THERMOCHEMICAL DECOMPOSITION OF TNT: RADICAL IDENTIFICATION AND...

L P Davis, A G Turner, W R Carper

UNCLASSIFIED

FUSRL-TR-81-0002

AD-0099 044
THERMOCHEMICAL DECOMPOSITION
OF TNT: RADICAL IDENTIFICATION
AND THEORETICAL STUDIES

Larry P. Davis           John S. Wilkes
Almon G. Turner          R. Cameron Dorey
W. Robert Carper         Henry L. Pugh
Kenneth E. Siegenthaler

PROJECT 2303

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This technical report has been reviewed and is approved for publication.

R. CAMERON DOREY, III, Captain, USAF
Project Scientist

KENNETH E. SIEGENTHALER, Lt Colonel, USAF
Director
Directorate of Chemical Sciences

WILLIAM D. SIURU, JR., Lt Colonel, USAF
Commander

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Thermochemical Decomposition of TNT: Radical Identification and Theoretical Studies

The thermochemical decomposition of 2,4,6-trinitrotoluene (TNT) has been studied experimentally with EPR and theoretically with the MNDO and MNDO/3 methods. Analysis of the EPR spectra of pure TNT and hexamethylbenzene (HMB)-doped TNT at elevated temperatures has afforded an identification of one of the initial radicals formed in these systems. In both cases the radical is the one formed by removal of a nitro group from TNT. MNDO and MNDO/3 results indicate that removal of a nitro group from 1-nitropropene, used as a model system for...
TNT, is likely. Comparisons of the theoretical activation energy and the deuterium kinetic isotope effect for this step are in excellent agreement with measured values for these quantities early in the TNT decomposition. The nitro group removal is apparently facilitated by interaction with a second molecule in the system (HMB in the binary system or another TNT in the pure TNT system), which weakens the C-NO₂ bond either by transferring a hydrogen atom or forming a charge-transfer complex. Other potential first steps in the mechanism have been investigated theoretically.
THERMOCHEMICAL DECOMPOSITION OF TNT:
RADICAL IDENTIFICATION AND THEORETICAL STUDIES

By

Larry P. Davis
Almon G. Turner
W. Robert Carper
J. S. Wilkes
R. Cameron Dorey
Henry L. Pugh
Kenneth E. Siegenthaler

APRIL 1981

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Directorate of Chemical Sciences
Frank J. Seiler Research Laboratory
Air Force Systems Command
US Air Force Academy, Colorado 80840
PREFACE

The research described here was performed largely under Project 2303-F3-05 (Chemical Structure/Bonding Decomposition Relationship) and Project 2303-F4-03 (Molecular Orbital Calculations of Excited Species). A portion of the funding was provided by the AF Armament Laboratory under Project Order ATL-1-68. Mr. Thomas Floyd was the AFATL Program Manager.
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INTRODUCTION

The thermochemical decomposition of 2,4,6-trinitrotoluene (TNT) and related nitroaromatics in the liquid phase has been studied by a variety of techniques for many years. Janzen first observed radical formation during the thermochemical decomposition of nitroaromatics with electron paramagnetic resonance (EPR) techniques. We have studied the thermal decomposition kinetics of TNT by EPR. This paper compares EPR studies of the initial TNT radicals with MNDO molecular orbital calculations on TNT and model systems to identify the initial radicals produced and propose plausible first steps in the thermal decomposition of TNT.

EXPERIMENTAL

EPR spectra were recorded on a Varian E-109 spectrometer operating in the X-band using a standard TE$_{102}$ rectangular cavity as previously described. A custom-built variable temperature apparatus utilizing a Scanlon Co. heater maintained temperatures in the cavity to $\pm 1^\circ$C as measured by a Doric trendicator. TNT and trinitrobenzene (TNB) were synthesized using previously described procedures. Hexamethylbenzene (HMB) was purchased from the Aldrich Chemical Co. All compounds were purified by recrystallization from ethanol prior to use. Mixtures of compounds were prepared for EPR analysis by mixing the components in the molten state in quartz EPR tubes at temperatures below 170°C.

Simulation of EPR spectra for comparison with the experimental spectra was accomplished with a program written and generously provided by Dr. Ira Goldberg of Rockwell Science Center. This program is capable of simulating overlapping spectra from multiple radicals.
The MNDO and MINDO/3 molecular orbital programs were used to calculate heats of formation and geometries of all suspected intermediates and transition states for mechanisms tested. A related force constant program, also developed by M. J. S. Dewar, et al., was used to examine suspected transition states and provide vibrational frequencies in order to calculate thermodynamic quantities. Deuterium kinetic isotope effects for some steps also were calculated using these thermodynamic quantities.

