

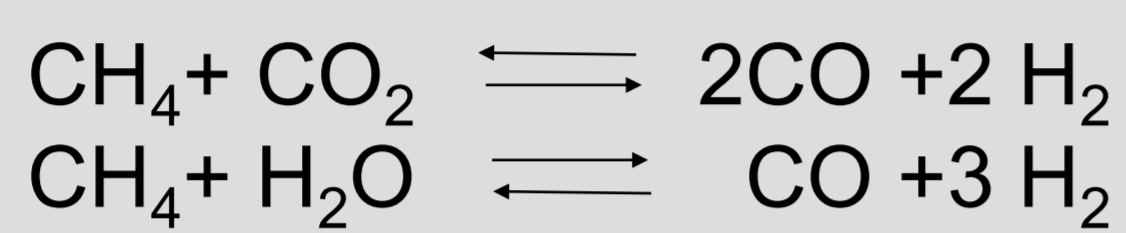
Development of a Combined Surface Reaction Mechanism for Steam- and Dry Reforming of Methane over Nickel

Karla Herrera Delgado¹, Lubow Maier², Olaf Deutschmann^{1, 2}

karla.herrera@kit.edu

Background

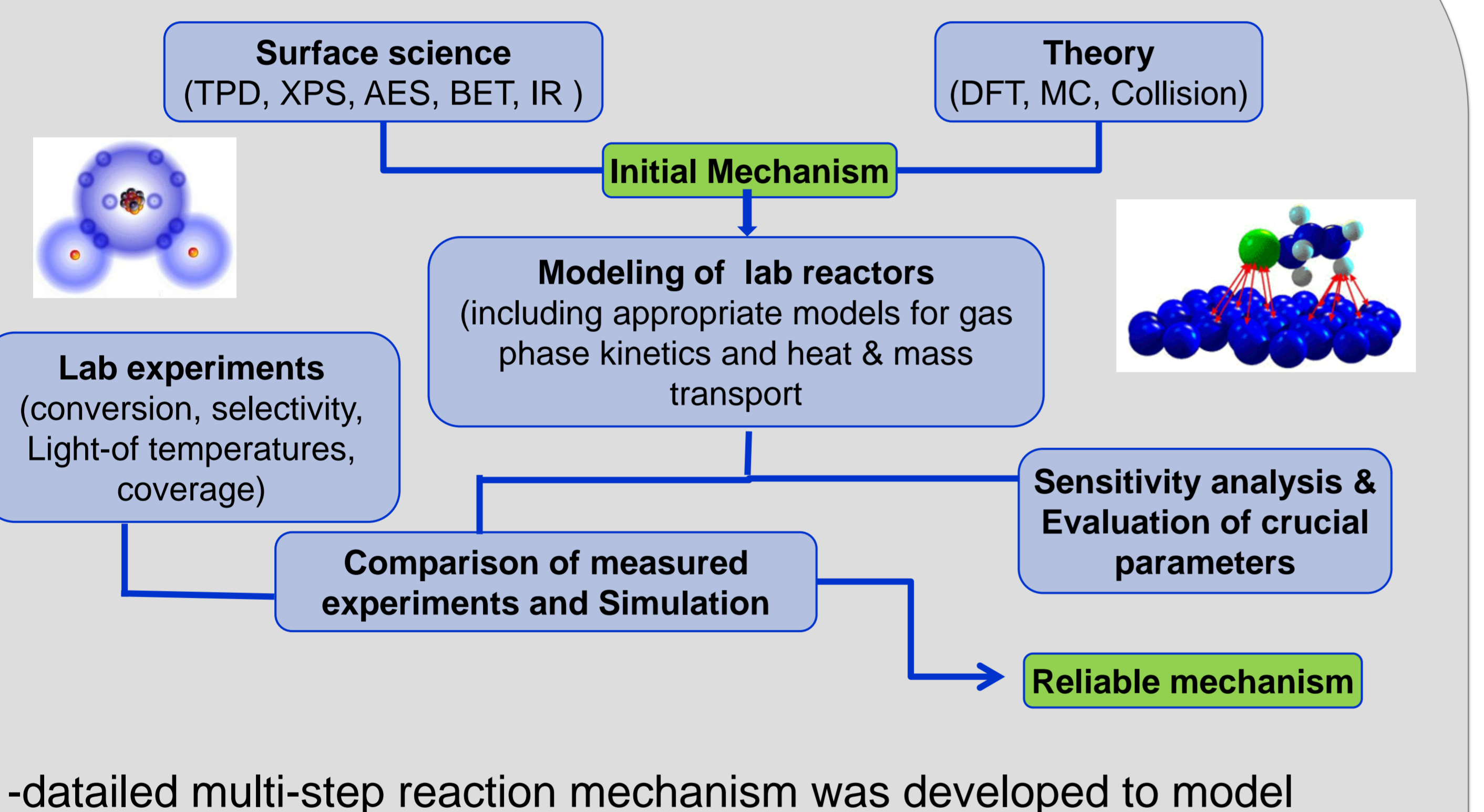
Steam reforming and dry reforming of methane play a key role in the production of syngas (H_2/CO), used in synthesis of chemicals and fuels (gas-to-liquids) [1]



