

Theoretical Gas Phase Chemical Kinetics

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Training in theoretical chemical kinetics

1. CEFRC – Summer School

Ab Initio Theoretical Chemical Kinetics

2. Peng Zhang (Ed Law graduate student; now a CEFRC Fellow)

Monomethylhydrazine decomposition

3. Franklin Goldsmith (Bill Green graduate student)

Allyl + HO₂

4. Mike Burke (Fred Dryer & Yiguang Ju graduate student)

H₂/O₂ Combustion

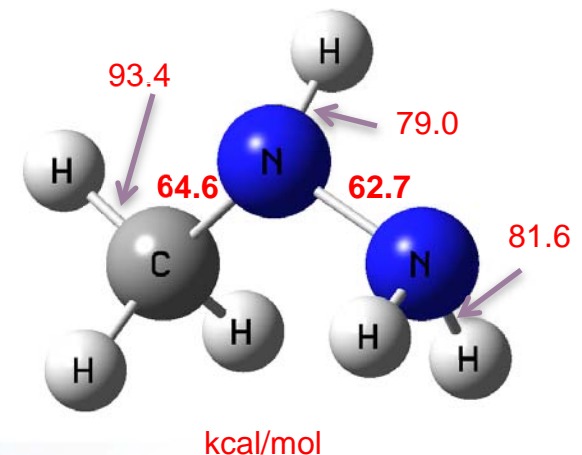


Monomethyl hydrazine decomposition

- Monomethylhydrazine (MMH)
 - Diamine-based rocket fuel
 - Commonly used as bipropellant and monopropellant
 - Exothermically decomposes upon contact with either a hot surface or an oxidizer
- Thermal decomposition of MMH
 - Fuel stability and storability
 - Necessary components of mechanism of MMH oxidation
 - Have not been sufficiently studied
- Dominant reactions of MMH decomposition: NN and CN bond fission



- Barrierless radical-radical recombination reactions



Potential Energy Surface

□ On-the-fly Calculation of Interaction Potential

– aug-cc-pVTZ: accurate but too computationally demanding → cc-pVDZ and aug-cc-pVDZ were used

– Reacting fragments have fixed internal geometries

□ Orientation-Independent Corrections

– One-dimensional basis set correction

– One-dimensional correction for geometric relaxation

Basis set Separation of fragments Relative orientation Reacting fragments have fixed internal geometries

$$V = V_{\text{CASPT2/ADZ}}(R, \Omega; \text{rigid})$$
$$+ \left[V_{\text{CASPT2/ATZ}}(R, \Omega^*; \text{rigid}) - V_{\text{CASPT2/ADZ}}(R, \Omega^*; \text{rigid}) \right]$$