RESULTS AND DISCUSSION

RADICAL IDENTIFICATION:

Previously we observed at least two radicals from the thermochemical decomposition of pure molten TNT. The final, or product, radical has been tentatively identified as a polymeric radical whose number of radical sites grows at an exponential rate throughout the decomposition. The initial radical, whose spectrum is composed of at least twenty lines, has eluded identification until now. By careful comparison in the wings of the spectrum of pure TNT with spectra generated by doping the TNT with a small amount of HMB, we find that the TNT spectrum is consistent with the radical structure below:

![Radical Structure](image)

Figure 1 shows a comparison of the spectra of the radical formed early in the decomposition of pure TNT at 245°C and a mixture of TNT and HMB (14:1 w:w) at 200°C. The spectrum exhibited by the HMB-doped TNT sample
EARLY TNT RADICAL SPECTRA

Figure 1

pure TNT

TNT/HMB

simulation
exhibits a large coupling constant of 10.3 G with an overall intensity pattern of 1:2:3:2:1, indicative of two equivalent nitrogens. Within each of these five groups, a six-line pattern is evident (coupling constant = 2.7 G) with intensity ratios of 1:5:10:10:5:1, indicative of five equivalent hydrogens. The computer simulated spectrum in Figure 1 using the measured couplings from the TNT/HMB spectrum shows a good match when two equivalent nitrogens and 5 equivalent hydrogens are assumed. The spectrum matches that of the initial radical from pure TNT decomposition in the wings. The simplest interpretation is that the five original hydrogens present in the TNT molecule are preserved in the radical, and one nitro group has been removed leaving two equivalent nitrogens. The apparent equivalence of the ring and methyl hydrogens has been noted for other radicals formed from TNT, in particular the radical anion\(^{23}\). It is also interesting to note that the HMB-doped sample produces the initial radical spectra at much lower temperatures than pure TNT (-200°C instead of 245°C) and the spectrum remains practically unchanged for over thirty minutes at 200°C.

The center portion of the neat TNT radical spectrum is not identical with the HMB-doped TNT spectrum, even though the wings are. Simulations which include a single line polymer radical spectrum with the HMB-doped TNT spectrum can be made identical with the neat TNT spectrum by varying the relative amounts of the two radicals (polymer + TNT radical), as shown in Figure 2. The neat TNT spectrum is thus apparently the superposition of the polymer radical spectrum on that of the TNT radical identified in the HMB-doped experiments.

A reaction analogous to the HMB-doped TNT decomposition apparently occurs during the thermal decomposition of 1,3,5-trinitrobenzene (TNB)
LATE TNT RADICAL SPECTRUM

Figure 2

pure TNT

simulation

5 G
doped with a small amount of HMB. TNB is thought to be very stable at temperatures at which TNT decomposes rapidly (-250°C). However, the HMB-doped TNB (TNB: HMB 82:78 w/w) produces a strong radical signal at 200°C which is quite stable as shown in Figure 3. An overall five-line pattern (coupling constant = 10.7 G) with a 1:2:3:2:1 intensity pattern again indicates two equivalent nitrogens. Within each of the five groups is a four-line pattern (coupling constant = 2.6 G) with intensity ratios of 1:3:3:1, indicative of three equivalent hydrogens. A simulation based on these values shows good agreement. Thus TNB doped with HMB appears to form a radical analogous to the TNT radical with HMB:

\[ \text{H}_2\text{ON} \]

MNDO CALCULATIONS:

Molecular orbital calculations using the MNDO and MNDO/3 methods developed by Dewar, et al.,19-22 have been carried out in this laboratory for the last several years. These methods have been shown to be useful in a variety of systems in the determination of plausible reaction mechanisms and kinetic parameters. We have applied these methods to the study of the initial steps of TNT and related nitroaromatics thermochemical decomposition. Calculated results for the activation energies and deuterium kinetic isotope effects of suspected mechanistic steps are compared with these experimentally-determined quantities using IDSC and EPR.
EARLY TNB RADICAL SPECTRUM

Figure 3
techniques. Also, structures of possible intermediates are compared with experimental spectra.

