ELECTROCHEMICAL STUDIES OF BENZOPHENONE
AND FLUORENONE IMINES, AMINES AND
DIPHENYLDIAZOMETHANE

by

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# TABLE OF CONTENTS

| LIST OF FIGURES | iii |
| LIST OF TABLES | iv |

## I. INTRODUCTION

- References | 1

## II. EXPERIMENTAL

- Electrochemical Instrumentation | 7
- Cells, Electrodes, and Electrolysis Procedures | 7
- Chromatography | 8
- Chemicals | 9
- References | 12

## III. ELECTROREDOX OF DIPHENYLDIAZOMETHANE (Ph₂CH₂)

- IN ACETONITRILE AT A HANGING MERCURY DROP ELECTRODE | 13
- Introduction | 13
- Results and Discussion | 14
- Conclusions | 19
- References | 21

## IV. ELECTROREDUCTION OF FLUORENONE IMINE, 9-AMINOFLUORENE,

AND N,N-DIMETHYLFUORENYLAMINE | 22

- Fluorenone Imine | 22
  - Cyclic Voltammetry | 22
  - Coulometry | 25
  - Chronoamperometry | 27
  - Effect of Diethyl malonate | 28
  - Comparison of the Redox Behavior of Fl=NH and Fl=NPh | 29

- Benzophenone Imine | 35
# TABLE OF CONTENTS (continued)

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>9-Aminofluorene</td>
<td>36</td>
</tr>
<tr>
<td>Cyclic Voltammetry</td>
<td>36</td>
</tr>
<tr>
<td>Proposed Reaction Scheme</td>
<td>36</td>
</tr>
<tr>
<td>N,N-Dimethyl-9-aminofluorene</td>
<td>42</td>
</tr>
<tr>
<td>Comparison of these results with those of previous workers</td>
<td>50</td>
</tr>
<tr>
<td>References</td>
<td>52</td>
</tr>
<tr>
<td>ACKNOWLEDGEMENTS</td>
<td>53</td>
</tr>
<tr>
<td>VITA</td>
<td>54</td>
</tr>
</tbody>
</table>
# LIST OF FIGURES

<table>
<thead>
<tr>
<th>Figure</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>III-1</td>
<td>Cyclic Voltammograms of Ph₂CN₂ in AN at Platinum and Hanging Mercury Drop Electrodes</td>
<td>18</td>
</tr>
<tr>
<td>IV-1</td>
<td>Cyclic Voltammograms of F1=NH in DMF and in the Presence of Diethyl malonate</td>
<td>24</td>
</tr>
<tr>
<td>IV-2</td>
<td>Cyclic Voltammograms of F1NPh in DMF and in the Presence of Added Base</td>
<td>31</td>
</tr>
<tr>
<td>IV-3</td>
<td>Cyclic Voltammograms of F1HNH₂ in DMF - Effect of Scan Rate</td>
<td>39</td>
</tr>
<tr>
<td>IV-4</td>
<td>Chronoamperimetric Data for F1HNH₂, ( E = -2.1 ) V</td>
<td>41</td>
</tr>
<tr>
<td>IV-5</td>
<td>Chronoamperometric Data for F1HN(CH₃)₂, ( E = -2.2 )</td>
<td>44</td>
</tr>
<tr>
<td>IV-6</td>
<td>Cyclic Voltammograms of F1HN(CH₃)₂ in DMF</td>
<td>46</td>
</tr>
<tr>
<td>IV-7</td>
<td>Cyclic Voltammogram of F1HN(CH₃)₂ in DMF - Effect of Added Azobenzene</td>
<td>49</td>
</tr>
</tbody>
</table>
# LIST OF TABLES

<table>
<thead>
<tr>
<th>Table</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>II-1</td>
<td>HPLC Eluting Solvent Systems</td>
<td>10</td>
</tr>
<tr>
<td>II-2</td>
<td>Compounds Synthesized for Study and the References for the Synthetic Procedures Followed</td>
<td>11</td>
</tr>
<tr>
<td>IV-1</td>
<td>Coulometric Data and Product Studies for the Controlled-Potential, Electrolytic Reductions of Fluorenone and Benzophenone Imines</td>
<td>26</td>
</tr>
</tbody>
</table>
I. INTRODUCTION

Beginning in 1976, our laboratory, in conjunction with the research group of Professor Richard N. McDonald, began an extensive study of a class of reactive intermediates termed "hypervalent radicals." The stepwise loss of $R$ groups from a tetravalent carbon substrate yields nine molecular fragments of neutral or unit. Four of these intermediates have received extensive study: carbanions ($R_3C^-$), carbonium ions ($R_3C^+$), free radicals ($R_3C^*$), and carbenes ($R_2C$). In contrast, very few studies of the other five possible intermediates (termed hypervalent radicals) have been undertaken.

One of these five, the carbene anion radical ($R_2C\cdot$), appeared the most promising source of new information for several reasons. First, this species could conceivably act as a base, a free radical, a nucleophile, or an electron transfer agent. Secondly, since all bonding and nonbonding molecular orbitals are occupied, the possibility of skeletal rearrangements should be minimized, if not eliminated. And finally, since both gas phase and solution studies were planned, reports of the generation of carbene anion radicals in the gas phase\textsuperscript{2} and by electrochemical reduction in solution\textsuperscript{3,4} were promising evidence of the feasibility of studying this intermediate.

The diphenylmethylene anion radical (Ph$_2$C\cdot) and the fluorenylidene anion radical (Fl\cdot) were chosen for the initial studies since it was felt that the electronic configuration at the divalent carbon center would be different for these two species. Experimental results\textsuperscript{2} and molecular orbital calculations\textsuperscript{2,5} describe the electronic configuration of the ground state of
simple carbene anion radicals, $\text{H}_2\text{C}^\cdot$, $\text{F}_2\text{C}^\cdot$, and $(\text{CH}_3)_2\text{C}^\cdot$, as $\sigma$-anions and
$\pi$-radical ($\sigma^2\pi^1$). It was felt that $\text{Ph}_2\text{C}^\cdot$ should also have this ground state configuration with the phenyl rings twisted about their $\text{C}_1-\text{C}$ bonds in order to relieve nonbonded repulsions. Both experiment$^6$ and calculations$^7$ showed that the ground state electronic configuration of cyclopentadienyldienide anion radical ($\sigma^2\text{C}_5\text{H}_4^-$) is $\sigma^1\pi^2$. Since $\text{Fl}^-$ is structurally similar to $\sigma^2\text{C}_5\text{H}_4^-$, the ground state electronic configuration at $\text{C}_9$ should also be $\sigma^1\pi^2$.

Substrate structures where $\text{R}_2\text{C}^-$ could result from a single one-electron step were deemed desirable. Possible precursors to $\text{R}_2\text{C}^-$ that require three successive one-electron reductions (i.e. gem-dihalo alkanes) were ruled out because they give intermediates that potentially would react before giving rise to $\text{R}_2\text{C}^-$. Generally, substrates that can be photolyzed or thermalized to give carbenes were felt to be potential precursors to $\text{R}_2\text{C}^-$ upon reduction (eq. I-1 and I-2).

$$\text{hv}, \Delta \rightarrow \text{R}_2\text{C}: + \text{N}_2$$  (I-1)

$$\text{R}_2\text{CH}_2 \rightarrow \text{R}_2\text{C}^- + \text{N}_2$$  (I-2)

To facilitate the fragmentation, the non-carbene anion radical fragment should be thermodynamically stable, such as $\text{N}_2$. Thus, diazooalkanes were a logical choice for potential $\text{R}_2\text{C}^-$ precursors.

Initial research on "hypovalent radicals" began in our laboratory with the electrochemical reduction of diazodiphenylmethane ($\text{Ph}_2\text{C}^-\text{N}_2$) and 9-diazofluorene ($\text{FlN}_2^-$) at a platinum cathode in dimethylformamide (DMF) with $0.1 \ F(\text{n-Bu})_4\text{NiClO}_4^-$ (TBAP) as the supporting electrolyte.$^8,^9$ McDonald and
coworkers have since reported on the chemical reductions of these species, the chemical generation of \( \text{Ph}_2\text{C}^- \) and \( \text{F}^- \) from certain epoxides, and the generation and ion-molecule reactions of anion radicals in a flowing afterglow apparatus. 

Our electrochemical studies of \( \text{Ph}_2\text{CN}_2^- \) concluded that (1) the anion radical \( \text{Ph}_2\text{CN}_2^- \) once formed is very short-lived, but its lifetime (at \( 0^\circ \text{C} \)) is sufficient to be partially trapped by proton donors to give \( \text{Ph}_2\text{C}^-\text{NNH}_2 \), (2) protonation of \( \text{Ph}_2\text{CN}_2^- \) occurs only on terminal nitrogen, thereby precluding "ambident" behavior of this species in this and other reactions, (3) the carbene anion radical, \( \text{Ph}_2\text{C}^- \), is formed by loss of nitrogen from \( \text{Ph}_2\text{CN}_2^- \) and is the principal, if not the only, precursor to \( \text{Ph}_2\text{CH}^- \), (4) the coupling of \( \text{Ph}_2\text{CH}^- \) with unreacted \( \text{Ph}_2\text{CN}_2 \) cannot involve initial electron transfer followed by coupling of \( \text{Ph}_2\text{CH}^- \) with \( \text{Ph}_2\text{CN}_2 \), (5) \( \text{Ph}_2\text{CH}^- \) is the principal chain-carrying species in the formation of benzophenone azine (the major reduction product), and (6) \( \text{Ph}_2\text{C}^-\text{NNH}^- \) also initiates azine formation, but the extent of reaction by the chain process is minor compared to that of \( \text{Ph}_2\text{CH}^- \).

