

Review of Condensed-Phase Reaction Kinetics

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There are Four Distinct Stages of Coal Conversion Chemistry

Devolatilization

- Source of all gaseous fuels and soot.
- Determines char yield, size, structure, and initial reactivity.

Volatiles Conversion

- Conversion of tars into soot
- Major heat source.
- Partial combustion of primary volatiles.
- Major source of CO, H₂, CO₂, and H₂O.
- Shifting/reforming chemistry throughout.

Char/Soot Oxidation

- Major heat source.
- Determines residual char yield for gasification.
- Some flyash production.

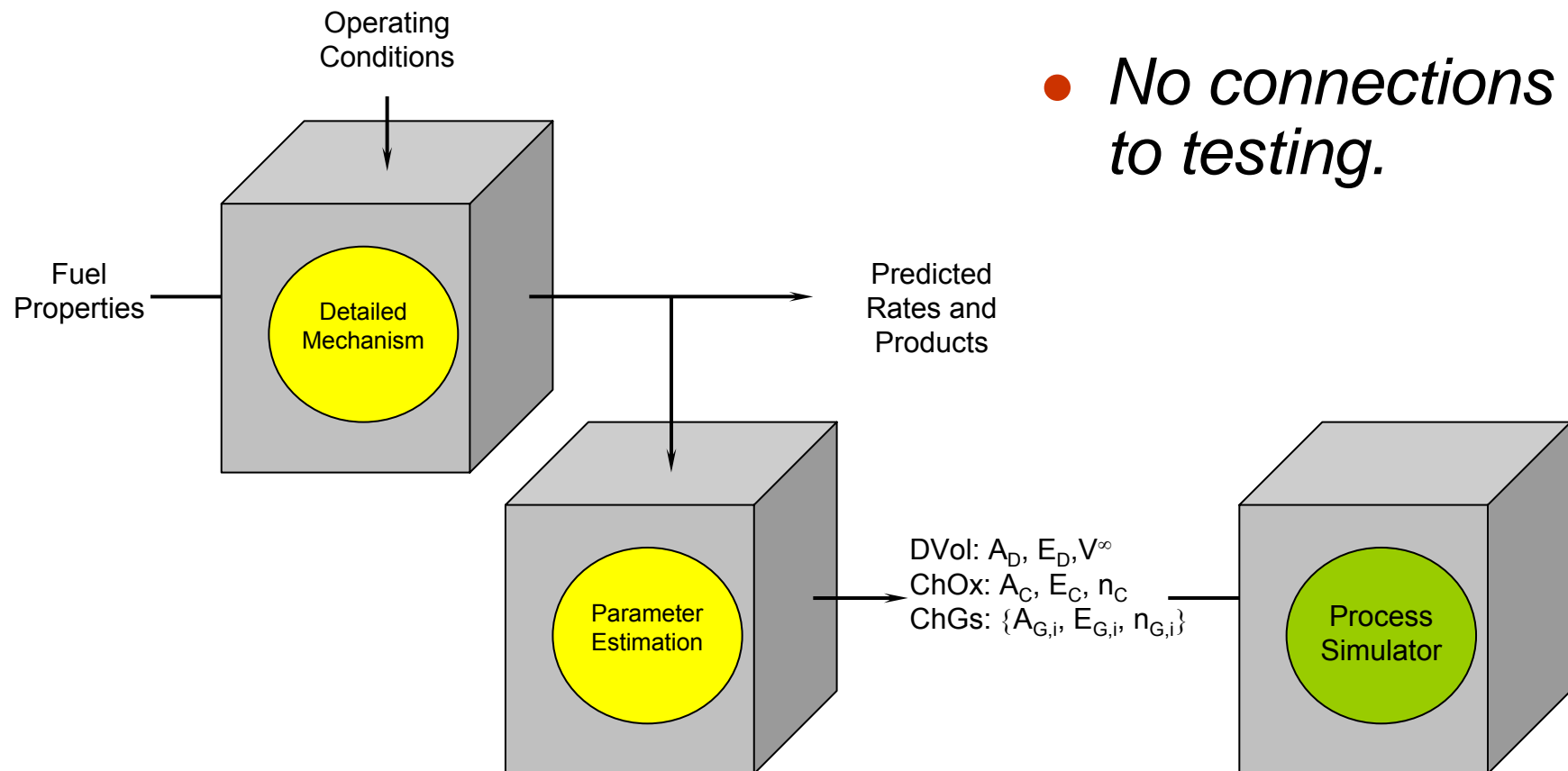
Char/Soot Gasification

- Determines overall conversion.
- Flyash production, via char particle fragmentation + ash agglomeration.

Forget “Understanding” and Focus On *Accuracy* in Applications

- *Fuel Science **SHOULD** specify all the rate parameters used in process simulations (CFD, AspenPlus, HySys, etc.)*
- *Simulation practitioners should **NOT** have to comb literature or resort to default values.*

All Rate Parameters Should Be Assigned From Readily Available Fuel Property Input