Dry reforming of methane with carbon dioxide has special interest due to the increasing concern of global warming and oil depletion since offers the opportunity to convert greenhouse gases into syngas with low H_2/CO ratio [2].

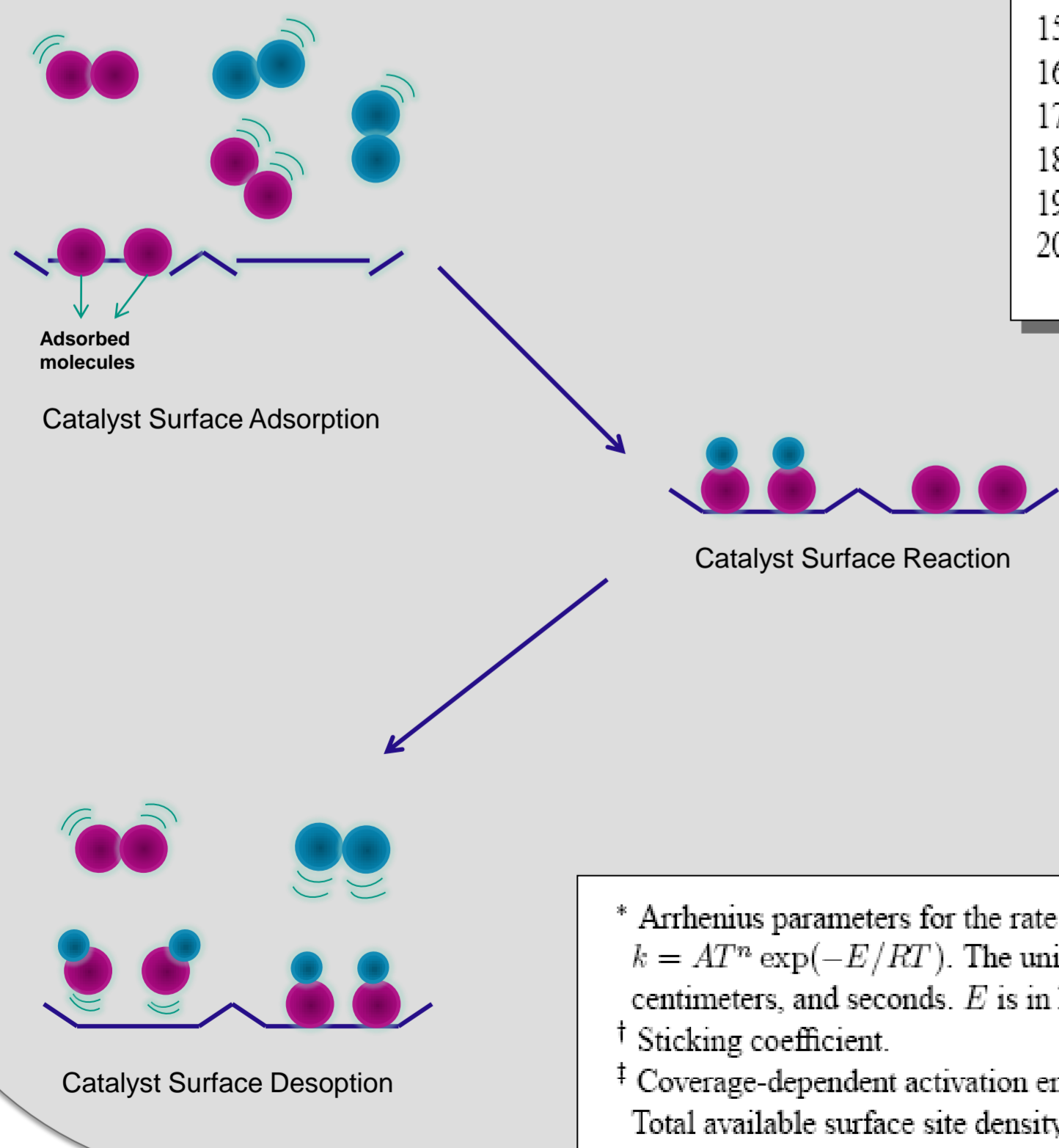
Nickel catalyst is widely used due to fast turnover rates, good availability and low cost, although it is more sensitive to coke formation and growth of carbon filaments than noble metals [3].