Recently, these conclusions have been questioned by other researchers. They have proposed that in the absence of hydrogen atom and proton donors, \( \text{Ph}_2\text{CN}_2^- \) is remarkably long-lived in solution, and that this anion radical displays low reactivity even toward added proton donors, e.g., diethyl malonate (DEM). These results will be contrasted in chapter III with our most recent studies of \( \text{Ph}_2\text{CN}_2 \) reduced at a hanging mercury drop electrode in acetonitrile (AN) with \( \text{Me}_4\text{NF} \) (saturated) as the supporting electrolyte.

Our electrochemical studies of the reduction of \( \text{F}^+ \text{N}_2^- \) showed a complex
mechanism involving numerous electroactive intermediates and products. Results of FLN$_2$ reductions in the presence and absence of proton donors in DMF, AN, and butyronitrile (BN) with various supporting electrolytes at room and reduced temperatures indicate that (1) although FLN$_2^-$ is very short-lived, it does not lose N$_2$ to give FL$^-$, (2) FLN$_2^-$ undergoes a series of reactions to give a dimeric dianion, (3) the dimeric dianion loses N$_2$ upon oxidation but cleaves to give FL=NNH$_2$ and (FL=N)$_2$ upon acidification, (4) reduction of FLN$_2$ in the presence of a strong proton donor, e.g., hexafluoroisopropanol (HFIP), results in protonation at C$_9$ of FLN$_2^-$ affording FLH$_2$ as the major product, and (5) FL=NNH$^-$ and FLH$^-$ both initiate azine formation, but the extent of reaction by these chain processes is minor compared to that of FLN$_2^-$, and they do not proceed through the unidentified dimeric dianion.$^9$

The two systems which have been discussed, Ph$_2$CN$_2$ and FLN$_2$, each afford several electroactive intermediates and products. In order to distinguish among the many processes that are observed for the electroreductions of Ph$_2$CN$_2$ and FLN$_2$, separate electrochemical studies of all proposed electroactive intermediates and products were deemed necessary. In previous papers from this laboratory the redox behavior of the carbanion intermediates (Ph$_2$CH$^-$ and FLH$^-$),$^{15}$ the hydrocarbon (FLH$_2$),$^{16}$ the hydrazones (Ph$_2$C=NNH$_2$, FL=NNH$_2$, Ph$_2$CNHH=CPh$_2$, and FLHNH=FL)$^{17}$ and the azines (Ph$_2$C=NN=CPh$_2$ and FL=NN=FL)$^{17}$ have been reported. In chapter IV species which arise from the stepwise reduction of the simple hydrazones, Ph$_2$C=NNH$_2$ and FL=NNH$_2$, are examined. The compounds studied include benzophenone imine (Ph$_2$C=NH) and fluorenone imine (FL=NH), 9-aminofluorene (FLHNH$_2$), and several N-substituted derivatives of these species.
REFERENCES

1. The term "hypovalent radicals" is used to describe a neutral or
charged radical species containing less than the number of attached
substituents found in the uncharged, free radical system normally
associated with the central atom in the radical; e.g., triplet H₂C:
and HCl are neutral carbon hypovalent radicals.


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II. EXPERIMENTAL

Electrochemical instrumentation. Cyclic voltammetric and chronoamperometric experiments were performed with three-electrode potentiostats which incorporated circuits for electronic correction of ohmic potential loss between the reference and working electrodes.\(^1\) Control of the potentiostat and the acquisition and processing of chronoamperometric data were performed with a laboratory computer (LSI 11/2, ADAC System 2000). The three-electrode potentiostat which was used for the exhaustive, controlled-potential electrolysises has also been described.\(^2\)

Cells, electrodes, and electrolysis procedures. All electrochemical experiments were performed on an all-glass vacuum line. The solvent was transferred into the cell on the vacuum line by trap-to-trap distillation. Traces of oxygen, if present, were removed by several freeze-pump-thaw cycles. To bring the cell up to atmospheric pressure, the system was filled with helium. A positive pressure of helium was maintained while the reference electrode was being transferred into the cell. Slush baths (8, 0, -22, and -51\(^\circ\)C), were used to maintain the cells at a constant temperature when studies were performed at temperatures other than room temperature. These were prepared using liquid nitrogen and the appropriate liquid following the directions given by Shriver.\(^3\)

Platinum, vitreous carbon, and hanging mercury drop electrodes were used. The area of the platinum planar button electrode which was used in all
chronoamperometric and some cyclic voltammetric experiments was 0.25 cm$^2$ while the area of the planar vitreous carbon electrode$^4$ which was used in some cyclic voltammetric experiments was 0.20 cm$^2$. The platinum electrode is a commercially available inlay (Corning). These electrodes are washed between experiments with trichloroethylene, 95% ethanol, and then dried in an oven at 90°C. They were occasionally polished (approximately once every two weeks) when the broad, reversible couple near 0 V in DMF-0.1 $\text{F} (n-$Bu)$_4\text{NCIO}_4$ exceeded $\approx 2 \mu\text{A}$ (or $\approx 8 \mu\text{A cm}^{-2}$) at a scan rate of 0.2 V/s. All potentials listed are with respect to a cadmium amalgam which is in contact with a dimethylformamide solution that is saturated with both sodium chloride and cadmium chloride (Type A-III)$^5$. The potential of this electrode is $-0.75$ V vs. SCE.$^6$ Dual reference electrodes were used in all rapid scan cyclic voltammetric and chronoamperometric experiments.$^7$ The second reference electrode, which was a platinum wire in series with a 0.1 μf capacitor, was placed in parallel with the cadmium amalgam electrode.

The extent of large-scale electrolyses was monitored periodically by cyclic voltammetry. At the conclusion of the experiment, the electrolysis mixture was protonated in a dry helium atmosphere with an appropriate proton donor (e.g., diethyl malonate). The mixtures were then analyzed directly by high performance liquid chromatography (HPLC).

Chromatography. The products of the electrolyzed solutions were separated by HPLC using a 6.35-mm diameter, 25-cm length stainless steel column packed with LiChrosorb RP8 or LiChrosorb RP18, 10-μm mean particle size. The eluting solvent was a mixture of methanol and water; the ratio of the mixture and the flow rate (1-3 mL/min) were dependent upon the nature of
the products being analyzed (Table II-1). A Waters Associates Inc. Model 6000A pump was used together with a Valco sample injector valve. The detector was a Schoeffel Instrument Corp., model SF-770 UV-VIS spectrophotometer; the wavelength used was 254 nm. Calibration curves for standards of all products were constructed daily.

Chemicals. N,N-Dimethylformamide (Burdick and Jackson) was purified by passage through a column of alumina (80-200 mesh, Brockman activity 1, activated at 600°C overnight) and was collected over a mixture of activated Davison 4A molecular sieves and alumina. This procedure was carried out in a dry, nitrogen-filled glovebag. Acetonitrile (Burdick and Jackson) was purified by the procedure of Walter and Ramaley.3

Whether synthesized (Table II-2) or purchased from a commercial supplier, all compounds studied were further purified by established procedures. Purity was checked by a melting point determination and/or by chromatography. 9-Aminofluorene was prepared from the hydrochloride (Aldrich Chemical Company). Heat and light-sensitive reagents were stored at -10°C in well-sealed brown glass bottles. All remaining reagents, except for solvents and supporting electrolytes, were stored in a desiccator over phosphorus pentoxide. Tetrabutylammonium perchlorate (Southwestern Analytical Chemicals), and tetramethylammonium hexafluorophosphate (Aldrich) were used as the supporting electrolytes. These salts were stored in a vacuum oven at 60°C with phosphorus pentoxide as the desiccant.
<table>
<thead>
<tr>
<th>Compound</th>
<th>%MeOH</th>
<th>%H₂O</th>
<th>Buffer</th>
</tr>
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<tr>
<td>Benzophenone Imine</td>
<td>45</td>
<td>55</td>
<td>0.085 M Formic Acid</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.015 M Sodium Formate</td>
</tr>
<tr>
<td>Diphenylmethylamine</td>
<td>45</td>
<td>55</td>
<td></td>
</tr>
<tr>
<td>Benzophenone</td>
<td>55</td>
<td>45</td>
<td></td>
</tr>
<tr>
<td>Fluorenone Imine</td>
<td>65</td>
<td>35</td>
<td>0.09 M Sodium Acetate</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.01 M Acetic Acid</td>
</tr>
<tr>
<td>9-Aminofluorene</td>
<td>65</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>Fluorenone</td>
<td>65</td>
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<td>N,N-Dimethyl-9-aminofluorene</td>
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<td>N-Phenylfluorenone Imine</td>
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<td>Bifluorenylidene</td>
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<tr>
<td>Fluorenone Azine</td>
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</table>
TABLE II-2. Compounds Synthesized for Study and the References for the Synthetic Procedures

<table>
<thead>
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</tr>
</thead>
<tbody>
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<td>Diazodiphenylmethane</td>
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<td>Fluorenone Imine</td>
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<td>N-Phenylfluorenone Imine</td>
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</tr>
<tr>
<td>N,N-Dimethyl-9-aminofluorene</td>
<td>13</td>
</tr>
</tbody>
</table>
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4. We thank Professor D. E. Bartak, The University of North Dakota, for a gift of this electrode.


