Experimental and Theoretical Studies of Reactions Important in Photochemical Smog: Aromatics and Alkanes

Thesis by
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Appearing below are two quotes that guided me in the past, and became even more meaningful during my time at Caltech.

*El que no busca, no encuentra.*
*El que no escucha, no aprende.*
*El que no trata, fracasa, con el alma y con la mente.*
*Cuando hay fe, hay con que hacer lo que hay que hacer.* - Rubén Blades

Knowledge of the effort required to carry us this distance should propel us in the right direction and help us bear up under our own burdens.

Continuing the struggle is the best way we have of saying thank you to those who struggled for us. - Eric V. Copage
Abstract

The development of effective ozone control strategies requires the use of atmospheric models. There is general agreement within the scientific community that several aspects of the chemistry within these models has yet to be fully elucidated, and is influential in their predictions. The work in this thesis is aimed at trying to determine some of the unknown aspects of the mechanisms of important atmospheric species. Specifically, the gas-phase reactions of two large alkanes, 2,2,4-trimethylpentane and 2,2,5-trimethylhexane are investigated. These two alkanes are present in urban air, and are potential aerosol presursors. The chemistry of several aromatic hydrocarbons are also studied using both theoretical and experimental techniques. The effects of NO$_2$ on the photooxidation of toluene, $m$-xylene, and $p$-xylene are examined, and mechanisms of each of these organics are thoroughly evaluated through closely coordinated laboratory work and computer modeling. In addition, product studies of the photooxidation of 1,2,4-trimethylbenzene and $m$-ethyltoluene are conducted. These studies provide the first identification of ring-retained products from 1,2,4-trimethylbenzene, and ring-retained and fragmented products from $m$-ethyltoluene.
A new indoor experimental reactor was designed to investigate gas-phase reaction kinetics and mechanisms. This new system has served to launch the atmospheric chemistry program at Caltech into a wide variety of new research topics.
Contents

1 Introduction ........................................... 1

2 Experimental System ................................ 9
   2.1 Photochemical Reactor ............................ 10
   2.2 Experimental System Characterization ......... 19
      2.2.1 Ultraviolet Light Intensity ............... 19
      2.2.2 Methyl Nitrite Chemistry .................. 22
      2.2.3 t-Butyl Alcohol Experiments .............. 31

3 Photooxidation of 2,2,4-Trimethylpentane and 2,2,5-Trimethylhexane: A Product Study 36
   3.1 Introduction ..................................... 38
   3.2 Experimental ...................................... 39
   3.3 Rate Constant of 2,2,5-Trimethylhexane-OH Reaction .... 44
   3.4 Photooxidation Products ......................... 44
      3.4.1 2,2,4 Trimethylpentane ...................... 46
      3.4.2 2,2,5 Trimethylhexane ....................... 58
   3.5 Atmospheric Reaction Mechanisms ................. 60
3.5.1 2,2,4-Trimethylpentane ........................................... 70
3.5.2 2,2,5-Trimethylhexane ........................................... 75
3.6 Conclusions .............................................................. 82

4 Mechanism of Atmospheric Photooxidation of Aromatics: A
Theoretical Study ........................................................ 86
4.1 Introduction .............................................................. 88
4.2 Calculation Methods .................................................. 89
4.3 Results of Calculations .............................................. 99
  4.3.1 Initial Hydroxyl Radical Attack ................................. 99
  4.3.2 Fate of the OH-Aromatic Adduct: NO₂ Reaction ............. 111
  4.3.3 Fate of the Peroxy Radical: Cyclization vs NO Reaction.... 119
  4.3.4 Bicyclic Peroxy Radicals ...................................... 125
  4.3.5 Ring Fragmentation Products .................................. 130
4.4 Conclusion ............................................................... 133

5 Photooxidation of Aromatic Compounds: Evidence for an
NO₂ Effect ................................................................. 138
5.1 Introduction .............................................................. 140
5.2 Experimental Procedure ............................................. 145
5.3 Effect of NO₂ on Product Yields in Aromatic Photooxidation .. 148
5.4 Aromatic Photooxidation Mechanisms ............................. 163
  5.4.1 Aromatic-OH Adduct Reaction with O₂ ...................... 174
  5.4.2 Aromatic-OH Adduct Reaction with NO₂ .................... 175
  5.4.3 Ring Fragmentation Products .................................. 178
5.5 Implications for Atmospheric Aromatic Photooxidation Mechanisms

6 Photooxidation of \( m \)-Ethyltoluene and 1,2,4 Trimethylbenzene: A Product Study 183
6.1 Introduction 185
6.2 Experimental 188
6.3 1,2,4-Trimethylbenzene Photooxidation 191
   6.3.1 Ring-Retained Products 191
   6.3.2 Ring-Fragmentation Products and the NO\(_2\) Effect 195
6.4 \( m \)-Ethyltoluene Photooxidation 201
   6.4.1 Ring-Retained Products 201
   6.4.2 Ring-Fragmentation Products and the NO\(_2\) Effect 204
6.5 Conclusions 205

7 Conclusions 212

A Calculation of the Photolysis Rate Constant, \( k_1 \) 218
A.1 Reactions of NO\(_2\) 219
A.2 Calculations 219

B Experimental Protocol 223
B.1 Before an Experiment 224
   B.1.1 NO\(_2\) Monitor Calibration 224
   B.1.2 O\(_3\) Analyzer Calibration 225
   B.1.3 GC-FID Calibration 225
B.2 Running an Experiment ........................................... 226
  B.2.1 Admitting Reactants ........................................... 226
  B.2.2 Sampling From the Bag ....................................... 228
B.3 Teflon Chamber .................................................... 229
  B.3.1 Deciding When to Change a Bag .............................. 229
  B.3.2 Making a New Chamber ....................................... 229
  B.3.3 Conditioning the Bag ......................................... 230
B.4 2,4 Dinitrophenylhydrazine Analyses ............................ 230
  B.4.1 DNPH-Carbonyl Derivatization ............................... 232
  B.4.2 Calculation of Carbonyl Yields .............................. 234
  B.4.3 Interferences with the DNPH Technique ................... 236

C Energies of Aromatic Intermediates ............................... 237

D Calculation of the Average $[\text{NO}_2]$ ....................... 253
## List of Tables

1.1 Tropospheric Lifetimes of Selected Organics ........................ 4

2.1 Equipment For Indoor Photochemical Reactor Experiments .... 17
2.2 Rate constants generated from the Carter program after normalization .................................. 22
2.3 Simplified CH$_3$ONO mechanism used in modeling ............ 24

3.1 OH Radical rate constants used to correct product yields ...... 56
3.2 Corrected Carbonyl Photooxidation Product Yields .......... 57
3.3 2,2,4-Trimethylpentane Mechanism .............................. 61
3.4 2,2,5-Trimethylhexane Mechanism ................................. 65

4.1 Toluene Photooxidation Products: Molar Yields ............... 90
4.2 $m$-Xylene Photooxidation Products: Molar Yields .......... 90
4.3 $p$-Xylene Photooxidation Products: Molar Yields .......... 91
4.4 Relative energies in kcal/mol for isomers of the methylhydroxycyclohexadienyl radical using various computational methods 94
4.5 A comparison of relative energies of different structural isomers based on experimental data and theoretical calculations 96
4.6 A comparison of experimentally and theoretically derived $\Delta H_{rxn}$ values for the Toluene + OH reaction. .................. 97
4.7 PM3 equilibrium structures for a representative selection of the most stable reaction products of aromatics with OH. ... 107
4.8 Calculated energies (in a.u.) of the most stable reaction intermediates for the pathway leading to the formation of the bicyclic peroxy radicals. ......................... 108
4.9 PM3 derived geometries for (a) equilibrium and (b) approximate transition state structures for the reactions of aromatic-OH adducts with NO$_2$. All bond distances are in Å, and angles are in degrees. ......................... 114
4.10 Calculated energy (in a.u.) of molecules involved in reactions with aromatic intermediates. ......................... 116
4.11 Calculated energies (in a.u.) of the products formed by the reaction of NO$_2$ with the toluene-OH adduct. ................. 117
4.12 Approximate transition state energies for the reactions of aromatic-OH adducts with O$_2$, represented in terms of the structure of the aromatic-OH adduct. ......................... 120
4.13 PM3 equilibrium structures for a representative selection of the most stable reaction products of aromatic-OH adducts with O$_2$ (Reaction 2b). All bond distances are in Å, angles are in degrees. ......................... 121
4.14 PM3 equilibrium structures for a representative selection of the most stable reaction products in the formation of bicyclic peroxyl radicals from peroxyl radical intermediates (Reaction 2b). .......................... 123

4.15 Approximate transition states for the addition reactions of bicyclic peroxyl radicals with O₂. ........................................ 124

4.16 Calculated energies (in a.u.) for the products of the reaction of NO with a representative selection of the most stable peroxyl intermediates and for the products of the reaction of O₂ with a representative selection of the most stable bicyclic intermediates. 128

4.17 Approximate transition states for the formation of the bicyclic peroxyl radical. .................................................. 129

5.1 Toluene Photooxidation Products: Molar Yields .............. 142

5.2 m-xylene Photooxidation Products: Molar Yields .............. 143

5.3 p-xylene Photooxidation Products: Molar Yields .............. 143

5.4 OH radical rate constants used to correct product yields. .... 161

5.5 Ranges of correction values, F, used for each quantified product. 162

5.6 Names and structures of species appearing in the aromatic mechanisms of Tables 5.7, 5.8, and 5.9. ......................... 166

5.7 Toluene Mechanism .................................................. 167

5.8 m-xylene Mechanism ............................................... 169

5.9 p-xylene Mechanism ............................................... 172

6.1 1,2,4 Trimethylbenzene Photooxidation Products: Molar Yields 186
B.1 Conversion Factors for Selected Carbonyls . . . . . . . . . . . . 236
List of Figures

2.1 Schematic of the Indoor Photochemical Reactor. 11
2.2 Photograph of the Indoor Photochemical Reactor. 12
2.3 Absorbance spectrum of 2 mil FEP teflon film. 13
2.4 Spectral distributions of black lamps and selected compounds. 15
2.5 Spectral distributions of black lamps compared to sunlight spectrum. 16
2.6 Schematic of the glass vacuum manifold which is used to admit reactants to the chamber. 18
2.7 Spectrum of methyl nitrite. 23
2.8 Experimental and theoretical data for NO for the photolysis of CH₃ONO. 25
2.9 Simulation of CH₃ONO system, no HCHO reactions. 27
2.10 Simulation of CH₃ONO system. HCHO included, initial NO > 0. 28
2.11 Simulation of CH₃ONO system. HCHO included, initial NO = 0. 29
2.12 Comparison of O₃ data from simulations of CH₃ONO chemistry. 30
2.13 Yield of acetone from the photooxidation of t-Butyl Alcohol... 33

3.1 Relative rate study to determine $k_{OH}$ for 2,2,5-trimethylhexane. 45
3.2 2,2,4-Trimethylpentane + OH Reactions, Tertiary . . . . . . . . . 47
3.3 2,2,4-Trimethylpentane + OH Reactions, Secondary . . . . . . . . 48
3.4 2,2,4-Trimethylpentane + OH Reactions, Primary . . . . . . . . . 49
3.5 2,2,4-Trimethylpentane + OH Reactions, Primary . . . . . . . . . 50
3.6 2,2,5-Trimethylhexane + OH Reactions, Tertiary . . . . . . . . . 51
3.7 2,2,5-Trimethylhexane + OH Reactions, Secondary . . . . . . . . . 52
3.8 2,2,5-Trimethylhexane + OH Reactions, Secondary . . . . . . . . . 53
3.9 2,2,5-Trimethylhexane + OH Reactions, Primary . . . . . . . . . 54
3.10 2,2,5-Trimethylhexane + OH Reactions, Primary . . . . . . . . . 55
3.11 Observed and predicted concentrations for 2-methyl-1-propanal from 2,2,4-trimethylpentane photooxidation. . . . . . . . . . . . . . . . 71
3.12 Observed and predicted concentrations for acetone from 2,2,4- trimethylpentane photooxidation. . . . . . . . . . . . . . . . . . . . . . 72
3.13 Observed and predicted concentrations for formaldehyde from 2,2,4-trimethylpentane photooxidation . . . . . . . . . . . . . . . . . . 73
3.14 Observed and predicted concentrations for 3,3-dimethylbutyraldehyde from 2,2,5-trimethylhexane photooxidation. . . . . . . . . . . . . 77
3.15 Observed and predicted concentrations for acetone from 2,2,5- trimethylhexane photooxidation. . . . . . . . . . . . . . . . . . . . . . 78
3.16 Observed and predicted concentrations for acetaldehyde from 2,2,5-trimethylhexane photooxidation. . . . . . . . . . . . . . . . . . . 79
3.17 Observed and predicted concentrations for formaldehyde from 2,2,5-trimethylhexane photooxidation. 80

4.1 Possible reactions in a generalized mechanism of Aromatic-OH photooxidation. 100

4.2 Reaction coordinate diagram for toluene. 101

4.3 Reaction coordinate diagram for m-xylene. 102

4.4 Reaction coordinate diagram for p-xylene. 103

4.5 Reaction coordinate diagram for 1,2,4-trimethylbenzene. 104

4.6 Reaction coordinate diagram for m-ethyltoluene. 105

4.7 $\Delta H_{rxn}$ values for Reactions 4 and 5 for different aromatic systems. 126

4.8 Subsequent reaction of the oxy radicals. 127

4.9 $\beta$-Scission of a bicyclic oxy radical of toluene to give ring fragmentation products. 131

4.10 Fragmentation products formed from the bicyclic oxy radicals. 132

5.1 Initial steps in the toluene-OH reaction. 141

5.2 Toluene Photooxidation- Glyoxal. 151

5.3 Toluene Photooxidation- Methyl Glyoxal. 152

5.4 Toluene Photooxidation- Cresols. 153

5.5 m-xylene Photooxidation- Glyoxal. 154

5.6 m-xylene Photooxidation- Methyl Glyoxal. 155

5.7 m-xylene Photooxidation- Dimethylphenols. 156

5.8 m-xylene Photooxidation- Nitro-m-xylene. 157

5.9 p-xylene Photooxidation- Glyoxal. 158
5.10  p-xylene Photooxidation- Methyl Glyoxal .......................... 159
5.11  p-xylene Photooxidation- Dimethylphenol .......................... 160
5.12  Reactions in a generalized mechanism of Aromatic-OH photooxidation. .............................................................. 164
5.13  Simulation of nitro-m-xylene formation ............................. 177
6.1  Initial reactions of the OH-initiated reaction of toluene ........ 187
6.2  Abstraction pathway for 1,2,4-trimethylbenzene .................. 192
6.3  GCD chromatogram of products formed from 1,2,4-trimethylbenzene photooxidation ................................................. 193
6.4  Pathways for the trimethylhydroxycyclohexadienyl radical reaction with O₂ ....................................................... 194
6.5  Pathway for ring-fragmentation formation in 1,2,4-trimethylbenzene photooxidation ................................................. 196
6.6  1,2,4 Trimethylbenzene Photooxidation Results: Methyl Glyoxal 198
6.7  1,2,4 Trimethylbenzene Photooxidation Results: Glyoxal ......... 199
6.8  1,2,4 Trimethylbenzene Photooxidation Results: CH₃CHO ..... 200
6.9  OH abstraction pathways for m-ethyltoluene photooxidation ...... 202
6.10 GCD chromatogram of products formed from m-ethyltoluene photooxidation ......................................................... 203
6.11 m-Ethyltoluene Photooxidation Results: CH₃CHO .................. 206
6.12 m-Ethyltoluene Photooxidation Results: Glyoxal .................. 207
6.13 m-Ethyltoluene Photooxidation Results: Methyl Glyoxal ..... 208
B.1  2,4 Dinitrophenylhydrazine Derivatization Reaction ............. 231
C.11 Energy differences for m-ethylphenylene Radicals.
C.10 Energy differences for 1,2,4-trimethylphenylene Radicals.
C.09 Energy differences for 1,2,4-trimethylphenylene Radicals.
C.08 Energy differences for 1,2,4-trimethylphenylene Radicals.
C.07 Energy differences for p-xylene Radicals.
C.06 Energy differences for m-xylene Radicals.
C.05 Energy differences for m-xylene Radicals.
C.04 Energy differences for m-xylene Radicals.
C.03 Energy differences for toluene Radicals.
C.02 Energy differences for toluene Radicals.
C.01 Energy differences for toluene Radicals.

R.2 Calibration curves for DNP-ethylglyoxal and DNP-methylglyoxal.
Chapter 1

Introduction
With the advent of industrialization came an increase in the emission of primary pollutants such as hydrocarbons and oxides of nitrogen. In the presence of sunlight, these primary pollutants react to form photochemical smog. To date, photochemical smog has been identified as the cause of adverse health effects in humans, phytotoxicity, the deterioration of buildings, and the reduction of visibility and solar radiation, among other things. For these reasons, it is important to fully understand and characterize the components leading to the formation of photochemical smog.

In order to control and abate the harmful effects of photochemical smog, its main components must be defined, and limits to their concentrations established. National ambient air quality standards (NAAQS) have been set for photochemical smog components such as ozone, nitrogen dioxide, and nitric oxide. The level and frequency of exceedences in particular areas of the country determine when air pollution control strategies are necessary. In the Los Angeles area, the main concern has focused particularly on extremely high levels of ozone. Agencies such as the Air Quality Management District and the California Air Resources Board have encouraged an increase in carpooling efforts and decreased use of items that result in high evaporative emissions of hydrocarbons (such as lighter fluid, commonly used in barbecues). The efforts of these two agencies have resulted in the improvement of air quality in Los Angeles, and a reduction in the number of exceedences of the NAAQS for ozone over the last decade.

Predictive computer modeling has been used to establish and evaluate proposed air pollution control strategies. However, many important issues related to these predictive models have yet to be fully elucidated. Mod-
els which currently exist include four components: chemistry, meteorology, transport mechanisms, and emissions data. In order to devise an effective air pollution control strategy, each of these components must reasonably represent the actual conditions present in the atmosphere. The work in this thesis is aimed at addressing two of the important, and as of yet unsettled issues in the chemistry portion of atmospheric models: the atmospheric chemistry of large alkanes and aromatic compounds.

The major loss processes for hydrocarbons in the atmosphere involve reaction with the oxidants $O_3$, $OH$, and $NO_3$. Rate constants for the reactions of many organics with these oxidants have been determined, and compilations of these values appear in Atkinson, 1989 and Atkinson, 1994. From these rates constants, and assuming concentrations of $O_3$, $OH$, and $NO_3$ of $7 \times 10^{11}$, $1.5 \times 10^6$, and $5 \times 10^8$ molecules cm$^{-3}$, respectively, lifetimes of organic compounds with respect to reaction with the three major oxidants can be determined [5]. Table 1.1, adapted from Seinfeld et al., 1995, details the tropospheric lifetimes of selected hydrocarbons. For the aromatics and the alkanes listed, it is evident that the major atmospheric loss process for these two classes of compounds are reactions with hydroxyl radicals. Thus, it is these reactions with hydroxyl radicals that are the focus of this thesis.

When hydrocarbons react in the atmosphere, they inevitably form per-oxy radicals. In the polluted troposphere, these peroxy radicals convert NO to $NO_2$. The formation of $NO_2$ is significant in that its subsequent photolysis is the principal mechanism for ozone formation. The reactions can be summarized as:
Table 1.1: Tropospheric Lifetimes of Selected Organics

<table>
<thead>
<tr>
<th>Organic</th>
<th>Lifetime due to reaction with:</th>
<th>OH</th>
<th>NO₃</th>
<th>O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methane</td>
<td>9.6 yr</td>
<td>&gt;120 yr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethane</td>
<td>60 d</td>
<td>&gt;12 yr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Propane</td>
<td>13d</td>
<td>&gt;2.5 yr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethene</td>
<td>1.8 d</td>
<td>225 d</td>
<td>9.7 d</td>
<td></td>
</tr>
<tr>
<td>Propene</td>
<td>7.0 h</td>
<td>4.9 d</td>
<td>1.5 d</td>
<td></td>
</tr>
<tr>
<td>Methanol</td>
<td>17 d</td>
<td>&gt;77 d</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethanol</td>
<td>4.7 d</td>
<td>&gt;51 d</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzene</td>
<td>12.5 d</td>
<td>&gt;6 yr</td>
<td>&gt;4.5 yr</td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>2.6 d</td>
<td>1.9 yr</td>
<td>&gt;4.5 yr</td>
<td></td>
</tr>
<tr>
<td>m-Xylene</td>
<td>7.8 h</td>
<td>200 d</td>
<td>&gt;4.5 yr</td>
<td></td>
</tr>
</tbody>
</table>

Hydrocarbon + •OH → R•
R• + O₂ → RO₂
RO₂ + NO → RO• + NO₂
NO₂ + hν → O• + NO
O• + O₂ → O₃

Thus NO₂ plays an important role in the formation of photochemical smog.

The kinetics and mechanisms of small alkanes have been widely characterized[2]. However, the only available literature data concerning the larger alkanes (> C₈) are kinetic data [3, 1]. The mechanisms of large alkane photooxidation have not been widely investigated. Many of these compounds are used in gasoline and industries. The potential for release of alkanes into the atmosphere is thus quite high. These alkanes can react in the atmosphere according to the reactions detailed above to eventually generate ozone and adversely affect air quality. The large alkanes are especially important. In
addition to their ozone forming potential, compounds with more than six
 carbons have been identified as potential aerosol precursors. It is therefore
 important to fully characterize the gas-phase mechanism of the large alkane
 reactions. This thesis focuses on the gas-phase OH radical reactions of two
 large, branched alkanes: 2,2,4-trimethylpentane and 2,2,5-trimethylhexane.

The atmospheric chemistry of several of the more common aromatics
(benzene, toluene, m- and p-xylene) have been investigated in the past[2].
The major atmospheric sink of this class of compounds is reaction with hy-
droxyl radicals via both the addition of OH to the ring and abstraction of
an H atom from the ring. However, the addition pathway dominates to form
a methyl substituted hydroxycyclohexadienyl radical. It is this radical and
its subsequent fate that are in question. Our approach to the question of
the fate of the radical was two-fold: experimental and theoretical in na-
ture. From the experimental perspective, our goal was to investigate the
effect of NO₂ on the reactions of the aromatic-oh adduct. Recent evidence
has suggested that the role of NO₂ may be more important than previously
believed [4]. We therefore undertook a systematic laboratory study of the
effects of NO₂ on the toluene, m-xylene, p-xylene, m-ethyltoluene, and 1,2,4-
trimethylbenzene systems. From the theoretical perspective, our goal was
to identify the intermediates believe to form from the susequent reactions
of the methyl substituted hydroxycyclohexadienyl radicals. Semi-empirical
and ab initio techniques were used to determine the energies of all possible
intermediates in the same aromatic systems which were studied experiment-
tally. The results of the theoretical calculations were compared to available
product data, and full mechanisms, from initial OH radical attack to stable
product formation, were developed for each aromatic.
Bibliography


Chapter 2

Experimental System
2.1 Photochemical Reactor

A new indoor photochemical reactor was designed and constructed to study detailed gas-phase kinetics and mechanisms. A schematic of the reactor appears in Figure 2.1, and a photograph of the chamber is in Figure 2.2. The system provides for excellent control of experimental parameters such as light intensity and temperature. The reaction chamber is a batch reactor composed of 2 mil Teflon FEP film, with a volume of approximately 1000 liters. Teflon film has a low absorbance in the ultraviolet wavelength region of critical importance to reactions in the troposphere (in the region around 350 nm), and is sufficiently versatile to allow for ease in cleaning or disposal. An absorbance spectrum of the teflon used appears in Figure 2.3. This spectrum was generated using a variable wavelength Fourier transform spectrometer (Nicolet SX 400).

Artificial light (two banks of 24 Sylvania F30T8BL black lamps) is used to initiate the chemistry within the chamber. The lamps are mounted on a specially designed reflective surface to provide uniformity in irradiation. There are many different types of lamps which could have been used to generate artificial light. These particular ultraviolet lamps were chosen because of the similarity of their spectral distribution to that of the radiation reaching the earth. The sun emits a variety of wavelengths of light. Radiation reaching the stratosphere contains wavelengths greater than 180 nm. Through the stratosphere, the radiation is filtered by ozone so that only wavelengths greater than 290 nm actually reach the earth. These particular black lamps emit most of their light in wavelength regions which are crucial for tropo-
Figure 2.1: Schematic of the Indoor Photochemical Reactor.
Figure 2.2: Photograph of the Indoor Photochemical Reactor.
Figure 2.3: Absorbance spectrum of 2 mil FEP teflon film.
spheric chemistry, thereby allowing for the simulation of atmospheric activity. The spectral distribution of a Sylvania F30T8BL black lamp appears in Figure 2.4, spectrum (A), and was obtained directly from Sylvania. The sharp features in the lamp spectrum are mercury absorption lines which can be attributed to the material used to construct the lamps. For comparison, the spectral absorption of NO₂, formaldehyde, and methyl glyoxal appear in Figure 2.4, (B), (C), and (D), respectively. The spectral data for these plots were obtained from Bass et al., 1983 and Moortgat et al., 1983 for formaldehyde, NASA, 1987 for NO₂, and Plum et al., 1993 for methyl glyoxal. These three compounds are significant to atmospheric processes, and are representative of the types of compounds that would be present in indoor chamber studies. A comparison of the spectral distribution of sun at two different zenith angles, 0 and 40 degrees, and the black lights appears in Figure 2.5. The light intensity within the chamber can be adjusted by varying the number of lamps from the minimum of 2 lamps to a maximum of 48 lamps.

A feedback-controlled cooling system which blows cool air around the chamber is used to control the temperature within the chamber. Temperature probes are located both inside of the bag and between the bag and the lamps. The temperature probe located on the outside of the bag is used to control the temperature of the system. The probe inside of the bag is used to record the actual bag temperature. The system is ordinarily set to operate at 298±2 K.

The reaction chamber is coupled to a glass vacuum manifold (input) and a glass sampling manifold (output). The glass vacuum manifold, pictured in Figure 2.6, has several detachable calibrated bulbs (only one is pictured)
Figure 2.4: (A) Spectral distribution of black lamps. (B) Absorbance Coefficient * Quantum Yield for NO₂. (C) Absorbance Coefficient * Quantum Yield for HCHO. (D) Absorbance Coefficient * Quantum Yield for methyl glyoxal.
Figure 2.5: Scaled spectral distribution of sun at zenith angles of 0 and 40 degrees as compared to Sylvania F30T8BL lamps.
which are used to admit known amounts of volatile reactants (such as methyl or ethyl nitrite) to the chamber. The evaporation bulb is used to admit organics by gently evaporating liquid samples with a heat gun while flowing air through the bulb. The sampling manifold has several ports (with stopcocks on them) which are used to deliver gas chamber samples to various on- and off-line analytical instruments. The schematic (Figure 2.1) shows how each port is used. Note that the dashed arrows indicate off-line analytical capabilities. In addition, because of the set up of our laboratory, it is possible to alter our system so that the GC-FID operates on-line rather than off-line (through the tenax packed tubes and the thermal desorber). Table 2.1 lists the manufacturers and model numbers of the equipment typically available for use with the indoor photochemical reactor.

An air purification system is used to deliver contaminant-free air to the chamber and the glass vacuum manifold. The system is made up of three stainless steel tubes. Tubes 1, 2, and 3 are filled with 50% molecular sieves/50% drierite, purafil, and activated carbon, respectively. Scrubbed air fed to the chamber typically contains less than approximately 1-2 ppb of ozone, NO₂, or hydrocarbons.

<table>
<thead>
<tr>
<th>Name</th>
<th>Manufacturer/Model Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO₂ monitor</td>
<td>Thermo Environmental / Model 42</td>
</tr>
<tr>
<td>Ozone Analyzer</td>
<td>Dasibi / Model 1003PC</td>
</tr>
<tr>
<td>Gas Chromatograph (ECD)</td>
<td>Shimadzu / Mini GC II</td>
</tr>
<tr>
<td>Gas Chromatograph (FID)</td>
<td>Hewlett Packard / Model 5890 Series II</td>
</tr>
<tr>
<td>Liquid Chromatograph (DAD)</td>
<td>Hewlett Packard / Model 1090</td>
</tr>
<tr>
<td>Gas Chromatograph (MS)</td>
<td>Hewlett Packard / Models 5890 and 5898</td>
</tr>
</tbody>
</table>
Figure 2.6: Schematic of the glass vacuum manifold which is used to admit reactants to the chamber.
2.2 Experimental System Characterization

2.2.1 Ultraviolet Light Intensity

One of the most important considerations in an indoor system lies in the determination of the ultraviolet intensity. This determination is crucial in ensuring that during the computer modeling of the experiments, all loss processes of photolyzing species are accurately accounted for. The ultraviolet intensity within the chamber is related to the photolysis rate constant through the following mathematical relationship:

\[ k(T) = \int \sigma(\lambda, T) \phi(\lambda, T) I(\lambda) d\lambda, \]  

(2.1)

where:  
\( \sigma(\lambda, T) \) = absorption coefficient  
\( \phi(\lambda, T) \) = quantum yield  
\( I(\lambda) \) = lamp intensity.

A calculation of the photolysis rate constants for various species can be made given the absorption coefficients and the quantum yields, but an experimental determination of the value is needed in order take into account the age and decreasing intensity of the lamps. The most widely accepted technique of determining the ultraviolet intensity of black lamps is a chemistry-based technique which relies on the photolysis of NO\(_2\) [3]. As mentioned earlier, NO\(_2\) is a critical component in the formation of ozone in the troposphere. In addition, nitrogen dioxide absorbs most of its energy in the wavelength region
were a large portion of tropospheric chemistry takes place (see Figure 2.4, B) and where the black lamp output is a maximum. Although other atmospheric species such as formaldehyde and ozone absorb in wavelength regions of crucial importance to tropospheric chemistry, their absorption coefficients are an order of magnitude lower than that of NO₂ in the same region.

By investigating the reaction of NO₂ in pure nitrogen diluent, a measure of the ultraviolet lamp intensity can be determined. The pertinent reactions in such a system are:

\begin{align*}
(1) & \quad NO₂ + hν \rightarrow O + NO \\
(2) & \quad O + O₂ + M \rightarrow O₃ + M \\
(3) & \quad O₃ + NO \rightarrow O₂ + NO₂ \\
(4) & \quad O + NO₂ \rightarrow O₂ + NO \\
(5) & \quad O + NO₂ + M \rightarrow NO₃ + M \\
(6) & \quad NO₃ + NO \rightarrow NO₂ + NO₂ \\
(7) & \quad O + NO + M \rightarrow NO₂ + M
\end{align*}

If O₂ is not initially present in the system, one can assume that [O], [NO₃], and [O₃] are at steady state. In addition, one can state that [NO] = [NO]₀ + [NO₂]₀ - [NO₂]. An expression for \( k_1 \), a measure of the light intensity, can then be obtained as [4]:

\[
k₁ = \frac{1}{2Δt} \left\{ [1 + R₁ + R₂] \ln \frac{[NO₂]₀}{[NO₂]} + R₂ \left( \frac{[NO₂]₀}{[NO₂]} - 1 \right) \right\}
\]

where

\[
Δt = t_{final} - t_{initial}
\]
assuming that [NO] and [O₂] are initially zero. (Note that if oxygen is initially present in the system, the above equation no longer holds since the steady-state assumptions do not necessarily apply. It has been estimated that concentrations above 20 ppm of oxygen will affect the determination of \( k_1 \) for the NO₂ system [4]). By measuring the concentration of NO₂ using a calibrated NOₓ monitor, and knowing the total diluent concentration, \( k_1 \) can be determined. A complete derivation of \( k_1 \) appears in Appendix 1.

