Detailed Kinetic Modeling of Gas-Phase Reactions in the Chemical Vapor Deposition of Carbon from Light Hydrocarbons

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The chemical kinetics of the pyrolysis of the hydrocarbons ethylene, acetylene, and propylene are modeled in detail under conditions relevant to the chemical vapor deposition of pyrolytic carbon. A mechanism that consisted of 227 species and 827 reactions (most of which are reversible) is developed and computed using a software package designed for computing time-dependent homogeneous reaction systems. Experimental results used for model validation are obtained using a vertical flow reactor at 900 °C; pressures of 2−15 kPa, and residence times of up to 1.6 s. The products are analyzed using on-line and off-line gas chromatography. Computational and experimental results are compared for more than 30 products, including hydrogen, small hydrocarbons (ranging from methane to C4), and aromatic hydrocarbons (ranging from benzene to coronene). The resulting reaction model predicts the profiles of the major pyrolysis products (mole fractions of >10−2) of the three hydrocarbons, as a function of both residence time and pressure, with satisfactory accuracy. It also predicts the mole-fraction profiles of minor compounds, ranging from polycyclic aromatic hydrocarbons (PAHs) to naphthalene, fairly well; however, it significantly underpredicts the larger PAHs. The deviation increases as the molecular mass of the PAHs increases. Sensitivity and reaction-rate analyses were also conducted to identify crucial reaction steps.

1. Introduction

Pyrolytic carbon production via chemical vapor deposition (CVD) is one of the most important processes in carbon materials technology, especially for carbon fiber-reinforced carbon (C/C) composites. Isobaric and isothermal chemical vapor infiltration (CVI) of carbon is an accepted process for the synthesis of C/C composites used in the aerospace industry for rockets, missile nozzles, and brake disks. In the CVI of carbon, the following processes proceed simultaneously and interdependently: (1) homogeneous gas-phase pyrolysis of light hydrocarbons (generally methane, propane, and propylene); (2) transportation of gaseous species inside a fibrous preform; and (3) heterogeneous reactions yielding solid deposits that densify the preform. Gas-phase reactions form a great variety of hydrocarbons and hydrocarbon radicals, and any of these species has the potential to chemisorb or physisorb at the growing pyrolytic carbon surface. The chemistry underlying carbon CVD and CVI has been obscured by the complexity of the mechanism, which involves both the gas phase and surface.

Mathematical models have been increasingly applied to understand the CVD and CVI of carbon more precisely. Appropriate models must incorporate chemical reaction schemes and all of the transport phenomena involved. Becker and Hüttner have proposed a simplified model that predicts the kinetics of carbon deposition from C2 hydrocarbons. In their model, seven parameters, including five adjustable reaction-rate constants, are used to fit only two observables: deposition rate and residence time. In addition, a large number of gas-phase species are lumped into three groups: C2, C3, and C4 hydrocarbons. Transport phenomena are not considered, and the deposition reactions are treated using simple first-order reaction constants. This simplified model is a good benchmark for further studies, including simulations of the CVI process; however, as it stands, it explains experimental results for only a limited range of conditions and gives little insight into the CVD of carbon at a molecular level.

To understand these chemical reaction processes more precisely, a detailed chemical kinetic modeling approach is needed, rather than the “lumped” approach of Becker and Hüttner. In the more-sophisticated approach, hundreds of elementary reactions are compiled, and the governing equations for a given system are solved numerically. The parameters that must be considered are the system parameters, such as temperature, pressure, time, and initial reactant concentrations, and the kinetic parameters for each chemical reaction. Assuming an Arrhenius-like rate expression, the pre-exponential, temperature exponent, and activation energy must be known for every elementary reaction. The kinetic parameters are intrinsic to the chemical reactions and can be derived experimentally or from theoretical calculations. Therefore, in principle, a model developed using this approach would involve no adjustable or fitting parameters and could be used under a wide range of conditions.