Modeling and Simulation



-detailed multi-step reaction mechanism was developed to model steam reforming

- it include
 - Adsorption- desorption reactions of reactants and products
 - Gas phase and surface species
 - Partial and total oxidation reactions
 - Thermodynamically consistent



- References:**
- V. F. Fischer, H.Tropsch, Brennstoff-Chemie, 3, 39 (1928)
 - J. Ma, N. Sun, X. Zhang, N. Zhao, Catal. Today 148 (2009)
 - S.Wang, G. Q. M. Lu, Appl.Catal. B: Environ, 16, 269 (1998)
 - O. Deutschmann, S. Tischer, et al, DETCHEM™ Software package, 2.3 ed., Karlsruhe 2010,
 - J.-H. Ryu, K.Y. Lee, J. Power Sources 171, 499 (2007)
 - E. Hecht, G. K. Gupta, H. Zhu, A. M. Dean, R. J. Kee, L. Maier, O. Deutschmann, Appl. Catal. A: General, 40, 295 (2005) 40.

Mechanistic Model (mean field approximation)

- The molecules are randomly distributed on the catalytic surface
- Surface is viewed as being uniform

Adsorption rate

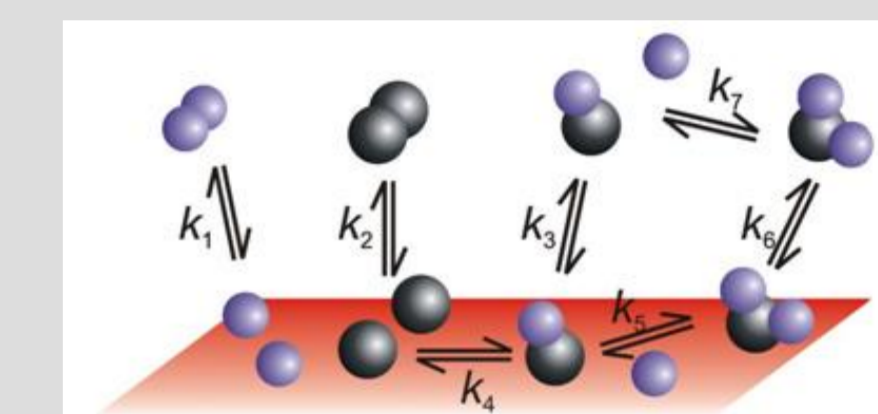
$$\dot{s}_i = s_i^{eff} \sqrt{\frac{RT}{2\pi M_i}} c_i$$

Surface reaction rate

$$\dot{s}_i = \sum_k v_{ik} k_{f_k} \prod_j^{N_g+N_s} c_j^{v_{jk}}$$

Sticking coefficient

$$s_{eff}^i = s_i^0 \prod_{j=1}^{N_s} \Theta_j^{v'_{jk} + \mu_{jk}}$$



Rate expression

$$\dot{s}_i = \frac{c_i \sigma_i}{\Gamma} \quad \frac{\partial \Theta_i}{\partial t} = \frac{\dot{s}_i \sigma_i}{\Gamma} \quad k_{f_k} = A_k T^{\beta_k} \exp\left[\frac{-E_{a_k}}{RT}\right] \prod_{i=1}^{N_s} \Theta_i^{\mu_{ik}} \exp\left[\frac{\epsilon_{ik} \Theta_i}{RT}\right]$$

Surface coverage

$$\Theta_i = \frac{c_i \sigma_i}{\Gamma} \quad \frac{\partial \Theta_i}{\partial t} = \frac{\dot{s}_i \sigma_i}{\Gamma}$$

Numerical Simulation - DETCHEM Software

- modeling and simulation of reacting flows field.
- coupled with the detailed surface chemistry model and transport phenomena.
- transport coefficients depend on mixture composition and temperature [4]

