazoacetate and cis-2-butene gave **a** mixture of two *meso* cyclopropanecarboxylic esters, whereas *trans*-2-butene gave only the corresponding racemic mixture:

9. Photochemistry of the diazoniurn and diazo groups **393**

Addition to cyclic or asymmetric double bonds gives the less hindered adduct¹⁸³⁻¹⁸⁵:

An attempt to compare singlet and triplet states of photolytically generated carboalkoxycarbenes has been made by Jones and his collaborators¹⁸⁶, ¹⁸⁷.

Direct photolysis of dimethyl diazomalonate has been shown by these investigators to afford a reactive intermediate whose chemical properties differ markedly from those of the corresponding species produced via benzophenone-sensitized photodecomposition of the same diazo ester. It is thought that the reacting species produced in the direct and sensitized photodecompositions are singlet and triplet biscarbomethoxycarbenes, respectively. The results with *cis-* and trans-4-rnethyl-2-pentene are shown in Table 18. Since the addition is only *90-92%* stereoselective, *(cis)* rather than **100%** stereospecific, it is possible that in the direct photolysis the carbene is not formed exclusively in one spin state, but predominantly in one state

TABLE 18. Product distribution in the reactions of dimethyl diazomalonate with cis-4-methyl-2-pentene and *trans-*4-methyl-2-pentene

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and to a lesser extent in the other. In an attempt to induce intersystem crossing from the singlet to the triplet, the photolysis of methyl diazomalonate in cis-4-methyl-2-pentene was studied in varying concentrations of hexafluorobenzene. Only when very large amounts of hexafluorobenzene are added does the amount of *trons*cyclopropane increase significantly. Indeed, initially, the amount of cis-cyclopropane increases s!Ight!y. **A** likely source of this increase is excited methyl diazomalonate, which could produce trans-cyclopropane by addition to the double bond followed by loss of nitrogen. **As** hexafluorobenzene is added fruitless collisions could deactivate the excited diazo compound to the ground state, producing more cis-cyclopropane until enough has been added so that all excited diazo compound has been deactivated and intersystem crossing begins.

FIGURE 14. Addition of dimethyl diazomalonate to cis-4-methyl-2-pentene in hexafluorobenzene. [Reprinted **with** permission from Jones, *J. Ainer. Chcni. Soc.,* **94,** 7469 (1972). Copyright by the American Chemical Society.]

In a recent study of the benzophenone-sensitized photolysis of dimethyl diazomalonate in the presence of cis-4-methyl-2-pentene, the ratio of products (108a)/ **(10Sb)** was larger in the presence of dimethyl sulphide than in its absence'88. Singlet carbenes generally act as electrophiles, although the bulkier ones are subject to steric hindrance. In contrast to the large amount of data reported on singlet relative rates, only fragmentary information is available for triplets. Diphenylcarbene is rcported to add to 1,3-butadiene and 1,l-diphcnylethylene 100-times faster than to cis -2-butene¹⁸⁹. This comparison is unsatisfactory since diphenylcarbene reacts with $cis-2$ -butene primarily by hydrogen abstraction, not addition¹⁹⁰. Table 19 shows that in **the** sensitized photolysis there is a three- to four-fold increase in the relative rate of addition to dienes over nionoolefins, and there is a dccrease in the relativc rates of addition to certain olefins, e.g. to 2,3-dimethyl-2-butene and to *cis*-4-methyl-2-pentene.

9. Photochemistry of the diazonium and diazo groups

Olefin	Direct	Sensitized
	0.88	0.33
	$1 - 00$	$1 - 00$
	0.47	0.46
	0.48	0.48
	0.55	0.15
	0.23	0.13
	$1-3$	$4 - 4$
		4.5

TABLE 19. Relative rates of addition of carbenes produced in direct and sensitized photolysis of dimethyl diazomalonate

Relative rate constants for addition of several carbenes to a series of allylic compounds were determined by means of competitive experiments¹⁹¹. Results summarized in Table 20 show that polar selectivities of carbenes are generally low but correlate with the polar nature of substituents in the carbenes. The selectivities of singlet carbenes were even lower than those of the corresponding triplet carbencs. These facts strongly support a mechanism of addition with singlet carbenes which does not involve any rate-determining process leading to polar intermediates, but involves transition states having slight contributions from polar structures.

				R		
Carbene precursor	Mode of decomposition	SiMe ₃	$n-Pr$	OMe	CI	CN
N,CHCOOMe	hν	2.07	1.47	1.23	$1 - 00$	0.97
N ₂ CHCOOMe	$hv + Ph2CO$	2.55	1.55	0.95	$1 - 00$	0.66
$N_2C(COOMe)_2$	hv	1.85	1.50	0.93	1.00	0.66
$N_2C(COOMe)_2$	$hv + Ph2CO$	3.66	2.15	1.25	$1 - 00$	$1 - 13$
N_2 CHPh	hv	1.13	1.06	1.00	---	0.80
N,=	hν	1.99	1.79	1.20	1.00	0.92
$NaOH + CHCl2$		80.14	14.04	3.44	1.00	0.75

TABLE 20. Competitive reaction of carbene with allylic compounds, $CH₂=CHCH₂R$

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However, Wulfman¹⁸², recently tried to establish the existence of bismethoxycarbonyl carbenoids by analysing the activation parameters for both catalytic and photolytic processes, but the data cast strong doubt on the existence of free carbenes during the cyclopropanation of olefins and during insertions into the allylic C-H bonds by the alleged singlet carbene. They are forced to conclude that a free singlet

carbene is not an intermediate during the non-sensitized photolysis of dimethyl diazomalonate. The cyclopropane and the allylic $C-H$ insertion products may be formed via the photoexcited diazo compound forming a complex with the olefin.

An alternative criterion was proposed recently by Shimizu and Nishida¹⁹³, who used **1 ,I-dicyclopropylethylene (109)** as the substrate. Here addition of a singlet carbene can proceed in the usual concerted manner to afford **110,** whereas the existence of a triplet carbene is revealed by the observation of rearranged products 111 and **112** as shown below.

9. Photochemistry of the diazonium and diazo groups 397

Unsensitized decomposition of dimethyl diazomalonate in allene gives **113 as** the only isolable product¹⁸⁶. One might have speculated that the sensitized reaction, proceeding through a triplet, would lead to trimethylenemethane **(114),** and then to **113** and **115. A** similar scrambling of label has been seen in the gas phase addition of dideuteromethylene to allene but the likely explanation for the scrambling involves hot molecule rearrangements, not spin state. The sensitized decomposition gave only **113** and 1.1-diphenylethylene in the ratio **12** : 1. The latter product probably arises via the oxetane (116). Hendrick¹⁹⁴ has observed furan formation in the

reactions of triplet biscarbomethoxycarbene with alkynes. The direct photolysis of methyl diazomalonate in 2-butyne afforded the expected cyclopropene **118** in 36% yield. Only trace amounts of other products, assumed to be insertion products, were detectable.

In sharp contrast, sensitization of the photolysis in 2-butyne with an equimolar amount of benzophenone gave as the major product the furan **119** in **43%** yield, accompanied by only 9% yield of **118.** The course of the triplet carbene reaction can be ascribed to cyclization of the diradical **120. It** is interesting to note that the substitution of bulky groups on the acetylene results in increased closure to cyclopropene but not nearly so dramatically as in the case of diphenylcarbene.

compositions of dimethyl diazomalonate in acetylenes **117a4 117 Mode** *of* decomposition **118 119**

TADLE 21. Relative yields of **118** and **119** from **de-**

In the photolyses of the diazo compound **121,** related to Meldrum's acid, direct irradiation in olefins produced very little product of any kind, while in the sensirized decomposition adducts can **be** isolated in relatively good yields. The stereochemistry of the sensitized decomposition is exactly what one would expect from a triplet intermediate. These results are summarized in Table 22¹⁸⁶.

Photolysis of ethyl bromo-, iodo- and chloro-diazoacetate affords the bromo-, iodo- and chloro-ethoxycarbenes, which add *cis* stereospecifically to olefins and insert to $C-H$ bonds of alkanes, though less readily than the carboethoxycarbene. The photosensitized decomposition of ethyl bromo- and chlorodiazoacetate in olefins leads to addition products with essentially complete *cis*stereospecificity (97%)¹⁹⁵.

Olefin	Conditions	Yield (%)	Cis $(\%)$	Trans (%)
	$h\nu$	$\overline{\mathbf{c}}$		
	$h\nu + Ph_2CO$	26	14	86
	$h\nu$	$\mathbf{2}$		
	$h\nu + Ph_2CO$	26	15	85
	$h\nu$	Very small		
	$h\nu + Ph_2CO$	17	4	96
	$h\nu$	Very small		
	$h\nu + Ph_2CO$		6	94

9. Photochemistry of the diazonium and diazo groups

TABLE *22.* **Decomposition of 121 in olefins**

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Addition of dicyanocarbene to olefins is largely but not completely stereospecific. Addition of cyclohexane as diluent evidently induces intersystem crossing since the stereochemistry of addition becomes independent of the configuration of the starting 2-butene at **100** : 1 (cyclohexane : olefin).

Addition to acetylenes gives cyclopropenes, and the carbon-hydrogen insertion reaction is common (tertiary/secondary/primary $= 12/4.6/1$) ¹⁹⁶.

2. Intramolecular reactions

Intramolecular reaction of diazoketones also give cyclic compounds. 3-Diazomethylketocyclopropenes **(122)** are the only unsaturated ketones so far where lighteffected intramolecular addition produces the tricyclo^{[1.1.1}.0^{4, 6}]pentane skeleton¹⁹⁷.

The intermediate diazoketone **(123)** obtained in the photolysis of **1** ,Zdiazine N-oxide has two competitive pathways open to it¹⁹⁹. One pathway is the intramolecular 1,3-dipolar cycloaddition resulting in the formation of benzoyl-5-phenylpyrazole **(127)** via the tautomeric 3H-pyrazole **(126);** this thermal isomerization is expected to be even faster than the known isomerization of vinyldiazomethane to pyrazoIe200. The other pathway involves photodecomposition of the diazoketone to a carbene which reacts intramolecularly to yield 2,5-diphenylfuran. Since there are competing thermal and photochemical reactions it would be expected that irradiation

with more intense light **source** or lowering the temperature during irradiation would increase the photochemical reaction, **i.e.** the conversion to 2,5-diphenylfuran.

Irradiation of pyridazinium dicyanomethylylide gave 3-(2,2-dicyanovinyl)pyrazoles **(128)** and **3-(2,2-dicyanovinyl)cyclopropenes (129) *01,** probably through a mechanism similar to that operating in the photolysis of pyridazine N-oxide. Photolysis of 3-acylpyrazolines yields an α , β -unsaturated ketocarbene, which rearranges mainly into ketene. However, intramolecular addition of the *a,p*ethylenic ketocarbene has been observed²⁰².

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F. Photolysis of Diazo Compounds i.n Aromatic Compounds

Photolysis of diazocyclopentadiene in benzene solution afforded a crystalline, rather sensitive, adduct of a spironorcaradiene structure. Thermal rearrangement of **132** !ed to **133,** while a mixture of phenylcyclopentadienes resulted from acidcatalysed rearrangement²⁰⁵. Hexafluorobenzene gives an adduct of a spiro cycloheptatriene structure²⁰⁶. Similarly, a norcaradiene-type adduct was isolated from the

photolysis of **I-diazo-4,4-dimethylcyclohexa-2,5-diene** in benzenezo7. Photolysis **of** dicyanodiazomethane in benzene gave, as the sole product in 82% yield, *7,7* dicyanonorcaradiene?O8. p-Xylene afforded two isomeric products, **135** and 136. The **14**

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photolysis of diazofluorenylidene in benzene afforded the equilibrating norcaradienecycloheptatriene system²⁰⁹. Tetraphenylcyclopentadienylidene reacts with benzene to give a product, presumably the norcaradiene, which rearranges further to **(140)**

and **(141).** The final product **(141),** which arises via ring expansion of the fivemembered ring, is somewhat unusual, since ring expansion of the seven-membered ring usually occurs21o. Spironorcaradienes **138** (in equilibrium with their valence isomers 139) can be isolated if the photolysis is carried out with a long wavelength filter $(\lambda > 360$ nm).

With pyrex filter ($\lambda > 290$ nm) the mixture 138 \rightleftharpoons 139 rearranges to benzocycloheptatrienes **(141)** and their *5H* isomers. Two possible mechanisms have been proposed for this rearrangement²¹¹; a symmetry-allowed [1,7]-sigmatropic shift (to **143), or** a di-x-methane rearrangement (to **143).** The successful isolation of **142,** provides strong evidence for the intermediacy of novel bisnorcaradiene **(143).**

7-Phenylcycloheptatriene was obtained by photolysis of phenyldiazomethane in benzene solution212. Similarly, **1,4-bis(diazornethyl)benzene** was converted into 1,4-bis(cycloheptatrienyl)benzene²¹³. Benzene also acts as an acceptor for biscarbomethoxycarbene, a report to the contrary notwithstanding²¹⁴. Photolysis of methyl diazomalonate in bcnzene gave **146** and **147** in **a** ratio of 2.7 : **1.** The photosensitized reaction gave the same two products in the ratio of 1.6 : 1. The increased amount of the phenylmalonate **(147)** is consistent with **a** mechanism involving **a** diradical which can either close to the norcaradiene related to **146** or undergo hydrogen shift to 147^{186, 215}. Photolysis of ethyl diazoacetate by means of pyrexfiltered light afforded **148** in reasonable yield and *SO-SO%* purity216, **217.** Short wavelength U.V. irradiation caused extensive isomerization of **148** to cycloheptatriene-3-, -4-carboxylate and bicyclo[3.2.0], and [2.2.1] heptadiene carboxylate²¹⁸. A Hammett treatment showed that the reaction was an electrophilic aromatic attack. The *p* value of **-0.38** suggested that not much positive charge had developed in the ring at the transition 219 .

Photolysis of 2-diazoacenaphthen-I -one in benzene gives spiro[acenaphthen-1,7'-cis-norcara-2',4'-diene]-2-one (149) which rapidly isornerizes to 2-phenylacenaphthen-1-one (150) under silver perchlorate catalysis²²⁰, and gives acenaphtho- $[1,2-b]$ benzofuran $(151, X = H)$ in the presence of dicyanobenzoquinone. Other **aromatic receptors give similar**

9. **Photochemistry of** the diazonium and diazo groups **407**

(Dimethoxyphosphoryl)(methoxycarbonyI)carbene (153) and benzene afford products that depend on the duration of the irradiation²²². If the photolysis of 152 is interrupted before the reaction is complete, the equilibrium system $154 \rightleftharpoons 155$ is produced. More prolonged irradiation leads next to a sequence of shifts of ester groups and of H atoms, producing the cycloheptatriene **157. On** still further irradiation, the photostable **bicyclo[3.2.0]heptadiene (156)** is formed from **157** by electrocyclic ring closure²²³.

G. insertion Reactions

The term 'insertion' is rescrved for reactions where cleavage of the **A-B** bond and formation of the **A-C** and **B-C** bonds occur in concert. The alternative

$$
A-B + CXY \longrightarrow \begin{matrix} X & Y \\ C & \longrightarrow & Y \\ C & \longrightarrow & A-C-B \\ A & B & Y \end{matrix}
$$

stepwise process, abstraction followed by recombination, may afford insertion type products via diradical processes:

$A \rightarrow B + C \rightarrow C$	$A \rightarrow B$	\uparrow
$A \rightarrow B$	\downarrow	
\downarrow	\downarrow	

When B is a heteroatom containing one or more unshared pairs of electrons, a third

mechanistic possibility may be operative involving intermediate ylide formation:
\n
$$
A-B + iCXY \longrightarrow A \xrightarrow{+} A \xrightarrow{+} B \xrightarrow{+} A \xrightarrow{+} A \xrightarrow{+} B
$$
\n
$$
\rightarrow A \xrightarrow{+} A \xrightarrow{+} A \xrightarrow{+} B
$$

1. C-H insertion of carbene in saturated hydrocarbons

The insertion of singlet methylene into hydrogen has been subjected to a very detailed analysis²²⁴. The results indicate that insertion can take place over a wider range of initial conditions and by a much more complex pathway than static 'minimum energy path' analysis would suggest. Thc reaction of methylene with hydrogen has also been the subject of other calculations²²⁵.

EHM calculations for concerted insertion of singlet methylene into $a C - H$ bond of methane²²⁶ indicate a nearly linear transition state, the methylene approaching the C-H bond in the direction shown in **158.** The hgdroger. is transferred from methane to methylene while the $C-C$ distance (2.5 Å) changes little; this is then followed by collapse to give ethane, with no evidence for **a** discrcte radical intermediate in the process.

MlNDO/2 calculations on the reaction of singlet and triplet methylene with methane predict²²⁷ that the singlet can approach along the axis of a $C-H$ bond of methane, either in an eclipsed **(159)** or in a staggered conformation **(160).** The triplet can also approach along the $C-H$ axis, or midway between two $C-H$ bonds of methane; thc latter approach is prcdicted to lend to insertion by thc triplet. Insertion of methylene has been analysed by Zimmerman, using his 'MO following' concept²²⁸.

Doering and Knox²²⁹ proposed the charge separation hypothesis for the rationalization of the selectivity of various substituted carbenes toward primary,

secondary and tertiary carbon-hydrogen bonds, as well as for the selectivity sequence $(CH_3OOC)_2CN_2 > CH_3OOCCHN_2 \geq CH_2N_2$.

TABLE 24. Relative insertion rates **of** carbenes into C-H bonds under photolytical conditions

Carbene precursor	Alkane	Secondary/primary	Tertiary/primary
N_2CH_2		$1-2$	1.5
N_2CH_2			$1-2$
N_2CHCF_3		$1-0$	
N_2CHCF_3			$1-3$
$N_2 =$			$7 - 3$
N_2 CHPh		$6-3$	
N_2 CHPh		8.3	
N ₂ CHCOOMe			3.1
N ₂ CHCOOMe			2.9
N_2 CHCOOMe		$2 - 3$	
$N_2C(COOMe)_2$		4.8	
$N_2C(COOMe)_2$			$13 - 1$
$N_3C(COOE1)_2$		8.4	
$N_2C(COOE1)_2$			12.5
$N_2C(COOE1)_2$			$21 - 1$

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Evidence supporting the postulated charge separation in the transition state comes from the reactivity of bridgehead hydrocarbons. Methylene reacts with bicyclo[2.2.1] heptane to give all of the possible $C-H$ insertion products in essentially statistical amounts²³⁰. The product mixtures obtained with methoxycarbonylcarbene and biscarbomethoxycarbene, however, reflected lower reactivities for the bridgehead positions. The bridgehead $C-H$ bonds of nortricyclene were found to be even less reactive toward ethoxycarbonylcarbene²³¹.

Diazo compound	Hydrocarbon	1-Position	All others	Tertiary/secondary
N_2 CHCO ₂ R		$1-0$	6.7	0.7
$N_2C(COOR)_2$		$1-0$	27.3	0.2
N_2CHCO_2R		$1-0$	6.0	$1-0$
$N_2C(COOR)_2$		$1-0$	6.7	0.7

TABLE 25. Relative rates of reaction with bicyclic hydrocarbons

Generally, the sensitized decomposition of diazomalonate affords a slightly more selective intermediate than does direct photolysis. Yields of insertion products are much lower in the sensitized decomposition. Rough composite values for the

ntermediate than does direct photolysis. Yields of insertion pro ver in the sensitized decomposition. Rough composite values TABLE 26. Relative rate of insertion of biscarbomethoxycarbene into					
	carbon-hydrogen bonds ¹⁸⁶				
Hydrocarbon	Ratio	Singlet	Triplet		
	Secondary/primary	$5-4$	6.9		
	Secondary/primary	4.8	5.2		
	Tertiary/primary	39	52		
	Tertiary/primary	13	20		

TABLE 26. Relative rate of insertion of biscarbomethoxycarbene into carbon-hydrogen bonds186

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tertiary-secondary-primary ratios of singlet and triplet biscarbomethoxycarbcnes are 15-40 : *5* : 1 and *20-50* : *6* : 1, respectively. This fits in general the data of Ring and Rabinovitch²³², ²³³ on the abstraction of hydrogen by triplet methylene, although the effects seem to be smaller. **A** hydrogen abstraction-recombination process would be expected *to* give new products derived from the radicals generated by abstraction. Such products do appear **as** shown below, with the yields of the sensitized reaction shown in brackets:

Direct irradiation produces only insertion products while the sensitized decomposition gives more of the products expected from free radical dimerization and hydrogen abstraction.