Detailed chemical kinetic modeling for light hydrocarbons pyrolysis were extensively performed, of which a few studies were concerned with the CVD of carbon. Ziegler et al. proposed a reaction mechanism for propane pyrolysis under conditions relevant to the CVI process. They evaluated their mechanism by comparing the computational and experimentally determined data for a perfectly stirred reactor at 900−1025 °C with a residence time of 0.5−3.0 s and a propane pressure of 0.35 kPa. Twenty-nine products were analyzed, ranging from hydrogen to pyrene. Considering the wide range of temperature and residence time, the agreement between the computational and experimental results was quite satisfactory. This mechanistic study was limited to propane pyrolysis at one pressure; however, the pressure or initial concentration of a reactant is an important operational variable in the CVD and CVI of carbon. In the present study, we attempted to develop...
a more-comprehensive mechanism that predicts the pyrolysis behaviors of three hydrocarbons, covering the formation profiles of a wide range of products. Rather than a manual mechanism model development, the computationally generating mechanism approaches\textsuperscript{15,17,20,30–35} would be suitable for modeling such a complex system of hydrocarbon pyrolysis. However, these automated methods are not mature enough to handle the chemistry of larger species such as polycyclic aromatic hydrocarbons (PAHs), which are important precursors of pyrolytic carbon.\textsuperscript{36}

This paper describes a detailed chemical kinetic model that simulates the gas-phase reactions in CVD of carbon from unsaturated light hydrocarbons. To validate this model, we used the experimental results from our previous analysis,\textsuperscript{37} in which we identified and quantified more than 40 gas-phase species in the CVD of carbon from ethylene, acetylene, and propylene. The computed and experimental mole fractions were compared for more than 30 products, including hydrogen and hydrocarbons ranging from methane to coronene. Reaction rate and sensitivity analyses were also conducted to determine the dominant reaction route and crucial steps in the mechanism.

2. Methods

2.1. Experimental Data. The experimental procedure and database used to validate the mechanism have been described in our previous work.\textsuperscript{37} A vertical flow reactor was used and operated at 900 °C, at a total pressure of 2–15 kPa and residence times up to 1.6 s. No inert diluent gas was used; in other words, the pressure means the initial concentration of the precursor. A 4-cm-long isothermal zone was located at the center of the reactor (total length of 44 cm), into which a honeycomb monolith made of cordierite with 400 channels per square inch was placed. Ethylene, acetylene, and propylene (purchased from Air Liquide) were used as the source hydrocarbons. Reaction products were analyzed using both on-line and off-line gas chromatography. The estimated uncertainties in determining the concentrations of gaseous, major condensing products, and minor condensing products (mainly PAHs) were ±9%, ±28%, and ±32%, respectively.

2.2. Mechanism. We developed a gas-phase reaction mechanism by compiling elementary reactions reported in the literature. This reaction mechanism consists of 227 species and 827 elementary reactions, 798 of which are reversible. The mechanism, the references for each of the elementary reactions, the thermodynamic data for all species, and the species abbreviations and corresponding chemical formulas are given in the Supporting Information and can also be downloaded from www.detchem.com.

Mechanism compilation began with extraction of hydrocarbon parts (non-oxygen-containing reactions and species) from the mechanism of aromatics formation in acetylene and ethylene flames reported by Wang and Frenklach,\textsuperscript{38} which was later extended by Appel et al.\textsuperscript{39} This mechanism is very comprehensive and covers the formation of PAHs up to pyrene, but it does not include the formation of hydrocarbons with odd numbers of carbon atoms, such as toluene and indene, and it does not distinguish the substituted positions of the PAHs. The former were obtained from the mechanism proposed by Marinov et al.,\textsuperscript{40} who used many reactions including C_5, C_7, and C_9 species. The detailed reactions for C_7 and C_9 species were added from the mechanism reported by Hidaka and co-workers\textsuperscript{16,41} and Tsang.\textsuperscript{42} The very detailed mechanism of the PAHs up to coronene, from Richter et al.,\textsuperscript{43} who also distinguished the substituted positions of the PAHs, was added to extend the model performance and was ultimately the largest contributor to our mechanism. Several reactions that involved acetone pyrolysis\textsuperscript{44,45} were included, because a significant amount of acetone was present in the acetylene cylinder used in the experiments. Thermochemical data (NASA polynomial form) were taken from several sources.\textsuperscript{38,43,46}

2.3. Computation. Simulations were conducted using the HOMREA software package that was designed for computational analysis of time-dependent homogeneous reaction systems.\textsuperscript{37} In this program, the input includes the forward reaction-rate parameters and thermodynamic polynomials for all of the participating species, in addition to temperature, pressure, and reactant concentrations. The program calculates the rate constant of the backward reaction for every given reversible reaction.