MONOLITH
Transient temperature profile of the solid structure

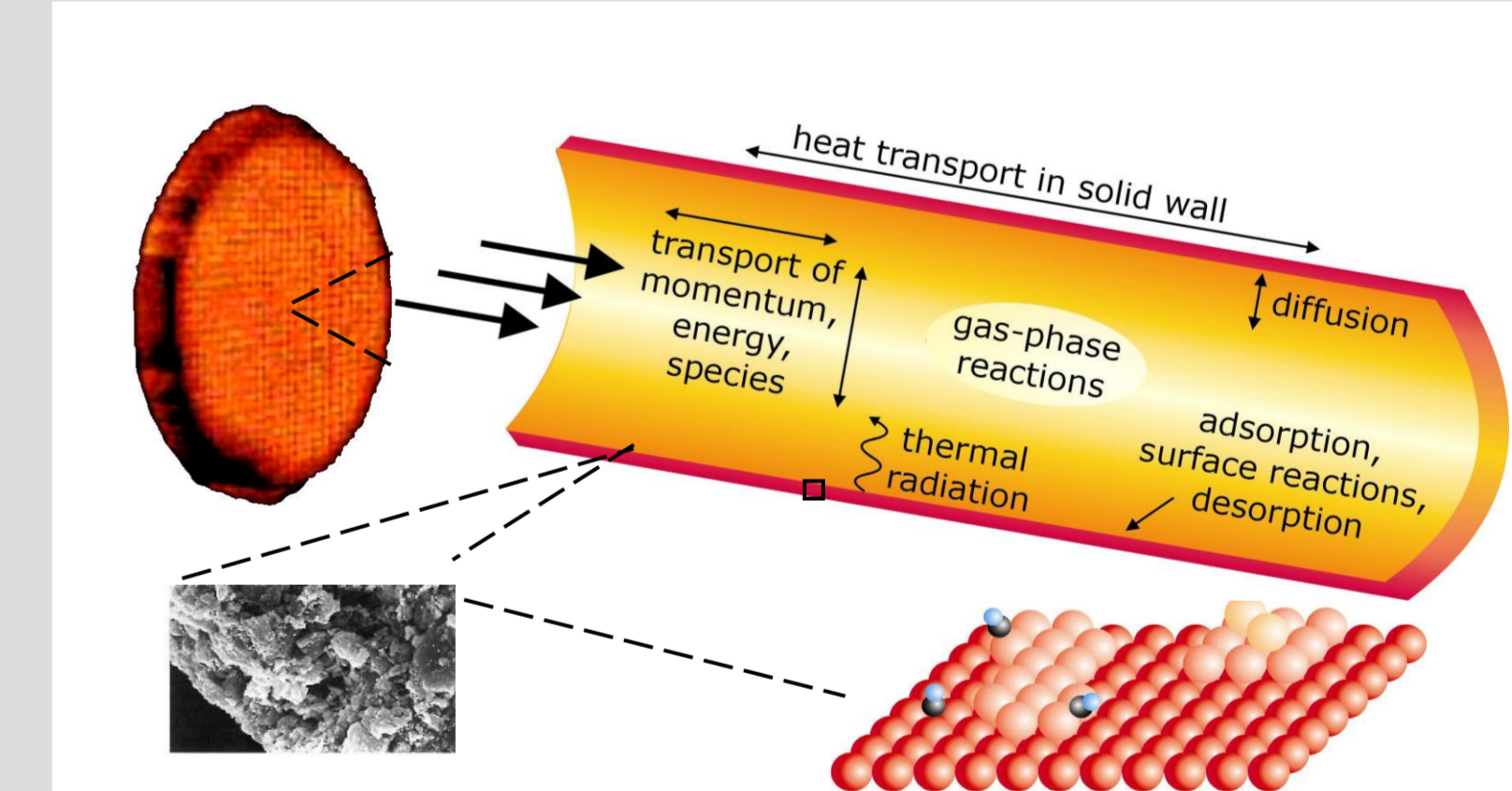
CHANNEL or PLUG
1D or 2D-flow field simulations for a representative number of channels using a boundary layer or plug flow equations

