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Kinetic modeling of Jet Propellant-10 pyrolysis

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Abstract

A detailed kinetic model for the thermal decomposition of the advanced fuel Jet-Propellant 10 (JP-10) was constructed using a combination of automated mechanism generation techniques and \textit{ab initio} calculations. Rate coefficients for important unimolecular initiation routes of exo-TCD were calculated using the multireference method CAS-PT2, while rate coefficients for the various primary decompositions of the exo-TCD derived monoradicals were obtained using CBS-QB3. Rate-of-production analysis showed the importance of four dominating JP-10 decomposition channels. The model predictions agree well with five independent experimental data sets of JP-10 pyrolysis that cover a wide range of operating conditions (T = 300-1500K, P = 300 Pa – 1.7 \times 10^5 Pa, dilution = 0.7 – 100 mol\% JP-10, conversion = 0-100 \%) without any adjustment of the model parameters. A significant part of the model comprises secondary conversion routes to aromatic and polyaromatic hydrocarbons (PAHs) and could thus be used to assess the tendency for deposit formation in fuel rich zones of endothermic fuel applications.

Keywords: JP-10, pyrolysis; thermal decomposition; kinetic modeling; polyaromatic hydrocarbon formation
1 Introduction

Mono- and polycyclic hydrocarbons represent an interesting class of molecules that can be used as fuels in advanced aviation applications \(^1\). Many of them have superior physical properties such as a high energy content, a low viscosity at low temperatures, an elevated flash point and a higher density compared to the classically used n-alkanes \(^2\)\(^3\). Cyclic alkanes also have higher thermal stability limits and a larger heat sink capacity than their acyclic counterparts (cf. n-decane vs. decalin in ref. \(^4\)), which are important properties in the propulsion technology. Endothermic cracking of the fuel prior to the combustion chamber \(^5\)\(^-\)\(^7\) not only amplifies the heat sink, it can also reduce ignition delays as illustrated by Colket et al. \(^8\). Hence, a thorough understanding of the decomposition chemistry of cyclic hydrocarbons is not only important for the design and optimization of the endothermic reaction technology itself, but also for the correct design of the combustor that converts the products of the endothermic decomposition.

This study focuses on the kinetic modeling of the thermal decomposition of the fuel Jet Propellant-10, “JP-10”, used as the fuel for air-breathing missiles \(^9\). JP-10 consists of a tricyclic hydrocarbon, exo-tricyclo[5.2.1.0\(^{2,6}\)]decane (exo-TCD), cf. Scheme 1.

![Scheme 1: Exo-tricyclo[5.2.1.0\(^{2,6}\)]decane (exo-TCD), the main constituent of JP-10.](image)

Although a fair number of experimental JP-10 pyrolysis studies identified and quantified light products containing up to five carbons \(^10\)\(^-\)\(^15\), less is known about the quantities of heavier intermediates and products. Several studies noted the significant formation of aromatic components such as benzene and toluene under pyrolysis conditions \(^10\)\(^,\)\(^12\)\(^-\)\(^16\). Even in the presence of an oxidizer, aromatic components represented an important share of the products as the comparative study between the flame structure of n-decane and JP-10 flames \(^17\) indicated. Vandewiele et al. \(^18\) obtained a new data set of JP-10 pyrolysis experiments using GCxGC-FID/ToF-MS in which over seventy species with mass-to-charge ratio up to 178 (C\(_{14}\)H\(_{10}\)) were identified and quantified. Validation of two available kinetic models of JP-10 thermal
decomposition, the San Diego model \(^{19}\) and the model by Magoon et al.\(^{20}\), revealed that they were largely inadequate in accurately predicting product distribution trends of the experiments of Vandewiele et al. \(^{18}\). This is not surprising since these kinetic models were designed for higher temperature combustion applications and not for the pyrolysis conditions investigated in that study. Herbinet et al. reported a pyrolysis model for JP-10, but this model is not publicly available \(^{10}\).

It is thus clear that an accurate kinetic model for the purpose of modeling JP-10 pyrolysis is currently not available in the open literature. Therefore, a new dedicated pyrolysis model was constructed with the help of the Reaction Mechanism Generator (RMG) \(^{21}\). To further elucidate the initial decomposition of exo-TCD, a large number of electronic structure calculations were carried out for reactions for which accurate reaction rate coefficients are missing. Five experimental data sets that encompass a wide range of operating conditions were used for the validation of the model.

## 2 Computational methods

### 2.1 Kinetic model construction

Fully automated kinetic model generation is not possible at present. Significant user involvement is still required to obtain an accurate model. Fortunately, automatic network generation programs can reduce part of the burden if the relevant reaction families are known and thermochemical and kinetic data is available. Reaction Mechanism Generator (RMG) \(^{22,23}\) is an open-source software package that automatically constructs kinetic models based on the evaluation of reaction rates and species concentrations. The procedure used by RMG to expand the kinetic model by successive addition of kinetically significant species \(^{24}\) is extensively discussed elsewhere (e.g. \(^{25-27}\) ) and therefore only the elements specific to the modeling of the pyrolysis of polycyclic hydrocarbons are highlighted here. Temperature, pressure and initial exo-TCD dilution, which are used to evaluate rates of production in RMG, are set to 1100K, 1 \(10^5\) Pa and 10 mol %, respectively. The main parameters determining the size of the generated reaction network, i.e. the time at which the integration of the ODEs is stopped and tolerance \(\varepsilon\) for the network generation, were set to 0.5s and 0.001, respectively. For these conditions the most important products observed in the
experiments are included in the reaction network. The use of tighter tolerances significantly increases the size of the model, but does not affect the model predictions. The effect of both parameters on the size of the reaction network for steam cracking of n-hexane was illustrated by Van Geem et al. 27.

RMG was previously employed to construct a model for combustion applications of JP-10 20. The model described in this work is not an update of this combustion model but was built from scratch for endothermic fuel applications of JP-10 and aims at elucidating a complementary part of the thermal decomposition chemistry of JP-10. The pyrolysis of exo-TCD is described mainly by three elementary reaction families: intermolecular hydrogen abstraction reactions by radicals, intra- and intermolecular radical addition reactions and intermolecular radical recombination reactions. Details of the rate rules of these reaction families can be found elsewhere 22, 23. Note that the reactions belonging to the reverse reaction families, i.e. β-scission reactions and unimolecular scission reactions, are allowed too; rate coefficients of these reactions are calculated based on thermodynamic consistency.

In addition to built-in databases for reaction families, a number of reactions from literature were added that contain relevant exo-TCD pyrolysis chemistry. First of all, two sub-models were added that contain chemistry of species with two carbon atoms or less using the seed mechanism option in RMG. The Leeds methane oxidation model v1.5 28, stripped of all oxygen chemistry, was used as a base model for high temperature pyrolysis reactions, resulting in 34 reactions between 22 species. An ethane steam cracking model by Sabbe et al. 26, stripped of all reactions containing four or more carbons, was used as a base model for the low temperature pyrolysis and contains 277 reactions between 35 species. Next, a number of literature sub-models were added to RMG through the so-called “Reaction Library” option of RMG. The reactions in these sub-models are considered during the iterative enlargement procedure and are added to the final model based on the kinetic significance of the species of these reactions. These literature sub-models can be divided into two categories. First of all, sources of cyclic C₅–C₆–C₇ chemistry 29, 30 were included, as it is believed that they are crucial in bridging the gap between the exo-TCD decomposition products and secondary products such as aromatics. Secondly, sources were included that contain pathways for the growth of polycyclic aromatic hydrocarbons such as
indene and naphthalene $^{31,32}$. Finally, thirty-three exo-TCD decomposition reactions were added, primarily originating from new transition state theory (TST) calculations performed in this study.