Computations were performed under isobaric and isothermal conditions. The temperature profile along the reactor length was not considered in detail, because its inclusion would have resulted in a large computational expense and made the development of the mechanism accompanying the numerous iterative comparisons of the computational outputs with the experimental data laborious. The calculations using the present mechanism and the software DETCHEM\textsuperscript{48} which incorporates the axial temperature profiles of the reactor, show almost the same outputs as the HOMREA results; therefore, pre-reactions/post-reactions before/beyond the hot zone can be neglected. The purities of the source hydrocarbons and the initial compositions used in the computations are summarized in Table 1. The compositions of the source hydrocarbons were based on the values reported by the manufacturer, with the exception of acetone, which was analyzed by gas chromatography coupled with mass spectroscopy (GC/MS) and was determined to contain 2% (v/v) acetone. Because acetylene is transported and stored dissolved in acetonitrile to prevent explosion, the high fraction of acetone is not surprising. In the computations for acetylene pyrolysis, the initial acetone volume fraction was fixed to 0.018, although the acetone concentration is dependent on the total pressure of acetylene bottle, as discussed by Bergmann et al.\textsuperscript{49} Notably, the calculation always led to incorrect predictions if pure (100%) hydrocarbons were used as the input. Small amounts of impurities, such as methane, ethane, and propane, have an important role in initializing the reactions. Although the addition of methane (0.2 vol %) in the acetylene computations was not validated, it was necessary to initiate the reactions for an acetylene feed. Those species predicted by the mechanism to have reasonable profiles were also analyzed for sensitivity and reaction rates, to identify the crucial reaction steps.

3. Comparison of the Model Prediction with Experimental Data for the Major Species and Reaction Rate Analysis

3.1. Ethylene Pyrolysis. The measured and predicted mole fractions of six major compounds (ethylene, hydrogen, acetylene, methane, 1,3-butadiene, and benzene) found in the CVD experiments conducted with an ethylene precursor at 900 °C and 8 kPa are shown in Figure 1. The experimental mole

<table>
<thead>
<tr>
<th>Source Hydrocarbon</th>
<th>Purity (Vol%)</th>
<th>Composition Used in the Computation (Mole Fraction)</th>
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</thead>
<tbody>
<tr>
<td>Ethylene</td>
<td>&gt;99.4</td>
<td>0.994 C_2H_4, 0.002 CH_4, 0.004 C_3H_6</td>
</tr>
<tr>
<td>Acetylene</td>
<td>&gt;99.6</td>
<td>0.980 C_2H_2, 0.002 CH_4, 0.018 acetone</td>
</tr>
<tr>
<td>Propylene</td>
<td>&gt;99.5</td>
<td>0.996 C_3H_6, 0.001 CH_4, 0.003 C_4H_8</td>
</tr>
</tbody>
</table>

* Company-reported values.
fractions are plotted as a function of time, which corresponds to the effective residence time in the isothermal zone in the reactor. (The term “effective residence time” was defined in our previous work.) The model predicts the consumption of ethylene and the formation of hydrogen, acetylene, and methane. The formation of 1,3-butadiene is overpredicted for the entire range of residence time, and benzene is slightly overpredicted for short residence times. Nevertheless, the model qualitatively predicts all of the trends observed experimentally for all species.

The reaction-rate analysis obtained for the major hydrocarbon species in ethylene pyrolysis under conditions of 900 °C, 8 kPa, and 0.5 s is shown in Figure 2. The thickness of the arrows represents the local rate of consumption or formation of the species at 0.5 s. The rates are expressed as C1-base (in units of mol m⁻³ s⁻¹). Below, we discuss the dominant reactions for the formation or consumption of the major species. Values given below in parentheses denote the contributions of each reaction to the formation or consumption of that species.