III. ELECTROREDUCTION OF DIPHENYLDAZOMETHANE IN ACETONITRILE
AT A HANGING MERCURY DROP ELECTRODE

INTRODUCTION

The electrochemical reduction of diphenyl Diazomethane (Ph₂CN₂⁻) in N,N-dimethylformamide (DMF) containing 0.1 M (n-Bu)₄NClO₄ at a platinum cathode was reported to involve the intermediacy of both Ph₂CN₂⁻ and Ph₂C⁺. Although Ph₂CN₂⁻ was not directly observed on the cyclic voltammetric time scale, its presence was inferred by the formation of Ph₂C=NNH₂ when electroreduction of Ph₂CN₂ was effected in the presence of added proton donors. From these and other results, Ph₂CN₂⁻ was concluded to react principally by rapid loss of molecular nitrogen to give the carbene anion radical, Ph₂C⁺. The principal reaction products, Ph₂C=N=N=CPh₂ and Ph₂CH₂, were proposed to arise by a chain reaction in which Ph₂CH⁻ is the chain-carrying species (eqs. III-1 thru -9):

\[
\text{Ph}_2\text{CN}_2^- + e^- \rightarrow \text{Ph}_2\text{CN}_2^- \quad \text{(III-1)}
\]

\[
\text{HA}^- + e^- \rightarrow \text{Ph}_2\text{C} = \text{NNH}^- + \text{A}^-
\]

\[
\text{Ph}_2\text{CN}_2^- \rightarrow \text{N}_2 \rightarrow \text{Ph}_2\text{C}^- \rightarrow \text{H}^+, e^- \rightarrow \text{Ph}_2\text{CH}^-
\]

\[
\text{Ph}_2\text{CH}^- + \text{Ph}_2\text{CN}_2^- \rightarrow \text{Ph}_2\text{CHNN=CPh}_2
\]

\[
\text{B} + \text{Ph}_2\text{CHNN=CPh}_2 \rightarrow \text{BH}^+ + \text{Ph}_2\text{C=N=N=CPh}_2
\]

\[
\text{Ph}_2\text{C=N=N=CPh}_2^2^- + 2 \text{Ph}_2\text{CN}_2^- \rightarrow \text{Ph}_2\text{C=N=N=CPh}_2 + 2 \text{Ph}_2\text{CN}_2^- \quad \text{(III-6)}
\]
\[
\begin{align*}
\text{Ph}_2\text{C} = \text{N}^+\text{H} + \text{Ph}_2\text{C} = \text{N}^+\text{CHPh}_2 & \rightarrow \text{Ph}_2\text{C} = \text{N}^-\text{N}^+\text{CHPh}_2 + \text{H}_2 \quad (\text{III}-7) \\
\text{Ph}_2\text{C} = \text{N}^+\text{H} + \text{HA} & \rightarrow \text{Ph}_2\text{C} = \text{N}^+\text{H}_2 + \text{A}^- \quad (\text{III}-8) \\
\text{Ph}_2\text{CH} + \text{HA} & \rightarrow \text{Ph}_2\text{CH}_2 + \text{A}^- \quad (\text{III}-9)
\end{align*}
\]

(where \( B = \text{Ph}_2\text{CH}^- \), \( \text{Ph}_2\text{C}^+ \) or \( \text{Ph}_2\text{CN}_2^- \), and \( \text{HA} = \text{Ph}_2\text{CH} = \text{N}^+\text{N}^-\text{CPh}_2 \) or \( (\text{n}-\text{Bu})_4\text{N}^+ \))

The stability of \( \text{Ph}_2\text{CN}_2^- \) and its principal reaction mode (eq. III-3) have been questioned.\(^3\) Whereas we found that \( \text{Ph}_2\text{CN}_2^- \) was too unstable (\( t^{1/2} < 1 \text{ ms} \)) to give a discernible anodic peak on the reverse, positive-going sweep in a cyclic voltammetric experiment under all conditions examined (-51^\circ\text{C} \leq T \leq 23^\circ\text{C}; \ 0.5 \text{ V/s} \leq v \leq 100 \text{ V/s}; \ \text{DMF}, \ \text{CH}_3\text{CN}, \ \text{and} \ C_3\text{H}_7\text{CN}; \ (\text{n}-\text{Bu})_4\text{NCIO}_4 \ \text{and/or} \ \text{Me}_4\text{NPF}_6 \)),\(^1,2\) Parker and Bethell\(^3\) claim that \( \text{Ph}_2\text{CN}_2 \) is reduced reversibly at a mercury cathode in acetonitrile containing \( \text{Me}_4\text{NPF}_6 \). We will demonstrate in this chapter that Parker and Bethell's observation is an artifact of their use of a mercury electrode surface for their electrochemical studies, and that their "reversible couple" is a fortuitous combination of two different redox processes.

RESULTS AND DISCUSSION

The cyclic voltammetric behavior of \( \text{Ph}_2\text{CN}_2 \) at a hanging mercury drop electrode is illustrated in Figure III-1, a-d. Under the conditions specified by Parker and Bethell for the observance of the reversible reduction of \( \text{Ph}_2\text{CN}_2 \) to \( \text{Ph}_2\text{CN}_2^- \) (\( T = 8^\circ\text{C} \), a scan rate of 100 V/s, and a saturated \( \text{CH}_3\text{CN} \) solution of \( \text{Me}_4\text{NPF}_6 \)), a cyclic voltammogram is obtained which resembles that expected for an electrochemically reversible one-electron process (Figure III-1a). However, although the separation of 60 mV for \( E_p \quad E_{p/2} \) for the cathodic wave is consistent with this interpretation, the 220 mV separation between the
anodic and cathodic peak potentials greatly exceeds the expected value of approximately 60 mV. The fact that \( \text{Ph}_2\text{CN}_2^- \) is not reversibly formed becomes obvious when the temperature is lowered. At \(-37^\circ\text{C}\) and a scan rate of 100 V/s, the shape of the anodic peak clearly reflects the effect of adsorption on the mercury surface. In addition, the separation between the anodic and cathodic peak potentials has increased to approximately 550 mV (Figure III-1b). Kinetic behavior also becomes evident as the scan rate is decreased. At \(-37^\circ\text{C}\) and a scan rate of 20 V/s, an adsorption peak that occurs at the same potential as that for the irreversible oxidation of \( \text{Ph}_2\text{CNH}=\text{CPh}_2 \) appears near 0.1 V on the second positive-going half cycle (Figure III-1c).

In addition to the claim by Parker and Bethell that \( \text{Ph}_2\text{CN}_2^- \) can be observed directly by cyclic voltammetry in \( \text{CH}_3\text{CN} \) in the presence of \( \text{Me}_4\text{N}^+ \), they reported that the cathodic peak potential for the reduction of \( \text{Ph}_2\text{CN}_2 \) is unaffected by the presence of electroinactive diethyl malonate (DEM). From this result, these workers concluded that the proton donor is not involved kinetically in the decomposition of \( \text{Ph}_2\text{CN}_2^- \) under the reaction conditions, and that \( \text{Ph}_2\text{CN}_2^- \) must undergo relatively slow, rate-determining hydrogen atom abstraction from acetonitrile. In our experiments, added DEM has a pronounced effect on the reduction of \( \text{Ph}_2\text{CN}_2 \) in \( \text{CH}_3\text{CN} \) as shown by comparing Figures III-1a and III-1d, the latter containing a 22-fold excess of DEM. The addition of DEM essentially eliminates the anodic peak at \(-0.7\) V on the reverse, positive-going sweep, and a reversible couple appears near 0.4 V. Since this same couple can be made to appear by scanning in the positive-going direction from an initial potential of 0 V if a base such as \( \text{Me}_4\text{NOH} \) is added to a solution of DEM in \( \text{CH}_3\text{CN}-\text{Me}_4\text{NPF}_6 \) (saturated), the conjugate base of DEM must be generated as a result of the electroreduction of \( \text{Ph}_2\text{CN}_2 \) in the
presence of DEM. The effects of temperature and proton donor not only disprove Parker and Bethell's assignment of the anodic peak near -0.7 V as being due to the reoxidation of Ph₂CN₂⁻, but it also invalidates their attempt to relate the disappearance of this anodic adsorption peak to the kinetics of Ph₂CN₂⁻ decomposition.