The photolysis of methyl diazoacetate in trimethylcthylsilane gave three major products²³⁴. The rates show that an α -secondary C-H bond (2.5) is more reactive than a primary α -C-H bond (1.0), which in turn is less reactive than a primary β -C-H bond (1.2). The differences may be explained by steric hindrance to carbene insertion. A competition reaction showed that the reactivities of $C-H$

$$
\begin{array}{cccc}\n\text{Me}_3\text{SiCH}_2\text{CH}_3 + \text{N}_2\text{CHCO}_2\text{Me} & \xrightarrow{\text{A}_{\text{P}}}\\
\text{Me}_2\text{SiCH}_2\text{CH}_3 + \text{Me}_3\text{SiCHCH}_3 + \text{Me}_3\text{SiCH}_2\text{CH}_2\text{CO}_2\text{Me} \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \text{CH}_2\text{CH}_2\text{CO}_2\text{Me} & \text{CH}_2\text{CH}_2\text{CO}_2\text{Me} & \overset{6\%}{\sim} \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow & \downarrow & \downarrow & \\
 & \downarrow & \downarrow &
$$

bonds towards carbomethoxycarbene insertion are higher in trimethylethylsilane than in 2,2-dimethylbutane $\frac{k(Si)}{k(C)} = 1.8$, possibly as a result of the inductive effect of the silicon substituent.

The relative rate of the carbene insertion into the primary γ -C-H bond in **trimethyl(tt-propy1)silane** and in **trimethyl(iso-buty1)silane** showed it to be less reactive than that of the primary β -C-H bond of trimethylethylsilane, probably because of the diminished effect of the silyl group.

The photolysis of mcthyl diazoacetate and diniethyl diazonialonatc in **1,1** -dimethyl-1-silacyclopcntane **(162)** gave carbene insertion products into the secondary *a.-* and P-C-H bonds as well **as** into primary C-H bonds in comparable cxtent.

The reaction of 1,1,3,3-tetramethyl-1,3-disilacyclobutane (163) gave carbene insertion into the $Si-C$ bond, as has been observed with dichlorocarbene²³⁵ as well as insertion into primary and secondary $C-H$ bonds in comparable rates.

2. C-H insertion of carbene in ethers

The insertion of singlet methylene into the **C-H** bonds of ethyl ether **was** reported **as long** ago as 1942, but the intervention **of** an ylide niechanism **was** ruled out only recently. Thus, attack of ¹⁴C-labelled methylene on ether gave only end-labelled propyl ethyl ether²³⁶:

*CH₂ + CH₃CH₂CH₂H₃
$$
\longrightarrow
$$
 *CH₃CH₂CH₂CH₂CH₂H₃ + CH₃CH₀CH₂CH₃

\n
$$
\downarrow
$$

Thus, methyl ethers often are obtained probably from ylides in decomposition of ethereal solutions of diazomethane²³⁶:

9. Photochemistry of the diagram and diazo groups
method for example, such as follows: 9. Photochemic²³⁶:
1 solutions of diagramthane²³⁶:
ROCH₂CH₃ + :CH₂ → R-
$$
\underset{CH_2}{\circ}
$$
 + $\underset{CH_2}{\circ}$ → ROCH₃ + CH₂=CH₂

The oxygen atom of aliphatic ethers has a small promoting effect on methylene insertion which is accompanied by minor side reactions which may **be** explained by an attack of methylene on oxygen. Triplet methylene abstracts liydrogen almost exclusively from the α position, giving rise to products of apparent α -insertion²³⁷⁻²³⁹: on which is accompanied by minor side reactions which may be explain
ack of methylene on oxygen. Triplet methylene abstracts liydrogen a
vely from the α position, giving rise to products of apparent α -insertion
CH₃

$$
CH_3CH_2OCH_2CH_3 + CH_2N_2 \xrightarrow{\hbar\nu} CH_3CH_2OCHCH_3 + CH_3CH_2O(CH_2)_2CH_3
$$
\n
$$
CH_3
$$
\n
$$
CH_3
$$
\n
$$
CH_3OCH_2CH_3 + CH_2N_2 \xrightarrow{\hbar\nu} CH_3OCHCH_3 + CH_3O(CH_2)_2CH_3 + CH_3CH_2OCH_2CH_3
$$
\n
$$
CH_3
$$
\n<math display="</math>

The small preference for insertion into the α -C-H bonds of ethers was rationalized in terms of polar resonance structures of the transition state.

In contrast to expectations based on this concept, the $O-CH₂-O$ hydrogens of methylal were found to be less reactive than the CH₃-O hydrogens. The reverse relation holds with 1,3-dioxolanc. Tlicsc results suggest that the conformation of ethers and acetals may be an important factor in determining the relative rates of methylene insertion²⁴⁰.

Insertion of carboalkoxycarbene into the $C-H$ bonds of cthers competes less efficiently with cleavage at the O-alkyl bond than does the insertion of methylcne. When methyl diazoacetate was photolysed 241 in the presence of diethyl ether, methyl ethoxyacetate, methyl β -ethoxybutyrate and γ -ethoxybutyrate were produced in the ratio of **4** : **4** : **1.**

hv CzH,OC,H,+N,CHCO,Me ___+ C,H,OCH,CO,Me+ C,H,0CHCH3+ C,H,OCH,CH,CH,CO,Me I C0,Me **4 4 1** *hv* C,H,OC,H,f N,C(COOMe), -> C,H,O CH (C OOMe), + C, H,CH C (CO 0 Me), (Reference **242)** CH, 39% yield **27%** yield

Cyclopentadienylidene inserts into both α -C-H and β -C-H bonds of diethyl ether, producing **164** and **165** in **27** and **2%** yields, respectively. The correction by statistical factor shows that the reaction occurred 19-times faster at an α -C-H bond than at a β -C-H bond²⁴³. Competition experiments, in which mixtures of an

ether and either cyclohexane or 2,3-dimethylbutane were allowed to react with cyclopentadienylidene, gave some indication that both resonance and inductive effects of the oxygen atom influenced the rate of the insertion reaction of the carbene. As in the other carbene reaction, cyclopentadienylidene inserts into $C-H$ bonds most favourably when a partial positive charge on the carbon atom involved can be stabilized. The inductive and resonance effects of the alkoxy group can be seen in the results shown in Table **27.**

TABLE 27. C-H insertion in the photolysis of diazocyclopentadiene in
ethers

	ethers		
Ether	α -C $-$ H insertion	β -C $-$ H insertion	γ -C $-$ H insertion
	5.9	0.32	
	$6-2$	0.55	
	$6-2$	2.3	0.9
	2.7	0.58	
	$4-2$	$1-3$	
			1.0 (standard, primary $C-H$ insertion)
		3.9 (secondary C-H insertion)	
riety of phenylcarbene precursors gave the same α -C-H insertion 4.			
PhCHN ₂ +C ₂ H ₃ OC ₂ H ₃ $\xrightarrow{h\nu}$ C ₂ H ₃ OCHCH ₃			
	λv	CH ₂ Ph 15% yield	

A variety of phenylcarbene precursors gave the same α -C-H insertion in alkyl ethers²⁴⁴.

 $PhCHN_2 + C_2H_2OC_2H_3 \xrightarrow{h\nu} C_2H_3OCHCH_3$ $PhCHN_2+PhCH_2OR \xrightarrow{h\nu} PhCHOR$ CH₂Ph 15% yield 3.9 (secondary C-H insertion)

cursors gave the same α -C-H in
 $\xrightarrow{h\nu}$ C₂H₃OCHCH₃

CH₂Ph 15% yield
 $\xrightarrow{h\nu}$ PhCHOR

CH₂Ph 20-30% yield I CH,Ph **2030%** yield

With allylic ethers, diphenylcarbene prefers insertion into the allylic C-H bond over addition to the C=C double bond by about **1.5-1-7** to 1, and with benzyl alkyl ethers, it inserts readily into the benzyl $C-H$ bond.

Photodecomposition of diphenyldiazomethane in the presence of aldehyde afforded the insertion product of diphenylcarbene into the aldehyde $C-H$ bond²⁴⁵.

H. Intramolecular **Reactions**

 γ -C-H bonds to give olefins and cyclopropanes. Alkyl and dialkylcarbenes react predominantly by self-insertion into β - and

The photolysis of diazoethane in the gas phase produced ethylene and acetylene as the major products. The ratio of ethylene to acetylene increased linearly with pressure²⁴⁶. The photolysis of $CD₃CHN₂$ showed that acetylene is formed from excited ethylene and not from CD₃CH:, directly²⁴⁷. Ethylcarbene illustrates two common reactions of alkylcarbenes, 1,2-shift and $1,3-C-H$ insertion²⁴⁸. For photolysis of CD_3CHN_2 showed that acetylene is formed
The photolysis of CD_3CHN_2 showed that acetylene is formed
ene and not from CD_3CH ; directly²⁴⁷. Ethylcarbene illustra
ctions of alkylcarbenes, 1,2-shift and

CH₃CH₂CHN₂
$$
\xrightarrow{h^{\circ}}
$$
 CH₃CH₂CH₂CH₃ $\xrightarrow{h^{\circ}}$ CH₃CH₂CH₂ $\xrightarrow{88\%}$ + 12%

Carbon- 14-labelled diazo-n-propane decomposes into nitrogen and singlet n-propylcarbene²⁴⁹, which rearranges²⁵⁰⁻²⁵² either to butene or to methylcyclopropane, both in a vibrationally excited state. The latter either decompose or are collisionally deactivated. The radical produced may recombine to give stable products. The yield of the product formed by collisional stabilization of 'hot' species should increase with increasing pressurc as indeed **was** observed while the fragmentation products (ethane, hexa-l,5-diene) should show opposite behaviour.

The 1,2-hydrogen shift in methylcarbene was subjected to detailed analysis²⁵³. In conformation **166,** the hydrogen atoms adjacent to the vacant p orbital can migrate with little or no activation, but movement of the third hydrogen is much less favourable; in cases where rotation about the carbon-carbon bond is inhibited,

for example, the axial hydrogen atom adjacent to the centre should migrate preferentially :

The 1,3-shift insertion can be formulated as follows:

The photolysis of **167** led to formation of the butadiene **(168)** arid **as** major product, **the** vinylcyclopropane **(169) 251.**

Hydrogen migration is **a** very eficient intramolecular trapping process **for** phenylmethylcarbene²⁵⁵. It was proposed that in the photolysis of 170 migration of the phenyl group takes place in the triplet carbene, whcreas hydrogen migration occurs exclusively in the singlet²⁵⁶. The diazo compounds 170 and 171 give the same ratio of stereoisomers of 1,2-diphcnylpropene **(172** and **173)** by migration of different groups: this points to the existence of **a** common intermediate. It is suggested that

this intermediate might best be formulated as the phantom singlet state **174** of α -methylstilbene. 1,2-Alkyl shift *(C-C* insertion) can also occur²⁵⁷:

Clearly, the more complicated the **alkyl** substituents in alkylcarbenes, the more products are expected from intramolecular insertion. **A** rclatively simple example is the following:

Kirmse and Grassmann²⁵⁸ found that 1,2-hydrogen shift dominates in alkylcarbenes. Some intramolecular cycloaddition may arise from decomposition of **176, but** this was slow compared to that of **175.** Saturated bicyclic systems undergo ring expansion as well **as** *C-C* insertion. Ring expansion also occurs in spiro systems and some unusual olefins have been synthesized by Kirmse and Pook²⁵⁹ and

later by Wiberg²⁶⁰. It seems remarkable that no hydrogen shift was observed:

Masamune and coworkers isolated the ring-expanded product **178** on the photolysis of **177 261.** Photolysis of *cis-* and **rrans-3-diazomethyIbicyclo[3.1** .Olhexane gave **a**

variety of products of carbon-hydrogen insertion but little ring expansion²⁶²:

Photoirradiation of the sodium salt of $2,3$ -homotropone p-toluenesulphonylhydrazone in tetrahydrofuran afforded a hydrocarbon **(180)** as the sole product in 70% yield, which is a dimer of **1,2,4,6-cycl.~octatetraene (179) 283.** The compound **180**

reacted smoothly with **N-phenyltriazoline-l,4-dione** at room temperature to afford, probably via **181,** the 1,l-adduct **182** in 65% yield. The intramolecular insertion of **(2-n-buty1phenyl)diazomethane (183)** was found to proceed with fairly indiscriminate attack on the α , β , γ and δ -C-+H bonds, both when the irradiation is effected directly or via photosensitization²⁶⁴. This is interpreted in terms of an equilibrium between the singlet **191** and the triplet **192,** equilibration being more rapid than the ensuing processes leading to intramolecular insertion (i.e. 184-187), intermolecular C-H insertion (Le. *to* form **188)** and hydrogen migration (i.e. to form **189** and **190).** Kirmse studied reactions of methoxy alkylcarbenes.

A methoxy group accelerates the shift of adjacent groups while itself being **slow to** move compared **to** hydrogen and also to alky12G5, **260.**

Substitution in the γ -position by a variety of groups X makes insertion more difficult. **The** data indicate a largely inductive effect of **X.** Stabilization of partial charge by resonance does not appear to be an important factor²⁶⁷. The small contribution of resonance suggests concerted hydrogen transfer and $C-C$ bonding in the γ -C- H insertion.

Direct irradiation of **193** and the tosylhydrazone anion **194** gave C-H insertion and vinyl group migration. Rearrangement of **195** might afford diphenyldimethyl-

butadiene268. Hydrogen shift was shown to be favoured over fluorine or Ruoroalkyl shift²⁶⁹:

CH₃
CHN₂
$$
\xrightarrow{hv} CH_2=CHCF_2C_2F_5
$$

C₂F₃CF₂

P-C-H insertion and alkyl migration predominate with alkyl(alkoxycarbonyl) carbene~?'~. In contrast *to* **p,P-dirnethyl-a-diazobutyrate,** r-butyl phenyldiazomethane prefers self-insertion to give cyclopropane rather than methyl shift²⁷¹.

Photolysis of ethyl diazoacetate, even in the vapour phase, did not produce butyrolactone. t-Butyl and t-amyl diazoacetate, however, affordcd small amounts of lactones on photolysis in cyclohexane²⁷². Diethyl diazomalonate gave both intermolecular and intramolecular reactions in the presence of thiobenzophenone²⁷³.

Several diazoamides were found to undergo intramolecular insertion into β -C-H bonds of the N-alkyl groups with formation of β -lactams^{274, 275}. This method was applied in a synthesis of methyl 6-phenylpenicillanate.

The 1,2-alkyl shift in carbenes has been used to generate highly unstable olefins as transient intermediates. a-Silylcarbene **(196)** rearranges in alcohols to the ester **(197)** containing an unstable silicon-carbon double bond which is then attacked by the solvent²⁷⁶. When trimethylsilyldiazomethane was photolysed with a high

pressure mercury lamp in various alcohols, **two** major products are obtained. Product **198** may be produced by **a** nucleophilic attack of alcohol on the silicon centre to release the diazomethane. The photolysis of silyldiazomethane in ethanol- d_1 showed that the deuterium was only introduced into the α -carbon of the ethyl group. The formation of *199* might be accounted for by 1,2-methyl migration producing a silicon--carbon double bond followed by addition of alcohol to the latter. While **Roof**
 Robush 198 may be produced by a nucleophilic attack of alcohol on

release the diazomethane. The photolysis of silyldiazomethane in

that the deuterium was only introduced into the α -carbon of the e

ation of

$$
\begin{array}{cccc}\n\text{Me}_{3}\text{SiCHN}_{2} & \xrightarrow{\text{A}\nu} & \text{Me}_{3}\text{SiCH}: \xrightarrow{\text{KOD}} & \text{Me}_{3}\text{SiOR} + \text{Me}_{2}\text{SiCHDCH}_{3} \\
& & \downarrow & & \downarrow \\
& & \text{OR} & & \text{OR} \\
& & & \text{(198)} & & \text{(199)}\n\end{array}
$$

niethoxy(trimethy1)silane is the major product in methanol, 1,2-methyl migration becomes more rapid in *t*-butanol^{277, 278}.

9. Photochemistry of thc diazonium and diazo groups

ROH	198 $(\%)$	199 $(\%)$	Total yield (%)		
MeOH	83	17	96		
E1OH	47	53	96		
<i>i</i> -PrOH	35	65	66		
t -BuOH		100	45		

TABLE 28. Photolysis of **trimethylsilyldiazomethane in** alcohols

Similarly, the photolysis of diniethylphenyl- and **dimethylbenzyl-silyldiazomethane** in methanol gave three major products, **200, 201** and **202.** Relative yields of **201** and **202** indicate that the migratory aptitude of the methyl group is larger than that of the phenyl or the benzyl group, even though statistical factors are considered. This tendency is quite different from that observed in cases when the migrating group is bound to carbon and not to silicon 277 : I initiarly, the photolysis of dimethylphenyl- and dimethylbenzyl-silyldiazomether methanol gave three major products, 200, 201 and 202. Relative yields of 1202 indicate that the migratory aptitude of the methyl group is

One	OMe	
$Me_2SICHN_2 \xrightarrow{h\nu} Me_2SICH: \xrightarrow{Me_2SICH} Me_2SiOMe + MesICH_2CH_3 + Me_2SiCH_2R$		

Photolysis of **203** in **204** gives **206** and **207,** which probably arise from the intermediate **205 278.**

1. Photolysis of Diazo Compounds in Compounds Containing Hetero Atoms

In this section we treat especially attacks by carbenes at $C-X$ bonds where X is a substituent having a lone-pair of electrons. This mechanism involves intermediate ylide formation prior to abstraction-recombination.