Although the use of NO₂ as a determination of the uv light intensity is a widely accepted practice, one should also note that it does not fully characterize the photolysis rates of other compounds in the system. Various other compounds which can potentially form and photolyze in the reaction chamber have absorption spectra and wavelength dependencies which are quite different from that of NO₂. A measure of \( k_1 \) is thus not necessarily an accurate measure of the photolysis rates of these other compounds. However, the Carter modeling program[2] used in this thesis incorporates the absorption spectra and quantum yields of these compounds in conjunction with a normalization factor (determined by ratioing the experimentally and mathematically determined values of \( k_1 \)) to determine the photolysis rate constants of the additional compounds. Rate constant derived from the Carter program for several compounds are presented in Table 2.2.
Table 2.2: Rate constants generated from the Carter program after normalization.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Rate Constant (min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO₂</td>
<td>4x10⁻²</td>
</tr>
<tr>
<td>HONO</td>
<td>1.119x10⁻²</td>
</tr>
<tr>
<td>HCHO</td>
<td>7.039x10⁻⁵</td>
</tr>
<tr>
<td>H₂O₂</td>
<td>2.994x10⁻⁵</td>
</tr>
<tr>
<td>O₃(O³P)</td>
<td>1.942x10⁻⁴</td>
</tr>
<tr>
<td>O₃(O¹D)</td>
<td>4.020x10⁻⁵</td>
</tr>
</tbody>
</table>

### 2.2.2 Methyl Nitrite Chemistry

Methyl nitrite, CH₃ONO, is commonly used as a source of OH radicals in chamber studies [3]. A spectrum of the absorbance of methyl nitrite (as compared to another important nitrite, t-butyl nitrite), taken from Pitts and Pitts, 1986, appears in Figure 2.7. (The quantum yield is taken as 1, so this figure is directly comparable to the spectra of Figure 2.4.) At wavelengths > 320 nm, methyl nitrite photolyzes according to the following:

\[
\text{CH}_3\text{ONO} + h\nu \rightarrow \text{CH}_3\text{O}^- + \text{NO}
\]

\[
\text{CH}_3\text{O}^- + \text{O}_2 \rightarrow \text{HCHO} + \text{HO}_2
\]

\[
\text{HO}_2 + \text{NO} \rightarrow \text{OH}^- + \text{NO}_2
\]

In order to adequately model the experimental studies presented in this thesis, it is important to characterize the photolysis rate constant for methyl nitrite (\(j_{\text{CH}_3\text{ONO}}\)). Thus, experiments were conducted by photolyzing CH₃ONO-air and CH₃ONO-NO-air systems and monitoring concentrations using a NOₓ monitor. The photolysis rate constant for methyl nitrite was determined by fits of experimental data for NO to theoretical time-concentration profiles.
Figure 2.7: Absorbance spectra at 300 K of methyl nitrite (1) and t-butyl nitrite (2), taken from Pitts and Pitts, 1986.
Table 2.3: Simplified CH$_3$ONO mechanism used in modeling.

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction</th>
<th>Rate Constant (@ 298 K) cm$^3$molecule$^{-1}$s$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MN1</td>
<td>CH$_3$ONO $+$ hv $\rightarrow$ CH$_3$O $+$ HCHO</td>
<td>Experiment</td>
</tr>
<tr>
<td>MN2</td>
<td>CH$_3$O $+$ O$_2$ $\rightarrow$ HCHO $+$ HO$_2$</td>
<td>$1.9 \times 10^{-15}$</td>
</tr>
<tr>
<td>MN3</td>
<td>HO$_2$ $+$ NO $\rightarrow$ OH $+$ NO$_2$</td>
<td>$3.7 \times 10^{-12}$</td>
</tr>
<tr>
<td>F1</td>
<td>HCHO $+$ hv $= 2$ HO$_2$ $+$ CO</td>
<td>PHOT $=$ HCHO</td>
</tr>
<tr>
<td>F2</td>
<td>HCHO $+$ OH $\rightarrow$ HO$_2$ $+$ CO $+$ H$_2$O</td>
<td>$1.125 \times 10^{-12}, -1.288, 2.0$</td>
</tr>
</tbody>
</table>

Methyl Nitrite Photolysis

**Formaldehyde Reactions**

1) NO$_2$ $+$ hv $\rightarrow$ NO $+$ O
2) O $+$ O$_2$ $+$ M $\rightarrow$ O$_3$ $+$ M
3) O $+$ NO$_2$ $\rightarrow$ NO $+$ O$_2$
4) O $+$ NO$_2$ $+$ M $\rightarrow$ NO$_3$ $+$ M

Inorganic Reactions

1) NO$_2$ $+$ hv $\rightarrow$ NO $+$ O
2) O $+$ O$_2$ $+$ M $\rightarrow$ O$_3$ $+$ M
3) O $+$ NO$_2$ $\rightarrow$ NO $+$ O$_2$
4) O $+$ NO$_2$ $+$ M $\rightarrow$ NO$_3$ $+$ M

Please note that the reactions are simplified and only the most important reactions are included. The photolysis rate constant for methyl nitrite was determined to be $0.06$ min$^{-1}$ for 24 lamps. Several additional modeling studies were conducted with the methyl nitrite mechanism to determine the importance of the initial NO concentration and the reactions of HCHO in the CH$_3$ONO experimental system. In all cases, the starting methyl nitrite concentration was 1.1 ppm.

Three different simulations were conducted:

- HCHO reactions not included in the mechanism, [NO]$_o > 0$. 

The generated using the Carter modeling program[2]. Table 2.3 is a simplified version of the mechanism used to model the CH$_3$ONO chemistry, including only the most important reactions. Figure 2.8 shows the agreement between the experimental and theoretical data for NO in one of the CH$_3$ONO systems. The photolysis rate constant for methyl nitrite was determined to be $0.06$ min$^{-1}$ for 24 lamps. Several additional modeling studies were conducted with the methyl nitrite mechanism to determine the importance of the initial NO concentration and the reactions of HCHO in the CH$_3$ONO experimental system. In all cases, the starting methyl nitrite concentration was 1.1 ppm. Three different simulations were conducted:

- HCHO reactions not included in the mechanism, [NO]$_o > 0$. 


Figure 2.8: Experimental and theoretical data for NO for the photolysis of CH₃ONO.
- HCHO reactions included in the mechanism, $[\text{NO}]_0 > 0$.

- HCHO reactions included in the mechanism, $[\text{NO}]_0 = 0$.

The time concentration profiles for these three conditions appear in Figures 2.9-2.11. In addition, Figure 2.12 shows the effect of NO on ozone concentrations. Comparison of Figures 2.9 and 2.10 indicates that the inclusion of HCHO reactions in a methyl nitrite mechanism has a significant effect on the NO and NO$_2$ concentrations predicted in the system. The NO concentration is decreased and the NO$_2$ concentration is increased by the reaction of HCHO. Comparison of Figures 2.10 and 2.11 indicates that the presence of an initial amount of NO increases the overall concentration of NO$_2$ in the system. However, the HCHO concentration remains essentially the same. This result may indicate that despite the presence of NO, which would enhance the conversion of HO$_2$ to OH and produce NO$_2$, the resulting OH preferentially reacts with compounds other than HCHO. Thus, the OH+HCHO is not a dominant reaction in the CH$_3$ONO system. Figure 2.12 indicates that O$_3$ concentrations are not significant in any of the three systems. The effect of NO is initially opposite to that which we might expect from examination of the mechanism alone, since the presence of NO should decrease the amount of O$_3$ in the system. After a while, though, there is a crossover in curves (b) and (c), indicating that NO is effectively scavenging the O$_3$ in the system. Thus, the addition of NO as a reactant to a methyl nitrite system does indeed result in the lowering of O$_3$ concentrations once a critical level of O$_3$ is reached.
Figure 2.9: Simulation of CH$_3$ONO system. HCHO reactions are excluded from the mechanism, and [NO]$_0$ > 0.
Figure 2.10: Simulation of CH$_3$ONO system. HCHO reactions are included in the mechanism, and [NO]$_o$ > 0.
Figure 2.11: Simulation of CH$_3$ONO system. HCHO reactions are included in the mechanism, and [NO]$_o = 0$. 

Methyl Nitrite Simulation
(HCHO rxns, [NO]$_o$=0)
Figure 2.12: Comparison of O₃ data from simulations of CH₃ONO chemistry (a) excluding HCHO chemistry with [NO]₀ > 0, (b) including HCHO chemistry with [NO]₀ > 0, and (c) including HCHO chemistry with [NO]₀ = 0.
2.2.3 t-Butyl Alcohol Experiments

To determine whether the new experimental system behaved similar to other indoor chambers, the photooxidation of t-butyl alcohol (tBA) was re-investigated. [5] Specifically, the goals of these experiments were to determine whether or not we could generate OH radicals by the photolysis of methyl nitrite, successfully react these radicals with a relatively well-studied compound, and reproduce the results of other research groups. In addition, these experiments served as a test of our analytical instruments.

The atmospheric chemistry of t-butyl alcohol has become quite important since this compound is being considered as a fuel additive in order to reduce the reactivity of vehicle emissions. [5, 8] The tBA-OH reaction mechanism proceeds via an abstraction mechanism:

1. \((\text{CH}_3)_3\text{COH} + \cdot \text{OH} \rightarrow \cdot \text{CH}_2\text{C(CH}_3)_2\text{OH}\)
2. \(\cdot \text{CH}_2\text{C(CH}_3)_2\text{OH} + \text{O}_2 + \text{M} \rightarrow \cdot \text{O}_2\text{CH}_2\text{C(CH}_3)_2\text{OH} + \text{M}\)
3. \(\cdot \text{O}_2\text{CH}_2\text{C(CH}_3)_2\text{OH} + \text{NO} \rightarrow \cdot \text{OCH}_2\text{C(CH}_3)_2\text{OH} + \text{NO}_2\)
4. \(\cdot \text{OCH}_2\text{C(CH}_3)_2\text{OH} \rightarrow \text{HCHO} + \cdot \text{C(CH}_3)_2\text{OH}\)
5. \(\cdot \text{C(CH}_3)_2\text{OH} + \text{O}_2 \rightarrow \text{HO}_2^\cdot + \text{CH}_3\text{C(O)CH}_3\)

The consumption of one molecule of tBA results in the formation of approximately one molecule of formaldehyde and one molecule of acetone.[5]

Four tBA-OH experiments were conducted in the indoor photochemical reactor, and had typical starting concentrations of 20-40 mtorr and 0.1-0.2 torr of tBA and methyl nitrite, respectively. The reaction was monitored using the GC-MS. Standards of tBA and acetone in methanol were generated
and used for calibration of the system. To account for the secondary loss of acetone via reaction with OH radicals, the following formula was used:

\[
F = \frac{k_1 - k_2}{k_1} \frac{1 - [\text{Hydrocarbon}]_e}{[\text{Hydrocarbon}]_o} - \frac{k_2}{k_1} \frac{[\text{Hydrocarbon}]_e}{[\text{Hydrocarbon}]_o}
\]  

(2.2)

The rate constants \(k_1\) and \(k_2\) correspond to the reactions of tBA and acetone with OH, respectively, and have values of \(1.12 \times 10^{-12}\) and \(2.26 \times 10^{-13}\). A plot of the corrected acetone concentration versus the loss of tBA appears in Figure 2.13. The plot is a straight line, with slope of \(0.910 \pm 0.051\), as expected from the reaction mechanism and previously published experimental results.[5] We were able to successfully react tBA with methyl nitrite, and, given that our results agreed with previously published data, found that there were no abnormalities within our system (such as unexplained losses within our chamber), or with our sampling methods.
Figure 2.13: Yield of acetone from the photooxidation of t-Butyl Alcohol.
Bibliography


Chapter 3

Photooxidation of 2,2,4-Trimethylpentane and 2,2,5-Trimethylhexane: A Product Study

Abstract

Hydroxyl radical-initiated photooxidation of 2,2,4-trimethylpentane and 2,2,5-trimethylhexane in the presence of NOx has been investigated. Primary products detected from the reaction of 2,2,4-trimethylpentane with OH radicals are 2-methyl-1-propanal, acetone, and formaldehyde, with molar yields of 0.190, 0.264, and 1.114, respectively. Acetaldehyde was also positively detected as a product.

Both the kinetics and mechanism of 2,2,5-trimethylhexane photooxidation were studied. The relative rate method, using 2,2,4-trimethylhexane as a reference, was used in order to make the first determination of the rate constant for OH radical reaction with 2,2,5-trimethylhexane. This rate constant was determined to be $(5.27 \pm 0.15) \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, in excellent agreement with the predicted value of $5.7 \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ as determined from Atkinson's structure reactivity relationship. Primary products detected from the reaction of 2,2,5-trimethylhexane with OH radicals were 3,3-dimethylbutyraldehyde, acetone, acetaldehyde, and formaldehyde, with molar yields of 0.208, 0.169, 0.014 and 1.111, respectively. 2,2-Dimethyl-1-propanal was also detected in the 2,2,5-trimethylhexane system. Based on the 2,2,4-trimethylpentane study, model predictions, and 3,3-dimethylbutyraldehyde + OH experiments, the formation of 2,2-dimethyl-1-propanal is, however, believed to be a secondary, rather than a primary product. We saw no evidence for the formation of 3-methyl-1-propanal or 2-methyl-1-propanal. The results obtained are discussed in terms of their implications for the atmospheric reactions of alkoxy radicals.
3.1 Introduction

Kinetics and mechanisms of the atmospheric reactions of a number of the smaller alkanes have been investigated[1]. Studies of larger (> C₈) alkanes, however, have focused primarily on rate constant determinations [1], and mechanisms have yet to be fully determined. We have undertaken a study of the photooxidation mechanism of two branched alkanes, 2,2,4-trimethylpentane (iso-octane) and 2,2,5-trimethylhexane. Both compounds were identified in the 1987 Southern California Air Quality Study (SCAQS)[2]. Of the total morning composition of non-methane organic compounds, 2,2,4-trimethylpentane and 2,2,5-trimethylhexane represented 1.414% and 0.170% of the total carbon, respectively. In addition, 2,2,4-trimethylpentane is listed as one of the 25 most abundant species in the fall episode of SCAQS, based on the fraction of non-methane organic carbon.

Hydroxyl radical reaction with alkanes proceeds via H-atom abstraction. The peroxy radicals formed after the initial H-atom abstraction are, in the presence of NO, rapidly converted to alkoxy radicals. The reaction pathways of the alkoxy radicals are crucial to the overall mechanisms for alkane reactions. These resulting alkoxy radicals can react with NO, NO₂, O₂, unimolecularly decompose, or isomerize [1, 3, 4, 5, 6, 7, 8]. While estimates of the reaction rates of the O₂ and decomposition pathways have been made [1, 4, 5, 8], alkoxy radical isomerization reactions, on the other hand, still have not been completely characterized. Alkoxy radicals with greater than
five carbons are believed to undergo 1,4- or 1,5- H atom shift isomerization reactions via 5- and 6-member rings, respectively [1, 3, 5, 7] to eventually form hydroxy carbonyls. Rate constant data exist for the 1,5-H atom isomerization reactions of alkoxy radicals, but have not been tested against a wide variety of alkanes [1].

We report here the investigation of the photooxidation of 2,2,4-trimethylpentane and 2,2,5-trimethylhexane. The first determination of the rate constant of the OH + 2,2,5-trimethylhexane reaction is presented and compared to the theoretical value derived from Atkinson’s structure reactivity relationship[16, 15]. Experimental studies of the photooxidation of 2,2,4-trimethylpentane and 2,2,5-trimethylhexane are presented, and product yields are compared to those predicted by photooxidation mechanisms.

3.2 Experimental

The Caltech Indoor Photochemical Reactor has been described in detail elsewhere¹. Briefly, the chamber is a 1 m³ reactor constructed of 2 mil FEP Teflon. Artificial light (two banks of 24 Sylvania F30T8BL black lamps) is used to initiate the chemistry within the chamber. The lamps are mounted on a cylindrical surface to provide for uniformity in irradiation. The light intensity within the chamber can be adjusted by varying the number of lamps that are used. These experiments were conducted using 50% of the total intensity available.

¹See Chapter 2.
Relative Rate Experiment

A relative rate experiment was conducted to determine the rate constant of hydroxyl radical reaction with 2,2,5-trimethylhexane. The relative rate method is a standard method for determining the rate constant of a compound given a known rate constant of a reference compound [12]. The reference used in this study was 2,2,4-trimethylpentane, with a known rate constant of $3.59 \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ [1]. Provided that the sole loss process for 2,2,4-trimethylpentane (Reference) and 2,2,5-trimethylhexane (Reactant) is reaction with OH radicals, the following relations hold:

$$\text{Reactant} + \cdot \text{OH} \rightarrow \text{Products}$$
$$\text{Reference} + \cdot \text{OH} \rightarrow \text{Products}$$

$$\ln \frac{[\text{Reactant}]_o}{[\text{Reactant}]_t} = \frac{k_1}{k_2} \ln \frac{[\text{Reference}]_o}{[\text{Reference}]_t}, \quad (3.1)$$

where $k_1$ and $k_2$ are the rate constants for reactions (1) and (2), respectively. If the logarithmic terms are plotted against each other, the slope is the ratio, $rac{k_1}{k_2}$. The rate constant, $k_1$ can then be placed on an absolute basis using the known rate constant, $k_2$.

Three experiments were conducted with initial mixing ratios of 2-5 ppm of 2,2,4-trimethylpentane and 2,2,5-trimethylhexane, respectively, 3-6 ppm of methyl nitrite, and 1-3 ppm of NO.
Product Determinations

Methyl nitrite was initially used as the OH radical precursor in these experiments. Formaldehyde cannot be determined exclusively as a product from alkane-CH$_3$ONO-NO experimental systems because HCHO is also produced in the photolysis of CH$_3$ONO. Thus, ethyl nitrite was used as the OH precursor in several experiments. Methyl and ethyl nitrite photolyze according to the following mechanism:

\[
\begin{align*}
RCH_2ONO + h\nu & \rightarrow RCH_2O^- + NO \\
RCH_2O^- + O_2 & \rightarrow RCHO + HO_2^- \\
HO_2^- + NO & \rightarrow OH^- + NO_2, \\
\end{align*}
\]

where R = H for methyl nitrite, or R = CH$_3$ for ethyl nitrite.

To minimize ozone production, and maximize the conversion of peroxy radicals to alkoxy radicals, NO was added to the chamber. Typical initial mixing ratios were approximately 2-4 ppm and 2-5 ppm for 2,2,4-trimethylpentane and 2,2,5-trimethylhexane, respectively, 3-10 ppm CH$_3$ONO (or C$_2$H$_5$ONO), and 2-5 ppm NO. 2,2,4-Trimethylpentane was obtained from Aldrich with a stated purity of 99+%, and 2,2,5-trimethylhexane was obtained from Wiley Organics with a stated purity of 99.8%. Nitric oxide was taken from a standard cylinder generated by Scott Marin. The nitrites were prepared by the dropwise addition of 50% sulfuric acid to a saturated alcohol (methanol for methyl nitrite and ethanol for ethyl nitrite) /sodium nitrite solution, and were vacuum purified and stored under liquid N$_2$[13].
The mixture was photolyzed and sampled periodically for analyses by an on-line GC-FID (HP 5890) and NO\textsubscript{x} analyzer (Thermo Environmental Model 42). The column used in the GC was a J&W Scientific DB-5 column of dimension 30m x 0.25mm x 0.25\textmu{}m. The temperature program used started at 35° C. The oven temperature was maintained at 35° C for 6 min., and then increased by 30° C/min. until 200° C. The final temperature of 200° C was maintained for 1 min. Retention times for 2,2,4-trimethylpentane and 2,2,5-trimethylhexane were 4.0 min. and 5.0 min., respectively. Typical irradiation times for the branched alkane study ranged from 5 to 20 min.

At the end of each experiment, a sample from the chamber was collected on a 2,4-dinitrophenylhydrazine (DNPH) impregnated C\textsubscript{18}-cartridge (Waters Co.). The samples were eluted using 2 ml of acetonitrile, and analyzed using a Hewlett Packard Model 1090 Series II liquid chromatograph, with a diode array detector. A reversed-phase Axxiom LC column of dimension 5 \textmu{}m x 150 cm was used to obtain adequate separation. HPLC grade acetonitrile and water were used as the solvents in a ratio of 55% to 45%, with a total flowrate of 1 ml/min. Typical observed pressures were between 109 and 112 bar. Authentic standards were compared to the experimental samples to identify and quantify the products. Formaldehyde-, acetaldehyde-, acetone-, and 3-methyl-1-butanal-DNPH hydrazone standards in acetonitrile were obtained from Radian corporation at a concentration of 3 \mu{}g/ml. Crystalline 2-methyl-1-propanal-DNPH hydrazone was obtained from Radian corporation, and was used to generate standard solutions using HPLC grade acetonitrile. Both the standard solutions and the crystalline hydrazones had stated purities of at least 99%. They were used without further purification.
2,2-Dimethyl-1-propanal and 3,3-dimethylbutyraldehyde DNPH hydrazones (crystalline forms) were synthesized in our laboratory according to the methods of Druzik et al. [14]. The hydrazones were combined with HPLC grade acetonitrile to form standard solutions of each. 2,2-Dimethyl-1-propanal and 3,3-dimethylbutyraldehyde were obtained from Aldrich, with stated purities of 97% and 95%, respectively. While the aldehydes themselves were not purified, the resulting hydrazones were recrystallized twice to remove any impurities. Based on LC-diode array analyses, these standard solutions had purities of at least 99%.

Interferences from Primary Carbonyl Product Reactions

Several experiments were performed to investigate the product yield interferences that may arise from carbonyl reaction with OH radicals or photolysis. 2,2-Dimethyl-1-propanal-, 3,3-dimethylbutyraldehyde-, 2-methyl-1-propanal-, and 3-methyl-1-butanal-CH$_3$ONO-NO mixtures were placed in the chamber and photolyzed. Samples were taken on DNPH impregnated cartridges for analysis by HPLC. The results were analyzed, and results indicate that the OH reactions with 3,3-dimethylbutyraldehyde and 3-methyl-1-butanal result in the formation of 2,2-dimethyl-1-propanal and 2-methyl-1-propanal, respectively. The OH reactions with 2,2-dimethyl-1-propanal and 2-methyl-1-propanal did not result in compounds that interfere with product analyses in the alkane studies. Separate experiments were conducted to investigate the photolysis lifetimes of aldehydes under the conditions used.
in these studies. Under typical time periods used in this study, significant losses in the aldehyde concentration were not detected.

## 3.3 Rate Constant of 2,2,5-Trimethylhexane-OH Reaction

Plotting the logarithmic terms of Equation 3.1 against each other results in a straight line with a slope of $1.467\pm0.041$ (see Figure 3.1). This slope corresponds to the ratio of rate constants, $\frac{k_2}{k_1}$, and the error corresponds to $2\sigma$. Using the known rate constant for 2,2,4-trimethylpentane of $3.59\times10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ [1], a rate constant for the OH-initiated photooxidation of 2,2,5-trimethylhexane of $(5.27\pm0.15)\times10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ is determined, and does not include errors due to the rate constant of the reference reaction. This determination of the rate constant for OH reaction with 2,2,5-trimethylhexane is in excellent agreement with the predicted value of $5.7\times10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ as determined from structure reactivity relationships [16, 15].

## 3.4 Photooxidation Products

The structure reactivity relationship [16, 15] predicts the following percentage distribution for OH attack sites on 2,2,4-trimethylpentane: tertiary carbon = 51%, secondary carbon = 29%, and primary carbon = 20%. Similarly, the structure reactivity relationship predicts the percentages of OH radical at-
Figure 3.1: Relative rate study to determine $k_{OH}$ for 2,2,5-trimethylhexane.
tack on the tertiary, secondary, and primary carbons of 2,2,5-trimethylhexane as 40%, 45%, and 15%, respectively. Proposed mechanisms for the tertiary, secondary, and primary OH reactions for each alkane, indicating the dominant pathways for decomposition, and excluding isomerization reactions, are shown in Figures 3.2-3.5 for 2,2,4-trimethylpentane and Figures 3.6-3.10 for 2,2,5-trimethylhexane. Additional decomposition pathways are included in the mechanisms which will be discussed subsequently.

3.4.1 2,2,4 Trimethylpentane

Products identified in the 2,2,4-trimethylpentane-OH-NO\textsubscript{x} system include formaldehyde, acetone, and 2-methyl-1-propanal. These aldehydes are themselves capable of photolyzing and reacting with OH radicals. The interference studies discussed earlier indicate the photolysis loss of these products is not significant over the photolysis time periods considered in these experiments. The yield of each of the products identified was corrected for losses due to reaction with OH radicals, according to the method of Atkinson et al. [15]. The correction factor, \( F \), was calculated using Equation 3.2 along with the rate constants in Table 3.1. Raw yields were multiplied by \( F \) to produce corrected yields,

\[
F = k_1 - k_2 \frac{1 - \frac{[\text{Hydrocarbon}]}{[\text{Hydrocarbon}]_0}}{k_1 \frac{[\text{Hydrocarbon}]}{[\text{Hydrocarbon}]_0}^{b_1} - \frac{[\text{Hydrocarbon}]}{[\text{Hydrocarbon}]_0^{b_2}}}
\]

where \( k_1 \) and \( k_2 \) are the rate constants of the following reactions:

\[
\text{Hydrocarbon} + \text{OH} \rightarrow \text{Product}
\]
Figure 3.2: 2,2,4-Trimethylpentane + OH Reactions, Tertiary
Figure 3.3: 2,2,4-Trimethylpentane + OH Reactions, Secondary
Figure 3.4: 2,2,4-Trimethylpentane + OH Reactions, Primary
Figure 3.5: 2,2,4-Trimethylpentane + OH Reactions, Primary
Figure 3.6: 2,2,5-Trimethylhexane + OH Reactions, Tertiary
Figure 3.7: 2,2,5-Trimethylhexane + OH Reactions, Secondary
Figure 3.8: 2,2,5-Trimethylhexane + OH Reactions, Secondary
Figure 3.9: 2,2,5-Trimethylhexane + OH Reactions, Primary
Figure 3.10: 2,2,5-Trimethylhexane + OH Reactions, Primary
Table 3.1: OH Radical rate constants used to correct product yields.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Rate constant ( \text{cm}^3\text{molecule}^{-1}\text{s}^{-1} )</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Alkanes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2,2,4-Trimethylpentane</td>
<td>3.59 x 10^{-12}</td>
<td>[1]</td>
</tr>
<tr>
<td>2,2,5-Trimethylhexane</td>
<td>5.267 x 10^{-12}</td>
<td>This work.</td>
</tr>
<tr>
<td><strong>Products</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>9.37 x 10^{-12}</td>
<td>[1]</td>
</tr>
<tr>
<td>Acetaldehyde</td>
<td>1.58 x 10^{-12}</td>
<td>[8, 1]</td>
</tr>
<tr>
<td>Acetone</td>
<td>2.19 x 10^{-13}</td>
<td>[1]</td>
</tr>
<tr>
<td>2-Methyl-1-propanal</td>
<td>2.63 x 10^{-12}</td>
<td>[8, 1]</td>
</tr>
<tr>
<td>3-Methyl-1-butanal</td>
<td>2.74 x 10^{-11}</td>
<td>[8]</td>
</tr>
<tr>
<td>2,2-Dimethyl-1-propanal</td>
<td>2.65 x 10^{-11}</td>
<td>[8, 1]</td>
</tr>
</tbody>
</table>

Product + OH $\rightarrow$ Secondary Products

The primary radical produced in the photooxidation of 2,2,4-trimethylpentane is the neopentyl peroxy radical. In the presence of significant NO, this radical will react with NO to form the corresponding alkoxy radical. The neopentyl alkoxy radical can react with O$_2$ to form 2,2-dimethyl-1-propanal or decompose to eventually form acetone and formaldehyde (see Figure 3.2). With the use of authentic samples of 2,2-dimethyl-1-propanal-DNPH, we were able to confirm that there was no evidence for the formation of 2,2-dimethyl-1-propanal in the 2,2,4-trimethylpentane photooxidation system. This finding is consistent with that of Wallington et al. [20] who saw no significant evidence for the formation of 2,2-dimethyl-1-propanal during their studies of the neopentyl radical in the presence of NO$_x$. They did, however, notice significant formation of this product in systems without NO$_x$. Our results
Table 3.2: Corrected Carbonyl Photooxidation Product Yields

<table>
<thead>
<tr>
<th>Alkane</th>
<th>Product</th>
<th>Molar Yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>2,2,4-Trimethylpentane</td>
<td>2-Methyl-1-propanal</td>
<td>0.190</td>
</tr>
<tr>
<td></td>
<td>Acetone</td>
<td>0.264</td>
</tr>
<tr>
<td></td>
<td>Formaldehyde</td>
<td>1.114</td>
</tr>
<tr>
<td></td>
<td>Acetaldehyde</td>
<td>Detected</td>
</tr>
<tr>
<td>2,2,5-Trimethylhexane</td>
<td>3,3-Dimethylbutyraldehyde</td>
<td>0.208</td>
</tr>
<tr>
<td></td>
<td>2,2-Dimethyl-1-propanal</td>
<td>See text.</td>
</tr>
<tr>
<td></td>
<td>Acetone</td>
<td>0.169</td>
</tr>
<tr>
<td></td>
<td>Acetaldehyde</td>
<td>0.014</td>
</tr>
<tr>
<td></td>
<td>Formaldehyde</td>
<td>1.111</td>
</tr>
</tbody>
</table>

indicate that the major pathway for the neopentyl alkoxy radical is decomposition rather than reaction with $O_2$. Thus, the major products obtained from the tertiary carbon reactions of 2,2,4-trimethylpentane are acetone and formaldehyde.

The major product from OH attack on the secondary carbon of 2,2,4-trimethylpentane is 2-methyl-1-propanal, whereas formaldehyde and acetone are major products from the primary carbon reactions. Acetaldehyde was also detected as a secondary product in a few of the experiments. Without standards, the larger aldehydes and ketones believed to be formed from the primary reactions could not be positively identified. However, several carbonyl compounds with relatively long retention times (consistent with large compounds) were detected by the LC. Yields of each of the quantified products are given in Table 3.2.
3.4.2 2,2,5 Trimethylhexane

The major products identified from 2,2,5-trimethylhexane photooxidation include formaldehyde, acetone, acetaldehyde, 2,2-dimethyl-1-propanal, and 3,3-dimethylbutyraldehyde. Yields for each of the detected carbonyls were corrected using Equation 3.2 and the rate constants in Table 3.1. The corrected yields are presented in Table 3.2.