In contrast, when the electroreduction of Ph₂CN₂ is effected on a platinum electrode surface under these conditions (T = 8°C, scan rate of 100 V/s, and CH₃CN-Me₄NPF₆ (saturated)), no adsorption is apparent and the cyclic voltammetric results are consistent with the interpretation which we presented previously for the reduction of Ph₂CN₂ in DMF.¹ ² As evidenced by the cyclic voltammogram in Figure III-1a, the reduction of Ph₂CN₂ on platinum is chemically irreversible at 8°C at our maximum scan rate of 100 V/s. Although the absence of an anodic wave for the oxidation of Ph₂CN₂⁻ precludes a kinetic study of Ph₂CN₂⁻ decomposition, it does establish a maximum half-life of 1 ms for Ph₂CN₂⁻.⁴ The only anodic wave discernible at this scan rate occurs near -0.35 V and is assigned to the irreversible oxidation of Ph₂CH⁻, the observed reaction product of Ph₂CN₂⁻.² The anodic peak for Ph₂CH⁻ is kinetically controlled and decreases in relative magnitude as the scan rate is decreased (Figure III-1f). Concomitantly, an irreversible anodic peak appears near 0.2 V that is due to the oxidation of the intermediate product, Ph₂CHNN=CPh₂, formed when Ph₂CH⁻ couples with Ph₂CN₂. The two principal products that were obtained by a coulometric reduction of Ph₂CN₂ at a platinum cathode in CH₃CN-Me₄NPF₆ (saturated) are Ph₂C=N=N=CPh₂ (42%) and Ph₂CH₂ (57%).
FIGURE III-1

Cyclic voltammograms for the reduction of Ph₂CN₂ in CH₃CN under the following conditions: (a) $T = 8^\circ$C, $v = 100$ V/s, Me₄NPF₆ (saturated), and hanging mercury drop electrode (HMDE); (b) $T = -37^\circ$C, $v = 100$ V/s, Me₄NPF₆ (saturated), and HMDE; (c) $T = -37^\circ$C, $v = 20$ V/s, Me₄NPF₆ (saturated), and HMDE; (d) $T = 8^\circ$C, $v = 100$ V/s, Me₄NPF₆ (saturated), DEM (89 mM), and HMDE; (e) $T = 8^\circ$C, $v = 100$ V/s, Me₄NPF₆ (saturated), and planar platinum electrode (0.25 cm²); and (f) $T = 8^\circ$C, $v = 20$ V/s, Me₄NPF₆ (saturated), and planar platinum electrode.
CONCLUSIONS

We suspect that Parker and Bethell's use of mercury may have contributed to the misinterpretation by these workers that Ph₂CN⁻ has an appreciable lifetime in CH₃CN. The formation of organomercurials when mercury is used as the electrode material is well known (e.g., \((\text{Ph}_2\text{CH}_2)_2\text{Hg}\) formation when \(\text{Ph}_2\text{CH}_2\text{Br}\) is reduced at a mercury cathode⁵) and may account for the anodic wave that appears near \(-0.7\) V at \(8^\circ\text{C}\) in Figure III-1a. A specific possibility is that \(\text{Ph}_2\text{CH}^-\) is oxidized to \(\text{Ph}_2\text{CH}^+\) and that this radical then rapidly reacts with the mercury surface to form an organomercurial species. The effect of this follow-up reaction would be to shift the \(\text{Ph}_2\text{CH}^-\) anodic wave to more negative potential.⁴ Since the cathodic peak potential for the irreversible reduction of \(\text{Ph}_2\text{CN}_2\) will be shifted negatively by decreasing temperature, Parker and Bethell perhaps arrived at a fortuitous set of conditions in which the irreversible reduction of \(\text{Ph}_2\text{CN}_2\) and the irreversible oxidation of \(\text{Ph}_2\text{CH}^-\) in the presence of mercury appeared to constitute a chemically reversible system.

It is important to note that our coulometric and product data preclude reaction pathways which involve extensive hydrogen atom abstraction by anion radical intermediates. For example, the electroreduction of \(\text{Ph}_2\text{CN}_2\) in DMF-0.1 \(\text{F}(\text{n-Bu})_4\text{NCIO}_4\) gives an \(n\) value of 0.30 and a product distribution of 87% Ph₂C=N=N=CH₂, 1% Ph₂C=O, 9% Ph₂CH₂, 2% Ph₂CHNNHN=CPh₂ and 0.6% Ph₂C=NNH₂.² If either \(\text{Ph}_2\text{CN}^-\) or \(\text{Ph}_2\text{C}^-\) were to hydrogen atom abstract, these hydrogens must ultimately be removed as protons from the coupled intermediate, \(\text{Ph}_2\text{CHNN}=\text{CPh}_2\), by the electrogenerated bases \(\text{Ph}_2\text{C}^-, \text{Ph}_2\text{CH}^-, \text{Ph}_2\text{C}=\text{NNH}^-,\) and \(\text{Ph}_2\text{CHNN}=\text{CPh}_2\) (see eqs. III-1 thru III-9). Because the combined yields of
the last three products is less than 12% under these conditions, the anion radicals \( \text{Ph}_2\text{CN}_2 \) and \( \text{Ph}_2\text{C}^- \) must instead proton abstract from \( \text{Ph}_2\text{CHNN}=\text{CPh}_2 \) (eq. III-5). Since \( \text{Ph}_2\text{CH}^- \), not \( \text{Ph}_2\text{C}=\text{NNH}^- \), is the first observable electroactive intermediate in the reduction of \( \text{Ph}_2\text{CN}_2 \), we conclude that \( \text{Ph}_2\text{CN}_2^- \) reacts principally by rapid loss of nitrogen, and that \( \text{Ph}_2\text{CH}^- \) is formed via the intermediacy of the carbene anion radical, \( \text{Ph}_2\text{C}^- \), as shown in eq. III-3.
REFERENCES


IV. THE ELECTROREDUCTION OF FLUORENONE IMINE, N-PHENYLFUORENONE IMINE, BENZOPHENONE IMINE, 9-AMINOFLUORENE, AND N,N-DIMETHYL-9-AMINOFLUORENE

Fluorenone Imine

Cyclic voltammetry. The cyclic voltammetric reduction of F1=NH in DMF-0.1 F (n-Bu)4 NC104 at a platinum cathode occurs in successive one-electron steps (Figure IV-1). Although the reduction of F1=NH to F1=NH+ at E_p,c = -0.92 V is nearly reversible (i_p,a/i_p,c = 0.93) at the relatively slow scan rate of 0.2 V/s (Figure IV-1a), the reduction of F1=NH+ to F1=NH2 at E_p,c = -1.68 V occurs irreversibly at all scan rates and temperatures in the range 0.05 V/s ≤ v ≤ 100 V/s and -51°C ≤ T ≤ 22°C. Reaction of F1=NH2 causes one or more anodic waves, depending on the temperature and scan rate, to appear on the reverse, positive-going sweep. At room temperature (Figure IV-1b), the most prominent of these is assigned to the irreversible oxidation of F1NH2- (E_p,a = -0.52 V). F1NH2-, which would arise if F1=NH2 were to proton abstract (eq. IV-2), has been shown previously1 to be oxidized irreversibly to give F1=NH. The protons that also are formed during the oxidation are then captured by unreacted F1NH2-.

The F1NH2 which is formed upon the oxidation of F1NH2- has a modest effect on the redox behavior of F1=NH on the second, negative-going scan. Protonation of F1=NH+ by F1NH2 affords F1NH2+ and F1NH2-; F1NH2+ is rapidly reduced, either electrochemically or chemically, by unreacted F1=NH+ (eq. IV-3). The protonation of F1=NH+ by F1NH2 thus facilitates the reduction of
FIGURE IV-1

Cyclic voltammograms of Fl=NH in DMF-0.1 \( \text{F} (\text{n-Bu})_4 \text{NCIO}_4 \) at a planar platinum electrode. The scans were initiated at 0 V in the negative-going direction at a rate 0.2 V/s. The temperature was 23°C; (a) 2.41 mM Fl=NH; (b) 9.46 mM Fl=NH; (c) 3.01 mM Fl=NH and 23.41 mM diethyl malonate.
F1=NH, thereby causing a shoulder to appear on the positive side of the F1=NH cathodic peak on the second, negative-going scan (Figure IV-1b). Since the concentration of F1=NH exceeds the concentration of F1HNH2 at the electrode surface, all F1HNH2 which was produced during the positive-going half cycle will be consumed when reduction of F1=NH is affected on the second, negative-going half cycle. As a result, no cathodic wave for the reduction of F1HNH2 is seen near -1.92 V (vide infra) on the second, negative-going scan.

\[
\text{F1=NH} + e^- \rightarrow \text{F1=NH}^+ \quad \text{(IV-1)}
\]

\[
\text{e}^- \rightarrow \text{F1=NH}^2^- \xrightarrow{\text{fast}} \text{S}^- + \text{F1NH}_2^- \quad \text{(IV-2)}
\]

\[
\text{F1HNH}_2 \rightarrow \text{F1NH}_2^+ + \text{e}^- \rightarrow \text{F1NH}_2^- \quad \text{(IV-3)}
\]

**Coulometry.** The controlled potential electrolysis of F1=NH was effected at a potential (E_{applied} = -1.0 V) which was sufficiently negative so as to reduce F1=NH to its anion radical, but not so negative so as to cause reduction of F1=NH\(^+\) to F1=NH\(^2^-\). HPLC analysis of a solution in which electrolysis was terminated arbitrarily at the point of 1.5 electrons per molecule of F1=NH showed that 72% of the F1=NH which was originally present had been converted into F1HNH2 (Table IV-2). This result is consistent with a nominal \(n\) value of 2.0 for the reduction of F1=NH to F1HNH2.

While F1=NH\(^+\) is quite stable on the cyclic voltammetric time scale, its reaction on the coulometric time scale is relatively rapid. Periodic monitoring of the electrolysis experiment by cyclic voltammetry showed that the F1=NH peak height remains almost constant for \(n \leq 1.0\), and then decreases
<table>
<thead>
<tr>
<th>Entry No.</th>
<th>Compound</th>
<th>Conc., mM</th>
<th>$E_{\text{applied}}$, V</th>
<th>n</th>
<th>Proton Donor; Conc., mM</th>
<th>Products, % yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F1=NH</td>
<td>4.43</td>
<td>-1.0</td>
<td>1.5</td>
<td>21</td>
<td>72</td>
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<tr>
<td>2</td>
<td>F1=NH</td>
<td>3.01</td>
<td>-0.9</td>
<td>2.3</td>
<td>DEM; 23.4</td>
<td>7</td>
</tr>
<tr>
<td>3</td>
<td>Ph$_2$C=NH</td>
<td>6.51</td>
<td>-1.5</td>
<td>2.0</td>
<td>2.5</td>
<td>97</td>
</tr>
<tr>
<td>4</td>
<td>Ph$_2$C=NH</td>
<td>9.63</td>
<td>-1.3</td>
<td>2.0</td>
<td>TFE; 101.1</td>
<td>13</td>
</tr>
</tbody>
</table>
linearly to zero in the range $1.0 \leq n \leq 2.0$. Concomitantly, the peak height for the reduction of $F_1^{-} = NH^{-}$ to $F_1^{-} = NH_2^{2-}$ decreases linearly to zero as $n$ approaches $1$. If $n > 1$, a cathodic wave for the reduction of $F_1NH_2^{-}$ is also present at $-1.92$ V.