1. Reaction with polyhalomethanes

The reaction of diazomcthane with polyhalomethanes in the liquid phase proceeds by **a** radical chain mechanism, which is initiated by halogen abstraction. The fast migration of chlorine in the intermediate radical is thought to prevent chain transfer at an early stage, and to produce **208,** the major product from carbon tetrachloride. The high quantum yield and the inhibiting effect of radical scavengers provide further support for the chain mechanism²⁸⁰. The course of free radical reactions of
 $:CH_2+CCI_4 \longrightarrow CH_2Cl+·CCI_3$ with polyhalomethanes

of diazomethane with polyhalomethanes in the liquid pha

hain mechanism, which is initiated by halogen abstractic

nlorine in the intermediate radical is thought to prevent ch

ge, and to produce 20

reaction of dizomethane with polyhalomethanes in the liquid phase p-
dical chain mechanism, which is initiated by halogen abstraction. T
on of chlorine in the intermediate radical is thought to prevent chain
arly stage, and to produce 208, the major product from carbon tetrac
gh quantum yield and the inhibiting effect of radical sexvengers
support for the chain mechanism²⁸⁰. The course of free radical react
:
$$
CH_2 + CCl_4
$$
 → $·CH_2Cl +·CCl_5$
 $·CL_2 + CH_2N_2$ → $N_2 +·CH_2Cl_1$ → CLH_2 Cl_2
 $ClCH_2$ $Cl_2 + CH_2N_2$ → $N_2 +·CH_2$ Cl_2 → $(ClCH_2)_2$ Cl
 $ClCH_2$ $Cl_2 + CH_2N_2$ → $N_2 +·CH_2$ Cl_2 → $(ClCH_2)_2$ Cl
 $ClCH_2$ $Cl_2 + CH_2N_2$ → $N_2 +·CH_2C(CH_2Cl_2$ → $ClCH_2)_2$ Cl
 $ClCH_2$ $Cl_2 + CH_2N_2$ → $·CH_2C(CH_2Cl)_3 + N_2$
 $·CH_2C(CH_2Cl)_3 + CCl_4$ → $ClCH_2$ $Cl_2 +·CL_5$
(208)
ethane with polyhaloalkanes depends upon the reactivity of the

diazomethane with polyhaloalkanes depends upon the reactivity of the organic halide, the stability of the intermediate and on the reaction conditions. With unreactive halides, polymethylene is the dominant product (cf. 2-chloro-2-methylpropanc²⁸⁰); with halides of low reactivity (cf. bromochloromethane), mixtures of products of the general formula $CICH_2(CH_2)_2$ Br are obtained²⁸¹. The highly reactive polybromo- and polyiodo-methanes give dibromo- or diiodo-methane and related alkenes via β -elimination of bromine or iodine atoms from intermediate radicals²⁸², while the light-promoted reaction of halides of comparable reactivity give products

of attack of intermediate free radicals upon the polyhaloalkanes:
\n
$$
CBr_4 + 2 CH_2N_2 \xrightarrow{h\nu} 2 N_2 + CH_2Br_2 + Br_2C = CH_2
$$
\n
$$
HCBrCl_2 + CH_2N_2 \xrightarrow{h\nu} BrCHCH_2Cl + (CICH_2)_2CHBr + (CICH_2)_2CHCH_2Br
$$
\n
$$
Cl
$$
\n
$$
7\%
$$
\n
$$
13\%
$$
\n
$$
BrC(Cl)_2Me + CH_2N_2 \xrightarrow{h\nu} CICH_2CCHBrMe + (CICH_2)_2CBrMe + (CICH_2)_2CMe
$$
\n
$$
15\%
$$
\n
$$
BrCCI_3 + CH_2N_2 \xrightarrow{h\nu} CICH_2CCH_2F + (CICH_2)_2CBr + (CICH_2)_2CH_2H
$$
\n
$$
10\%
$$
\n
$$
20\%
$$
\n
$$
13\%
$$
\n
$$
BrCCI_3 + CH_2N_2 \xrightarrow{h\nu} CICH_2CCl_2Br + (CICH_2)_2CCH_2H + (CICH_2)_3CH_2Br
$$
\n
$$
at_4^{15-15\degree C} C 1\%
$$
\n
$$
16\%
$$
\n
$$
15\%
$$
\n
$$
16\%
$$
\n
$$
15\%
$$
\n
$$
14\%
$$
\n
$$
16\%
$$
\n
$$
15\%
$$
\n
$$
16\%
$$
\n
$$
1
$$

The following is the postulated mechanism in the reaction of diazomethane with bromodichloromethane:

$$
CH_{2}N_{2}+hv \longrightarrow :CH_{2}
$$
\n
$$
:CH_{2}+HCCI_{2}Br \longrightarrow \cdot CH_{2}Br + \cdot CHCl_{2}
$$
\n
$$
\cdot CHCI_{2}+CH_{2}N_{2} \longrightarrow HCCI_{2}CH_{2}+N_{2} \longrightarrow HCCICH_{2}Cl
$$
\n
$$
HCCICH_{2}CH_{2}H + HCCI_{2}Br \longrightarrow HCCICH_{2}CH + HCCI_{2}
$$
\n
$$
HCCICH_{2}CH + HCCI_{2}Br \longrightarrow HCCICH_{2}CH + HCCI_{2}
$$
\n
$$
Br
$$

The gas-phase reactions of methylene with methyl chloride and methylene chloride take place predominantly by chlorine and hydrogen abstraction. in the case of methyl chloride, both abstraction processes give rise to the same intermediates, methyl and chloromethyl radicals. At higher pressures the major products are

TABLE 29. Reaction products of :CH₂ with CH₃Cl

ethane, chloroethane and 1,2-dichloroethane²⁸³. These products are consistent with the following set of reactions:

 $:CH₂+CH₃Cl \longrightarrow CH₃+CH₂Cl$ *CH,+*CH, - C,H, \cdot CH₁+ \cdot CH₂CI \longrightarrow CH₃CH₂CI \cdot CH₂CI+ \cdot CH₂CI \longrightarrow CICH₂CH₂CI

The products above could also be consistent with **C-H** abstraction as the initial reaction of **:CH,.** This possibility was checked by experiments with dichloromethane where C-H abstraction would yield ethane, 1,1-dichloroethane, 1,1,2-trichloroethane, chloroethane and 1,1,2,2-tetrachloroethane in addition to 1,2-dichloroethane. Only 1,2-dichloroethane was produced in significant quantities. Further evidence for the abstraction mechanism was obtained by adding **a** small percentage of oxygen, when ethane, chloroethane and 1,2-dichloroethane were virtually eliminated from the reaction products. Mechanisms involving the formation of an ylide intermediate, however, cannot be eliminated:

$$
\begin{array}{ccccccc}\n & & & & & \\
-\text{C}-\text{C1}+\text{C} & & & & \\
-\text{C}-\text{C} & &
$$

Doering has found that the reaction of methylene with the carbon-chlorine bond of optically active 2-chlorobutane proceeds with a high degree of racemization²⁸⁴:

These experiments indicate the possibility of an ylide intermediate followed by radical cleavage and recombination. More recently, the use of CIDNP has implicated the singlet state in chlorine abstraction and the triplet state in hydrogen abstraction²⁸⁵, ²⁸⁶.

It was suggested that reactions of alkyl diazoacetates with polyhalomethanes may proceed via a free radical chain mechanism²⁶⁷. Urry and Wilt found that photolytic reaclions of methyl diazoacetate with chloroform, carbon tetrachloride and bromotrichloromethane produced the corresponding methyl polyhalopropionates²⁸⁸, 289 . **A** free radical chain mechanism was postulated on the basis of the identity of the products and by analogy to the reaction with diazomethane. Recently, arguments

N,CHCO,R :CHCO,R :CHCO,R+CCI,-X ---+ ~CI,+-CHCO,R **I X** *CCI,+N,CHCO,R - CI,C~HCO,R+N, CI,C~HCO,R ---+ CI,~CHCICO,R CI,~CHCICO,R+CI,C-X - *CCI,+CI,CXCHCICO,R (X = *CI,* Br **or** H)

against the operation of **a** radical chain mechanism in the photolytic reaction of diazoacetate with carbon tetrachloride, chloroform and methylene chloride have appeared. When ethyl diazoacetate was decomposed in each of these substratcs in the presence of radical chain initiators (such as benzoyl peroxide), different products were obtained from those reported earlier by Urry and Wilt. These results suggest²⁹⁰ that the photochemical decomposition of ethyl diazoacetate affords a different species than that produced when the diazo ester is decomposed in radical fashion. The preferred mechanism to account for the formation of the methyl propionates involves carbene attack at halogen with consequent ylide formation, followed by S_Ni type displacement of the halogen involved in ylide formation:

The pliotolysis of ethyl trimcthylsilyldiazoacetatc in polyhalomethanes afforded ethyl acrylate derivatives?", considered to be formed froni **a** chloronium ylicie intermediate followed by cleavage and recombination.

9. Photochemistry of the diazonium and diazo groups

Diazo ester	Halide	Products	Yield $(\%)$
N_2 CHCO ₂ Et	CCI,	$CCl_3CHCICO_2Et$	42
	CHCI,	CCI2HCHCICO2Et	58
		CICH ₂ CO ₂ Et	$15 - 2$
	CH ₂ Cl ₂	CCIH ₂ CHCICO ₂ Et	42.4
		$CICH_2CO_2Et$	3.7
$N_2C(CO_2Me)_2$	$\overline{\text{CCl}_4}$	$CCl_3CCl(CO_2Me)_2$	49.8
		$(MeO_2C)_2C=C(CO_2Me)_2$	$13-2$
	CHCI,	$CHCI2CCI(CO2Me)2$	43.3
		$(MeO2C)2C = C(CO2Me)2$	8·6
		$H_2C(CO_2Me)_2$, HCIC $(CO_2Me)_2$	trace
	CH_2Cl_2	$CH2ClCCl(CO2Me)2$	44.5
		$(MeO2C)2C=C(CO2Me)2$	4.6
		$H_2C(CO_2Me)_2$	4.9
		HCIC(CO,Me),	$5-1$

TARLE 30. The photolysis of diazo ester in polyhalomethanes

CIDNP studies have confirmed the cage recombination of radical pairs from the decomposition of halonium ylide intermediate, but are inconsistent with radical chain mechanism²⁹²⁻²⁹⁴.

Photolysis of diethyl mercurybisdiazoacetate with chloroalkanes may yield chlorocarboethoxycarbene $\left(CI - C - CO₂Et\right)$ by chlorine abstraction, with the resulting products then arising from reaction of either carboethoxycarbene **or** chlorocarboethoxycarbene with the chloroalkane²⁹⁵. However, the marked differences

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in reaction products from diethyl mercurybisdiazoacetate as compared with those from ethyl diazoacetate give strong support for the absence of carboethoxycarbene in the photolysis of the mercurybisdiazoacetate. The results suggest that carbyne **209** may be produced from mercurybisdiazoacetate in yields from **63** to **83%:**

By analogy with the suggested mechanism for the carbon-halogen insertion reaction of carbenes, one can speculate about a novel ylide-radical intermediate **(210)** in the reaction of **209** with chloroalkanes:

 $Hg(N_zCCO_2Et)$, \longrightarrow 2 N₂+Hg+2 \cdot CCO₂Et **(209)** $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ \blacksquare (210) cl

uggested mechanism for the carbon-

speculate about a novel ylide-radio

ith chloroalkanes:
 $g(N_2CCO_2Et)_2 \xrightarrow{\hbar\nu} 2 N_2 + Hg + 2 \cdot C$

(20
 $-c-CI + 209 \xrightarrow{\hbar} -c-CI - \bar{C}I - \bar{C} - C$

(210)

Halogen abstraction and recombination of the radicals thus produced appears to be the major path of diarylcarbene reactions with polyhalomethanes²⁰⁶, 297. On the other hand, ylide formation plays an important part in the formation of

diphenoquinones by photolysis of 211 in the presence of halogen compounds²⁹⁸. The isolation of the bromonium salt **212** lends support to the ylide intermediate which **is** converted into the diphenoquinone **213** by triethylamine.

The reaction between singlet diphenylcarbene and bromotrichloromethane could also lead to the formation of the transition state **214** which could then eliminate bromine chloride to give **215.**

Generally, alkyl halides appear to react whenever possible with carboalkoxycarbene to give **olefins** via **HX** elimination, when there is a P-carbon-hydrogen bond present as indicated below:

In neopentyl bromide, carbon-bromine insertion predominates over carbonhydrogen insertion by a factor of 4. It is especially interesting to note that no rearranged products were observed in this reaction²⁹⁹:

1. The image is a product of the following matrices:

\n
$$
Me_{3}CCH_{2}Br + N_{2}CHCO_{2}Et
$$
\n
$$
Me_{3}CCH_{2}HCO_{2}Et
$$
\n
$$
O_{2}F1
$$
\n<math display="</p>

This argues against the operation of a mechanism whereby an intermediate ylide breaks down to form the carbon-bromine insertion product via dipolar abstraction of bromine with subsequent recombination of ionic fragments. The reaction is therefore envisaged as an initial attack of singlet carbene on the halogen atom to give an ylide intermediate 216 followed by β -elimination. Insertion into the C-Cl

bond *of* chlorotetramethylsilane by singlet biscarbornethoxycarbene or carboethoxy-

\n Carbene may proceed via a chloronium ylide which undergoes homolytic fission at the original C–Cl bond and recombination³⁰⁰:\n

\n\n
$$
Me_3SiCH_2Cl + N_2C(COOMe)_2 \xrightarrow{h\nu} Me_3SiCH_2Cl(COOMe)_2
$$
\n

\n\n $Me_3SiCH_2Cl + N_2CHCO_2Et \xrightarrow{h\nu} Me_3SiCH_2CHClCO_2Et + Me_3SiCH_2ClCO_2Et$ \n

\n\n $CH_2CH_2CO_2Et$ \n

\n\n 24% \n

\n\n 8% \n

A chloronium ylide intermediate has recently been suggested to account for the reactions of carboethoxycarbene and biscarbomethoxycarbene with allylic halides. Direct photolysis of dimethyl diazomalonate yields singlet biscarbomethoxycarbene which reacts with allyl chloride to give 53% of allyl chloromalonate and **23%** of the cyclopropane derivative³⁰¹. With γ -methylallyl chloride the formation of α -methylallyl chloromalonate and the cyclopropane was observed, but no γ -methylally1 chloromalonate was observed. The formation of 'insertion' products may **be** explained by the formation of a halonium ylide followed by a symmetry allowed, thermal [2,3]sigmatropic rearrangement³⁰². With allyl bromide, an excellent yield

CH,CH=CHCH,CI

$$
N_{2}C(COOME)_{2}
$$
\n
$$
N_{2}C(COOME)_{2}
$$
\n
$$
H C f C C C C O C C P_{2}
$$
\n
$$
H C f C C C C C O C P_{2}
$$
\n
$$
H C f C C C C C O C P_{2}
$$
\n
$$
C H_{2} = C H C H C C I (C O O Me)_{2}
$$
\n
$$
C H_{3}
$$
\n
$$
H H_{4}
$$
\n
$$
H H_{5}
$$
\n
$$
H H_{5}
$$
\n
$$
H H_{6}
$$
\n
$$
H H_{7}
$$
\n
$$
H H_{8}
$$
\n
$$
H H_{9}
$$
\n
$$
H H_{10}
$$
\n
$$
H H_{11}
$$
\n
$$
38\%
$$

was obtained by the insertion of biscarbomethoxycarbene into the carbon-bromine bond, whereas only a trace of addition product was formed (Table **31).**

The most marked change between direct and sensitized photolysis is in the ratio of the insertion and addition products.

~ ~~ ~

TABLE **3 1.** The photolysis of diazocarbonyl compounds in allvl halides

Dimethyl diazomalonate.

 S_{Et}

 \curvearrowleft

Ethyl diazoacetate.

Ethyl **trimethylsilyldiazoacetate.**

Benzophenone sensitized decomposition.

The proposed mechanism for the formation of insertion products through halonium ylide intermediates may be supported by the following evidence. In direct photolysis, the insertion reaction with ally1 chloride was about **1-2.5** times and with the bromide about **8** times faster than the addition. Since the reactivities of the double bonds toward the attacking carbene species are not considered to be very different in these two allylic substrates, the change in the ratio of insertion to addition may be due to the difference in nucleophilicity between chlorine and bromine atoms. This consideration is also supported by results obtained in competition experiments using pairs of allylic compounds (Table **32).** From these

 $1 - 2$ **6.3**

1 .o *0.5*

Addition Insertion

results it was concluded that singlet carboalkoxycarbene produced via direct photolysis of diazo ester preferentially attacks the halogen atom in allylic halides to afford predominantly **C-X** bond insertion products, whereas the corresponding triplet carbenes generated by benzophenone-sensitized photolysis preferentially add to the carbon-carbon double bonds in these substrates.

Support for this interpretation of the differing behaviour of the carboalkoxycarbenes produced via direct photolyses *us* sensitized photolyses of diazo ester was obtained by the observation of a solvent effect on the insertion/addition ratio in the photolytic reaction of dimethyl diazomalonate. It can immediately be seen that the ratio of the insertion to addition products decreases with dilution with methylene halides or cyclohexane 303 . This was interpreted as indicating collisioninduccd conversion **of** singlet to triplet. In the heavily diluted runs the yields are severely reduced, and the scavenging by methylene halide leaves **a** mixture of carbenes richer in the triplet.

Solvent	Insertion/Addition	Yield $(\%)$	
None	2.35	76	
$None + Ph2CO$	0.056	93	
10% C_6H_{12}	3.39	64	
50%	2.52	26	
90%	1.95	5	
10% CH ₂ Cl ₂	2.38	60	
50%	$2 - 20$	35	
90%	$1 - 17$	10	
50% CH ₂ Br ₂	1.60	14	
90%	0.06	4	
10% CH ₂ I ₂	1.94	27	

TABLE 33. The effect of added solvents on the photodecomposition of dimethyl diazomalonate in ally1 chloride

2. Reaction with nitrogen compounds

Insertion into carbon-hydrogen bonds is the predominant reaction of methylene304 and of ethoxycarbonylcarbene^{305, 306} with triethylamine. Products possibly resulting from β -elimination within the hypothetical nitrogen ylide are formed only to a minor extent:

$$
Et_{3}N + N_{2}CH_{2} \xrightarrow{h_{1}} Et_{2}NMe + Et_{2}NCHMe_{2}
$$
\n
$$
Et_{3}N + N_{2}CHCO_{2}Et \xrightarrow{h_{1}} Et_{2}NCHCH_{2}CO_{2}Et + Et_{2}N(CH_{2})_{3}CO_{2}Et
$$
\n
$$
45\%
$$
\n
$$
Et_{3}N + N_{2}C(COOMe)_{2} \xrightarrow{h_{1}} Et_{2} - N \xrightarrow{+} CH_{2}CH_{2} \xrightarrow{+} Et_{2}NC(COOMe)_{2}
$$
\n
$$
(MeOOC)_{2}C \xrightarrow{+} H + CH_{2}CH_{2}
$$
\n
$$
H + CH_{2} = CH_{2}
$$

Benzyldimethylamine reacts with ethyl diazoacetate and with diazofluorene by apparent insertion of the carbene into the N-benzyl bond³⁰⁷:

Most likely these products arise from a Stevens rearrangement of the ylide intermediate.

Relatively little is known about the isolation of stable nitrogen ylides in these reactions. Substituted pyridinium **tetraphenylcyclopentadienylides** were produced in almost theoretical yields on refluxing tetraphenyldiazocyclopentadiene in
pyridine, the picolines or 2,6-lutidine under an atmosphere of nitrogen for
10 min $308-311$:
Ph
Ph
Ph
Ph
Ph
Ph
Ph
Ph pyridine, the picolines or 2,6-lutidine under an atmosphere **of** nitrogen **for**

Cyclopentadienylides of other hetero atoms were similarly_x readily obtainable by decomposition of diazo compounds with reagents containing thcse elements.

The **triphenyldiazocyclopentadiene was** irradiated under nitrogen in the 2,5-dimethylpyridine with a high-pressure mercury lamp through a pyrex filter. Recrystallization afforded the 2,5-dimethylpyridiniumtriphenylcyclopentadienylide **in** 48% yield312:

Pyridinium and isoquinolinium biscarbomethoxymethylides could be isolated from the photodecomposition of dimethyl diazomalonate in pyridine and isoquinoline³¹³. Isoquinolinium carboethoxymethylide has also been obtained by photodecomposition of ethyl diazoacetate in the presence of isoquinoline³¹⁴.

Isoquinolium dialkoxycarbonyl ylides (217) and dimethyl acetylene dicarboxylate react in methanol to give the $1,10-b$ -dihydropyrrole $[2,1-a]$ isoquinolines $(217a)$;

9. Photochemistry of the diazonium and diazo **groups 435**

217 in methanol alone yields the more reactive monoalkoxycarbonyl ylide and dialkyl carbonate, the former dimerizes to **217b** or reacts *in siru* with olefins to yield **1,2,3,10-b-tetrahydropyrroIo[2,1-a]isoquinolines (217c) 316.** The isolation of stable pyridinium ylides suggests also the intervention of nitrogen ylides in reactions in which one **group** bound to nitrogen **is** unsaturated. Here the ylide cannot be isolated but rearranges *in situ³¹⁷*. Very similar results were reported for the reaction of carbomethoxycarbene with ally1 chloride and ally1 sulphides:

virgin territory. In one such study, the reaction of carboethoxycarbene with f-butylarnine has been reported to afford the nitrogen-hydrogen bond insertion product along with a product resulting from WolfT rearrangement (see Section **J)** of the carbene³¹⁸:

> N_2 CHCGOEt \longrightarrow CHCOOEt $\stackrel{t\cdot\text{BuNH}}{\longrightarrow}$ *t*-BuNHCH₂COOEt **Wolff renrnrriurrnent** II *0* EtOCH=C=O I f-BUNH-C-CH,OEt

The photolysis of a hydrazine or **a** hydrazone with w-diazoacetophenone in **dry** tetrahydrofuran yields the corresponding phenylacctylhydrazine or phenylacetylhydrazone in fairly good yields³¹⁹, probably via Wolff rearrangement. The benzhydrylamines produced from diphenyldiazornethane were accompanied by tetraphenylethane suggesting hydrogen abstraction as another possibility³²⁰:

The photolysis of ethyl diazoacctate and of ethyl diazomalonate in acetonitrile gave oxazoles. It seems likely that the carboethoxycarbene formed by decomposition **of** the diazo ester undergoes 1,3-dipolar addition to the nitrile to form the oxazole **219** directly. The photolysis of ethyl diazoacetate in the presence of benzophenone gave **an** oxazoline incorporating the dements **of** carboethoxymethylcne, benzophenone and acetonitrile, presumably by the addition of triplet carboethoxyrnethylene **IS**

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 $R-C-CO_2Et$ \xrightarrow{hv} $R-C-CO_2$
 $\xrightarrow{N_2}$ ii
N, \downarrow OEt **(219)**

to benzophenone and further addition of the diradical intermediate **220** to acetonitrile³²¹.