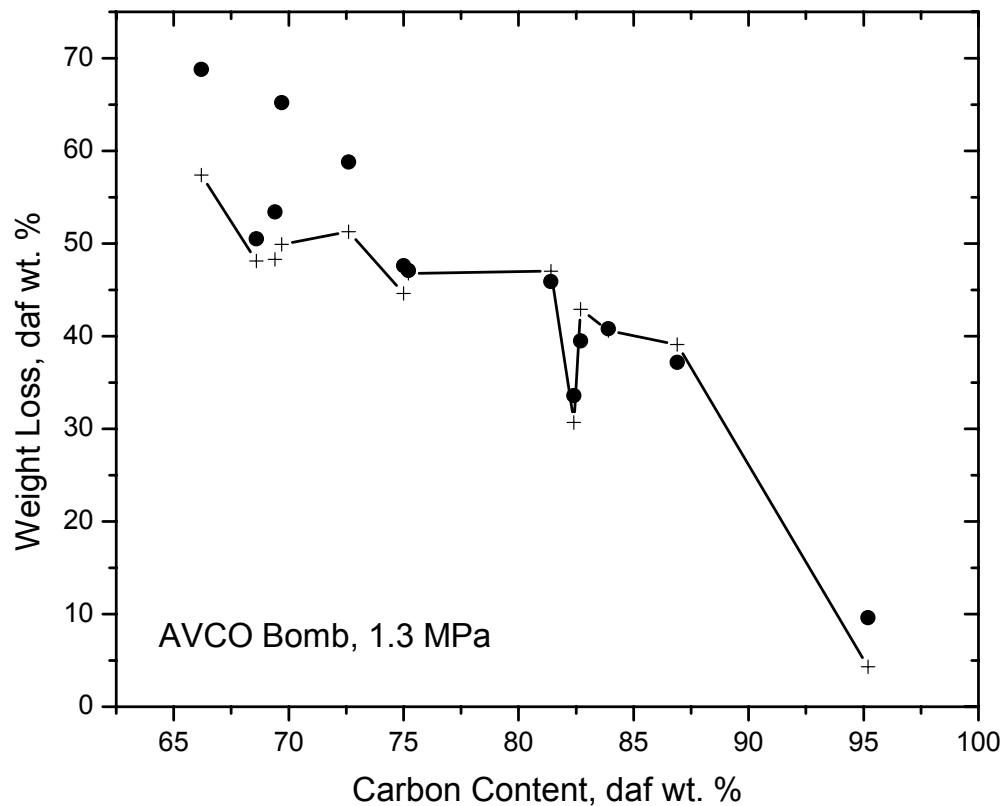


- *No connections to testing.*

FLASHCHAIN[®] for Devolatilization

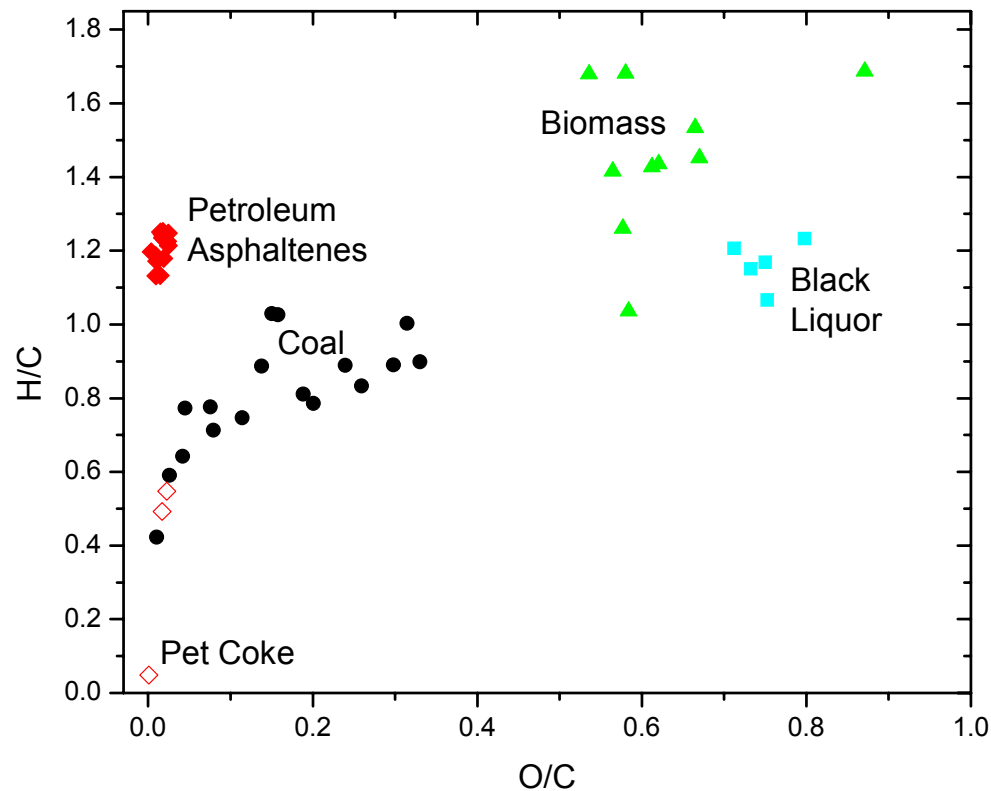
- *FLASHCHAIN[®] was recently validated against a database of 332 independent tests involving 99 coals and broad ranges of heating rates, temperatures, and pressures to 16.7 MPa.*
- *Predicts the **complete distribution** of all volatile products plus tar and char properties.*
- ***Proximate and ultimate analyses** are the only sample-specific fuel properties.*
- *Already used to predict the devolatilization behavior of over 2000 coals.*
- *Versions available for any coal, biomass, pet coke, black liquor, and petroleum asphaltenes.*

Accurate Predictions for Any Coal Type



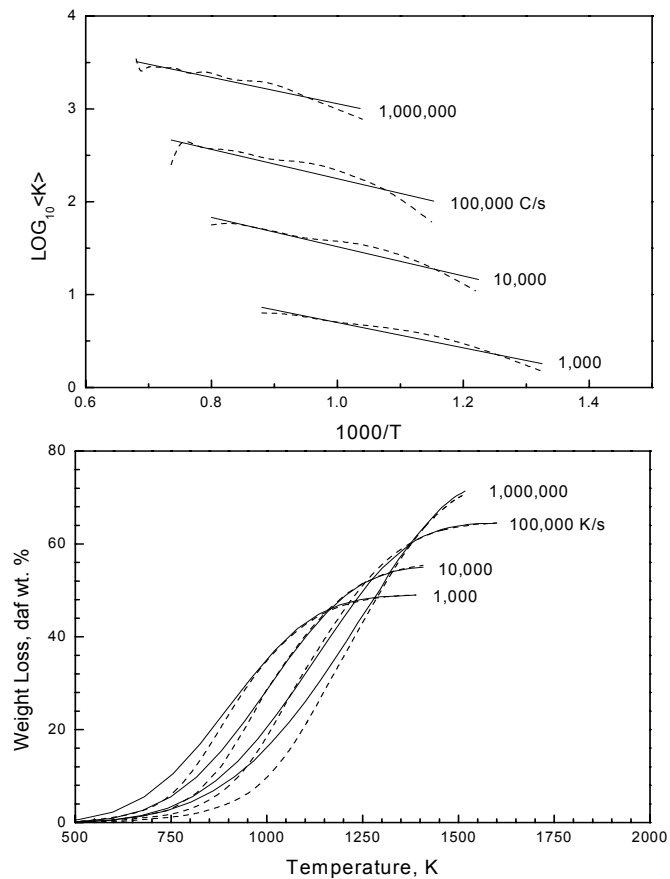
- Depicts the distinctive yields of individual samples of even the same coal rank.
- Based on only the proximate and ultimate analyses.

One Framework Covers All P.F.



- Only coal exhibits a continuous rank dependence.
- Petroleum derivatives & biomass determine $(H/C)_{MAX}$.
- Black liquor & biomass determine $(O/C)_{MAX}$.

