

# Chemical Kinetics and Thermal Properties of Ablator Pyrolysis Products during Atmospheric Entry

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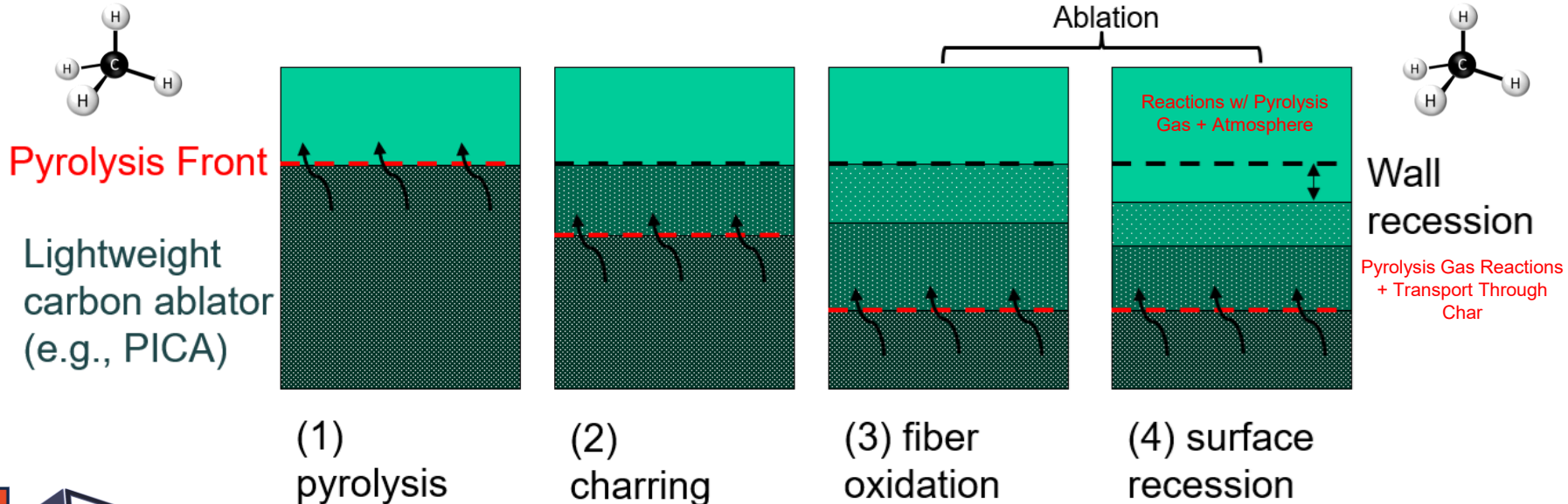
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# Background – Thermal Protection System Modeling

- Interest in modeling thermochemical response of ablative TPS materials
  - Focus: Modeling of reacting pyrolysis gas in char layer; Surface reactions with pyrolysis gas + atmospheric species
  - Interests: **thermal transport, gas-phase chemistry**



# Modeling Ablator Energy Balance

- **Pyrolysis gas enthalpy is critical**
- Different composition -> Different thermal response

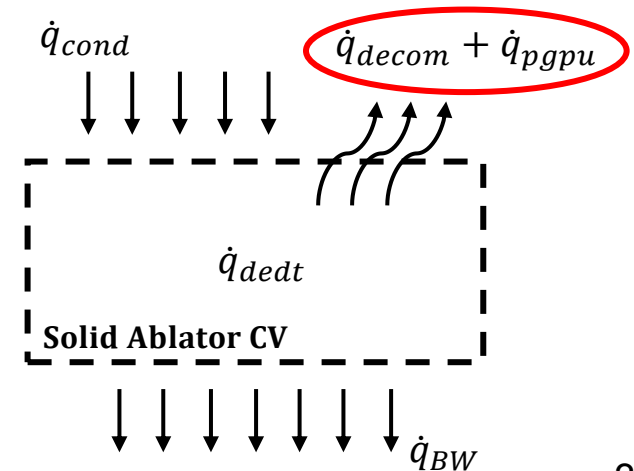
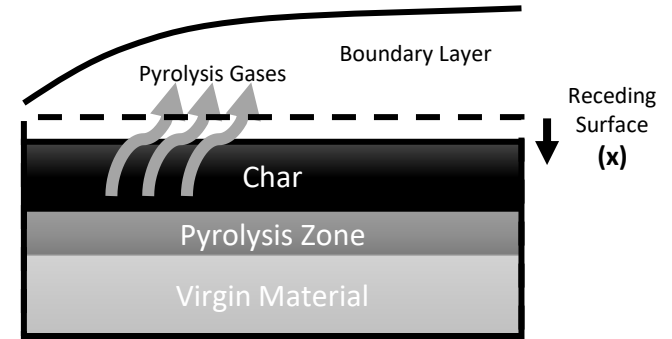
## ➤ Thermal Qols:

- $\dot{q}_{decom}$  = Heat of Decomposition

$$\dot{q}_{decom} = h_{decom} \dot{w}_g = (h_{g,\omega} - \bar{h}) \dot{w}_g$$

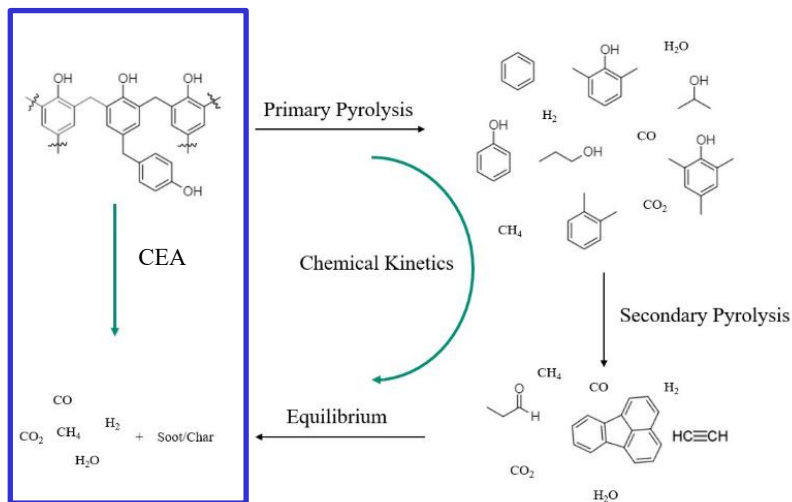
- $\dot{q}_{pgpu}$  = Pyrolysis Gas Pick Up (Transpiration Cooling)