Ph I Ph
 $-\frac{1}{C}$ -O1 $\frac{MeC-N}{C}$
 \leftarrow Ph₁O

HC1

CO₂Et EtO₂C
 \leftarrow N²
Me . I C0,Et **(220)** $HCCO_2Et + Ph_2C=O \xrightarrow{hr} \xrightarrow{Ph-C-O1} \xrightarrow{MeC-N}$

Irradiation of diazoacetate **(221)** in acetonitrile gave the oxazole **222** initially which in the presence of benzophenone rearranges to the isoxazole **224.** The azirine **223** has been proposed as the intermediate but could not be detected. When molecular oxygen, **a** triplet quencher, is introduced into the reaction vessel, the conversion of **222** to the isoxazole **224** is completely inhibited. The photolysis of trifluoroacetyldiazoacctate in acetonilrile gives **225** in *50%* yield. The photolysis of thc latter

affords **a 10%** yield of the dimeric species **226** which is thought to arise from a formal $[2+2]$ cycloaddition of two oxazole rings. Weygand and coworkers have

developed **a** general procedure for preparing **P-perfluoroalkylalanines** from perfluoroalkyloxazole synthesized **by** the 1,3-dipolar addition of perfluoroalkyl diazoacetic esters *to* acetonitrile. Over-all yields of **227** are *35-50%* **322.**

9. Photochcmistry of thc diazoniurn and diazo **groups 437**

$$
(R_{f}CO)_{2}O + N_{2}CHCO_{2}Et \longrightarrow R_{f}CO_{C}CCO_{2}Et \xrightarrow{MeC \to sN} N_{e}CO_{f}CO_{f}E1
$$
\n
$$
\xrightarrow[N_{f}]{H_{2}/P1O_{r}} R_{f}CH_{2}CHCO_{2}Et \xrightarrow{HC} R_{f}CH_{2}CHCO_{2}H
$$
\n
$$
\xrightarrow[NH_{2}]{H_{2}/P1O_{r}} R_{f}CH_{2}CHCO_{2}Et \xrightarrow{HC} R_{f}CH_{2}CHCO_{2}H
$$
\n
$$
R_{f} = CF_{3}, C_{2}F_{5}, C_{3}F, \qquad (227)
$$

Diphenyldiazomethane and phenylisocyanate react under influence of u.v. light to afford 2,2-diphenylindoxyl³²³, presumably via diphenylcarbene addition to the $C=N$ bond and rearrangement of the α -lactam intermediate:

Irradiation of N-sulphurylaniline and diphenyldiazomethane **gave** N-diphenylmethyleneaniline as the major product³²⁴. There is evidence that $S=O$ was evolved:

$$
Ph
$$
\n
$$
Ph
$$
\n
$$
Ph
$$
\n
$$
Sh
$$
\n
$$
Ph
$$
\n
$$
Sh
$$
\n
$$
Ch
$$
\n
$$
Ph
$$

3. Reaction with oxygen compounds

a. *Molecular oxygen*. The photolysis of diphenyldiazomethane in the presence of oxygen afforded benzophenone, which became the major product under suitable conditions³²⁵. The isolation of the cyclic peroxide from the photooxidation of diphenyldiazomethane suggests a carbonyl oxide as the intermediate³²⁶.

Experiments with ${}^{18}O_2$ indicate that no decomposition of 230 to give oxygen and benzophenone occurred under the conditions of the photooxidation. The processes leading to benzophenone appear to proceed in a parallel fashion rathcr than consecutively³²⁷. When the photooxidation was carried out in the presence of

acetaldehyde, propionaldehyde or benzaldehydc, the ozonides of **1 ,I** -diphenylpropene, 1,1-diphenyl-1-butene or triphenylethylene, respectively, could be isolated³²⁸. In the last case, no diperoxide was formed and the major product was benzophenone. The ozonide is presumably formed from reaction of the carbonyl oxide and the aldehyde, which reaction competes successfully with diperoxide

formation. Results to date indicate that ozonide formation is limited to diazo compounds containing two aryl substituents.

Some saturated hydrocarbons are oxidizcd at room temperature when carbenes are generated in them in the presence of oxygen³²⁹. The reaction seems to involve

Diazo compound	Irradiation time (min)	Cyclohexanol $\binom{6}{6}$	Cyclohexanone $\left(\frac{9}{6}\right)$
N,	300	29	$19(55)^a$
None	300		
Ph ₂ CN ₂	88	21	$13(81)^a$
$N2C(COOE1)2$	300		
	300		

TABLE 34. The oxidation products obtained **from** the photodecomposition of various diazo compounds in cyclohexanc saturated with oxygen

^a Yield of ketone from diazo compound by gas chromatography.

photolysis to give a triplet carbene which reacts with oxygen to give **a** diradical of the type **228.** The carbonyl oxide diradical then abstracts hydrogen from the alkane to give an alkyl radical and $R_2\overset{\bullet}{C}$ -O -OH; the alkyl radical may react with 228, RCOOH or oxygen to give eventually alcohols and carbonyl compounds by steps

Triplet carbenes react with oxygen more readily than singlet carbenes, apparently yielding a triplet carbene-oxygen adduct. This method has been extended to cases where diazo compounds can be converted to ozonides by allowing them to react with singlet oxygen in the presence of an aldehyde³³⁰. This reaction is observed with both aryl and alkyl diazo compounds. A possible mechanism involves attack by the electrophilic singlet oxygen at an electron-rich centre of the diazo compound. The resulting intermediate **231** may decompose directly to give nitrogen and the zwitterion **229** or it may cyclize **first** to the heterocycle **232.**

The photochemical reaction of diazo ketones **233** with molecular oxygen in aromatic solvents gave tetraphenylglycolide **(2343)** and **3,6-dimethyI-3,6-diphenyl**glycolide (234b), respectively, instead of the peroxide (235) or diketone (236)³³¹.

By contrast, diazo ketones **(237)** afforded products resulting from 1,2-hydrogen shift and 1,3-dipolar additicn of the keto-carbene intermediate.

b. *Reaction with alcohols*. When diazomethane and *t*-butanol mixtures were photolysed, the major products were t-butyl mcthyl ether and 2-methylbutan-2-ol³³². From the amounts formed, the $O-H$ bond of *t*-butanol is 10.9 times as reactive as a $C-H$ bond. Competitive reactions with roughly equimolar mixtures of alcohols gave the following relative reactivities of $O-H$ bonds toward attack by methylene :

ROH+CH₂N₂ ^{*n*_µ} → ROCH₃+C-H insertion products

Reactivities of RO-H bonds for :CH,:

MeO-H EtO-H **i-Pro-H** 1-BuO--H 2.01 **1 a95 1.37** 1.00

This is the ordcr of decreasing acidity of the alcohols as well as the order **of** increasing steric hindrance. Photolysis **of** diphenyldiazomethane in methanol afforded benzhydryl methyl ether in 70% yield³³³. The quantum yields of the photolyses in methanol and in nprotic solvents wcrc identical within experimental error.334

Photolysis of diphenyldiazomethane in **a** methanolic solution of lithium azide produced benzhydryl methyl ether and benzhydryl azide in virtually the samc ratio as that obtained in the solvolysis of benzhydryl chloride. These results suggested a diphenylcarbonium ion intermediate, arising by protonation of diphenylcarbene:

Ph,CHOMe M *c 03~* Ph,CN, + Ph,C: - > Ph,CH+ *I1* **I'** McOH * Ph,CHN,

From studies of isotope effects it has bcen deduced that diphenylcarbene reacts with the oxygen atom of ROH, giving an ylide, subsequent rearrangement of which gives Ph₂CHOR. The kinetics were rather complex^{355, 336}. With *i*-propanol, hydrogen abstraction to give diphenylmethane and acetone competed with formation of benzhydryl i-propyl ether. If hydrogen abstraction and autooxidation are attributed to triplet carbene, the kinetics require the reversible interconvcrsion of the singlet and triplet states, indicating a delicate balance of mechanisms.

$$
Ar_{r}CN_{r} \xrightarrow{hr} Ar_{r}C: \xrightarrow{ROH} \xrightarrow{R \rightarrow C} \xrightarrow{1}_{r} Ar_{r}CHOR
$$
\n
$$
Ar_{r}CN_{r} \xrightarrow{hr} Ar_{r}C: \xrightarrow{ROH} \xrightarrow{R \rightarrow C} Ar_{r}CHOR
$$

The reaction of ally1 alcohol with biscarbomethoxycarbenc produced by direct photolysis of dirncthyl diazomalonate afforded two principal products, that of 0-H insertion **(238)** and bicyclo[3.1 .O]lactonc **(239).** The product **239** is presumably formed by lactonization with elimination of methanol from the initially formed cyclopropylcarbinol arising by addition of the carbene to the double bond337. **338.**

	Direct $(\%)$		
Alcohol	(238)	(239)	Sensitized $(\%)$ (239)
OН	24.5	$31 - 9$	38.7
HO.	14.9	$33 - 2$	$19 - 7$
OН	34.1	21.9	39.4
ЮН	$11-5$	69.2	7.2
ЮH	21.2	$11-8$	$50-1$

TABLE 35. Photolysis of dimethyl diazomalonate in allylic alcohols

9. Photochemistry of **the** diazonium **and diazo groups 441**

Photolysis **of** ethyl diazoacetate in allylic alcohols gave similar products, **240** and **241,** together with **trans-cyclopropylcarbinol (242)** and Wolff rearrangement product (243). *trans-Cyclopropylcarbinol* cannot be converted to bicyclo^{[3.1.0]-} lactone. The most marked difference between direct and sensitized photolysis is in

$$
R^1R^2C = CR^3CR_1^4OH + N_2CHCO_2Et
$$

$$
R^1R^2C = CR^3R_1^4OCH_2CO_2Et
$$

(240)

the relative ratio **of** the products **of** insertion and addition. The benzophenonesensitized photolysis of dimethyl diazomalonate gave bicyclo[3.1 .O]lactone, but no 0-H insertion product. Similarly, the sensitized photodecomposition of ethyl diazoacetate in allylic alcohols gave mainly the carbon-carbon double-bond addition products.

The most probable mechanism of the $O-H$ insertion involves an electrophilic singlet carbene attack on the oxygen atom of the alcohol to give an oxygen ylide followed by the proton migration from oxygen to carbon. Distribution of the products depends on the structure of the allylic alcohols and seems to be controlled mainly by steric factors.

The photolysis of dimethyl diazomalonate in alcohols produced substantial amounts of dimethyl malonate, and the efficiency of hydrogen abstraction varies with the hydrogen donor properties of the alcohol³³⁹.

n₂C(COOMe)₂ + Me₂CHOH - ^h^b Me₂CHOCH(COOMe)₂ **²¹**% C0,Me $+$ MeOCH $+$ CH₂(COOMe)₂ + Me₂C-CMe₃
+ MeOCH + CH₂(COOMe)₂ + Me₂C-CMe₃ C0,C H MeZ **41** % HO **OH 12%**

The formation of pinacol in *i*-propanol fits a scheme in which the triplet abstracts hydrogen to give **a** radical pair followed by another hydrogen abstraction and radical recombination. Both the photosensitized and **benzoyl-peroxide-induced** decomposition of dimethyl diazomalonate in ethanol and in i-propanol gave dimethyl malonate in 74-80% yields. $N_2C(COOMe)_2 + RR'CHOH + Ph_2CO \xrightarrow{A\gamma} CH_2(COOMe)_2$

$$
N_{2}C(COOMe)_{2} + RR'CHOH + Ph_{2}CO \xrightarrow{\Lambda_{\nu}} CH_{2}(COOMe)_{2}
$$

c. *Reaction with ethers.* The major reaction of methylene with ethers is nearly random insertion into the various $C-H$ bonds. Two minor reactions are displacement, leading to methyl ethers, and apparent insertion into the **C-0** bond. The displacement reaction with methyl alkyl ethers resuits in the formation of dimethyl ether and an olefin²³⁹. It is not suppressed by the presence of oxygen nor does it occur when the methylene is produced by mercury photosensitization. The reaction is therefore envisaged as resulting from the initial attack of singlet methylene on the oxygen atom to give an ylide intermediate, followed by β -elimination.

ROCH₂CH₂R' + CH₂N₂
$$
\xrightarrow{h_V}
$$
 $\xrightarrow{R \rightarrow O \atop O \rightarrow C} CH_2$

\nICHR'

\nCH₂CHR'

\nCH₂ $\xrightarrow{h_V}$ $\xrightarrow{2}$ $\xrightarrow{P \atop O} CH_2$ \xrightarrow{H}

\nRCH₂CHR'

\nCH₂ \xrightarrow{H} \xrightarrow{H}

TABLE 36. The relative amounts **of** displacement and C-0 insertion of methylene with ethers

Ethers	Phase	Displacement $(\%)$	C —O Insertion $\binom{9}{0}$
MeO M eO.	Gas Liquid Gas Liquid Gas	4.0 0 3.2 0 $10-0$	0 0 0
MeO OMe MeCH OMe	Liquid Gas Liquid	5.6 3.9 1.6	0 4.6 $11-3$

Photolysis of ethyl diazoacetatc in dicthyl ether Icd to **244** and **245 as** the principal products 3^{310} . The product 244 in the observation of CIDNP emission can be rationalized in tcrrns of oxygen ylide intermediate which then undergoes Stevens rearrangements via a homolysis-recombination mechanism³⁴¹. Formation of a different ylide intermediate followed by Hoffman type β -elimination of ethylene by **a** non-radical path is a reasonable explanation for the formation of **245.** Many substitutcd anisols and silyl-substituted methyl ethers gave rise to methyl elimination products when treated with ethyl diazoacetate and diphenyldiazomethane. These reactions may also bc rationalizcd by **an** oxygen ylidc intermediate from which,

$$
442
$$

Insertion into the $C-O$ bond is the predominant reaction of ethyl trimethylsilyldiazoacetate²⁹¹ and dimethyl diazomalonate¹⁸⁸ with dimethyl ether. The insertion reaction probably involves the formation of an oxygen ylide intermediate. However, the sensitized reaction produces no such products. Apparently complexes of the type 246 either are not formed or do not rearrange to products.

Similarly, $C-O$ bond insertion was observed in the reaction with carboethoxycarbene with cyclic ethers. The reaction of ethyl diazoacetate with 2-phenyloxirane gave an oxetane, styrene and glyoxalate³⁴³. 2-Phenyloxetane produced a mixture of *cis-* and **rrans-2-ethoxycarbonyl-3-phenyltetrahydrofurans3".**

Addition of biscarbomethoxycarbene to the C=C double bond of allyl ethers **was** accompanied by apparent 0-ally1 bond insertion products which can be considered to form through the intermediate oxygen ylides by allylic rearrangement 317 . The bulky 1-butyl group **on** the oxygen atom seems to interfere with the formation of the ylide. The corresponding reaction with ethyl trimethylsilyldiazoacetate²⁹¹ results in

Allyl ethers	Mode of decomposition Insertion $\binom{9}{0}$		Addition $(\%)$
OMe.	$h\nu$	31	20
OMe	$\hbar\nu$	37	17
,OEt	$h\nu$	15	38
.OEt	$h\nu$	16	25
.OEt	$h\nu$ + Ph ₂ CO	trace	52
OEt	hν	15	38
OEt	$h\nu$ + Ph ₂ CO	trace	62
OBu-t	hv	$\bf{0}$	39

TADLE 37. Yield of products from the photolysis of dimethyl diazomalonate in **allyl** ethers -~ ~ ~ ~ ~~~~~~ ~ ~

the formation of a **C-0** insertion product via [2,3]-sigmatropic rearrangement of the oxonium ylide and of addition products. However, no *C-0* insertion product was observed when diphenyl diazomethane was photolysed in allyl ether.

4. Reaction with sulphur compounds

Insertion of carbencs into the **S-H** bond of thiols has been achieved by the photodecomposition of ethyl diazoacetate³¹⁵ and of bis(phenylsulphonyl)diazomethane³¹⁶. Probably, singlet carbethoxycarbene reacts with the sulphur N_2 CHCO₂Et+EtSH \longrightarrow EtSCH₂CO₂Et+EtOCH₂COSEt

$$
N_{2}CHCO_{2}Et + EtSH \xrightarrow{\phantom{N_{2}N_{2}}} EtSCH_{2}CO_{2}Et + EtOCH_{2}COSEt
$$
\n
$$
53\% \qquad (247)
$$
\n
$$
6\%
$$

atom to give **a** sulphur ylide intermediate, followcd by proton migration as with alcohols. The product **247** involves the Wolff rearrangement of the carbene to a

 \overline{a}

$$
N_{2}CHCO_{2}Et \xrightarrow{\hbar\nu} \text{CHCO}_{2}Et \xrightarrow{EtSLH} H \xrightarrow{\text{Et}S} \text{CHCO}_{2}Et \xrightarrow{\text{Et}S} \text{Et}SCH_{2}CO_{2}Et
$$
\n
$$
EtOCH = O \xrightarrow{EtSLH} (247)
$$
\n
$$
\text{cetane. The corresponding benzophenone-sensitivee irradiation of ethyl diazc\nccetate in ethyl mercaptan gives products in similar yields. The radical mechanism\nis probably involved in the S-H insertion even in direct photosis as follows:\n
$$
N_{2}CHCO_{2}Et \xrightarrow{\hbar\nu} \text{CHCO}_{2}Et \xrightarrow{EtSLH} \text{Et}S\cdot + \cdot CH_{2}CO_{2}Et
$$
\n
$$
EtS\cdot + N_{2}CHCO_{2}Et \xrightarrow{\text{Et}S} N_{2}+EtSCHCO_{2}Et
$$
$$

ketene. The corresponding benzophenone-sensitized irradiation of ethyl diazoacetate in ethyl mercaptan gives products in similar yields. The radical mechanism is probably involved in the S-H insertion even in direct photolysis as follows:

$$
\begin{array}{cccc}\n & & \\
 \downarrow & & \\
 \hline\n \text{EtoCH} = 0 & \xrightarrow{\text{EtSL}} (247) \\
 \text{ne corresponding benzophenone-sensitive d irradiation of e}\n \\ \text{ethyl mercaptan gives products in similar yields. The radical } \times \text{involved in the S-H insertion even in direct photosis as f}\n \\ \text{N}_{2}\text{CHCO}_{2}\text{Et} & \xrightarrow{\hbar\nu} \text{Et} \text{CHCO}_{2}\text{Et} & \xrightarrow{\text{Et} \text{S}\text{H}} \text{Et} \text{S} + \cdot \text{CH}_{2}\text{CO}_{2}\text{Et}\n \\ \text{Et} \text{S} + \text{N}_{2}\text{CHCO}_{2}\text{Et} & \xrightarrow{\text{Et} \text{S}\text{H}} \text{N}_{2} + \text{Et} \text{SCHCO}_{2}\text{Et}\n \\ \text{Et} \text{SCHCO}_{2}\text{Et} + \text{Et} \text{SH} & \xrightarrow{\text{Et} \text{S}} + \text{Et} \text{SCH}_2\text{CO}_{2}\text{Et}\n \\ \text{radical reaction of ethyl diazoacetate can be achieved the azoisobutyronitrile (AIBN) as a radical initiator. Radi.
$$

Thc same radical reaction of ethyl diazoacetate can be achieved thermally by employing azoisobutyronitrile **(AIBN)** as **a** radical initiator. Radical-induced decomposition of diazo compounds was also proposed by Horner and coworkers^{347, 348}. The formation of the S-H insertion product was rapid in the presence of, and very slow in the absence of, the radical initiator. No other product was found except **a** trace of diethyl disulphide:

$$
EtSH + N2CHCO2Et + AIBN \xrightarrow{60^{\circ}C} EtSCH2CO2Et + (EtS)2
$$

Since no rearrangement product was detected in the aromatic thiol (Table **38),** the carbene may **not** be involved in these reactions, and the insertion into the sulphur-hydrogen bond might occur only by radical chain reactions.

Dialkyl sulphides afforded stable sulphur ylides by adding carbenes with strongly electron-withdrawing substituents. Biscarbomethoxycarbene, generated photochemically from dimethyl diazomalonate, reacts with alkyl and aryl sulphides and sulphoxides to form stable sulphonium and sulphoxonium biscarbomethoxymethylides^{319, 350}. The ability of sulphur to stabilize an adjacent negative charge remains an interesting phenomenon.