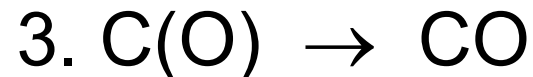
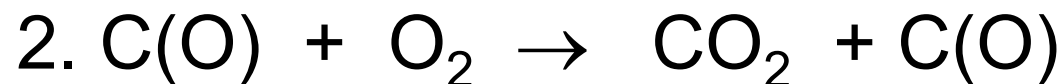
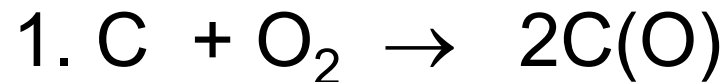
Automatically Assign All Devolatilization Rate Parameters



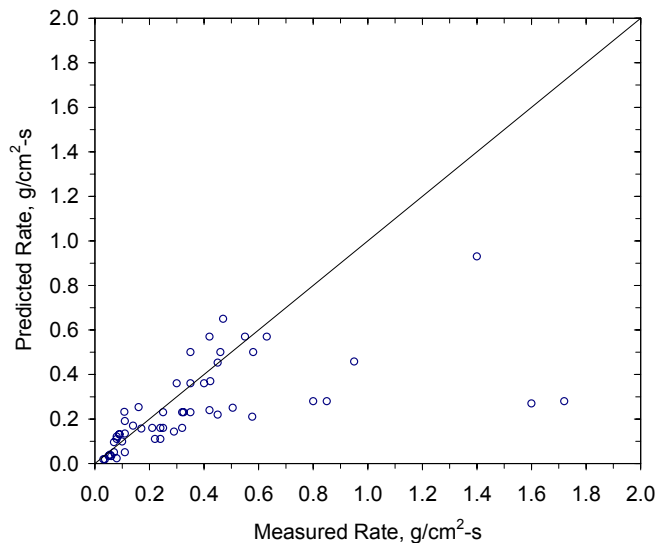
- Even the SFOR can match the FC predictions for devolatilization during heatup.
- Assigned activation energies are constant over a broad range of heating rate.
- Assigned frequency factors expressed as a function of heating rate, along with ultimate yields.
- Rate laws can be specified for any product predicted by FC, including volatile-N.

The Carbon Burnout Kinetics Model/Extended Version (CBK/E) for Char Oxidation

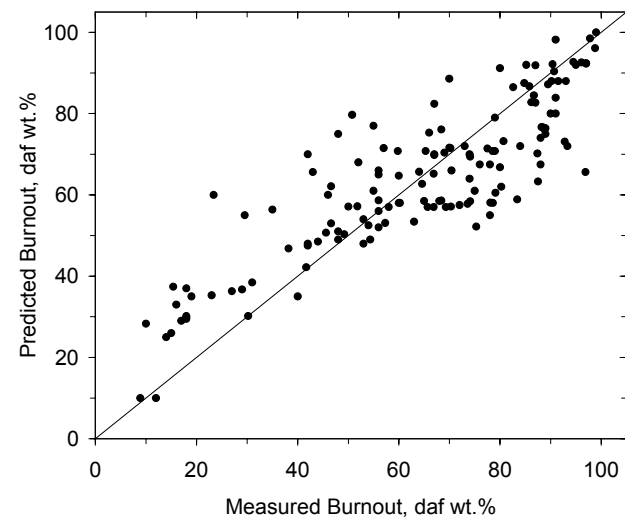
- CBK/E includes single-film char combustion, intraparticle reaction/diffusion, thermal annealing, and ash inhibition.
- Three-step intrinsic kinetics resolves the problems in the reaction order for conventional char oxidation kinetics.



No Systematic Discrepancies for Shock Tube or EFT Databases



Parity plot for burning rate predictions for the shock tube database based on the best-fit assignment to A_{30} for each coal.



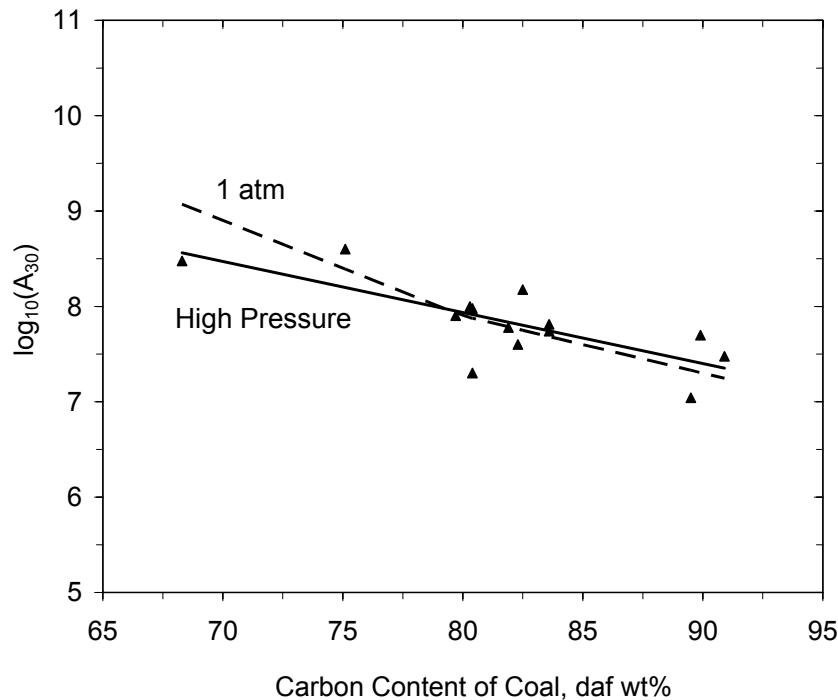
Parity plot of burnout predictions for the EFR database based on the best-fit parameter assignment for each coal.

CBK/E was validated against a database of 235 independent tests that characterized 11 coals, 2 coal chars, and a graphite, heating rates approaching 10^6 °C/s, furnace temperatures to 1527 °C, pressures to 2.0 MPa, and O_2 levels to 100 %.

A One-Point Calibration is Needed for Every Fuel Sample

- Adjust one frequency factor, A_{70} , to fit the measurements for each coal.
- Use default values/correlations for all other modeling parameters.
- Correlate A_{70} with rank to estimate default rate parameters.

The Rank Dependence of Char Burnout is Similar At Elevated Pressure



- Same reactivity for subbituminous coals through low volatility coals.
- Low rank chars have diffusion-limited burning rates.
- Enhanced plasticity of low-rank coals at elevated pressure may lower the reactivity.

Fundamental Difference Between Combustion and Gasification:

- *Chemistry in the gas phase determines the levels of the char gasification agents (CO_2 , H_2O , CO , H_2).*