Ethylene is consumed by two main reactions, hydrogen abstraction [C₂H₂ + H → C₂H₃ + H₂ (44%)][38] and vinyl radical addition, which yields 1,3-butadiene [C₂H₂ + CH₃ → 1,3-C₄H₆ + H (31%)][38]. Hydrogen addition [C₂H₂ + H → C₂H₃ + M (6%)] and methyl radical attack [C₂H₂ + CH₃ → C₂H₃ + CH₄ (6%)] also contribute to the consumption of ethylene. Acetylene is formed primarily by vinylacetylene decomposition [C₂H₂ → C₂H₂ + C₂H₂ (40%)][39] and secondarily by C₂H₂ + M → C₂H₂ + H + M (16%)[38] 1,3-C₄H₆ → C₂H₂ + C₄H₂ (13%) and C₂H₂ + H → C₂H₂ + C₂H₂ (7%)[31].

The main source of methane is the methyl radical, which reacts with hydrogen, ethylene, and 1,3-butadiene to form methane, i.e., CH₃ + H₂ → CH₄ + H (31%)[38] C₂H₄ + CH₃ → C₂H₅ + CH₄ (25%)[38] 1,3-C₄H₆ + CH₃ → C₂H₅CH₂ + CH₄ (9%)[52] and 1,3-C₄H₆ + CH₃ → 1,3-C₄H₆ + CH₄ (5%)[52] Ethyl radical decomposition [C₂H₂ + CH₃ → CH₃ + CH₄ (25%)] also contributes to methane formation. The primary and secondary sources of methyl radicals are the decomposition of 1,2-butadiene [1,3-C₄H₆ + CH₃ → 1,3-C₄H₆ + CH₄ (50%)] and ethyl radical decomposition [C₂H₅ + H → CH₄ + CH₃ (28%)][38].

The mole-fraction profiles of 1,3-butadiene in both the experiments and model predictions are maximal at ~0.1 s, when the formation and consumption rates are in competition. The subsequent decrease in mole fractions shows that consumption becomes dominant. 1,3-Butadiene is formed almost solely by the reaction C₂H₄ + CH₂ → 1,3-C₄H₆ + H (94%)[38] whereas it is consumed by various reactions, primarily isomerization to 1,2-butadiene [1,3-C₄H₆ → 1,2-C₄H₆ (22%)] [54] hydrogen abstraction by radical attack [1,3-C₄H₆ + H → 1,3-C₄H₅ + H₂ (18%)[38] 1,3-C₄H₆ + H → 1,3-C₄H₅ + H₂ (9%)[38] and 1,3-C₄H₆ + C₂H₂ → 1,3-C₄H₅ + C₂H₂ (8%)[38] vinyl addition forming hexatriene [1,3-C₄H₆ + C₂H₂ → C₆H₆ + H (9%)[52] and decomposition [1,3-C₄H₆ → C₂H₂ + C₆H₆ (9%)[41].

Benzene is formed primarily from 1,3-cyclohexadiene [1,3-C₄H₆ → C₆H₆ + H₂ (52%)] and secondarily by cyclization of 1,3-hexadiene-5-yne [L-C₆H₈ + H → C₆H₆ + H (11%)] [51] Toluene decomposition [1,3-C₄H₆ + H → C₆H₆ + CH₃ (7%)[57] and hydrogen removal from 1,3-cyclohexadienyl [C₆H₆ + H → C₆H₆ + H (6%)] also produce benzene. Most of the benzene comes from C₆ species. The direct formation of benzene from

![Figure 1. Comparison of model predictions (lines) with experimental mole fraction profiles (symbols) of major species during ethylene pyrolysis in a flow reactor at 900 °C and 8 kPa.](image1.png)

![Figure 2. Reaction-rate analysis in ethylene pyrolysis under conditions of 900 °C, 8 kPa, and 0.5 s. The thickness of the arrow represents the local rate of consumption or formation of the species expressed as C1-base (in units of mol m⁻³ s⁻¹).](image2.png)
3.2. Acetylene Pyrolysis. The measured and predicted mole fractions of the six major compounds observed in the CVD experiments using acetylene as a precursor are shown in Figure 3; these compounds were acetylene, hydrogen, benzene, vinylacetylene, methane, and ethylene. The consumption of acetylene was well-predicted, and product formation was predicted fairly well, although vinylacetylene was significantly overpredicted. Again, the simulation qualitatively predicted all of the trends observed experimentally.