The automated calculation of pressure-dependent rate coefficients of reactions by RMG was not applied for this study. As a result, the assigned rate coefficients of the generated reactions in the model should be treated as high-pressure limits. Nevertheless, 20 reactions originating from \textit{ad hoc} added reaction libraries have pressure-dependent rate expressions.

### 2.2 Thermochemistry

A detailed decomposition model necessitates the inclusion of thermochemical parameters for all intermediate species. This can be achieved on the basis of group and bond additivity methods $^{33}$ but the accuracy of these estimates may be doubtful for species with structures that fall beyond the scope of the regressed group additive parameters. More specifically, the influence of ring strain on the thermochemistry estimates of polycyclic intermediates remains a point of concern. Magoon et al. $^{34,35}$ addressed this issue by incorporating on-the-fly thermochemical calculations for polycyclic intermediates using three-dimensional atomic coordinates in molecules $^{20}$. In the present work, several important modifications in the calculation of thermochemical parameters for polycyclic intermediates were incorporated. First of all, a database of polycyclic ring strain corrections (RSC) was added containing ca. one hundred polycyclic structures $^{36-40}$. If an appropriate polycyclic RSC is found for a component, it is used in the estimation of the thermochemical parameters. If not, the molecule is passed-on to the framework that generates 3D atomic coordinates on-the-fly using a distance-geometry algorithm implemented in the chemoinformatics toolkit RDKit $^{41}$ to convert the chemical graph representation into a 3D geometry. Secondly, the database of hydrogen bond increments (HBIs), a method developed by Lay et al. $^{42}$ to estimate the thermochemistry of radicals, was expanded with groups for cyclic and polycyclic radicals found in literature $^{43-46}$. The radical atom in strained species cannot adopt the energetically favored planar geometry because of the constriction of the rigid carbon frame and therefore significantly alters the value of the corresponding HBI in comparison with the acyclic analog. Table 1 shows a comparison of the thermochemistry estimates for a selected set of polycyclic strained species relevant for exo-TCD decomposition. Three values are reported: the Benson group additivity method including polycyclic ring strain corrections (RMG-GA), the
semi-empirical PM3 method using 3D atomic coordinates that are generated on-the-fly (RMG-PM3) and finally the best available benchmark value. It was observed that neither RMG-GA nor RMG-PM3 reproduces all benchmark data well. In general, RMG-GA scores better for enthalpies of formation than for entropies or heat capacities because more data is available in literature. In the worst case, S and $C_p$ contributions of polycyclic RSCs were estimated as a sum of contributions of smaller rings. The example of norbornene illustrates this principle. The value of the standard enthalpy of formation at 298 K, estimated through RMG-GA corresponds well with the benchmark value, whereas significantly differences are observed for the standard entropy at 298 K. The values for the enthalpy contribution to the norbornene RSC originate from Wiberg\textsuperscript{37}. Due to the lack of values for the entropy contribution of the norbornene RSC, the value is taken equal to the cyclopentane RSC. Note that the thermochemistry of species in the model is always based on the best available estimates, e.g. CBS-QB3 values for the six tricyclopentyl $C_{10}H_{15}$ radicals from\textsuperscript{47}. 
Table 1: Comparison of the thermochemical properties for selected polycyclic species from the following methods: 1. PM3 method implemented through RMG with on-the-fly generation of atomic coordinates (RMG-PM3); 2. Benson group additivity method implemented in RMG including polycyclic ring strain corrections (RMG-GA); 3. Benchmark values either from ab initio calculations or experimental data.

| Component | Component | \( \Delta H^\circ \) (298K) | S\(^\circ\) (298K) | C\(_p^\circ\) (298K) | C\(_p^\circ\) (800K) | C\(_p^\circ\) (1500K) |
|-----------|-----------|----------------------------|-------------------|-----------------|----------------|-----------------|
| 2-Norbornene | Benchmark | 80
| | RMG-PM3 | 92 | 306 | 100 | 260 | 343 |
| | RMG-GA | 80 | 188 | 121 | 272 | 352 |
| Norbornane | Benchmark | -54.9±1.1
| | RMG-PM3 | -59 | 306 | 105 | 289 | 381 |
| | RMG-GA | -50 | 193 | 134 | 301 | 389 |
| 3-cyclopentyl-cyclopentene (MA110) | THERGAS | 4
| | RMG-PM3 | -54 | 422 | 163 | 406 | 532 |
| | RMG-GA | -13 | 423 | 163 | 402 | 527 |
| BR1 | UB3LYP/cbsb7 | 243 | 410 | 180 | 410 | 536 |
| | RMG-PM3 | 218 | 467 | 159 | 398 | 527 |
| | RMG-GA | 251 | 481 | 155 | 389 | 523 |
| tricyclodecyl radical (TCDR8) | CBS-QB3 | 113 | 364 | 155 | 389 | 511 |
| | RMG-PM3 | 105 | 377 | 142 | 385 | 511 |
| | RMG-GA | 134 | 377 | 142 | 385 | 511 |

*Based on B3LYP calculation, this study.
2.3 Ab initio calculations

High level thermochemical data for species involved in the JP-10 thermal decomposition is very scarce. Rate coefficients are even less available, in particular regarding the initial decomposition of exo-TCD. Therefore in this work new high-level *ab initio* calculations were carried out on a set of reactions that are believed to be important for the decomposition of exo-TCD. Three types of reactions were considered.

Unimolecular C-C scission reactions of exo-TCD leading to biradicals and subsequent intramolecular H-abstraction reactions of the biradicals were explored using density functional theory (DFT) and multireference methods. In the DFT methodology, geometries were optimized using UB3LYP (unrestricted B3LYP) and energies were calculated using CBS-QB3 to explore the PES and obtain initial refinements of kinetic parameters. Using the DFT geometries as initial guess, a multireference methodology CASPT2 $^{51, 52}$, a second-order perturbation theory, multireference counterpart of the MP2 method, as implemented in MOLPRO $^{53}$, was applied. A four-electron, four-orbital active space, generally corresponding to the electrons and orbitals of interest for these reactive transformations, as described in $^{20}$, was employed. In contrast to the previous study $^{20}$, which relied on CASSCF reaction path geometries for performing higher-level single-point calculations, the saddle points and minima considered here were obtained directly on the higher-level, in this case CASPT2. A 6-311G(d,p) basis set, as used by Sirjean et al in CASSCF studies of similar reactions $^{54, 55}$, was applied here. Hessian/frequency calculations at the minima and saddle points were performed using finite differencing of analytic CASPT2 gradients, as calculated by MOLPRO. Partition functions were computed with rigid-rotor, harmonic oscillator (RRHO) approximation and Arrhenius parameters were obtained using conventional transition-state theory (TST).

A third type of reaction is the unimolecular C-C and C-H β-scission of tricyclodecyl (C$_{10}$H$_{15}$) monoradicals. These reactions were studied on the CBS-QB3 level of theory using Gaussian03 and Gaussian09. Tricyclodecyl reactant structures were optimized with “verytight” convergence criteria; correspondingly, a very fine integration grid was used (“int=ultrafine”). Vibrational frequencies, calculated at B3LYP/CBSB7 level of theory, confirmed that the transition state structures were saddle points of first order and have a single
imaginary frequency. Geometries were visually inspected to confirm that the structure was intermediate between the intended reactant and product(s). Conventional transition-state theory (TST) calculations were performed using results for the saddle point and corresponding tricyclodecyl reactant. A table with reaction barriers at 0K for the reactions calculated in this work is added in S2 of the supporting information. Quantum tunneling coefficients were calculated by employing the symmetric Eckart tunneling scheme with energetic information from tricyclodecyl reactant and the saddle point. Frequencies were corrected by a scaling factor of 0.99. These TST calculations were performed using the software package CanTherm. Rate coefficients were computed at ten different temperatures ranging from 298K to 3000K. Based on these sampled points, a three-parameter least squares fit was performed to determine the parameters for a modified Arrhenius rate coefficient expression. A discussion of the uncertainties associated with the 3-parameter fit of the modified Arrhenius parameters is added in Section SXX of the supporting information.