Thiol	Mode of decomposition $S-H$ insertion $(\%)$		Rearrangement $(\%)$
EtSH	hν	53	o
EtSH	$hv + Ph2CO$	70	o
<i>i</i> -PrSH	hv	31	
t -BuSH	hν	35	b
PhSH	hν	74	0
PhSH	$h\nu + Ph2CO$	90	
p-MeOPhSH	hν	58	

TABLE 38. Photodecomposition of ethyl diazoacetate in thiols

Comparison of the sulphur ylides to nitrogen ylides indicates that factors beyond electrostatic stabilization must be involved. The simplest explanation involves delocalization of electron density into low-lying d orbitals of sulphur³⁵¹. It has been pointed out that a combination of only s and p type orbitals can also accommodate

such a valence shell expansion³⁵² (ylene structure 250). A large shift (\sim 110-170 cm⁻¹) to lower energy in the carbonyl stretching frequency in the infrared spectrum of the biscarbomethoxymethylides compared to the analogous sulphonium salts also demonstrates extensive delocalization of charge by such

TABLE 39. Formation of sulphonium ylides from bis(carbonyl)diazo compounds and sulphides^{319, 350}

$(R = COMe, R^1 = COOEt, R^2 = COMe, R^3 = PPh2)$				
Sulphides	Ylides	Melting point $(^{\circ}C)$	i.r. $(C=0)$	Yield $(\%)$
Me ₂ S	$Me2 \overline{S} - \overline{C}R_2$	169-170	1625, 1675	88
Et ₂ S	$Et_2S-\overline{C}R_2$	150	1625, 1655	57
Ph_2S	Ph_2S-CR_2	$127 - 128$	1650, 1675	12
Me ₂ S	$Me2 S - \overline{C}R2$	$134 - 135$	1625, 1655	87
Me ₂ S	$Me2S - \overline{CR}22$	166–167	1570, 1600	52
Me ₂ S ³⁵³	$Me3 - \overline{C}R2R3$		1572	
$Me3SiCH2SMe291$	$Me3SiCH2\searrow \frac{4}{C}$ $-CR$ Me	$132 - 134$		

О \mathbf{I}

carbanion stabilizing groups. An increase in the ylide carbon-carbonyl carbon π -bond order accompanies the decrease in the π -bond order of the carbonyl group (betaine structure **249).**

In the reaction of dimethyl diazomalonate with saturated cyclic sulphides, the corrcsponding sulphonium ylides were **also** obtained as stable colourless solids. Irradiation of dimethyl diazomalonate in pentamethylene sulphide gave **41%** of sulphonium ylide. Thiethane did not give sulphonium ylides because of its photo-
 *polymerization*³⁵⁴:
 S + N₂C(COOMe)₂ $\xrightarrow{h_r}$ $S-\overline{C}(\text{COOMe})_2$ polymerization 354 : *e* reaction of dimetriy dazomatoriate

onding sulphonium ylides were also

tion of dimetriyl diazomalonate in pe

nium ylide. Thiethane did not give sulphare
 $\frac{hc}{s}$ + $N_2C(COOMe)$, $\frac{hc}{s}$
 $+ N_2C(COOR)$, $\frac{hc}{s}$

$$
\begin{matrix} & + N_2C(COOME) & \xrightarrow{h\nu} & S-\bar{C}(COOME) & \\ & & \searrow & \end{matrix}
$$

High stereoselectivity is observed with 4-t-butyltetrahydrothiopyran³⁵⁵:

$$
+\overbrace{\hspace{1.5cm}}^{S} + N_{r}C(COOR)_{r} \xrightarrow{h_{r}} + \overbrace{\hspace{1.5cm}}^{S} \overbrace{\hspace{1.5cm}}^{C(COOR)_{r}}
$$

Even vinylic³⁵⁶ and acetylenic sulphides³⁵⁴, thiophene and benzothiophene³⁵⁴, in which the lone pair of sulphur is highly delocalized, may trap the carbene produced in the photolysis of dimethyl diazomalonate. These ylides are quite stable at room temperature:

The photolysis of dimethyl diazomalonate in Δ^3 -dihydrothiopyran³¹⁷ gave the sulphonium bis(carbomethoxy)methylide (251), which rearranged to 252 in 83% yield **on** heating at 180°C for 30min. The ylidc **251** could be isolated and rearranges only at higher temperatures, while ylides derived from open-chain allylic sulphides gave rearranged products at room temperature. This low susceptibility of the ylide **(251)** toward rearrangement is probably due to the transition state of the concerted allylic rearrangement requiring the unfavoured boat form of the dihydropyran ring bearing the COOMe group *endo* to the sulphur atom.

A clear preference for the *exo* orientation was also seen in the rearrangement of **253 358.**

Photolysis of diazocyclopentadiene in alkyl sulphides also gives stable cyclopentadienylides in high yields³⁵⁹. Dimethylsulphonium cyclopentadienylide is considered to be the resonance hybrid of the ylide structures **254, 255** and the

ylene structure 256, with the ylide structures predominating³⁶⁰. Stabilized oxosulphonium ylides, unavailable by the direct condensation of sulphoxides and α -halocarbonyl compounds³⁶¹, are suited for the carbene approach^{350, 362}. Both

Example 2. The sum of the original methods and photographs have been employed.

\n
$$
Me_{2}S = O + N_{2}C(R)CO_{2}R' \xrightarrow{h\nu} Me_{2}S - \overline{C}(R)CO_{2}R'
$$
\n
$$
R = H \text{ or } CO_{2}Me
$$
\n
$$
R' = Me, Et
$$
\n
$$
Ph_{2}S = O + N_{2}C(COOMe)_{2} \xrightarrow{h\nu} Ph_{2}S - \overline{C}(COOMe)_{2}
$$
\n
$$
10
$$
\n
$$
12\%
$$

The relative reactivity of the sulphur atom of dimethyl sulphide toward carbene was estimated from the product distribution in the photolysis of dimethyl diazomalonate in mixture of cyclohexene and dimethyl sulphide³⁴⁹. It was suggested that

TABLE 40. Yields of products obtained from dimethyl diazomalonate and dimethyl sulphide in the presence **of** an equimolar amount of a second substrate

Yield of ylide $(\%)$	Yield of adducts $(\%)$
61	15ª
49	8.5^{b}
	5 ^b
	10 ^c
73	12 ^d
	60 44 78

^aMixture **of 7,7'-dicarbomethoxynorcarane** and 2-cyclohexenyl malonate in the ratio of 2.1 : 1.

* Olefin adduct only.

0-H insertion product.

 d Ethylene elimination and C-H insertion products.

the biscarbomethoxycarbene reacts with dimethyl sulphide about *6* timcs faster than with the double bond of cyclohexene. Similar high reactivity of sulphur toward carbenes was also observed in the photolysis of equimolar mixtures of dimcthyl sulphide and other nucleophiles.

Ethyl diazoacetate and aryl diazomethanes did not yield stable sulphonium Ethyl diazoacetate and aryl diazomethanes did not yield
ylides, but afforded decomposition products of the latter^{363, 364}:
RSR + N₂CHCO₂R' $\frac{hv}{m}$

$$
RSR + N_2CHCO_2R' \xrightarrow{nv}
$$

$$
\begin{bmatrix} R \setminus \begin{matrix} 1 \\ 1 \\ 2 \end{matrix} & C \end{bmatrix} \longrightarrow \begin{matrix} R \setminus C \\ 1 \\ 2 \end{matrix} \longrightarrow \begin{matrix} 1 \\ 1 \end{matrix} \longrightarrow \begin
$$

$$
R = Et, i\text{-}Pr, or t-Bu
$$

The reactions of alkyl sulphides bearing a β -hydrogen atom with ethyl diazoacetate yielded, by @-climination, alkyl thioacetates formed by thermal intramolecular olefin elimination from the intermediate sulphonium ylides via a five-member cyclic transition state.

Additional results concerning the behaviour of sulphonium ylides are available. Thermal decomposition of biscarbomethoxymethylides gave dimethyl alkylthiomalonate in quantitative yield³⁶⁵. This supports the proposed ylide mechanism in the reactions of carbethoxycarbene with hetero atoms bearing β -hydrogens.

-% EtSCH(COOMe), + CH,=CH, ,C;L H MeOOC COOMe

Photolysis of phenyl- and diphenyl-diazomethanes in diethyl and benzyl methyl sulphides gave mainly α -C-H insertion, with some β -elimination products, but phenyldiazomethane with i -propyl and i -butyl sulphides gave the β -elimination products³⁶⁴. The C-H insertion has been suggested to proceed via sulphur ylides followed by benzyl or benzhydryl migration from sulphur to carbon.

Some $C-S$ insertion of carbene was also observed. The photolysis of diphenyldiazomethane in diisopropyl sulphide gave *i-*propyl benzhydryl sulphide (257) and *i*-propyl(*i*-propylthio)diphenylmethane (258) in 20 and 15% yields, respectively. The sulphur ylide could, by thermal Stevens rearrangement, yield 258.

Such 1,2-shifts are forbidden in **a** concerted reaction mechanism. In fact, the accumulated evidence points to a diradical pathway³⁶⁶. A CIDNP enhancement of the n.m.r. signal of the methylene group is observed if the reaction is carried out in an n.m.r. spectrometer³⁵⁴.

Hence, C-S insertion of carbenes is considered to involve homolytic cleavage of the sulphonium ylide yielding a radical pair, which collapses to the product. Intervention of radical pair intermediates has been suggested in the reaction of sulphonium salts with bases³⁶⁶⁻³⁶⁸.

Stevens rearrangement was also observed in the photolysis of trimethylsilyldiazoacetate and trimethylsilyl(phenyl)diazomethane in dimethyl, diethyl and di-i-propyl sulphides at room temperature³⁶⁹.

TABLE 41. Photolysis of ethyl trimethylsilyldiazoacetate and trimethylphenyldiazomethane in alkyl sulphides at room temperature

The α -C-H and β -elimination reactions were found to yield minor products. **A** sulphonium ylide was also isolated when the reaction was carried out below 0 *"C.* However, this ylide does not rearrange to the **C-S** insertion product **260.**

The corresponding reaction with the vinyl sulphide **(261)** affords **262** and **263;** product **263** results formally via carbene insertion into the *C-S* bond of **261.** No addition product was observed³⁷⁰. Thermal decomposition of ethyl diazoacetate in

vinyl sulphides and acetylenic sulphides gave mainly @-elimination products. No *C-S* insertion product has been observed3sG. **357.** In the photolysis of diazocyclopentadiene in di-r-butyl sulphide, *C-S* insertion, @-elimination and the **C-H** insertion products were obtained³⁵⁹.

In the photolytic reactions of diazo carbonyl compounds with dialkyl disulphides, the products arising from thiocarbonyl elimination have been observed. Thus, the reaction of dimethyl diazomalonate with dimethyl disulphide afforded dimethyl

methylthiomalonate **(265)** and dimethyl diniethylthiomalonate **(266)** via **an** ylide intermediate **(264) 354s 364.**

$$
N_{2}C\left\{\n\begin{array}{ccc}\n\text{SiMe}_{3} & + \text{RSSR} & \xrightarrow{h_{1'}} & \text{RSCHSiMe}_{3} \\
\text{COOEt} & | & \text{COOEt} \\
\text{R} = \text{Me}, 49\% & R = \text{Et}, 95\% \\
\text{N}_{2}\text{CPh}_{2} + \text{MeSSMe} & \xrightarrow{h_{1'}} & \text{MeSCHPh}_{2} \\
& 49\% & \n\end{array}\n\right.
$$

Photolysis of diphenyl and phenyldiazomethanes in dimethyl sulphide **led** to 267, 268 and 269. Product 267 can be rationalized in terms of Stevens rearrangement of the intermediate sulphur ylide; products **268** and **269** do not conform to this scheme. but rather to [2,3]sigmatropic rearrangement of sulphur ylide³⁶⁴ (Sommelet-Hauser rearrangement).

The ratio of Sommelct-Hauser to Stevens rearrangement varied when the photolysis of diphenyldiazornethane in dimethyl sulphide **was** carried out in different

types of solvents. The striking result is the formation of a norrnal Sommelet-Hauser type product (orbital symmetry-allowed [2,3]sigmatropic process) in polar solvent conditions, whereas in non-polar conditions only a Stevens type product (orbital symmetry forbidden process) is formed. Possibly, in the ylide **270,** the methylene carbanion attacks the benzene ring followed by proton migrationhence, the necessity for **a** more polar medium. In non-polar solvents this requirement is not met, so that homolytic cleavage competes successfully with carbanion attack $(k_1 > k_2)$ and the product is derived from the net orbital symmetry forbidden 1,2-shift.

TABLE 42. Solvent effects on the reaction of diphenyldiazomethane with dimethyl sulphide⁴

Solvents	Product ratio, 268/267
Cyclohexane	0.2
Benzene	0.3
Acetone	3.2
Mesityl oxide	3.5

Mole **ratio** of sulphidc **to** solvent **is 0.3.**

1,2-Alkyl shifts of the trialkylsilyl group have also been reported in the reaction of diazo compounds with silyl sulphides and silyl ethers³⁷¹. Carbethoxycarbene produced in the photolysis of diazoacetate reacts with cthylthio(trimethyl)silane **(271)** to yield ethy! **triniethylsilyl(cthylthio)acetate (273)** in **38%** yield and ethyl ethylthioacetate **(274)** in 29% yield. The photolysis of diphenyl and phenyldiazomethanes in ethylthio(trimelhyl)silane afforded products similar to **273** and

274. The formation of **273** may involve the sulphoniuni ylide **(272)** followed by 1,2-migration of the silyl group involving a geminate radical-pair intermediate **275,** which collapses to **273.** The radical **276** through hydrogen abstraction, yields **274.** Whilc it may account for the products, the geminate ion-pair mechanism has never been established for 1,2-migration of the silyl group. A concerted 1,2-migration of the trinicthylsilyl group might be involved since silicon may expand its valence shell, and the decomposition of the sulphonium ylide via a concerted, five-member, cyclic transition state could give 274 and $Si=C$, the fate of which is unknown^{372, 373}.

9. Photochcrnistry of the diazonium and diazo **groups 455**

1,2-Migration of a silyl group from the oxonium site to a carbon may also take place in the reaction between carbenes and alkoxytriinethylsilanes **(277).**

Since the carbene attacks a sulphur atom more rapidly than the carbon-carbon double bond, the reaction with allylic sulphides may also procced by attack on the sulphur atom. The ylide may rearrange with complete allylic inversion as demanded by 2,3-sigmatropic symmetry control. Thus, the photolysis of dimethyl diazomalonate in n-butyl ally1 sulphide gave **278** as the major and **279 as** the minor product :

 $n_{\rm e}$, $n_{\rm e}$ -BuSCH₂CH-CH ie in *n*-butyl allyl sulphide gave 278 as the major and 279 as the

:

n-BuSCH₂CH=CH₂ + N₂C(COOMe)₂ $\xrightarrow{h_r}$ n-BuSCH₂CH-CH₂

C(COOMe)₃ (279) *(279)* Jh **I.** *J.* COOMe I ^{1"}
^{1"}
COOMe
¹

C₅ − C(COOMe), ---> n-BuSC-CH₂CH=CH₂

n-Bu COOMe
¹ *(278)*

In the bcnzophcnone-scnsitized photolysis, addition products were favoured indicating that electrophilic singlet carbene is capable of accepting the electron from sulphur more easily than the triplet. Copper salt or coppcr powder catalysis, on the other hand, yielded exclusively the products of the 2,3-sigmatropic process³⁷⁴.

Sulphide	$C-S$ insertion $(\%)$	Cyclopropane $(\%)$
∫SEt	51 $(18)^a$	$10(37)^a$
SEt_	46 (7)	12 (29)
SEtبر	53 (31)	7 (19)
_SEt	43 (19)	12(27)

TABLE 43. Product yields in the photolysis of dimethyl diazomalonate in allylic sulphides

" Benzophcnone photosensitized reaction.

In the rcaction of allylic sulphidcs with ethyl diazoacetate, a greater lack **of** specificity was observed³⁶³. In the photolysis of phenyl and diphenyldiazomethanes, *on* the other hand. the *C-S* insertion product is observed in *20-30%* yields, but only a trace of cyclopropane or none at all was obtained³⁶⁴:

Introduction of a bulky group at **C(31** leads to preferential formation of the *trans* isomer at the newly created double bond. Of the two orientations at C₍₃₎, the larger C₍₃₎ substituent should preferentially occupy the pseudo-equatorial orientation³⁷⁵:

Cyclopentadienylidene gave, with allylic sulphides, insertion into the C-S bond as well as addition. The former could arise from an intermediate sulphonium ylide

9. Photochemistry of the diazonium and diazo groups followed by 3,3-sigmatropic rearrangement³⁷⁶:

Intramolecular carbene reaction on the sulphur atom was also found. α -Allyl and α -arylthioall: ylcarbenes generated by a modification of the Bamford–Stevens method gave labile episulphonium ylides, which then rearranged to vinyl sulphides^{377, 378}. The episulphonium ylides could be trapped by diethyl maleate or dimethyl fumarate. An *a*-allylthioalkylcarbene generated by the photolysis of allylthioacetophenone tosylhydrazone afforded α -allylthiostyrene, which gave a smooth thio-Claisen rearrangement. 3-Thiabicyclo[3.1.0]hexane was obtained when allylthioacetophenone tosylhydrazone was decomposed photochemically in the presence of bases.

J. Wolff Rearrangement

The Wolff rearrangement of *a*-diazocarbonyl compounds has received much mechanistic attention. Thus, photolysis of α -diazo ketones in protic solvents yields carboxylic acid derivatives. Ketenes (280) have been proposed as reaction intermediates^{80, 81, 85}. In aprotic media ketenes are often obtainable in preparative

$$
\begin{array}{ccc}\nN_1 \\
R-C-C-R' & \xrightarrow{h_r} & \xrightarrow{R} & \xrightarrow{R} & \text{RRCHCO}_2R' \\
& 0 & & R' \\
& & & (280)\n\end{array}
$$

quantities. Many studies have attempted to determine whether loss of nitrogen precedes migration to give an intermediate ketocarbene (281) and/or oxirene (282).

I. Synthetic utility

The reaction has considerable synthetic use, e.g. for chain lengthening³⁷⁹:

In cyclic systems ring contraction occurs and the reaction is useful in synthesis of strained small-ring compounds:

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When **283** in benzene solution was irradiated at room temperature, in the presence of an alcohol, the product (284) was obtained via the Wolff rearrangement³⁸². The

photolysis of 285 gave in 65% yield a 2:5 mixture of β -lactam (286) and its epimerization product **(287).** The reaction yiclds some compounds whose production

by other methods is either impossible or very difficult. For instance, the photolysis of 288 is a convenient method of obtaining 6-aminoindene (289)³³⁵. Horner and

coworkers irradiated the 0-quinonoid diazoketone *(290)* in alcohol or in the presence of aromatic amines, and separated derivatives of the corresponding ketene in the form of esters or anilides of indenecarboxylic acid³⁸⁶. In boiling xylene, the product was an adduct of the ketene identified as *293.* Diazokctoncs *294-296* undergo smooth **Wolff** rearrangement, but only 2% ring contraction occurs with *296,* presurnably owing to an increase of *c.* **50** kcal mol-' in strain energy.

 (296)

Similarly pyridine and quinoline derivatives yield pyrrole- and indole-carboxylic **392.** Photolysis of **298,** through ring contraction, gives 2,5-dimethylpyrrole-3-carboxylic acid **(299)** when the reaction is carried out at low temperatures.

Without cooling, *298* reacts with *299* to give the azo dye **300a** which readily loses carbon dioxide leaving the azo compound **300b 302.**

Quinoline derivatives decompose similarly on irradiation:

This reaction can be used to obtain the corresponding indoles including the unsubstituted indole, the 5-chloro- and 5,6-dimethoxyindoles and their carboxylic The photolysis of the diazo compound **301a** gives **4-azaindole-3-carboxylic** acid which is characterized by a high stability and **is** incapable of decarboxylating. **Its** isomer, the **6-azaindole-3-carboxylic** acid from **(301b),** is very easily decarboxylated, yielding 6-azaindole³⁹³. Wolff rearrangement of α -diazoesters to alkoxyketenes, and products obtained from these were reported in a study of the photolysis of diazoacetyl enzymes³⁹⁴⁻³⁹⁶. A diazoester is attached at the active site of the enzyme and subsequently reacts with neighbouring amino acid residues. In particular, photolysis of diazoacetyl chymotrypsin presumably gencrates **a** carbcne that reacts by a rearrangement analogous to that of diazo ketones, either by insertion into water or by insertion into amino acid residues. Histidine and tyrosinc residues have so far been identified as sites of attack.