Figure 4 shows the result of the reaction-rate analysis obtained according to the proposed mechanism for major hydrocarbon species in acetylene pyrolysis under conditions of 900 °C, 8 kPa, and 0.5 s. The chemistry of acetylene pyrolysis is simpler than ethylene and propylene pyrolysis. Acetylene is consumed primarily by dimerization to form vinylacetylene ($2\text{C}_2\text{H}_2 \rightarrow \text{C}_2\text{H}_4$ (68%)) and secondarily by diacetylene formation ($2\text{C}_2\text{H}_2 \rightarrow \text{C}_6\text{H}_6 + \text{H}$ (17%)). Combination of acetylene and vinylacetylene, $\text{C}_3\text{H}_4 + \text{C}_2\text{H}_2 \rightarrow \text{benzene}$, contributes to 7% of acetylene consumption but the 95% of benzene formation. Ninety-eight percent of vinylacetylene is formed by $2\text{C}_2\text{H}_2 \rightarrow \text{C}_2\text{H}_4$. Ethylene is primarily formed by the decomposition of styrene, $\text{C}_8\text{H}_8 + \text{H} \rightarrow \text{C}_2\text{H}_4 + \text{C}_2\text{H}_2$ (51%). The second source of ethylene is the reaction of vinyl radical with hydrogen (27%). The decomposition of 1,3-butadiene, $1,3\text{-C}_4\text{H}_6 + \text{H} \rightarrow \text{C}_2\text{H}_4 + \text{C}_2\text{H}_2$ (8%), also contributes to the ethylene formation. Methane is formed predominantly from methyl radicals that abstract the H radical from other molecules via reactions such as $\text{CH}_3 + \text{H}_2 \rightarrow \text{CH}_4 + \text{H}$ (76%). Sixty-four percent of methyl is yielded from acetone, which is a contamination (2 vol%) in the acetylene cylinder. The reaction paths are acetone $\rightarrow \text{CH}_3\text{CO} + \text{H}_2$ (32%) and $\text{CH}_3\text{CO} \rightarrow \text{CH}_2 + \text{CO}$ (32%). The decompositions of 1-butyne and 1,2-butadiene, such as 1-butyne $\rightarrow \text{C}_4\text{H}_6 + \text{CH}_3$ (14%) and 1,2-C4H6 $\rightarrow \text{M} \rightarrow \text{C}_2\text{H}_4 + \text{CH}_3 + \text{H}_2 + \text{M}$ (10%), also produce methyl radicals.

3.3. Propylene Pyrolysis. The nine major compounds (propylene, methane, ethylene, hydrogen, benzene, acetylene, propyne, propadiene, and 1,3-butadiene) observed in the CVD experiments using propylene as a precursor are shown in Figure 5. The propylene mole fraction is markedly overpredicted at residence times of $<0.5$ s and is underpredicted at longer residence times. The methane and ethylene profiles are in excellent agreement, and the formation of the other products is predicted fairly well. The prediction curves also capture the experimental trends well, except for those of acetylene and 1,3-butadiene.