Because of the large amounts of computer time to completely optimize the geometries of the TNT species involved in suspected decomposition steps, we chose to use 1-nitropropene as a model system:

\[
\text{TNT} \quad \text{1-nitropropene}
\]

\[
\begin{align*}
\text{TNT} & \quad \text{1-nitropropene} \\
\begin{array}{c}
\text{CH}_3 \\
\text{O} \\
\text{N} \\
\text{N} \\
\text{2NO}_2 \\
\end{array} & \quad \\
\begin{array}{c}
\text{CH}_3 \\
\text{O} \\
\text{H} \\
\text{C=CH} \\
\text{N} \\
\text{O} \\
\end{array}
\end{align*}
\]

Previous experimental evidence\textsuperscript{15} had indicated involvement of a methyl group hydrogen in the rate-determining step. Based on this, six potential rate-determining steps were selected for calculations of \( \Delta H^\circ \) and \( E_a \):

\[
\begin{align*}
(1) \quad \text{TNT} & \rightarrow \text{X} \\
\text{X} = \text{NO}_2
\end{align*}
\]

\textit{Intramolecular H Transfer}
(2) TNT \[\rightarrow\] 

\text{Intramolecular O Insertion}

(3) 2 (TNT) \[\rightarrow\] 

\text{Intermolecular H Transfer}

\(X = \text{NO}_2\)
Step (5) is a removal of an \( \text{NO}_2 \) group (either para or ortho). Step (6) is a nitro group rearrangement to form a nitrite species.

In order to test the usefulness of the 1-nitropropene model in simulating TNT behavior, \( \Delta H^\circ \)'s for all six steps were calculated for both TNT and the 1-nitropropene model system, all geometries being optimized. The results in Table I are MNDO results. MINDO/3 results gave the same agreement between the two molecular systems that the MNDO results did.

**TABLE I**

Results of MNDO Calculations: Reaction \( \Delta H^\circ \)'s

<table>
<thead>
<tr>
<th>Step</th>
<th>( \Delta H^\circ ) (TNT System) (Kcal/mole)</th>
<th>( \Delta H^\circ ) (1-nitropropene system) (Kcal/mole)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) Intramolecular H Transfer</td>
<td>+16.6</td>
<td>+14.5</td>
</tr>
<tr>
<td>(2) Intramolecular O Insertion</td>
<td>-49.9</td>
<td>-49.3</td>
</tr>
<tr>
<td>(3) Intermolecular H Transfer</td>
<td>+16.8</td>
<td>+19.5</td>
</tr>
<tr>
<td>(4) Intermolecular O Insertion</td>
<td>-52.4</td>
<td>-48.5</td>
</tr>
<tr>
<td>(5) NO(_2) Removal</td>
<td>+32.3</td>
<td>+26.9 (ortho-NO(_2))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+32.5 (para-NO(_2))</td>
</tr>
<tr>
<td>(6) Nitro Rearrangement</td>
<td>-38.4</td>
<td>-41.1 (para-NO(_2))</td>
</tr>
</tbody>
</table>
Thus it appears that the nitropropene system is at least a potential model for simulating TNT decomposition steps. It is a good model if the calculated activation energies are also similar.

We simulated the reaction pathway for each of the six steps by defining a reaction coordinate, fixing it at various intermediate points along the pathway, and then allowing all other geometric variables to optimize at each value of the reaction coordinate. If a maximum was located along the pathway, the transition state optimization program was run. Finally, the vibrational force constant program was employed on the transition state geometry to support the existence of a transition state by the presence of one negative vibrational force constant, which is identified as the reaction coordinate. Each of the six steps will be discussed in turn.

The intramolecular hydrogen transfer (1) calculations for 1-nitropropene produced a species which proved to be a transition state with an activation energy of 63.8 kcal/mole. The most recently measured value (IPR) for the activation is 40.9 kcal/mole\(^{17}\). The optimized geometry from the calculations was intermediate between that of the reactant and product, with the O-H bond order being 0.48 and the C-H bond order 0.39. The geometry is shown in Figure 4. A calculation of a deuterium kinetic isotope effect utilizing the calculated force constants and moments of inertia yielded a \(k_D/k_H\) value of 2.2 at 227°C. The measured value of \(k_D/k_H\) for the rate-determining step of TNT decomposition near this same temperature is 1.15. Thus the calculations do not rule out this intramolecular hydrogen transfer as the rate-determining step for TNT thermochemical decomposition, particularly since MNDO usually overestimates activation energies for hydrogen transfer reactions.
TRANSITION STATE FOR INTRAMOLECULAR HYDROGEN TRANSFER

Figure 4

TRANSITION STATE FOR INTERMOLECULAR OXYGEN INSERTION

UNLABELED ATOMS ARE HYDROGEN

Figure 5
On the other hand, the intramolecular oxygen insertion (2) calculations for 1-nitropropene gave very high energy barriers even though $\Delta H^\circ$ for this step is quite negative. In one case for which the methyl group was not allowed to rotate while the oxygen was moved toward the carbon, the molecule dissociated into acetylene and methyl nitrite with an activation energy of 96 kcal/mole. If complete relaxation of the methyl group was allowed, no stable product was formed. Thus it appears that this step is not likely during the thermochemical decomposition of TNT.

The intermolecular hydrogen transfer (3) transition state has not yet been located, but approximations point to an activation energy of 82 kcal/mole. This step also appears to have too high an activation energy to be a viable candidate for the first step of TNT decomposition.