basis set correction from CASPT2/aug-cc-pVTZ

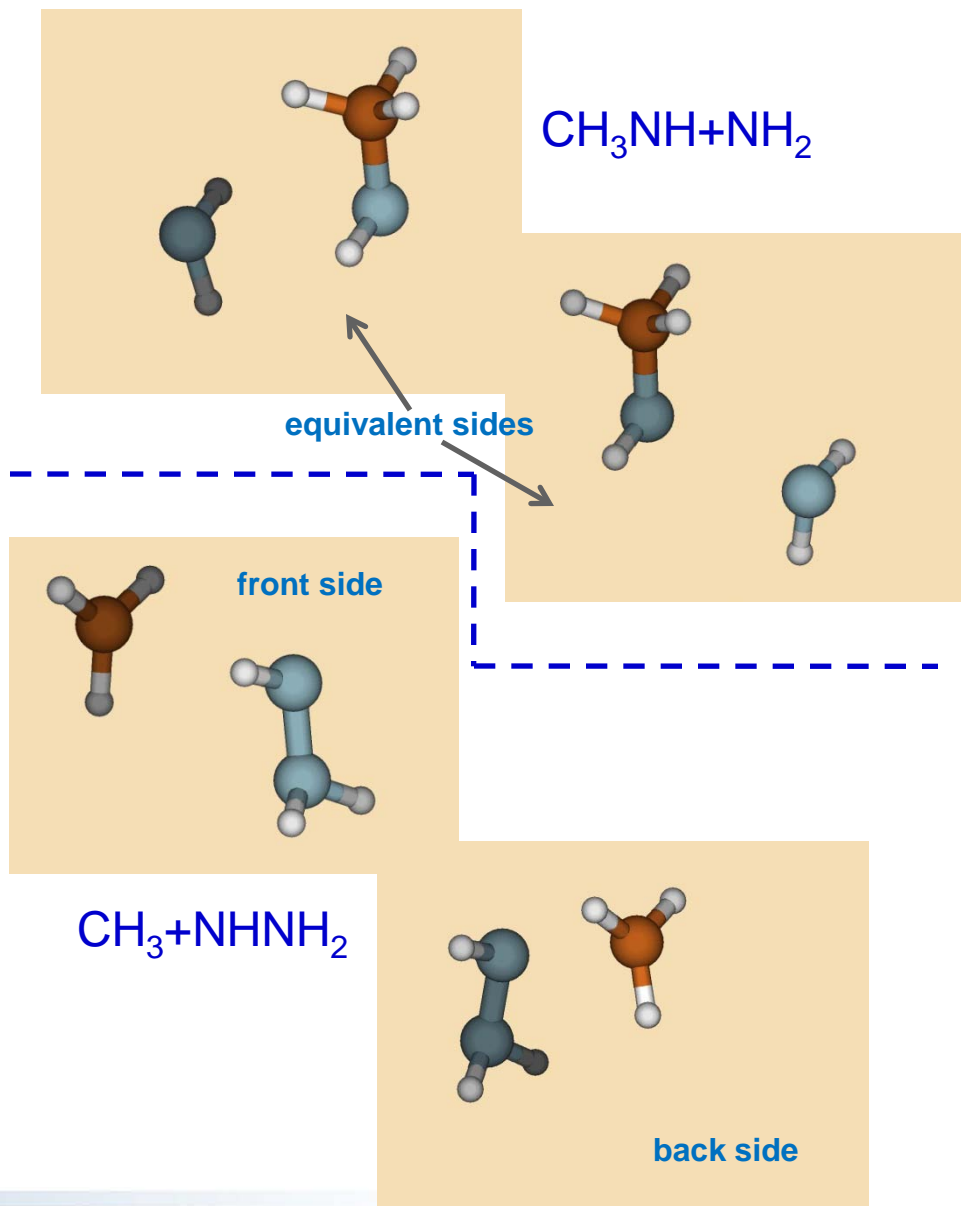
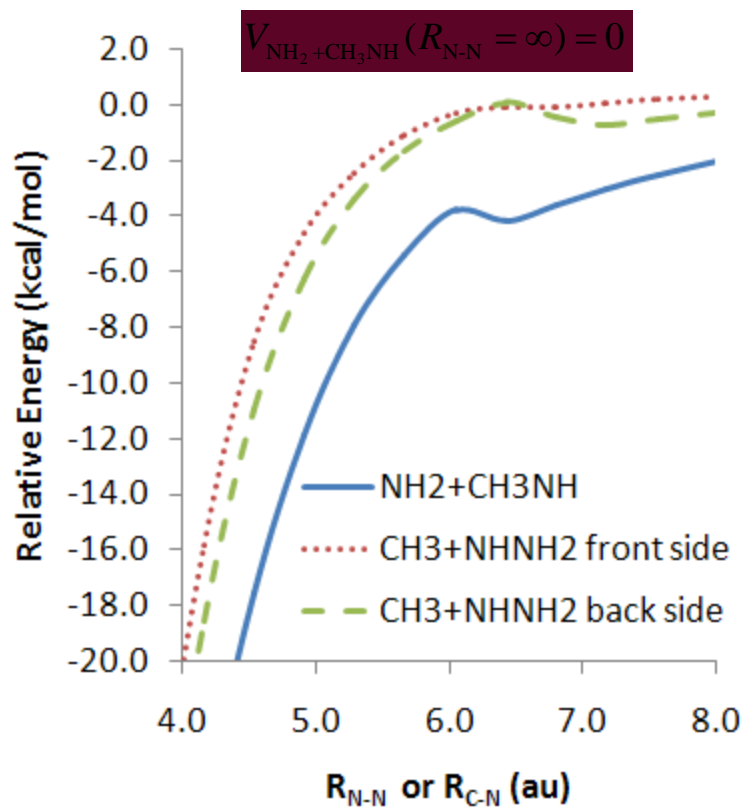
$$+ \left[V_{\text{CASPT2/ADZ}}(R, \Omega; \text{relaxed}) - V_{\text{CASPT2/ADZ}}(R, \Omega; \text{rigid}) \right]$$