The most abundant radicals formed in the 2,2,5-trimethylhexane system are those resulting from OH attack on the tertiary carbon of 2,2,5-trimethylhexane. Based on the mechanism presented in Figure 3.6, the possible products formed from attack on the tertiary carbon include 3,3-dimethylbutyraldehyde, 2,2-dimethyl-1-propanal, formaldehyde, and acetaldehyde. Each of these products was detected in the photooxidation studies of 2,2,5-trimethylhexane. However, the 2,2-dimethyl-1-propanal formed is believed to be a secondary, rather than a dominant, product. The only pathway for formation of the 2,2-dimethyl-1-propanal is via the neopentyl alkoxy radical. The 2,2,4-trimethylpentane study presented in this paper and the neopentyl peroxy study of Wallington et al.[20] indicate that neopentyl radicals are not a precursor to 2,2-dimethyl-1-propanal in the presence of significant NOx. In addition, the interference study reported in this work indicates that 2,2-dimethyl-1-propanal is formed as a product of OH reaction with 3,3-dimethylbutyraldehyde. Thus, the 2,2-dimethyl-1-propanal identified is a result of secondary reactions of 3,3-dimethylbutyraldehyde, and has been included in the calculation of the total 3,3-dimethylbutyraldehyde concentration. It is important to note that the $F$ factor was not used to correct
the 3,3-dimethylbutyraldehyde yield, since the addition of the corrected 2,2-dimethyl-1-propanal yield to the 3,3-dimethylbutyraldehyde yield accounts for the losses due to OH reaction. More detail will be given in the next section regarding the direct formation of 2,2-dimethyl-1-propanal.

Based on Figure 3.7, OH attack on the secondary carbon of 2,2,5-trimethylhexane could result in the formation of 3-methyl-1-butanal, formaldehyde, and acetone from the alkoxy radical decomposition route, or the ketone, (CH₃)₃CC(O)-CH₂C(CH₃)₂, from the alkoxy radical reaction with O₂. Authentic DNPH derivatized samples were not available to investigate the formation of the C₉ ketone. However, with the use of authentic DNPH derivatized samples of 3-methyl-1-butanal, we were able to confirm that, under the conditions of the study presented in this paper, there was no evidence for the formation of 3-methyl-1-butanal from the photooxidation of 2,2,5-trimethylhexane.

Attack on the primary carbon centers of 2,2,5-trimethylhexane can result in the formation of a variety of carbonyl products. From Figure 3.9, possible photooxidation products include 2-methyl-1-propanal, acetaldehyde, formaldehyde, acetone, and 3-methyl-1-butanal from the decomposition pathway of the initial C₉ alkoxy radical formed, or the C₉ carbonyl from reaction of this same alkoxy radical with O₂. Authentic DNPH derivatized samples of the C₉ carbonyl depicted in Figure 3.9 were not available. However, authentic DNPH derivatized samples were available for the other possible carbonyl products. There was no evidence for the formation of 2-methyl-1-butanal or 3-methyl-1-butanal in OH-initiated photooxidation of 2,2,5-trimethylhexane in the presence of NO₂.
3.5 Atmospheric Reaction Mechanisms

2,2,4-Trimethylpentane and 2,2,5-trimethylhexane photooxidation mechanisms were developed using the SAPRC Atmospheric Photochemical Mechanism Preparation Program[16]. To simplify the mechanisms slightly, we included all species leading to nitrite (with the exception of CH$_3$ONO) or nitrates in two general categories entitled "Nitrites" and "Nitrates", respectively, and did not include the secondary reactions of the major products formed in the two studies (see Table 3.2). Since all of the major products were corrected for losses using Equation 3.2, the exclusion of secondary loss processes in the mechanisms facilitated the direct comparison between observed and predicted yields of major products. The mechanisms used for 2,2,4-trimethylpentane and 2,2,5-trimethylhexane appear in Tables 3.3 and 3.4, respectively. Note that while the inorganic reactions (those involving NO$_x$) were included in the actual mechanisms used for simulating the experiments, to conserve space, they are not included in Tables 3.3 and 3.4. Rate constants used for all reactions except those involving the alkoxy radicals were obtained directly from Atkinson [1]. The rate constants for the alkoxy radical reactions were derived using $\Delta$H estimates obtained from Stein et al. [25] in conjunction with the estimation technique of Atkinson [1]. Units for all of the rate constants with the exception of those corresponding to the isomerization and decomposition reactions are cm$^3$ molecule$^{-1}$ s$^{-1}$. The isomerization and decomposition reaction rate constants are in units of s$^{-1}$. All rate constants are at 298 K.
Table 3.3: 2,2,4-Trimethylpentane Mechanism
(Units: cm³, molecule, s)

<table>
<thead>
<tr>
<th>No.</th>
<th>k @ 298K</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><strong>2,2,4-Trimethylpentane Reactions</strong></td>
</tr>
<tr>
<td>1A)</td>
<td>1.83x10⁻¹²</td>
<td>TMP + OH → TMPR1 + H₂O</td>
</tr>
<tr>
<td>1B)</td>
<td>1.04x10⁻¹²</td>
<td>TMP + OH → TMPR2 + H₂O</td>
</tr>
<tr>
<td>1C)</td>
<td>4.314x10⁻¹³</td>
<td>TMP + OH → TMPR3 + H₂O</td>
</tr>
<tr>
<td>1D)</td>
<td>2.87x10⁻¹³</td>
<td>TMP + OH → TMP3A + H₂O</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Tertiary Radical Reactions</strong></td>
</tr>
<tr>
<td>2)</td>
<td>1x10⁻¹²</td>
<td>TMPR1 + O₂ → TMPR1-O₂</td>
</tr>
<tr>
<td>3)</td>
<td>2.0x10⁻¹⁷</td>
<td>TMPR1-O₂ + TMPR1-O₂ → TMPR1-O + TMPR1-O + O₂</td>
</tr>
<tr>
<td>4)</td>
<td>7.65x10⁻¹²</td>
<td>TMPR1-O₂ + NO → TMPR1-O + NO₂</td>
</tr>
<tr>
<td>4A)</td>
<td>1.25x10⁻¹²</td>
<td>TMPR1-O₂ + NO → Nitrate</td>
</tr>
<tr>
<td>5)</td>
<td>6.7x10⁴</td>
<td>TMPR1-O → Isomerization</td>
</tr>
<tr>
<td>6)</td>
<td>3.8x10⁻¹¹</td>
<td>TMPR1-O + NO → Nitrite</td>
</tr>
<tr>
<td>7)</td>
<td>3.8x10⁻¹¹</td>
<td>TMPR1-O + NO₂ → Nitrate</td>
</tr>
<tr>
<td>8)</td>
<td>3x10⁴</td>
<td>TMPR1-O → ACETONE + NPT</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Secondary Radical Reactions</strong></td>
</tr>
<tr>
<td>9)</td>
<td>1x10⁻¹²</td>
<td>TMPR2 + O₂ → TMPR2-O₂</td>
</tr>
<tr>
<td>10A)</td>
<td>2.25x10⁻¹⁵</td>
<td>TMPR2-O₂ + TMPR2-O₂ → TMPR2-O + TMPR2-O + O₂</td>
</tr>
<tr>
<td>10B)</td>
<td>2.75x10⁻¹⁵</td>
<td>TMPR2-O₂ + TMPR2-O₂ → ALCOHOL1 + P₃O</td>
</tr>
<tr>
<td>11)</td>
<td>5.79⁻¹²</td>
<td>TMPR2-O₂ + NO → TMPR2-O + NO₂</td>
</tr>
<tr>
<td>11A)</td>
<td>3.11x10⁻¹²</td>
<td>TMPR2-O₂ + NO → Nitrate</td>
</tr>
<tr>
<td>12)</td>
<td>3.8x10⁻¹¹</td>
<td>TMPR2-O + NO → Nitrite</td>
</tr>
<tr>
<td>13)</td>
<td>3.8x10⁻¹¹</td>
<td>TMPR2-O + NO₂ → Nitrate</td>
</tr>
</tbody>
</table>
14) \[ 2.48 \times 10^{-14} \text{ TMPR}_2\text{-O} + \text{O}_2 \rightarrow \text{P}_3\text{O} + \text{HO}_2 \]
15) \[ 5.66 \times 10^5 \text{ TMPR}_2\text{-O} \rightarrow (\text{CH}_3)_3\text{C} + \text{MPROP} \]
15A) Varied \[ \text{TMPR}_2\text{-O} \rightarrow 1,4\text{-Isomerization} \]

Primary Radical Reactions

16) \[ 1 \times 10^{-12} \text{ TMPR}_3 + \text{O}_2 \rightarrow \text{TMPR}_3\text{-O}_2 \]
17A) \[ 1.125 \times 10^{-13} \text{ TMPR}_3\text{-O}_2 + \text{TMPR}_3\text{-O}_2 \rightarrow \text{TMPR}_3\text{-O} + \text{TMPR}_3\text{-O} + \text{O}_2 \]
17B) \[ 1.375 \times 10^{-13} \text{ TMPR}_3\text{-O}_2 + \text{TMPR}_3\text{-O}_2 \rightarrow \text{ALCOHOL}_2 + \text{ALDEHYDE}_2 \]
18) \[ 7.3 \times 10^{-12} \text{ TMPR}_3\text{-O}_2 + \text{NO} \rightarrow \text{TMPR}_3\text{-O} + \text{NO}_2 \]
18A) \[ 1.60 \times 10^{-12} \text{ TMPR}_3\text{-O}_2 + \text{NO} \rightarrow \text{Nitrate} \]
19) \[ 3.8 \times 10^{-11} \text{ TMPR}_3\text{-O} + \text{NO} \rightarrow \text{Nitrite} \]
20) \[ 3.8 \times 10^{-11} \text{ TMPR}_3\text{-O} + \text{NO}_2 \rightarrow \text{Nitrate} \]
21) \[ 8.82 \times 10^{-15} \text{ TMPR}_3\text{-O}_2 + \text{O}_2 \rightarrow \text{HO}_2 + \text{ALDEHYDE}_2 \]
22) \[ 6.3 \times 10^4 \text{ TMPR}_3\text{-O} \rightarrow \text{HCHO} + \text{R}_3\text{I} \]
23) \[ 4.1 \times 10^9 \text{ TMPR}_3\text{-O} \rightarrow 1,5\text{-Isomerization} \]
24) \[ 1 \times 10^{-12} \text{ R}_3\text{I} + \text{O}_2 \rightarrow \text{R}_3\text{I}\text{-O}_2 \]
25) \[ 2.0 \times 10^{-17} \text{ R}_3\text{I}\text{-O}_2 + \text{R}_3\text{I}\text{-O}_2 \rightarrow \text{R}_3\text{I}\text{-O} + \text{R}_3\text{I}\text{-O} + \text{O}_2 \]
26) \[ 7.97 \times 10^{-12} \text{ R}_3\text{I}\text{-O}_2 + \text{NO} \rightarrow \text{R}_3\text{I}\text{-O} + \text{NO}_2 \]
26A) \[ 9.34 \times 10^{-13} \text{ R}_3\text{I}\text{-O}_2 + \text{NO} \rightarrow \text{Nitrate} \]
27) \[ 3.8 \times 10^{-11} \text{ R}_3\text{I}\text{-O} + \text{NO} \rightarrow \text{Nitrite} \]
28) \[ 3.8 \times 10^{-11} \text{ R}_3\text{I}\text{-O} + \text{NO}_2 \rightarrow \text{Nitrate} \]
29) \[ 4.9 \times 10^4 \text{ R}_3\text{I}\text{-O} \rightarrow \text{ACETONE} + \text{MP} \]
30) \[ 2.9 \times 10^{-12} \text{ MP} + \text{O}_2 \rightarrow \text{MP}\text{-O}_2 \]
31) \[ 1.125 \times 10^{-13} \text{ MP}\text{-O}_2 + \text{MP}\text{-O}_2 \rightarrow \text{MP}\text{-O} + \text{MP}\text{-O} + \text{O}_2 \]
31A) \[ 1.375 \times 10^{-13} \text{ MP}\text{-O}_2 + \text{MP}\text{-O}_2 \rightarrow \text{ALCOHOL}_3 + \text{MPROP} + \text{O}_2 \]
32) \[ 8.9 \times 10^{-12} \text{ MP}\text{-O}_2 + \text{NO} \rightarrow \text{MP}\text{-O} + \text{NO}_2 \]
33) \[ 3.8 \times 10^{-11} \text{ MP}\text{-O} + \text{NO} \rightarrow \text{Nitrite} \]
34) \[ 3.8 \times 10^{-11} \text{ MP}\text{-O} + \text{NO}_2 \rightarrow \text{Nitrate} \]
35) \[ 1.1 \times 10^3 \text{ MP}\text{-O} \rightarrow \text{HCHO} + \text{P} \]
36) \[ 8.5 \times 10^{-15} \text{ MP}\text{-O} + \text{O}_2 \rightarrow \text{MPROP} + \text{HO}_2 \]
<table>
<thead>
<tr>
<th>Reaction Number</th>
<th>Rate Coefficient</th>
<th>Reaction Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>37</td>
<td>$1.1 \times 10^{-12}$</td>
<td>$P + O_2 \rightarrow P-O_2$</td>
</tr>
<tr>
<td>38</td>
<td>$5.5 \times 10^{-16}$</td>
<td>$P-O_2 + P-O_2 \rightarrow P-O + P-O + O_2$</td>
</tr>
<tr>
<td>38A</td>
<td>$5.4 \times 10^{-16}$</td>
<td>$P-O_2 + P-O_2 \rightarrow \text{ALCOHOL}4 + \text{ACETONE}$</td>
</tr>
<tr>
<td>39</td>
<td>$8.9 \times 10^{-12}$</td>
<td>$P-O_2 + NO \rightarrow P-O + NO_2$</td>
</tr>
<tr>
<td>40</td>
<td>$3.4 \times 10^{-11}$</td>
<td>$P-O + NO \rightarrow \text{Nitrite}$</td>
</tr>
<tr>
<td>41</td>
<td>$3.5 \times 10^{-11}$</td>
<td>$P-O + NO_2 \rightarrow \text{Nitrate}$</td>
</tr>
<tr>
<td>42</td>
<td>$8.0 \times 10^{-15}$</td>
<td>$P-O + O_2 \rightarrow \text{ACETONE} + HO_2$</td>
</tr>
<tr>
<td>43</td>
<td>$1.5 \times 10^{-1}$</td>
<td>$P-O \rightarrow CH3 + CH3CHO$</td>
</tr>
<tr>
<td><strong>Primary Radical Reactions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>44</td>
<td>$1 \times 10^{-12}$</td>
<td>$\text{TMP3A} + O_2 \rightarrow \text{TMP3A-}O_2$</td>
</tr>
<tr>
<td>45</td>
<td>$1.125 \times 10^{-13}$</td>
<td>$\text{TMP3A-}O_2 + \text{TMP3A-}O_2 \rightarrow \text{TMP3A-O} + \text{TMP3A-O} + O_2$</td>
</tr>
<tr>
<td>45A</td>
<td>$1.375 \times 10^{-13}$</td>
<td>$\text{TMP3A-}O_2 + \text{TMP3A-}O_2 \rightarrow \text{ALCOHOL5} + \text{ALDEHYDE3}$</td>
</tr>
<tr>
<td>46</td>
<td>$7.30 \times 10^{-12}$</td>
<td>$\text{TMP3A-}O_2 + NO \rightarrow \text{TMP3A-O} + NO_2$</td>
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<tr>
<td>46A</td>
<td>$1.60 \times 10^{-12}$</td>
<td>$\text{TMP3A-}O_2 + NO \rightarrow \text{Nitrate}$</td>
</tr>
<tr>
<td>47</td>
<td>$3.8 \times 10^{-11}$</td>
<td>$\text{TMP3A-O} + NO \rightarrow \text{Nitrite}$</td>
</tr>
<tr>
<td>48</td>
<td>$3.8 \times 10^{-11}$</td>
<td>$\text{TMP3A-O} + NO_2 \rightarrow \text{Nitrate}$</td>
</tr>
<tr>
<td>49</td>
<td>$1.1 \times 10^{-14}$</td>
<td>$\text{TMP3A-O} + O_2 \rightarrow HO_2 + \text{ALDEHYDE3}$</td>
</tr>
<tr>
<td>50</td>
<td>$1.6 \times 10^{4}$</td>
<td>$\text{TMP3A-O} \rightarrow HCHO + R32$</td>
</tr>
<tr>
<td>51</td>
<td>$1 \times 10^{-12}$</td>
<td>$R32 + O_2 \rightarrow R32-O_2$</td>
</tr>
<tr>
<td>52</td>
<td>$2.0 \times 10^{-17}$</td>
<td>$R32-O_2 + R32-O_2 \rightarrow R32-O + R32-O + O_2$</td>
</tr>
<tr>
<td>53</td>
<td>$6.39 \times 10^{-12}$</td>
<td>$R32-O_2 + NO \rightarrow R32-O + NO_2$</td>
</tr>
<tr>
<td>53A</td>
<td>$2.51 \times 10^{-12}$</td>
<td>$R32-O_2 + NO \rightarrow \text{Nitrate}$</td>
</tr>
<tr>
<td>54</td>
<td>$3.8 \times 10^{-11}$</td>
<td>$R32-O + NO \rightarrow \text{Nitrite}$</td>
</tr>
<tr>
<td>55</td>
<td>$3.8 \times 10^{-11}$</td>
<td>$R32-O + NO_2 \rightarrow \text{Nitrate}$</td>
</tr>
<tr>
<td>56</td>
<td>$4.5 \times 10^{3}$</td>
<td>$R32-O \rightarrow CH3CHO + \text{NPT}$</td>
</tr>
<tr>
<td><strong>Neopentyl Radical Reactions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP1</td>
<td>$2.4 \times 10^{-12}$</td>
<td>$\text{NPT} + O_2 \rightarrow \text{NPTO2}$</td>
</tr>
<tr>
<td>NP2A</td>
<td>$6.0 \times 10^{-13}$</td>
<td>$\text{NPTO2} + \text{NPTO2} \rightarrow \text{DMP} + \text{DMPA} + O_2$</td>
</tr>
<tr>
<td>NP2B</td>
<td>$4.0 \times 10^{-13}$</td>
<td>$\text{NPTO2} + \text{NPTO2} \rightarrow \text{NPT} + \text{NPT} + O_2$</td>
</tr>
</tbody>
</table>
\begin{align*}
& \text{Primary Reactions} \\
& \text{Secondary Reactions} \\
& \text{Tertiary Hydrogen Abstraction Reactions} \\
& \text{Secondary Hydrogen Abstraction Reactions} \\
& \text{Tertiary Hydrogen Transfer Reactions} \\
& \text{Secondary Hydrogen Transfer Reactions}
\end{align*}

\text{Table 3.4: 2,2'-Dimethylthiophene Mechanism}
Primary Radical Reactions

1. \( \text{TMA} + \text{NO} + \text{O}_2 \) → 1.2 \( \text{TMA} + \text{O}_2 \) → 1.2

2. \( \text{TMA} + \text{CO} + \text{HCHO} + \text{ACETONE} \) → 2.8 \( \text{TMA} + \text{O} + \text{CO} \) → 2.8

3. \( \text{TMA} + \text{O} + \text{HCHO} + \text{ACETONE} \) → 3 \( \text{TMA} + \text{O} + \text{HCHO} + \text{ACETONE} \) → 3

Secondary Radical Reactions

4. \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 4.1 \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 4.1

5. \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 5 \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 5

6. \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 6 \( \text{TMA} + \text{O}_2 \) + \( \text{HCHO} + \text{ACETONE} \) → 6

Secondary Radical Reactions

7. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 2 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 2

8. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 3 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 3

9. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 4 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 4

10. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 5 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 5

11. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 6 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 6

12. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 7 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 7

13. \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 8 \( \text{CH}_3 + \text{NO} + \text{O}_2 \) → 8
31a) $1.7 \times 10^{-12}$ TMHIA-O$_2$ + NO $\rightarrow$ Nitrate
32) $8.82 \times 10^{-15}$ TMHIA-O + O$_2$ $\rightarrow$ HO$_2$ + C$_8$H$_{17}$CHO
33) $2.16 \times 10^4$ TMHIA-O $\rightarrow$ HCHO + C$_8$H$_{17}$
33A) $8.4 \times 10^6$ TMHIA-O $\rightarrow$ 1,5-Isomerization
34) $3.8 \times 10^{-11}$ TMHIA-O + NO $\rightarrow$ Nitrite
35) $3.8 \times 10^{-11}$ TMHIA-O + NO$_2$ $\rightarrow$ Nitrate
36) $1 \times 10^{-12}$ C$_8$H$_{17}$ + O$_2$ $\rightarrow$ C$_8$H$_{17}$-O$_2$
37) $1.87 \times 10^{-17}$ C$_8$H$_{17}$-O$_2$ + C$_8$H$_{17}$-O$_2$ $\rightarrow$ C$_8$H$_{17}$-O + C$_8$H$_{17}$-O + O$_2$
38) $7.75 \times 10^{-12}$ C$_8$H$_{17}$-O$_2$ + NO $\rightarrow$ C$_8$H$_{17}$-O + NO$_2$
38a) $1.15 \times 10^{-12}$ C$_8$H$_{17}$-O$_2$ + NO $\rightarrow$ Nitrate
39) $4.9 \times 10^4$ C$_8$H$_{17}$-O $\rightarrow$ ACETONE + C$_5$H$_{11}$
39) $4.1 \times 10^6$ C$_8$H$_{17}$-O $\rightarrow$ 1,5-Isomerization
40) $3.8 \times 10^{-11}$ C$_8$H$_{17}$-O + NO $\rightarrow$ Nitrite
41) $3.8 \times 10^{-11}$ C$_8$H$_{17}$-O + NO$_2$ $\rightarrow$ Nitrate
42) $1 \times 10^{-12}$ C$_5$H$_{11}$ + O$_2$ $\rightarrow$ C$_5$H$_{11}$-O$_2$
43) $5 \times 10^{-14}$ C$_5$H$_{11}$-O$_2$ + C$_5$H$_{11}$-O$_2$ $\rightarrow$ C$_5$H$_{11}$-O + C$_5$H$_{11}$-O + O$_2$
43a) $5 \times 10^{-14}$ C$_5$H$_{11}$-O$_2$ + C$_5$H$_{11}$-O$_2$ $\rightarrow$ VALALD + ALCOHOL5
44) $8.43 \times 10^{-12}$ C$_5$H$_{11}$-O$_2$ + NO $\rightarrow$ C$_5$H$_{11}$-O + NO$_2$
44a) $4.74 \times 10^{-13}$ C$_5$H$_{11}$-O$_2$ + NO $\rightarrow$ Nitrate
45) $1.3 \times 10^{-14}$ C$_5$H$_{11}$-O + O$_2$ $\rightarrow$ HO$_2$ + VALALD
46) $2.9 \times 10^2$ C$_5$H$_{11}$-O $\rightarrow$ HCHO + MP
46A) $6.7 \times 10^4$ C$_5$H$_{11}$-O $\rightarrow$ 1,5-Isomerization
47) $3.8 \times 10^{-11}$ C$_5$H$_{11}$-O + NO $\rightarrow$ Nitrite
48) $3.8 \times 10^{-11}$ C$_5$H$_{11}$-O + NO$_2$ $\rightarrow$ Nitrate
49) $2.9 \times 10^{-12}$ MP + O$_2$ $\rightarrow$ MP-O$_2$
50) $1.125 \times 10^{-13}$ MP-O$_2$ + MP-O$_2$ $\rightarrow$ MP-O + MP-O + O$_2$
50a) $1.375 \times 10^{-13}$ MP-O$_2$ + MP-O$_2$ $\rightarrow$ ALCOHOL3 + MPROP + O$_2$
51) $8.9 \times 10^{-12}$ MP-O$_2$ + NO $\rightarrow$ MP-O + NO$_2$
52) $3.8 \times 10^{-11}$ MP-O + NO $\rightarrow$ Nitrite
p-O + NO → Nitrite
p-O + NO → p-O Nitro
p-NO + p-NO → Acetone
p-O + p-O + Acetone
p-NO + p-NO + Nitric

Reactions of the p-Propyl Radical

3:8x10-11
C8-O + NO2 → Nitrate
3:8x10-11
C8-O + NO → Nitrite
4:3x10-9
C8-O + CH3CHO + CH3I
1:8x10-14
C8-O + H2O + CH3I
2:9x10-12
C8-O + NO2 + CH3I
2:9x10-12
C8-O + NO + CH3I
4:9x10-12
C8-O + NO2 + CH3I

4:9x10-16
C8-O + NO2 + C8-O + O2
1:1x10-12
C8 + O2 + C8-O2

3:8x10-11
T MHIB-O + NO2 → Nitrate
3:8x10-11
T MHIB-O + NO → Nitrite
8:4x10-6
T MHIB-O + I-2 Isomerization
1:9x10-6
T MHIB-O + HCHO
1:1x10-14
T MHIB-O + NO2 + H2O + C8-O2
1:7x10-12
T MHIB-O + NO2 + H2O + C8-O2
7:2x10-12
T MHIB-O2 + T MHIB-O2 → C8-O + Alcohols
5:1x10-14
T MHIB-O2 + T MHIB-O2 + O2
1:1x10-12
T MHIB-O2 + T MHIB-O2

Primary Radical Reactions

8:3x10-16
MPO + O2 → NO2 + H2O
3:8x10-11
MPO + NO2 → Nitrite

89
\[ \begin{align*}
\text{j41)} & \quad 3.5 \times 10^{-11} \quad \text{P-O + NO} \rightarrow \text{Nitrate} \\
\text{j42)} & \quad 8.0 \times 10^{-15} \quad \text{P-O + O}_2 \rightarrow \text{ACETONE} + \text{HO}_2 \\
\text{j43)} & \quad 1.5 \times 10^{-1} \quad \text{P-O} \rightarrow \text{CH}_3 + \text{CH}_3\text{CHO} \\
\end{align*} \]

**Neopentyl Radical Reactions**
Same as in 2,2,4-Trimethylpentane mechanism.

**Methyl Radical Reactions**
Same as in 2,2,4-Trimethylpentane mechanism.
3.5.1 2,2,4-Trimethylpentane

A mechanism for 2,2,4-trimethylpentane that does not include alkoxy radical isomerization reactions overpredicts, by a factor of 2, the yield of acetone, the major carbonyl formed from attack on the tertiary carbon of 2,2,4-trimethylpentane. Since the majority of the acetone is generated from the tertiary reactions of 2,2,4-trimethylpentane, this overprediction indicates that alkoxy radical isomerization pathways are important. The major isomerization pathway is via 1,5-H atom shift, and in the case of 2,2,4-trimethylpentane, the only radicals that are likely to undergo this shift are the tertiary and primary alkoxy radicals. Rate constants for 1,5-H atom isomerizations are available in Atkinson [1]. Incorporating the alkoxy radical isomerization reactions into the mechanism produces predicted yields that agree reasonably well with measured concentrations of acetone and 2-methyl-1-propanal, errors of +1.5% and +16%, as compared to observed results, respectively. The mechanism consistently underpredicts formaldehyde formation. Observed and predicted concentrations of acetone, 2-methyl-1-propanal, and formaldehyde are presented in Figures 3.11, 3.12, and 3.13, respectively.

The elevated experimental formaldehyde yields relative to those predicted by the mechanism may be attributed to reactions of isomerization products for which standards were not available. Each product formed from 1,5-H atom isomerization is expected to result in the formation of at least one HCHO molecule. Examination of the mechanisms in Figures 3.2–3.4 shows that each 1,5-isomerization for an alkoxy radical either bypasses or delays the
Figure 3.11: Observed data (points) and predicted concentrations excluding alkoxy radical isomerization and including 1,5-isomerization for 2-methyl-1-propanal from 2,2,4-trimethylpentane photooxidation.
Figure 3.12: Observed data (points) and predicted concentrations excluding alkoxy radical isomerization and including 1,5-isomerization for acetone from 2,2,4-trimethylpentane photooxidation.
Figure 3.13: Observed data (points) and predicted concentrations excluding alkoxy radical isomerization and including 1,5-isomerization for formaldehyde from 2,2,4-trimethylpentane photooxidation.
formation of one molecule of HCHO. Thus, the line labeled "No Isomerization" in Figure 3.13 actually represents the HCHO production resulting from isomerization when HCHO formation is accounted for from further reactions of isomerization products.

A 1,5-H atom isomerization step for the secondary alkoxy radical from 2,2,4-trimethylpentane is not possible. However, another type of isomerization, 1,4- H atom shift, may be possible for the secondary radical in the 2,2,4-trimethylpentane system. Given the reasonably good agreement between the predicted yield, including isomerization pathways for the tertiary and primary radicals, and the experimental yield for 2-methyl propanal, it is unlikely that there is a significant 1,4- H atom shift pathway for the secondary alkoxy radical. Such a pathway would result in poor agreement between observations and predictions for the yield of 2-methyl propanal. An estimate for the rate constant for a 1,4-H atom shift with abstraction from a -CH₃ group can be made from comparisons of observed and predicted concentrations of 2-methyl propanal. 1,4-Isomerization reactions for the secondary alkoxy radical, TMPR2-O, and the primary alkoxy radical, TMPR3-O, resulting in abstraction of H-atoms from a -CH₃ and a -CH₂ group, respectively, were included in the 2,2,4-trimethylpentane mechanism. The rate constant for 1,4-isomerization corresponding to H-atom abstraction from a -CH₃ group was varied from 1x10⁷-1x10⁸ s⁻¹. From previous studies, the ratio of rate constants for 1,5-isomerization corresponding to abstraction from a -CH₃ group is approximately a factor of 100 smaller than the corresponding abstraction from a -CH₂ group [1, 18]. Thus, the rate constant for 1,4-isomerization of the primary alkoxy radical was varied from 1x10³-1x10⁷ s⁻¹, simultaneous
to the variation of the rate constant for 1,4-isomerization of the secondary alkoxy radical. The resulting 2-methyl-1-propanal yield was compared to the experimentally derived yield, and an estimate for this isomerization step was obtained by calculating the error between the yields. Upper limits to the rate constants for 1,4-isomerization with H-atom abstraction from a -CH₃ and a -CH₂ group were determined to be $1 \times 10^3$ and $1 \times 10^5$ s⁻¹, respectively. Initial comparisons of the 1,4- isomerization rate constant for abstraction from a -CH₃ group determined in this work to literature data [7] show that the value determined here is approximately an order of magnitude larger than the value reported in the literature. However, the uncertainty factor of 60 reported by Baldwin et al. [7] places the value for 1,4-isomerization with abstraction from a -CH₃ group reported in this work in reasonable agreement with previous estimates.