BATCH + CSTR
Batch and stirred tank

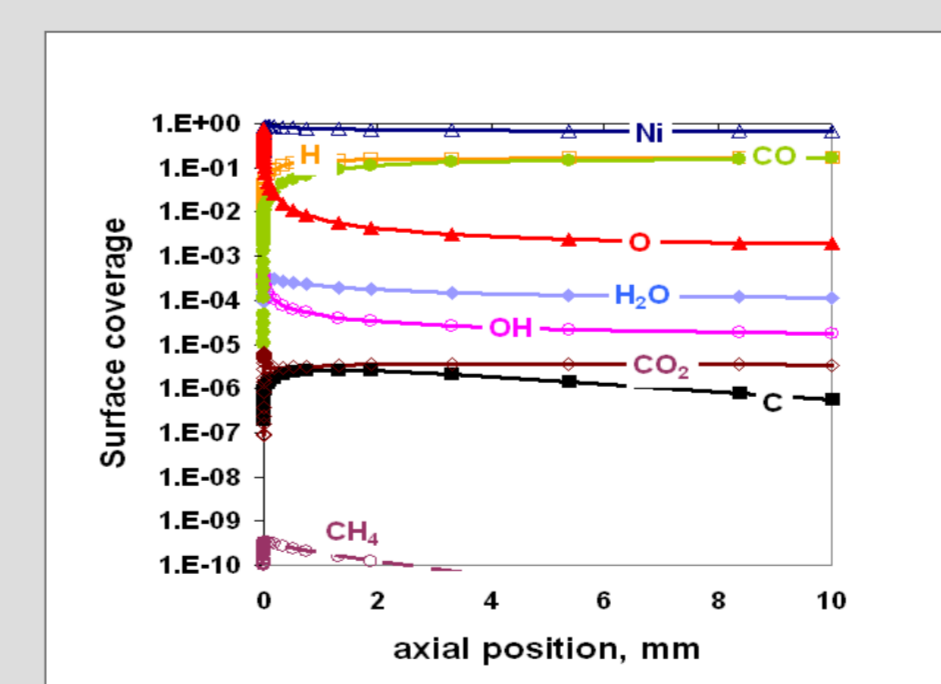
SENSITIVITY TEST

PACKEDBED
1D Bed Reactor

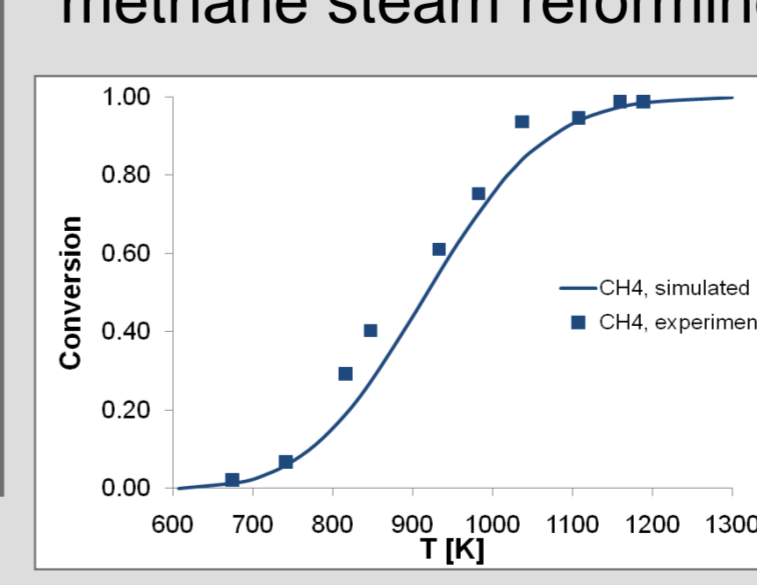
DETACHEM- Library
Detailed Reaction Mechanisms
Thermodynamic and Transport Data
Mol Data
Species