3 Reactor modeling and experimental data

A wide variety of experimental data on the thermal decomposition of exo-TCD published in open literature were used for validating the model. The kinetic model was compared to the product distribution data obtained by Herbinet et al. (hereafter referred to as the “Nancy data”), Nakra et al. (“Nakra data”), Van Devener and Anderson (“Van Devener data”), Rao and Kunzru (“Rao data”), Xing et al. (“Xing data”) and Vandewiele et al., referred to as “Ghent data” in this work. Table 2 shows an overview of the operating conditions of each experimental data set.
Table 2: Summary of operating conditions of the experimental data sets used for model validation.

|                     | Nakra data | Vandevener data | Nancy data | Rao data | Xing data | Ghent |
|---------------------|------------|-----------------|------------|----------|-----------|-------|
|                     | 13         | 14              | 10         | 12       | 15        | 18    |
| Pressure / Pa       | 300-400    | 1500 - 1900     | 110⁵       | 110⁵     | 1-37 10⁵  | 1.7 10⁵|
| Residence time / ms | 2 – 10     | 3 – 10          | 500-6000   | 700-6400 | 480-26400 | 300-500|
| mol % exo-TCD in inert | 2.6:Ar    | 3.9:Ar          | 0.7-4:He   | 100⁶     | 100⁶      | 7/10; N₂ |
| Temperature range / K | 300-1500 | 300 - 1400      | 848-933    | 903-968  | 823-903   | 930-1080|
| Conversion range / % | 0-100      | 0-100           | 0.01-25    | 10-60    | 0-25      | 4-94   |
| Reactor type        | Plug flow  | Plug flow       | CSTR       | Plug flow | Plug flow  | Plug flow|
| Endo-TCD content / % | N/R⁷       | N/R⁷            | 2.5        | 3.6⁷     | 0.23      | 0.6    |

⁷ No diluent present
⁶ Residence time based on the total (including diluent) molar flow rate at the entrance of the reactor, at the mean temperature inside the reactor.
⁵ Based on reported 96.4% purity of reactant
⁴ Not reported

Reactor simulations were systematically conducted with the Chemkin Pro package. The experiments of Herbinet et al. performed in a jet-stirred reactor were modeled with a perfect mixed reactor model, similar to what was done in their comparison with the Nancy model. The steady-state reactor model was solved using a reactor volume of 90 cm³ and a given temperature and pressure. The data from Nakra et al. and Van Devener and Anderson were collected at millisecond time scale under reduced pressure. Both reactors were modeled as plug flow reactors specifying the given diameter and length of the hot zone in the reactor. Isothermal and isobaric conditions with the pressure based on the provided hot zone midpoint pressure were used. In the continuous flow tubular reactor of Rao and Kunzru the axial temperature profile along the reactor was not reported. Instead, the equivalent reactor temperature, i.e. the temperature at which the same exo-TCD conversion would be attained as in the non-isothermal case, was provided. Therefore, the reactor was modeled using isothermal, isobaric boundary conditions. Xing et al. reported JP-10 pyrolysis data up to pressures of 37 10⁵ Pa. Because species fractions were reported for the gaseous and liquid phase separately, without any indication of the relative ratio of the two phases, only exo-TCD conversion was compared. The flow reactor was modeled as a plug flow reactor using isothermal, isobaric boundary conditions. Vandewiele et al. reported experiments in a continuous flow tubular reactor. Radial temperature and concentration gradients were shown to be negligible using two-dimensional reactor models for this reactor, so it was also modeled as an ideal plug flow reactor. For all the flow reactor experiments modeled in this work, the integration was stopped at the reported space time.
JP-10 feedstock contains a number of impurities, such as endo-tricyclo[5.2.1.0\(^{2.6}\)]decane (endo-TCD), adamantane and decalin in decreasing order of importance. While the cumulative content of these impurities typically does not exceed 3.5% of JP-10 feedstock \(^{60}\), the presence of endo-TCD may be of importance for the reactivity of exo-TCD. Hudzik et al. \(^{47}\) reported that the standard enthalpy of formation of endo-TCD is 17 kJ mol\(^{-1}\) higher than the value for exo-TCD, suggesting that the endo-isomer may be less stable than the exo-isomer. Information on the initial decomposition of endo-TCD is non-existing in literature; therefore, only one reaction was added that converts endo-TCD into BR1, similar to the initiation of exo-TCD.

\[
\text{endo-TCD} \rightarrow \text{BR1}
\]

The activation energy of this reaction is lowered by 17 kJ mol\(^{-1}\), relative to the value of the reaction with exo-TCD, to reflect the higher enthalpy of formation of the endo-isomer. The JP-10 feedstock of the modeled experiments was treated as a mixture of exo- and endo-TCD whenever information on the content of endo-TCD was available, cf. Table 2. If not, JP-10 was considered as 100% exo isomer.

4 Results and Discussion

The final RMG-generated, kinetic model for the pyrolysis of exo-TCD contains 5261 reactions between 384 species, i.e. 234 radicals and 150 molecular species and can be found in Section S1 of the Supplementary Information. Thermochemistry of 18% of the cyclic species in the model was estimated via PM3 calculations, 19% from literature while the remainder was estimated via group contribution methods. The model consists of 4308 H-abstraction reactions, 316 radical addition reactions to molecules with double or triple bonds, 158 radical recombination reactions, the remaining 479 reactions originated from specific libraries, as was previously discussed; including 7 ring-opening reactions, 3 intramolecular H-abstraction reactions, and 33 β-scission reactions calculated in this work.
4.1 Primary decomposition chemistry

4.1.1 Exo-TCD initiation
It is generally assumed that the pyrolysis of hydrocarbons such as exo-TCD proceeds through a free radical mechanism. The initiation of exo-TCD involves the creation of radicals through unimolecular bond scission reactions of exo-TCD. Although the rupture of C–H bonds is possible as an exo-TCD initiation route, these bonds are stronger than C–C bonds and were consequently neglected. The unimolecular initiation of exo-TCD can proceed through unimolecular C-C bond scission reactions resulting in the creation of seven distinct biradicals named BRx, x=1..7, cf. Scheme 2.

![Scheme 2](image)

Scheme 2: Exo-TCD initiation routes through unimolecular C–C scission reactions resulting in seven biradicals, BRx, x=1..7. Nomenclature for the biradicals is adopted from Herbinet et al. \(^{10}\). Values represent standard reaction enthalpies (kJ mol\(^{-1}\)) at 298K, based on thermochemistry used in this model.

The resulting highly unstable biradicals can further isomerize to stable molecules via intramolecular H-abstraction reactions (disproportionation reactions), proposed by Herbinet et al. \(^{10}\) upon detecting 3-cyclopentylcyclopentene, further referred to as “MA110”, as a major primary decomposition product.