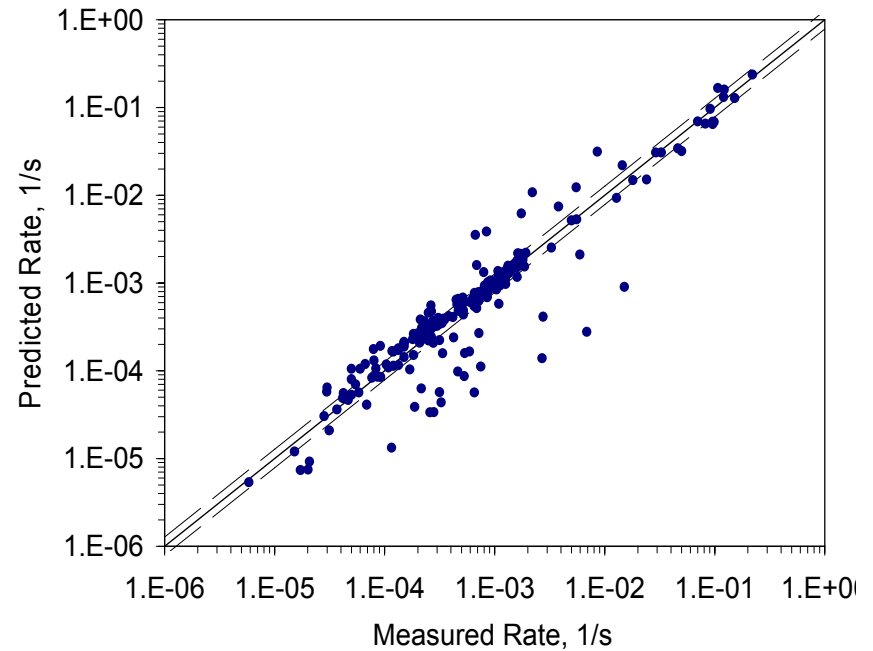
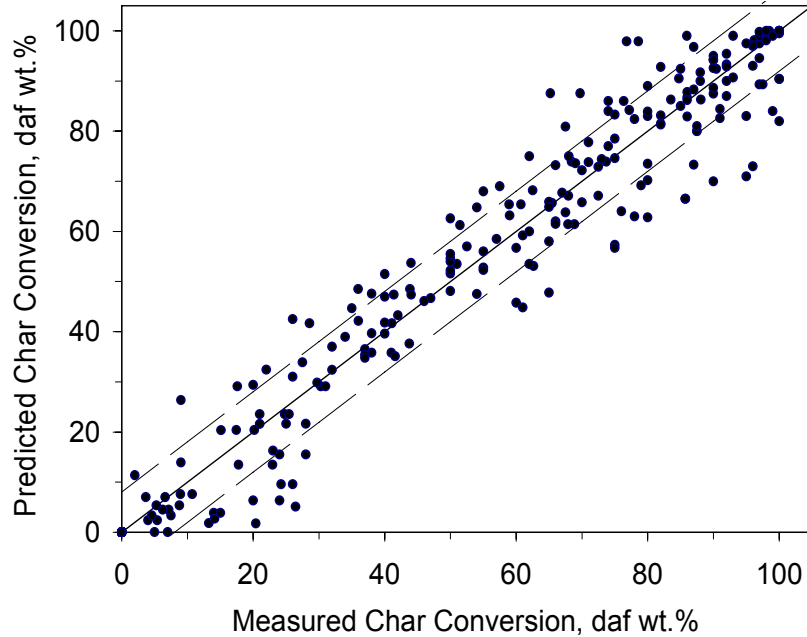
In a **p. c. flame**, all O_2 enters the furnace through burners & OFA ports, so mixing limited extents of conversion determine the local O_2 concentration.

In a **gasifier**, injected O_2 only partly determines the conversion levels of volatiles/soot/char, whereas the **levels of CO , H_2 , CO_2 , and H_2O are variable**. Gas mixtures are not stable at 1600°C , so gas phase chemistry is important **throughout the entire gasifier**.

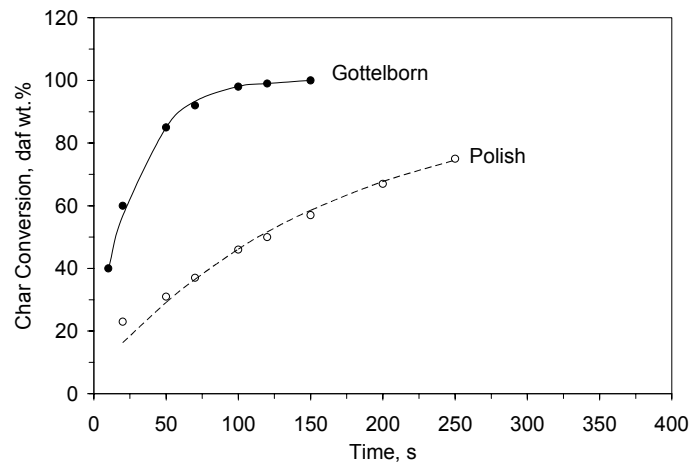
Use CBK/G to Predict Gasification Rates

- Combustion
$$2\text{C} + \text{O}_2 \rightarrow \text{C}(\text{O}) + \text{CO}$$
$$\text{C} + \text{C}(\text{O}) + \text{O}_2 \rightarrow \text{C}(\text{O}) + \text{CO}_2$$
$$\text{C}(\text{O}) \rightarrow \text{CO}$$
- Gasification
$$\text{C} + \text{CO}_2 \leftrightarrow \text{C}(\text{O}) + \text{CO}$$
$$\text{C}(\text{O}) \rightarrow \text{CO}$$
$$\text{C} + \text{H}_2\text{O} \leftrightarrow \text{C}(\text{O}) + \text{H}_2$$
$$\text{C} + 2\text{H}_2 \rightarrow \text{CH}_4 \quad (\text{slow})$$
- CBK/G was validated against a database of 452 independent tests that characterized 26 coals, heating rates approaching 10^5 °C/s, furnace temperatures to 1500 °C, pressures to 3.0 MPa, and broad ranges of CO₂, H₂O, CO, and H₂ levels.
- Separate surface oxide pools for the combustion and gasification reactions.
- Separate surface oxide pools for CO₂ and H₂O gasification.
- Currently neglecting CC(O) chemistry and CO chemisorption as marginal.

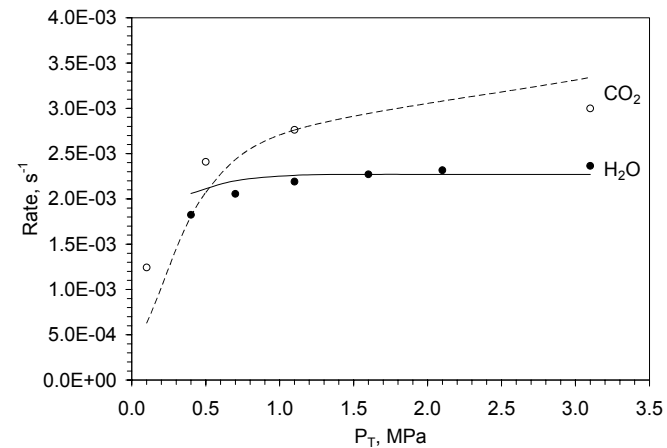
No Systematic Discrepancies in Predicted Extents of Conversion or Gasification Rates



CBK/G Performs Well Over A Broad Domain of Operating Conditions

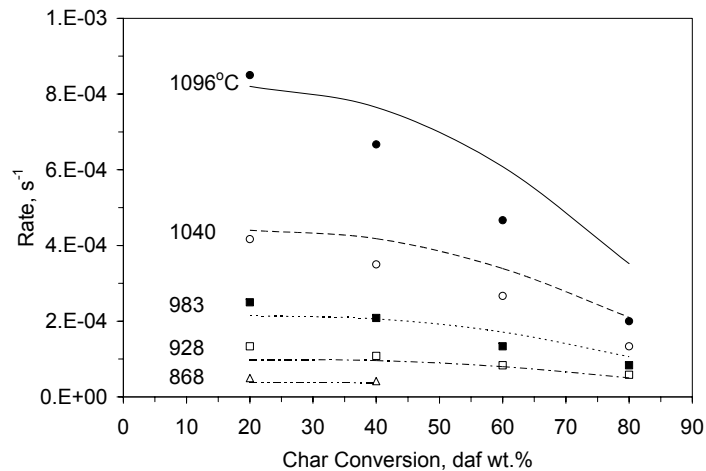


Predicted (curves) and measured (data points) char conversion histories for (● and solid line) Polish and (○ and dashed line) Gottelborn chars at 1500°C and 0.1 MPa pure CO₂ in a WMR (Moors, 1998).