The fact that the magnitude of the $F_1^{-} = NH$ cathodic wave remains constant for $n \leq 1$ also requires that $F_1NH_2^{-}$ rather rapidly abstract a proton from a component of the solvent-electrolyte system. A constant peak height for $F_1^{-} = NH$ reduction is then predicted as long as $[F_1^{-} = NH] \geq [F_1NH_2^{-}]$, and the sum of the instantaneous concentrations of $F_1^{-} = NH$ and $F_1NH_2^{-}$ equals the initial concentration of $F_1^{-} = NH$. The pathway by which $F_1^{-} = NH$ reacts cannot be ascertained from these data, but presumably it involves either disproportionation of $F_1^{-} = NH$ or a half-regeneration mechanism in which $F_1^{-} = NH$ is protonated by a component of the solvent-electrolyte system, and the resulting $F_1NH_2^{-}$ is reduced by unreacted $F_1^{-} = NH$.

**Chronoamperometry.** Inspection of the cyclic voltammogram of $F_1^{-} = NH$ (Figure IV-1b) shows that the magnitude of the cathodic peak for the reduction of $F_1^{-} = NH$ to $F_1^{-} = NH_2^{-}$ is approximately two-thirds of that for the reduction of $F_1^{-} = NH$ to $F_1NH^{-}$. Since $i_p, a/i_p, c = 0.93$ at $y = 0.2$ V/s for the reduction of $F_1^{-} = NH$ to $F_1NH^{-}$ (Figure IV-1a), the possibility that the smaller-than-expected peak height for the reduction of $F_1^{-} = NH$ to $F_1^{-} = NH_2^{2-}$ may be due to causes other than a follow-up reaction involving $F_1^{-} = NH$ was examined. First, $F_1^{-} = NH_2^{-}$ may be reacting with $F_1^{-} = NH$, e.g., proton transfer, to give a product or products which are electroinactive at the potential at which $F_1^{-} = NH$ is reduced to $F_1^{-} = NH_2^{2-}$. Second, if the rate of the follow-up reaction involving $F_1^{-} = NH_2^{2-}$ is very rapid, the $F_1^{-} = NH$ cathodic peak height
will be a function of the transfer coefficient for heterogeneous electron
transfer. In order to test the merit of these possibilities, the stepwise
reduction of F1=NH to F1=NH
2
−
 was examined by single potential step
chronoamperometry. The reduction of F1=NH to F1=NH
−
 (E
applied
 = -1.40 V) afforded
\textit{iK}{1/2/c} = 41 \pm 3 \mu A \cdot s^{1/2} \cdot mM^{-1}
for the time range 3 ms \leq \tau \leq 4s.

When the potential instead was stepped sufficiently negative so as to effect
the reduction of F1=NH to F1=NH
2
−
 (E
applied
 = -1.95 V), the values for \textit{iK}{1/2/c}
were approximately 1.8 times larger than the corresponding values at the less
negative potential. In order to relate these values to the number of
electrons involved, the reduction of fluorenone (F1=O) to its stable anion
radical was studied under similar conditions. If it is assumed that the
diffusion coefficients of F1=NH and F1=O are equal, then the \textit{iK}{1/2/c} value of
37 \mu A \cdot s^{1/2} \cdot mM^{-1}
that was obtained for the one-electron reduction of F1=O
indicates that the number of electrons transferred in the first and second
steps of the reduction of F1=NH are approximately 1.1 and 0.9 (2.0 overall),
respectively. These results are consistent with a minor reaction channel
involving the slow protonation of F1=NH
−
, a cathodic peak shape for the
reduction of F1=NH
−
 to F1=NH
2
−
 which is controlled by slow electron transfer,
and little, if any, reaction of F1=NH
2
−
 with F1=NH to give an electroinactive
product.

\textbf{Effect of diethyl malonate (DEM).} The addition of DEM, a weak proton
donor, causes the cyclic voltammetric peak height for the reduction of F1=NH
to F1=NH
−
 (E
p,e
 = -0.92 v) to increase approximately 1.8 times, eliminates
the cathodic peak attributed to the reduction of F1=NH
−
 to F1=NH
2
−
 at -1.6 V,
causes a new peak to appear near -1.92 V which is due to the concomitant
reduction of F1=NH
2
 and its two-electron reduction product, fluorene (F1H
2
),
and eliminates all anodic peaks on the positive-going half cycle (Figure IV-1c). The chronomeperometric n value which was obtained at a potential (E = -1.35 V) that was sufficiently negative so as to reduce F1=NH to F1=NH\textsuperscript{2-}, but not so negative so as to cause the reduction of F1=NH\textsuperscript{2-} to F1=NH\textsuperscript{2-}, is diffusion controlled and equal to 2 in the time range 8 ms < t < 8 s. This result is consistent with rapid protonation of all anionic intermediates and the formation of F1NHNH\textsubscript{2} as the final product. Product studies of a partially electrolyzed solution confirmed that F1NHNH\textsubscript{2} was the principal product (85% relative yield) (Table IV-1, entry 2). Minor amounts of F1NH\textsubscript{2} and F1=O were also found. The former species is the expected two-electron reduction product of the more difficultly reduced F1NHNH\textsubscript{2} (vide infra), while the latter species is expected if fluorenlyl anions are formed in the presence of adventitious oxygen.\textsuperscript{2}

**Comparison of the redox behavior of F1=NH and F1=NPh.** Inspection of Figure IV-1b will show that reduction of F1=NH\textsuperscript{+} to F1=NH\textsuperscript{2-} affords a relatively large anodic peak for the oxidation of F1NH\textsubscript{2}\textsuperscript{-} at -0.52 V and a smaller, unidentified anodic peak at -0.62 V. The latter peak shifts positively with increasing scan rate, merging with the F1NH\textsubscript{2}\textsuperscript{-} anodic peak at scan rates in excess of 5 V/s at both room temperature and -51°C.

The possibility that a small, but discernible, amount of electrogenerated F1=NH\textsuperscript{2-} reacts with F1=NH to give F1NH\textsubscript{2} and F1=N\textsuperscript{-} and that the latter species is oxidized at -0.62 V was examined. In order to eliminate the possibility of proton transfer, the reduction of the N-substituted fluorenone imine, F1=NPh, was studied in the presence and absence of potassium tert-butoxide (K-t-BuO). When electroreduction of F1=NPh was effected in DMF-0.1 M (n-Bu)\textsubscript{4}NClO\textsubscript{4} at a
FIGURE IV-2

Cyclic voltammograms of F1=NPh in DMF-0.1 M (n-Bu)4ClO4 at a planar platinum electrode. The scans were initiated at 0 V in the negative-going direction at rates of (a) 0.2 V/s, (b) 0.2 V/s, and (c) 5.0 V/s. The temperature was 23°C. (a) 9.46 mM F1=NPh; (b) 9.46 mM F1=NPh and 30 mM K-t-BuO; (c) 9.46 mM F1=NPh and 30 mM K-t-BuO.
The image contains a series of electrochemical cyclic voltammograms. Each graph shows the current density (I) as a function of the potential (E) in volts (V). The current is labeled with subscripts 'c' and 'a', representing cathodic and anodic currents, respectively. The potential range is from 0.8 to -1.6 V. The scale for current density is marked at 20 μA, 50 μA, and 150 μA. The graphs are labeled 'a', 'b', and 'c'.
platinum electrode, two anodic waves, corresponding to the oxidation of
\( \text{F1=NHPh}^- (E_{p,a} = -0.67 \text{ V}) \) and \( \text{F1NHPh}^- (E_{p,a} = -0.18 \text{ V}) \), were observed on the
reverse positive-going sweep (Figure IV-2a). However, when \( K-\text{BuO}^- \) was added
in excess, the oxidation of \( \text{F1NHPh}^- \) occurred in either one or two steps,
depending upon the sweep rate. At scan rates less than approximately 1 V/s,
the anodic peak ascribed to the oxidation of \( \text{F1NHPh}^- (E_{p,a} = -0.18 \text{ V}) \)
disappeared while the anodic peak ascribed to the oxidation of \( \text{F1=NHPh}^- \) nearly
doubled in magnitude (Figure IV-2b). At scan rates in excess of 1 V/s, the
anodic peak for \( \text{F1NHPh}^- \) reappeared and grew in relative magnitude at the
expense of the \( \text{F1=NHPh}^- \) anodic peak with increasing scan rate (Figure IV-2c).
The anodic peaks were approximately equal in magnitude under these solution
conditions when the scan rate exceeded 50 V/s.

