

Reaction kinetics modelling

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Recommendation of special courses of Tamás Turányi

Chemistry and Physics of Flames (in English)

in each spring semester

- combustion chemistry
- experimental methods for the investigation of combustion reactions
- experimental determination of gas-phase rate coefficients
- physics of flame spread

Analysis of Kinetic Reaction Mechanisms (in English)

in each autumn semester

- reaction kinetics modelling
- sensitivity analysis
- uncertainty analysis
- time scale analysis
- mechanism reduction

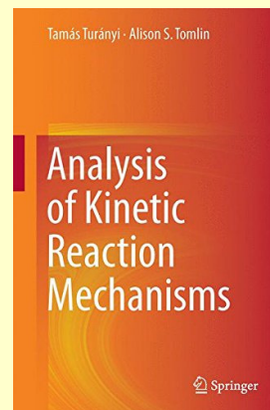
Analysis of kinetic reaction mechanisms – the book

Tamás Turányi and Alison S. Tomlin:

Analysis of kinetic reaction mechanisms

Springer, 2014

(with 1025 references)



Web page:

<http://garfield.chem.elte.hu/Turanyi/KineticReactionMechanisms.html>

- table of contents
- download the chapters
- references
- typos found

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Practical applications of reaction kinetics

- **modelling atmospheric chemical processes**
 - forecast of air pollution (weather forecast is needed!)
 - determination of emission limits
- **modelling of ignition and combustion**
 - modelling power stations, furnaces, engines
 - improving efficiency
 - elaboration of methods for the decrease of pollutant emission
- **process engineering; modelling of chemical engineering processes**
 - considering efficiency and the aspects of environment protection
- **systems biology: modelling biochemical processes within living organisms**
 - metabolic networks (*e.g.* medical drug decomposition in the body)
 - molecular signal transfer
 - modelling the cell cycle
- **non-chemical models using reaction kinetic formalism**
 - predator-prey models
 - description of ecological systems

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Simulation of spatially homogeneous systems

calculation of concentration changes
(solution of the kinetic system of differential equations) :

$$\frac{d\mathbf{Y}}{dt} = \mathbf{f}(\mathbf{Y}, \mathbf{p}) \quad \mathbf{Y}(t_0) = \mathbf{Y}_0$$

calculation of temperature changes in an adiabatic system:

$$C_p \frac{dT}{dt} = \sum_{i=1}^{N_R} \Delta_r H_i^\ominus r_i \quad T(t_0) = T_0$$

rate of reaction step i
↓
↑ standard reaction enthalpy of reaction step i

heat capacity of the mixture ↑

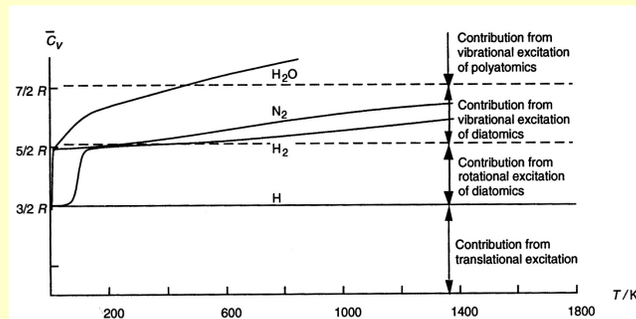
Temperature dependence of thermodynamic data

NASA polynomials

$$\frac{H^\ominus}{RT} = a_1 + \frac{a_2}{2} T + \frac{a_3}{3} T^2 + \frac{a_4}{4} T^3 + \frac{a_5}{5} T^4 + \frac{a_6}{T}$$

$$\frac{c_p}{R} = a_1 + a_2 T + a_3 T^2 + a_4 T^3 + a_5 T^4$$

$$\frac{S^\ominus}{R} = a_1 \ln T + a_2 T + \frac{a_3}{2} T^2 + \frac{a_4}{3} T^3 + \frac{a_5}{4} T^4 + a_7$$



Temperature dependence of rate coefficient k

Described by the Arrhenius equation:

$$k = A \exp\left(-\frac{E_a}{RT}\right) \qquad \ln k = \ln A - \frac{E_a}{RT}$$

A preexponential factor
E_a activation energy

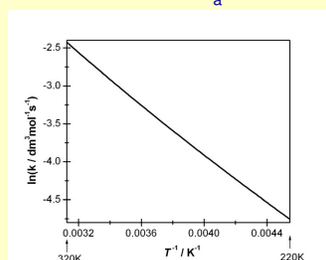
If the rate coefficient k is measured at several T temperatures and