The rearrangement encountered in these investigations was also illustrated with azaserine³⁹⁶, which reacted in part as follows:

The product of photolysis of a diazothiolester protein derivative **(304)** in water, after hydrolysis, gave **305** as derived by Wolff rearrangement of the thialkoxy $group^{397}$:

Ethyl diazoacctate **(306),** phenyl diazoacetate **(307)** and N-methyl diazoacetamide **(308)** also give products from Wolff rearrangement when photolysed in protic

9. Photochcmistry of the diazonium and diazo **groups 463**

solvents. The major competing reaction is insertion into the $O-H$ bond of the solvent.

$$
N_{2}CHCO_{2}R \xrightarrow{\text{MeOH}} ROCH_{2}CO_{2}Me + \text{MeOCH}_{2}CO_{2}R
$$
\n
$$
(306) \quad R = Et \quad 20-25\% \quad 75-80\%
$$
\n
$$
(307) \quad R = Ph \quad 45-60\% \quad 40-55\%
$$
\n
$$
N_{2}CHCONHMe \xrightarrow{\hbar\nu} \to :CHCONHMe \xrightarrow{\Pi_{1}O} HOCH_{2}CONHMe
$$
\n
$$
(308) \quad 70\%
$$
\n
$$
O=C=CHNHMe \xrightarrow{\Pi_{1}O} HO_{2}CCH_{2}NHMe
$$
\n
$$
30\%
$$

The ketenes formed in the **WolR** rearrangement have been frequently subjected *in situ* to cycloaddition with C=C, C=N and N=N double bonds^{398, 399}:

.

Horner obtained **309** from azibenzil and azobenzene, **310** from benzoyl diazomethane and azobenzene, **311** from diphenylketene and azodibenzoyl, and **312** from azibenzil and α, α' -azotoluene $400, 401$. There is no doubt that the α -diazocarbonyl compounds used are first converted to the corresponding ketenes which then react with the azo compounds to form the corresponding derivatives of 1,2-diazetidine.

Photolysis of **5-acetyl-3,3-dimethyI-3H-pyrazole (313)** gives rise to a ketene intermediate **314,** which reacts with **313** to produce the adduct **315 402.** The ketene

obtained from the photolysis of 283 reacts with azobenzene to give spiro compounds³⁸². Ketenes formed by photolysis of diazoketones can also add to azomethines to give β -lactams⁴⁰³:

The scope of this cycloaddition is limited above all by reaction of the starting compound itself or its primary fragments (carbenes) prior to rearrangement. With diazoketones intramolecular cycloadditions occur preferentially at the ketene stage in the photochemical reaction^{404a}, but at the carbene stage in the catalytic process^{404b}.

Methyl diazoacetate upon photolysis in benzene affords two tetrameric isomers 316 and 317 (overall yield $55\frac{\%}{100}$, which apparently arise from dimerization of the cyclopropane (319) formed by addition of methoxyketene (318) to carbomethoxycarbene.

Irradiation of α , β -epoxydiazomethylketones (320) in benzene afforded compounds to which the butenolide structure 322 was attributed⁴⁰⁷.

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γ-hydroxy-α, β-unsaturated esters⁴⁰⁷. The same reaction in the presence of an excess of methanol or ethanol yielded
hydroxy- α , β -unsaturated esters⁴⁰⁷.
 R^1 *OH*
 R^2 *R*² *OR*⁴
 R^3 *OR⁴*

$$
321 \xrightarrow{R^*OH} R^2
$$

On rearrangement of unsymmetrically-substituted 2-diazo-l,3-dicarbonyl compounds, both **R1** and **R2** may migrate in the diacylcarbene providing an indication of the relative migration tendencies of various groups $408-410$.

TABLE 44. The relative migration tendencies of various groups

Cyclic 2-diazo-1,3-dicarbonyl compounds can also be included in these investigations. The arrows in structures 325^{411} , 326^{412} and 327^{413} indicate the centres of selcctive migratory aptitude.

The photolytic dccornposition of **6-diazo-2,4-cyclohexadien-l** -one in methanol affords a guaiacol **(329),** phenol, a dinier **(330)** of methyl **cyclopentadienecarboxylate** and a reddish pigment 331 ¹¹⁴.

2. The mechanism of Wolff rearrangement

To date, the most thorough study of the Wolff rearrangement of diazo esters has been conducted by Strausz and coworkers⁴¹⁵⁻⁴¹⁹. From the photolysis of ethyl diazoacetate in i-propanol they obtained **332** *(9%),* **333** *(29%),* **334** *(25%)* and **335 (12%).** The ethoxy ether **333** results from ethoxy migration to form ethoxy ketene which is trapped by *i*-propanol. The hydroxy ester stems from C-H insertion into

Mich is trapped by *i***-propanol. The hydroxy ester sterned in the CH₂CH₂CO₂Et + EtOCH₂CO₂CHM
M₂CHCO₂Et** $\frac{\text{Me}_1 \text{C} \text{F} \text{O} \text{F}}{h_F}$ **Me**₂CCH₂CO₂Et + EtOCH₂CO₂CHM ^I**(333)** OH *(332)* **+Me,CHOCH,COzEt+Me,CHOCH~COzCHMe, (334) (335)**

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the solvent; the ether **334** arises from **0-H** insertion and may also be a product of transesterification. Control studies indicated that the formation of **335** is a photoinduced process which does not involve carbenes. Instead, a mechanism involving light-induced heterolysis of the diazo ester **was** suggested :

The mechanism of the Wolff rearrangement of α -diazo ketones and esters in the gas phase has been studied by '3C-labelling of diazo derivatives **336.** The results are attributed to the formation of two different ketenes **337** and **339,** involving also an oxirene intermediate 338 . Franzen applied the ¹⁴C-labelling technique for the photolysis of azibenzil (340). He concluded that oxirene was not an intermediate¹²⁰.

However, more recently, Strausz and coworkers found in the photolysis of ¹³C-labelled azibenzil in cyclopentane that the isotopic composition of CO indicated scrambling of the oxygen atom to an extent corresponding to **54%** oxirene

By the same technique, **28-32%** oxirene participation (path 2) was indicated in the gas-phase photolytic decomposition of methyl and ethyl diazoacetate (336). Molecular orbital calculations suggest that ketene is more stable than the isomeric oxirene or formyl carbene by c . 70 kcal mol^{-1 421}.

An experiment with ¹⁴C-labelled α -diazohomoadamantanone (341) yielded only the acid **342.** No radioactivity could be found in the carbon dioxide upon oxidative decarbonylation, and hence the oxirene **343** did not intervene in any way in this reaction⁴²². The photolysis of 2-diazo-1-¹³C-naphthalen-1(2H)-one (344) yields indene-1-carboxylic acid containing all the label in the carboxy group¹²³. This result shows that no isomerization of the oxocarbene 345 to 347 takes place. Compound **348** may be formed directly via Wolff rearrangement of **344** by concerted **N2** expulsion and ring contraction.

A test for the transient formation of oxirenes was based on the observation that when a methylene group is adjacent to the diazo function, decomposition to the oxocarbene is followed by **a** 1,2-shift to the carbene carbon atom, resulting in formation of an α , β -unsaturated ketone. Thus, for unsymmetrically substituted diazoketones, different *a,* P-unsaturated ketones would form if the oxocarbene initially produced equilibrated with an oxirene intermediate 424 .

Proof of oxirene formation that is completely independent of the Wolff rearrangement was obtained with unsymmetrical α -diazo dialkyl ketones⁴²⁵ and some aryl substituted α -diazo ketones⁴²⁶. The trapping reaction yielding 349 strongly suggests that oxirenes are true intermediates, and not transition states⁴²⁷.

Photolysis of dimethyl diazomalonate in the presence of methanol- $d_{\bf{a}}$ affords **350** and **351** in the ratio **9** : **4.** The absence of **352** suggests that no oxirene intermediate **is** involved, and the Wolff rearrangement to **351** proceeds with methoxy

group migration, as indicated below 428 :

Photolysis of the diazoketone in the presence of sufficient benzophenone, so that the latter absorbed > *95%* of the light, yielded only traces of the enones obtained by direct excitation. Thus, it can be assumed that triplet species do not rearrange via oxirene intermediates. There seems to be general agreement that singlets must be involved. Padwa and Layton 429 report that photosensitized decomposition of diazoacetophenone in i-propanol results in a decrease in yield of Wolff rearrangement product and an increase in reduced product as compared to direct photolysis :

O	∥	
Ph-C–CHN ₂ +Me ₂ CHOH	Åν	PhCH ₂ CO ₂ CHMe ₂ +PhCOMe
(A)	(B)	
A/B	Direct $hν$	2.5
Sensitized $hν$	0.003	

Diazodibenzoylmethane **(353)** undergoes two primary photochemical processes leading to **354** and to dibenzoylmethane **(355) 530.** The formation of **354** is related to the lowest excited singlet state of **353** and the formation of **355** is related to the lowest triplet state of 353. The quantum yields of both processes, Φ_{354} and Φ_{355} , are strongly wavelength dependent. It is unambiguously demonstrated that the population of two excited states depends on the energy of the excited light, thus causing a wavelength effect.

Photosensitized decomposition differs from photolytic decomposition of diazot hiolacetate in that rearrangement is suppressed and products derived from radical-

Wavelength of irradiation (nm, 297 K)	Ratio	
	355	354
366		0.80
313		0.31
254		0.17
P yrex (77 K)		20
Sensitized (Michler ketone)	19	

?-ABLE 45. Wavelength **effect** on the photolysis of **353**

like abstraction and insertion are obtained⁴³¹. Photosensitized decomposition of diazothiolacetate (356) is a triplet process and the various reactions of triplet ethyl α -diazothiolacetate (359) and/or triplet carbene (360) are summarized in Scheme 9.

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It is apparent that there can be only minor intersystem crossing in the photosensitized decomposition of **356.** Nevertheless, Wolff rearrangement products **(358)** are formed in photosensitized decomposition of **356.** Jones and Ando that aliphatic diazoketones such as diazocyclohexanone underwent Wolff rearrangement on unsensitized irradiation. The corresponding carbene could not be trapped with olefins. Photosensitized decomposition gave an intermediate that did give **Wolff** rearrangement and was trapped by olcfins:

The sensitized decomposition of diazoacetone in olefins yields similar adducts while none could be found in the unsensitized reactions:

It is tempting to describe these changes in the most simple terms, that is, singlet carbenes are involved in the unsensitized and triplet in the sensitized decompositions. The singlet finds an accessible pair of electrons in the adjacent carbon-carbon single bond and reacts with these at **a** faster rate than with the more distant π -electrons of the external olefin.

Carbcnes that are formed by photolysis of diazo compound **362** are rapidly deactivated to their ground states, probably because of **an** internal 'heavy atom cffect'. This modifies their chemistry and provides a useful probe for determining the spin multiplicity of the ground state⁴³³⁻⁴³⁵.

Use of methanol- d_4 shows that it is the methyl group of the ether which comes from the alcohol, and no product of Wolff rearrangement is cvident :

Since photolysis of mercury-free diazoacetates in alcohol gives rise to products of oxygen-hydrogen insertion, carbon-hydrogen insertion, Wolff rearrangement and 'exchange' reactions, the presence of a mercury atom adjacent to the carbene site enhances the selectivity of the intermediate⁴³⁶.

Irradiation of **363** in methanol leads to competition between the phosphorylcarbene and methylene phosphorane oxide rearrangement : **364-365-369,** and the equally possible classical Wolff rearrangement **364-368-372,** the methyl esters **369** and **372** are formed in the ratio of approximately **4** : 1.

It is possible, however, to reverse this ratio by introducing a p -dimethylamino group into the benzoyl group of **363.** Furthermore, **364** reacts with methanol by 0-H insertion although the phosphorane oxide **366** cannot be isolated; it undergoes intramolecular photochemical reduction of the carbonyl group to form the diradical **370,** which then cyclizes to the oxctane **371.** Intermediate **365** afforded dienes on reaction with conjugated ketones⁴³⁷.

K. Biodiazo Compounds

cyclic olefins: Bisdiazo compounds are potential intermediates for the synthesis **of** strained

Vicinal bisdiazo compounds yield acetylenes^{438–440}. This route has led to the successful synthesis of a cycloheptyne and of cyclohexyne^{439, 440}.

In 1959, Kirmse reported the ring contraction of **2,6-bis(diazo)cyclohexanone** and interpreted the result on the basis of a Wolff rearrangement⁴¹¹. Later studies gave similar results⁴⁴²⁻⁴⁴⁴. However, a recent alternative explanation is that in the proposed carbene intermediate obtained from **373** the interaction of a carbanion with a carbene centre should compete favourably with σ -bond migration. While **374** may yield the ketene **375,** the exceedingly low activation energy associated with nitrogen elimination from aliphatic diazonium salts suggests that formation of the cyclopropenone could compete with σ -bond cleavage⁴⁴⁵. A methanolic solution of

373 was irradiated at -40° C to yield methyl cyclopentene-3-carboxylate (376, 15%), methyl cyclopentene-1-carboxylate (377, 61%), methyl *trans-2*-methoxycyclopentane carboxylate **(378,** *5%)* and methyl cis-2-niethoxycyclopentanecarboxylate **(379,** 10%). Irradiation of **(373)** in toluene or furan at -40 "C or benzene at 10°C containing a trace of water gave the ring-contracted cyclopentene-1-carboxylic acid **(380)** and the corrcsponding anhydride **(381)** in a combined 65% yield. Irradiation of 1,3-bisdiazo-1,3-diphenyl-2-propanone (382) at -40 °C in a

9. Photochemistry of the diazonium and diazo groups

4:1 methanol-tetrahydrofuran (v/v) solution gave methyl threo-2,3-diphenyl-3-methoxypropionate (383, 41%), methyl erythro-2,3-diphenyl-3-methoxypropionate (384, 13%), methyl trans-2-phenylcinnamate (385, 5%) and diphenylacetylene (24%). Irradiation of 382 under conditions at which diphenylcyclopropenone was photostable, indeed resulted in the isolation of the latter. Photolysis of 387 at -40° C in an inert solvent such as toluene resulted in a 65% yield of the acetylene (386) as the only characterized product.

In the photolytic decompositions, cyclopropenones are established as the major primary products, at least in the acyclic cases. This also provides support for the intermediacy of the cyclopropenenone 388 in the photolysis of 373.

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CHAPTER **10**

The electrochemistry of the diazo and diazonium groups

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1. INTRODUCTION

This review includes a discussion of the electrochemical oxidation and reduction of diazo compounds **(1)** and diazoniuni salts **(2),** where R can bc any of a wide variety of alkyl or aryl substituents, or, with **1,** hydrogen. Diazo compounds have been both oxidized and reduced electrochemically; on the other hand, the positive charge in *2* inhibits anodic electron removal, and hence diazonium salts have only becn rcduccd electrochemically.

> **R,CN, RN:X-** (1) (2)

Many mechanistic fcalures of the reactions to be discussed herein are similar to thosc observed in the electrochemistry of a numbcr of other simple functional groups. The rcadcr may wish to explore such analogies by referring to onc or more reference works on organic electrochemistry, of which there are several texts¹⁻³ and review articles¹⁻⁷ explicitly written for the organic chemist unfamiliar with this area. The rcvicw by **Fry** and Reed' rcvicws elementary considerations involved in interpreting the clectrochemical behaviour of systems containing multiple bonds, and much of what is said there could be applied in the present context. Most of the rcferences cited above also elaborate upon the principles and techniques employed in organic electrochemistry, and hence the present article will presuppose a working knowledge of the appropriate concepts. The emphasis herein is, at any rate, heavily upon the chemical reactions undergone by 1 and 2 at electrodes, rather than upon clectrochcmical details.

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A. Reduction

Most studies of the electrochemical reduction of diazo compounds have been carried out with diazocarbonyl compounds (3). Since they constitute the mechanistically best-understood class of diazo compounds, their elcctrochemical behaviour will be discussed in some detail. It is probably true that many of the same considerations can be applied to an understanding of thc electrochemistry of simple alkyl and aryl diazo compounds, but thcre are probably differences, as we shall see.

> **R'** I RCOCN, C,H,COCHN, **(3) (4)**

The electrochemical reduction of diazocarbonyl compounds is nicely understood by reference to diazoacetophenone **(4)** This compound exhibits three polarographic waves in aqueous buffer at pH **6.** The heights of the three polarographic waves correspond to uptake of six, two and two electrons, respectively, suggesting stepwise reduction of **4** to a-aminoacetophenone *(S),* then acetophenone *(6)* and finally a-phenylethanol **(7)** (Scheme 1).

Controlled-potential coulometry confirmed the numbers of clectrons consumed in each step as six, two and two, respectively⁸, and the suggestion that the second and third waves in thc polarogram of **4** correspond to reduction of intermediates **5** and *6* was supported by thc observation that *5* docs exhibit two polarographic waves corresponding to the last two of **4,** and that *6* exhibits a single wavc at the potential of thc third **wave** of **4** (or second wave of **S)\$.** Thc reduction of **4** to **5** is clearly particularly complex in nature, involving as it docs overall addition of six electrons and six protons. Although onc could imagine a largc number of sequences in which these electron and proton transfers could take place. other investigations have thrown considerable light upon this conversion. For example, diazocaniphor **(8)** also exhibits **a** six-electron wave over the pH range 6-8.5, and the aminoketone (9) (stereochemistry uncertain) is the product of preparative electrolysis9. In **alkali,** Iiowever, the six-clcctron wave of **S** splits into **two** wavcs, a two-electron wave followed by a four-electron wave (Scheme 2)⁹. Controlledpotential elcctrolyscs at thc potentials of thc first and sccond waves of **8** in alkali afford hydrazone **10** and aniinokctone **9,** respectivcly. I'resumably **10** is also an intermediate in the reduction of $\bf{8}$ in the pH range 6-8.5. The conversion of $\bf{8}$ to

10, like many other organic electrode processes, especially reductions of multiple bonds', probably involves an alternating sequence of electron and proton transfcrs (Scheme 3), i.e. a so-called ECE process. One might imagine an alternate sequence

(10)
\nSCHEME 2
\nthere organic electrode processes, especially reduct
\ninvolves an alternating sequence of electron and
\na so-called ECE process. One might imagine an al
\n
$$
R_2CN_2 \xrightarrow{e^-} R_2\tilde{C}-N=\bar{N} \longleftrightarrow R_2\bar{C}-N=N
$$

\n(1)
\n(1)
\n11 $\xrightarrow{H^+} R_2\tilde{C}-N=NH$
\n(12)
\n12 $\xrightarrow{e^-} R_2\bar{C}-N=NH \xrightarrow{H^+} R_2C=NNH_2$
\n(13)
\nSCHEME 3

to 12 involving initial proton transfer (equation 1). A number of lines of evidence favour Scheme 3, however: (a) diazo camphor is unstable in acid $(pH < 6)$ ⁹; (b) the radical anion **(11)** has been observed by e.s.r. spectroscopy in the reduction of

$$
1 \xrightarrow{\text{H}^+} R_2 \overset{\star}{C} - N = NH \xrightarrow{\text{e}^-} 12 \tag{1}
$$

diazoacetophenone (although under quite different experimental conditions)¹⁰; and (c) diazoacetophenone **(4)** is reduced to acetophenone *(6)* in **a** rwo-electron process at pH *5* **11,** demonstrating that protonation of **1** leads to a different reduction pathway involving protonation on carbon (equation **2). 1EXECT:** (a) diazo camphor is unstable in acid (pH < 6)⁹; (b) the anion (11) has been observed by e.s.r. spectroscopy in the reduction of $1 - \frac{H^+}{2}$, $R_z\bar{C} - N = NH - \frac{e^-}{2}$, 12 (1) diazo camphor is unstable in aci

$$
1 \xrightarrow{\text{H}^+} R_2 \text{CHN}_2^+ \xrightarrow{\text{c}^-} R_2 \text{CH} \cdot \xrightarrow{\text{c}^-} R_2 \text{CH} \cdot \xrightarrow{\text{c}^-} R_2 \text{CH}^- \xrightarrow{\text{H}^+} R_2 \text{CH}_2
$$
 (2)

The conversion of hydrazone **10** to amine **9** involves no ncw chemistry; it is known that hydrazones^{12, 13}, including those α - to a carbonyl group¹⁴, are reduced electrochemically in a four-electron step, and that this involves initial cleavage of the $N-N$ bond, followed by reduction of the resulting imine. Thus the overall sixelectron reduction of diazo conipounds **1** may now be seen to involve a sequence of three two-electron reductions (equation 3). Each of the latter in turn presumably is itself an ECE process, analogous to the conversior. of **1** to **13.**

$$
1 \xrightarrow[2]{2e^-} 13 \xrightarrow[2]{2e^-} R_2C = NH \xrightarrow[2]{2e^-} R_2CHNH_2
$$
 (3)

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The electrochemical behaviour of 3-diazooxindole **(14)** is very similar to that of **8.** In base **@H** > *8.5),* a two-electron reduction to hydrazone **15** is observed, followed by a four-electron reduction to amine 16 at more negative potentials¹⁵ (equation 4). acetate **(17)** is reduced cleanly to ethyl phenylacetatc in aqueous dioxane at pH 7

surprisingly, the diazophosphonate **18** is reduced in similar fashion in 90% yield even in alkaline media, in contrast with the behaviour of **4, 8** or **14** under such conditions¹⁶. The reasons for this difference are not clear.