3.5.2 2,2,5-Trimethylhexane

Initially, the mechanism for the 2,2,5-trimethylhexane system included all of the reactions except the isomerization reactions. Once again, comparisons of the predicted and observed yields of the major products formed (acetone and 3,3 dimethylbutyraldehyde) indicate that the mechanism without alkoxy radical isomerization overpredicts the observed yields. Once isomerization reaction steps corresponding to 1,5-H atom shifts are added to the mechanism, the experimental results are more closely represented. The predicted yield for 3,3-dimethylbutyraldehyde is approximately 22% larger than the observed yield. However, the mechanism predictions for acetone are
still approximately 150% larger than the experimental results. This observation indicates either that additional and significant isomerization reactions other than 1,5-H shifts are possible with the radicals formed in the photooxidation of larger alkanes, rate constants corresponding to 1,5-H atom shifts for branched compounds are too low, or decomposition and O$_2$ reaction rate constants for the alkoxy radicals are too high. Because there was excellent agreement between observed and predicted concentrations for the case of 2,2,4-trimethylpentane using rate constant data available for 1,5-isomerization, decomposition, and O$_2$ reactions of the alkoxy radicals, we must conclude that there are additional isomerization pathways for 2,2,5-trimethylhexane that are insignificant for 2,2,4-trimethylpentane. Therefore, 1,4-isomerization alkoxy radical reactions were included in the mechanisms, and the results compared to the observed data. The rate constants used for the 1,4-isomerizations were those determined from fits of the 2-methyl-1-propanal data from the photooxidation of 2,2,4-trimethylpentane (see the previous section). Inclusion of 1,4-isomerization steps, where applicable, provide a better fit to the acetone data. The predicted acetone yield is only 18% larger than the observed yield. The 3,3-dimethylbutyraldehyde data remained unchanged. In all cases, however, the formaldehyde and acetaldehyde yields are underpredicted. Experimental data and mechanism predictions for 3,3-dimethylbutyraldehyde, acetone, acetaldehyde, and formaldehyde appear in Figures 3.14-3.17, respectively.

The underpredicted yields for acetaldehyde and formaldehyde, as in the case of 2,2,4-trimethylpentane, is likely a result of reactions of isomerization products from the alkoxy radicals. To examine this, we used the predicted
Figure 3.14: Depicted are the observed and the predicted concentrations for 3,3-dimethylbutyraldehyde from 2,2,5-trimethylhexane photooxidation. The predicted concentrations are based on mechanisms that exclude isomerizations and include both 1,4- and 1,5- isomerizations.
Figure 3.15: Observed and predicted concentrations for acetone from 2,2,5-trimethylhexane photooxidation. The predicted concentrations are based on mechanisms that exclude all alkoxy radical isomerizations, include only 1,5-isomerizations, and include both 1,4- and 1,5-isomerizations.
Figure 3.16: Observed and predicted concentrations for acetaldehyde from 2,2,5-trimethylhexane photooxidation. The line labeled "Correction Estimated" takes into account the formation of acetaldehyde from the reactions of alkoxy radical isomerization products.
Figure 3.17: Observed and predicted concentrations for formaldehyde from 2,2,5-trimethylhexane photooxidation. The line labeled "Correction Estimated" takes into account the formation of formaldehyde from the secondary reactions of alkoxy radical isomerization products.
concentrations of products resulting from the alkoxy radical 1,4- and 1,5-isomerization steps to calculate the amount of HCHO that would result from continued reaction of these products. Most of the pathways that are capable of undergoing 1,4- and 1,5-isomerizations lead to the direct formation of at least one molecule of HCHO. In addition, acetaldehyde is generated from the reaction of the 1,5-isomerization product resulting from the alkoxy radical, TMH2A-O (see Figure 3.7). Assuming that only one HCHO molecule is formed in any of the reactions of the products of alkoxy radical isomerizations, a lower bound estimate for the production of HCHO is possible, and included in Figure 3.17. The yield of the isomerization products from the single alkoxy radical isomerization reaction producing CH$_3$CHO as a product was used to correct the CH$_3$CHO yield, and the result appears in Figure 3.16. The predicted yields based on the formation of only one HCHO or CH$_3$CHO molecule per reaction of alkoxy radical isomerization product are still lower than the observed yields, but it is important to note that we have not taken into account the full reaction mechanisms of these products.

As discussed earlier, the neopentyl radicals are not expected to react to form 2,2-dimethyl-1-propanal in the presence of NO$_x$. Regardless of whether or not isomerization reactions are included in the overall mechanisms for 2,2 4-trimethylpentane or 2,2,5-trimethylhexane, the predicted yields of 2,2-dimethyl-1-propanal are negligible. This result lends credence to the supposition that the 2,2-dimethyl-1-propanal detected in the photooxidation of 2,2,5-trimethylhexane is a result of secondary reaction of OH with 3,3-dimethylbutyraldehyde, rather than primary formation from the neopentyl radical.
3.6 Conclusions

Experiments have been performed to investigate the photooxidation of two relatively large alkanes, 2,2,4-trimethylpentane and 2,2,5-trimethylhexane. The first kinetic study of the OH + 2,2,5-trimethylhexane reaction resulted in a rate constant of \((5.267 \pm 0.147) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\), in excellent agreement with a predicted value of \(5.7 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\), using Atkinson’s structure reactivity relationship[15]. Carbonyl products have been identified from laboratory experiments with each alkane, and the major products observed are consistent with those expected from the mechanisms. Mechanisms for each of the alkanes were developed to examine the effect of alkoxy radical isomerization reactions on predicted carbonyl yields. Using the mechanism for 2,2,4-trimethylpentane photooxidation, upper limits for the rate constants of 1,4-H atom isomerization with abstraction from -CH$_3$ and -CH$_2$ groups have been determined as \(1 \times 10^3\) and \(1 \times 10^5\) s$^{-1}$, respectively.

Acknowledgement

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Bibliography


Chapter 4

Mechanism of Atmospheric Photooxidation of Aromatics: A Theoretical Study

Abstract

The mechanisms of atmospheric photooxidation of aromatic compounds are of seminal importance in the chemistry of the urban and regional atmosphere. It has been difficult to experimentally account for the full spectrum of oxidation products in laboratory studies. In an effort to fully elucidate the atmospheric reaction pathways for the aromatic-OH reaction, we have conducted theoretical calculations on aromatic intermediates. Energies have been determined for these intermediates using semi-empirical UHF/PM3 geometry optimizations combined with ab initio calculations using density functional theory (DFT). A hybrid DFT model, the Becke 3 parameter functional with the non-local correlation functional of Lee, Yang and Parr, was used in conjunction with the 6-31G(d,p) basis set to study the intermediate structures. Full mechanisms for the OH-initiated photooxidation of toluene, m-xylene, p-xylene, 1,2,4- trimethylbenzene and m-ethyltoluene are developed. The lowest energy intermediates have been determined and predicted products from these structures are compared to available experimental product data. These studies serve to refine proposed mechanisms currently available for toluene, m-xylene and p-xylene, while providing new information on the 1,2,4- trimethylbenzene and m-ethyltoluene reaction pathways.
4.1 Introduction

Aromatic compounds are of great interest in atmospheric chemistry because of their abundance in motor vehicle emissions and because of their reactivity with respect to ozone and organic aerosol formation. Understanding the atmospheric oxidation mechanisms of aromatics has long been cited as the most critical need in further development of reaction mechanisms for the urban and regional atmosphere\(^1\). The major atmospheric sink for aromatics is reaction with the hydroxyl radical. Whereas rate constants for the OH reaction with aromatics have been well characterized\(^2\), mechanisms of aromatic oxidation following the initial OH attack have been highly uncertain. In experimental photooxidation studies of toluene, m-xylene, and p-xylene (summaries of which are available in \([2, 3, 4]\)), typically less than 50% of the reacted carbon has been identified as products. A difficulty in accounting for the remaining fraction of reacted carbon lies in the fact that the intermediates involved in aromatic-OH oxidation have not been unambiguously identified. Consequently, theoretical studies can be extremely useful in evaluating the possible mechanisms of oxidation. We present here theoretical calculations on possible structural intermediates formed in the OH-initiated reactions of toluene, m-xylene, p-xylene, 1,2,4-trimethylbenzene, and m-ethyltoluene. The aromatics chosen correspond to ones which our group has studied in the laboratory. From these energy determinations, the most likely intermediate species have been identified, and overall mechanisms for the photooxidation of these aromatics are proposed.

The aromatic-OH reaction proceeds by both abstraction and addition
pathways (see [2] and [3] and references therein). The addition pathway, occurring roughly 90% of the time, is the more prevalent route. Products identified in the OH addition reactions of toluene, m-xylene, and p-xylene in previous laboratory studies appear in Tables 4.1-4.3. The yields presented in the tables are defined as the ratio of the molar concentration of the product to that of reacted aromatic. As noted above, generally less than 50% of the carbon in these systems has been accounted for experimentally. In the current study, we compare available experimental product data for the toluene, p-xylene, and m-xylene systems to our proposed mechanisms to evaluate the extent of agreement between the theoretical mechanisms and observed product data. The theory is then used to predict additional products that might be found in these three systems. Since only two studies on products resulting from 1,2,4-trimethylbenzene photooxidation are available \cite{5, 6}, neither considering the full mechanism for photooxidation, and no product data on the reactions of m-ethyltoluene are available, we develop plausible atmospheric reaction mechanisms for both of these species based on theory alone.

4.2 Calculation Methods

The essential element of the theoretical approach is to employ semi-empirical and \textit{ab initio} quantum mechanical techniques to determine the most energetically favored intermediates in the reaction mechanisms, taking into account transition state complexes. These methods are used to determine the lowest energy structures in the most computationally efficient manner, balancing the desire for reasonable computational times with the highest level of the-
Table 4.1: Toluene Photooxidation Products: Molar Yields

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>0.105 ± 0.019</td>
<td>[6]</td>
</tr>
<tr>
<td></td>
<td>0.058</td>
<td>[7]</td>
</tr>
<tr>
<td></td>
<td>0.150 ± 0.040</td>
<td>[8]</td>
</tr>
<tr>
<td></td>
<td>0.080</td>
<td>[9]</td>
</tr>
<tr>
<td>Methyl Glyoxal</td>
<td>0.146 ± 0.006</td>
<td>[6]</td>
</tr>
<tr>
<td></td>
<td>0.077</td>
<td>[7]</td>
</tr>
<tr>
<td></td>
<td>0.140 ± 0.04</td>
<td>[8]</td>
</tr>
<tr>
<td></td>
<td>0.075</td>
<td>[9]</td>
</tr>
<tr>
<td>Methylbutenedial</td>
<td>0.058</td>
<td>[7]</td>
</tr>
<tr>
<td>Hydroxymethylbutenedial</td>
<td>0.034</td>
<td>[7]</td>
</tr>
<tr>
<td>Oxoheptadienal</td>
<td>0.027</td>
<td>[7]</td>
</tr>
<tr>
<td>Methylhydroperoxide</td>
<td>0.018</td>
<td>[7]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>0.010</td>
<td>[7]</td>
</tr>
<tr>
<td>Hexadienyl</td>
<td>0.010</td>
<td>[7]</td>
</tr>
<tr>
<td>Hydroxyoxoheptadienal</td>
<td>0.010</td>
<td>[7]</td>
</tr>
<tr>
<td>Maleic Anhydride</td>
<td>0.040 ± 0.004</td>
<td>[8]</td>
</tr>
<tr>
<td>*-Cresol</td>
<td>0.204 ± 0.027</td>
<td>[10]</td>
</tr>
<tr>
<td>m- and p-Cresol</td>
<td>0.048 ± 0.009</td>
<td>[10]</td>
</tr>
<tr>
<td>*-Nitrotoluene</td>
<td>(0.0135±0.0029) + (1.90±0.25)* 10^-16*[NO2]</td>
<td>[10]</td>
</tr>
<tr>
<td>o-Nitrotoluene</td>
<td>0.07*[m-nitrotoluene]</td>
<td>[10]</td>
</tr>
<tr>
<td>p-Nitrotoluene</td>
<td>0.35*[m-nitrotoluene]</td>
<td>[10]</td>
</tr>
</tbody>
</table>

Table 4.2: m-Xylene Photooxidation Products: Molar Yields

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>0.13 ± 0.03</td>
<td>[11]</td>
</tr>
<tr>
<td></td>
<td>0.086</td>
<td>[12]</td>
</tr>
<tr>
<td></td>
<td>0.086 ± 0.011</td>
<td>[6]</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>0.42 ± 0.05</td>
<td>[11]</td>
</tr>
<tr>
<td></td>
<td>0.375</td>
<td>[12]</td>
</tr>
<tr>
<td></td>
<td>0.319±0.009</td>
<td>[6]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>0.17 ± 0.02</td>
<td>[11]</td>
</tr>
<tr>
<td>2,4 Dimethylphenol</td>
<td>0.099</td>
<td>[13]</td>
</tr>
<tr>
<td>2,6 Dimethylphenol</td>
<td>0.111</td>
<td>[13]</td>
</tr>
<tr>
<td>4-Nitro-<em>m</em>-xylene</td>
<td>0.0018</td>
<td>[13]</td>
</tr>
<tr>
<td>5-Nitro-<em>m</em>-xylene</td>
<td>0.0032 + 1.6<em>10^-17</em>[NO2]</td>
<td>[13]</td>
</tr>
</tbody>
</table>
Semi-empirical optimizations require less computational time, and, given that all of the structures studied are similar in nature (aromatic structures with C, N, O, and H atoms), semi-empirical optimizations are adequate for screening and optimizing the structures, while *ab initio* calculations provide accurate energy determinations.

To examine the effectiveness of semi-empirical optimizations followed by *ab initio* calculations, studies were conducted on a variety of compounds, including methylhydroxycyclohexadienyl radicals resulting from toluene photooxidation, alkyl radicals, alkoxy radicals and nitroalkenes, using different geometry optimization and single point energy calculation schemes. The first study compared the use of high and low level computational techniques to examine methylhydroxycyclohexadienyl radical formation from OH addition to toluene. Since experimental data for the energies of these radicals are not available, comparisons of theoretical and experimental results are not possible. However, these calculations were used in order to find a balance between the highest level of theory and the computational time required to accurately predict the stability of radicals that are similar to those studied.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>0.24 ± 0.02</td>
<td>[11]</td>
</tr>
<tr>
<td></td>
<td>0.225 ± 0.039</td>
<td>[6]</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>0.12 ± 0.02</td>
<td>[11]</td>
</tr>
<tr>
<td></td>
<td>0.105 ± 0.034</td>
<td>[6]</td>
</tr>
<tr>
<td>3-Hexene-2,5 Dione</td>
<td>Detected</td>
<td>[11]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>0.17 ± 0.02</td>
<td>[11]</td>
</tr>
<tr>
<td>2,5 Dimethylphenol</td>
<td>0.188</td>
<td>[13]</td>
</tr>
<tr>
<td>2-Nitro-p-xylene</td>
<td>0.0120+2.8*10^{-10}[NO2]</td>
<td>[13]</td>
</tr>
</tbody>
</table>
in this work, assuming that the most rigorous computational method ade-
quately reflects the true energy of the species. The second study compared
the results of the chosen computational technique to experimental values to
provide a calibration of the technique.

Techniques used for geometry optimization include density functional
theory (DFT)[14] using the hybrid density functional, Becke3LYP, of Becke
[15, 16] and Lee et al. [17, 18] and a 6-31G(d,p) basis set, restricted open-
shell Hartree Fock (ROHF) optimization using a 6-31G(d,p) basis set, a semi-
empirical unrestricted Hartree-Fock PM3 (UHF/PM3) optimization[19], and
a molecular mechanics optimization using the MM+ functional form[20]. Of
the semi-empirical techniques, the PM3 parameterization, a Modified Neglect
of Diatomic Overlap (MNDO) method, is most often cited as producing the
best optimizations for hydrocarbon systems and complexes containing nitro-
gen and oxygen[21]. In addition to providing an accurate geometry, PM3
has also been shown to provide accurate vibrational frequencies that can be
compared directly to experiment without post hoc correction[22]. These vi-
brational frequencies allow for the calculation of zero-point energies. The
MM+ functional form of the potential was used in the molecular mechan-
ics optimizations since it is most appropriate for small organic molecules[20].
The DFT and ROHF optimizations were performed on HP 9000/735 worksta-
tions using Gaussian 92[18] and PSGVB, version 2.13[23], respectively, while
the UHF/PM3 and molecular mechanics optimizations were performed on
a personal computer using Hyperchem, release 4.0[24]. Single point calcu-
lations were performed following geometry optimizations utilizing either the
ROHF method or density functional theory[14] using the Becke3LYP hybrid
density functional. Both types of calculations employed a 6-31G(d,p) basis set, and were performed on HP 9000/735 workstations using Gaussian 92[18]. The results for the methylhydroxycyclohexadienyl radicals are given in Table 4.4, with relative energy defined as the difference in energy between any given structural isomer and the lowest energy structural isomer. Table 4.4 indicates that the MM+ optimization, followed by the DFT calculation, does not reproduce the results of the high-level DFT optimization followed by the DFT calculation. The UHF/PM3 geometry optimization method, however, gives relative energies that are quite close to the DFT values. Of the techniques studied, a semi-empirical UHF/PM3 geometry optimization method, followed by a single point calculation using the Becke3LYP density functional and a 6-31G(d,p) basis set, exhibits the best combination of computational efficiency and accuracy. This method was therefore chosen for all of the calculations that are presented in this study.

To calibrate the chosen computational scheme, a second study was performed. Geometry optimizations using the UHF/PM3 method followed by a single point calculation using the Becke3LYP density functional and a 6-31G(d,p) basis set were performed on radicals and stable species for which experimental energy data exist. The relative energies of isomers of the structures were compared, and are tabulated in Table 4.5. From this table it is evident that the PM3/Becke3LYP technique adequately reproduces the relative energies between structural isomers for several different classes of compounds. In addition, we can obtain an estimate of our accuracy by comparing experimentally derived and theoretical relative energies. These values differ at most by 1.6 kcal/mol (which can be rounded to 2 kcal/mol). As a further
Table 4.4: Relative energies in kcal/mol for isomers of the methylhydroxy- cyclohexadieny radical using various computational methods. The 6-31G\((d,p)\) basis set was used for all \textit{ab initio} calculations.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Becke3LYP / Becke3LYP</td>
<td>[\text{a}]</td>
<td>1.1</td>
<td>1.8</td>
<td></td>
</tr>
<tr>
<td>ROHF / Becke3LYP</td>
<td>[\text{b}]</td>
<td>0.9</td>
<td>1.7</td>
<td></td>
</tr>
<tr>
<td>ROHF / ROHF</td>
<td>[\text{c}]</td>
<td>0.8</td>
<td>0.7</td>
<td></td>
</tr>
<tr>
<td>UHF-PM3 / Becke3LYP</td>
<td>[\text{d}]</td>
<td>1.2</td>
<td>1.6</td>
<td></td>
</tr>
<tr>
<td>MM+ / Becke3LYP</td>
<td>0.1</td>
<td>0 [\text{e}]</td>
<td>0.4</td>
<td></td>
</tr>
</tbody>
</table>

Absolute energies: (a) -347.3406 a.u., (b) -347.3364 a.u., (c) -345.1331 a.u., (d) -347.3374 a.u., (e) -347.3318 a.u.
calibration, $\Delta H_{rxn}$ values derived from literature data [25] and PM3/DFT calculations for the toluene + OH reaction resulting in the formation of cresol isomers are compared. These data appear in Table 4.6, and show agreement between experiment and theory to within 2 kcal/mol. Although the $\Delta H_{rxn}$ comparisons between experiment and theory are within 2 kcal/mol agreement for the case presented in Table 4.6, it is difficult to say that the agreement between experiment and theory will be at the level of 2 kcal/mol for all cases studied in this work. However, because literature data were not available to indicate a reasonable level of error for $\Delta H_{rxn}$ values, and we have not performed a variety of $\Delta H_{rxn}$ calculations on reactions involving the alteration of carbon centers, interpretations of inclusions/exclusions of reaction pathways are limited to those which involve large differences between $\Delta H_{rxn}$ values for different reaction pathways.

Because molecular mechanics provides a fast, first approximation to the optimized structure, this technique was used before employing the UHF/PM3 method in the present study of aromatic mechanisms. To examine whether the individual structures investigated were at global rather than local minima, several starting geometries of each structure were subjected to geometry optimization calculations. The heats of formation (obtained at the PM3 level) of the different geometries were compared, and the lowest energy structure was chosen for subsequent single point ab initio calculations. In cases where the semi-empirically derived heats of formation were similar, several conformations were subjected to single point ab initio calculations to locate the lowest energy conformation. All structures were characterized as being genuine minima (i.e. having no imaginary frequencies) at the PM3 level. The
Table 4.5: A comparison of relative energies of different structural isomers based on experimental data and theoretical calculations.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Relative Energy - Experiment $^a$ (kcal/mol)</th>
<th>Relative Energy - Calculated $^b$ (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Structure Image]</td>
<td>3.8</td>
<td>4.3</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>4.9</td>
<td>3.3</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>6.6</td>
<td>6.1</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>2.6</td>
<td>3.3</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>5</td>
<td>5.1</td>
</tr>
<tr>
<td>![Structure Image]</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

(a) Experimental data from NIST Standard Reference Database 25 [28].
(b) Relative energies determined using UHF/PM3 geometry optimization followed by Becke3LYP single point energy calculation.
Table 4.6: A comparison of experimentally and theoretically derived $\Delta H_{rxn}$ values for the Toluene + OH reaction.

<table>
<thead>
<tr>
<th>Reactants</th>
<th>Products</th>
<th>$\Delta H_{rxn}$, Theoretical (kcal/mol)</th>
<th>$\Delta H_{rxn}$, Experimental (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathcal{C}$ + OH</td>
<td>$\mathcal{C}$OH + H</td>
<td>+1.38</td>
<td>+0.8 $\pm$ 0.4</td>
</tr>
<tr>
<td></td>
<td>$\mathcal{C}$OH + H</td>
<td>+1.25</td>
<td>-0.8 $\pm$ 0.7</td>
</tr>
<tr>
<td></td>
<td>$\mathcal{C}$OH + H</td>
<td>+1.94</td>
<td>+0.9 $\pm$ 0.4</td>
</tr>
</tbody>
</table>
Becke3LYP energy was subsequently corrected to 298 K using the normal mode vibrational frequencies and thermal corrections obtained at the PM3 level. Thus, it is this corrected energy that is used to determine the lowest energy intermediate species in a particular step of the reaction mechanism.

Transition states were calculated with a constrained optimization procedure at the UHF/PM3 level using Gaussian 92. Conventional direct optimization schemes available through Gaussian 92 were attempted. However, because of the complexity of the structures studied, it was difficult to locate the structure with an initial geometry corresponding to a single negative eigenvalue of the second derivative matrix, a condition which is necessary before a conventional transition state optimization can proceed. The constrained optimization method relies on the assumption that the transition states are governed by the formation of a single bond in each step (for example, the C-O bond in the formation of the methylhydroxycyclohexadienyl radicals). Thus, the bond distance is adjusted by small increments (0.1 Å steps, in this case), and frozen while the geometry of the molecule is optimized. At the end of each optimization, the PM3-derived energy of the structure is noted. This procedure is repeated until a point of maximum energy is obtained. The constrained optimization procedure at the UHF/PM3 level was used to obtain an initial estimate of the transition state. Once this estimated transition state structure was found, DFT single point calculations were performed on this structure and small geometrical perturbations of the structure to ensure that the UHF/PM3 constrained optimization was successful in determining an estimate for the transition state geometry. This estimate of the true transition state is assumed to be an upper limit to the
first-order saddle point due to the elementary means of accounting for electron correlation. Because of the large number of structures considered in this study, transition states were located for representative cases and subsequently applied to homologous transition state complexes.

4.3 Results of Calculations

A generalized summary of the steps considered in the OH-aromatic mechanism appears in Figure 4.1. Each reaction is subsequently treated individually. Figures 4.2 - 4.6 show the reaction coordinate diagrams (Relative energy in kcal/mol versus reaction path) for the lowest energy structures found for each of the five aromatic-OH reactions. Energy and geometry data for the various structures studied in this work are available throughout this manuscript for comparison. Additional data are available from the authors upon request.

4.3.1 Initial Hydroxyl Radical Attack

There are several possible sites of attack for the OH radical in each of the aromatics considered. Some sites are less sterically hindered than others or are favored because of stabilizations as a result of group interactions. If we consider the transition states leading to the formation of the substituted hydroxycyclohexadienyl radicals, the aromatic-OH adducts formed in Reaction 1b, we find that the barrier heights are essentially negligible. Several different geometries were considered for each aromatic. Shown in Table 4.7 are
Figure 4.1: Possible reactions in a generalized mechanism of Aromatic-OH photooxidation. “R” represents either an H atom or a methyl/ethyl group, depending on the aromatic considered.
Figure 4.2: Reaction coordinate diagram for toluene. Indicated are the three favored pathways. Shown below the plots are the corresponding aromatic-OH adduct, peroxy radical, and bicyclic radical corresponding to the pathways. The structures are labeled P1, P2, P3 corresponding to paths 1, 2, and 3, respectively.
Figure 4.3: Reaction coordinate diagram for \(m\)-xylene. Indicated are the energies of the two favored pathways. Shown below the plots are the corresponding aromatic-OH adduct, peroxo radical, and bicyclic radical corresponding to the pathways. The structures are labeled P1 and P2 corresponding to paths 1 and 2, respectively.
Figure 4.4: Reaction coordinate diagram for $p$-xylene. Shown below the path are the corresponding aromatic-OH adduct, per oxy radical, and bicyclic radical corresponding to the favored pathway.
Figure 4.5: Reaction coordinate diagram for 1,2,4-trimethylbenzene. Indicated are the energies of the three favored pathways. Shown below the diagram are the corresponding aromatic-OH adduct, peroxyl radical, and bicyclic radical corresponding to the pathways. The structures are labeled P1, P2, and P3 corresponding to paths 1, 2 and 3, respectively.
Figure 4.6: Reaction coordinate diagram for $m$-ethyltoluene. Indicated are the energies of the three favored pathways. Shown below the diagram are the corresponding aromatic-OH adduct, peroxy radical, and bicyclic radical corresponding to the pathways. The structures are labeled P1, P2, P3 corresponding to paths 1, 2, and 3, respectively.
the PM3-derived equilibrium structures for a representative sample of the most stable aromatic-OH adducts. Total energies for aromatic-OH adducts are tabulated in Table 4.8. Indicated are single point energies at 0 K derived using density functional theory, and energies corrected to 298 K using vibrational information generated at the PM3 level. In general, the preferred place of OH addition, based on total energies of the aromatic-OH adducts, is a position ortho to a substituent methyl group. In considering the total energies of the each of the aromatic-OH isomers, we did, however, find that there were some cases where isomers had energies within 2 kcal/mol of the lowest energy structures. So, whereas the structures with OH attachment at the ortho position are lowest in energy, pathways that include aromatic-OH structures whose relative energy falls within ±2 kcal/mol of the lowest energy isomer will also be considered. Throughout our discussion, we compare available experimental product data to results predicted by the theoretical calculations. This comparison serves two purposes: to lend credence to our uncertainty limit of 2 kcal/mol and to show that the theoretical predictions are adequate representations of atmospheric and/or experimental conditions.

Toluene

The lowest energy aromatic-OH structure for toluene is that resulting from addition to the ortho position. However, OH additions to the meta and para positions yield structures that are only 1.6 and 1.2 kcal/mol higher in energy than addition at the ortho site. Thus, based on theoretical considerations, additions to the meta and para positions cannot be excluded from reaction
Table 4.7: PM3 equilibrium structures for a representative selection of the most stable aromatic-OH adducts formed from OH addition to the aromatic ring (Reaction 1b). Bond distances are in Å, and angles are in degrees.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Toluene</th>
<th>M-xylene</th>
<th>P-xylene</th>
<th>M-ethyltoluene</th>
</tr>
</thead>
<tbody>
<tr>
<td>r(C6,C7)</td>
<td>1.505</td>
<td>1.505</td>
<td>1.505</td>
<td>1.507</td>
</tr>
<tr>
<td>r(C6,H)</td>
<td>1.414</td>
<td>1.416</td>
<td>1.415</td>
<td>1.416</td>
</tr>
<tr>
<td>r(C2,C3)</td>
<td>1.498</td>
<td>1.507</td>
<td>1.497</td>
<td>1.507</td>
</tr>
<tr>
<td>r(C6,C8)</td>
<td>1.370</td>
<td>1.379</td>
<td>1.377</td>
<td>1.379</td>
</tr>
<tr>
<td>r(C6,C9)</td>
<td>1.411</td>
<td>1.407</td>
<td>1.416</td>
<td>1.408</td>
</tr>
<tr>
<td>r(C6,C10)</td>
<td>1.406</td>
<td>1.407</td>
<td>1.405</td>
<td>1.407</td>
</tr>
<tr>
<td>r(C6,C11)</td>
<td>1.381</td>
<td>1.379</td>
<td>1.381</td>
<td>1.379</td>
</tr>
<tr>
<td>r(C6,C12)</td>
<td>1.479</td>
<td>1.479</td>
<td>1.478</td>
<td>1.479</td>
</tr>
<tr>
<td>r(C6,C13)</td>
<td></td>
<td>1.479</td>
<td>-</td>
<td>1.490</td>
</tr>
<tr>
<td>r(C6,C14)</td>
<td></td>
<td>-</td>
<td>1.486</td>
<td>-</td>
</tr>
<tr>
<td>θ(C6,C12,C7)</td>
<td>128</td>
<td>127</td>
<td>127</td>
<td>128</td>
</tr>
<tr>
<td>θ(C6,C8,C9)</td>
<td>128</td>
<td>127</td>
<td>127</td>
<td>128</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>1,2,4-TMB</th>
</tr>
</thead>
<tbody>
<tr>
<td>r(C6,C7)</td>
<td>1.388</td>
</tr>
<tr>
<td>r(C6,H)</td>
<td>1.417</td>
</tr>
<tr>
<td>r(C2,C3)</td>
<td>1.507</td>
</tr>
<tr>
<td>r(C6,C8)</td>
<td>1.506</td>
</tr>
<tr>
<td>r(C6,C9)</td>
<td>1.378</td>
</tr>
<tr>
<td>r(C6,C10)</td>
<td>1.406</td>
</tr>
<tr>
<td>r(C6,C11)</td>
<td>1.413</td>
</tr>
<tr>
<td>r(C6,C12)</td>
<td>1.486</td>
</tr>
<tr>
<td>r(C6,C13)</td>
<td>1.480</td>
</tr>
<tr>
<td>r(C6,C14)</td>
<td>1.479</td>
</tr>
<tr>
<td>θ(C6,C8,C9)</td>
<td>127</td>
</tr>
</tbody>
</table>
Table 4.8: Calculated energies (in a.u.) of the most stable reaction intermediates for the pathway leading to the formation of the bicyclic peroxy radicals.