$$\dot{q}_{pgpu} = -\frac{\partial h_g}{\partial x} \bar{m}_g'' = (h_{g,o} - h_{g,\omega}) \dot{w}_g + \frac{h_{g,o} - h_{g,i}}{\Delta x} \dot{m}_{g,i}''$$



# Pyrolysis Gas Chemistry – Legacy Assumption

- In modeling of pyrolysis gas in ablation codes, two assumptions are common:
  - **Equilibrium Chemical Composition**
  - **Uniform Elemental Composition**



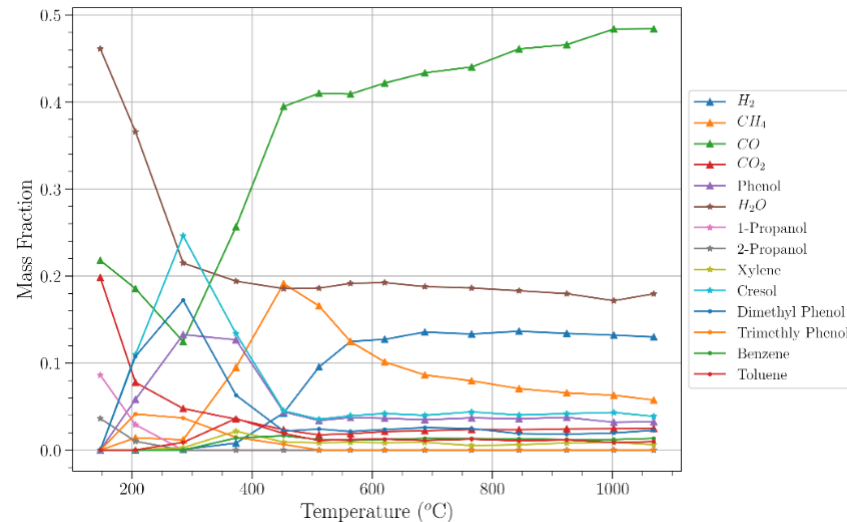
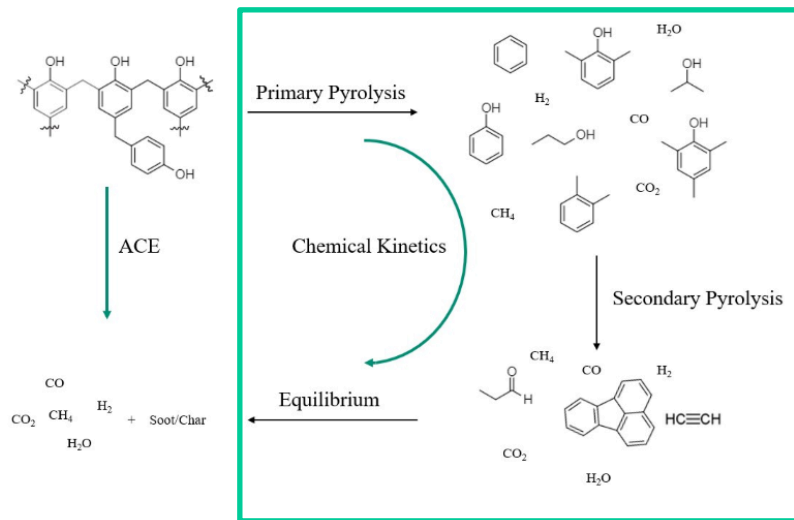
➤ **Advantage: Significantly easier to simulate in Material Response Codes**



# Pyrolysis Gas Chemistry – Legacy Assumption

## ➤ Is this assumption accurate?

- Experimental results show large variation in Primary Pyrolysis elemental composition
- **We will examine the kinetics of these species with a coupled simulation approach**



Source: Brody K. Bessire and Timothy K. Minton. Decomposition of phenolic impregnated carbon ablator (pica) as a function of temperature and heating rate. *ACS Applied Materials & Interfaces*, 9(25):21422–21437, 201

# Analysis – Coupled Aria/Cantera Simulation

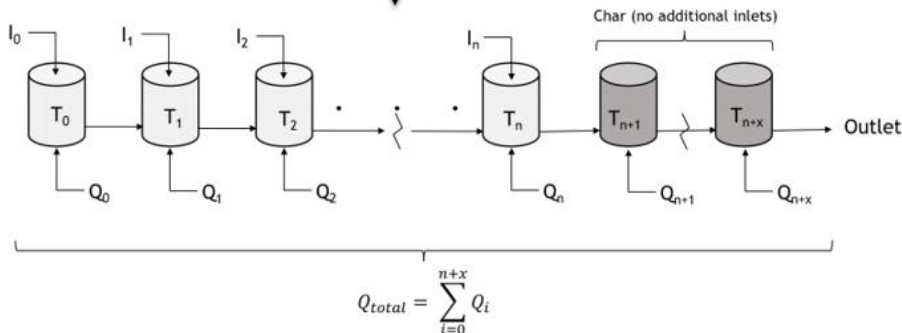
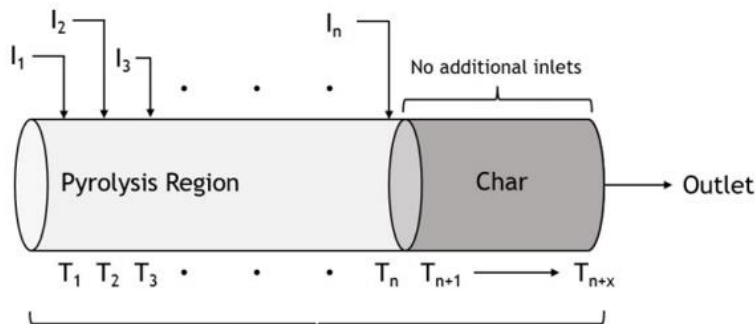
## Material Response Code (Aria)