The intermolecular oxygen insertion reaction (4) produced a transition state with an activation energy of 110 kcal/mole. The geometry of this transition state is shown in Figure 5.

We also investigated rearrangement of an $\text{NO}_2$ group and removal of an $\text{NO}_2$ group from the molecule, even though these steps would not produce a primary deuterium kinetic isotope effect as indicated by the previous experimental evidence. We found that the calculated dissociation energy for 1-nitropropene of 32.3 kcal/mole agreed well with the experimental value for the activation energy, particularly since MNDO typically underestimates $\text{C-NO}_2$ bond energies. In addition, the calculated deuterium kinetic isotope effect at 227°C for $\text{NO}_2$ removal was 1.7, even though it is a secondary kinetic isotope effect. This, too, agrees well with the experimental value of approximately 1.7. The nitro group rearrangement to nitrite for 1-nitropropene had a calculated activation energy of 52.4 kcal/mole.
slightly more than the experimental number for TNT. The geometry of the
transition state is shown in Figure 6. This step cannot be ruled out as a
potential first step. In addition, if NO then dissociates from the molecule,
the resulting radical, with an oxygen on the para position, could give
qualitatively the experimentally observed EPR spectrum.

Table II summarizes activation energies and deuterium kinetic isotope
effects for both 1-nitropropene and TNT. The activation energy given for
TNT for the intramolecular hydrogen transfer step is a lower bound, and
the one for the intermolecular hydrogen transfer is an upper bound.

TABLE II
RESULTS OF MNDO CALCULATIONS:

<table>
<thead>
<tr>
<th>PROCESS</th>
<th>$E_a$ (1-NP)</th>
<th>$E_a$ (TNT)</th>
<th>$K_{H}/K_D$ (1-NP)</th>
</tr>
</thead>
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<tr>
<td>Intramolecular H Transfer</td>
<td>63.8</td>
<td>&gt;55</td>
<td>2.2</td>
</tr>
<tr>
<td>Intramolecular O Insertion</td>
<td>96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NO$_2$ Removal</td>
<td>32.3</td>
<td>26.9 (o-NO$_2$)</td>
<td>1.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>32.5 (p-NO$_2$)</td>
<td></td>
</tr>
<tr>
<td>Nitro Rearrangement</td>
<td>52.4</td>
<td>&lt;122</td>
<td></td>
</tr>
<tr>
<td>Intermolecular H Transfer</td>
<td>-82</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermolecular O Insertion</td>
<td>110</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CONCLUSIONS

An initial radical in the thermochemical decomposition of TNT and
related nitroaromatics is one in which a nitro group or an NO group has
been removed from the TNT molecule. MO calculations for removal of a
nitro group from the molecule are consistent with experimental evidence
TRANSITION STATE FOR

NITRO-NITRITE REARRANGEMENT

Figure 6
both for the activation energy and the deuterium kinetic isotope effect.
The enhancement of radical production upon addition of HMB indicates that
the interaction of a second species results in much easier NO₂ removal.
Two likely mechanisms arise:

1. The donor species (HMB) transfers a hydrogen atom to the acceptor
   species (TNT or TNB). The removal of HONO can then be possibly accomplished
   more easily than the original nitro group.

2. A charge-transfer complex (HMB/TNT or HMB/TNB) is formed which
   weakens in some manner the C-nitro group bond. Thus NO₂ can be removed
   more easily.

The calculations indicate that oxygen atom insertions are unlikely due
to the high energies of the transition states involved. Also, the activa-
tion energy for intermolecular hydrogen transfer has a value which is
probably inconsistent with the experimental value. Intramolecular hydrogen
transfer or nitro group rearrangement to the nitrite cannot be ruled out as
the initial step, although they appear to be less likely than simple nitro
group removal.
REFERENCES


ABBREVIATIONS, ACRONYMS, AND SYMBOLS

TNT - 2,4,6 Trinitrotoluene
EPR - Electron Paramagnetic Resonance
MNDO - Modified Neglect of Diatomic Overlap
MNDO/3 - Modified Intermediate Neglect of Differential Overlap/3
TE\textsubscript{102} - Transverse Electric Mode 102
TNB - 1,3,5 Trinitrobenzene
HMB - Hexamethylbenzene
G - Gauss
w:w - Weight to Weight Ratio
°C - Degrees Centigrade
AH° - Change of enthalpy for the reaction
E\textsubscript{a} - Activation energy for the reaction
IHSCE - Isothermal Differential Scanning Calorimeter
X - Nitro group
K\textsubscript{H}/K\textsubscript{D} - Deuterium Kinetic isotope effect
ATELMED 8