Figure 6 shows the result of the reaction-rate analysis obtained according to the proposed mechanism for major hydrocarbon species in propylene pyrolysis under conditions of 900 °C, 8 kPa, and 0.5 s. The chemistry of propylene pyrolysis is most complex. Propylene is consumed primarily by the decomposition that yields methyl and ethylene: $\text{C}_3\text{H}_4 + \text{H} \rightarrow \text{C}_2\text{H}_4 + \text{CH}_3$ (32%). Radical attacks such as $\text{C}_3\text{H}_4 + \text{C}_2\text{H}_4 \rightarrow \text{AC}_3\text{H}_5 + \text{C}_2\text{H}_2$ (28%), $\text{C}_3\text{H}_4 + \text{H} \rightarrow \text{AC}_3\text{H}_5 + \text{H}_2$ (14%), and $\text{C}_3\text{H}_4 + \text{CH}_3 \rightarrow \text{AC}_3\text{H}_5 + \text{CH}_4$ (6%) convert propylene into AC3H5 (allyl radical, CH2=CHCH2•). Hydrogen addition ($\text{C}_3\text{H}_4 + \text{H}$...
N-C₃H₇ (7%) also consume propylene. More than 80% of methane is formed through methyl radicals. The direct formation of methane, via C₃H₆ → C₂H₂ + CH₄ (3%), contributes to only 5% of methane formation. Methyl radicals are primarily formed by C₃H₆ → H → C₂H₂ + CH₃ (50%). The decompositions of SC₃H₅ (2-methyl vinyl radical, ΚHC=CHCH₃) and N-C₃H₇ (CH₃CH₂CH₂Δ), such as SC₃H₅ → C₂H₂ + CH₃ (29%) and N-C₃H₇ → C₂H₂ + CH₃ (12%), also yield methyl radicals. SC₃H₅ is predominantly produced via the isomerization of allyl radicals.

Ethylene is formed directly by C₃H₆ → C₂H₂ + CH₃ (56%) and N-C₃H₇ → C₂H₂ + CH₃ (13%). There is an indirect route through vinyl radicals, C₂H₃ + C₅H₆ → C₂H₄ + C₅H₅ (25%). Acetylene is yielded primarily by SC₃H₅ → C₂H₂ + CH₃ (70%) and secondarily by C₃H₆ → C₂H₂ + CH₄ (11%). Various routes are found in 1,3-butadiene formation. Forty-seven percent of 1,3-C₄H₆ is formed from hydrogen abstraction of C₄H₅ radicals (N-C₄H₅ and I-C₄H₅). Another major route is the decomposition of C₄H₇ radicals (I-C₄H₇ and N-C₄H₇), which accounts for 28% of 1,3-butadiene formation. Metathesis of two olefins such as C₃H₆ + C₃H₆ → 1,3-C₆H₈ + CH₃ (19%) also contributes to the formation of 1,3-butadiene.

4. Sensitivity Analysis

The time-dependent changes in the relative sensitivity coefficients computed for the three major species of interest in the pyrolysis of ethylene (upper), acetylene (middle), and propylene (bottom), under conditions of 900 °C and 8 kPa, are shown in Figure 7. The sensitivity coefficients were calculated for each source hydrocarbon, benzene, and important intermediates, such as 1,3-butadiene, vinylacetylene, and propyne for the pyrolyses of ethylene, acetylene, and propylene, respectively. A positive sensitivity coefficient indicates that the reaction enhances the species production rate or slows its consumption rate, whereas a negative sensitivity coefficient indicates the opposite effect. Although our mechanism consists of more than 800 reactions, the reactions that significantly affect the formation or consumption of the species of interest are rather limited in number. Only a few sensitive reactions are found for ethylene and acetylene pyrolysis, whereas 16 sensitive reactions are found for propylene pyrolysis.

In ethylene pyrolysis, ethylene consumption is greatly affected by ethylene reactions 7 (reaction E07) (vinyl formation from hydrogen attack of ethylene) and 8 (reaction E08) (vinyl addition to ethylene yielding 1,3-butadiene). These reactions are also sensitive in regard to the formation of 1,3-butadiene, especially at short residence times. Reactions E13 (isomerization of 1,3-butadiene to 1,2-butadiene) and E10 (decomposition of 1,2-
butadiene) are sensitive in regard to 1,3-butadiene consumption at longer residence times. Benzene formation is affected greatly by reactions E07, E10, and E13, as well as by reaction E11 (vinyl addition to 1,3-butadiene yielding hexatriene).

Acetylene pyrolysis is rather simple. Acetylene consumption is greatly affected by acetylene reactions 2 (reaction A02) (combination of acetylene to form diacetylene) and 4 (reaction A04) (combination of acetylene to form vinylacetylene). Reaction A04 is also sensitive in regard to vinylacetylene formation, especially at short residence times. Reaction A11 (the combination of vinylacetylene to form cyclo-octatetraene) is sensitive in regard to vinylacetylene consumption. Benzene formation is affected greatly by reactions A04 and A09 (acetylene addition to vinylacetylene, yielding benzene).