- Temperature/Pressure
- Mass Flow Rate
- Primary Pyrolysis Rate
- Recession

Update Every  
Timestep

## Chemistry Solver (Cantera)

- Pyrolysis Speciation
- Finite-Rate Kinetics
- Species/Element Mole Fractions
- Thermal Properties



## Aria Model Assumptions

- TACOT Ablator Material
- Equilibrium Chemistry
- Constant Element Comp.
- Legacy Pyrolysis Model
  - Goldstein

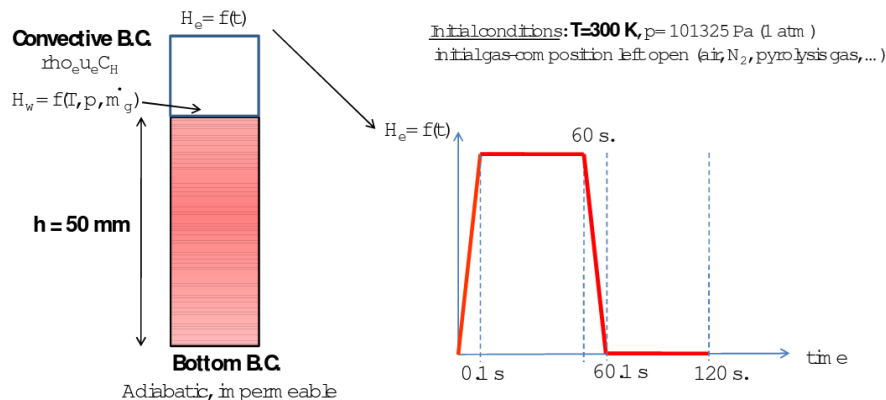
## Cantera Model Assumptions

- Neglect porous flow, GSI effects
  - No coking
- Neglect Transport
- Pyrolysis Speciation
  - Bessire-Minton (6.1 C°/s) Data
- Kinetics
  - Blanquart Mech.
  - 148 species, 1600+ reactions



# Aria – Ablation Workshop Test Cases 2.2 & 2.3

- Aria – Galerkin finite element porous flow solver
- Part of Sandia's SIERRA Multitool
- Used to run Lachaud's 1D Ablation Test Cases using the TACOT Material:



| Case | Time, s | $T_\infty$ , K | $T_{surf}$ , K | $h \rho_e u_e C_H$ , $W m^{-2}$ |
|------|---------|----------------|----------------|---------------------------------|
| 1    | 120.1   | 300            | 3200           | $7.5 \times 10^6$               |
| 2    | 120.1   | 300            | 1600           | $4.5 \times 10^5$               |

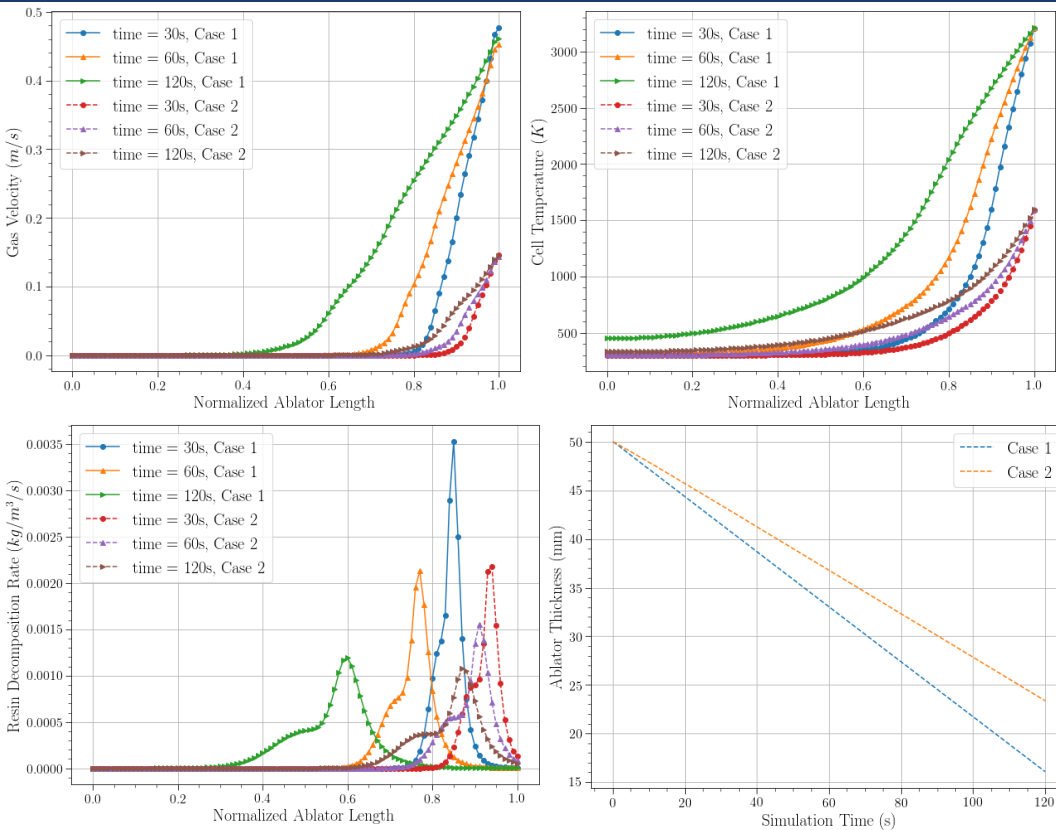


Source: "Sierra Multimechanics module: Aria User Manual - Version 5.8.," 2022.