In the only study carried out to date under aprotic conditions, Elofson and coworkers explored the electrochemical behaviour of several diazo compounds in sulfolane¹⁷. Like 17, ethyl diazoacetate suffered clean reduction to ethyl acetate and nitrogen. Diazomethane was not reducible electrochemically. Reduction of diazodiphenylniethane **(19)** was more complex : diphenylmet hane **(24,** 20%) and benzhydrylamine (23, 20%) were the major products, but azodiphenylmethane (22, yield unspecified) could also be identified as a product by the use of mass spectrometry. Formation of **22-24** was rationalized by the reactions shown in Scheme **4.** It would appear that successive electron and proton transfers to **21** should

Contentane was not reducible electrochemically. Reducone (19) was more complex: diphenylmethane (24, 2(23, 20%) were the major products, but azodipheny
\nied) could also be identified as a product by the
\nFormation of 22–24 was rationalized by the reacti
\nould appear that successive electron and proton transfer
\n(C₆H₃)₂CN₂
$$
\xrightarrow{e^-}
$$
 (C₆H₃)₂CN₂ $\xrightarrow{H^+}$ (C₆H₃)₂CN₂ $\xrightarrow{(20)}$ (20)
\n20 $\xrightarrow{(C_6H_3)_2$ CH++N₂ (21)
\n20+21 $\xrightarrow{(C_6H_3)_2}$ CHN=NCH(C₆H₃)₂ (22)
\n22 $\xrightarrow{\text{reduction}}$ (C₆H₃)₂CHNH₂ (23)
\n21 $\xrightarrow{\text{H} \text{-n} \text{bstraction}}$ (C₆H₃)₂CH₂ (24)
\nSCHEME 4

be at least as likcly as hydrogen abstraction as a route to **24,** and that **23** might be formed by electrochemical reduction of 20, not 22, but the scheme seems essentially sound. Judging from these results, the electrochemistry of diazo compounds in aprotic mcdia differs considcrably from that in protic media. Further experimentation to ciarify these differences would appear to be in order.

B. Oxidation

Anodic oxidation of a variety of diazo compounds, e.g. 25–27, in acetonitrile has been shown to proceed by initial formation of a short-lived intermediate.

presumably the corresponding radical cation¹⁸⁻²⁰. The oxidation potentials of each class of compounds were found to be linearly correlated with Hammett σ^+ and $\sigma^{18, 20}$. Furthermore, the rates of reaction of members of a given class of diazo compound with benzoic acid were found to be linearly correlated with their oxidation potentials^{18, 20}. For example, the rates of reaction of substituted diazodiphenylmethanes (25) with benzoic acid were found to obey the following relationship (equation 6) 18 :

$$
\log k_{\rm R} = -6.13E_{k,\rm R} + 5.59\tag{6}
$$

where k_R and $E_{\text{A},R}$ are the rate of reaction and oxidation potential, respectively of a given diazo compound. Similar correlations were observed for other classes of diazo compounds, e.g. 26 and 27. The success of such correlations was ascribed to the fact that both processes occur at the diazo group, the position of highest electron density.

The radical cations from oxidation of diazo compounds decompose in complex fashion. From diazodiphenylmethane (19), for example, tetraphenylethylene (30) is the major product $(80\%$ yield) but benzophenone, benzpinacolone and

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benzhydrol¹⁹ are also formed. An important mechanistic clue was provided by coulometric experiments, which demonstrate that the anodic process involves a chain reaction: at an initial concentration of **19** of **0.05 M, 25** molecules of **19** react for every electron passed; when the initial concentration of **19** is **0.004 M,** only seven molecules of 19 react per electron. The products were rationalized as in Scheme 5 ^{19, 21}.

Oxidation of 19 in the presence of n -propylamine, on the other hand, consumed two electrons per molecule of 19 and afforded benzophenone *n*-propyl imine (32)

$$
(C_6H_5)_2C=NC_3H_7
$$
\n(32)

as the product¹⁰. This reaction may have synthetic utility as a route to imines (Schiff bases). Other diazo compounds which were found to undergo electrochemical oxidation to afford the corresponding olefin include the following (yields of olefin in parentheses) 21 :

A different anodic pathway appears to intervene when the diazo group is alpha to a carbonyl group²² (33) or one of its phosphorus analogues¹⁶ (34). With such compounds a complex set of products is formed, some of which apparently arise by

coupling of two molecules of substrate at the *para* position. This was established by permanganate oxidation of the neutral fraction from the electrochemical reaction, af;er removal of acidic materials. For example, anodic oxidation of benzoylphenyldiazomethane **(35)** in acetonitrile containing lithium perchlorate and sodium bicarbonate afforded benzoic acid **(27%),** benzilic acid (trace), benzil **(9%),** benzoin (? %), benzaldehyde (? %) and **a** neutral fraction which afforded diphenic acid $(17%)$ and benzoic acid $(35%)$ upon treatment with permanganate²². The coupling was rationalized as illustrated in Scheme *6.* All of the observed products can be rationalized in terms of intermediates **36** and **39** (which could also be written as a quinoid structure), though other mechanisms could be imagined. **(It** should be noted, incidentally, that there are many precedents for *porn* coupling of anodically generated radical cations²³.) The difference between the anodic behaviour of aryldiazo compounds, e.g. 19, and species such as **35,** appears to lie in the site at which the corresponding radical cations couple: coupling of two 28 species occurs at the diazo carbons, while with **36** coupling is between two para sites. Apparently the electron-withdrawing carbonyl and phosphoryl groups in **33** and **34,** respectively, inhibit coupling at the diazo carbon. while with 36 coupling is between to
rawing carbonyl and phosphoryl grou
g at the diazo carbon.
 $C_6H_5COCN_2C_6H_5$ $\xrightarrow{ee^-} C_6H_5COCN_2C$
(35) (36)

The anodic conversion of the diazo group to an n-hydroxycarbonyl radical²² (equation **7)** *(cf.* the conversion of 38 to 39) can not only explain the formation of

 $\ddot{}$

$$
N_2 \longrightarrow \begin{array}{c} N_2 \longrightarrow 0 & N_2 \longrightarrow 0 \\ \parallel 0 & -C \longrightarrow 0 \\ \parallel 0 & 0 \end{array} \longrightarrow \begin{array}{c} N_2 \longrightarrow 0 & 0 \\ \parallel 0 & 0 \\ \parallel 0 & 0 \end{array} \longrightarrow \begin{array}{c} \parallel 0 \\ \parallel 0 \\ \parallel 0 \end{array} \tag{7}
$$

benzoin and benzil from oxidation of **34,** but also explains the anodic conversion of diazoketone 40 to the dicarboxylic acid 43 (along with a trace of diketone 42)²⁴.

SCHEME 7

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40 appears to be the only aliphatic diazoketone whose anodic electrochemistry has been examined. The generality of the conversion exemplified by $40 \rightarrow 43$, and its possible synthetic utility as a means of carbon-carbon bond cleavage alpha to the carbonyl group of aliphatic ketones (via the corresponding diazoketones) remain to be established.

Elofson and coworkers obtained N-niethylpyridinium perchlorate **(44)** by clectrochemical oxidation of diazomethane in sulfolane in the presence of pyridine (equation 8) **17.** These workers suggestcd a mechanism involving successive generation

of diazomethane radical cation **(49,** niethylene radical cation **(46)** and finally methyl cation (equation **9).** It is more likely that the product **44** derives from nuclcophilic attack upon **45** (or possibly **46)** by pyridine.

$$
\begin{array}{ccc}\nCH_{2}N_{2} & \xrightarrow{-c^{-}} CH_{2}N_{2}^{\frac{1}{2}} & \xrightarrow{-N_{2}} CH_{2}^{\frac{1}{2}} & \xrightarrow{\text{II}^{*}} CH_{2}^{\frac{1}{2}} & \xrightarrow{\text{I}^{*}} CH_{3}^{\frac{1}{2}} & \xrightarrow{\text{Pyr}} & \text{44}\n\end{array}\n\tag{9}
$$
\n
$$
\begin{array}{ccc}\n(45) & (46)\n\end{array}
$$

111. THE DlAZONlUM GROUP

Electrochemical reduction of arcnediazonium salts in aqueous media exhibits some confusing features. Two polarographic \vaves arc observed, the height of the first being controlled by adsorption. Microcoulometry at thc dropping mercury electrode shows that the two waves correspond to uptake of one and four (overall) electrons, respectively. The products of preparative electrolysis at a mercury pool are either aryl mercury compounds or phenylhydrazines; which is formed depends not only on the clectrode potential, but also the rate of stirring²⁶. In aprotic media, on the other hand, arencdiazonium salts arc electrochemically better behaved. **A** one-electron wave is observed in aprotic media, *e.g.* sulfolane or acetonitrile, corresponding to the electrode process described by equation $(10)^{17}$, 27 . There is a tures. Two polarographic waves are observed,
ed by adsorption. Microcoulometry at the d
ed by adsorption. Microcoulometry at the d
at the two waves correspond to uptake of one a
ely. The products of preparative electrolys

$$
ArN_{z}^{+} + e^{-} \longrightarrow ArN^{+} \longrightarrow Arr + N_{z}
$$
\n(10)

good linear correlation²⁷ between the polarographic half-wave potentials of substituted benzenediazonium ions and Hammett σ substituent constants (σ^+ for *para* substituents capable of donating electrons by rcsonnncc). Free phenyl radicals formed by decomposition of intermediate **47** have been identified in several ways: (a) electrolysis in the presence of α -phenyl-N-t-butyl nitrone, a well-known radical trap, affords radical **48** (equation 1 I) which can be readily detected by e.s.r. spectroscopy $2⁹$, and (b) isomer ratios, total rate ratios and partial rate factors for arylation (47)

relation²⁷ between the polarographic half-wave potentials of sub-

diazonium ions and Hammett σ substituent constants (σ^+ for *para*

able of donating electrons by resonance). Free phenyl radicals formed

i

$$
C_{6}H_{5}+C_{6}H_{5}CH=N-Bu-t \longrightarrow (C_{6}H_{5})_{2}CHN-Bu-t
$$
\n(11)
\n0
\n0
\n(48)

by electrochemically-generated phenyl radicals coincide with those for phenyl radicals generated by thermolysis of benzoyl peroxide or N-nitrosoacetanilide, indicating that there is no electrophilic component to the arylation (from

10. The electrochemistry of the diazo and diazonium groups **497**

decomposition of the parent diazonium ion) and that the phcnyl radical reactivity is not influenced by the electrode surface²⁹. Yields of phenylated products are somewhat lower than with benzoyl peroxide or N-nitrosoacetanilide, but the only radical generated is the phenyl radical, unlike other routes. (Thus, benzoyl peroxide thermolysis affords products derived from both phenyl and benzoyloxy radicals³⁰.) Other advantages of the electrochemical reduction of arene diazonium ions as a Preparative route to aryl free radicals include the facts that aryl radicals may in primiple be generated ovcr a wide tempcrature range, and that diazonium ions are among the most easily reducible organic functional groups, so that a wide variety of substituents may **bt:** accommodatcd in the aromatic ring. Elofson, Gadallah and Schulz found not only that benzenediazonium tetrafluoroborate is converted to phenylpyridines in 81% yield $(\alpha : \beta : \gamma = 56 : 27 : 17)$ upon electrolysis in pyridine as solvent, but that metallic mercury alone can effect reduction of the diazonium ion (87% yield of phenylpyridines)³¹. In fact, phenylation of pyridine could be effected (though more slowly) by simply allowing the diazonium tetrafluoroborate to react directly with pyridine (92% yield of phenylpyridines). It was also found possible to phenylate pyridine N-oxide with electrochemically generated phenyl radicals. Yields of phenylpyridine N-oxides were better $(35\%, \alpha : \beta : \gamma = 89 : 1 : 10)$ than with conventional methods of phenyl radical generation³¹. hulz found not only that benzenediazonium tetrafluorobor

emylpyridines in 81% yield ($\alpha : \beta : \gamma = 56 : 27 : 17$) upon elso

solvent, but that metallic mercury alone can effect reduction

1 (87%, yield of phenylpyridines)³¹

Elofson and Gadallah have shown that it is possible to effect electrochemical cyclization of the appropriate diazonium α -phenylcinnamic acids in nearly quantitative yield 32 (equation 12). They also found it possible to effect electrochemical

cyclization of diazotized 2-aminobenzophenones *(49)* to fluorenones **(51)** (equation 13). Surprisingly, the same process was found to occur thermally, and it was suggested

that this occurs by intramolecular reduction of the diazonium group by the other ring (an intramolecular charge-transfer process). Depending on the experimental conditions, greater or lesser amounts of benzophenones **(52)** and diarylmercury

compounds **(53)** accompanied products from decomposition of **49.** Yields of **52** were highest in aprotic solvents, where hydrogen abstraction by radical *50* is easiest ; conversely, yields of fluorenones were best in protic media, or in aprotic thermolyses of **49,** which therefore apparently do not involve radicals *50* as discrete intermediates. Finally, reduction of **49** by iodide ion in aprotic media afforded 2-iodobenzophenones **(54)** in good yield ; electron transfer from iodide ion affords *50* and an iodine atom, which then couple in the solvent cage.

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CHAPTER 11

The influence of the diazo and diazoniurn groups

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1. INTRODUCTION

A major interest in organic chemistry is the effect of structural change on reactivity, especially the effect of change in structure of one part of a molecule on reactivity at some other part of the molecule. The influence of such a change in structure is rathcr conventionally divided into steric effects and electronic effects, although these are neither totally suficient nor completely separable. An example of the insufficiency lies in the existence of secondary isotope effects, where the effect arises exclusively from a difference in mass of two substituents having identical electronic effects and sizes. An example of the inseparability of steric and electronic effects can be found in the efforts to understand the effect of *orrho* substituents on aromatic side chain reactivity, where, despite strong efforts, no reasonable and consistent electronic effects have been found by incthods often capable of separating stcric and electronic effects in other cases.

Wc can nevertheless begin to understand the electronic part of substituent effects by combining qualitative considerations of inductive effects from electronegativities and formal or real charges with resonance effects, alternatively estimated by various molecular orbital methods, of which the semi-empirical methods now appear to **be**

the most quantitative. Similarly, steric effects can be related to Van der Waals' interactions by qualitative or quantitative methods.

On these bases, the diazo **(I)** and diazonium ion **(Xi)** groups would be expected to have **very** strong influences, because of the formal cliarges in the first, the real positive charge in the second, as well as the unsaturation which gives the possibility of extensive resonance interaction in both. The direction of the effect is somewhat equivocal in the diazo compounds, since even the sign of the charge nearest to the rest of the molecule is uncertain, depending on the relative weights of contributing structures **Ia** and Ib. Structure Ic is unimportant partly because it has the wrong

$$
\sum_{\substack{\\n=1\\n \text{ (la)}}}^{\lambda} \frac{1}{n} = \overline{N} \iff \sum_{n=1}^{\infty} \frac{1}{n} = N \iff \sum_{n=1}^{\infty} \frac{1}{n} = \overline{N} \quad \text{C}-\overline{N} = N
$$
\n
$$
\sum_{n=1}^{\infty} \frac{1}{n} = N \quad \text{(Ib)} \quad \text{(Ic)} \quad \text{(II)}
$$

geometry. In the diazonium ion the substituent effect is unequivocally that of electron withdrawal; it will be acid strengthening, it can be expected to have a positive Hammett *o* or Taft *o*₁.

The experimental realization of these expectations is another problem. In order to determine the effect of the diazo or diazonium groups on a reaction elsewhere in the molecule, it is necessary to have such **a** reaction occur, leaving these nitrogen functions untouched. The extraordinary reactivity of these two functional groups makes all such studies difficult, for the majority of the reagents will attack these very reactive functional groups before reacting elsewhere in the molecule. The low thermal stability of diazo and diazonium compounds sometimes niakes it uncertain whether or not the loss of nitrogen has occurred before, after or in concert with a reaction elsewhere. We therefore lack the wide range of information on substituent effects for the groups, and for the diazo groups there is virtually no quantitative information. However, there are somc available data, mostly qualitative, and these will be presented. The effect of the diazo group will be discussed first, then that of the diazonium ion group. Three sourccs have been especially useful in this compilation; the valuable book of Zollinger', the review on diazo compounds in acid by More O'Ferrall², and a review by Regitz³ which covers several reactions of diazo compounds without loss of this functional group.

II. INFLUENCE OF THE DIAZO GROUP

A. The Carbonyl Group as a Model

The first question we can attack on the relative importance of structures Ia and **Ib** is to **ask** whether a doubly bonded structure rather like a Schiff base or a ketone is a better representation than a carbanion. We shall see that the carbanion structure Ib is more descriptive of the behaviour than any carbonyl-like one. In fact, only one reaction turned up to suggest the carbonyl-like structure. The reaction (equation 1) was postulated⁴ on the basis of the formation of PhCHOHCN₂CO₂Et and ethyl

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\nated⁴ on the basis of the formation of PhCHOHCN₂CO₂Et and ethyl
\n
$$
N^{-}
$$
\n
$$
HO
$$
\n
$$
N^{+}
$$
\n
$$
PhC-CO2Et + PhCO2Et (1)
$$
\n
$$
Ph
$$

benzoate from the attcmptcd condcnsation in basic ethanol of benzil with ethyl diazoacetatc. Here the diazo group may have stabilized the intermediate carbanion, **111.**

B. The Carbanion Model; Protonation

The analogy of diazo compounds to carbanions is relatively better supported. Diazo compounds rcact very generally with acids?, usually however giving diazonium salts too unstable to allow the drawing of quantitative conclusions. Other electrophiles also attack diazo compounds, and these reactions also contribute to our information. diazo compounds to carbanions is relatively better supported.
eact very generally with acids², usually however giving diazonium
o allow the drawing of quantitative conclusions. Other electro-
diazo compounds, and these

One reaction showing the protonation unequivocally but casting little light on the equilibrium constant is thc reaction of trifluorodiazoethane with fluorosulfonic acid (equation 2). The conversion to the diazonium salt, which is stable at -60° C

$$
F_{J}CCHN_{2}+HO_{3}SF \longrightarrow F_{3}CCH_{2}N_{2}+O_{3}SF
$$
 (2)

is quantitative'. Some other diazo compounds are acid stable to some extent, but it has not been established whether these correspond to stable diazonium salts or very weakly basic diazo compounds. They include diethyl diazomalonate, stable *to* methanolic H_2SO_4 ⁶, and $(RSO_2)_2CN_2$, which decompose at rate slow enough to measure in chlorosulphonic acid?.

At a more familiar level ethyl diazoacctate hydrolysis rates correlate with *H"* up to the fastest rates measureds. This suggcsts that the equilibrium extent **of** protonation does not cxcecd a few percent in the most highly acidic media used. The rate of deprotonation of \overrightarrow{N}_2 CH₂CO₂Et has not been measured, but the relative rates of proton and nitrogen loss come out of studies on the acid-catalysed exchange of tritium-labelled diazoacetic ester⁹. One can attempt to estimate the acidity of N₂CH₂CO₂Et, either by using a substituent constant derived from the aromatic series (discussed below), or by correcting the pK_a of $N=C-CH_2CO_2Et$ for an electronegativity and clcctrostatic changc. This leads, depending on how the various corrections are applied, to $pK_a = -7 \pm 5$, a not very useful or firmly based number, but at least consistent with the available data. Kreevoy¹⁰ has estimated from kinetic $\frac{1}{2}$ considerations based on the decomposition of diazoacetate ion that for $N_2CH_2CO_2^-,$ $pK_a \geq 0$.

The protonation of diazo compounds which do not have these base weakening and carboniuni ion destabilizing substituents is even more uncertain insofar as the equilibrium constants arc conccrned. Thc details of protonation of diazo compounds and the subsequent reactions are discussed by More O'Ferrall²; in general, the expected stability of the carbonium ion determines whether or not the protonation is significantly reversible. Thus under ordinary circumstances only diazomethane shows exchange in acidic D₂O competing with nitrogen loss¹¹. Clearly R₂CHN⁺ and R,CN2 arc interconvertible, but **we** have not bcen able to establish an equilibrium constant even in one casc within orders of magnitude. In summary, the protonation does occur, though hardly with the facility that would be associated with carbanions.