$\ln k$ is plotted as a function of $1/T$

the data fit to a line, if the (original) Arrhenius equation is valid

slope is $m = -E_a/R$ \Rightarrow determination of E_a

Arrhenius plot:



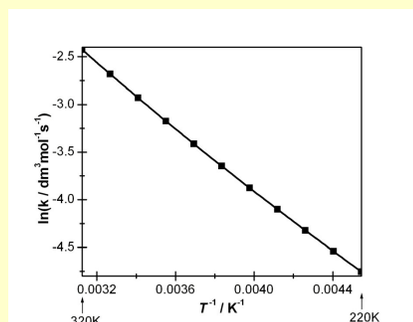
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Example: reaction $\text{CH}_4 + \text{OH} \rightarrow \text{CH}_3 + \text{H}_2\text{O}$

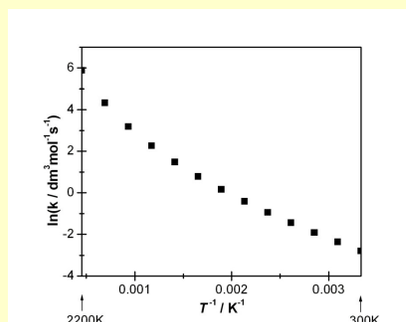
- the most important methane consuming reaction step in the troposphere
- one of the most important steps at methane combustion

Arrhenius plot between 220 K (– 53 °C)
and 320 K (+ 47 °C)

Arrhenius plot between 300 K (27 °C)
and 2200 K (\approx 1930 °C)



the Arrhenius equation is usually
very accurate in a small
(few times 10 K) temperature range.
(solution phase and atmospheric chemistry)

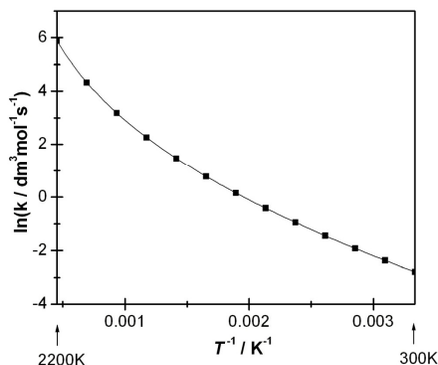


the original Arrhenius equation
is usually not applicable in a wide
temperature range
(combustion and pyrolytic systems)

Temperature dependence of the rate coefficient 2

$$k = BT^n e^{-\frac{C}{RT}}$$

extended Arrhenius equation



Important!

If $n \neq 0$, then $A \neq B$ and $E_a \neq C$

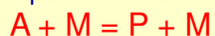
General definition of activation energy:

$$E_a = -R \frac{\partial \ln k}{\partial (1/T)}$$

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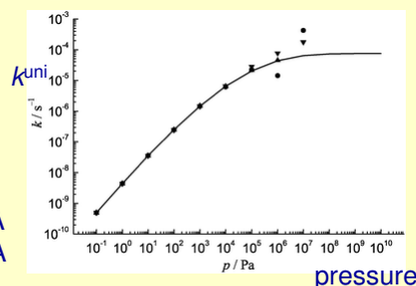
Pressure dependence of the rate coefficients 1 unimolecular decomposition

decomposition or isomerization of a species



A reactant
P product
M colliding partner for energy transfer

low pressure 2nd order decay of A
high pressure 1st order decay of A



„pseudo first order” rate coefficient:

high pressure limit: $k^{\text{uni}} = k_{\infty}$
low pressure limit: $k^{\text{uni}} = k_0 [M]$

The temperature dependences of k_0 and k_{∞} are described independently by extended Arrhenius equations:

$$k_0 = A_0 T^{n_0} \exp\left(\frac{-E_0}{RT}\right) \quad k_{\infty} = A_{\infty} T^{n_{\infty}} \exp\left(\frac{-E_{\infty}}{RT}\right) \quad 12$$

Lindemann – Hinshelwood model



high pressure: $[M] \rightarrow \infty$ $k_{\text{uni}} \approx \frac{k_3 k_1}{k_2}$ $k_{\infty} = \frac{k_3 k_1}{k_2}$

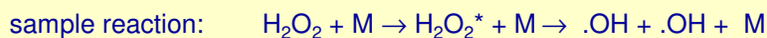
low pressure: $[M] \approx 0$ $k_{\text{uni}} \approx k_1 [M]$ $k_0 = k_1$

unimolecular decomposition rates:

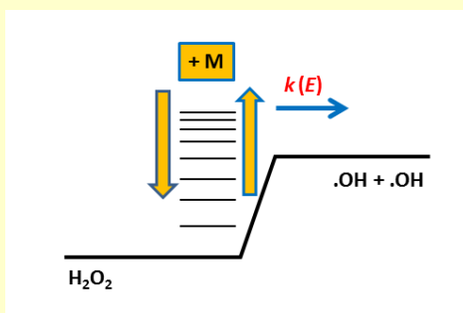
k_{uni} „pseudo first order” rate coefficient (s^{-1})
 k_{∞} high pressure limit (first order) rate coefficient (s^{-1})
 k_0 low pressure limit (second order) rate coefficient ($\text{mol}^{-1} \text{dm}^3 \text{s}^{-1}$)

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Refined Lindemann – Hinshelwood model of unimolecular reactions



- instead of assuming a single excited species, H_2O_2^* exists in a range of different rovibrationally excited states.
- the rate of the decomposition step increases with the increasing rovibrational energy of H_2O_2^*



low pressure:

- the rate limiting is the collision with M
- the overall decomposition rate is proportional with pressure

high pressure:

- the rate limiting is the decomposition of H_2O_2^*
- the overall decomposition rate is independent of pressure



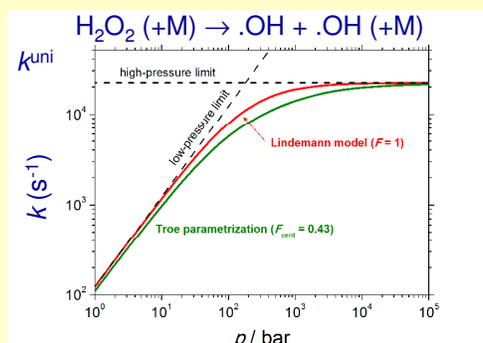
2nd order at low pressure, 1st order at high pressure¹⁴

Unimolecular decomposition: calculation of the rate coefficient at an intermediate pressure

$$k = k_{\infty} \left(\frac{P_r}{1 + P_r} \right) F$$

$$P_r = \frac{k_0[M]}{k_{\infty}}$$

P_r reduced pressure
 F controls the shape of the $k(p)$ curve



in the Lindemann model $F = 1$

F can be defined as a function of pressure and temperature
(e.g. using Troe parameterization):

$$\log F = \log F_{\text{cent}} \left[1 + \left[\frac{\log P_r + c}{n - d(\log P_r + c)} \right]^{2n} \right]^{-1}$$

$$\begin{aligned} c &= -0.4 - 0.67 \log F_{\text{cent}} \\ n &= -0.75 - 1.271 \log F_{\text{cent}} \\ d &= 0.14 \end{aligned}$$

$$F_{\text{cent}} = (1 - \alpha) \exp\left(-\frac{T}{T^{***}}\right) + \alpha \exp\left(-\frac{T}{T^*}\right) + \exp\left(-\frac{T^{**}}{T}\right) \quad 15$$

Collision efficiency parameters

M any species present in the mixture
 BUT some species are more effective colliders

good collider: removes much energy from the excited species in each collision

Which are the good colliders?

- species with similar energy levels to those of the excited species
- large molecules with many energy levels

poor collider: e.g. noble gases:
 no rotational or vibrational energy levels
 only the translational mode can be excited

calculation of the effective concentration of M:

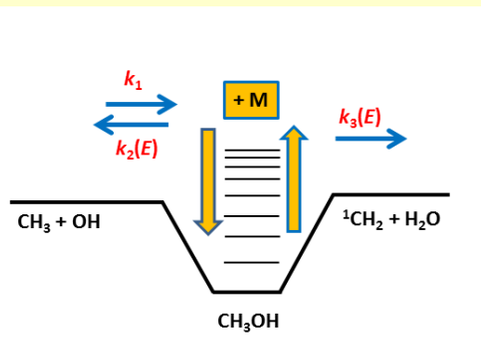
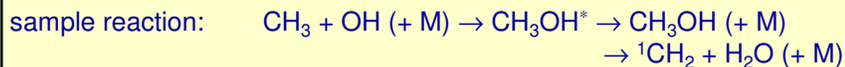
m_i : collision efficiency parameter

$$[M] = \sum_i m_{y_i} [Y_i]$$

calculation for reaction $\text{H}_2\text{O}_2 (+M) \rightarrow \cdot\text{OH} + \cdot\text{OH} (+M)$:

$$[M] = 5[\text{H}_2\text{O}] + 5.13[\text{H}_2\text{O}_2] + 0.8[\text{O}_2] + 2.47[\text{H}_2] + 1.87[\text{CO}] + 1.07[\text{CO}_2] + 0.67[\text{Ar}] + 0.43[\text{He}] + [\text{all others}]$$