correction for relaxing of the internal structure

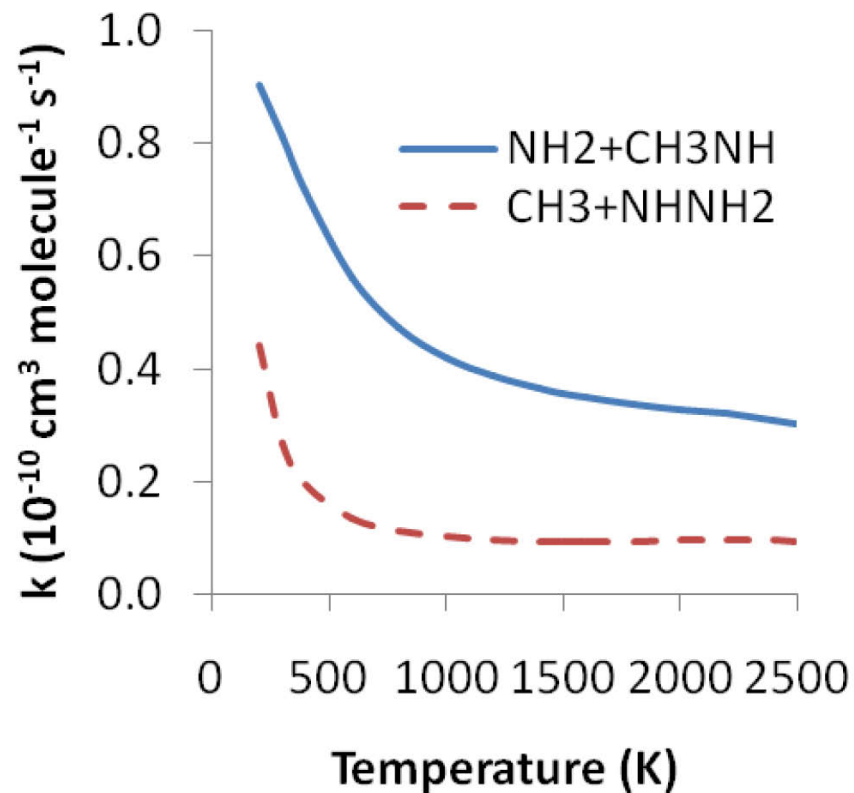


Potential Energy Surface

Potential curves



VRC-TST Capture Rates

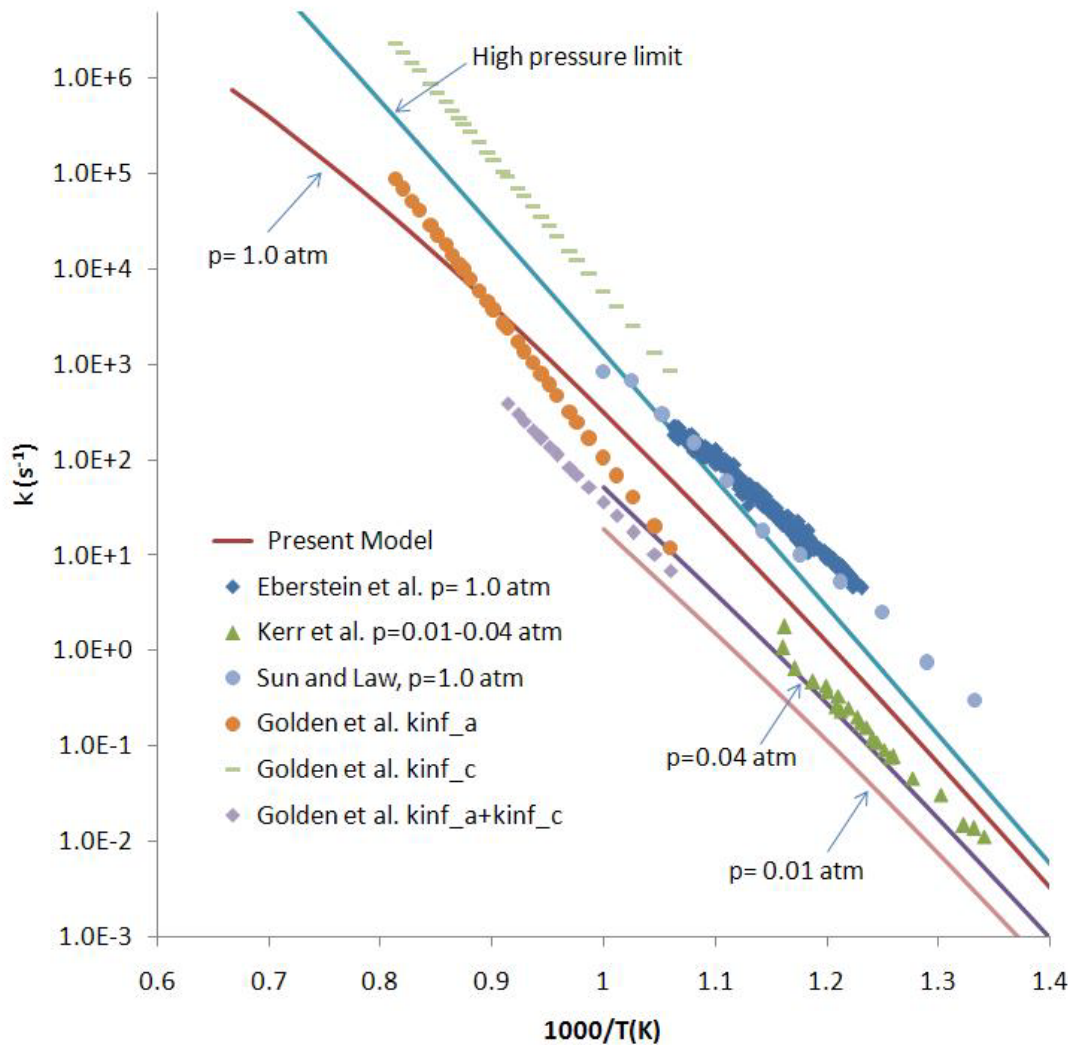


$$k_{-1}^{\infty} = 8.34 \times 10^{-10} T^{-0.429} \exp(20.1/T) \text{ cm}^3 \cdot \text{molecule}^{-1} \cdot \text{s}^{-1}$$

$$k_{-2}^{\infty} = 3.99 \times 10^{-12} T^{0.085} \exp(404.2/T) \text{ cm}^3 \cdot \text{molecule}^{-1} \cdot \text{s}^{-1}$$



MMH Dissociation: Theory-Experiment Comparison



- Kerr et al (1963): the first-order rate coefficient for the homogeneous dissociation of NN bond of MMH.

- Very good agreement with the present theory

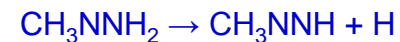
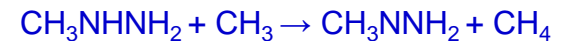
- Golden et al (1972): very low pressure reactor → complicated by gas-wall interactions

- Not appropriate for a direct comparison with the theory

- Eberstein et al (1965): total thermal decomposition rate of MMH

- Modeled by a mechanism containing 43 species and 160 reactions (Sun and Law 1007)