Source: Lachaud, Jean R.; Martin, Alexandre; van Eekelen, Tom; and Cozmuta, Ioana, "Ablation Test-Case Series #2. Numerical Simulation of Ablative-Material Response: Code and Model Comparisons" (2012).



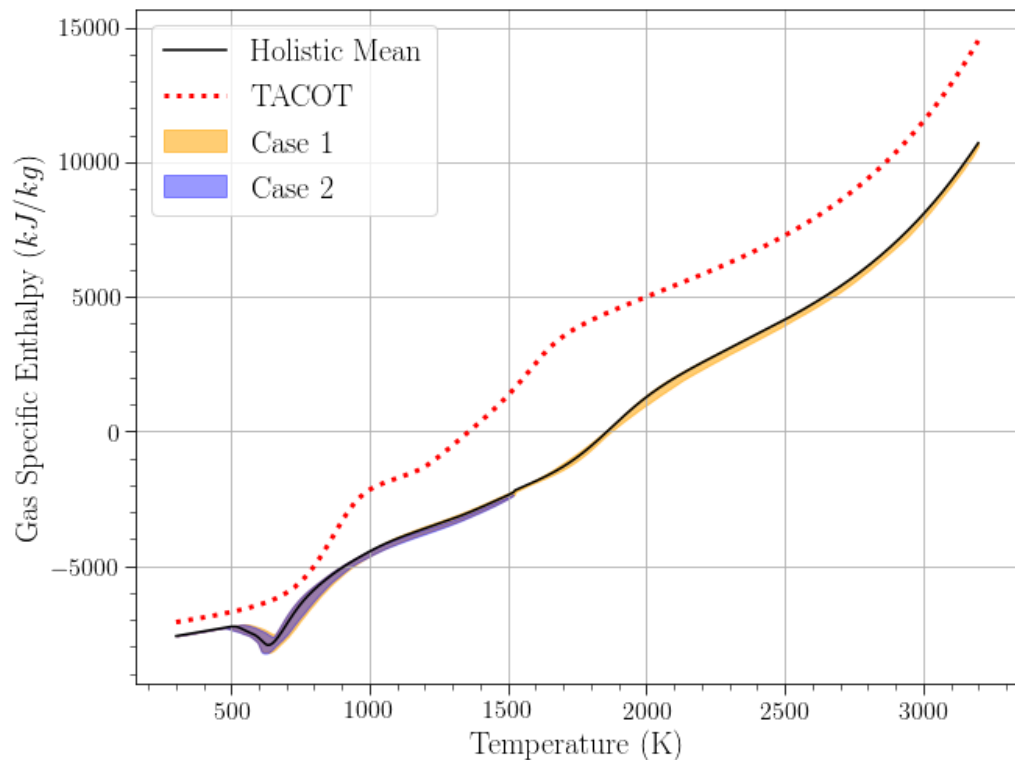
# Aria Simulation Results



- Resulting temperature and recession profiles are nominal to published PATO results of Lachaud
- Results are then used to run Cantera reactor network



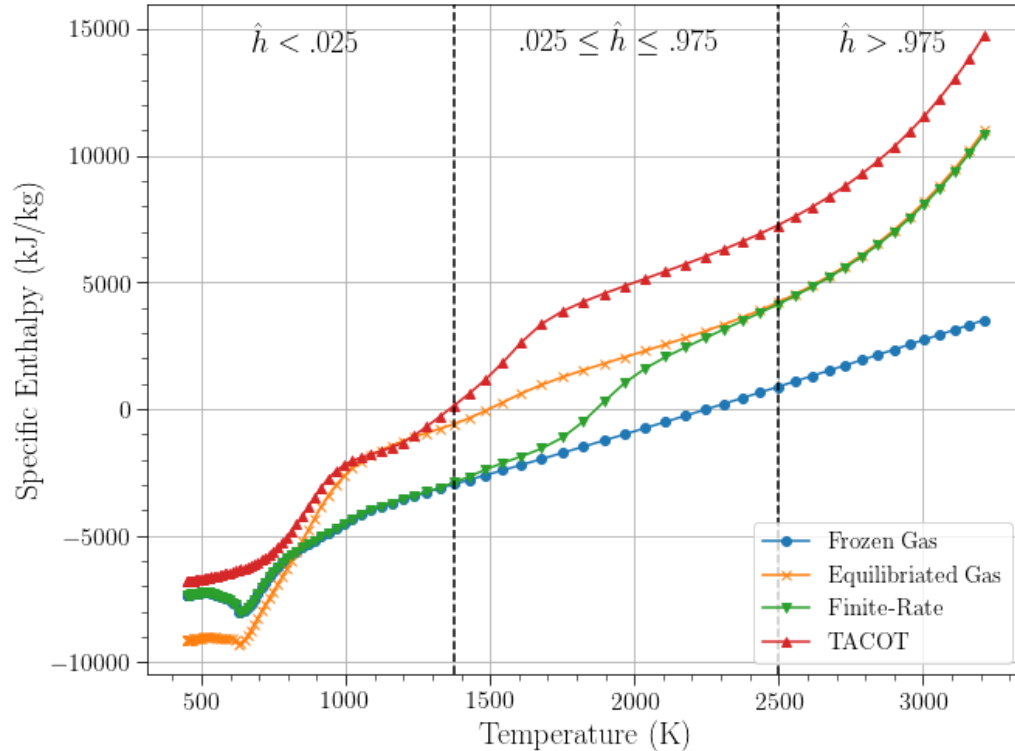
# Holistic Cantera Results – Gas Enthalpy over T & Time



- Shaded regions are bounded by recorded maximum and minimum values.
- Little variation is seen transiently or between the two test cases.