Pressure dependence of the rate coefficients 2 complex-forming bimolecular reactions



low pressure:
mainly CH_3OH^* decomposition to
 $\text{}^1\text{CH}_2 + \text{H}_2\text{O}$
($\text{}^1\text{CH}_2 = \text{singlet CH}_2 =$
electronically excited CH_2)

high pressure:
mainly CH_3OH^* stabilization,
giving CH_3OH

$\text{CH}_3 + \text{OH} \rightarrow \text{CH}_3\text{OH}$ 3rd order at low pressure, 2nd order at high pressure

$\text{CH}_3 + \text{OH} \rightarrow \text{}^1\text{CH}_2 + \text{H}_2\text{O}$ 3rd order at low pressure, 2nd order at high pressure

Formation of the stabilization product

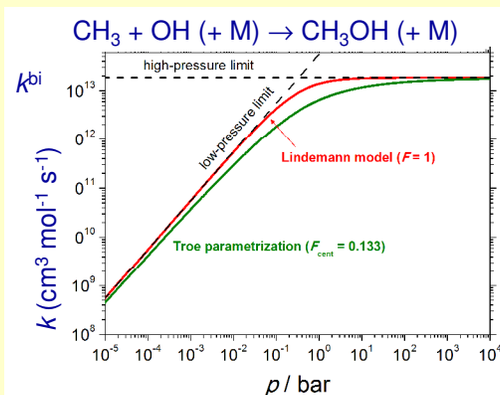
$$k^{\text{bi}} = k_{\infty} \left(\frac{P_r}{1 + P_r} \right) F$$

$$P_r = \frac{k_0 [\text{M}]}{k_{\infty}}$$

P_r reduced pressure
 F controls the shape of the
 $k(p)$ curve

in the Lindemann model $F = 1$

$F(p, T)$ can be defined by e.g. Troe parameterization



k^{bi} „pseudo second order” rate coefficient

at low pressure: $k^{\text{bi}} = k_0 [\text{M}]$

at high pressure: $k^{\text{bi}} = k_{\infty}$

Formation of the decomposition products

$$k^{\text{tri}} = k_0 \left(\frac{1}{1 + P_r} \right) F$$

$$P_r = \frac{k_0 [M]}{k_\infty}$$

P_r reduced pressure
 F controls the shape of the
 $k(p)$ curve

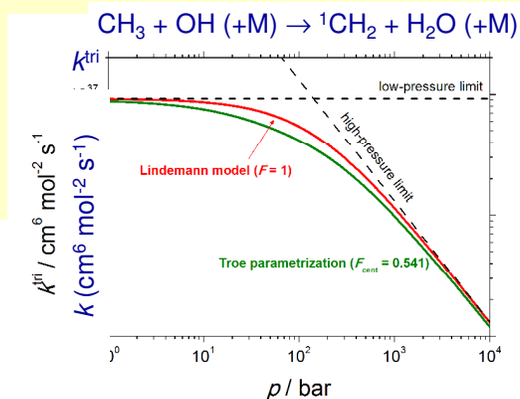
in the Lindemann model $F = 1$

$F(p, T)$ can be defined by e.g. Troe parameterization

k^{tri} „pseudo third order” rate coefficient

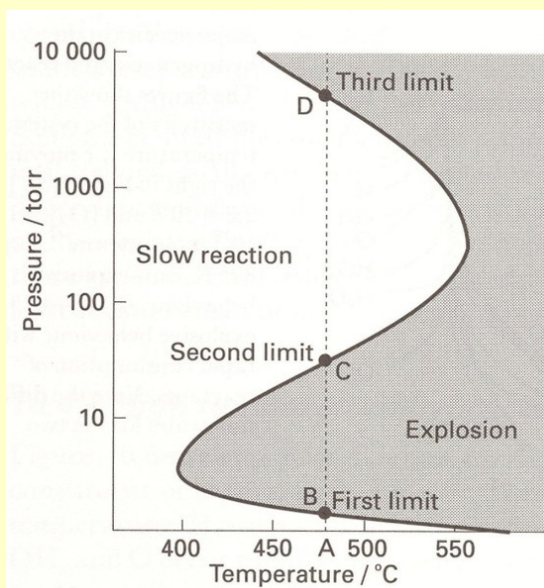
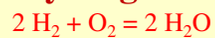
at low pressure: $k^{\text{tri}} = k_0$

at high pressure: $k^{\text{tri}} = k_\infty / [M]$



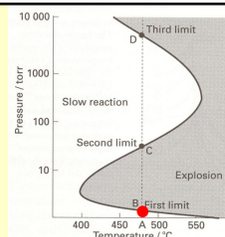
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Explosion of a hydrogen-oxygen mixture