(a) Addition sites in parenthesis indicate comparable low energy structures.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Aromatic</th>
<th>OH Addition (Rxn. 1b)</th>
<th>O₂ Addition (Rxn. 2b)</th>
<th>O₂ Bridge Formation (Rxn. 4)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Site</td>
<td>E°</td>
<td>E°²⁶⁸</td>
<td>Site</td>
</tr>
<tr>
<td>toluene</td>
<td>3</td>
<td>-347.3348</td>
<td>-347.1676</td>
<td>(4,6)</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>-347.3355</td>
<td>-347.1683</td>
<td>(1)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>-347.3374</td>
<td>-347.1902</td>
<td>(3,1.5)</td>
</tr>
<tr>
<td>m-xylene</td>
<td>4</td>
<td>-366.6598</td>
<td>-366.4826</td>
<td>(3,1)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>-366.6603</td>
<td>-366.4946</td>
<td>(3)</td>
</tr>
<tr>
<td>p-xylene</td>
<td>2</td>
<td>-386.6553</td>
<td>-386.4482</td>
<td>(3,1.5)</td>
</tr>
<tr>
<td>1,2,4-</td>
<td>6</td>
<td>-425.9739</td>
<td>-425.7671</td>
<td>1</td>
</tr>
<tr>
<td>trimethyl-</td>
<td>3</td>
<td>-425.9773</td>
<td>-425.7705</td>
<td>2</td>
</tr>
<tr>
<td>benzene</td>
<td>5</td>
<td>-425.9774</td>
<td>-425.7706</td>
<td>(2,2)</td>
</tr>
<tr>
<td>m-ethyl-</td>
<td>4</td>
<td>-425.9740</td>
<td>-425.7675</td>
<td>(3,1)</td>
</tr>
<tr>
<td>toluene</td>
<td>6</td>
<td>-425.9742</td>
<td>-425.7677</td>
<td>(1,1.3)</td>
</tr>
</tbody>
</table>
mechanism considerations. Following formation of the adduct, molecular oxygen can abstract a hydrogen atom to form a cresol. There is experimental evidence for o-, m-, and p-cresol, with the o-cresol yield dominating cresol formation (Table 4.1), consistent with the theoretical predictions.

m-Xylene

The lowest energy aromatic-OH structure for m-xylene also corresponds to OH addition to the 2- position. Additions to the 4- and 5- positions are 0.88 and 3.5 kcal/mol higher in energy than addition to the 2- position. Additions to the 6- and 4- positions create identical structures. Based on these results, the lowest energy adducts for m-xylene correspond to OH addition to the 2- and 4- positions, both of which are ortho to one of the substituent methyl groups. The reaction coordinate diagram for m-xylene, Figure 4.3, depicts the lowest energy adducts formed. Comparing the theoretical predictions to experimentally determined products in Table 4.2, we note experimental evidence for 2,6- and 2,4- dimethylphenol (with the 2,6- dimethylphenol yield dominating), consistent with OH addition to the 2- and 4- positions. There is no experimental evidence for 3,5- dimethylphenol, consistent with our finding that OH addition to the 5- position can be excluded.

p-Xylene

Because of the high degree of symmetry of p-xylene, only one structure was considered. The experimental evidence for O₂ reaction with the p-xylene-OH adduct leading to the formation of only 2,5-dimethylphenol (see Table
indicates that there is only one possible pathway for OH addition to the ring. The reaction coordinate diagram for \( p \)-xylene (Figure 4.4) and Table 4.7 show the structure for the OH-aromatic adduct.

**1,2,4-Trimethylbenzene**

The lowest energy aromatic-OH adducts for trimethylbenzene correspond to OH additions at the 3- and 5- positions. The energies of these two adducts are identical. Addition to the 6- position is 2.1 kcal/mol higher in energy than the other two adducts. The reaction coordinate diagram for 1,2,4-trimethylbenzene, Figure 4.5, details the path of the favored OH-aromatic adduct structures. Given that there are no experimental data available on the ring-retaining products of 1,2,4-trimethylbenzene, the given energy calculations can be used to predict the phenolic compounds formed. Thus, we predict the formation of 2,3,6-trimethylphenol, 2,3,5-trimethylphenol and 2,4,5-trimethylphenol from the \( O_2 \) reaction with the adducts formed from OH addition to the 3-, 6- and 5- positions, respectively (according to Reaction 2a).

**\( m \)-Ethyltoluene**

The lowest energy aromatic-OH adduct for \( m \)-ethyltoluene corresponds to OH addition to the 2- position. The energies for the adducts formed from addition to the 4-, 5-, and 6- positions are 0.69, 3.0, and 0.56 kcal/mol higher than that for addition to the 2- position. From these values, we conclude that the aromatic-OH adducts to be included in the reaction mechanism for \( m \)-
ethyltoluene include addition of OH to the 2-, 4-, and 6- positions. Note that these results are very similar to those of m-xylene; addition of OH to the 5-position is the least favored radical. Substituent methyl groups have a considerable effect on the favored addition sites for OH. Figure 4.6 details the paths of the three lowest energy aromatic-OH structures. As with the case of 1,2,4-trimethylbenzene, there are no experimental data available for ring-retaining products formed from the m-ethyltoluene-OH reaction. Based on our energy determinations, we predict the formation of 2-ethyl-6-methylphenol, 2-ethyl-1-methylphenol, and 4-ethyl-6-methylphenol from the O₂ reaction with the adducts formed from OH addition to the 2-, 4-, and 6- positions, respectively.

4.3.2 Fate of the OH-Aromatic Adduct: NO₂ Reaction

It has been suggested that aromatic-OH adducts react with NO₂ to yield nitroaromatics according to Reaction 3a of the generalized mechanism.²⁹ To explain the observed variation of aromatic photooxidation product yields with varying NO₂ concentration, alternative paths for aromatic-OH adduct reaction with NO₂ that results in the formation of oxy type radicals (Reaction 3e)²⁷, or the formation of a phenolic type compound and HONO (Reaction 3d)²⁷ have been proposed. Two additional pathways can be suggested for Reaction 3, pathway 3c forming the hydroxyaromatic nitrite, and 3b forming the hydroxy nitroaromatic (nitrophenol, nitro-dimethylphenol, etc.) .

Experimental evidence has been found for the presence of nitroaromatic compounds in the photooxidation of toluene, m- xylene, and p-xylene (see
Tables 4.1-4.3). While calculations were not performed on all of the nitroaromatic compounds, it will be shown in the discussion that follows that $\Delta H_{rxn}$ for Reaction 3a is -26.3 for the case of toluene. Thus, the reaction of NO$_2$ with the OH-aromatic adduct is assumed to be exothermic for all aromatics in this study. The nitroaromatics that are predicted to form involve NO$_2$ reaction with the most stable OH-aromatic adducts, using the conventional wisdom that NO$_2$ addition to an alkyl hydroxycyclohexadienyl radical will be at a site ortho to the hydroxyl substituent group.$^{[29]}$. Consistent with the theoretical prediction that all three possible toluene-OH adducts form, experimental evidence exists for three different nitrotoluene products. Similarly, consistent with the prediction that only two of the possible m-xylene-OH adducts form, experimental evidence exists for only two nitro-m-xylene products. Hydroxyl radical addition to the 2-position of m-xylene, with subsequent reaction with NO$_2$, yields 4-nitro-m-xylene, whereas addition to the 4-position yields 5-nitro-m-xylene. Observations of the nitroaromatics from the p-xylene photooxidation systems are also consistent with theory. Only one nitroaromatic, 2-nitro-p-xylene, is detected in laboratory studies. Since experimental data are not available for the nitroaromatics resulting from the 1,2,4-trimethylbenzene and m-ethyltoluene systems, the results of the previous section, indicating the favored aromatic-OH adducts, can predict the nitroaromatic species expected to form in photooxidation studies of 1,2,4-trimethylbenzene and m-ethyltoluene. From OH addition to the 3-position and 5-positions, 4-nitro-1,2,4-trimethylbenzene and 6-nitro-1,2,4-trimethylbenzene are predicted to form, and 5-nitro-1,2,4-trimethylbenzene should not form. For m-ethyltoluene, the formation of 4-nitro-m-ethyltoluene from OH
addition to the 2-position, 5-nitro-\textit{m}-ethyltoluene from OH addition to the 4-position, and 2-nitro-\textit{m}-ethyltoluene from addition to the 6-position are predicted.

To examine the stabilities of the five competing pathways for aromatic-OH adduct reaction with NO$_2$, theoretical calculations were performed for the case of toluene. Energies were calculated for nitrotoluene (Reaction 3a), nitrophenol (Reaction 3b), phenol nitrite (Reaction 3c), phenol (Reaction 3d), and oxy radicals (Reaction 3e). Only one structural isomer for each reaction (3a-3d) was considered. Transition states were determined for the first steps of Reactions 3a-3c, and for Reactions 3d and 3e. Geometrical information for these reaction intermediates as well as the transition states appear in Table 4.9. The second steps of reactions 3a-3c were not subjected to transition state analyses since these reactions involve processes (rearrangement and the subsequent loss of water or molecular hydrogen) which are difficult to analyze using the constrained optimization procedure, and are subject to interpretation. Clearly, much additional work is required to completely elucidate the transition states of the second steps of Reactions 3a-3c. Even though transition states are not included for the second steps of Reactions 3a-3c, valuable information can be gained regarding the relative stability of intermediates as well as the overall heats of reactions for the reactions considered.

The compounds, NO, NO$_2$, H$_2$O and HONO were subjected to the same type of calculations as each of the intermediate species. A PM3 geometry optimization was conducted followed by a Becke3LYP single point calculation. The single point energy was subsequently corrected to 298 K using
Table 4.9: PM3 derived geometries for (a) equilibrium and (b) approximate transition state structures for the reactions of aromatic-OH adducts with NO$_2$. All bond distances are in Å, and angles are in degrees.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Opt. Value</th>
<th>TS Value</th>
</tr>
</thead>
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<td></td>
</tr>
<tr>
<td>r(C$_2$O$_1$)</td>
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<td></td>
</tr>
<tr>
<td>r(C$_2$C$_2$)</td>
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</tr>
<tr>
<td>r(C$_2$O$_2$)</td>
<td>1.436 2.2</td>
<td></td>
</tr>
<tr>
<td>r(C$_2$O$_4$)</td>
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<tr>
<td>r(O$_3$N)</td>
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<td>r(N$_3$O$_3$)</td>
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<td>$\angle$(C$_2$C$_2$C$_2$O$_2$)</td>
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<table>
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</thead>
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<td>r(H$_2$O$_2$)</td>
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<tr>
<td>$\angle$(H$_2$O$_2$N)</td>
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the normal mode vibrational frequencies and thermal corrections obtained at the PM3 level. Table 4.10 shows the uncorrected energy at 0 K (from the Becke3LYP single point calculation) and the corrected energy at 298 K for NO, NO₂, H₂O HONO, and each of the other species participating in a reaction with an aromatic intermediate. Table 4.11 contains the energies (0 K and 298 K) of all of the products formed in the aromatic-OH adduct reaction with NO₂.

For the case of toluene as an example, the ΔH_rxn for the first steps of pathways 3a-3c, and overall pathways 3d and 3e are -26.3, -26.3, -24.5, -52.8, and +38.3 kcal/mol, respectively. The barrier heights for pathways 3a,b (first step), 3c (first step), 3d are 5, 1, 2 kcal/mol, respectively. Note that Reactions 3c and 3e are assumed to have the same transition state. Based on this information for the initial steps of Reactions 3a-3c, and the overall Reactions 3d and 3e, the only pathway that can be unambiguously excluded is Reaction 3e, because of its endothermic nature. In the absence of additional transition state data regarding the second steps of Reactions 3a-3c, it impossible to recommend the exclusion of one of these pathways. Therefore, until additional data can be obtained for Reactions 3a-3c, we recommend the inclusion of Reactions 3a-3d in overall mechanisms for aromatic-OH photooxidation. As mentioned previously, it is clear that additional computational work is required to fully elucidate the pathways of aromatic-OH reaction with NO₂.
Table 4.10: Calculated energy (in a.u.) of molecules involved in reactions with aromatic intermediates. Structures optimized using PM3 and energies determined using Becke3LYP/6-31g(d,p). (a) Electronic energy at 0 K. (b) Corrected energy at 298 K, which includes zero point energies and thermal corrections determined using PM3 normal mode vibrations.

<table>
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<th>Molecule</th>
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<th>$E^{298}$ b</th>
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</thead>
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<tr>
<td>OH</td>
<td>-75.7268</td>
<td>-75.7153</td>
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<tr>
<td>O$_2$</td>
<td>-150.3165</td>
<td>-150.3093</td>
</tr>
<tr>
<td>H$_2$O</td>
<td>-76.4190</td>
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</tr>
<tr>
<td>HONO</td>
<td>-205.6981</td>
<td>-205.6736</td>
</tr>
<tr>
<td>NO$_2$</td>
<td>-205.0700</td>
<td>-205.0573</td>
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<tr>
<td>NO</td>
<td>-129.8879</td>
<td>-129.8803</td>
</tr>
</tbody>
</table>
Table 4.11: Calculated energies (in a.u.) of the products formed by the reaction of NO₂ with the toluene-OH adduct.

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<th>Molecule</th>
<th>$E_a^\circ$</th>
<th>$E^{298}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-nitrotoluene</td>
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<td>-475.9356</td>
</tr>
<tr>
<td>2-cresol</td>
<td>-346.7967</td>
<td>-346.6581</td>
</tr>
<tr>
<td></td>
<td>-552.4537</td>
<td>-552.2865</td>
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<td>-552.2894</td>
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<tr>
<td></td>
<td>-551.2802</td>
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</tr>
<tr>
<td></td>
<td>-551.2802</td>
<td>-551.1342</td>
</tr>
</tbody>
</table>
Aromatic Peroxy Radical

Under atmospheric conditions, oxygen is expected to rapidly add to the aromatic-OH adduct, forming a peroxy radical (Reaction 2b in the generalized mechanism)\textsuperscript{[26]}. Semi-empirical geometry optimizations and \textit{ab initio} calculations were performed on all of the resulting peroxy radicals for each of the five aromatics considered. Transition state analyses indicate that the transition state energies are governed by consideration of the type of carbon atom being bonded to (secondary versus tertiary) and the steric nature of the site of attachment. The energies of the approximated transition states for the reactions of aromatic-OH adducts with O\textsubscript{2} are available in Table 4.12. From this table, it can be seen that addition of O\textsubscript{2} to a site adjacent to the OH group (such as O\textsubscript{2} addition to the 3-position with OH at the 2-position for \textit{p}-xylene) results in a transition state energy that is roughly 10 kcal/mol higher than the parent aromatic-OH adduct + O\textsubscript{2} energy. Additions of O\textsubscript{2} to a tertiary carbon give a transition state energy that is roughly 7 kcal/mol higher than the parent aromatic-OH + O\textsubscript{2} energy. Additions to a secondary carbon site not adjacent to the OH radical addition site give a transition state energy that is roughly 5 kcal/mol higher than the parent aromatic-OH + O\textsubscript{2} energy. These barrier heights are consistent with the low rate constants observed by Knispel \textit{et al.}\textsuperscript{[26]}. Table 4.13 contains the geometrical information for the structures of a representative set of the peroxy radicals. Energies for the most stable peroxy radical structures appear in Table 4.8. Examination of the relative energies of each of the peroxy radicals for a given aromatic reveals that the energies of the intermediate peroxy radicals are all similar.
Given this fact, all structural isomers of the aromatic peroxy radicals are equally likely to form.

4.3.3 Fate of the Peroxy Radical: Cyclization vs NO Reaction

Alkyl peroxy radicals present in polluted atmospheres generally react with NO to form alkoxy radicals\textsuperscript{[2, 28]}. Aromatic peroxy radicals, in contrast, were postulated to cyclize forming bicyclic radicals in order to rationalize the observed formation of α-dicarbonyls from the aromatic hydrocarbon reactions \textsuperscript{[29]}. To test this hypothesis, we examined the ΔH\textsubscript{rxn} for reactions 4 and 5 in the generalized mechanism for aromatic-OH reaction.

Calculations were performed for Reaction 4 on all possible bicyclic radicals, regardless of whether or not they originated from the lowest energy aromatic-OH adduct, with the exception of those radicals containing a four-membered oxygen ring. The ring strain energies associated with the formation of such radicals were assumed to be much higher than those of the corresponding five- and six-membered rings. Structural information for selected bicyclic peroxy radicals resulting from the aromatic-OH reaction appear in Table 4.14. Energies of the most stable bicyclic peroxy radical intermediates appear in Table 4.8, under the heading of “O\textsubscript{2} Bridge formation”. In all of the cases studied, the allylically stabilized five-membered bicyclic radicals are the lowest energy bicyclic radicals formed. In fact, all other non-allylically stabilized bicyclic radicals are about 20 kcal higher in energy than the lowest energy allylically stabilized bicyclic radical, and are approximately
Table 4.12: Approximate transition state energies for the reactions of aromatic-OH adducts with O₂, represented in terms of the local structure of the aromatic-OH adduct, (a) at the addition site (Add’n Site), where all tertiary carbons (C₃) are bound to methyl (Me) groups unless otherwise indicated, and (b) in the two positions ortho to the addition site. (c) Transition state bond distance for the C-O bond which is formed during the addition of O₂ to the aromatic-OH adduct.

<table>
<thead>
<tr>
<th>Aromatic</th>
<th>Add’n Site</th>
<th>Ortho Groups</th>
<th>( E_s^0 ) (a.u.)</th>
<th>( E^{360} ) (a.u.)</th>
<th>( r(C-O) ) (Å)</th>
<th>( E^{360} ) (Reactants) (a.u.)</th>
<th>Barrier Ht. (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>p-xylene</td>
<td>C₁</td>
<td>H, OH</td>
<td>-536.9635</td>
<td>-536.7780</td>
<td>1.90</td>
<td>-536.7904</td>
<td>7.8</td>
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<tr>
<td></td>
<td>C₂</td>
<td>H, Me</td>
<td>-536.9669</td>
<td>-536.7814</td>
<td>1.85</td>
<td>-536.7904</td>
<td>5.4</td>
</tr>
<tr>
<td>toluene</td>
<td>C₃</td>
<td>H, OH</td>
<td>-497.6406</td>
<td>-497.4857</td>
<td>1.85</td>
<td>-497.4995</td>
<td>8.7</td>
</tr>
<tr>
<td></td>
<td>C₁</td>
<td>H, OH</td>
<td>-497.6454</td>
<td>-497.4905</td>
<td>1.85</td>
<td>-497.4995</td>
<td>5.6</td>
</tr>
<tr>
<td>m-xylene</td>
<td>C₁</td>
<td>H, OH</td>
<td>-536.9686</td>
<td>-536.7833</td>
<td>1.90</td>
<td>-536.8023</td>
<td>11.9</td>
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<td>m-ethyl-</td>
<td>C₁(ET)</td>
<td>H, OH</td>
<td>-576.2861</td>
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<td>1.80</td>
<td>-576.0779</td>
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<td>C₂</td>
<td>H, H</td>
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<td>-576.0742</td>
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<td>-576.0779</td>
<td>2.3</td>
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</table>
Table 4.13: PM3 equilibrium structures for a representative selection of the most stable reaction products of aromatic-OH adducts with O$_2$ (Reaction 2b). All bond distances are in Å, angles are in degrees.
10 kcal/mol higher in energy than the corresponding peroxy radicals from which they are formed. Thus, the reactions forming the allylically stabilized bicyclics are the only exothermic reactions determined in this step of the mechanisms. The barrier heights for bicyclic ring closure seem to be governed by the attachment sites of the O$_2$ bridge. Representative transition state energies for the bicyclic radical intermediates appear in Table 4.15. A transition state that results in the attachment of an O$_2$ bridge to two tertiary carbons results in a lower energy transition state than that which results in the attachment to one tertiary and one secondary carbon. The energy of the transition state which results in an O$_2$ bridge between two tertiary carbons is approximately 10 kcal/mol higher than that of the parent peroxy radical. A transition state that results in O$_2$ attachment to a secondary carbon, with O$_2$ originating from a tertiary site, results in a transition state energy that is approximately 12 kcal/mol higher than the parent peroxy radical. A transition state that results in O$_2$ attachment to a tertiary carbon, with O$_2$ originating from a secondary carbon site, results in a transition state energy that is approximately 17 kcal/mol higher than the parent peroxy radical.

Energy data for a representative selection of aromatic oxy radicals formed in the reaction of NO with the aromatic peroxy radicals appear in Table 4.16. Representative aromatic peroxy radicals, bicyclic radicals, and aromatic oxy radicals, along with the $\Delta H_{rxn}$ values for Reactions 4 and 5 are presented in Figure 4.7. Not all of the lowest energy aromatic peroxy radicals (which appear in the reaction coordinate diagrams) appear in Figure 4.7. The peroxy radicals chosen for Figure 4.7 are representative cases chosen to give the reader an indication that the two pathways for peroxy radical reaction
Table 4.14: PM3 equilibrium structures for a representative selection of the most stable reaction products in the formation of bicyclic peroxy radicals from peroxy radical intermediates (Reaction 2b). Bond distances are in units of Å, and angles are in degrees.
Table 4.15: Approximate transition states for the addition reactions of bicyclic peroxy radicals with $O_2$, represented in terms of the local structure of the bicyclic peroxy radical (a) at the addition site, and (b) in the two positions ortho to the addition site. “Me” represents a methyl group bond to a tertiary carbon, whereas “CO” represents an attachment site for the $O_2$ bridge. (c) Transition state bond distance for C-O bond which is formed in the addition of $O_2$.

<table>
<thead>
<tr>
<th>Aromatic</th>
<th>Add'n Site</th>
<th>Ortho Groups</th>
<th>$E_s^{0}$(TS) (a.u.)</th>
<th>$E_{2m}^{0}$(TS) (a.u.)</th>
<th>$r$(C-O) (Å)</th>
<th>$E_{2m}$(reactants) (a.u.)</th>
<th>Barrier Ht. (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>toluene</td>
<td>$C_s$</td>
<td>Me, H</td>
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<td>-647.7973</td>
<td>1.9</td>
<td>-647.8262</td>
<td>18.2</td>
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<td></td>
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<td>CO, H</td>
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<td>-647.7975</td>
<td>1.9</td>
<td>-647.8262</td>
<td>18.0</td>
</tr>
<tr>
<td>p-xylene</td>
<td>$C_t$</td>
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</table>
are both possible, and that Reaction 5 cannot be unambiguously excluded from a generalized reaction mechanism. Inclusion of Reaction 5 results in the formation of oxy radicals. These radicals can go on to react according the scheme in Figure 4.8 to yield ring-retained hydroxy carbonyl compounds or decompose by C-C bond scission to ultimately form a di-unsaturated di-carbonyl. These carbonyl compounds can serve to account for part of the carbon unaccounted for in laboratory studies of aromatic-OH systems.

4.3.4 **Bicyclic Peroxy Radicals**

After bicyclic radical formation, oxygen quickly adds to the radical, forming a bicyclic peroxy radical. Transition state energies for a representative selection of bicyclic peroxy radicals appear in Table 4.17. Energy calculations were performed on each of the resulting possible bicyclic peroxy radicals. Note that O₂ addition to rings that result in bicyclic peroxy structures with two radical sites are unlikely to form, and were thus not subjected to single point calculations. The energies determined for the bicyclic peroxy radicals are all similar in magnitude; thus, all bicyclic peroxy structures resulting from the lowest energy bicyclic radicals must be considered in an overall mechanism. Given the steric hindrance problems associated with another cyclization, the bicyclic peroxy radical reacts with NO to form a bicyclic oxy radical and NO₂.
Figure 4.7: $\Delta H_{rxn}$ values for Reactions 4 and 5 for different aromatic systems.
Figure 4.8: Subsequent reaction of oxy radicals formed from Reaction 5 of the generalized mechanism.
Table 4.16: Calculated energies (in a.u.) for the products of the reaction of NO with a representative selection of the most stable peroxy intermediates and for the products of the reaction of O₂ with a representative selection of the most stable bicyclic intermediates.

<table>
<thead>
<tr>
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<th>O' Site</th>
<th>( E^* )</th>
<th>( E^{\text{AM1}} )</th>
<th>OH Site</th>
<th>O₂ Bridge Sites</th>
<th>O₂ Addition Site</th>
<th>( E^* )</th>
<th>( E^{\text{AM1}} )</th>
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<td>5,1</td>
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Table 4.17: Approximate transition states for the formation of the bicyclic peroxy radical, represented in terms of the type of carbon that is bound to an oxygen (a) at the original site of O$_2$ addition to the peroxy radical, and (b) at the bond which is created when the O$_2$ bridge is formed. (c) Transition state bond distance for the bond which is created when the O$_2$ bridge is formed.

<table>
<thead>
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<th>Peroxy</th>
<th>Bicyclic Peroxy</th>
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<th>$E^{298}$(TS)</th>
<th>$r$(C-O)</th>
<th>$E^{298}$(peroxy)</th>
<th>Barrier Ht.</th>
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<tbody>
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<td>Attachment</td>
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<td>(a.u.)</td>
<td>(Å)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>m-xylene</td>
<td>$C_1$</td>
<td>$C_1$</td>
<td>-536.9672</td>
<td>-536.7800</td>
<td>2.20</td>
<td>-536.7986</td>
<td>11.7</td>
</tr>
<tr>
<td>toluene</td>
<td>$C_9$</td>
<td>$C_1$</td>
<td>-497.6318</td>
<td>-497.4764</td>
<td>2.10</td>
<td>-497.5040</td>
<td>17.3</td>
</tr>
<tr>
<td></td>
<td>$C_1$</td>
<td>$C_9$</td>
<td>-497.6396</td>
<td>-497.4842</td>
<td>2.05</td>
<td>-497.5040</td>
<td>12.4</td>
</tr>
<tr>
<td>p-xylene</td>
<td>$C_1$</td>
<td>$C_6$</td>
<td>-536.9625</td>
<td>-536.7775</td>
<td>2.05</td>
<td>-536.7968</td>
<td>12.1</td>
</tr>
</tbody>
</table>
4.3.5 Ring Fragmentation Products

The only path for the resulting bicyclic oxy radicals is fragmentation via favorable $\beta$-scission reactions. These fragmentation reactions yield an acyclic radical and eventually dicarbonyl products (see Table 4.1 - 4.3. Shown in Figure 4.9 is an example of the $\beta$-scission process for a bicyclic oxy radical formed in the photooxidation of toluene. The key issue is the method of fragmentation of the resulting acyclic radical. Calculations of the $\Delta H_{rxn}$ for the decomposition of the acyclic radical were conducted using the NIST Structures and Properties estimation program \cite{25}. This program is based on the group additivity theory of Benson \cite{30}. Based on the calculations, each acyclic radical fragments at the bond between the carbon attached to the oxygen atom and the saturated carbon center rather than the unsaturated carbon center. This point is clearly shown in Figure 4.9. Decomposition path A, illustrating fragmentation between the carbon with the oxygen atom attached to it and the unsaturated carbon center, is the favored path based on group additivity calculations. Decomposition path B is the unlikely method of fragmentation based on group additivity considerations. Shown in Figure 4.10 are the bicyclic oxy radicals and the products that are predicted to form following the $\beta$-scission process analogous to that depicted in Figure 4.9. It is well known that $\alpha$-hydroxy carbonyls subsequently react with $O_2$ to form $HO_2$ and the corresponding dicarbonyl product\cite{29}. Note from Tables 4.1 - 4.3 that a number of the ring fragmentation products theoretically predicted for toluene, $m$-xylene, and $p$-xylene have been identified experimentally.
Figure 4.9: $\beta$-Scission of a bicyclic oxy radical of toluene to give ring fragmentation products. Decomposition path A is the favored path for fragmentation of the acyclic radical.
Figure 4.10: Fragmentation products formed from the bicyclic oxy radicals.
4.4 Conclusion

We have elucidated the intermediates in five atmospheric aromatic-OH photooxidation systems, using theoretically-based calculations. In general we find that the predicted site of initial OH attack is ortho to a substituent group attached to the ring. In addition, we have investigated the NO$_2$ reaction with the aromatic-OH adduct, and have shown that this reaction proceeds via four pathways, forming the nitroaromatic (Reaction 3a), hydroxy nitroaromatic (Reaction 3b), hydroxy aromatic nitrite (Reaction 3c), and the phenolic compound (Reaction 3d). Reaction 3e is not energetically favored, and is thus excluded from a general aromatic-OH mechanism. All peroxy radicals formed from the reaction of the aromatic adduct with O$_2$ have approximately the same energy. Therefore, it is difficult to exclude a particular structure. However, it is clear that if the original aromatic-OH adduct is not favored, then the resulting peroxy radicals will not be formed.

The allylically stabilized five-membered ring bicyclic radicals are, in all cases, approximately 20 kcal/mol more stable than any other bicyclic radical, including those with six-membered rings. Thus, these bicyclic radicals should be included in overall aromatic reaction mechanisms. The aromatic peroxy radical reaction with NO (Reaction 5) is determined to be a plausible reaction to include in aromatic-OH reaction mechanisms. Subsequent reaction of the oxy radicals formed in Reaction 5 can lead to ring-retaining hydroxy carbonyl compounds. These additional ring-retaining compounds can account for part of the remaining carbon unaccounted for in laboratory studies of aromatics. When compared to available product data, the theoretical results
for the intermediates formed and the subsequent reactions for toluene, \textit{m}-xylene, and \textit{p}-xylene are found to be consistent with experimental results.

The theoretical technique has been applied to two important, yet relatively unstudied, aromatics, 1,2,4- trimethylbenzene and \textit{m}-ethyltoluene, to predict their reaction intermediates and stable products.

**Acknowledgments**

This work was supported by U.S. Environmental Protection Agency Center for Airborne Organics (R-819714-01-0), National Science Foundation grant ATM-9307603, the Coordinating Research Council (A-5-1), and the Chevron Corporation. The authors thank J.K. Perry and T. Jungkamp for helpful discussions.
Bibliography


Chapter 5

Photooxidation of Aromatic Compounds: Evidence for an NO$_2$ Effect

Abstract

Atmospheric reaction pathways in the NO$_x$-air photooxidation of aromatics remain one of the most uncertain elements of urban atmospheric chemistry. This work examines the effect of NO$_2$ on the product distribution formed after the initial aromatic-OH reaction of toluene, $m$-xylene, and $p$-xylene. Results of experiments conducted in a 1 m$^3$ photochemical reactor are used to evaluate key branching steps in atmospheric aromatic photooxidation mechanisms.
5.1 Introduction

Aromatics are an important class of compounds in urban air. Notably, in the Southern California Air Quality Study of 1987, aromatics were found to compromise approximately 25% of the carbon in the South Coast Air Basin[1]. Despite the prevalence of these compounds in the atmosphere and their importance in ozone formation, the reaction mechanisms of these species are one of the greatest uncertainties in urban atmospheric chemistry[2].

The major atmospheric sink of aromatics is reaction with the OH radical [3, 4]. Approximately 10% of the hydroxyl radical reaction with an aromatic proceeds via the abstraction route, the remainder by addition. The abstraction route produces ring-retaining products such as aromatic aldehydes and nitrated compounds. The first step in the OH reaction with toluene (as an example) is shown in Figure 5.1. The addition route results in ring fragmentation products such as dicarbonyls and ring-retaining products such as nitroaromatics and phenols. In general, the preferred site of OH addition to the aromatic ring is at positions ortho to the carbon that is substituted with an electron donating group[5].

Molecular oxygen and NO$_2$ can react with the substituted methylcyclohexadienyl (MHCD) radical that results from the OH addition pathway. Present indications are that the O$_2$ pathway proceeds via two routes: addition to form a peroxy radical and H-atom abstraction to form a phenolic type compound. The peroxy radical stabilizes itself by cyclicizing. The cyclicized radical adds oxygen, forming another peroxy radical which subsequently undergoes a series of favorable $\beta$-scission reactions to form ring
Figure 5.1: Initial steps in the toluene-OH reaction.
fragmentation products. Recently, through \textit{ab initio} calculations, Andino et al. [5] identified the energetically favored intermediate species formed in the OH-initiated photooxidation of aromatic compounds, and examined their ring fragmentation pathways. NO$_2$ reaction with the substituted MHCD radical leads to the formation of nitroaromatic compounds. Atkinson and Aschmann [6] have suggested that the NO$_2$ pathway also results in the formation of phenolic type compounds. Several of the products identified in laboratory studies of toluene, \textit{m}-xylene, and \textit{p}-xylene photooxidation are listed in Tables 5.1-5.3; these products are consistent with the theoretical study of Andino et al.[5].