The kinetically controlled cyclic voltammetric behavior of \( \text{F1NHPh}^- \) and
\( \text{F1=NHPh}^- \) in the presence of added base must be due to electrocatalysis. The
concomitant oxidation of an anion radical and the corresponding anion has
ample precedent, e.g., \( \text{F1=O}^- \) and \( \text{F1OH}^- \) and the conjugate base of a
hydroquinone, \( \text{QH}^- \), and \( \text{Q}^- \), and would occur if \( \text{F1=NPh} \) were to accept an
electron from the more-difficultly-oxidized \( \text{F1NHPh}^- \) in the rate determining
step (eq. IV-5).

\[
\begin{align*}
\text{F1=NHPh}^- & \rightarrow \text{F1=NPh}^- \quad \text{(IV-4)} \\
\text{F1=NPh}^- + \text{F1NHPh}^- & \rightarrow \text{F1=NPh}^- + \text{F1NHPh}^- \\
\text{F1NHPh}^- + \text{t-BuO}^- & \rightarrow \text{F1=NPh}^- + \text{t-BuOH} \quad \text{(IV-5)}
\end{align*}
\]

Electron transfer is then rendered irreversible by the action of the added
base, \( \text{t-BuO}^- \), on \( \text{F1NHPh}^+ \) (eq. IV-6). The failure of \( \text{F1NHPh}^- \) to act in the
same capacity as \( \text{t-BuO}^- \) is presumably the result of slow proton transfer from
\( \text{F1NHPh}^+ \) to \( \text{F1NHPh}^- \). The relatively slow rate of proton transfer to and from
the C₉ center in several fluorene compounds has been noted previously.¹

Since an anodic peak which is analogous to the one in question did not appear in the Fl=NPh system, attempts were then made to prepare Fl=N⁻ directly from either the addition or electrogeneration of a strong base. (CH₃)₄NOH in methanol acted only as a proton donor toward Fl=NH⁺, and neither K-t-BuO nor electrogenerated PhN=NPh₂⁻ produced any discernible reaction with Fl=NH. When Fl=NH was reduced to Fl=NH₂⁻ in the presence K-t-BuO, the anodic peak assigned to the oxidation of FlNH₂⁻ was eliminated while the magnitude of the anodic peak at -0.62 V became approximately twice as large as the FlNH₂⁻ peak in the absence of base. Surprisingly, unlike the Fl=NPh system, the anodic wave for the oxidation of Fl=NH⁺ was not enhanced when Fl=NH was reduced to either Fl=NH⁺ or Fl=NH₂⁻ in the presence of K-t-BuO. The apparent absence of electrocatalysis in the Fl=NH system is puzzling, since the separation between the Fl=NH⁺ (Eₚ,a = -0.92 V) and FlNH₂⁻ (Eₚ,a = -0.52 V) anodic peaks is approximately 0.1 V less than the corresponding separation of the anodic peaks for Fl=NPh⁺ (Eₚ,a = -0.67 V) and FlNHPh⁻ (Eₚ,a = -0.18 V).

While these data appear to be consistent with a pathway which involves the strong-base assisted oxidation of FlNH₂⁻ (eq. IV-7 thru 10), confirmation of the reaction sequence could not be verified by independent experiment.

\[
\text{FlNH₂⁻} \rightarrow \text{FlNH₂⁻} + e^- \\
\text{FlNH₂⁻} \rightarrow \text{FlNH₂} + \text{Fl=NH⁺} \quad \text{(IV-7)}
\]

\[
\text{FlNH₂⁻} \rightarrow \text{FlNH₂} + \text{Fl=NH⁺} \\
\text{FlNH₂} \rightarrow \text{FlNH₂} + \text{Fl=NH⁺} \quad \text{(IV-8)}
\]

\[
\text{FlNH₂⁻} \rightarrow \text{t-BuOH} + \text{Fl=NH⁺} \quad \text{(IV-9)}
\]

\[
\text{Fl=NH⁺} \rightarrow \text{Fl=NH} + e^- \quad \text{(IV-10)}
\]
If oxidation of $\text{F1NH}_2^-$ were facilitated by the presence of a strong base such as either $\text{OH}^-$ formed by action of $\text{F1=NH}^2-$ on adventitious $\text{H}_2\text{O}$ or added $\text{t-BuO}^-$, theory predicts that the $\text{F1NH}_2^-$ anodic peak would be shifted in the negative direction because of the increased rate of proton loss from $\text{F1NH}_2^-$ (eq. IV-7 and -8) and that the $\text{F1NH}_2^-$ anodic peak height could be increased by as much as 100%. Although the addition of K-$\text{t-BuO}$ to a solution of $\text{F1=NH}$ does cause these changes to occur when $\text{F1=NH}$ is reduced to $\text{F1=NH}^2-$, the addition of excess K-$\text{t-BuO}$ to a solution of $\text{F1NHNH}_2^-$ affords only a single anodic peak for the oxidation of $\text{F1NH}_2^-$ at its normal potential of $-0.52$ V. The expected single anodic peak at $-0.62$ V could be made to arise only by first oxidizing $\text{F1NH}_2^-$ to $\text{F1=NH}$ and then reducing the electrogenerated $\text{F1=NH}$ to $\text{F1=NH}^2-$; no anodic wave at $-0.62$ was seen in either the $\text{F1=NH}$ or the $\text{F1NH}_2^-$ systems unless the reduction of $\text{F1=NH}^-$ to $\text{F1=NH}^2-$ was first made to occur.

Finally, since the unidentified anodic peak at $-0.62$ V occurs at the same potential as that for the reversible one-electron oxidation of fluorenone azine dianion, $\text{F1}=\text{N}=\text{N}=\text{F1}^2-$, the possibility that this species may be formed in small amounts when $\text{F1=NH}$ is reduced to $\text{F1=NH}^2-$ and that $\text{F1=N}=\text{F1}^-$ then catalyzes the oxidation of $\text{F1NH}_2^-$ when an added base is present was examined. However, when $\text{F1=N}=\text{F1}^-$ was added in a small amount to a solution of $\text{F1=NH}$, no apparent change in the cyclic voltammetric behavior occurred. Although the reason why electrocatalysis is not observed in the $\text{F1=NH}$ system under these conditions is not clear, the result is consistent with the similar absence of electrocatalytic oxidation of $\text{F1NH}_2^-$ by $\text{F1=NH}$ when K-$\text{t-BuO}$ has been added (vide supra).
Benzophenone Imine (Ph₂C=NH)

The cyclic voltammetric behavior of Ph₂C=NH in DMF-0.1 \( \mathcal{F}(\text{Bu}_4N)\text{ClO}_4 \) closely resembles that of Fl=NPh. At a scan rate of 5 V/s, clearly defined cathodic peaks corresponding to the nearly reversible reduction of Ph₂C=NH to Ph₂C=NH⁻ and the irreversible reduction of Ph₂C=NH⁻ to Ph₂C=NH²⁻ are observed at -1.47 V and -1.96 V, respectively, while anodic peaks are assigned to the oxidation of Ph₂C=NH⁻ and Ph₂CNH₂⁻ arise at -1.40 and -0.88 V, respectively, on the reverse, positive-going sweep. When the scan rate is decreased, loss of reversibility for the Ph₂C=NH/Ph₂C=NH⁻ couple occurs, as evidenced by a value of 0.52 for \( \frac{i_p,a}{i_p,c} \) at \( v = 0.2 \) V/s. Concomitantly, the relative magnitude of the Ph₂C=NH cathodic wave is increased, with its increase occurring at the expense of the more-negative cathodic wave for the reduction of Ph₂C=NH⁻ to Ph₂C=NH²⁻. In addition, the anodic peak assigned to the irreversible oxidation of Ph₂CNH₂⁻ is no longer discernible at a scan rate of 0.1 V/s. The latter result is consistent with the known instability of benzhydryl anions and their propensity to proton abstract. The former result is consistent with the relatively slow abstraction of a proton from a component of the solvent-supporting electrolyte system by Ph₂C=NH⁻ and the subsequent reduction of Ph₂CNH₂⁺ at the applied potential to give Ph₂C=NH²⁻. The formation of Ph₂CNH₂ upon reduction of Ph₂C=NH to its anion radical was confirmed by product studies from controlled potential electrolyses and occurred when reduction was effected in either the presence or absence of an added proton donor (entries 3 and 4, Table IV-1).
9-Aminofluorene (F1HNNH$_2$)

Cyclic voltammetry. The cyclic voltammetric reduction of F1HNNH$_2$ to F1HNNH$_2$$^-$ (E$_{p,c} =$ -1.92 V) is nearly reversible at T = -51°C and v $>$ 20 V/s. As the temperature is increased and/or the scan rate is decreased, the decomposition of F1HNNH$_2$$^-$ first affords two equal-sized, irreversible, anodic processes which appear near -0.48 V and 0.12 V (Figure IV-3a). The more-negative process has been assigned above to the irreversible oxidation of F1NH$_2$$^-$ to give F1=NH while the more-positive peak has been shown in earlier work$^2$ to be due to the irreversible oxidation of 9-fluorenyl anion, F1H$^-$. As the scan rate is decreased further (Figure IV-3b), the relative magnitude of the F1NH$_2$$^-$ anodic peak is increased at the expense of the F1H$^-$ anodic peak. Finally, at room temperature and a scan rate of 0.2 V/s, both the reversibility for the reduction of F1HNNH$_2$ and the anodic peak for the oxidation of F1H$^-$ disappear. Subsequent negative-going scans show, in addition to the F1HNNH$_2$ cathodic peak, a single cathodic peak (E$_{p,c} =$ -0.84 V) for the reduction of the F1=NH which is electrogenerated by the oxidation of F1NH$_2$$^-$ on the previous positive-going half-cycle. Since the sum of the F1HNNH$_2$ and F1H$_2$ concentrations at the electrode surface exceeds the concentration of electrogenerated F1=NH, the reduction of F1=NH on the second negative-going scan occurs in a single, irreversible, two-electron step to regenerate F1HNNH$_2$.