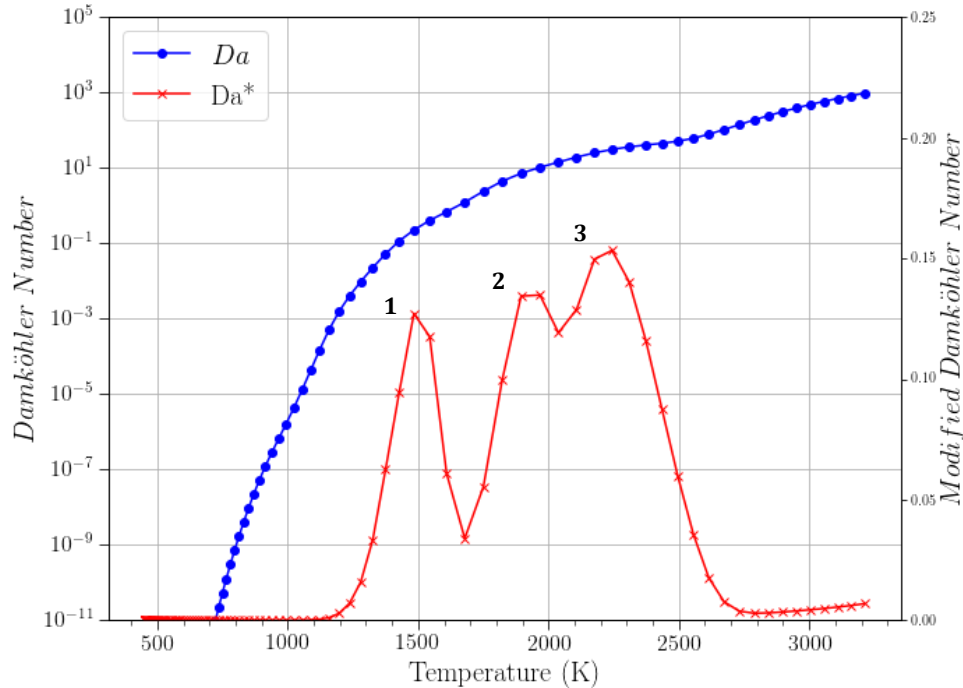
# Case 1 – T = 120 seconds



$$\hat{h} = \frac{|h - h_{frzn}|}{|h_{equil} - h_{frzn}|}$$



# Case 1 – T = 120 seconds



$$Da = \frac{\text{Rate of Reaction}}{\text{Rate of Advection}} = \frac{\tau_R \sum_{i=0}^n r_i^+}{\rho}$$

$$Da^* = \frac{\tau_R \sum_{i=0}^n (r_i^+ - r_i^-)}{\rho}$$

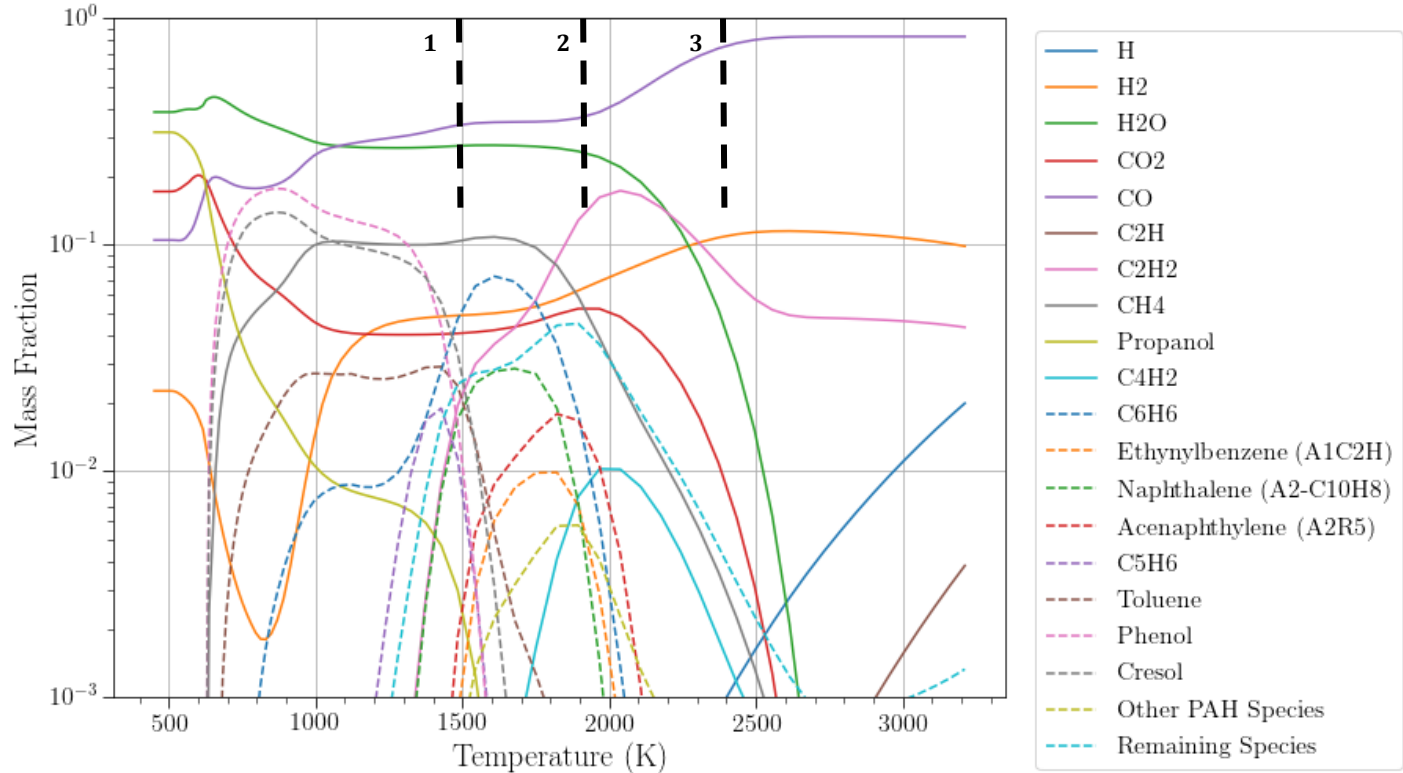
- The standard Damköhler Number shows the onset of chemical kinetics, but not equilibrium
- The modified  $Da^*$  shows finite-rate/equilibrium transition