Intramolecular H-abstraction reactions are thought to proceed via multiple distinct transition states that differ by the polycyclic entity that is formed through the course of the reaction, cf. Scheme 3.
Scheme 3: The unimolecular C-C scission of exo-TCD yielding biradicals and subsequent intramolecular H-abstraction reactions yielding molecular intermediates taken into account in this work.

Herbinet et al. $^{10}$ were among the first to estimate both the ring opening reactions as well as the intramolecular H-abstraction reactions via semi-empirical correlations based on observations by O’Neal and Benson $^{61}$ in which the ring strain of reactants, products and transition states played a central role. These qualitative estimations were combined with thermochemistry estimated via Benson group additivity and showed that the initiation route via BR1 to MA110 was significantly faster than the other competing initiation routes. More recently, Magoon et al. $^{20}$ discussed the difficulty in accurately quantifying reaction barriers for the intramolecular disproportionation reactions. *Ab initio* calculations performed by Magoon et al. $^{20}$ suggested there may be a very small barrier or even no barrier at all. So, Magoon et al. $^{20}$ suggested that initiation of exo-TCD towards MA110 may proceed as a concerted step rather than via a distinct biradical intermediate $^{20}$. In the present work, rate coefficients for 10 ring-opening and intramolecular H-abstraction reactions were calculated. The latter are particularly useful in the re-evaluation of the relative importance of the different exo-TCD initiation routes. Table 3 shows values for the Arrhenius parameters for these reactions.

**Table 3: Rate coefficients for exo-TCD initiation reactions and intramolecular H-abstraction reactions.** Rate
coefficient $k$ is expressed as $k = AT^n \exp\left(-\frac{E_a}{RT}\right)$. Arrhenius parameters are fitted for a temperature range from 298 - 3000K. Unit for $A$ is s$^{-1}$, for $E_a$ kJ mol$^{-1}$.

| Reaction | $A$ | $n$ | $E_a$ | Method used |
|----------|-----|-----|-------|-------------|
| Exo-TCD = BR1 | $1.80 \times 10^{13}$ | 0.83 | 326 | CAS-PT2* |
| Exo-TCD = BR1 | $7.09 \times 10^{12}$ | 0.97 | 319 | UB3LYP/CBS-QB3 |
| Exo-TCD = BR1 | $5.0 \times 10^{15}$ | 0 | 322 | Herbinet et al. $^{10}$ |
| Exo-TCD = BR2c* | $8.32 \times 10^{10}$ | 1.87 | 333 | CAS-PT2* |
| Exo-TCD = BR2d* | $1.09 \times 10^{13}$ | 0.80 | 325 | CAS-PT2* |
| Exo-TCD = BR2c* | $5.06 \times 10^{14}$ | 0.96 | 328 | UB3LYP/CBS-QB3 |
| Exo-TCD = BR2d* | $3.87 \times 10^{12}$ | 0.95 | 321 | UB3LYP/CBS-QB3 |
| Exo-TCD = BR2 | $5.0 \times 10^{15}$ | 0 | 297 | Herbinet et al. $^{10}$ |
| Exo-TCD = BR3 | $6.09 \times 10^{12}$ | 0.92 | 335 | CAS-PT2* |
| Exo-TCD = BR3 | $3.40 \times 10^{12}$ | 1.09 | 334 | UB3LYP/CBS-QB3 |
| Exo-TCD = BR3 | $5.0 \times 10^{15}$ | 0 | 322 | Herbinet et al. $^{10}$ |
| Exo-TCD = BR4 | $8.05 \times 10^{12}$ | 1.24 | 334 | UB3LYP/CBS-QB3* |
| Exo-TCD = BR4 | $5.0^{10}$ | 0 | 322 | Herbinet et al. $^{10}$ |
| Exo-TCD = MA110 | $5.76 \times 10^{10}$ | 6.88 | 416 | UB3LYP/CBS-QB3* |
| Exo-TCD = MA111 | $1.85 \times 10^{11}$ | 1.23 | 336 | UB3LYP/CBS-QB3* |

**Intramolecular H-abstraction reactions**

| Reaction | $A$ | $n$ | $E_a$ | Method used |
|----------|-----|-----|-------|-------------|
| BR1 = MA110 | $8.64 \times 10^{11}$ | 0.08 | -1 | CAS-PT2* |
| BR1 = MA110 | $2.90 \times 10^{10}$ | 0.44 | 15 | UB3LYP/CBS-QB3 |
| BR1 = MA110 | $1.91 \times 10^{10}$ | 1 | 32 | Herbinet et al. $^{10}$ |
| BR2 = BR2_I | $4.5 \times 10^{6}$ | 0.65 | 20 | UB3LYP/CBS-QB3* |
| BR3 = BR3_I | $2.87 \times 10^{10}$ | 0.40 | 23 | UB3LYP/CBS-QB3* |

*For BR2, two reactions are identified corresponding to two different BR2 product conformers.  
* denotes the rate coefficient used in the current model.

Activation energies for the ring-opening reactions are similar to those reported by Herbinet et al. $^{10}$. The good agreement between the *ab initio* and semi-empirical values also shows that the estimation of activation barriers based on the difference in ring strain between the reactant and product is acceptable as a first approximation. Values for the initiation of exo-TCD leading to the formation of BR2 are the exception where the current calculations indicate an activation barrier that is comparable to the other scission possibilities while the Herbinet et al. estimates $^{10}$ are ca. 30 kJ mol$^{-1}$ lower. Good agreement for the ring-opening reactions between the CBS-QB3 values and the CAS-PT2 values for activation energies show the usefulness of the less expensive CBS-QB3 method for these kinds of calculations. Re-evaluation of the global decomposition rates of exo-TCD via the initiation and corresponding subsequent disproportionation channels confirms that the initiation route via BR1 to MA110 is two orders of magnitude faster than the competing
channels via BR2 and BR3, implying that this channel is the principal unimolecular initiation route for exo-TCD at temperatures around 1000K. This is in line with conclusions by Herbinet et al. The concerted pathway from exo-TCD to MA110, using the kinetics by Magoon et al. is significantly slower than the route via biradical BR1. Finally, despite the lesser stability of endo-TCD, and the faster ring-opening kinetics of endo-TCD relative to exo-TCD, the presence of endo-TCD in the JP-10 feedstock did not have a significant influence on the initiation of exo-TCD at the experimental conditions studied in this work.

4.1.2 Tricyclodecyl decomposition reactions
It is expected that a large part of exo-TCD conversion is attributed to H-abstraction reactions by radicals. The abstraction of a hydrogen atom from exo-TCD yields six distinct C_{10}H_{15} tricyclodecyl radicals, cf. Scheme 4. The kinetics of these H-abstractions not only greatly affect the exo-TCD conversion, but they also determine selectivity of the primary products since each tricyclodecyl radical leads to distinct primary products. The activation energy of the H-abstraction reactions by H atoms from exo-TCD were calculated using an Evans-Polanyi relationship proposed by Dean and Bozzelli. The instability induced by the strained nature of the resulting tricyclodecyl radicals was thus taken into account in the kinetics of these hydrogen abstraction reactions.

Scheme 4: Six possible C_{10}H_{15} tricyclodecyl radicals created by H-abstraction reactions from exo-TCD. Nomenclature for the tricyclodecyl monoradicals is adopted from Hudzik et al.

The tricyclodecyl radicals subsequently react via various C-H and C-C β-scission reactions leading to either C_{10}H_{14} molecules or to new C_{10}H_{15} radicals. A single monoradical can react
through multiple parallel decomposition channels. The kinetics for these elementary reactions were not available and therefore they were calculated.