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>0.105 ± 0.019</td>
<td>[15]</td>
</tr>
<tr>
<td></td>
<td>0.058</td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td>0.150 ± 0.040</td>
<td>[17]</td>
</tr>
<tr>
<td></td>
<td>0.080</td>
<td>[18]</td>
</tr>
<tr>
<td>Methyl Glyoxal</td>
<td>0.146 ± 0.006</td>
<td>[15]</td>
</tr>
<tr>
<td></td>
<td>0.077</td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td>0.140 ± 0.04</td>
<td>[17]</td>
</tr>
<tr>
<td></td>
<td>0.075</td>
<td>[18]</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>0.0645 ± 0.0080</td>
<td>[19]</td>
</tr>
<tr>
<td></td>
<td>0.05</td>
<td>[16]</td>
</tr>
<tr>
<td></td>
<td>0.054</td>
<td>[18]</td>
</tr>
<tr>
<td>Benzyl Nitrate</td>
<td>0.0084 ± 0.0017</td>
<td>[19]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>0.010</td>
<td>[16]</td>
</tr>
<tr>
<td>\textit{o}-Cresol</td>
<td>0.204 ± 0.027</td>
<td>[19]</td>
</tr>
<tr>
<td>\textit{m}- and \textit{p}-Cresol</td>
<td>0.048 ± 0.009</td>
<td>[19]</td>
</tr>
<tr>
<td>\textit{m}-Nitrotoluene</td>
<td>0.0135±0.0029 + 1.90±0.25x 10^{-16}[NO$_2$]</td>
<td>[19]</td>
</tr>
<tr>
<td>\textit{o}-Nitrotoluene</td>
<td>0.07x[m-nitrotoluene]</td>
<td>[19]</td>
</tr>
<tr>
<td>\textit{p}-Nitrotoluene</td>
<td>0.35x[m-nitrotoluene]</td>
<td>[19]</td>
</tr>
</tbody>
</table>
Table 5.2: \textit{m}-xylene Photooxidation Products: Molar Yields

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>$0.13 \pm 0.03$</td>
<td>[20]</td>
</tr>
<tr>
<td></td>
<td>0.086</td>
<td>[21]</td>
</tr>
<tr>
<td></td>
<td>0.086 \pm 0.011</td>
<td>[15]</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>$0.42 \pm 0.05$</td>
<td>[20]</td>
</tr>
<tr>
<td></td>
<td>0.375</td>
<td>[21]</td>
</tr>
<tr>
<td></td>
<td>0.319\pm0.009</td>
<td>[15]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>$0.17 \pm 0.02$</td>
<td>[20]</td>
</tr>
<tr>
<td>\textit{m}-Tolualdehyde</td>
<td>0.0331</td>
<td>[22]</td>
</tr>
<tr>
<td></td>
<td>0.12 \pm 0.059</td>
<td>[21]</td>
</tr>
<tr>
<td>3-Methylbenzyl nitrate</td>
<td>0.0061</td>
<td>[22]</td>
</tr>
<tr>
<td>2,4 Dimethylphenol</td>
<td>0.099</td>
<td>[22]</td>
</tr>
<tr>
<td>2,6 Dimethylphenol</td>
<td>0.111</td>
<td>[22]</td>
</tr>
<tr>
<td>Dimethylphenol</td>
<td>$0.178 \pm 0.065$</td>
<td>[21]</td>
</tr>
<tr>
<td>4-Nitro-\textit{m}-xylene</td>
<td>0.0018</td>
<td>[22]</td>
</tr>
<tr>
<td>5-Nitro-\textit{m}-xylene</td>
<td>See Note\textsuperscript{2}</td>
<td>[22]</td>
</tr>
<tr>
<td>Nitro-\textit{m}-xylene</td>
<td>$0.033 \pm 0.025$</td>
<td>[21]</td>
</tr>
</tbody>
</table>

Table 5.3: \textit{p}-xylene Photooxidation Products: Molar Yields

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>$0.24 \pm 0.02$</td>
<td>[20]</td>
</tr>
<tr>
<td></td>
<td>0.225 \pm 0.039</td>
<td>[15]</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>$0.12 \pm 0.02$</td>
<td>[20]</td>
</tr>
<tr>
<td></td>
<td>0.105 \pm 0.034</td>
<td>[15]</td>
</tr>
<tr>
<td>3-Hexene-2,5 Dione</td>
<td>Detected</td>
<td>[20]</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>$0.17 \pm 0.02$</td>
<td>[20]</td>
</tr>
<tr>
<td>\textit{p}-Tolualdehyde</td>
<td>0.0701</td>
<td>[22]</td>
</tr>
<tr>
<td>4-Methylbenzyl nitrate</td>
<td>0.0082</td>
<td>[22]</td>
</tr>
<tr>
<td>2,5 Dimethylphenol</td>
<td>0.188</td>
<td>[22]</td>
</tr>
<tr>
<td>2-Nitro-\textit{p}-xylene</td>
<td>$(0.0120 + 2.8x10^{-17}[NO_2])$</td>
<td>[22]</td>
</tr>
</tbody>
</table>
Recent evidence has suggested that the level of NO$_2$ will significantly affect the yields of aromatic-OH reaction products [6]. While there have been a series of investigations of the photooxidation of toluene, m-xylene, and p-xylene, the only systematic study of effects of NO$_2$ on aromatic systems is that of Atkinson and Aschmann [6], examining the reactions of toluene, o-xylene, and 1,2,3-trimethylbenzene.

Examination of the product yields in Tables 5.1-5.3 shows that, in several cases, discrepancies exist between reported yields. As examples, note the yields of methyl glyoxal from toluene and m-xylene and glyoxal from m-xylene. Even if the error limits on the yields are considered, these discrepancies can be significant, and may be attributed to the neglect of secondary reactions or differences in NO$_2$ concentrations in the experimental systems. When considering the mechanism of aromatic photooxidation, there are several junctures where the NO$_2$ level might affect observed product yields. The most influential reaction involving NO$_2$ is that with the substituted MHCD radical. The branching ratio between the reactions of O$_2$ and NO$_2$ with this radical is of crucial importance to product yields. We present here the results of a systematic laboratory study on the effects of NO$_2$ on selected product yields in toluene, m-xylene, and p-xylene photooxidations. Based on these results, we have developed plausible reaction mechanisms for the photooxidation of these aromatics, and have estimated the branching ratio between the MCHD radical reaction with NO$_2$ and O$_2$. 

5.2 Experimental Procedure

The Caltech Indoor Photochemical Reactor has been described in detail elsewhere\textsuperscript{4}. Briefly, the chamber is a 1 m\textsuperscript{3} reactor composed of 2 mil FEP Teflon. Artificial light (two banks of 24 Sylvania F30T8BL black lamps) is used to initiate the chemistry within the chamber. The lamps are mounted on a cylindrical surface to provide for uniformity in irradiation. The light intensity within the chamber can be adjusted by varying the number of lamps that are used (minimum of 2 lamps, and maximum of 48 lamps). Typical irradiation times for toluene, \textit{m}-xylene, and \textit{p}-xylene were 1-6 min., 1-5 min., and 1-10 min., respectively.

To study the hydroxyl radical-aromatic reactions in the presence of NO\textsubscript{x}, methyl nitrite was introduced to the chamber along with the aromatic of interest via a calibrated glass vacuum manifold. Methyl nitrite generates OH radicals according to the following reactions:

\begin{align*}
\text{CH}_3\text{ONO} + \text{hv} & \rightarrow \text{CH}_3\text{O}^\cdot + \text{NO} \\
\text{CH}_3\text{O}^\cdot + \text{O}_2 & \rightarrow \text{HCHO} + \text{HO}_2^\cdot \\
\text{HO}_2^\cdot + \text{NO} & \rightarrow \cdot\text{OH} + \text{NO}_2
\end{align*}

In order to minimize ozone production, and maximize the conversion of peroxy radicals to alkoxy radicals, NO was added to the chamber. Typical starting mixing ratios were 4-5 ppm aromatic, 1-5 ppm CH\textsubscript{3}ONO, and 1 ppm NO. Toluene, \textit{m}-xylene, and \textit{p}-xylene were obtained from Aldrich with stated purities of 99+\%, 99+\%, and 99+\%. All were used without further

\textsuperscript{4}See Chapter 2.
purification. Nitric oxide was taken from a standard cylinder (generated by Scott-Marin). Methyl nitrite was prepared by the dropwise addition of 50% sulfuric acid to a saturated methanol/sodium nitrite solution. Using high purity N₂, the resulting gas was bubbled through a NaOH solution to neutralize the acid, and then passed through CaCl₂ to dry it. The nitrite was trapped in a bulb which was immersed in a dry ice/ethanol slush bath, purified using the glass vacuum manifold, and stored at liquid N₂ temperatures.

To study aromatic-OH reactions in the absence of NOₓ, ozone-olefin reactions were used to generate OH radicals. The olefin-ozone reaction has been shown to yield significant amounts of hydroxyl radicals [7]. α-Pinene was chosen for the current study because its photooxidation products have been reasonably well characterized, and it produces a high yield of OH radicals[7, 8, 9]. Hydroxyl radical yield from the ozone reaction with α-pinene is 0.85, much higher than other readily available alkenes and monoterpenes. Dark reactions of ozone, α-pinene, and aromatic were carried out to generate OH radicals for NOₓ-free experiments. The ozone was produced by an ozone generator (Ennnett Inc.), and α-pinene was purchased from Aldrich with a stated purity of 99%. It was used without further purification.

The chamber was sampled periodically for analyses by the on-line ozone analyzer (Dasibi 1008) and NOₓ analyzer (Thermo Environmental Model 42). Samples were also collected onto Tenax packed stainless steel tubes for desorption and subsequent analysis by GC-FID. Desorption was accomplished using a Tekmar Model 6000 Aerotrap desorber with an attached cryofocusing unit. The column used in the GC-FID was a J&W Scientific DB-5 column of dimension 30 m x 0.25 mm x 0.25 µm. The GC-FID oven temperature
was held at 50°C for 1 min and increased by 20°C/min. until the final temperature of 275°C was reached. The oven was then held at 275°C for 1 min.

The GC-FID was calibrated for the aromatics by using standard cylinders (generated by Scott-Marrin) of each aromatic. Standard concentrations of the aromatic were introduced into a small (4 liter) teflon bag. Tenax packed tube samples were taken from this bag and analyzed in the same manner as the experimental samples. Calibrations of the products were carried out in a slightly different manner. Liquid samples of the products were vaporized into the chamber along with an aromatic. The calibration amount of aromatic introduced into the chamber was used to calculate the bag volume, and the product concentration was obtained. Response factors were obtained for each product identified by evaporating a known amount of the product, in liquid form, into the chamber along with an aromatic. The aromatic concentration was calibrated, and the value obtained was used to determine the bag volume. This volume was then used to calculate the product concentration. Response factors were subsequently generated from these calculated concentrations.

At the end of each experiment, a sample from the chamber was collected on a 2,4-dinitrophenylhydrazine (DNPH) impregnated C\textsubscript{18} cartridge. The DNPH method was the only method used to analyze for aldehydes in these studies. The samples were eluted using 3 ml of acetonitrile, and analyzed using a Hewlett Packard Model 1090 Series II liquid chromatograph with a diode array detector. A reversed-phase ODS Axxiom LC column of dimension 0.5 \( \mu \)m x 150 cm was used to obtain adequate separation. HPLC grade acetonitrile and water were used as the solvents in a ratio of 55% to 45%,
with a total flowrate of 1 ml/min. Typical observed pressures were between 109 and 112 mbar.

Authentic DNPH derivatized carbonyl standards for the LC were purchased from Radian Corporation for comparison to the experimental samples. Additional standard samples for glyoxal and methyl glyoxal were synthesized in our laboratory according to the method of Druzik et al. [10].

5.3 Effect of NO₂ on Product Yields in Aromatic Photooxidation

The primary products detected and quantified from the OH-addition pathway in the toluene photoxidation system are o-, m-, p-cresol, methyl glyoxal, and glyoxal. Average NO₂ concentrations before correction for the OH-NO₂ reaction were determined using the following equation, incorporating the contribution of methyl nitrite photolysis and NO to NO₂ conversion:

$$[\text{NO}_2] = \frac{1}{n} \sum_{i=1}^{n} ([\text{CH}_3\text{ONO}]_0 (1 - e^{-kt_i}) + \Delta [\text{NO}]_i)$$

where: $k =$ photolysis rate constant for methyl nitrite, as measured experimentally, $\Delta [\text{NO}]_n = [\text{NO}]_0 - [\text{NO}]_t$, and $n =$ number of data points. This expression assumes that the initial NO₂ concentration is zero. Because a NOₓ monitor was used as the method of monitoring the inorganic compound concentrations, the actual amount of NO₂ initially present was not quantifiable. Use of a NOₓ monitor does not allow for the direct measurement of NO₂.
In addition, organic nitrites and nitrates can (and do) effect a response on the \( \text{NO}_2 \) channel. The average concentration of \( \text{NO}_2 \) obtained in the above formula was corrected to account for the OH-\( \text{NO}_2 \) reaction. This correction term was derived using modeling predictions determined with the aid of the mechanisms appearing in Tables 7-9. The mechanisms will be discussed subsequently. For completeness, the OH reactions of products appearing in Table 5.4 were also included in these simulations. Mechanism predictions of the \( \text{NO}_2 \) concentrations resulting from both the inclusion and exclusion of the OH + \( \text{NO}_2 \) reaction for each aromatic were plotted against each other, and regression analyses yielded straight lines with excellent correlation coefficients (\( R^2 > 0.999 \)). The average \( \text{NO}_2 \) concentration determined by taking into account the formation of \( \text{NO}_2 \) from methyl nitrite photolysis and NO to \( \text{NO}_2 \) conversion (denoted as \([\text{NO}_2]_{\text{average}}\)) was then corrected using the following linear relationships for the given aromatics:

Toluene: \([\text{NO}_2]_{\text{corrected}} = (0.8487)([\text{NO}_2]_{\text{average}}) + 0.0157\)

\(m\)-Xylene: \([\text{NO}_2]_{\text{corrected}} = (0.9175)([\text{NO}_2]_{\text{average}}) + 0.0358\)

\(p\)-Xylene: \([\text{NO}_2]_{\text{corrected}} = (0.8717)([\text{NO}_2]_{\text{average}}) + 0.056\)

From these relationships, it is clear that the correction term increases as the concentration of \( \text{NO}_2 \) in the system increases.

The range of \( \text{NO}_2 \) mixing ratios used in the toluene study was 200-1650 ppb. Yields of major species versus average \([\text{NO}_2]\) for toluene are presented in Figures 5.2-5.4. The data points in Figures 5.2-5.4, and subsequent figures represent the measured data from experiments with and without \( \text{NO}_2 \). The
lines in these figures will be discussed subsequently.

The primary OH-addition pathway products that were detected and quantified from the photooxidation of \textit{m}-xylene are glyoxal, methyl glyoxal, nitro-\textit{m}-xylene, 2,4-dimethylphenol, and 2,6-dimethylphenol. The average NO\textsubscript{2} mixing ratios ranged from 200 ppb to 1100 ppb. Yields of major species versus average [NO\textsubscript{2}] for \textit{m}-xylene are presented in Figures 5.5-5.8.

The primary products resulting from the addition pathway of the photooxidation of \textit{p}-xylene that were detected and quantified are glyoxal, methyl glyoxal, and 2,5-dimethylphenol. Average NO\textsubscript{2} mixing ratios in these experiments ranged from 200 ppb to 1050 ppb. Yields of major species versus average [NO\textsubscript{2}] for \textit{p}-xylene are presented in Figures 5.9-5.11.

Because the detection limit of the GC-FID/thermal desorber towards nitroaromatics was not adequate to see these species, only limited data are available for nitroaromatics, in particular, only those resulting from the \textit{m}-xylene system.

Since all products identified are capable of undergoing secondary reactions with hydroxyl radicals, raw product yields must be corrected. Using the method of Atkinson et al. [11], the following formula,

\[
F = \frac{k_1 - k_2}{k_1} \frac{1 - \frac{[\text{Aromatic}]_t}{[\text{Aromatic}]_o}}{\frac{[\text{Aromatic}]_t}{[\text{Aromatic}]_o} - \frac{[\text{Aromatic}]_t}{[\text{Aromatic}]_o}}
\]

(5.1)

where \(k_1\) and \(k_2\) are the rate constants of the following reactions:

Aromatic + OH \(\xrightarrow{1} \) Product

Product + OH \(\xrightarrow{2} \) Secondary Products
Figure 5.2: Toluene Photooxidation- Glyoxal
Figure 5.3: Toluene Photooxidation- Methyl Glyoxal
Figure 5.4: Toluene Photooxidation - Cresols
Figure 5.5: \textit{m}-xylene Photooxidation - Glyoxal
Figure 5.6: $m$-xylene Photooxidation - Methyl Glyoxal
Figure 5.7: m-xylene Photooxidation- Dimethylphenols
Figure 5.8: m-xylene Photooxidation- Nitro-m-xylene
Figure 5.9: p-xylene Photooxidation- Glyoxal
Figure 5.10: $p$-xylene Photooxidation- Methyl Glyoxal
Figure 5.11: \( p \)-xylene Photooxidation - Dimethylphenol
Table 5.4: OH radical rate constants used to correct product yields.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Rate constant @ 298K [3, 4] (cm³/molecule⁻¹s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aromatics</td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>5.96 x 10⁻¹²</td>
</tr>
<tr>
<td>m-xylene</td>
<td>2.36 x 10⁻¹¹</td>
</tr>
<tr>
<td>p-xylene</td>
<td>1.43 x 10⁻¹¹</td>
</tr>
<tr>
<td>Products</td>
<td></td>
</tr>
<tr>
<td>Glyoxal</td>
<td>1.14 x 10⁻¹¹</td>
</tr>
<tr>
<td>Methyl Glyoxal</td>
<td>1.72 x 10⁻¹¹</td>
</tr>
<tr>
<td>o-Cresol</td>
<td>4.2 x 10⁻¹¹</td>
</tr>
<tr>
<td>m-Cresol</td>
<td>6.4 x 10⁻¹¹</td>
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<td>p-Cresol</td>
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<td>2,4-Dimethylphenol</td>
<td>7.15 x 10⁻¹¹</td>
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<tr>
<td>2,6-Dimethylphenol</td>
<td>6.59 x 10⁻¹¹</td>
</tr>
<tr>
<td>2,5-Dimethylphenol</td>
<td>8 x 10⁻¹¹</td>
</tr>
<tr>
<td>Nitro-m-xylene</td>
<td>0.95 x 10⁻¹²5</td>
</tr>
</tbody>
</table>

is used to derive a correction factor. Raw yields are multiplied by this correction factor, $F$ to obtain a corrected yield. Rate constants used in Equation (5.1) for compounds identified in this work are given in Table 5.4. Presented in Table 5.5 are the ranges determined for $F$ for each of the products quantified in the studies presented in this work. It should be noted that the correction factor, $F$, increased with increasing NO₂ concentration, and that uncertainties in $F$ due to uncertainties in the rate constants, $k_1$ and $k_2$, were not taken into account.

For the three aromatics, several general trends can be discerned. As the NO₂ concentration increases, yields of ring fragmentation products and phenolic compounds decrease. In the case of m-xylene, an increase in the nitroaromatic yield is noticed. These observations suggest that the pathways
Table 5.5: Ranges of correction values, $F$, used for each quantified product.

<table>
<thead>
<tr>
<th>Aromatic</th>
<th>Product</th>
<th>$F$ Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toluene</td>
<td>Glyoxal</td>
<td>1.03 - 1.95</td>
</tr>
<tr>
<td></td>
<td>Methyl Glyoxal</td>
<td>1.05 - 2.59</td>
</tr>
<tr>
<td></td>
<td>Cresols</td>
<td>1.12 - 6.16</td>
</tr>
<tr>
<td>$m$-Xylene</td>
<td>Glyoxal</td>
<td>1.01 - 1.13</td>
</tr>
<tr>
<td></td>
<td>Methyl Glyoxal</td>
<td>1.02 - 1.21</td>
</tr>
<tr>
<td></td>
<td>Dimethylphenols</td>
<td>1.06 - 1.94</td>
</tr>
<tr>
<td></td>
<td>Nitro-$m$-Xylene</td>
<td>1.003 - 1.006</td>
</tr>
<tr>
<td>$p$-Xylene</td>
<td>Glyoxal</td>
<td>1.02 - 1.25</td>
</tr>
<tr>
<td></td>
<td>Methyl Glyoxal</td>
<td>1.02 - 1.39</td>
</tr>
<tr>
<td></td>
<td>Dimethylphenols</td>
<td>1.11 - 3.48</td>
</tr>
</tbody>
</table>

Forming ring fragmentation and phenolic compounds are in competition with those forming nitroaromatic compounds. Yields of ring fragmentation products and phenolic compounds formed in the absence of NO$_2$ are lower than the product yield corresponding to the lowest NO$_2$ concentration. This is likely a result of the fact that peroxy radical-radical interactions in NO$_2$-free systems produce additional products as compared to those in peroxy radical-NO reactions.

Previous yield data reported in the literature fall in the high NO$_2$ concentration regime relative to the present experiments. It should be noted that while the present study extends the data concerning the aromatic product yields to lower NO$_2$ concentrations, these concentrations are still, at their lowest value, approximately a factor of two higher than that of NO$_2$ in urban areas. Difficulties were encountered in performing even lower level NO$_2$ experiments with the CH$_3$ONO-NO-aromatic systems because of the difficulty in accurately distinguishing small changes in aromatic concentration.
5.4 Aromatic Photooxidation Mechanisms

Reaction mechanisms (excluding the reactions of products) for toluene, m-xylene and p-xylene were developed to provide a direct comparison between the experimental product data (corrected for losses due to OH reaction) and the mechanisms. The mechanisms were executed using the SAPRC Atmospheric Photochemical Mechanism Preparation Program[12]. A detailed description of the modeling program, its development and implementation, is available [12].

A generalized mechanism for aromatic reactions following OH radical addition is given in Figure 5.12. The major issues faced in preparing detailed mechanisms for each of the aromatics are determination of the following:

1. OH + Aromatic branching ratio (addition versus abstraction).
2. Branching ratio of the MHCD radical reaction with O₂ and NO₂.
3. Branching ratio of the MHCD + O₂ reaction (phenolic compounds vs. peroxy radical formation leading to ring-fragmentation products).
4. Branching ratio of the MHCD + NO₂ reaction.

Each of these issues, except the first, is addressed by analysis of product data and is discussed subsequently. Branching ratios and absolute rate constants for the OH-abstraction versus OH-addition pathway for each of the aromatics studied are available [4].

Mechanisms for the primary reactions of toluene, m-xylene, and p-xylene, are presented in Tables 5.7 - 5.9, and structures for the species used in the mechanisms are presented in Table 5.6. The names and structures used in the p-xylene mechanism are identical to those in the m-xylene mechanism, with
Figure 5.12: Reactions in a generalized mechanism of Aromatic-OH photooxidation. "R" represents either an H atom or a CH₃ group, depending on the aromatic considered.
the exception that the substituent methyl groups are in the para rather than meta position. While the mechanisms listed do not include the secondary reactions of the primary products, in full atmospheric models these additional reactions would normally be included.
Table 5.6: Names and structures of species appearing in the aromatic mechanisms of Tables 5.7, 5.8, and 5.9.

<table>
<thead>
<tr>
<th>Name</th>
<th>Structure</th>
<th>Name</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOL</td>
<td><img src="image1" alt="Structure" /></td>
<td>MXYL</td>
<td><img src="image2" alt="Structure" /></td>
</tr>
<tr>
<td>TOL-</td>
<td><img src="image3" alt="Structure" /></td>
<td>DMHCD</td>
<td><img src="image4" alt="Structure" /></td>
</tr>
<tr>
<td>BENZALD</td>
<td><img src="image5" alt="Structure" /></td>
<td>DMHCD-02</td>
<td><img src="image6" alt="Structure" /></td>
</tr>
<tr>
<td>MHCD</td>
<td><img src="image7" alt="Structure" /></td>
<td>NMXYL</td>
<td><img src="image8" alt="Structure" /></td>
</tr>
<tr>
<td>MHCD-02</td>
<td><img src="image9" alt="Structure" /></td>
<td>DMPHEN</td>
<td><img src="image10" alt="Structure" /></td>
</tr>
<tr>
<td>CRES</td>
<td><img src="image11" alt="Structure" /></td>
<td>DMHCD2-02</td>
<td><img src="image12" alt="Structure" /></td>
</tr>
<tr>
<td>NTOL</td>
<td><img src="image13" alt="Structure" /></td>
<td>MXY-</td>
<td><img src="image14" alt="Structure" /></td>
</tr>
<tr>
<td>BCYLRAD</td>
<td><img src="image15" alt="Structure" /></td>
<td>MXY-02</td>
<td><img src="image16" alt="Structure" /></td>
</tr>
<tr>
<td>MHCD2-02</td>
<td><img src="image17" alt="Structure" /></td>
<td></td>
<td><img src="image18" alt="Structure" /></td>
</tr>
</tbody>
</table>
Primary Reactions of Toluene

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.7: Toluene Mechanism

167
\begin{align*}
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (16) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (15) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (14) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (13) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (12) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (11) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (10) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (9) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (8) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (7) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (6) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (5) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (4) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (3) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (2) \\
\text{HOME} & \rightarrow \text{HOME} + \text{HOME} \quad (1) \\
\end{align*}
17) HONO + HV → OH + NO
18) OH + NO₂ → HNO₃

19) OH + HNO₃ → H₂O + NO₃
20) HNO₃ + HV → OH + NO₂
21) OH + CO → HO₂ + CO₂
22) OH + O₃ → HO₂ + O₂
24) HO₂ + NO₂ → HNO₄

25) HNO₄ + RCON₂ → HO₂ + NO₂
27) HNO₄ + OH → H₂O + NO₂ + O₂
28) HO₂ + O₃ → OH + 2 O₂
29A) HO₂ + HO₂ → HO₂H + O₂
29B) HO₂ + HO₂ + M → HO₂H + O₂
29C) HO₂ + HO₂ + H₂O → HO₂H + O₂ + H₂O
29D) HO₂ + HO₂ + H₂O → HO₂H + O₂ + H₂O
30A) NO₃ + HO₂ → HNO₃ + O₂
30B) NO₃ + HO₂ + O₂ → HNO₃ + O₂
30C) NO₃ + HO₂ + H₂O → HNO₃ + O₂ + H₂O
30A) NO₃ + HO₂ + H₂O → HNO₃ + O₂ + H₂O
31) HO₂H + HV → 2 OH
32) HO₂H + OH → HO₂ + H₂O
33) OH + HO₂ → H₂O + O₂

K₀: 2.514x10⁻⁶² 0.00 -4.600
Kᵢ: 2.202x10⁻⁶⁴ 0.00 -1.500
PHOT. = HONO
FALLOFF F = 0.00, N = 1.000

K₀: 9.337x10⁻⁶² 0.00 -5.200
Kᵢ: 3.523x10⁻⁶⁴ 0.00 -2.300
6.425x10⁻¹⁵
PHOT. = HNO₃

2.384x10⁻¹³
1.584x10⁻¹²
FALLOFF F = 0.00, N = 1.000

K₀: 6.464x10⁻⁶³ 0.00 -5.200
Kᵢ: 6.899x10⁻⁶³ 0.00 -2.400
4.558x10²⁶ x k₂₄
1.293x10⁻¹²
1.091x10⁻¹⁴
2.190x10⁻¹³
1.894x10⁻³³
3.109x10⁻³⁴
2.682x10⁻⁵⁴
SAME K AS 29A
SAME K AS 29B
SAME K AS 29C
SAME K AS 29D
PHOT. = H₂O₂

3.276x10⁻¹²
4.573x10⁻¹¹
Table 5.8: \textit{m}-xylene Mechanism

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction</th>
<th>Rate Constant, 298 K</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>(cm$^3$, molecule, s units)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Experiment</td>
</tr>
<tr>
<td>MN1</td>
<td>CH$_3$ONO + HV $\rightarrow$ CH$_3$O + HCHO</td>
<td>1.9x10$^{-15}$</td>
</tr>
<tr>
<td>MN2</td>
<td>CH$_3$O + O$_2$ $\rightarrow$ HCHO + HO$_2$</td>
<td>3.7x10$^{-12}$</td>
</tr>
<tr>
<td>MN3</td>
<td>HO$_2$ + NO $\rightarrow$ OH + NO$_2$</td>
<td></td>
</tr>
</tbody>
</table>

*Primary Reactions of \textit{m}-xylene*

| XB1 | OH + MXYL $\rightarrow$ DMHCD                                           | 2.3x10$^{-11}$      |
| XB2a| DMHCD + O$_2$ $\rightarrow$ HO$_2$ + DMPHEN                              | 2x10$^{-16}$        |
| XB2b| DMHCD + O$_2$ $\rightarrow$ DMHCD-O$_2$                                 | 1x10$^{-12}$        |
| XB3a| DMHCD + NO$_2$ $\rightarrow$ NMXYL + H$_2$O                             | See text            |
| XB3b| DMHCD + NO$_2$ $\rightarrow$ Products                                   | See text            |
| XB4 | DMHCD-O$_2$ + NO $\rightarrow$ DMHCD-O + NO$_2$                          | 8.9x10$^{-12}$      |
| XB5 | DMHCD-O + O$_2$ $\rightarrow$ PRODUCT1 + HO$_2$                          | 7x10$^{-15}$        |
| XB6 | DMHCD-O + NO$_2$ $\rightarrow$ DMHCD-ONO$_2$                            | 3.8x10$^{-11}$      |
| XB7 | DMHCD-O$_2$ + DMHCD-O$_2$ $\rightarrow$ 2 DMHCD-O + O$_2$               | 5x10$^{-15}$        |
| XB8 | DMHCD-O$_2$ $\rightarrow$ BCYLRAD                                       | 8x10$^{3}$          |
| XB9 | BCYLRAD + O$_2$ $\rightarrow$ DMHC2-O$_2$                               | 1x10$^{-12}$        |
| XB10| DMHC2-O$_2$ + NO $\rightarrow$ NO$_2$ + DMHC2-O                          | 8.9x10$^{-12}$      |
| X11a| DMHC2-O $\rightarrow$ METHGLY + C5H6O2                                 | 9x10$^{2}$          |
| X11b| DMHC2-O $\rightarrow$ GLYOXAL + C6H8O2                                 | 2.45x10$^{2}$       |
| XB12| DMHC2-O + NO$_2$ $\rightarrow$ DMHC2-ONO$_2$                            | 3.8x10$^{-11}$      |
| XB13| DMHC2-O$_2$ + DMHC2-O$_2$ $\rightarrow$ 2 DMHC2-O + O$_2$               | 5x10$^{-15}$        |
| XA1 | OH + MXYL $\rightarrow$ MXY- + H$_2$O                                   | 7.2x10$^{-13}$      |
| XA2 | MXY- + O$_2$ $\rightarrow$ MXY-O$_2$                                    | 1.0x10$^{-12}$      |
XA3)  $\text{MXY-O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{MXY-O}$  \hspace{1cm} 8.01x10^{-12}  
XA4)  $\text{MXY-O}_2 + \text{NO} \rightarrow \text{MXY-ONO}_2$  \hspace{1cm} 0.89x10^{-12}  
XA5)  $\text{MXY-O} + \text{O}_2 \rightarrow \text{HO}_2 + \text{MTOLALD}$  \hspace{1cm} 9.5x10^{-15}  
XA6a)  $\text{MXY-O} + \text{NO}_2 \rightarrow \text{MXY-ONO}_2$  \hspace{1cm} 3.8x10^{-11}  
XA6b)  $\text{MXY-O} + \text{NO} \rightarrow \text{MXY-ONO}$  \hspace{1cm} 3.8x10^{-11}  
XA7)  $\text{MXY-O}_2 + \text{MXY-O}_2 \rightarrow 2 \text{MXY-O} + \text{O}_2$  \hspace{1cm} 2.5x10^{-13}  