Proposed reaction scheme. The scheme described by eqs. IV-11 thru -16 is proposed to account for the kinetically controlled behavior of the F1HNNH$_2$$^-$, F1NH$_2$$^-$, and F1H$^-$ anodic peaks.

\[
\begin{align*}
F1HNNH_2 + e^- & \rightarrow F1HNNH_2^- \quad \text{(IV-11)} \\
F1HNNH_2^- & \overset{\text{slow}}{\rightarrow} F1H^-/ + NH_2^-/ . \quad \text{(IV-12)}
\end{align*}
\]
\[
\text{FLNH}_2^+ + \text{FlH}^+ \text{ or } \text{NH}_2^+ \rightarrow \text{FLNH}_2 + \text{FlH}^- \text{ or } \text{NH}_2^-
\]  \hspace{1cm} (IV-13)

\[
\text{NH}_2^- + \text{FLNH}_2 \xrightarrow{\text{fast}} \text{NH}_3 + \text{FLNH}_2^-
\]  \hspace{1cm} (IV-14)

\[
\text{FlH}^- + \text{FLNH}_2 \xrightarrow{K} \text{FlH}_2 + \text{FlNH}_2^-
\]  \hspace{1cm} (IV-15)

\[
\text{FlH}_2 + e^- \rightarrow \text{FlH}_2^-
\]  \hspace{1cm} (IV-16)

The reduction of FLNH$_2$ as a diffusion controlled, overall one-electron step was ascertained by single-potential-step chronoamperometry. Although double-potential-step chronoamperometry established that the decomposition of FLNH$_2^+$ was first order in this species ($k = 0.8$ s$^{-1}$ at $-22^\circ$C; see Figure IV-4), it could not be determined unequivocally which fragment of the carbon-nitrogen bond cleavage bears the negative charge (eq. IV-12). The $-$/.$ superscript which is written for each of the fragments reflects this uncertainty. However, the chronoamperometric data do require that the radical fragment be reduced by unreacted FLNH$_2^+$ (eq. IV-13) and that NH$_2^-$ rapidly abstract a proton from FLNH$_2$ (eq. IV-14). The disappearance of the FlH$^-$ anodic peak with decreasing scan rate is the result of relatively slow proton transfer from FLNH$_2$ to electrogenerated FlH$^-$ (eq. IV-15) and the subsequent oxidation of the more-readily-oxidized FLNH$_2^-$. Because the FlH$_2$ which is formed in this reaction is also reduced to its relatively stable anion radical at the same potential at which FLNH$_2$ is reduced,$^2$ the experimental double-potential-step chronoamperometric data for this process deviate from the theoretical curve for the simple, first order EC model at larger values of time, $t$. A more complicated EC model which includes the effect of eqs. IV-15 and -16 was not developed. If the knowledge of the proton transfer rate constant in eq. IV-15 were of interest, a double-potential-step chronoamperometric method which involves the reduction of FLNH$_2$ on the forward step and the oxidation of both unreacted FLNH$_2^+$ and FLNH$_2^-$ on the reverse step should be amenable experimentally. The value of the equilibrium
FIGURE IV-3

Cyclic voltammograms of FLHNNH₂ in DMF-0.1 F (n-Bu)₄ClO₄ at a spherical platinum electrode. The scans were initiated at 0 V in the negative-going direction at rates of (a) 100 V/s and (b) 5 V/s. Concentration of FLHNNH₂ is 5.95 mM. The small "2" indicates the second scan in the negative direction.
Double potential step chronoamperometry data for the reduction of \( \text{Fl} \text{HNNH}_2 \) in DMF-0.1 \( \text{F} \text{(m-Bu)}_4 \text{ClO}_4 \) at a planar platinum electrode at \(-22^\circ \text{C}\). Concentrations are 2.46 mM (closed symbols) and 8.70 mM (open symbols). For the circles \((t-\tau_f)/\tau_f\) is 0.3; for the triangles 0.5; and for the squares 0.7. Theoretical curves are for \( k = 0.8 \text{ s}^{-1} \). The applied potential for \( t = 0, t < \tau \) and \( t > \tau \) is \(-1.2 \text{ V}, -2.1 \text{ V} \) and \(-1.2 \text{ V}\), respectively.
constant for the reaction described by eq. IV-15 has not been reported but
should be approximately one; pK values of 22.9 and 23.0 have been reported
for FLH₂ and FLHN(CH₃)₂, respectively, in DMSO.⁶

**N,N-Dimethyl-9-aminofluorene**

Although FLHNMe₂⁻ (k = 1.1 s⁻¹ at T= -51°C, see Figure IV-5) is slightly
less stable than FLHNH₂⁻, analogous pathways for FLHNMe₂ and FLHNH₂
reductions may be written. The principal differences in the cyclic
voltammetric behavior of these compounds occur when the corresponding
conjugate bases of FLNMe₂⁻ and FLNH₂⁻ are oxidized. Whereas oxidation of
FLNH₂⁻ to give Fl=NH involves the relatively slow loss of a proton from FLNH₂⁺
to FLNH₂⁻ and other indirectly electrogenerated bases, the absence of a rapid
follow-up chemical reaction involving FLNMe₂⁺ causes FLNMe₂⁻ (E°,pc = -0.16 V)
to be oxidized with considerably greater difficulty than FLNH₂⁻ (E°,pa = -0.52
V). If FLH⁻ has not been permitted sufficient time to react completely by
abstracting a proton from FLHNMe₂, then the anodic processes which occur on
the reverse, positive-going half cycle appear to resemble that of FLNH₂⁻,
*viz.*, FLNMe₂⁻ and FLH⁻ afford single irreversible peaks at -0.16 and 0.16 V,
respectively (Figure IV-6). However, if reaction of FLH⁻ with FLHNMe₂ is
complete on the cyclic voltammetric time scale, then oxidation of FLNMe₂⁻
occurs stepwise and reasonably reversibly, affording first the neutral
radical, FLNMe₂⁺, at -0.16 V and then the cation, FLNMe₂⁺, at 0.34 V. The
stepwise oxidation of FLNMe₂⁻ is most readily seen when FLNMe₂⁻ is prepared in
situ by action of electrogenerated PhN=NPh⁺ on FLHNMe₂ (Figure IV-7). The
cathodic waves which appear on the first negative-going half cycle arise from
the reversible reduction of PhN=NPh to PhN=NPh⁻ (E°,pc = -0.60 V) and the
FIGURE IV-5

Double potential step chronoamperometric data for the reduction of 
$\text{FLHN(CH}_3\text{)}_2$ in DMF-0.1 F (n-Bu)$_4$ClO$_4$ at a planar platinum electrode at
$-51^\circ\text{C}$. Concentrations are 1.67 mM (closed symbols) and 6.71 mM (open
symbols). For the circles $(t-t_f)/t_f$ is 0.3; for the triangles 0.5; and
for the squares 0.7. Theoretical curves are for $k = 1.1 \text{ s}^{-1}$. The applied
potential for $t = 0$, $t \leq \tau$ and $t > \tau$ is $-1.2 \text{ V}$, $-2.2 \text{ V}$ and $-1.2 \text{ V}$, respectively.
FIGURE IV-6

Cyclic voltammogram of 5.95 mM FlHNC(CH$_3$)$_2$ in DMF-0.1 F (n-Bu)$_4$ClO$_4$ at a planar platinum electrode. The scan was initiated in the negative-going direction at 0 V and a rate of 0.2 V/s. The small "2" indicates the second scan in the negative-going direction.
irreversible reduction of PhN=NPh\(^{-}\) to PhN=NPH\(^{2-}\) at \(-1.4\) V. The latter species and/or PhNH\(\text{H}^\text{N}^\text{Ph}\) then rapidly abstract a proton from FlHNMe\(_2\) to give FlNMe\(_2\)\(^{-}\); successive one-electron oxidations of FlNMe\(_2\)\(^{-}\) account for the two reversible couples that appear at \(-0.16\) and \(0.34\) V in Figure IV-7.