# Case 1 – T = 120 seconds

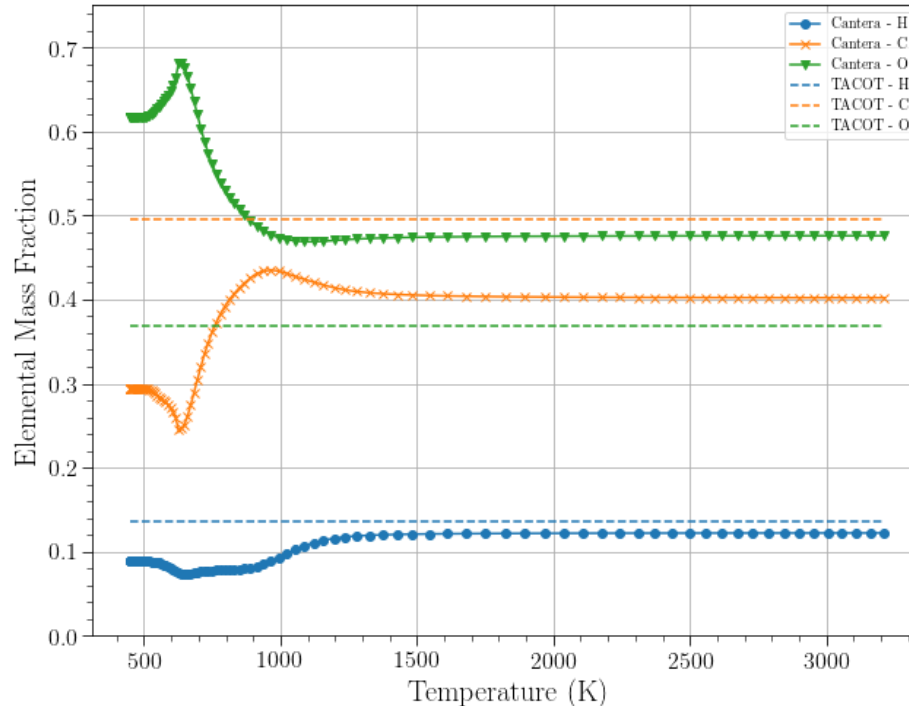
➤ Strong variations in chemical composition seen, only approaching equilibrium at very high temperatures; PAH peaks at around 5% mass fraction

Low Temperature +  
Low Free Oxygen =  
**Non-Equilibrium**



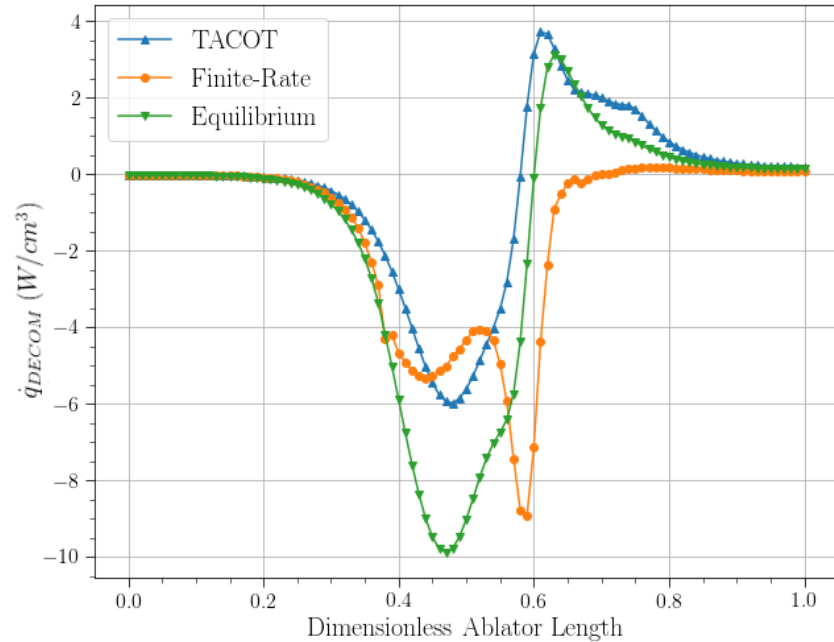
# Case 1 – T = 120 seconds

- Elemental composition is seen to stabilize at around 1400K, once primary pyrolysis is mostly completed
- **Molar Ratios: 18.2% carbon, 16.1% oxygen, 65.7% hydrogen**



# $\dot{q}_{decom}$

- Both cases are similar, so only **Case 1** is shown here
- As expected, equilibrium and TACOT results behave similarly
- Finite-rate results show exothermic behavior for both reaction peaks