*Inorganic Reactions*

Same as in the Toluene mechanism.
### Table 5.9: \( p \)-xylene Mechanism

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction</th>
<th>Rate Constant, 298 K (cm(^3), molecule, s units)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Methyl Nitrite Photolysis</strong></td>
</tr>
<tr>
<td>MN1</td>
<td>( \text{CH}_3\text{ONO} + \text{HV} \rightarrow \text{CH}_3\text{O} + \text{HCHO} )</td>
<td>Experiment 1.9x10(^{-15})</td>
</tr>
<tr>
<td>MN2</td>
<td>( \text{CH}_3\text{O} + \text{O}_2 \rightarrow \text{HCHO} + \text{HO}_2 )</td>
<td>3.7x10(^{-12})</td>
</tr>
<tr>
<td>MN3</td>
<td>( \text{HO}_2 + \text{NO} \rightarrow \text{OH} + \text{NO}_2 )</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>( p )-xylene Mechanism</strong></td>
</tr>
<tr>
<td>XB1</td>
<td>( \text{OH} + \text{PXYL} = \text{DMHCD} )</td>
<td>1.36x10(^{-11})</td>
</tr>
<tr>
<td>XB2</td>
<td>( \text{DMHCD} + \text{O}_2 \rightarrow \text{HO}_2 + \text{DMPHEN} )</td>
<td>2x10(^{-16})</td>
</tr>
<tr>
<td>XB3a</td>
<td>( \text{DMHCD} + \text{NO}_2 \rightarrow \text{NPXYL} + \text{H}_2\text{O} )</td>
<td>4x10(^{-11})</td>
</tr>
<tr>
<td>XB3b</td>
<td>( \text{DMHCD} + \text{NO}_2 \rightarrow \text{DMPHEN} + \text{HONO} )</td>
<td>3x10(^{-11})</td>
</tr>
<tr>
<td>XB4a</td>
<td>( \text{DMHCD} + \text{O}_2 \rightarrow \text{DMHCD-O}_2 )</td>
<td>1x10(^{-12})</td>
</tr>
<tr>
<td>XB4b</td>
<td>( \text{DMHCD-O}_2 \rightarrow \text{DMHCD} + \text{O}_2 )</td>
<td>6x10(^6)</td>
</tr>
<tr>
<td>XB5</td>
<td>( \text{DMHCD-O}_2 + \text{NO} \rightarrow \text{DMHCD-O} + \text{NO}_2 )</td>
<td>8.9x10(^{-12})</td>
</tr>
<tr>
<td>XB6</td>
<td>( \text{DMHCD-O} + \text{O}_2 \rightarrow \text{PRODUCT1} + \text{HO}_2 )</td>
<td>7x10(^{-15})</td>
</tr>
<tr>
<td>XB15</td>
<td>( \text{DMHCD-O} + \text{NO}_2 \rightarrow \text{DMHCD-ONO}_2 )</td>
<td>3.8x10(^{-11})</td>
</tr>
<tr>
<td>XB7</td>
<td>( \text{DMHCD-O}_2 + \text{DMHCD-O}_2 \rightarrow \text{DMHCD-O} + \text{DMHCD-O} )</td>
<td>5x10(^{-15})</td>
</tr>
<tr>
<td>XB8</td>
<td>( \text{DMHCD-O}_2 \rightarrow \text{BCYLRAD} )</td>
<td>8x10(^4)</td>
</tr>
<tr>
<td>XB9</td>
<td>( \text{BCYLRAD} + \text{O}_2 \rightarrow \text{DMHC2-O}_2 )</td>
<td>1x10(^{-12})</td>
</tr>
<tr>
<td>XB10</td>
<td>( \text{DMHC2-O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{DMHC2-O} )</td>
<td>8.9x10(^{-12})</td>
</tr>
<tr>
<td>X11a</td>
<td>( \text{DMHC2-O} \rightarrow \text{METHGLY} + \text{C5H6O}_2 )</td>
<td>9x10(^2)</td>
</tr>
<tr>
<td>X11b</td>
<td>( \text{DMHC2-O} \rightarrow \text{GLYOXAL} + \text{C6H8O}_2 )</td>
<td>2.45x10(^2)</td>
</tr>
<tr>
<td>XB12</td>
<td>( \text{DMHC2-O} + \text{NO}_2 \rightarrow \text{DMHC2-ONO}_2 )</td>
<td>3.8x10(^{-11})</td>
</tr>
<tr>
<td>XB13</td>
<td>( \text{DMHC2-O}_2 + \text{DMHC2-O}_2 \rightarrow 2 \text{DMHC2-O} + \text{O}_2 )</td>
<td>5x10(^{-15})</td>
</tr>
<tr>
<td>XA1</td>
<td>( \text{OH} + \text{PXYL} \rightarrow \text{PXY-} + \text{H}_2\text{O} )</td>
<td>7.2x10(^{-13})</td>
</tr>
</tbody>
</table>
XA2) \( \text{PXY}^- + \text{O}_2 \rightarrow \text{PXY-O}_2 \) 
XA3) \( \text{PXY-O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{PXY-O} \)  
XA4) \( \text{PXY-O}_2 + \text{NO} \rightarrow \text{PXY-ONO}_2 \) 
XA5) \( \text{PXY-O} + \text{O}_2 \rightarrow \text{HO}_2 + \text{PTOLALD} \)  
XA6) \( \text{PXY-O} + \text{NO}_2 \rightarrow \text{PXY-ONO}_2 \) 
XA7) \( \text{PXY-O}_2 + \text{PXY-O}_2 \rightarrow 2 \text{PXY-O} + \text{O}_2 \)

*Inorganic Reactions*

Same as in the Toluene mechanism.
5.4.1 Aromatic-OH Adduct Reaction with O$_2$

The overall rate constant, $k_2$, for O$_2$ reaction with the MHCD radical has been estimated [6, 13]. Direct measurements of the O$_2$ reaction of hydroxycyclohexadienyl and methylhydroxycyclohexadienyl radicals have been made, and estimates of the rate constant are $2 \times 10^{-16}$ and $5 \times 10^{-16}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, respectively [13]. Direct measurements of the dimethylhydroxycyclohexadienyl (DMHCD) radical reaction with O$_2$ have not been conducted, but an indirect estimate of the rate constant of this reaction is $2.3 \times 10^{-16}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ [6].

To determine the overall rate constant for the DMHCD + O$_2$ reaction, product yields predicted by the mechanism for each aromatic were compared to those determined experimentally as the value for $k_2$ was varied between $1 \times 10^{-16}$ - $1 \times 10^{-15}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. It should be noted that the value used for $k_3$, the rate constant for the aromatic-OH adduct reaction with NO$_2$, was $3 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, as will be discussed in the subsequent section. Resulting yield data were plotted versus average NO$_2$ concentration. The branching ratio, $R_1 = \frac{k_{2a}}{k_{2b}}$, was estimated from experimental data as the ratio of the yield of phenolic compounds to the sum of the methyl glyoxal and glyoxal yields. Average branching ratios for the rate constant of reaction 2 for toluene, m-xylene, and p-xylene, determined by averaging the $R_1$ values of all of our experimental data for each aromatic, were 0.78, 0.20, and 0.36, respectively. The lines in Figures 5.5-5.7 and 5.9-5.11 are the yields predicted by the mechanisms using the best fit DMHCD radical + O$_2$ rate constant of approximately $2 \times 10^{-16}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. This value is in agreement with
the estimate of $k_2$ of Atkinson et al. [6] for o-xylene. The rate constant of the MHCD + O$_2$ reaction from toluene photooxidation, determined by best fits to the experimental data, is approximately 1x10$^{-16}$ cm$^3$molecule$^{-1}$s$^{-1}$. This value is somewhat lower than the direct measurements of Knispel et al. [13] of 5x10$^{-16}$ cm$^3$molecule$^{-1}$s$^{-1}$. Figures 5.2- 5.4 show the results of the mechanism predictions for products formed from toluene photooxidation.

5.4.2 Aromatic-OH Adduct Reaction with NO$_2$

The NO$_2$ pathway, reaction 3 in Figure 5.12, was assumed to have an overall rate constant of 3x10$^{-11}$ cm$^3$molecule$^{-1}$s$^{-1}$[4]. If it is assumed that the NO$_2$ pathway proceeds to exclusively form the nitroaromatic compound, then nitroaromatic yields are obtained that are an order of magnitude higher than those measured in the present or in previous studies. This can clearly be seen in Figure 5.13 where the results of the mechanism predicting the exclusive formation of the nitroaromatic ($k_3=3x10^{-11}$ cm$^3$molecule$^{-1}$s$^{-1}$) are compared to that which produces the best fit of the nitro-m-xylene data, with $k_3=3x10^{-12}$ cm$^3$molecule$^{-1}$s$^{-1}$. Because this value has been determined experimentally, we must conclude that there is an additional pathway involving the substituted MHCD radicals and NO$_2$. This conclusion is consistent with the results of the ab initio calculations of Andino et al. [5], and the data of Atkinson et al.[6]. The branching ratio, $\frac{k_{3a}}{k_{total}}$, is determined by fitting the experimental and predicted values of nitro-m-xylene formed in the photooxidation of m-xylene while varying $k_{3a}$. Results indicate a branching ratio, $\frac{k_{3a}}{k_{total}}$ of approximately 0.1, consistent with the data of Atkinson et al.
Several pathways have been suggested for the reaction of the aromatic-OH adduct with NO$_2$ [5, 6]. The recent theoretical study of Andino et al. [5] has narrowed the possible pathways down to the four that are represented in Figure 5.12 as reactions 3a-d, based on calculations using toluene as an example. Possible pathways for the aromatic-OH adduct-NO$_2$ reaction were evaluated based on barrier heights and overall $\Delta H_{rxn}$ [5]. The $\Delta H_{rxn}$ for the first steps of pathways 3a-3c, and overall pathways 3d and 3e for toluene are -26.3, -26.3, -24.5, -52.8, and +38.3 kcal/mol, respectively. The barrier heights for pathways 3a,b (first step), 3c (first step), and 3d are 5, 1, 2 kcal/mol, respectively. Note that Reactions 3c and 3e are assumed to have the same transition state. Based on this information for the initial steps of Reactions 3a-3c, and the overall Reactions 3d and 3e, the only pathway that can be unambiguously excluded is Reaction 3e, because of its endothermic nature. The decreasing yield of phenolic compounds observed in the present experimental study as average [NO$_2$] increases is not consistent with a significant pathway for phenolic compound formation from aromatic-OH adduct reaction with NO$_2$, namely, reaction 3d. The study of Atkinson et al. [6], which examined the effect of NO$_2$ on the phenolic compound yield only from the toluene reaction, suggest a slight increase in phenolic compound yield as [NO$_2$] increases. The hydroxy nitroaromatics are products of phenolic compound reaction with OH radicals, thus, it is impossible to determine the contribution of hydroxy nitroaromatics resulting from Reaction 3 by experiment alone.
Figure 5.13: Nitro-m-xylene formation simulated based on two assumed values of the rate constant $k_{3a}$.
5.4.3 Ring Fragmentation Products

Andino et al. [5] showed that each bicyclic oxy radical (generalized structure shown in Reaction 8 of Figure 5.12) isomer for toluene, m- and p-xylene leads to the formation of either glyoxal and a larger ring fragmentation product, or methyl glyoxal and another large ring fragmentation product. The relative formation yields of each of these bicyclic oxy radical isomers is not known, but since each isomer is believed to lead either to glyoxal or methyl glyoxal, a rate constant ratio can be determined for the decomposition step from the glyoxal and methyl glyoxal yields. An estimate of the decomposition rate constant for the bicyclic oxy radicals (Reaction 8 of Figure 5.12) leading to the formation of methyl glyoxal is made by analogy with alkoxy radicals that decompose to form a secondary alkyl radical and a carbonyl compound[14]. On this basis, the rate constant for \( k_{8a} = 8.8 \times 10^3 \) s\(^{-1}\). The ratio, \( R_2 = k_{8a} / k_{8b} \), is assumed to be equal to the ratio, \([\text{Methyl Glyoxal}]/[\text{Glyoxal}]\). The \( R_2 \) values for toluene, m-xylene, and p-xylene are determined from the data as 0.68, 3.72, and 0.45, respectively, and the resulting rate constants for each step appear in the mechanisms. These values for \( R_2 \) are in excellent agreement with those calculated from literature data [4].

5.5 Implications for Atmospheric Aromatic Photooxidation Mechanisms

The present work serves to support the recent work of Atkinson et al.[6] and Knispel et al.[13] concerning the kinetics of the reaction of the aromatic-OH
adduct with O₂. In addition, we have shown a clear effect of NO₂ on the products formed in the addition pathway of aromatic reactions. However, if the relative atmospheric concentrations of NO₂ and O₂ are considered, it is clear that the aromatic-OH adduct reaction with NO₂ is negligible under atmospheric conditions. Under the typically high NO₂ conditions of laboratory chamber studies, the reactions of NO₂ with the aromatic-OH adduct are, however, significant. Nitroaromatic formation only accounts for approximately 10% of the aromatic-OH reaction with NO₂. This point is reflected in Tables 7, 8, and 9 for each of the aromatics studied in this work.

Acknowledgements

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Bibliography


Chapter 6

Photooxidation of m-Ethyltoluene and 1,2,4 Trimethylbenzene: A Product Study
Abstract

The OH-initiated photooxidation of 1,2,4-trimethylbenzene and m-ethyltoluene have been investigated in the Caltech Indoor Photochemical Reactor. These studies are the first studies of the products resulting from m-ethyltoluene photooxidation, and are the only studies reporting ring retaining products for the 1,2,4-trimethylbenzene system. Products identified from the m-ethyltoluene system include acetaldehyde, glyoxal, methyl glyoxal, 2-ethyl-4-methylphenol, 4-ethyl-2-methylphenol, 1-(3-methylphenol) ethanone, 3-methyl benzaldehyde, and ethylbenzaldehyde. Products identified from 1,2,4-trimethylbenzene photooxidation include acetaldehyde, glyoxal, methyl glyoxal, 2,4- dimethylbenzaldehyde, 3,4- dimethylbenzaldehyde, 2,5- dimethylbenzaldehyde, 2,3,6- trimethylphenol, and 2,4,5-trimethylphenol.
6.1 Introduction

Aromatics are an important class of compounds in urban air. Notably, in the Southern California Air Quality Study of 1987, aromatics were found to compromise approximately 25% of the carbon in the south coast air basin.[1] The aromatics that are focused on in this study, m-ethyltoluene and 1,2,4-trimethylbenzene, were found to be in the top 25 of the most abundant species (based on carbon) during the summer and fall episodes, respectively, of the SCAQS study. Despite the prevalence of these compounds in the atmosphere and their importance in ozone formation, the reaction mechanisms of these species are one of the greatest uncertainties in atmospheric chemical models[2]. Aromatics are capable of reacting with OH radicals, O₃ and NO₃. However, the major atmospheric sink of these species is reaction with the OH radical. The OH rate constants for m-ethyltoluene and 1,2,4-trimethylbenzene are 1.92x10⁻¹¹ and 3.25x10⁻¹¹ cm³molecule⁻¹s⁻¹, respectively [4].

Approximately 10% of the hydroxyl radical reaction with an aromatic proceeds via the abstraction route, the remainder by addition[4]. The abstraction route produces ring-retaining products such as aromatic aldehydes and nitrated compounds. The first step in the abstraction channel of toluene (as an example) is shown in Figure 6.1. The addition route results in ring fragmentation products such as dicarbonyls and ring-retaining products such as nitroaromatics and phenols. The results of two previous studies identifying some of the ring fragmentation products resulting from the photooxidation of 1,2,4-trimethylbenzene are presented in Table 6.1. In general, addition to
the aromatic ring occurs at positions meta to the carbon that is substituted with an electron donating group[5]. The first step in the OH addition channel of toluene, as an example, is shown in Figure 6.1.

Molecular oxygen and NO2 can react with the resulting substituted methylhydroxycyclohexadienyl (MHCD) radical. Present indications are that the O2 pathway proceeds via two routes: addition to form a peroxy radical and abstraction to form a phenolic type compound. The peroxy radical stabilizes itself by cyclicizing. The cyclicized radical adds oxygen, forming another peroxy radical which subsequently undergoes a series of favorable β scission reactions to form ring fragmentation products. Recently, through ab initio calculations, Andino et al. [5] identified the favored intermediate species formed in the OH-initiated photooxidation of several aromatic comounds, including m-ethyltoluene and 1,2,4-trimethylbenzene, and examined their ring fragmentation pathways. The experimental results presented in this paper are directly compared to those predicted by the ab initio calculations of Andino et al. [5] for the addition pathways of aromatics, and to proposed mechanisms, based on current knowledge [4], for the abstraction pathways.

Table 6.1: 1,2,4 Trimethylbenzene Photooxidation Products: Molar Yields

<table>
<thead>
<tr>
<th>Compound</th>
<th>Yield</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formaldehyde</td>
<td>0.18 ± 0.01</td>
<td>[6]</td>
</tr>
<tr>
<td>Glyoxal</td>
<td>0.078 ± 0.005</td>
<td>[6]</td>
</tr>
<tr>
<td></td>
<td>0.048 ± 0.005</td>
<td>[7]</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>0.37 ± 0.01</td>
<td>[6]</td>
</tr>
<tr>
<td></td>
<td>0.357 ± 0.017</td>
<td>[7]</td>
</tr>
<tr>
<td>Biacetyl</td>
<td>0.11 ± 0.01</td>
<td>[6]</td>
</tr>
<tr>
<td>3-Hexene-2,5 dione</td>
<td>Detected</td>
<td>[6]</td>
</tr>
</tbody>
</table>
Figure 6.1: Initial reactions of the OH-initated reaction of toluene.
6.2 Experimental

The Caltech Indoor Photochemical Reactor has been described in detail elsewhere. Briefly, the chamber is a 1 m$^3$ reactor composed of 2 mil FEP Teflon. Artificial light (two banks of 24 Sylvania F30T8BL black lamps) is used to initiate the chemistry within the chamber. The lamps are mounted on a cylindrical surface to provide for uniformity in irradiation. The light intensity within the chamber can be adjusted by varying the number of lamps that are used (minimum of 2 lamps, and maximum of 48 lamps). Typical irradiation times for 1,2,4-trimethylbenzene and $m$-ethyltoluene were 1-3 min. and 0.5-4 min., respectively.

To study the hydroxyl radical-aromatic reactions in the presence of NOx, methyl nitrite was introduced to the chamber along with the aromatic of interest via a calibrated glass vacuum manifold. Methyl nitrite generates OH radicals according to the following reactions:

\[
\text{CH}_3\text{ONO} + h\nu \rightarrow \text{CH}_3\text{O}^- + \text{NO}
\]
\[
\text{CH}_3\text{O}^- + \text{O}_2 \rightarrow \text{HCHO} + \text{HO}_2
\]
\[
\text{HO}_2 + \text{NO} \rightarrow \text{OH}^- + \text{NO}_2
\]

In order to minimize ozone production, and maximize the conversion of peroxy radicals to alkoxy radicals, NO was added to the chamber. Typical starting mixing ratios were 4-5 ppm aromatic, 1-5 ppm CH$_3$ONO, and 1 ppm NO. 1,2,4-Trimethylbenzene and $m$-ethyltoluene were obtained from Aldrich with stated purities of 99+%, 99+. All were used without further purification.

\footnote{See Chapter 2.}
Nitric oxide was taken from a standard cylinder (generated by Scott-Marin). Methyl nitrite was prepared by the dropwise addition of 50% sulfuric acid to a saturated methanol/sodium nitrite solution [8]. Using high purity N₂, the resulting gas was bubbled through a NaOH solution to neutralize the acid, and then passed through CaCl₂ to dry it. The nitrite was trapped in a bulb which was immersed in a dry ice/ethanol slush bath, purified using a vacuum manifold, and stored at liquid N₂ temperatures.

To study the aromatic-OH reactions in the absence of NOₓ, ozone- olefin reactions were used to generate OH radicals. The olefin-ozone reaction has been shown to yield significant amounts of hydroxyl radicals.[9] α-Pinene was chosen because its photooxidation products have been reasonably well characterized, and it produces a high yield of OH radicals. [9] [10] [11]. Hydroxyl radical yield from the ozone reaction is 0.85, much higher than other readily available alkenes and monoterpenes. Thus, dark reactions of ozone, α-pinene, and aromatic were conducted. The ozone was produced by an ozone generator (Enmett Inc.), and α-pinene was purchased from Aldrich with a stated purity of 99%. It was used without further purification.

The chamber was sampled periodically for analyses by the on-line ozone analyzer (Dasibi 1008) and NOₓ analyzer (Thermo Environmental Model 42). Samples were also collected onto Tenax packed stainless steel tubes for desorption and subsequent analysis by GC-FID. Desorption was accomplished using a Tekmar Model 6000 Aerotrap desorber with an attached cryofocusing unit. The column used in the GC-FID was a J&W Scientific DB-5 column of dimension 30 m x 0.25 mm x 0.25 µm. The GC-FID oven temperature was held at 50° C for 1 min. and increased by 20° C/min. until a final tempera-
ture of 275°C. The oven was then held at 275°C for 1 min. The GC-FID was calibrated for the aromatics by using standard cylinders (generated by Scott-Marrin) of each aromatic. Standard concentrations of the aromatic were introduced into a small (4 liter) teflon bag. Tube samples were taken from this bag, and analyzed in the same manner as the experimental samples.

At the end of each experiment, a sample from the chamber was collected on a 2,4-dinitrophenylhydrazine (DNPH) impregnated C18 cartridge. The samples were eluted using 3 ml of acetonitrile, and analyzed using a Hewlett Packard Model 1090 Series II liquid chromatograph, with a diode array detector. A reversed phase Axxiom LC column of dimension 0.5 μm x 150 cm was used to obtain adequate separation. HPLC grade acetonitrile and water were used as the solvents in a ratio of 55% to 45%, with a total flowrate of 1 ml/min. Typical observed pressures were between 109 and 112 mbar.

Authentic DNPH derivatized carbonyl standards for the LC were purchased from Radian Corporation for comparison to the experimental samples. Standards for DNPH-glyoxal and DNPH-methyl glyoxal were synthesized in our laboratory according to the method of Druzik et al. [12].

Additional experiments were conducted to positively identify additional products formed during the photoxidation of m-ethyltoluene and 1,2,4 trimethylbenzene from mass spectral data. Experiments were conducted in the same manner as indicated above. However, the tenax tube samples collected were desorbed into a Hewlett Packard GCD (Model G1800A). The column used was a HP5, of dimensions 30 m x 0.25 mm x 0.25 μm.
6.3 1,2,4-Trimethylbenzene Photooxidation

6.3.1 Ring-Retained Products

Each of the three substituent methyl groups of 1,2,4-trimethylbenzene are possible sites for OH abstraction of an H atom. The resulting radicals are expected to react with O₂ to form the corresponding ring-retained aromatic aldehydes[4]. Figure 6.2 details the abstraction channel for 1,2,4-trimethylbenzene. The LC chromatograms did not show any evidence for high molecular weight carbonyls. However, the GCD analyses, which were found to be more sensitive to the detection of products than the DNPH technique or the GC-FID analyses, showed positive evidence for the presence of 2,4-dimethylbenzaldehyde, 3,4-dimethylbenzaldehyde, and 2,5-dimethylbenzaldehyde. Shown in Figure 6.3 is a selected portion of a typical GCD chromatogram for 1,2,4-trimethylbenzene photooxidation. The mass spectral fingerprints of the detected dimethylbenzaldehydes were compared to the spectral library of the GCD (Wiley) for positive identification.

The results of ab initio calculations performed by Andino et al. [5] indicate that the preferred sites for OH addition to the ring of 1,2,4 trimethylbenzene are the 3- and 5-positions. Shown in Figure 6.4 are all of the possible reaction pathways for the trimethylhydroxycyclohexadienyl radical reaction with O₂ to form the trimethylphenols. Figure 6.3 shows positive evidence for the formation of only two phenolic compounds, 2,3,6-trimethylphenol and 2,4,5-trimethylphenol, consistent with the predictions based on theory of Andino et al. [5].
Figure 6.2: Abstraction pathway for 1,2,4-trimethylbenzene. Depicted are all of the possible radicals formed from the abstraction pathway of 1,2,4-trimethylbenzene and their reactions with O$_2$ to form the corresponding aromatic aldehydes.
Figure 6.3: GCD chromatogram of products formed from the photooxidation of 1,2,4-trimethylbenzene. Depicted are the benzaldehydes and the trimethylphenols.
Figure 6.4: Pathways for the reaction of isomers of the trimethylhydroxycyclohexadienyl radical with O₂. Depicted are all of the possible trimethylphenols formed from each path.
6.3.2 Ring-Fragmentation Products and the NO$_2$ Effect

LC data show positive evidence for the formation of acetaldehyde, glyoxal, and methyl glyoxal. Methyl glyoxal and glyoxal have previously been identified as products of 1,2,4-trimethylbenzene photooxidation (see Table 6.1 and references within). Both products are ring fragmentation products of the bicyclic peroxy radicals formed after initial OH addition to the ring [5]. Figure 6.5 shows the mechanisms of an initial 1,2,4-trimethylbenzene-OH adduct leading to the formation of methyl glyoxal. An analogous path exists for the formation of glyoxal. A path does not exist for the primary formation of acetaldehyde from 1,2,4-trimethylbenzene photooxidation, thus, it must be a result of secondary reaction.

The yields of glyoxal, methyl glyoxal, and acetaldehyde were determined by comparison with authentic DNPH-derivatized samples. Product yields were corrected for secondary reactions with OH according to the method of Atkinson et al. [13], using the following equation:

$$ F = \frac{k_1}{k_1} - \frac{k_2}{k_1} \frac{1 - \frac{[Aromatic]_i}{[Aromatic]_o}}{\frac{[Aromatic]_i}{[Aromatic]_o} - \frac{[Aromatic]_i}{[Aromatic]_o}} $$

(6.1)

where $k_1$ and $k_2$ are the rate constants of the following reactions:

1. Aromatic + OH → Product
2. Product + OH → Secondary Products.

Corrected yields were determined by multiplying the raw yields by the factor,
Figure 6.5: Depicted is the proposed mechanism for the formation of ring-fragmentation products from the photooxidation of 1,2,4-trimethylbenzene.
F. The rate constants for the ring fragmentation products, acetaldehyde, glyoxal, and methyl glyoxal are 1.58x10^{-11} [4], 1.14x10^{-11} [14], and 1.72x10^{-11} [14], respectively, with units of cm^3 molecule^{-1} s^{-1}.

Average NO_2 concentrations were determined using the following equation, incorporating the contribution of methyl nitrite photolysis and NO to NO_2 conversion to the total NO_2 concentration in the system:

\[
[\text{NO}_2] = \frac{1}{n} \sum_n ([\text{CH}_3\text{ONO}]_0 (1 - e^{-kn}) + \Delta [\text{NO}]_n)
\]

where: \( k \) = photolysis rate constant for methyl nitrite, as measured experimentally, \( \Delta [\text{NO}]_n = [\text{NO}]_0 - [\text{NO}]_t \), and \( n \) = number of data points. Average NO_2 concentrations for 1,2,4-trimethylbenzene ranged from 200 to 800 ppb. Plots of yields of the ring-fragmentation products versus average [NO_2] formed in 1,2,4-trimethylbenzene photooxidation are presented in Figures 6.6-6.8. Note that the points at [NO_2] = 0 are from the ozone-\( \alpha \)-pinene-1,2,4-trimethylbenzene experiments. From Figures 6.6-6.8, a decrease in the yield of ring-fragmentation products is noticed as the average concentration of NO_2 increases, indicating the existence of a competing pathway for NO_2 reaction.
Figure 6.6: 1,2,4 Trimethylbenzene Photooxidation Results: Methyl Glyoxal
Figure 6.7: 1,2,4 Trimethylbenzene Photooxidation Results: Glyoxal
Figure 6.8: 1,2,4 Trimethylbenzene Photooxidation Results: CH$_3$CHO
6.4  \textit{m-}Ethyltoluene Photooxidation

6.4.1  Ring-Retained Products

As mentioned previously, the studies presented here are the first product studies of the photooxidation of \textit{m-}ethyltoluene. Abstraction of a hydrogen from \textit{m-}ethyltoluene can occur at a variety of sites. The resulting radicals can react with $O_2$ to eventually form aromatic aldehydes. These reactions are depicted in Figure 6.9. As with the case of 1,2,4-trimethylbenzene, high molecular weight carbonyls were not detected in the LC chromatograms. However, GCD results indicate the formation of 1-(3-methylphenyl)-ethanone, ethylbenzaldehyde, and 3-tolualdehyde from comparisons with library spectra. As indicated in Figure 6.9, 3-tolualdehyde formation may be the result of two different reaction pathways for radical (c).

The results of \textit{ab initio} calculations performed by Andino et al. [5] indicate that the preferred sites for OH addition to the ring of \textit{m-}ethyltoluene are the 2-, 4- and 6-positions. The reactions forming the ethyl-methylphenols from \textit{m-}ethyltoluene are analogous to those depicted in Figure 6.4 for 1,2,4-trimethylbenzene photooxidation. Comparisons of mass spectra to library spectra indicated the presence of 2-ethyl-4-methylphenol, 4-ethyl-2-methylphenol, and 2-ethyl-6-methylphenol from OH addition to the 4-, 6-, and 2-positions of \textit{m-}ethyltoluene, respectively. A section of a typical GCD total ion chromatogram depicting the reaction products for the abstraction and addition pathways for \textit{m-}ethyltoluene appears in Figure 6.10.
Figure 6.9: Depicted are the radicals resulting from the OH abstraction pathways for $m$-ethyltoluene and the subsequent formation of aromatic aldehydes.
Figure 6.10: GCD chromatogram of products formed from the photooxidation of $m$-ethyltoluene. Depicted are the ring-retained carbonyls and the phenolic compounds.
6.4.2 Ring-Fragmentation Products and the NO$_2$ Effect

LC data show positive evidence for the formation of acetaldehyde, glyoxal, and methyl glyoxal. Both methyl glyoxal and glyoxal are formed as a result of ring fragmentation of the bicyclic peroxy radicals formed after initial OH addition to the $m$-ethyltoluene [5]. The pathways are analogous to those depicted in Figure 6.5 for 1,2,4-trimethylbenzene. Acetaldehyde is believed to be formed as a secondary product from one of the C$_7$ ring fragmentation products, or from ethyl glyoxal, another possible ring fragmentation product predicted by Andino et al. [5]. The yields of glyoxal, methyl glyoxal, and acetaldehyde were determined by comparison with authentic DNPH-derivatized samples. Product yields were corrected for secondary reactions with OH according to the method of Atkinson et al. [13], using Equation 6.1.