It should be noted that reduction of PhN=NPh gives substantially different results when FlHNMe\(_2\) is present than when FlHNH\(_2\) is present. Whereas PhN=NPH\(^{2-}\) proton abstracts from FlHNMe\(_2\) to give the expected products, PhNH\(\text{H}^\text{N}^\text{Ph}\) and FlNMe\(_2\)\(^{-}\), reduction of PhN=NPh in the presence of FlHNH\(_2\) causes a chain reaction to ensue in which PhN=NPh is reduced to PhNH\(\text{H}^\text{N}^\text{Ph}\) and FlHNH\(_2\) is oxidized to Fl=NH.\(^3\) Two important steps in the propagation cycle are oxidation of FlNH\(_2\)\(^{-}\) by PhN=NPh and the reduction of PhNH\(\text{H}^\text{N}^\text{Ph}\) by Fl=NH\(^{7-}\). A similar chain process occurs when PhN=NPh is reduced in the presence of 9-fluorenyl, FlHOH.\(^3\)

Finally, a comparison of Figures 6 and 7 will show that oxidation of FlNMe\(_2\)\(^{-}\) in the presence of FlH\(^{-}\) renders the FlNMe\(_2\)\(^{-}\)/FlNMe\(_2\)\(^{+}\) couple at \(-0.16\) V irreversible and eliminates the anodic wave at \(0.34\) V which is attributed to the reversible oxidation of FlNMe\(_2\)\(^{+}\) to FlNMe\(_2\)\(^{+}\). In addition, the oxidation of FlNMe\(_2\)\(^{-}\) in the presence of FlH\(^{-}\) affords a small, irreversible, cathodic peak on the second, negative-going half cycle at \(-1.72\) V. Since (1) the reduction of bifluorenyl, (FlH)\(_2\), to its anion radical occurs irreversibly with carbon-carbon bond cleavage near this potential,\(^2\) (2) the oxidation of FlH\(^{-}\) occurs irreversibly to give both (FlH)\(_2\) and FlH\(_2\)\(^{2-}\), and (3) FlNMe\(_2\)\(^{+}\) reacts rapidly in the presence of FlH\(^{-}\), we suspect that FlHFlNMe\(_2\) may be formed by the coupling of FlNMe\(_2\)\(^{+}\) with FlH\(^{+}\). It is important to add that relatively rapid reaction of FlNMe\(_2\)\(^{+}\) is observed even if the direct electrochemical
FIGURE IV-7

Cyclic voltammogram of 5.42 mM FLHN(CH₃)₂ and 1.21 mM azobenzene in DMF-0.1 M (n-Bu)₄NClO₄ at a planar platinum electrode. The scan was initiated at 0 V in the negative-going direction at a rate of 10 V/s. The temperature was 23°C.
oxidation of \( \text{FLH}^- \) is not effected. Because the rapid carbon-carbon cleavage of \((\text{FLH})_2^-\) probably precludes the coupling of \(\text{FLNMe}_2^-\) with \(\text{FLH}^-\) to give \(\text{FLHFLNMe}_2^-\), \(\text{FLH}^-\) must be generated by an electrocatalytic pathway in which electrogogenerated \(\text{FLNMe}_2^-\) is the electron acceptor (eqs. IV-17 thru -19).

\[
\begin{align*}
\text{FLNMe}_2^- & \longrightarrow \text{FLNMe}_2^- + e^- \quad \text{(IV-17)} \\
\text{FLNMe}_2^- + \text{FLH}^- & \longrightarrow \text{FLNMe}_2^- + \text{FLH}^+ \quad \text{(IV-18)} \\
\text{FLH}^+ + \text{FLNMe}_2^- & \longrightarrow \text{FLHFLNMe}_2 \quad \text{(IV-19)}
\end{align*}
\]

We have attempted to prove the feasibility of reactions IV-17 thru IV-19 by electrochemically oxidizing \(\text{FLNMe}_2^-\) in the presence of \(\text{FLH}^-\). Unfortunately, either the \(t\)-BuO\(^-\) which is used in excess to prepare \(\text{FLNMe}_2^-\) and \(\text{FLH}^-\) and/or the \(\text{FLH}^-\) itself must abstract a proton from the coupling product, \(\text{FLHFLNMe}_2\), to give \(\text{FL(NMe}_2)\text{Fl}^-\). The latter species is oxidized irreversibly near 0 V to give unidentified products. The anodic wave which is due to the oxidation of \(\text{FL(NMe}_2)\text{Fl}^-\) can be seen in the cyclic voltammograms of \(\text{FLH}NMe_2\) at certain combinations of temperature and scan rate as a shoulder on the \(\text{FLH}^-\) anodic peak.

Comparison of these results with those of previous workers. The electrochemical reduction of several N-substituted fluorenone and benzophenone imines has been reported to involve two steps of unequal magnitude in solvent systems where proton availability is low and a single two-electron step in protic media.\(^6,7\) Although the corresponding amine is the assumed imine reduction product in protic media, dimeric products have been postulated in the reduction of certain sterically unhindered imines in aprotic media.\(^7,8\) No evidence was obtained in this study for dimer formation upon reduction of either \(\text{FL} = \text{NH}\) or \(\text{Ph}_2\text{C} = \text{NH}\). A small amount of fluorene was obtained when \(\text{FL} = \text{NH}\)
was reduced in the presence of diethyl malonate, but this is presumed to be the result of inadvertent reduction of the more-difficultly-reduced amine product, $\text{FLNH}_2\text{H}$. 

Since neither the oxidation of $\text{FLNH}_2\text{H}^-$ and $\text{FLNMe}_2\text{H}^-$ nor the reduction of $\text{FLNH}_2\text{H}$ and $\text{FLNMe}_2\text{H}$ has been reported previously, we can only compare the redox behaviors of these compounds to those of the corresponding 9-hydroxy- and 9-methoxyfluorenes. While parallels exist, for example, $\text{FLOH}^-$ and $\text{FLNHPh}^-$ are oxidized catalytically by electrogenerated $\text{Fl}=\text{O}$ and $\text{Fl}=\text{NPh}$, respectively, significant differences are also seen, for example, electrocatalysis is only observed in the $\text{FLNHPh}^-$ system when a strong base is also present and no electrocatalysis could be detected for the $\text{FLNH}_2^-$ system under any solution conditions. The decomposition pathways for the anion radicals present another interesting contrast. Whereas hydroxide and alkoxide which are formed in the reductive cleavage of the carbon-oxygen bond of $\text{FLOH}$ and $\text{FLOCH}_3$, respectively, are unreactive with respect to the $\text{C}_9$ proton, amide formed by the reductive cleavage of the carbon-nitrogen bond in $\text{FLNH}_2\text{H}$ reacts only by abstracting the $\text{C}_9$ proton from unreacted starting material. The selectivity of amide is unexpected, since the $pK_a$'s of $\text{FlH}_2\text{H}$, $\text{H}_2\text{O}$, and $\text{NH}_3$ in dipolar, aprotic solvents are approximately $22, 8, 31, 9$ and $34, 10$ respectively.
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ELECTROCHEMICAL STUDIES OF BENZOPHENONE
AND FLUORENONE IMINES, AMINES AND
DIPHENYLDIAZOMETHANE

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ABSTRACT

The redox behavior of several nitrogen containing derivatives of fluorenone and benzophenone has been studied. In this continuation of the study of possible precursors to carbene anion radicals, diphenyldiazomethane (Ph$_2$CN$_2^+$) has been studied in acetonitrile (AN) containing Me$_4$NPF$_6$ (saturated) as the supporting electrolyte. This was done to confirm that the lifetime of the anion radical (Ph$_2$CN$_2^-$) is less than 1 ms in this electrolyte system. Parker and Bethell have recently reported that electroreduction of Ph$_2$CN$_2$ in AN-Me$_4$NPF$_6$ (saturated) at a hanging mercury drop electrode is electrochemically reversible at T = 8°C and v = 100 V/s. We have shown that their interpretation is in error, and that it is an artifact of unrecognized adsorption effects on the mercury electrode. The results reported here are consistent with previous findings using platinum as the cathode, i.e., the Ph$_2$CN$_2^-$ initially formed undergoes rapid loss of molecular nitrogen ($t^{1/2} \leq 1$ ms) to yield the carbene anion radical.

Other compounds reported are fluorenone imine (Fl=NH), N-phenylfluorenone imine (Fl=NPh), benzophenone imine (Ph$_2$C=NH), 9-aminofluorene (FlHNH$_2$) and 9-N,N-dimethylaminofluorene (FlHNMe$_2$). The imines have been shown to be reduced in two successive one-electron steps in aprotic media, first to the anion radical which is stable on the cyclic voltammetric time scale and then to the unstable dianion. However, on the coulometric time scale the anion radical was also shown to undergo a series of reactions leading to the amine in high yield. Oxidation of FlNHPh$^-$ to Fl=NPh in the presence of potassium-t-butoxide involves a kinetically controlled anodic peak ($E_{p,a} = -0.67$ V) which arises from the catalytic oxidation of FlNHPh$^-$ by
electrogenerated \( \text{Fl=NPh} \) and a second, irreversible, anodic peak at more positive potential \( E_{p,a} = -0.18 \text{ V} \) which is attributed to the direct electrochemical oxidation of \( \text{FlNHPh}^- \). Although electrocatalysis by \( \text{Fl=NH} \) is not observed in the corresponding \( \text{FlNH}_2^- \) system, the ease of \( \text{FlNH}_2^- \) oxidation is facilitated by the presence of \( \text{K-t-OBu} \) if reduction of \( \text{Fl=NH} \) to \( \text{FlNH}_2^2^- \) is also made to occur. The reductions of \( \text{FlHNH}_2 \) and \( \text{FlHNMe}_2 \) result in anion radicals which have half-lives in excess of one second at \(-51^\circ\text{C}\) and which react by unimolecular carbon-nitrogen bond cleavage. The amide anion formed by the decomposition process rapidly abstracts the \( C_9 \) proton from unreacted starting material; the other fragment anion, \( \text{FlH}^- \), also abstracts the \( C_9 \) proton from \( \text{FlHNR}_2 \), but at a rate which can be followed by cyclic voltammetry. The oxidation of \( \text{FlNMe}_2^- \) to the corresponding cation occurs in successive, one-electron steps in the absence of \( \text{FlH}^- \). If reaction of the electrogenerated \( \text{FlH}^- \) with \( \text{FlHNMe}_2 \) is incomplete when oxidation of \( \text{FlNMe}_2^- \) is effected, the intermediate radical, \( \text{FlNMe}_2^\cdot \), is interdicted by electrocatalytically formed \( \text{FlH}^\cdot \), presumably by coupling of \( \text{FlH}^\cdot \) and \( \text{FlNMe}_2^\cdot \) to give the 9-N,N-dimethylamino-9',9'-bifluorenyl, \( \text{FlHFlNMe}_2^\cdot \).