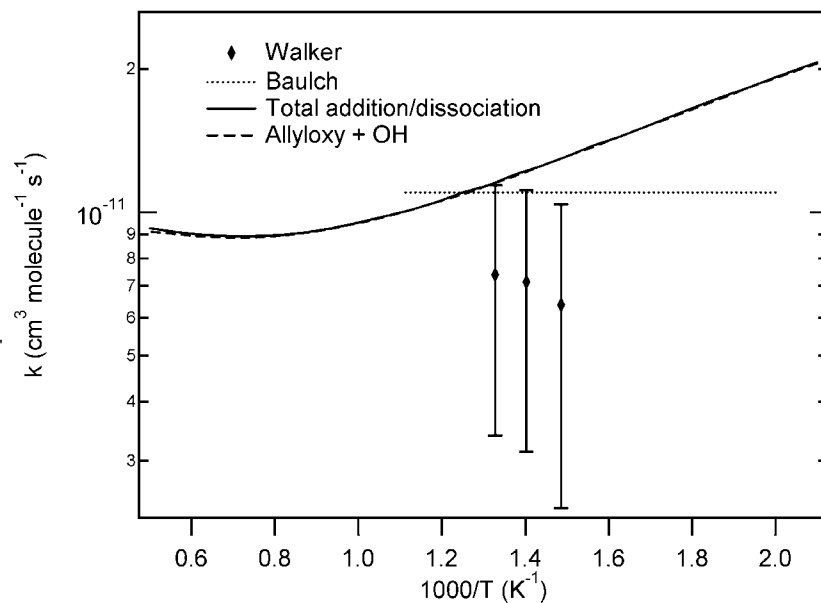
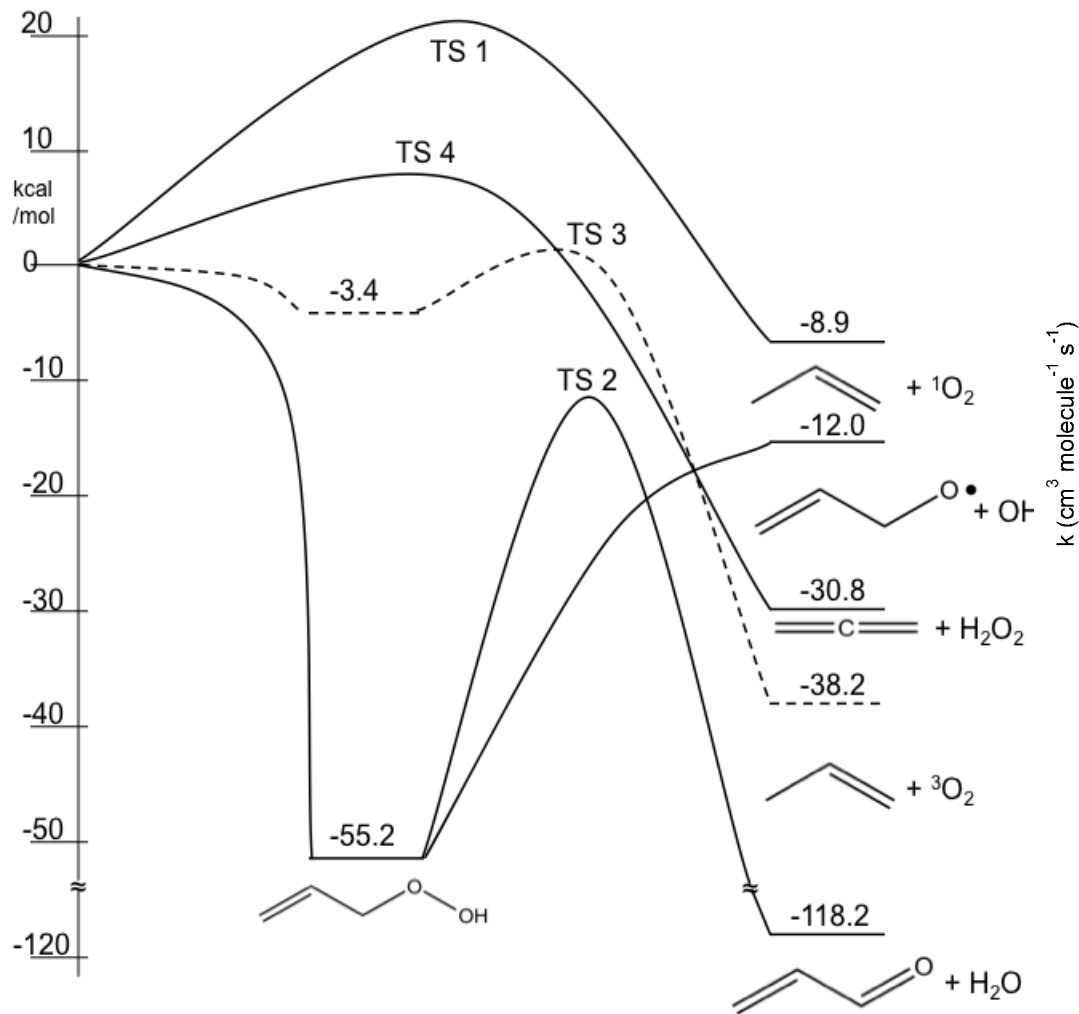
- Sensitivity analysis identified two qualitatively important reaction channels:



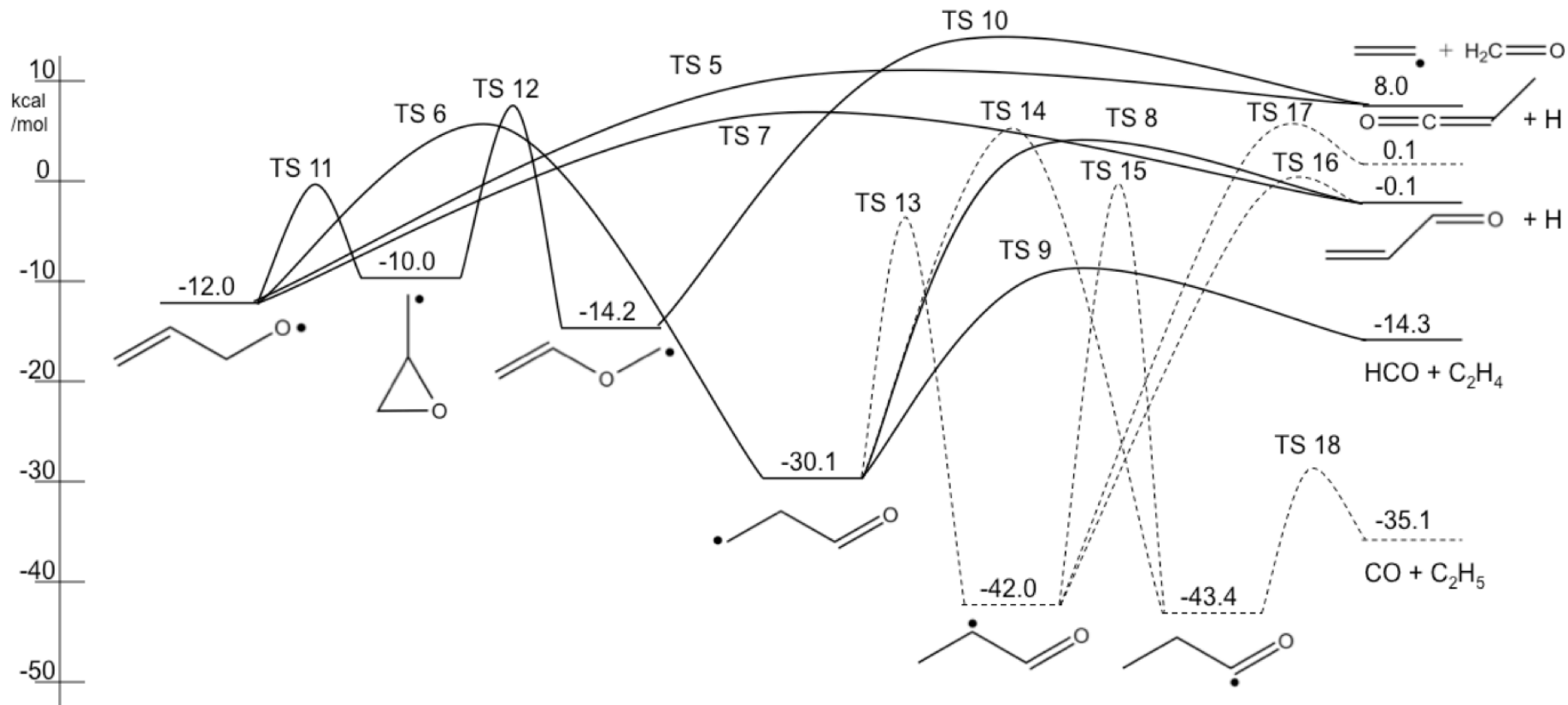
- Disagreement with the theory might be due to the absence of the two reaction channels:



Allyl + HO2



Allyloxy Decomposition



Allyloxy Decomposition