Average NO$_2$ concentrations were determined using the method presented in the 1,2,4-trimethylbenzene section, and ranged from 250 to 650 ppb. Plots of ring-fragmentation product yields resulting from $m$-ethyltoluene photooxidation versus average [NO$_2$] are presented in Figures 6.11-6.13. The plots for glyoxal and acetaldehyde indicate that there is no clear effect of NO$_2$ on product formation within the range of NO$_2$ concentrations considered in this work. However, the methyl glyoxal data show a slight decrease in yield with increasing average [NO$_2$]. These results, when compared with the results from the 1,2,4-trimethylbenzene system and the results for toluene, $m$- and $p$-xylene from Andino et al. [5] may indicate that as the order of substitution
at a particular site increases (i.e. in going from a methyl substituted carbon to an ethyl substituted carbon), the effect of NO$_2$ on ring-fragmentation product yields decreases. This suggestion is plausible given that in the case of methyl substituted aromatics (such as 1,2,4-trimethylbenzene), only two possible pathways for ring fragmentation exist, one forming glyoxal (and a C$_7$ carbonyl), and the other forming methyl glyoxal (and a C$_6$ carbonyl). m-Ethyltoluene, on the other hand, has three different possible pathways: formation of glyoxal and a C$_7$ carbonyl, methyl glyoxal and a C$_6$ carbonyl, and ethyl glyoxal and a C$_5$ carbonyl. Thus, the effect of NO$_2$ on any one possible pathway would be mitigated by the existence of several different pathways.

6.5 Conclusions

1,2,4-Trimethylbenzene and m-ethyltoluene are important aromatics which have been detected in urban air. As such, their photooxidation products are important. This study provides the first information on the ring-retained and ring-fragmented products observed during the photooxidation of m-ethyltoluene. In addition, the first evidence for ring-retained products from the 1,2,4-trimethylbenzene system are reported. The yields of ring fragmentation products identified in the 1,2,4-trimethylbenzene system are in excellent agreement with those of previous studies. These product data serve to increase the available information for the reactions pathways of substituted aromatics.

The effect of NO$_2$ on the ring-fragmentation products of 1,2,4- trimethyl-
Figure 6.11: m-Ethyltoluene Photooxidation Results: CH₃CHO
Figure 6.12: $m$-Ethyltoluene Photooxidation Results: Glyoxal
Figure 6.13: \textit{m}-Ethyltoluene Photooxidation Results: Methyl Glyoxal
benzene and \textit{m}-ethyltoluene was also examined. The yields of the ring-
fragmentation products formed from the 1,2,4-trimethylbenzene system de-
creased with increasing average NO\textsubscript{2} concentration. However, only the yield
of methyl glyoxal from \textit{m}-ethyltoluene photooxidation decreased with in-
creasing NO\textsubscript{2}, and the decrease was almost negligible.

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Bibliography


Chapter 7

Conclusions
A new indoor photochemical reactor was designed and constructed to study detailed gas-phase kinetics and mechanisms. This system was used to study the photooxidation of branched alkanes and aromatic compounds, the results of which have been presented in this thesis. The development of an indoor system for detailed gas-phase studies serves to complement the outdoor system which has traditionally been available at Caltech for studies of gas to particle conversions. The development of this new and smaller indoor system launches the Air Pollution/Aerosols group at Caltech into a wide variety of new research areas involving gas-phase atmospheric chemistry.

Studies of the branched alkanes, 2,2,4-trimethylpentane and 2,2,5-trimethylhexane were conducted, and the results presented in Chapter 3. The first determination of the rate constant for 2,2,5-trimethylhexane was made, and the result is in excellent agreement with the value predicted from the structure-reactivity relationship[1, 2]. The product studies were aimed at elucidating the gas-phase mechanism of 2,2,4-trimethylpentane and 2,2,5-trimethylhexane in particular, and providing additional information on the reactions of alkoxy radicals in general. Experimentally observed product yields were compared to those predicted from mechanisms developed for each alkane studied in this work. The results of these comparisons indicated that the 1,5-H atom isomerization reaction pathway is a significant atmospheric sink for many of the large alkoxy radicals formed in the two alkane systems. Rate constant estimates were made for the 1,4 H-atom isomerization pathways based on the reactions of 2,2,4-trimethylpentane, and are consistent with previous estimates based on data from straight chain alkanes. These studies serve to expand the available database for the reactions of large alkanes, and indicate
that currently available kinetic data for alkanes provide reasonable fits to experimental results.

This thesis also focused on the photooxidation of aromatic hydrocarbons. Theoretical and experimental studies were carried out on the OH-initiated photooxidation of toluene, m-xylene, p-xylene, m-ethyltoluene, and 1,2,4-trimethylbenzene. The theoretical studies presented in Chapter 4 provide the first, unambiguous determination of the intermediates formed in aromatic-OH systems. *Ab initio* calculations were performed on proposed reaction intermediates, and the lowest energy structures were identified. Where possible, predictions based on theory were compared to available experimental data. Particular reactions were examined in detail to determine the most likely paths. In particular, the aromatic-OH adduct reaction with NO₂ was examined. Based on the results, three pathways for this reaction were found to be likely. Previous indications were that the aromatic-OH adduct reaction proceeded by one pathway, forming only the nitroaromatic[3]. In addition, the reactions of the aromatic peroxy radical were investigated. Previous indications were that the peroxy radical preferentially cyclicized rather than reacting with NO to form the oxy radical. Theoretical results indicated that both of these reactions should be included in overall mechanisms for aromatic-OH reaction. Thus, Chapter 4 elucidated the structures of the intermediates formed, and suggested additional reaction pathways.

Experimental studies on toluene, m-, and p-xylene presented in Chapter 5 focused on the effect of NO₂ on the products formed during the OH-initiated photooxidation of the aromatics. Results showed a positive indication for an effect of NO₂ on products in all three systems. Full mechanisms for the
aromatics were developed and tested against experimental data. While a NO2
effect was noticed in the experimental systems, and predicted results show
the same phenomenon, rate constant fits indicate that, under atmospheric
conditions, the reaction of the aromatic-OH adduct with NO2 is insignificant
as compared to reaction with O2. The mechanisms developed also support
the data presented in Chapter 4 concerning the NO2 reactions of aromatics.
Based on comparisons of observed and predicted data, additional pathways
for the NO2 reaction with the aromatic-OH adduct were necessary in order to
explain the observed nitroaromatic yields. The results of this chapter extend
the available information regarding the effect of NO2 on product yield, and
provide an updated and detailed mechanism for aromatic photooxidation.

The gas-phase photooxidations of m-ethyltoluene and 1,2,4-trimethyl-
benzene were examined in Chapter 6. These studies represent the first prod-
uct study for the m-ethyltoluene system, and only the second study for the
1,2,4-trimethylbenzene system. Both studies provide the first identifications
of ring-retaining products formed, and are in excellent agreement with those
predicted from the theoretical study of Chapter 4. Ring fragmentation prod-
ucts identified in the 1,2,4-trimethylbenzene system in these studies are con-
sistent with those found in previous studies. The information obtained from
these studies serves to further validate the theoretical predictions of Chapter
4, and to extend the available information on aromatic-OH photooxidation
mechanisms. Additional work is, however, required to fully determine the
effects of NO2 on ring-retained compounds.

Although this thesis provides insight into the photooxidation of large
alkanes (alkoxy radical reactions, in particular) and aromatics, there is still
much information to be gained in both cases. Alkoxy radical isomerizations have been found to be important for branched and straight chain alkanes. However, alkoxy-type radicals formed from other classes of compounds have yet to be investigated, and should be focused on in the future. While much information has been gained about aromatic-\(\text{OH}\) photooxidation from the data of Chapter 4 in particular, additional experimental evidence is needed in order to confirm the mechanism of photooxidation. Techniques that are capable of directly monitoring the reactions of the short-lived intermediates should be extremely helpful in confirming the aromatic reaction mechanism and identifying additional products of photooxidation. These new analytical techniques are currently being developed within the analytical chemistry field, and should prove to be quite useful to the atmospheric chemistry field.
Bibliography


Appendix A

Calculation of the Photolysis Rate Constant, $k_1$
A.1 Reactions of NO$_2$

(1) NO$_2$ + h$\nu$ → O + NO
(2) O + O$_2$ + M → O$_3$ + M
(3) O$_3$ + NO → O$_2$ + NO$_2$
(4) O + NO$_2$ → O$_2$ + NO
(5) O + NO$_2$ + M → NO$_3$ + M
(6) NO$_3$ + NO → NO$_2$ + NO$_2$
(7) O + NO + M → NO$_2$ + M

A.2 Calculations

$$\frac{d[NO_2]}{dt} = -k_1[NO_2] + k_3[O_3][NO] - k_4[O][NO_2]$$
$$-k_5[O][NO_2][M] + 2k_6[NO_3][NO] + k_7[O][NO][M] \quad (A.1)$$

If we assume that [O$_3$], [NO$_3$], and [O] are at pseudo-steady state, then:

$$\frac{d[O_3]}{dt} = k_2[O][O_2][M] - k_3[O_3][NO] = 0$$

Therefore:

$$[O_3] = \frac{k_2[O][O_2][M]}{k_3[NO]} \quad (A.2)$$
\[
\frac{d[NO_3]}{dt} = k_5[O][NO_2][M] - k_6[NO_3][NO] = 0
\]

Therefore:

\[
[NO_3] = \frac{k_5[O][NO_2][M]}{k_6[NO]} \quad (A.3)
\]

\[
\frac{d[O]}{dt} = k_1[NO_2] - k_2[O][O_2][M] - k_4[O][NO_2] - k_5[O][NO_2][M] - k_7[O][NO][M] = 0
\]

\[
[O] = \frac{k_1[NO_2]}{k_2[O_2][M] + k_4[NO_2] + k_5[NO_2][M] + k_7[NO][M]}
\]

\[
[O] = \frac{k_1[NO_2]}{D} \quad (A.4)
\]

Substituting equations A.2, A.3 and A.4 into equation A.1 gives:

\[
\frac{d[NO_2]}{dt} = \frac{-k_1[NO_2]D + k_1k_2[NO_2][O_2][M] - k_1k_4[NO_2]^2}{D}
+ \frac{k_1k_5[NO_2]^2[M] + k_1k_7[NO_2][NO][M]}{D}. \quad (A.5)
\]

Simplifying equation A.5 and rearranging gives:

\[
\frac{d[NO_2]}{dt} = \frac{-2k_1k_4[NO_2]^2}{D} \quad (A.6)
\]
and:

\[
\frac{d[NO_2]}{d[NO_2]} = \frac{D}{k_4[NO_2]} \quad (A.7)
\]

where:

\[
\frac{D}{k_4[NO_2]} = 1 + \frac{k_2[O_2][M]}{k_4[NO_2]} + \frac{k_5[M]}{k_4} + \frac{k_7[NO][M]}{k_4[NO_2]} \quad (A.8)
\]

Making the following definitions:

\[
R_1 = \frac{k_5[M]}{k_4}
\]

\[
R_2 = \frac{k_7[M]}{k_4}
\]

\[
R_3 = \frac{k_2[M]}{k_4}
\]

Then:

\[
\frac{-2k_1}{d(ln[NO_2])} = 1 + \frac{R_3[O_2]}{[NO_2]} + R_1 + \frac{R_2[NO]}{[NO_2]} \quad (A.9)
\]

Defining: \([NO] = [NO]_p + [NO_2]_o - [NO_2] \]

\[
\Delta t = t_{final} - t_{initial}
\]

Integrating equation A.9, and assuming that the initial NO and O_2 con-
centrations are zero gives:

\[ k_1 = \frac{1}{2\Delta t} \left\{ [1 + R_1 - R_2] \ln \frac{[NO_2]_o}{[NO_2]} + R_2 \left( \frac{[NO_2]_o}{[NO_2]} - 1 \right) \right\} \] (A.10)

This last equation is a measure of the light intensity.
Appendix B

Experimental Protocol
This appendix is intended as a general guide for future users of the Indoor Photochemical Reactor. It is not intended to be an exhaustive, step-by-step manual for performing an experiment. Instead, it is written to provide the user with helpful suggestions before and during the running of experiments in the indoor system. Before reading this, it would be helpful to have reviewed the following reference materials:

- Thermo Environmental Inc., Manual for NO\textsubscript{x} monitor, Model 42.
- Dasibi Inc., Manual for O\textsubscript{3} monitor, Model 1003-PC.
- Hewlett Packard, Manuals for GC-FID, Model 5890.
- Tekmar, Manual for Aerotrap 6000 and Cryofocusing unit.
- Waters Inc., Waters Sep-Pak DNPH-Silica Cartridge Care and Use Manual

**B.1 Before an Experiment**

There are several procedures which should be completed before running an experiment. These procedures are listed in the subsequent subsections.

**B.1.1 NO\textsubscript{x} Monitor Calibration**

A detailed description of the calibration procedure for the NO\textsubscript{x} monitor appears in the manual for the instrument. The reader is referred to section
III of the manual for details. After calibration, the results should be plotted and calibration factors should be entered into Labtech Notebook’s data acquisition program (blocks 1,2, and 3). Full calibrations should be repeated whenever the monitor begins to drift. Span and zero checks should be conducted at the start of each day of experiments. Comparison of the results of the daily zero and span checks to the full calibration curve will determine when the instrument has drifted beyond a reasonable amount. The NO$_x$ monitor manual lists $<10\%$ as a reasonable drift, but $<5\%$ is a better value to follow. Typical drifts for the NO$_x$ monitor attached to the indoor system have been less than 1-3\% over as long as a two month period.

**B.1.2 O$_3$ Analyzer Calibration**

The O$_3$ analyzer is taken to the Air Quality Management District for full calibration once a year. They will provide a calibration curve which can then be used in Labtech Notebook’s data acquisition program (block 4).

**B.1.3 GC-FID Calibration**

Every organic compound studied should be calibrated on the GC-FID. A full calibration should be performed at least once to determine whether the compound is linear over the range of concentrations considered. If the GC-FID is left on over the course of the study of the hydrocarbon of interest, one need only conduct a one or two point calibrations (using duplicate points)
on a daily basis. Leaving the GC-FID on over the course of an experiment ensures that all FID gas flows and response factors remain essentially the same over the course of the experimental study. We have found that the GC-FID Series II attached to the indoor system is quite stable over time.

B.2 Running an Experiment

B.2.1 Admitting Reactants

There are three primary methods of admitting reactants to the chamber. Reactants can be admitted from a standard gas cylinder, from the vacuum manifold, and from what has been termed as the "evaporation" bulb (see Figure 2.6). The most important point to remember when admitting reactants is to make sure that the teflon lines from the gas cylinders, vacuum manifold or the evaporation bulb are fully flushed with scrubbed air after the reactant has been admitted. This added flushing ensures that the reactant has reached the chamber and that there are no remnants of reactant in the teflon lines, vacuum manifold or evaporation bulb.

*Nitric Oxide.*

Nitric oxide is admitted using the high concentration NO cylinders available in the laboratory. It is important to measure the flowrate of the NO and the time taken to admit the reactant. An estimate of the concentration admitted to the chamber can be made by using the following formula:

\[ [\text{NO}] \text{ ppm} = \frac{F \times [\text{NO}]_{cylinder}}{V_{bag}} \]
where:

\[ F = \text{flowrate of NO from the cylinder (liters/min)}, \]
\[ t = \text{time (in min.) taken to admit the reactant}, \]
\[ [NO]_{cylinder} = \text{concentration of the standard cylinder of NO, and} \]
\[ V_{bag} = \text{volume of the teflon chamber (in liters)}. \]

**Methyl Nitrite.**

Methyl nitrite is stored in a cold finger in liquid nitrogen, and is admitted through the vacuum manifold. Before admitting methyl nitrite to the chamber it is important to utilize the manifold to purify the methyl nitrite. Using the technique of freezing-pumping-thawing the cold finger one or two times works well. Before admitting the methyl nitrite, it is important that it be sufficiently thawed. The pressure gauge on the manifold can be used to admit a known amount of reactant to one of the calibrated bulbs on the manifold. Assuming ideal gas behavior, the partial pressure of methyl nitrite in the bag can be determined using the following relationship:

\[
P_{\text{chamber}} = \frac{P_{\text{manifold}} \cdot V_{\text{manifold}}}{V_{\text{chamber}}} \tag{B.1}
\]

where \( P \) and \( V \) are the partial pressures and volumes of methyl nitrite in either the chamber or the glass bulb on the vacuum manifold. (The volume of the glass bulb on the vacuum manifold can be calibrated using a gravimetric technique, i.e. placing water in the bulb, weighing the bulb, and computing
the volume.) By flowing scrubbed air to the manifold and through the bulb containing the reactant, the methyl nitrite is admitted to the chamber. It is important to let the air flow through the manifold for a while to ensure that the reactant has been completely admitted to the chamber.

**Hydrocarbon**

Hydrocarbons are admitted using the evaporation bulb. A known amount of pure, liquid hydrocarbon is injected to the evaporation bulb. The bulb is connected to the scrubbed air supply. While flowing air over the liquid, the bulb is gently heated using a heat gun. Again, it is important to continue flowing scrubbed air through the bulb even after all of the liquid has evaporated.

### B.2.2 Sampling From the Bag

Once the sampling valve on the chamber and the appropriate valves on the glass sampling manifold are opened, gas samples from the chamber can be obtained. The only caution to be given during sampling concerns the start of the experiment. It is extrememly important that the user wait until the system is fully stabilized and well mixed before proceeding with an experiment. The user will have to perform several experiments to determine exactly how long of a delay is needed to ensure stabilization. Two things to monitor are the NO<sub>x</sub> monitor values and the hydrocarbon concentrations. It is important that the t=0 hydrocarbon peak areas for the GC-FID be within a few percent. If the user finds that the concentrations of reactants decrease over
time, and the system has had time to mix, radical sources within the chamber have probably reached high concentrations. It is then necessary to either re-condition the bag or change the bag. It is often best to simply change the bag.

**B.3 Teflon Chamber**

**B.3.1 Deciding When to Change a Bag**

One cannot really place a time limit on the frequency of bag changing since the frequency depends upon what has been placed in the bag and how often the bag has been used. As mentioned previously, though, it is important to change the bag when radical sources become too high. To test for the presence of radical sources, the user should irradiate a bag containing only air. The analytical instruments should be used to monitor the concentrations of $\text{O}_3$, $\text{NO}_x$, and hydrocarbon present in the bag. If there are significant radical sources in the bag, one should see an increase in the $\text{O}_3$ concentration over time. The bag should be changed if radical sources are found to be significant. In addition, one can make a visual inspection of the bag. Obviously, if there are any damages to the bag, it is important to immediately replace and condition the bag.

**B.3.2 Making a New Chamber**

Teflon (2 mil, FEP) is purchased directly from duPont in rolls that are 58 in. wide. Two sheets, each 6 ft. long, are cut from the 58 in. roll of teflon.
These two sheets of teflon are heat sealed along the long edges with the 14 in. Vertrod Inc. heat sealer available in the roof lab. It is important to place the seals within one inch from the edge of the teflon, or the user may encounter problems placing the teflon over the circular plates that form the top and bottom of the chamber. To strengthen the seals, it is wise to either make double seals on each edge or to reinforce the edges with teflon tape.

B.3.3 Conditioning the Bag

It is important to fully condition the chamber before performing experiments, since FEP teflon is a source of CO and volatile residues (Kelly, 1982; Killus and Whitten, 1990). To condition the chamber, fill the bag with NO$_2$ and air, and photolyze. Alternatively, the system can be filled with O$_3$ and air and photolyzed. It is crucial to fully flush the bag with scrubbed air after the conditioning process. The bag is considered to be fully conditioned when radical sources are insignificant (see the section entitled “Deciding when to change a bag”).

B.4 2,4 Dinitrophenylhydrazine Analyses

The 2,4 dinitrophenylhydrazine (DNPH) method has been used extensively to collect and identify gas-phase carbonyl compounds. Gas samples from the indoor chamber are collected onto cartridges impregnated with an acidic DNPH solution. DNPH selectively reacts with aldehydes and ketones according to the reaction pictured in Figure B.1.
Figure B.1: 2,4 Dinitrophenylhydrazine Derivatization Reaction
The DNPH cartridge samples which are collected are eluted with 3 ml of acetonitrile, and analyzed using high performance liquid chromatography. To determine product yields, DNPH derivatized standards for all identified carbonyl compounds are required. These DNPH derivatized carbonyls are not always readily available. Radian corporation has many of the simpler carbonyls such as formaldehyde, acetone, and acetaldehyde, but they do not have many other monocarbonyls, nor do they carry dicarbonyls such as glyoxal and methyl glyoxal.

**B.4.1 DNPH-Carbonyl Derivatization**

Several monocarbonyls are readily available from Aldrich chemical company. They can be used without further purification, since many of them have stated purities of 95% or greater. The method for derivatization presented here is adapted from the Standard Operating Procedure for Carbonyl Analysis by HPLC from DGA, Inc., Ventura California. An additional procedure is available in the Waters Sep-Pak DNPH-Silica Cartridge Care and Use Manual. The first step in the generating the DNPH derivative is to create an acidic DNPH solution. This is done by combining 0.5g of DNPH in 2ml of H$_2$SO$_4$ and 3ml of water. Add 10 ml of ethanol to this solution. Next, prepare the carbonyl solution by adding 0.5 g of the monocarbonyl to 20 ml of ethanol. Add the DNPH acidic solution to the carbonyl solution. A precipitate will form within a few minutes. This precipitate is the hydrazone of the monocarbonyl. Purification of the derivative may be necessary. A qualitative standard of derivative and acetonitrile can be synthesized and tested
on a liquid chromatograph to determine the purity of the hydrazone. To
purify the derivative, recrystallize it using a minimum amount of methanol.
If necessary, heat the solution to make it supersaturated. Once the solution
begins to cool, fine, needlelike crystals of the derivative will form. Vacuum
filter the crystals from the solution with methanol. A bit of ether can be used
to speed the drying process of the crystals. Care should be taken during the
crystallization process to avoid contact with sources of carbonyls.

Dicarbonyls such as glyoxal and methyl glyoxal are also available from
Aldrich. These can be used without further purification. The same method
for generating monocarbonyl derivatives can be used to generate 1,2 dicar-
bonyl DNPH derivatives. In this case, however, 1.25 gm of the dicarbonyl
should be used in the carbonyl solution. It was found that the dicarbonyls
required recrystallization. The procedure mentioned above can be used to
recrystallize the dicarbonyl derivative.

Standard solutions, stable for up to several months, can be formed with
acetonitrile. To prepare the standard solutions, it is best to start with a
highly concentrated solution (approximately 200 $\mu g/mL$, as carbonyl) and di-
lute with acetonitrile until the desired concentration range is reached. The
following formula can be used to calculate the concentration of the standard:

$$\text{Concentration, } \mu g/mL \text{ as carbonyl} = \frac{mg \text{ hydrazone}}{mL \text{ of acetonitrile}} \times \frac{MW \text{ carbonyl}}{MW \text{ hydrazone}} \times 10^3.$$  

Note that dicarbonyls will be double derivatized by the DNPH. This fact must
be taken into account when calculating the concentrations of the standards.

To check the quality of the standards, calibrations curves should be gener-
ated for each DNPH carbonyl standard synthesized in the laboratory. These
curves should be linear. As an example, the calibration curves for several glyoxal and methyl glyoxal standards are presented in Figure B.2.

**B.4.2 Calculation of Carbonyl Yields**

Once the carbonyl standards have been synthesized, experimental yields can be calculated. To do so, use the following formula:

\[
\text{Concentration}(\text{ppb}) = \frac{\text{PeakHeight}_{\text{exp}} - \text{PeakHeight}_{\text{blank}}}{x} \times \frac{V_{\text{eluate}}}{V_{\text{gas}}} \times \frac{1}{y}
\]

where:

- \(\text{PeakHeight}_{\text{exp}}\) = Peak Height of compound from the experimental sample
- \(\text{PeakHeight}_{\text{blank}}\) = Peak Height of compound from the blank sample
- \(x\) = Response factor for 1 \(\mu g\) of carbonyl standard
- \(V_{\text{eluate}}\) = Volume of eluate (ml)
- \(V_{\text{gas}}\) = Volume of gas collected onto the cartridge (m³)
- \(y\) = Conversion factor from \(\mu g\) m³ to ppb for the carbonyl

Conversion factors have been calculated for several carbonyls, and are tabulated in Table B.1.
Figure B.2: Calibration curves for DNPH-glyoxal and DNPH-methyl glyoxal standards synthesized at Caltech
Table B.1: Conversion Factors for Selected Carbonyls

<table>
<thead>
<tr>
<th>Carbonyl</th>
<th>Conversion Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaldehyde</td>
<td>1.80</td>
</tr>
<tr>
<td>Acetone</td>
<td>2.37</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>4.33</td>
</tr>
<tr>
<td>3,3-Dimethylbutyraldehyde</td>
<td>4.10</td>
</tr>
<tr>
<td>2,2-Dimethyl-1-propanal</td>
<td>3.52</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>1.23</td>
</tr>
<tr>
<td>Glyoxal</td>
<td>2.37</td>
</tr>
<tr>
<td>Methyl Glyoxal</td>
<td>2.94</td>
</tr>
<tr>
<td>2-Methyl-1-propanal</td>
<td>2.95</td>
</tr>
<tr>
<td>3-Methyl-1-butanal</td>
<td>3.52</td>
</tr>
<tr>
<td>Tolualdehyde</td>
<td>4.91</td>
</tr>
</tbody>
</table>

**B.4.3 Interferences with the DNPH Technique**

High levels (approximately 100 ppb) of ozone have been found to interfere with the accurate collection of DNPH samples. In order to eliminate the effect of O₃ on the DNPH cartridges, it is important to install a KI trap in-line between the glass sampling manifold and the DNPH cartridge. Experiments with and without the trap in place should be conducted by the user to convince oneself of the effectiveness of the KI trap. A full description of the construction of a KI ozone scrubber appears in the reference, “Waters Sep-Pak DNPH Silica Cartridge Care and Use Manual”, published by Waters Inc., and available with every box of DNPH silica cartridges.
Appendix C

Energies of Aromatic Intermediates
In order to provide the reader with a more complete view of the steps leading to bicyclic radical formation, we have included energy “trees”, indicating the possible radicals from each of the steps leading to bicyclic radical formation for the five aromatics studied in chapter 4. Toluene data appear in Figures C.1- C.3, m-xylene data in Figures C.4- C.6, p-xylene data in Figure C.7, 1,2,4 trimethylbenzene data in Figures C.8- C.10, and m-ethyltoluene data in Figures C.11- C.14.
Figure C.1: Energy diagrams for toluene radicals.
Figure C.2: Energy diagrams for toluene radicals.
Figure C.3: Energy diagrams for toluene radicals.
Figure C.4: Energy diagrams for m-xylene radicals.
Figure C.5: Energy diagrams for m-xylene radicals.
Figure C.6: Energy diagrams for m-xylene radicals.
Figure C.7: Energy diagrams for p-xylene radicals.
Figure C.8: Energy diagrams for 1,2,4 trimethylbenzene radicals.
Figure C.9: Energy diagrams for 1,2,4 trimethylbenzene radicals.
Figure C.10: Energy diagrams for 1,2,4 trimethylbenzene radicals.
Figure C.11: Energy diagrams for m-ethyltoluene radicals.
Figure C.12: Energy diagrams for m-ethyltoluene radicals.
Figure C.13: Energy diagrams for m-ethyltoluene radicals.
Figure C.14: Energy diagrams for m-ethyltoluene radicals.
Appendix D

Calculation of the Average

\([\text{NO}_2]\)
The expression used to calculate the average value of NO₂, \([\overline{NO}_2]\), was determined as :

\[
[NO_2] = \frac{1}{n} \sum_{i=1}^{n} ([CH_3ONO]_0 (1 - e^{-kt}) + \Delta [NO]_i)
\]

where: \(k\) = photolysis rate constant for methyl nitrite, as measured experimentally
\[
\Delta [NO] = [NO]_0 - [NO]_t
\]
\(n\) = number of data.

This appendix presents a derivation of this formula.

In a system with methyl nitrite and NO, the NO₂ concentration at any point in time can be approximated by the formula,

\[ [NO_2] = \text{Contribution from CH}_3\text{ONO} + \text{contribution from NO} \]

Methyl nitrite photolyzes according to the following reactions:

1. \(CH_3ONO + h\nu \rightarrow CH_3O^+ + NO\)
2. \(CH_3O^+ + O_2 \rightarrow HCHO + HO_2\)
3. \(HO_2 + NO \rightarrow OH^+ + NO_2\)

Given these reactions, it is evident that the concentration of NO₂ from this source is simply the difference between the initial and final concentrations of CH₃ONO. While it is easy to determine the initial concentration of CH₃ONO, it is difficult to determine the final concentration by experiment alone. Thus, it is helpful to relate the final concentration of CH₃ONO to the initial concentration. This can be done by developing the rate equations for methyl
nitrite photolysis using Reactions 1-3. The rate expression for the photolysis of methyl nitrite can be stated as follows:

\[
\frac{d[CH_3ONO]}{dt} = -k_1[CH_3ONO]. \tag{D.1}
\]

Rearranging this equation and integrating gives:

\[
\int_{t_o}^{t} \frac{d[CH_3ONO]}{[CH_3ONO]} = \int_{t_o}^{t} -k_1 dt, \tag{D.2}
\]

and

\[
\ln \frac{[CH_3ONO]_t}{[CH_3ONO]_{t_o}} = -k_1 t, \tag{D.3}
\]

which leads to

\[
[CH_3ONO]_t = [CH_3ONO]_{t_o} \exp^{-k_1 t} \tag{D.4}
\]

Substituting Equation D.4 into the expression for \([NO_2]\) gives the following expression:

\[
NO_2 = [CH_3ONO]_o(1 - \exp^{-k_1 t}) + \text{contribution from NO.}
\]

If we make the assumption that all of the NO is used to reduce peroxy radicals (RO_2) to oxy radicals (RO^-), forming one NO_2 radical in the process, then the contribution to the concentration of NO_2 from NO is simply the difference between the starting and final concentrations of NO, \(\Delta[NO]\). Thus, the expression for \([NO_2]\) becomes:
Note that the assumption that all of the NO is used to reduce peroxy radicals to oxy radicals as opposed to forming the nitrate is reasonable given rate constant considerations for each path (see J.Phys. Chem Ref. Data, Monograph 2, 1995).

Summing over all of the readings, and dividing by the number of data points determined in time, $t$, results in the expression for the average NO$_2$ concentration,

$$\overline{[NO_2]} = \frac{1}{n} \sum_{i=1}^{n} ([CH_3ONO]_i (1 - e^{-kt}) + \Delta [NO]) .$$