Literature providing kinetic data for cycloalkyl decomposition reactions is very scarce, which makes a comparison of the calculated rate coefficients with previously reported data for these reactions impossible. Instead, the question arises whether the obtained rate coefficients are comparable to the decomposition rates of the smaller units that are present in the tricyclodecyl radicals. Exo-TCD can be regarded as the joining of a cyclopentane ring and a norbornane bicyclic ring, cf. Scheme 1. Cyclopentyl, derived from cyclopentane, has two decomposition routes, one C-C β-scission reaction leading to 1-penten-5-yl and one C-H β-scission reaction leading to cyclopentene and H. Norbornane has three different types of carbon atoms, leading to three possible norbornyl radicals, each of these norbornyl radicals possess C-C β-scission possibilities. Sirjean et al. provided kinetic data at the CBS-QB3 level of theory for these reactions\textsuperscript{63, 64} which can be used in comparing the rate coefficients with the tricyclodecyl decomposition reactions.

Rate coefficients for tricyclodecyl reactions with the analogous reactions of cyclopentyl (Figure 1a) and norbornyl radicals (Figure 1b-d) are compared. Rate coefficients of the tricyclodecyl decomposition reactions are smaller than for the analogous reactions of the smaller ring units, up to one order of magnitude of difference between β-scission reactions of TCDR4 and cyclopentyl, cf. Figure 1a. In some cases, the joining of the cyclopentane and norbornane ring in tricyclodecyl radicals leads to reactions whose TS structure is not comparable to TS’s of the smaller units. Therefore, Figure 1 only compares tricyclodecyl reactions whose TS’s are located in one of the two rings.
Figure 1: Rate coefficients as a function of temperature. Cyclopentyl and norbornyl decomposition reaction rate coefficients from Sirjean et al. 63, 64. Tricyclodecyl decomposition reaction rate coefficients calculated in this work. Tricyclodecyl decomposition reactions are represented by full symbols.

a) TCDR4  

b) TCDR7  

c) TCDR10  

d) TCDR8
4.2 Kinetic model validation

The generated kinetic model was validated against five data sets of exo-TCD pyrolysis experiments. None of the rate coefficients or thermochemical parameters were adjusted by regression of the experimental data, i.e. these comparisons are a test of the predictive ability of the model.

Figure 2a shows the exo-TCD conversion as a function of the mean reactor temperature for the Ghent data. It can be seen that good agreement is found for all temperatures and both levels of dilution, considering the reported uncertainty on the measurement data.
Figure 2: a) JP-10 conversion as a function of temperature for two dilution levels of JP-10 in N$_2$ at reactor inlet for the Ghent data: ● 10 mol% and ○ 7 mol % JP-10 in N$_2$, — 10 mol% and - - 7 mol % JP-10 in N$_2$, P = 1.7 10$^5$ Pa. b) JP-10 conversion as a function of equivalent space time for the Rao data at different temperatures: ♦ 968K, ▲ 948K, ■ 923K, ● 903K. Conditions: 100% JP-10 at the reactor inlet, P = 1 10$^5$ Pa, T = 903-968K. c) JP-10 conversion as a function of reactor pressure for the Xing data at different temperatures: ▲ 903K, ♦ 883K, ▼ 863K, ■ 843K, ● 823K. Conditions: 100% JP-10 at the reactor inlet, T = 823-903K. d) JP-10 conversion as a function of residence time for the Nancy data at different temperatures: ♦ 933K, ▼ 913K, ■ 893K, ● 873K. Lines are model predictions. Conditions: 4 mol% exo-TCD in He at the reactor inlet, T = 848-933K, P = 1 10$^5$ Pa, conversion = 0.01-25%. Symbols are experimental data, lines are model predictions.

Figure 2b shows the exo-TCD conversion as a function of the equivalent space time for the Rao data. The good agreement between experiments and model predictions indicates that the generated kinetic model performs well for high exo-TCD partial pressures. Figure 2c shows the exo-TCD conversion as a function of the reactor pressure for the Xing data, the only dataset at elevated pressures up to 37 10$^5$ Pa. While exo-TCD conversion is predicted to increase with increasing pressure by the model, the experiments show little influence of pressure on exo-TCD conversion at a given temperature. In the experiments of Xing et al., the reported space time increases by a factor of 50 from the lowest reported pressure relative to the highest pressure and explains why the model predictions for exo-TCD conversion at a given temperature increase as a function of pressure. The model predictions are in reasonable agreement at pressures up to 1.8 10$^5$ Pa and tend to slightly deviate at more elevated pressures. While the Xing experiments are very valuable since they are the only available JP-10 pyrolysis experiments recorded at high pressures, it remains unclear what causes the insensitivity of the experimentally measured JP-10 conversion for the reactor space time and pressure. Figure 2d shows the exo-TCD conversion as a function of the residence time for the Nancy data. Good agreement is found between model predictions and experiments, indicating the validity of the model at low conversions under diluted conditions.

Figure 3a shows the mass fractions of the major products as a function of the mean reactor temperature for the Ghent data. Model predictions of propene, CPD and cyclopentene are in good agreement with the data. The predicted maxima of CPD concentrations are at slightly higher
temperatures than experimentally observed. Ethene is consistently underpredicted, which is analyzed in more detail in Section 4.3.1 of this work.

Figure 3: a) Mass fractions as a function of temperature for the Ghent data\textsuperscript{18}. Left: ● dihydrogen, ▲ methane, ■ ethene, —: dihydrogen, – -: methane, – --: ethene. Right: ♦ 1,3-cyclopentadiene, ▼ propene, ⃰ cyclopentene, —: 1,3-cyclopentadiene, – -: propene, – --: cyclopentene. Conditions: 10 mol % JP-10 in N\textsubscript{2} at reactor inlet. P = 1.7 \times 10^5 Pa. b) Mass fractions as a function of exo-TCD conversion for the Rao data\textsuperscript{12}. Left: ▲ methane, ▼ propene, ■ ethene, — methane, – - propene, – -- ethene. Right: ⃰ cyclopentene, × benzene, ♦ 1,3-cyclopentadiene, – – cyclopentene, – - benzene, — 1,3-cyclopentadiene. Conditions: 100 mol % JP-10 at
reactor inlet, T=903-968K, P = 1 \times 10^5 \text{ Pa}. Symbols are experimental data, lines are model predictions.

Figure 3b shows mass fractions of the major products as a function of the equivalent space time for the Rao data. In this case the agreement between model predictions and experimental data is better than for the Ghent data. The main difference is that in this case model predictions of methane agree with experiments, while methane was slightly underpredicted for the Ghent experiments.

Figure 4 shows the mole fractions of the major products as a function of the mean reactor temperature for the Nancy data. Again, good agreement is found for the majority of the major products and confirms the trends seen in Figure 3.
Figure 4: Mole fractions as a function of residence time of a) dihydrogen, b) propene, c) 1,3-cyclopentadiene, d) ethene, e) methane, f) cyclopentene at four temperatures: ♦ 933K, ▼ 913K, ■ 893K, ● 873K. Lines are model predictions, symbols are Nancy data\textsuperscript{10}. Conditions: 4 mol\% exo-TCD in He at the reactor inlet, T = 848-933K, P = 1 \texttimes 10^5 Pa, conversion = 0.01-25\%. 
Mass fractions of the most important formed aromatic species as a function of the mean reactor temperature for the Ghent data are shown in Figure 5a. Besides benzene, the four most significantly formed aromatic species are toluene, styrene, indene and naphthalene. Benzene and toluene are slightly underpredicted at temperatures below 1000K. Indene and naphthalene are overpredicted over the entire temperature range and could suggest that channels to indene and naphthalene are too much favored over the channels to monoaromatics.
Figure 5. a) Mass fractions as a function of temperature for the Ghent data\textsuperscript{18}. Right: ● benzene, ▲ styrene, ■ toluene, —: benzene, – -: styrene, – - –: toluene. Left: ♦ indene, ▼ naphthalene, —: indene, – -: naphthalene. Conditions: 10 mol% JP-10 in N\textsubscript{2} at reactor inlet. P = 1.7 \times 10\textsuperscript{5} Pa. b) Mole fractions as a function of residence time for the Nancy data\textsuperscript{10}. Left: benzene, right: toluene at four temperatures: ♦ 933K, ▼ 913K, ■ 893K, ● 873K. Conditions: 4 mol% exo-TCD in He at the reactor inlet, T = 848-933K, P = 1 \times 10\textsuperscript{5} Pa, conversion = 0.01-25%. Symbols are experimental data, lines are model predictions.