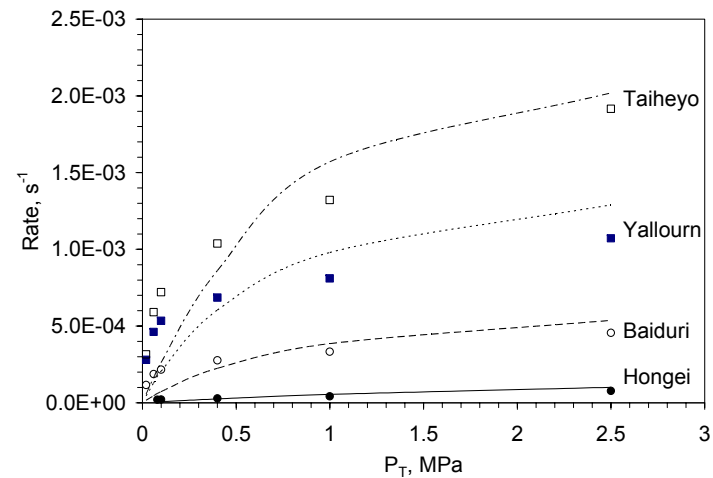


Predicted (curves) and measured (data points) rates of Xiao Long Tan lignite char gasification at (● and solid line) 850°C in 80 % H₂O, 10 % H₂, and 10 % CO, and at (○ and dashed line) 900°C in 90 % CO₂ and 10 % CO (Sha et al., 1990).

CBK/G Performs Well Over A Broad Domain of Operating Conditions

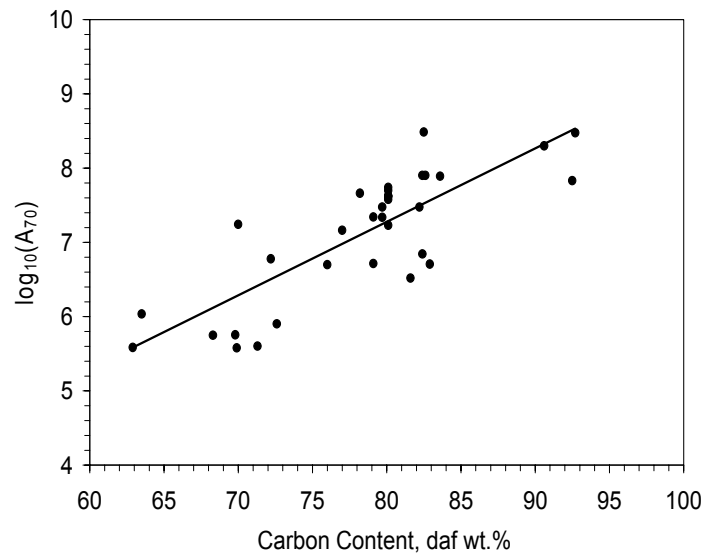


Predicted (curves) and measured (data points) reaction rate profiles for Jincheng anthracite under 0.1 MPa steam at (● and solid line) 1096, (○ and dashed line) 1040, (■ and dotted line) 983, (□ and dotted-dashed line) 928, and (△ and double dotted-dashed line) 868°C (Ma et al., 1992).

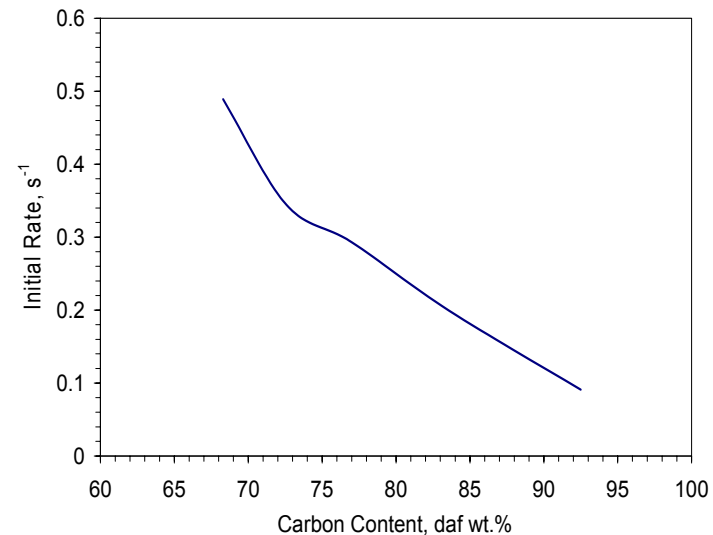


Initial CO₂ gasification rates of (● and solid line) Hongei, (○ and dashed line) Baiduri, (■ and dotted line) Yallourn, and (□ and dotted-dashed line) Taiheyo at 850°C in a PTGA (Nozaki et al., 1992).

Plausible Correlations for the Reactivity, *BUT* Wide Dispersion



Rank dependence of corrected values for A_{70} . Solid circles denote the best fit values for each coal. The solid line represents the correlation between A_{70} and a coal's carbon content.



Rank dependence of the initial burning rate of a 90- μm coal in an 1200°C EFR, at 2.0 MPa and 1400°C and with 10 % CO_2 , 30 % H_2O , 10 % CO and 10 % H_2 , based on the correlation in Eq. 64.

Simple L-H/Nth-Order Rate Laws Reproduce the Rates from CBK/G

$$R_{CO_2} = \mathcal{G} \cdot \frac{A_{CO_2} \cdot \exp(-E_{CO_2} / RT) P_{CO_2,S}^{n_{CO_2}}}{1 + K_{CO} P_{CO,S}}$$

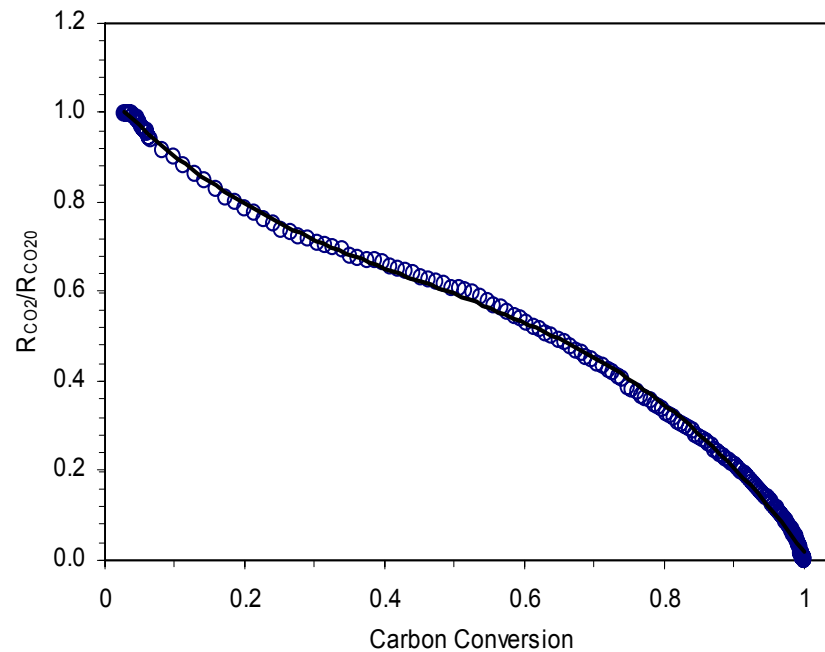
$$R_{H_2O} = \mathcal{G} \cdot \frac{A_{H_2O} \cdot \exp(-E_{H_2O} / RT) P_{H_2O,S}^{n_{H_2O}}}{1 + K_{H_2} P_{H_2,S}}$$