Simulation Results

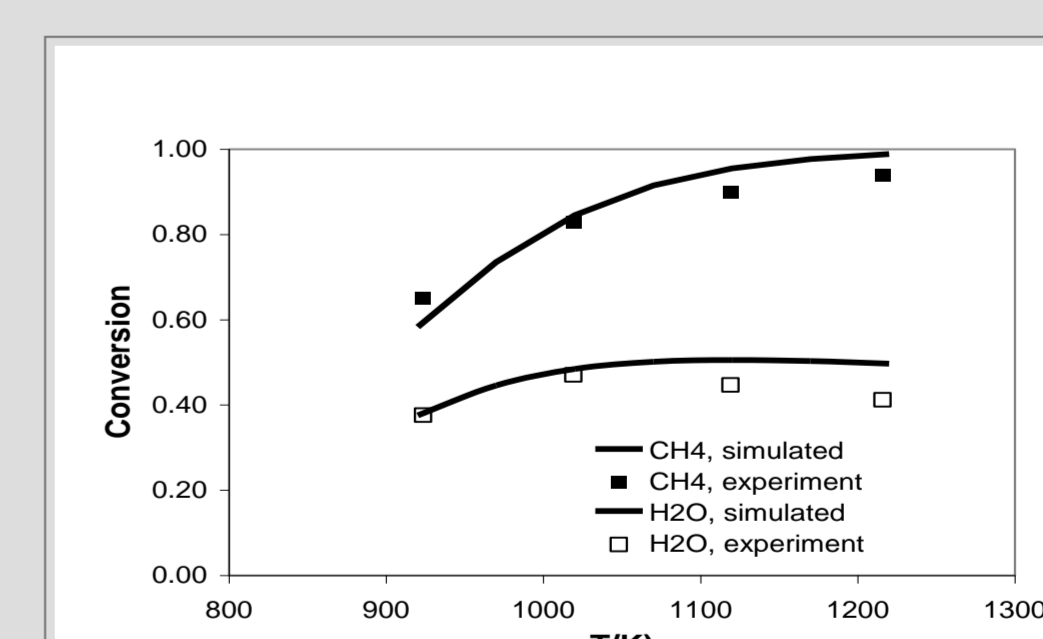


Computed surface coverage of adsorbed species along the catalytic channel wall in methane steam reforming at 1120K, S/C=2.77



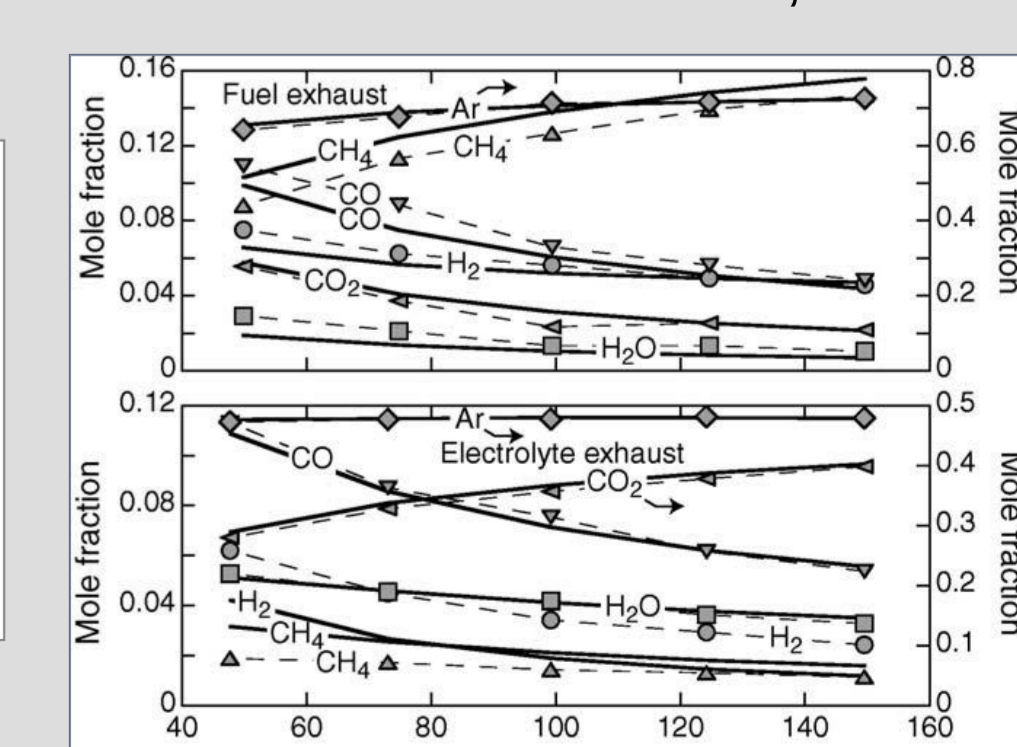
Steam reforming (metal monolith) [5]

The model developed is able to properly describe steam reforming of methane for wide ranges of experimental conditions, simulation results of the mechanism are in good agreement with experiments .



Steam reforming (packed-bed) [5]

CH4 and H2O conversion as a function of temperature in CH4 SR: S/C=2.77, 75% Ar



Dry Reforming of CO2 over Ni/YSZ anodes in Solid-oxide fuel cells (SOFC) [6]