The same trends are observed in Figure 5b that shows mole fractions of benzene and toluene as a function of the mean reactor temperature for the Nancy experiments.

Figure 6 shows the model predictions versus the Nakra and Van Devenen data as a function of temperature. Overall, the good agreement between model and experiment indicates that the current model performs well at low pressures and elevated temperatures despite being generated with RMG at significantly different conditions: T = 1100K, P = 1 \times 10\textsuperscript{5} Pa, 10 mol% exo-TCD in N\textsubscript{2}. Good agreement is observed for exo-TCD conversion. Although both data sets were recorded at similar operating conditions, CPD and ethyne concentrations are underpredicted in the Van Devenen data and overpredicted in the Nakra data while reverse trends are seen for ethene. Less discordant trends are found for benzene. The latter is systematically underpredicted. Benzene model predictions also show a small decrease in benzene mole fractions starting at T > 1500K which was not observed in the experimental data.
Figure 6: Mole fractions as a function of reactor temperature for model predictions versus Nakra data\textsuperscript{13} (left column) and Van Devener data\textsuperscript{14} (right column). a) and b) experiments: ● exo-TCD, ■ ethyne, model predictions: ―: exo-TCD, - - : ethyne, c) and d) experiments: ● ethene, ■ propyne, model predictions: ―: ethene, - - : propyne, e) and f) experiments: ● benzene, ■ 1,3-cyclopentadiene, model predictions: ―: benzene, - - : 1,3-cyclopentadiene. Conditions for the Nakra\textsuperscript{13} and Van Devener\textsuperscript{14} data respectively: 2.6 / 3.9 mol% JP-10 in Argon, P = 300-400 / 1500-1900 Pa, conversion = 0-100%.

The Ghent and Nancy experiments allowed detecting a number of species with very low concentrations. First of all, the Ghent experiments isolated a number of products containing a five-membered ring, such as methylcyclopentadiene (C\textsubscript{6}H\textsubscript{8}) and ethenylcyclopentene (C\textsubscript{7}H\textsubscript{10})\textsuperscript{65}, cf. Figure 7. Since the position of the methyl or ethenyl substituent in methylcyclopentadiene and ethenylcyclopentene could not be determined, they are compared to model predictions of 1-methyl-1,3-cyclopentadiene and 3-ethenylcyclopentene respectively. Good agreement between experiments and model predictions is found for the components.
Tricyclo[5.2.1.0²⁶]dec-4-ene is another higher molecular weight component that was identified as an important primary decomposition product and which also appears in the current model. As can be seen on Figure 7, the temperature of peak concentration is well predicted, but concentrations are slightly mispredicted.

**Figure 7:** Mass fractions as a function of temperature for the Ghent data\(^{18}\) for a) ● 1-methyl-1,3-cyclopentadiene, ■ 3-ethenylcyclopentene, —: 1-methyl-1,3-cyclopentadiene, – – –: 3-ethenylcyclopentene, b) ● tricyclo(5.2.1.0²⁶)dec-4-ene, —: tricyclo(5.2.1.0²⁶)dec-4-ene. Conditions: 10 mol % JP-10 in N\(_2\) at reactor inlet. P = 1.7 \(10^5\) Pa. Symbols are experimental data, lines are model predictions.

The overall good agreement between the model predictions and experimentally measured concentrations for a large number of experimental data sets indicates the usefulness of the current model. None of the model parameters were fitted, despite the many uncertainties in kinetics of reactions and thermochemistry of species. This gives an indication of the overall quality of the kinetic model. Finally, remaining discrepancies between model and experiment should be interpreted carefully. Estimated error bars associated with the reported experimental data were rarely reported, making it difficult to judge to which extent the discrepancies are significant. Also, in some of the experiments the boundary conditions are uncertain and the ideal reactor models employed here are only approximate. Most importantly, despite the best efforts of several researchers including this present work, many of the parameters in this large kinetic model are significantly uncertain, so one would expect significant uncertainties due just to the model. Considering all these factors, the level of agreement between the model predictions and the
experiments is very satisfactory, sufficient to justify some more detailed analysis of the reaction paths below.

4.3 Main reaction pathways

A schematic representation of the main decomposition pathways of exo-TCD predicted by the model for the conditions of the Ghent experiments can be found in Scheme 5. The rates of the important reactions are shown as a percentage relative to the total decomposition rate of exo-TCD.
Scheme 5: Reaction pathway analysis for the decomposition of exo-TCD. The reported percentages represent the predicted reaction rate relative to the total exo-TCD decomposition rate. T=1100K, P=1.7 $10^5$ Pa, Conversion$_{\text{exo-TCD}} = 19\%$, 10 mol% exo-TCD in N$_2$ corresponding to conditions of the Ghent data set. Only major exo-TCD decomposition pathways are shown.
4.3.1 Major decomposition pathways of exo-TCD

While the thermal decomposition of exo-TCD is initiated by the scission of a C-C bond yielding BR1, at the conditions corresponding to Scheme 5, 85% of the decomposition rate of exo-TCD is accounted for by H-abstraction reactions by H. The main decomposition channels involve TCDR5, TCDR8 and TCDR4 with contributions of 33%, 26% and 14% respectively. Decomposition pathways through TCDR7 and TCDR10 are negligible due to the significantly higher bond dissociation energies of the corresponding C-H bonds. The important C_{10} radicals lead to different primary products: TCDR8 leads primarily to cyclopentene and 1,3-cyclopentadiene, with a minor pathway forming benzene and ethyl, TCDR5 leads to 1,4-pentadiene, 1,3-cyclopentadiene, 3-ethenylcyclopentene and propene. TCDR4 leads to ethene and 1,3-cyclopentadiene via a retro-Diels-Alder reaction of 2-norbornene. A secondary decomposition channel of TCDR4 leads to tricyclo[5.2.1.0^{2,6}]dec-4-ene and accounts for 45% of the decomposition rate of TCDR4. The channel through TCDR4 is the only significant source of tricyclo[5.2.1.0^{2,6}]dec-4-ene. One of the major uncertainties affecting tricyclo[5.2.1.0^{2,6}]dec-4-ene predictions is the ring strain present in unsaturated species and derived radicals. For example, H-abstraction reactions from tricyclo[5.2.1.0^{2,6}]dec-4-ene result in resonance-stabilized C_{10}H_{13} species containing rings whose ring strain is unknown.

Ethene is initially formed via various primary decomposition routes starting from BR1, TCDR4, TCDR6 and TCDR8. Once propene attains high enough concentrations, the reaction of propene with H atoms resulting in methyl and ethene forms another major ethene formation channel.