Several sensitive reactions were found for propylene pyrolysis. Propylene consumption is greatly affected by propylene reactions 10 (reaction P10) (isomerization of allyl to 2-methyl vinyl radical), 19 (reaction P19) (cyclopentadiene decomposition yielding cyclopentadienyl radical), 26 (reaction P26) (allyl formation from cyclopentadienyl radical), and 12 (reaction P12) (decomposition of propylene to form allyl). The cyclopentadienyl radical is likely a key species in propylene pyrolysis.
Table 2. Rate Constant Parameters for the Selected Reactions That Exhibit Global Sensitivity Coefficients of >0.2 (Mole Fractions of \(10^{-2}\)) for the Formation and Consumption of Major Compounds in the Pyrolysis of Ethylene, Acetylene, and Propylene

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Value</th>
<th>Error</th>
<th>(k) at 900°C</th>
</tr>
</thead>
</table>

The rates of formation (reaction P19) and hydrogen abstraction (reaction P26) affect propylene consumption. The isomerization of propadiene to propyne (reaction P04) is the most sensitive reaction for propyne formation, whereas its reverse reaction (reaction P05) is most sensitive for propyne consumption. Reactions P08 (direct formation of propyne from the decomposition of 2-methyl vinyl radical) and P16 (allyl formation from methyl attack on propylene) are sensitive for propyne formation at short residence times, whereas reaction P20 (propadiene formation from allyl by cyclopentadienyl attack) becomes sensitive at longer residence times. The isomerization of allyl to the 2-methyl vinyl radical (reaction P10) initially has a positive effect on propyne formation, but later has a negative effect. Propyne conversion to propargyl by hydrogen abstraction

\(\text{Conditions: } 900^\circ\text{C}, 8\text{ kPa, and } \tau = 0.5 \text{ s, where } k = AT^a \exp(-E_a/(RT))\). \(^b\) CYPD = cyclopentadiene; CYPD\(^d\) = cyclopentadienyl. \(^a\) Values are for a pressure of 8 kPa.
by methyl (reaction P06) and the conversion of the 2-methyl vinyl radical into acetylene and methyl (reaction P11) are sensitive in propyne consumption. Benzene formation is affected by reactions P19 and P07 (the combination of propargyl and allyl to form benzene), whereas reactions P08, P12, and P26 have positive effects on benzene formation. Cyclopentadienyl also likely has an important role in benzene formation as a source of 1-methyl-cyclopentadiene, which converts benzene (see Figure 6). In addition, the hydrogen abstraction of C3 species, creating important intermediates for benzene formation, such as allyl, propadiene, propyne, and propargyl, are also important.

Table 2 lists rate constant parameters for the selected reactions that exhibit global sensitivity coefficients of >0.2 for formation and consumption of major compounds (mole fractions larger than 10^{-2}) in the pyrolysis of ethylene, acetylene, and propylene under the conditions of 900 °C, 8 kPa, and 0.5 s.

5. Pressure Dependence

The predictive performance of our mechanism for ethylene pyrolysis at varying pressures (initial concentrations of the precursor hydrocarbons) is shown in Figure 8. Similar figures for the other two source hydrocarbons (acetylene and propylene) are given in the Supporting Information. Generally, the pressure effect on the consumption of the source hydrocarbons, as well as on formation of the major products, is modeled very well. Although some noticeable disagreements between the experimental and computational results are present (e.g., 1,3-butadiene in ethylene pyrolysis), our kinetic model adequately captures the trends of the pressure effect; an increase in pressure enhances consumption of source hydrocarbons and also enhances product formation. However, this simple rule is not valid for acetylene formation in ethylene pyrolysis. The acetylene mole fraction increases with pressure at residence times up to 0.2 s, whereas it is maximal at 8 kPa at longer residence times. Acetylene consumption becomes dominant at the highest pressure tested (15 kPa). This rather complicated trend, which has been observed experimentally, is traced correctly by the computational results.