$$R_{H_2} = \mathcal{G} \cdot A_{H_2} \cdot \exp(-E_{H_2} / RT) P_{H_2,S}^{n_{H_2}}$$

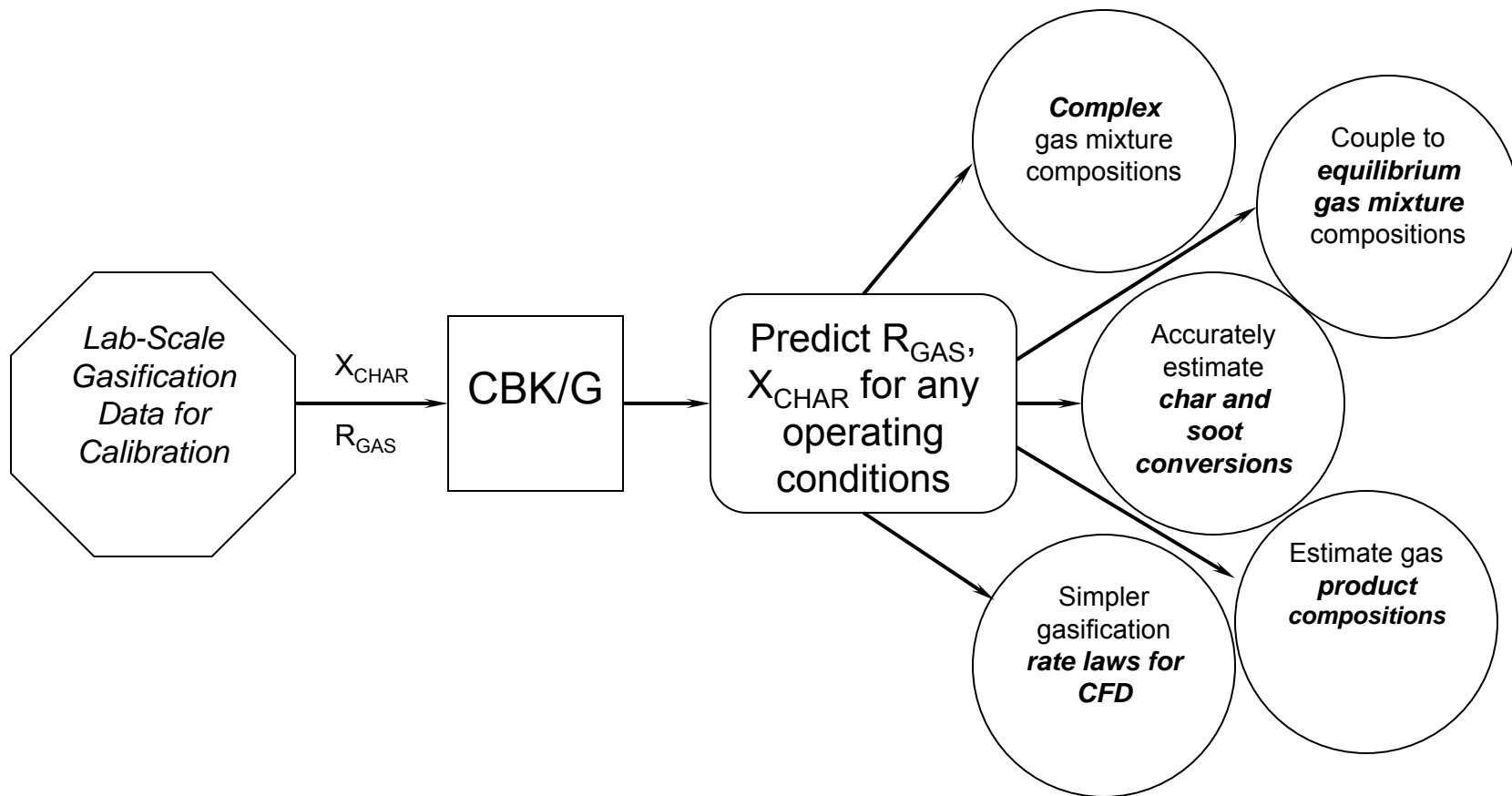
$$\text{where } \mathcal{G} = a_0 + a_1 X + a_2 X^2 + a_3 X^3 + a_4 X^4 + a_5 X^5$$