The observation that ethene concentrations are underestimated over most of the data sets at lower temperatures raises questions on the source of the discrepancy. One potential source is the lack of pathways leading towards ethene, e.g. because of the exclusion of reaction families such as intramolecular 1,4- and 1,5-H-abstraction reactions. It is possible that the tricyclodecyl or other derived radicals that do not lead to ethene may isomerize to other C_{10} radicals that do eventually decompose into ethene. Secondly the accuracy of rate coefficients can be questioned. Sensitivity analysis was used to quantitatively assess the importance of rate coefficients. Normalized sensitivity coefficients S_{ij} for pre-exponential factors A_j of reaction j on the mole fraction of species X_i were calculated using Chemkin-Pro.
Figure 8 shows ethene has a very large positive sensitivity coefficient for the primary decomposition channel through BR1. This channel is the major exo-TCD initiation pathway through which a pool of H atoms and allyl radicals is established. These radicals are the most important moieties that subsequently consume exo-TCD. Not surprisingly, ethene is also sensitive to reactions that influence the branching of tricyclodecyl radicals, such as the consumption of exo-TCD by H yielding TCDR6. These reactions directly influence the fraction of exo-TCD molecules that lead to ethene.

Since 1,3-Cyclopentadiene is slightly overpredicted, it is possible that the deviating ethene concentration can be attributed to small errors on the rate coefficients for the H-abstractions by H from exo-TCD. Moreover, since reverse rate coefficients are calculated through thermodynamic consistency, uncertain estimates of thermochemistry may play a role too.

Furthermore, the overprediction of tricyclo[5.2.1.0^{2,6}]-dec-4-ene is correlated with the underprediction of ethene, as less TCDR4 is decomposing through the competing channel that produces ethene. A major factor of uncertainty is the ring strain in the unsaturated tricyclo[5.2.1.0^{2,6}]decene rings. At the conditions in Scheme 5, benzene is primarily formed via two channels of primary decomposition of exo-TCD, which account for 90% of the total rate of production of benzene. 46% of benzene is formed through the primary decomposition channel starting from TCDR6 through α-H-fulvenyl, cf. Scheme 5. The second important benzene formation pathway starts from TCDR8 and proceeds via a series of rearrangements of C_{8}H_{11}, eventually resulting in benzene and ethyl.
4.3.2 Secondary chemistry and aromatics formation

Many studies of JP-10 pyrolysis indicated the presence of significant quantities of aromatic components, with benzene, toluene, styrene, indene and naphthalene as the most prominent species. The current model suggests that major exo-TCD decomposition pathways lead to hydrogen, ethene, propene, CPD, cyclopentene and ethenylcyclopentenes. These primary products represent the hydrocarbon pool from which secondary products such as aromatics are formed. CPD and the derived CPDyl radical play a central role in the formation of aromatic species. A schematic overview of important secondary pathways is depicted in Scheme 6.

Scheme 6: Pathways of secondary conversion of primary products relevant at \( T=1100\, \text{K}, \, P=1.7\times 10^5\, \text{Pa} \).

At higher exo-TCD conversion H-abstraction reactions convert CPD into CPDyl with allyl and H the most important abstracting radicals. Routes from CPD/CPDyl to indene and naphthalene form the two most important ways of consumption of CPD, accounting both for 30% of the total decomposition rate of CPDyl. Naphthalene formation kinetics were based on Lindstedt et al. It involves a two-step sequence in which CPDyl radicals recombine to the stabilized \( \text{C}_5\text{H}_5\text{C}_5\text{H}_4 \) moiety, which rearranges to naphthalene. Kinetics of the indene formation route by addition of CPDyl to CPD originate from Cavallotti et al. reported at \( P=1.1\times 10^5\, \text{Pa} \).

Secondary conversion of cyclopentene leads to 1,3-cyclopentadiene, with the direct dehydrogenation pathway accounting for ca. 80% of the total cyclopentene decomposition rate.
The conversion of propene produces methyl radical and is the second major source for methyl radicals. The latter recombine with CPDyl yielding 5-methylcyclopentadiene. This C₆H₈ molecule can rearrange into the other methylcyclopentadiene isomers or can undergo H-abstraction resulting in methyl-1,3-cyclopentadien-5-yl and isomeric H-fulvenyl radicals. The latter rearrange via various exit channels to fulvene, 1,3-cyclohexadienyl and eventually benzene. The kinetics and thermochemistry for these reactions and species were taken from Sharma and Green 30.

Another important aspect of the kinetic model at high exo-TCD conversion, is the secondary conversion of ethenylcyclopentenes. The latter produce fulvene and represents an important pathway to α-H-fulvenyl, cf. Scheme 6. Next to the primary decomposition channels from TCDR6 and TCDR8, the hydrogen assisted fulvene isomerization and the decomposition of 5-methyl-1,3-cyclopentadien-5-yl 30 become importance sources of benzene too. A sensitivity analysis of benzene predictions at 1100K is shown in Figure 9 and illustrates again the importance of the unimolecular ring opening reaction of exo-TCD yielding BR1, initiating the chemistry.

Figure 9: Normalized sensitivity coefficients for benzene at T = 1100K, P = 1.7 × 10⁵ Pa, corresponding to an exo-TCD conversion = 36%. “BS” indicates that the species represents is formed through a β-scission reaction of the reactant.

The majority of the reactions shown in Figure 9 were based on estimations by analogy from RMG’s databases. More accurate rate coefficients for these reactions based on ab initio techniques may therefore be valuable.
5 Conclusions

A detailed kinetic model was developed to simulate JP-10 pyrolysis using the RMG. New elements of this model compared to previously developed JP-10 models are the inclusion of more accurate kinetics for initial exo-TCD decomposition chemistry, improved fused ring thermochemistry and the incorporation of many secondary conversion routes to aromatics.

The good agreement between model predictions and five independent sets of experimental data was surprising given the uncertainty in many parameters in the model and without adjustment of any of the model parameters to experimental data. Significant differences exist between the newly calculated CBS-QB3 rate coefficients of the tricyclodecyl radical decomposition reactions with literature values for the rate coefficients of the smaller analogous units of norbornane and cyclopentyl, illustrating that the use of rate coefficients from analogous reactions to model JP-10 decomposition reactions could lead to inaccurate results. It not only indicated the usefulness and validity of the kinetic model over a wide range of conditions but also confirmed the value of RMG as a tool for the kinetic modeling of complex decomposition chemistry of polycyclic hydrocarbons.

Analysis of the kinetic model showed the importance of four important H-abstraction reactions from exo-TCD besides the unimolecular initiation route leading to 3-cyclopentylcyclopentene and indicated that each of these routes lead to primary products such as hydrogen, ethene, propene, 1,3-cyclopentadiene, cyclopentenes, ethenylcyclopentenes and benzene. Next to the primary decomposition reactions, a large part of the model involves detailed kinetics for the secondary conversion of primary products towards aromatics such as toluene, styrene. These components are formed through the secondary conversion of unsaturated five-membered rings such as ethenylcyclopentenes and 1,3-cyclopentadiene-derived moieties. PAHs such as indene and naphthalene are formed by molecular growth reactions of cyclopentadienyl/cyclopentadiene moieties rather than direct dehydrogenation routes from exo-TCD.

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7 Supporting Information

Section S1: Kinetic model in Chemkin format
Section S2: Ab initio reaction barriers at 0K
Section S3: Modified Arrhenius parameter fit

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Section S1 Kinetic model in Chemkin format
Cf. separate text files.

Section S2 Ab initio reaction barriers at 0K

Tricyclodecyl beta-scission reactions

The structure and nomenclature of the tricyclodecyl radicals is shown in Scheme 1.