6. Comparison of the Model Prediction with Experimental Data for the Minor Species

6.1. Minor Species. The mole fractions of minor compounds found in the gas phase for propylene pyrolysis at 900 °C and 8 kPa are shown as a function of residence time in Figure 9. The minor compounds are classified as C3 and C4 hydrocarbons (upper), benzene substitutes (middle), and compounds with two rings and acenaphthylene (bottom). Similar figures for the remaining two hydrocarbon sources (ethylene and acetylene) are given in the Supporting Information. Although some gaps in the agreement between the experimental and computational results are noted, especially for acenaphthylene, the mechanism generally predicts the mole fractions of the minor species well, with mole fractions ranging from 10^{-5} to 10^{-2}.

6.2. Polycyclic Aromatic Hydrocarbons (PAHs) Larger than Phenanthrene. The mole fractions of polycyclic aromatic hydrocarbons (PAHs) larger than phenanthrene found in the gas phase from acetylene pyrolysis at 900 °C and 15 kPa are shown as a function of residence time in Figure 10. Similar figures for ethylene and propylene are found in the Supporting Information. The experimental mole fractions of the PAHs range from 10^{-3} to 10^{-5} and generally decrease as the molecular size of the PAHs increases. PAH formation is underpredicted, and the deviations generally become larger as the molecular size of the PAH increases. The computed mole fractions of coronene are 4 orders of magnitude lower than the experimental mole fractions. The experimental uncertainty is ±32%, even for the PAHs formed at low concentration levels. The deviations between the model predictions and experimental data for PAHs larger than pyrene vary from an order of magnitude to several orders of magnitude and are much larger than the experimental uncertainties. This
indicates that the experimental uncertainties in determining the concentration of the large PAHs cannot fully explain the gaps between the computations and experiments.

Our model does not satisfactorily predict many PAHs. Our computations are based on simple chemical reactions in a completely mixed reactor. The lower mass transportability of the large PAHs might induce an increase in the effective residence time in the reactor, yielding more PAHs than were observed experimentally.

The kinetic parameters of the reactions involved in the chemistry of the PAHs were not obtained experimentally and theoretically but rather estimated, based on their prototype reactions of small molecules. Accurate evaluation of the rate of these reactions, as well as further exploration of new routes to PAHs, will be helpful to improve the model predictions of the PAH profiles. Nevertheless, the increasing gap between experimental and computational results with increasing size of the PAH strongly suggests the need for a transport model that describes the flow characteristics in the reactor. Numerical simulations with a transport model will identify the reason for the disagreements found in the PAH profiles.

7. Summary

The kinetics of gas-phase reactions during the chemical vapor deposition (CVD) of carbon from ethylene, acetylene, and propylene were modeled using elementary reactions consisting of 227 species and 827 reactions and computed using a time-dependent homogeneous reaction system. Experimental data obtained using a conventional flow reactor were used to validate the model. For the pyrolysis of all three hydrocarbons, satisfactory agreement between the computations and experiments was observed for the major species with mole fractions of $>10^{-2}$. Information about the crucial reaction steps in the formation of the large PAHs will be helpful to improve the model predictions.
and consumption of the major species was obtained by sensitivity and reaction-rate analyses. The influences of pressure and concentration also were predicted well.

Our work describes a single mechanism that has a comprehensive capability to predict the major components found in the pyrolysis of three different hydrocarbons with an acceptable level of accuracy. However, disagreements between the computations and experiments are observed for minor species; in particular, the formation of polycyclic aromatic hydrocarbons (PAHs) is significantly underpredicted. Our lack of understanding of the formation chemistry of PAHs and the lack of flow simulation in our model are likely the major sources of the disagreements. Future work will include studies at different temperatures and surface (deposition) chemistry and with different transport models.

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Supporting Information Available: The reaction mechanism and references for each of the elementary reactions (Table S1), species abbreviations and corresponding chemical formulas (Table S2), and thermodynamic data for all species (Table S3), Results of sensitivity analysis (Figures S1–S3) and comparison of model predictions with experimental mole fractions (Figures S4–S9). This material is available free of charge via the Internet at http://pubs.acs.org.

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