Must Also Apply a Rate-Reduction Polynomial

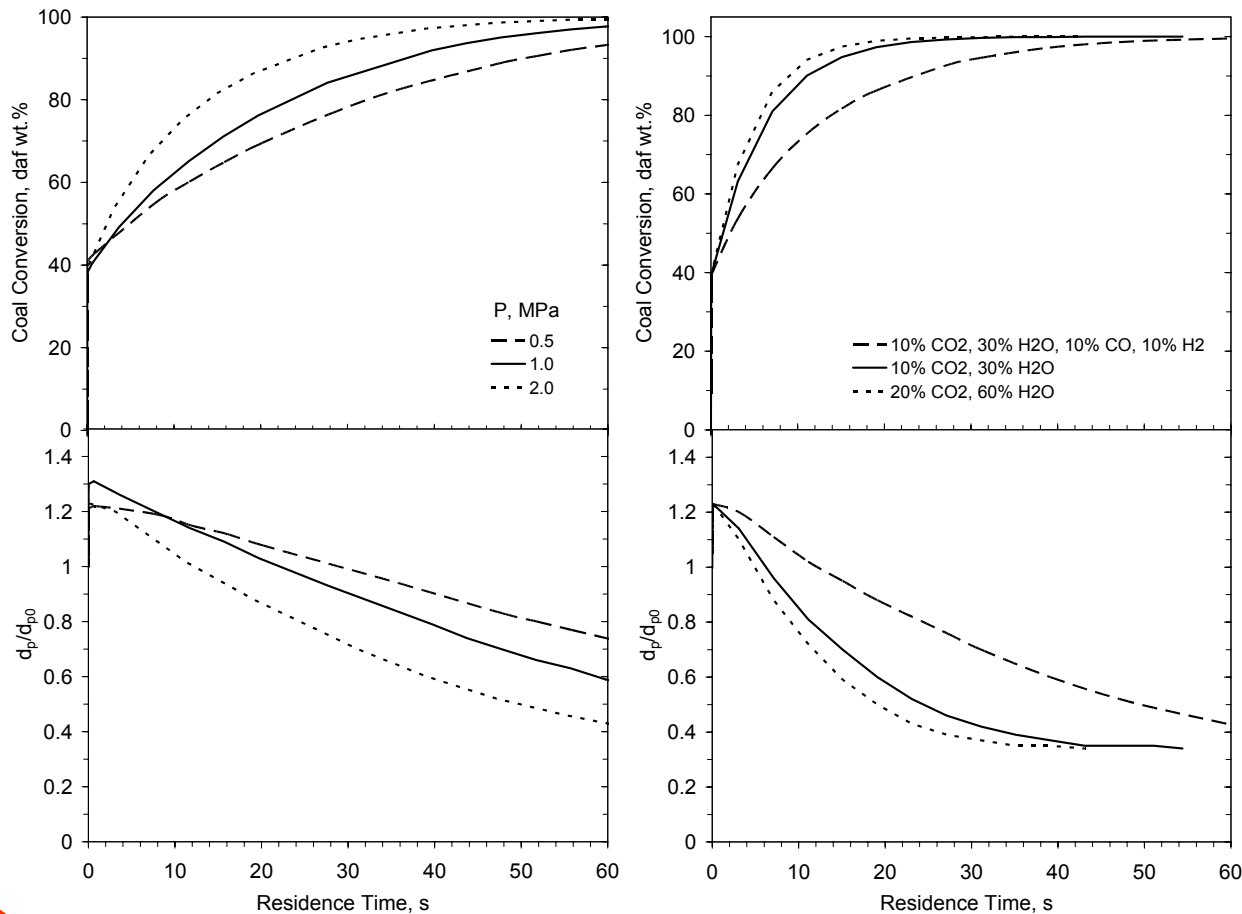
$$\frac{R_{\text{CO}_2}}{R_{\text{CO}_2_0}} = -1.7825X^3 + 2.4287X^2 - 1.679X + 1.0 \text{ where } X = \text{Extent of conversion}$$