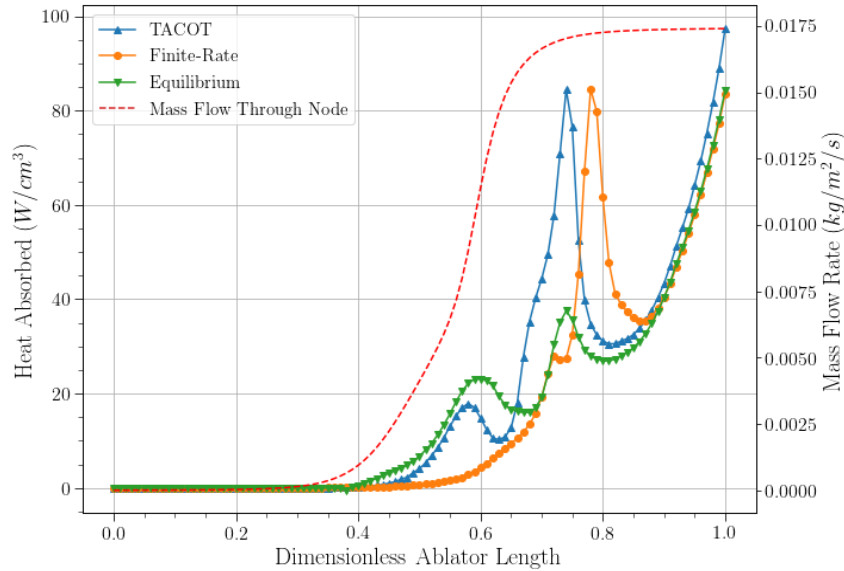


$$\dot{q}_{decom} = h_{decom}\dot{\omega}_g = (h_{g,\omega} - \bar{h})$$

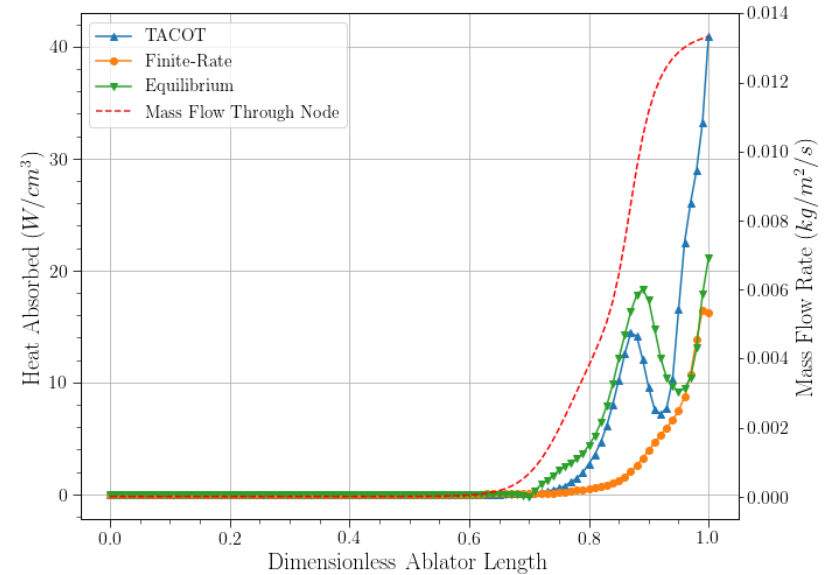


$$\dot{q}_{pgpu}$$

Case 1



Case 2



$$\dot{q}_{pgpu} = -\frac{\partial h_g}{\partial x} \bar{m}_g'' = (h_{g,o} - h_{g,\omega}) \dot{\omega}_g + \frac{h_{g,o} - h_{g,i}}{\Delta x} \dot{m}_{g,i}''$$