C. The Carbanion Model; Electrophiles

Diazo compounds also react with other electrophiles. Condensations using base catalysis will be discussed later, but some react without any base catalysis. Ethyl diazoacetate, diazoacetonitrile, trifluorodiazoethane¹² and nitrodiazomethane¹³ are
nitrated by N_2O_5 , presumably by way of the stable but strongly acidic diazonium salts (equation 3). Diazomethane is chlorinated¹⁴ by *t*-butyl hypochlorite at -100 °C,

Edward S. Lewis

\nEdward S. Lewis

\n[₂O₅, presumably by way of the stable but strongly acidic diagram

\n13). Diazomethane is chlorinated¹⁴ by *t*-butyl hypochlorite at
$$
-100^{\circ}
$$
C, NO_2

\nXCHN₂+N₂O₃ — \rightarrow X- $C-N_2^+$ +NO₃ — \rightarrow XCN₂

\n1

\nX = CO₂Et, CN, CF₃, NO₂

\numed chloromethanediazonium ion is protected from decomposition

\now temperature and the very basic counter ion, *t*-BuO-
\nelectrophilic attack on diazomethane and other diazo alkanes is the acid chlorides¹⁵ or anhydrides¹⁶, yielding diazoketones (equation 4).

\nRCOCl+2 CH₂N, — \rightarrow RCOCHN₂+CH₃Cl+N₂

\n[4]

\n[e of diazomethane is believed to serve as a sink for the proton of the um salt, RCOCH₂N⁺, rather than as a base catalyst at an earlier stage.

here the presumed chloromethanediazonium ion is protected from decomposition both by the low temperature and the very basic counter ion, t -BuO⁻.

A familiar electrophilic attack on diazomethane and other diazo alkanes is the reaction with acid chlorides¹⁵ or anhydrides¹⁶, yielding diazoketones (equation 4).

$$
RCOCI + 2 CH2N2 \longrightarrow RCOCHN2 + CH3Cl + N2
$$
 (4)

The extra mole **of** diazomethane is believed to serve as a sink for the proton of the initial diazonium salt, $RCOCH₂N⁺$, rather than as a base catalyst at an earlier stage.

The formation **of** polymethylene by the reaction of diazomethane with boron fluoride¹⁷ is also evidence for electrophilic attack on the diazo centre, even though the immediate product is not observed. Similarly the reaction of diazo compounds in the absence of basic catalysts, with ketones or aldehydes, is an example of electrophilic attack, but with a non-isolable diazonium ion product¹⁸.

D. **Acidity** *of* **Diazo Compounds**

Diazo compounds not only act as bases but they are also pcrceptibly, although not strongly, acidic. Diazomethane reacts with methyl lithium to give a lithium derivative¹⁰. Ionization is also possible with triphenylmethylsodium²⁰, which puts a lower limit on the acidity of diazomethane, since pK_a of triphenylmethane = 33^{21} . Ethyl diazoacetate has also been converted to the lithium salt²², and ionization under relatively mild conditions is suggested by the facile hydrogen isotope exchange in base9, compared to ethyl acetate which suffers hydrolysis much faster than exchange. Furthermore, ethyl diazoacetate and other acyldiazo compounds undergo aldol condensation with a variety of compounds with dilute ethanolic sodium hydroxide⁴. Diazosulphones even add to enamines²³, presumably via an initial proton transfer. **Example 12**
 XCOV
 XCOV

The perceptible stability of the anions $N_2 \overline{C} \overline{C}$ is also indicated by reverse condensations observed in a number of cases, mostly of diacyldiazomethanes (equation *5).* The mechanism is presumably that shown, and the diazo compound

0

formed is often the one which would be expected to give the most stable anion. **A** number of examples are listed by Regitz3, but cxcept for showing that acyldiazomethane anions have a stability comparable or greater than that of ordinary enolate ions, quantitative conclusions are impossible.

The chemical evidence is then that the diazo carbon is definitely nucleophilic, and is perceptibly, although not strongly, basic. **It** is also significantly more acidic than a comparable methyl group, although the magnitude of the enhancement of acidity is not known.

E. Remote Effects of the Diazo Group

Before going on to physical evidence for the charge distribution about the diazo compounds, it is appropriate to comment briefly about reactivity more remote from the diazo group. An acid strengthening effect on the β carbon is suggested although not convincingly demonstrated by the cleavage mentioned earlier⁴. A few reactions at positions remote from the diazo group have been observed, but without rate measurements they are hard to interpret. Thus, potassium diazoacetate is prepared by saponification of the ethyl ester²⁴ by concentrated potassium hydroxide under conditions which would also be expected to saponify ethyl acetate, but relative rates are not known.

F. Effect of the Diazo Group **on** *lJC-n.m.r., Spectra*

A possible approach to electronic effects in diazo compounds is the chemical shift in the n.m.r., for extensive electron withdrawal should reduce the nuclear shielding, and supply should increase it. There are few proton spectra for diazo compounds available, but a number of **I3C** spectra of diazo compounds have been taken²⁵. Several compounds, all diazo ketones or esters, are here presented, with chemical shifts (in p.p.m. downfield from **TMS),** together with some model compounds, taken from Breitmeyer and Voelter²⁶.

These chemical shifts allow us to draw some conclusions, limited however by the small size of the sample. The important effects are on the diazo carbon, on the adjacent carbonyl carbon, and there is one example of a perceptible effect on thc other carbon flanking the diazo group. More distant groups, **as** the ethyl group in **VI** and **VII,** or the phenyl group in **VIII** and **IX,** seem to be relatively uninfluenced (hcre dilferences of less than **2** p.p.rn. are ignored, largely because the author is not competent to judge the sources of error in the various spectra). The carbonyl frequency is uniformly at higher field for the diazo compound than for the models. Since (in the one case of the tertiary alcohol, **X,** shown) the shift on the other side

of the diazo group is in the other direction, this is probably not a consequence of an inductive effect of the diazo group; instead it probably represents a specific conjugation similar to that in α, β -unsaturated ketones, which also causes a higher field shift of the carbonyl frequency. Alternatively, the extra shielding could be a consequence of the formal negative charge on the diazo carbon (structure **Ib).**

The chemical shift of the diazo carbon is especially interesting; any model with neutral carbon bonded with sp² hybridization to carbon, oxygen or nitrogen would predict chemical shift more than 100 p.p.m. from TMS. The observed shift of only **40-60** p.p.m. appears to require the addition of extra shielding, such as would bc associated with a substantial negative charge associated with this carbon, for example through the important contribution of **Ib.**

In sum, the c.m.r. chemical shifts are in agreement qualitatively with the chemical evidence, namely that diazo compounds have a significant partial negative charge on carbon.

G. Special Diazo Structures

Certain diazo compounds have such important contributions of diazonium-like structures that they can really be thought of as members of one class as **well** *2s* of the other. These include the diazo oxides, XI, which are the conjugate bases of *ortho-* or para-hydroxydiazonium ions, which we shall consider briefly in a later section. Another example is diazocyclopentadiene, XII, which appears to be virtually an aromatic molecule, and undergoes electrophilic substitution with facility²⁷.

111. THE DlAZONlUM ION GROUP

A. Coverage

The effect of the diazonium ion group will here be limited to discussion of aromatic diazonium salts. The rclatively few aliphatic cases of any stability contribute little to our knowledge. There are also sonie inorganic 'diazonium salts' of tlie form L_nMN_2 or metal nitrogen complexes, which share little of the chemistry of the aromatic compounds; they will not be considered hcre.

Aromatic diazonium ions, ArN_2^+ , have been extensively studied since their discovery more than a hundred years ago. Nevcrthelcss, most of the chemistry involves either loss of N_2 , and replacement by various groups (such as OH in the hydiolytic dccomposition, halogen in the Sandmcyer reaction among many), or alteration of the N_2^+ group as in the reduction to phenylhydrazine or coupling to form ArN2X (where **X** may be carbon, oxygen, nitrogen. sulphur, phosphorus, among others). With the one exception of partial reaction of tetrazotized aromatic diamines, discussed below in Section **11I.E, all** these reactions are outsidc thc scope **of** this chapter.

B. **The Hammett Substituent Constants**

Among the reactions suitable for discussing the effcct of thc diazonium ion group as a substituent are acidities and basicities of acids and bases containing the N_2^+ substituent, and the rates of various rcactions, including electrophilic and nuclcophilic substitution on the aromatic ring as long as the diazoniuni ion group remains attached after the reaction. The diazonium ion group was recognized as a powerfully electron-withdrawing substituent for as long as that phrase has had meaning, but the quantitative evaluation of the cffect was not attempted until a number of acid strengths and reaction rates were measured with the goal of establishing the Hammett substituent constant σ^{28} . Table 1 shows some of the results obtained for various

Group	Reaction	σ_m	σ_{τ}	σ_n^-
$+N_2C_6H_4OH$	Acid ionization	< 2.1		3.03
$+N_{2}C_{6}H_{4}NH_{3}+$	Acid ionization			$3-43$
$+N_2C_6H_4CO_2H$	Acid ionization	1.76	1.91	
$+$ N ₂ C ₆ H ₁ CH ₂ CO ₂ H	Acid ionization		2.18	
$+N$ ₂ C ₆ H ₃ COCH ₃	Bromination rate	> 0.2	>1.1	

TAnLE 1. Substitucnt constants for the diazonium ion group2'

cases. Thc absence of entries or estimatcs of one limit only reflects experimental problems, almost always due to side reactions which destroy the diasoniuni group. The nature of the vzrious uncertainties is discussed in the original papcr.

The large difference between σ and σ ⁻ implies a large resonance interaction between the N_2^+ and the p-O⁻ (as shown in **XI**). This extra stabilization is confirmed by the low reactivity of these 'diazo oxides' in coupling reactions²⁹ (among others), as well as by the rather quinoid bond distanccs shown in **XIII3O.** The distances shown are averages of chemically equivalent distances in two independent molecules.

(XIII)

The very large substituent effect is also evident from the acidity of o-carboxybenzenediazonium ion²⁸, which has a $pK_a = 1.47$; it is almost ten times stronger than the *para* isomer, but of course no substituent constant should be assigned.

In spite of the fact that the results in the Table weye determined more than **15** years ago, there has been very little revision. The value for $\sigma_m = 1.76$, based on a single measurement, has been reported to be confirmed on the basis of studics of ionization equilibria of some diazodiazoles and other heterocycles³¹. A study of the ¹³C-n.m.r. of diazonium salts is reported³² to give chemical shifts simply related to σ and consistent with the old values.28 **A** study33 of the **35CI** nuclear quadrupole resonance spectra of some chlorobenzenediazonium ions is reported to show substantial electron withdrawal by N_2^+ , but in the order $o > m > p$. This order is stated to be consistent with the chemically determined substituent effects since the nqr chemical shifts are bclieved to be insensitive to resonance effects, but sensitivc to inductive effects. A further confirmatory value of σ_p is presented in Section III.E.

The substituent effects in the Table can be subjectively weighted to give $\sigma_m = 1.7$, $\sigma_p = 2$ and $\sigma_p = 3$, but it should be emphasized that they are only valid in aqueous solution, sincc charged and uncharged substituents will behave differently as the medium is changed. It is nevertheless clear that the N_{τ}^{+} group has a more powerful electron-withdrawing effect than any other single faniiliar group, and this is reasonable, for it combines the resonance effect of thc isoelectronic cyano group $(\sigma_p = 0.66)$ with the pure charge effect of the trimethylammonium substituent $(\sigma_p = 0.82)$. It is perhaps the plausibility of the very large σ values for N₂⁺ which has led to a general acceptance of these values, rather than conviction from an overwhelming mass of data.

C. Efectrophific Substitution on Diozoniurn Salts

Benzenediazonium ion has been nitrated with difficulty, and nitration occurs mostly in the *meta* position. This is consistent with Hammett's observation³⁴ that substituents with $\sigma_p > \sigma_m$ are *meta* directing. There are no direct data on the rates of electrophilic substitution on benzenzdiazoniuni ion, although the following, rather flawed, argument suggests that in one case it may be slower than the attack on nitrobenzene. When benzenediazoniuni ion is decomposed in nitrobenzene, m -nitrobiphenyl is an important product. When benzenediazonium fluoroborate is heated in trifluoroethanol and nitrobenzene³⁵, only 6% of the product is nitrobiphenyl, but most of the remainder is the expected fluorobenzene and phcnyltrifluoroethyl ether. If this electrophilic arylation had been able to attack benzenediazonium ion as easily as nitrobenzene, then fluorobiphenyl or trifluoroethoxybiphenyl should have been observed. The absence of these products is not convincingly demonstrated, but again it is plausible and hence believable. The argument is somewhat strengthened because arylation with dccomposing diazonium salts is believed to be very unselective, as would be expected for the postulated phenyl cation intermediate,[†] although a small discrimination among neutral organic solvents is $observed³⁵$.

The much more facile electrophilic attack on diazocyclopcntadiene has been mentioned above.

D. Nucleophilic Substitution Activoted by the Diozonium Group

In contrast to the rare and difficult electrophilic attacks on diazonium salts there are numerous examples of nucleophilic attack. Only those leaving thc diazonium group intact, which still comprise a large body of data, arc relevant here. The activation of nucleophilic aromatic substitution by an o - or $p-N_2^+$ group had been observed very early, especially by Hantzsch⁴² and by Meldola⁴³. Miller⁴⁴ identified the N_{τ}^{+} group as more strongly activating than any other substituent. Quantitative studics on the rates of attack of thiocyanate *orflio* and *porn* to thc diazonium ion group were undertaken⁴⁵. The reaction is facile for halogen and the second-order ratc constants arc much faster in lower dielectric constant medium, as expccted for this anion--cation reaction. The logarithm of the ratc constants correlated well with

t The actual intermediacy of the phenyl cation in the solvolysis of diazonium ions is somewhat controversial and is complicated but illuminated by the simultaneous occurrence of several reactions. The controversy centres around whether^{35, 36} or not³⁷ nucleophiles are present in the rate-determining step, **but** there is no controversy over the description of all transition states as closely resembling the phenyl cation, and the consequent unselectivity of the reaction, which discriminates poorly between halide ions and water^{36, 38}, and allows attack on very poor nucleophiles such as chlorobenzene³⁹, fluoroborate ion⁴⁰ or molecular nitrogen⁴¹.

the reciprocal of the diclectric constant for a range of aqueous f-butyl alcohol solvents, and also correlated well with the solubility of **p-chlorobenzenediazonium** fluoroborate in the same solvents. This strong and well-understood solvent dependence emphasizcs the problem of a substituent constant for a charged group and makes a direct comparison of the activation by N_2^+ with that of one or more nitro groups generally futile.

In the mostly aqueous systems usually studied, the diazonium group is, however, very powerfully activating by comparison with any other group or combination of groups. In this qualitative vein, we note that in the reaction of thiocyanate ion with p -nitrobenzenediazonium ion in aqueous t -butyl alcohol, the replacement of nitro is more important than that of the diazonium group⁴⁵, behaviour which it would be unreasonable to attribute to the nitro group being a better leaving group. **A** similar but even more striking observation is that **2,4,6-trinitrobenzenediazonium** ion and related substanccs suffer hydrolysis (at sufficiently high pH) with loss of nitrite by attack on carbon *orfho* to one diazonium ion group, rather than losing nitrogen by attack on the carbon activated by as many as three nitro groups⁴⁶.

It is necessary to be careful about the nature of the mechanism involved in these substitutions. Thus in apparent contradiction to the results with thiocyanate, at higher temperatures **p-nitrobenzenediazoniuni** ion reacts with aqueous bromide ion to give a mixture of p -nitrophenol and p -nitrobromobenzene. No isolated product was assignable to nitro group loss⁴⁷. This behaviour is now believed³⁶ to be associated with the acidity-independent solvolysis rather than to an activated nucleophilic attack. The perceptible acceleration¹⁷ by bromide ion is not much greater than that observed with other diazonium salts, and has been explained³⁶ using transition states strongly resembling the phenyl cation (see footnote above). Thus there is no unequivocal evidence of nucleophilic substitution of N_2^+ activated by a nitro group. Substitution of N_2^+ activated by a second N_2^+ group is, however, well established and is discussed below.

E. Chemistry **of** *Bisdiazoniom* **Salts**

It was mentioncd in Section **1II.A** that the vcry rich chemistry *of* aromatic diazonium salts nearly always involved destruction of the N_z^+ group, and could not therefore give information about the influence or substituent effect *of* this group. This statement has t\vo exccptions. The first *of* these is that the reciprocal relation between ρ and σ first pointed out by Hine⁴⁸ has been developed by Sager and Ritchie¹⁹ to the point that the measurement of ρ for a reaction is related to the change (in substituent effect) of the reaction site. Thus the rather extensive study of substituent effects on diazoniuni salt cquilibria has been rendered understandable by using a large positive σ for the diazonium ion group⁵⁰. The second is that for any reaction of the diazonium group, for example the conversion of N_2^+ to *Y* (equation *6),* one of the possible substitucnts X is a second diazonium group. This

$$
XC_{6}H_{4}N_{2}^{+} \longrightarrow XC_{6}H_{4}Y
$$
 (6)

allows a host of reactions to be studicd, subject only to the limitation that only one of the two diazonium groups must react. This limitation is not very restrictive: of the three most accessible tetrazonium salts, tetrazotized p-phenylenediamine allows in nearly all cases the isolation of the reaction of one group only; tetrazotized m -phenylenediamine is complex⁵¹ in its behaviour and has not been extensively studied, and tetrazotized benzidine is, in the dye industry, regularly coupled at one end before the other⁵², and would therefore be suitable for quantitative study, although it has not yet been done.

Tetrazotized p -phenylenediamine is subject to nucleophilic displacement of onc diazonium ion group by CI-, Br- and SCN- **53;** sccond-order kinetics arc followed and the products, p-chlorobcnzenediazonium ion, etc., are quite stable under the reaction conditions. Iodide ion also reacts, but a very fast free-radical mechanism is involved, and we can only say that the $p-N_2^+$ group accelerates the nicchanism to allow it to bc scen and to overwhelm nuclcophilic attack. Azide ion also attacks very rapidly, but the mechanism probably does not involve carbon attack⁵⁴. The powerful activating effect of the $p-\mathbb{N}_2^+$ group is clearly illustrated, and the fact that N_2^+ can be a leaving group in an activated nucleophilic substitution is also demonstrated.

The tetrazonium salt is also convertible to a monodiazotate.⁵⁵ Because this is bclieved to have thc *arzti* configuration, the equilibrium is not readily compared to the litcrature values *56, 5i* for cquilibrium diazotate formation which involvc the *syn* isomers. However, the protonation equilibrium between the monodiazotate and its conjugate acid is directly comparable to mcasurenicnts of this reaction with other anti-diazotates. One determination gives⁵⁸ for this equilibrium $\rho = 1.17$, another gives⁵⁹ $\rho = 1.45$, and combining these with the measured equilibrium constant⁵⁵ give then for σ_p the values 2.0 and 1.6 respectively. This is in reasonable agreement with the substitutcd benzoic acid values prcsented above.

is at first sight incompatible with the extreme electron-withdrawing character of the N_2^+ group, since electron-withdrawing substituents normally retard the rate of this phenol-forming reaction. However, the hydrolysis turns out to be pH dependent⁵⁵. In fairly strong acidic media this special mechanism is suppressed and the tetrazonium salt now has thc expected high stability. The facile 'dediazonization' or hydrolysis of the p -phenylene-bis diazonium salt^{55, 60}

The tetrazonium salt shows several very facile free radical reactions⁶¹, notably reduction by alcohols, and it is clear (and not surprising) that the second diazonium group facilitatcs one-electron rcduction of the first, although we have little to compare this with quantitativcly, cspccially since in this reaction there is considerable net reduction of both diazonium groups. The details of chain initiation and termination proposed in this study of **the** alcohol reduction have been shoun to bc subjcct *to* a more reasonable interpretation by introducing a new termination mechanism⁶².

Finally, we should interpret again the rather low σ (1.3) derived from the rate of coupling of the tetrazonium salt with 2-aminonaphthalene-6-sulphonic acid²⁸, based upon the value of ρ for this coupling of $+4.26$ ⁶³. Some coupling rates of diazonium salts with various compounds have given curved Hammett plots⁶¹, and this deviation may correspond to the predicted concavc downward shape of sutliciently wide-range Hammett plots⁶⁵. The deviation may also arise from a shift of rate-determining step in this known two-step reaction⁶⁶.

IV. CONCLUSIONS

The effect of the diazo group as a substituent has been shown to be not very large and of ambiguous dircction, since it is both acid strengthening and base strengthening. The cffccts have not bcen detected cxcept very close to the diazo group.

The effect of the diazonium ion group is quantitatively established, it has a very strong acid-strengthening, electron-withdrawing effect. It is, by virtue of its charge, a substituent with a strong solvent dependence of its substituent effect and one can therefore imagine some effects of unprecedented magnitude, but even in water it has more influence than any other known substituent.

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