Use CBK/G to Extrapolate to Actual Gasifier Conditions

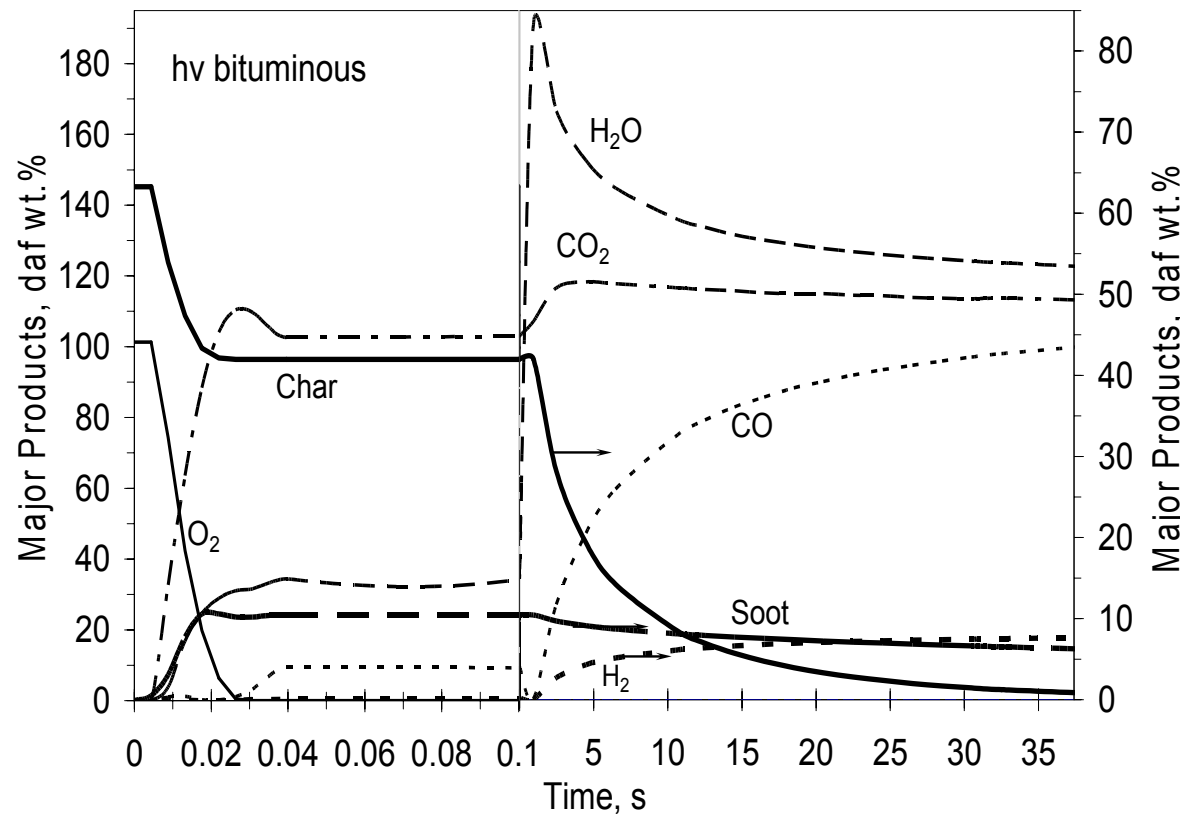


Extrapolations Based on CBK/G



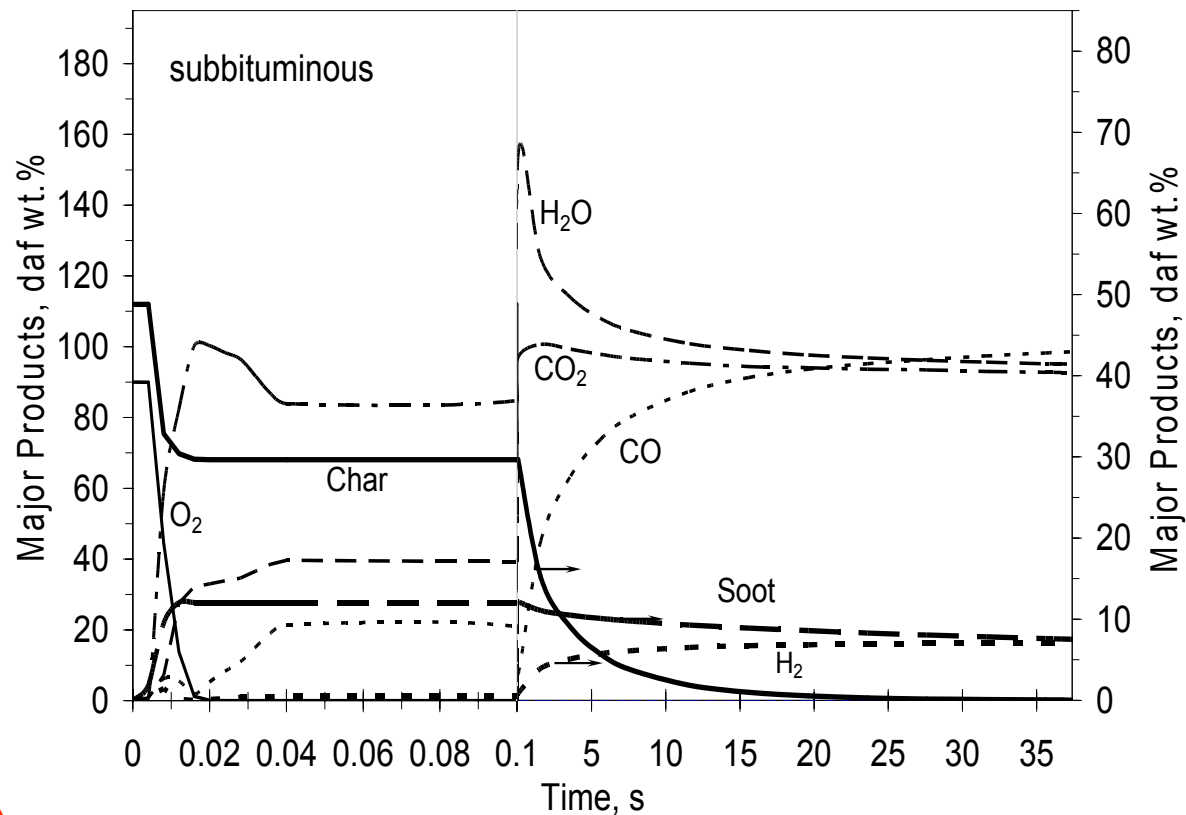
- *Faster gasification for higher pressures.*
- *Inhibition by CO & H₂ stronger than the impact of doubling the CO₂ and H₂O levels.*

The FQ Impacts are Evident in a 1D Gasifier Simulation With Detailed Chemistry



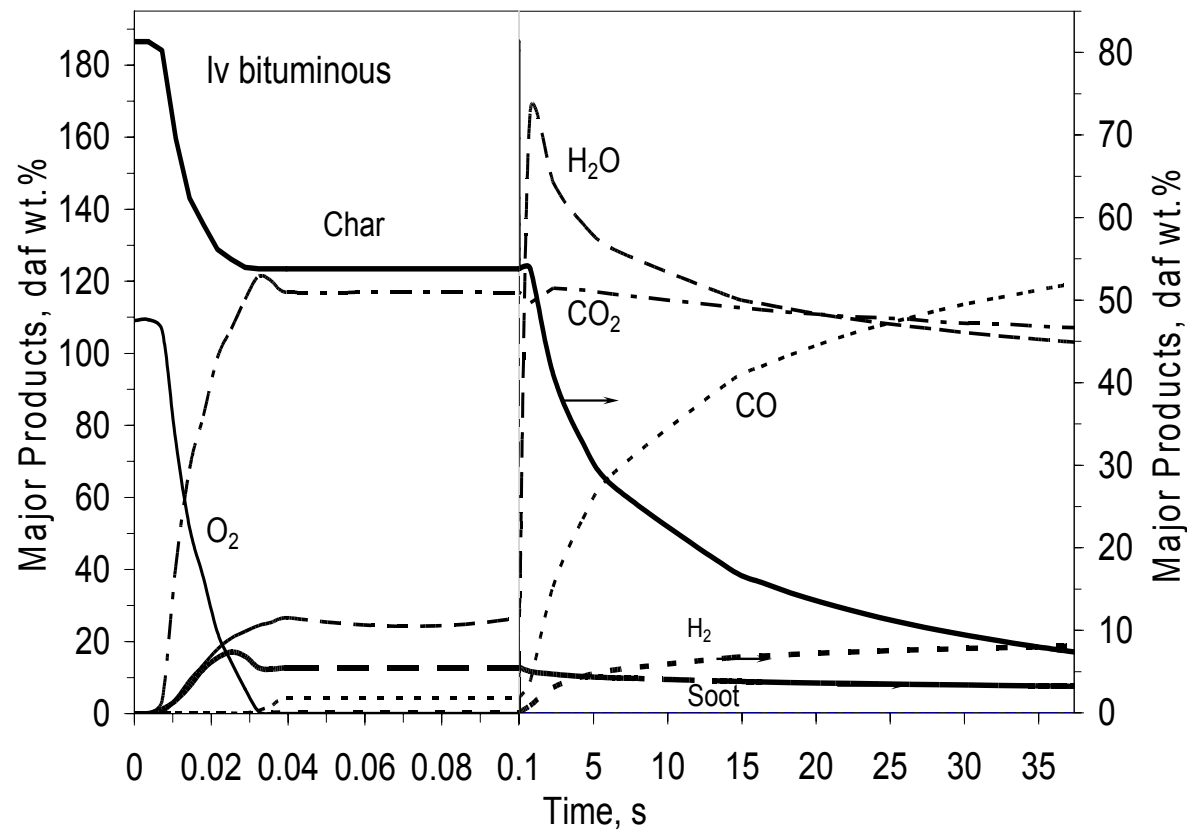
- *First-stage calculation based on full kinetics to determine X_{char} and X_{soot} .*
- *Steam injection into a reducing second stage.*
- *Equilibrium gas compositions shift throughout the second stage.*
- *Steam gasification with strong CO inhibition.*
- *Soot persists.*

The FQ Impacts are Evident in a 1D Gasifier Simulation With Detailed Chemistry



- *Similar trends with subbituminous coal.*
- *Comparable product gas quality.*
- *Soot persists.*

The FQ Impacts are Evident in a 1D Gasifier Simulation With Detailed Chemistry



- *Similar trends with Iv bituminous coal.*
- *Richer product gas quality.*
- *Soot and char persist.*

Where's the Soot ?!

Coal	Soot Yield, daf wt. %	% of Wt. Loss
Pit. #8	22.9 - 29.1	43 - 57
Ill. #6	21.0	40
PRB	9.1	19

- **Most of the Volatile-C incorporated into soot. Soot competes for available O₂, scavenges radicals. Omitted in every reported gasifier simulator.**
- **No equilibrium until the soot is consumed.**
- **Kinetics determine the gas composition in, perhaps, the first half of a gasifier.**
- **Without the correct gas composition, predicted gasification rates will be incorrect.**
- **Must work with multiple gasification agents (CO, CO₂, H₂O, H₂, CH₄).**

Critical Needs

- Tests with >20 coals under standard conditions are needed to develop improved correlations between coal properties and the initial char gasification reactivity. Monitor loadings of alkali and alkaline earth cations.
- Divert lab testing away from cases with a single gasification agent to characterize (i) inhibition by CO and H₂ and (ii) the complex mixtures that arise in gasifiers.
- Monitor gasification rates for coal-derived soot.
- Characterize the coupling among secondary volatiles pyrolysis, gas phase chemistry, and the conversion of char and soot throughout gasification at realistic suspension loadings.