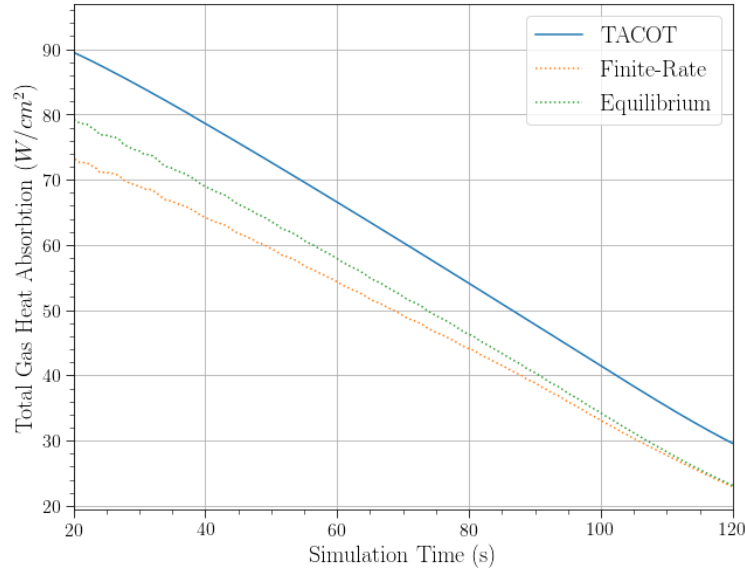


# Total Heat Transfer: Case 1 vs 2

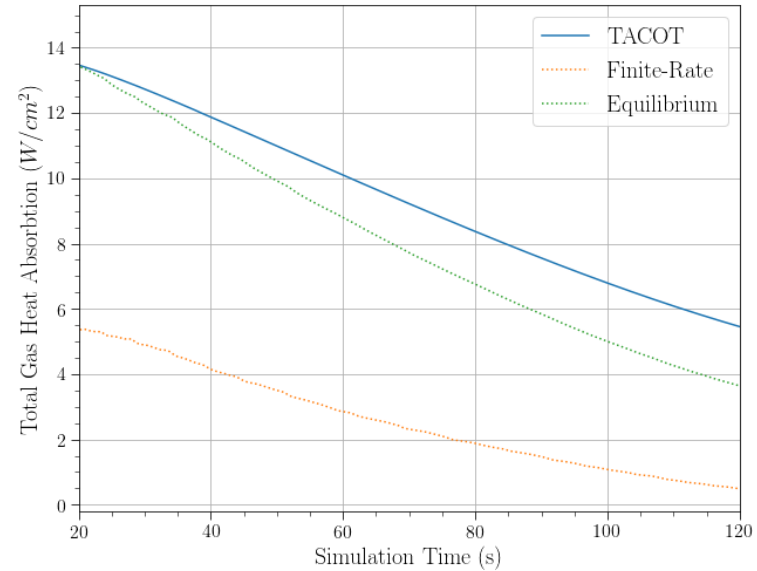
Previous curves integrated over length vs simulation time.

➤ **Typical surface aeroheating values:  $\sim 600 \text{ W/cm}^2$  (1),  $\sim 40 \text{ W/cm}^2$  (2)**

## Case 1



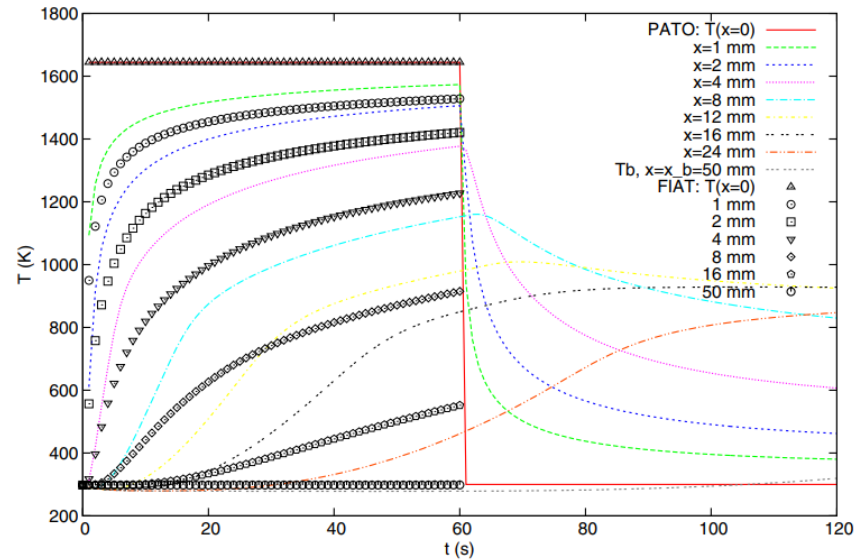
## Case 2



# Conclusion

- Non-equilibrium pyrolysis gas conditions in ablator is common <2500K
- Differences between Eq. and Finite-rate gas cause substantial deviations in thermal Qols
- These Qols potentially have substantial impact on Ablator Energy Balance
  - See Lachaud et al ->

Finite-Rate; 22-species reduced Blanquart mech.



# Acknowledgements

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## Questions?



# Backup – Batch Reactor Tests

- Used to validate achievement of equilibrium with Blanquart mechanism & identify relevant timescales for chemical kinetics
- Initial mixture: Gas composition taken from Minton 25 C/s experiment at various  $T_s$

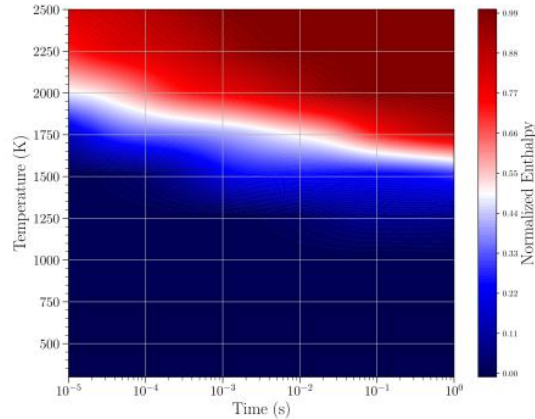


Figure 4-3. Normalized enthalpy of pyrolysis gases with respect to time. Species correspond to Bessire and Minton data at 147.5 °C.

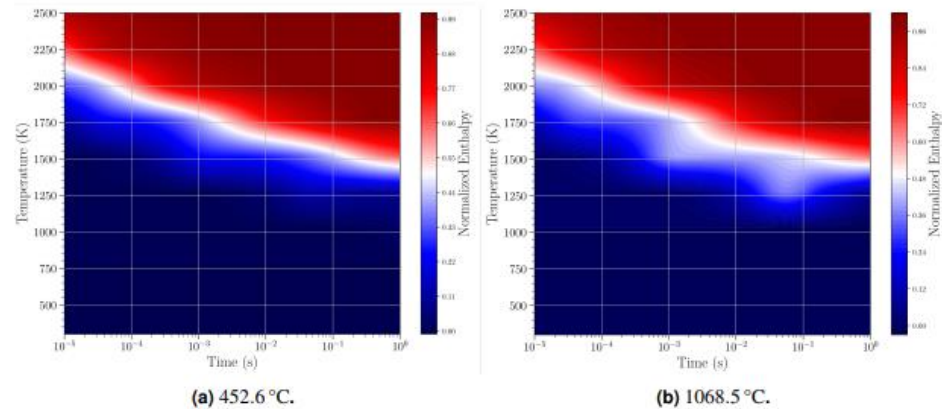


Figure 4-4. Normalized enthalpy of pyrolysis gases with respect to time. Species correspond to Bessire and Minton data at 452.6 °C and 1068.5 °C