![Scheme 1](image)

**Scheme 2**: Six possible $C_{10}H_{15}$ tricyclodecyl radicals created by H-abstraction reactions from exo-TCD. Nomenclature for the tricyclodecyl monoradicals is adopted from Hudzik et al. [1].

Structures were optimized with “verytight” convergence criteria and, correspondingly, a very fine integration grid was used (“int=ultrafine”). In the case of TCDR10, there are two minima corresponding to inversion of the radical center; the lower energy conformation was considered here. Saddle points were confirmed to have a single imaginary frequency and the geometries were visually inspected to confirm that the structure was intermediate between the intended reactant and product(s) and intrinsic reaction coordinate calculations were performed to confirm that the saddle point connected the intended reactant and product. Transition state theory calculations were performed using results for the saddle point and corresponding tricyclodecyl reactant.

Reaction barriers at 0K (including zero-point energy effects) are tabulated in Table 1. Decomposition channels for which saddle point could not be successfully located are
indicated in Table 1. The few unsuccessful calculations comprise channels that involve components with double bonds at the bridgehead of the bicycle[2.2.1]heptane ring system, and as a result violate Bredt’s rule [2].

Table 1: reaction barrier at 0K (including zero-point energy effects) for the tricyclodecyl beta-scission reactions.

| Tricyclodecyl radical | \( E_{TS} - E_{reactant} \) (kJ mol\(^{-1}\)) |
|-----------------------|------------------------------------------|
|                       |                                          |
| TCDR4                 |                                          |
| ![Reaction R1](image) | 136                                      |
| ![Reaction R2](image) | 142                                      |
| TCDR5                 |                                          |
| ![Reaction R3](image) | 89                                       |
| ![Reaction R4](image) | 122                                      |
| ![Reaction R5](image) | 133                                      |
| ![Reaction R6](image) | 170                                      |
| ![Reaction R7](image) | 141                                      |
| TCDR6                 |                                          |
| ![Reaction R8](image) | 127                                      |
| ![Reaction R9](image) | 139                                      |
| ![Reaction R10](image)| 127                                      |
| ![Reaction R11](image)| 101                                      |
| Reaction | Chemical Structure | Value |
|----------|--------------------|-------|
| R12     | ![Chemical Structure](image1) | 126   |
| R13     | ![Chemical Structure](image2) | N/A   |
| R14     | ![Chemical Structure](image3) | 178   |
| R15     | ![Chemical Structure](image4) | 159   |

| Reaction | Chemical Structure | Value |
|----------|--------------------|-------|
| R16     | ![Chemical Structure](image5) | 104   |
| R17     | ![Chemical Structure](image6) | 79    |
| R18     | ![Chemical Structure](image7) | N/A   |
| R19     | ![Chemical Structure](image8) | 97    |
| R20     | ![Chemical Structure](image9) | N/A   |
| R21     | ![Chemical Structure](image10) | N/A   |
| R22     | ![Chemical Structure](image11) | N/A   |

| Reaction | Chemical Structure | Value |
|----------|--------------------|-------|
| R23     | ![Chemical Structure](image12) | 98    |
| R24     | ![Chemical Structure](image13) | 114   |
| Reaction | Reaction Equation | Page |
|----------|------------------|------|
| R25     | $\text{Comp} \overset{\text{Cr}^*}{\rightleftharpoons} \text{Comp}$ | 141  |
| R26     | $\text{Comp} \overset{\text{Cr}^*}{\rightarrow} \text{Comp}^+ + H^*$ | N/A  |
| R27     | $\text{Comp} \overset{\text{Cr}^*}{\rightarrow} \text{Comp}^+ + H^*$ | 160  |
| R28     | $\text{Comp} \overset{\text{Cr}^*}{\rightleftharpoons} \text{Comp}'$ | 102  |
| R29     | $\text{Comp} \overset{\text{Cr}^*}{\rightleftharpoons} \text{Comp}'$ | 103  |

Exo-TCD initiation reactions and intramolecular H-abstraction reactions
Table 2: reaction barrier at 0K (including zero-point energy effects) for exo-TCD initiation reactions and intramolecular H-abstraction reactions.

| Reaction                  | $E_a$ | $E_{TS} - E_{reactant}$ |
|---------------------------|-------|-------------------------|
| **Exo-TCD initiation reactions** |       |                         |
| Exo-TCD = BR1             | 326   | CAS-PT2                 |
| Exo-TCD = BR1             | 319   | UB3LYP/CBS-QB3          |
| Exo-TCD = BR1             | 339\* | CR-CC(2,3)/CASSCF(4,4) ref. [3] |
| Exo-TCD = BR2c\*         | 333   | CAS-PT2                 |
| Exo-TCD = BR2d\*         | 325   | CAS-PT2                 |
| Exo-TCD = BR2c\*         | 328   | UB3LYP/CBS-QB3          |
| Exo-TCD = BR2d\*         | 321   | UB3LYP/CBS-QB3          |
| Exo-TCD = BR3             | 335   | CAS-PT2                 |
| Exo-TCD = BR3             | 334   | UB3LYP/CBS-QB3          |
| Exo-TCD = BR4             | 334   | UB3LYP/CBS-QB3          |
| Exo-TCD = MA110           | 416   | UB3LYP/CBS-QB3          |
| exo-TCD = MA111           | 336   | UB3LYP/CBS-QB3          |
| **Intramolecular H-abstraction reactions** |       |                         |
| BR1 = MA110               | -1    | CAS-PT2                 |
| BR1 = MA110               | 15    | UB3LYP/CBS-QB3          |
| BR1 = MA110               | 0     | CR-CC(2,3)/CASSCF(4,4) ref. [3] |
| BR2 = BR2\_1             | 20    | UB3LYP/CBS-QB3          |
| BR3 = BR3\_1             | 23    | UB3LYP/CBS-QB3          |

# based on the 0 K (including ZPE) bond energy associated with carbon–carbon bond breaking to form the BR1 biradical associated with TS1A geometry.

**Section S3: Modified Arrhenius parameter fit using Cantherm**

Rate coefficients $k$ are obtained using the software package CanTherm [4] using the results of the ab initio calculations. They are expressed as a function of temperatures through a so-called modified Arrhenius expressions, involving three parameters, $A$, $n$, $E_a$:

$$k_{fit} = A T^n \exp\left(-\frac{E_a}{RT}\right)$$

The modified Arrhenius parameters are obtained through a least-squares fit using calculated rate coefficients $k_{training}$, at ten equidistant temperatures between 298 - 3000K. The value of $k_{training}$ is obtained by multiplying the ab initio contribution $k_{AI}$ at temperature $T$ and the Eckart tunneling coefficient $\kappa$ at temperature $T$:

$$k_{training}(T) = \kappa(T) \cdot k_{AI}(T)$$
In general, the uncertainties associated with the parameter fitting are significantly smaller than the uncertainties associated with the level-of-theory of the employed ab initio methods. For example, uncertainties of CBS-QB3 derived reaction barriers typically amount to 4-8 kJ mol\(^{-1}\). Statistical analysis has shown that the 95\% confidence interval of the E\(_a\) parameter in the modified Arrhenius expression typically amounts to ± 0.05 kJ mol\(^{-1}\), for the A parameter a multiplicative uncertainty of 10\% and for the n parameter an uncertainty of ±0.01. Nevertheless, it should also be noted that for reactions with substantial tunneling coefficients at lower temperatures (< 500K) a less optimal correspondence between the value of the fitted rate coefficient k\(_{\text{fit}}\) and the rate coefficient k\(_{\text{training}}\) from the training set can be expected.

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