1	$\text{H}_2 + \text{O}_2 \rightarrow \cdot\text{H} + \cdot\text{HO}_2$	chain initiation
2	$\cdot\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$	chain branching
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$	chain continuation
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{OH} + \cdot\text{H}$	chain branching
5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
6	$\cdot\text{H} \rightarrow \text{wall}$	chain termination
7	$\cdot\text{O} \rightarrow \text{wall}$	chain termination
8	$\cdot\text{OH} \rightarrow \text{wall}$	chain termination
9	$\cdot\text{HO}_2 + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}_2$	chain initiation*
10	$2 \cdot\text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	chain termination
11	$\text{H}_2\text{O}_2 + \text{M} \rightarrow 2 \cdot\text{OH} + \text{M}$	chain initiation
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination

1	$\text{H}_2 + \text{O}_2 \rightarrow \cdot\text{H} + \cdot\text{HO}_2$	chain initiation
2	$\cdot\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$	chain branching
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$	chain continuation
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{OH} + \cdot\text{H}$	chain branching
5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
6	$\cdot\text{H} \rightarrow \text{wall}$	chain termination
7	$\cdot\text{O} \rightarrow \text{wall}$	chain termination
8	$\cdot\text{OH} \rightarrow \text{wall}$	chain termination
9	$\cdot\text{HO}_2 + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}_2$	chain initiation*
10	$2 \cdot\text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	chain termination
11	$\text{H}_2\text{O}_2 + \text{M} \rightarrow 2 \cdot\text{OH} + \text{M}$	chain initiation
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination



below the 1st explosion limit:

chain termination reactions 6, 7, 8 remove the radicals
(radical loss on the wall)

→ no explosion

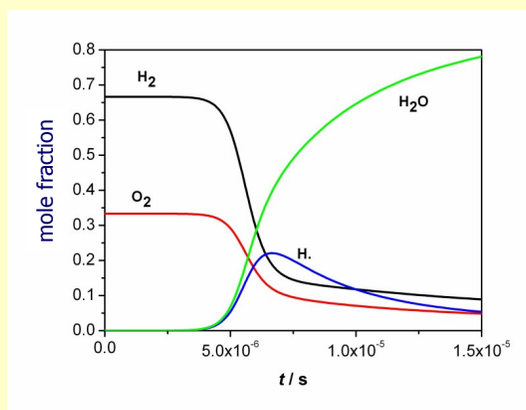
1	$\text{H}_2 + \text{O}_2 \rightarrow \cdot\text{H} + \cdot\text{HO}_2$	chain initiation
2	$\cdot\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$	chain branching
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$	chain continuation
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{OH} + \cdot\text{H}$	chain branching
5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
6	$\cdot\text{H} \rightarrow \text{wall}$	chain termination
7	$\cdot\text{O} \rightarrow \text{wall}$	chain termination
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9	$\cdot\text{HO}_2 + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}_2$	chain initiation*
10	$2 \cdot\text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	chain termination
11	$\text{H}_2\text{O}_2 + \text{M} \rightarrow 2 \cdot\text{OH} + \text{M}$	chain initiation
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination

between explosion limits 1 and 2:

2-3-4 chain branching steps

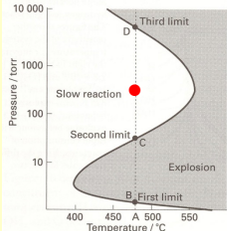
2	$\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{H} + \cdot\text{OH}$
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$
+ _____	
	$\cdot\text{H} + \text{O}_2 + 3 \text{H}_2 \rightarrow 3 \cdot\text{H} + 2 \text{H}_2\text{O} \rightarrow \text{explosion}$

Hydrogen-oxygen explosion between limits 1 and 2



H atom is a very reactive intermediate
 its concentration first increases quickly + starts to decay
 when the concentrations of H_2 and O_2 decreases
 concentration of H_2O : saturation curve

1	$\text{H}_2 + \text{O}_2 \rightarrow \cdot\text{H} + \cdot\text{HO}_2$	chain initiation
2	$\cdot\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$	chain branching
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$	chain continuation
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{OH} + \cdot\text{H}$	chain branching
5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
6	$\cdot\text{H} \rightarrow \text{wall}$	chain termination
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11	$\text{H}_2\text{O}_2 + \text{M} \rightarrow 2 \cdot\text{OH} + \text{M}$	chain initiation
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination



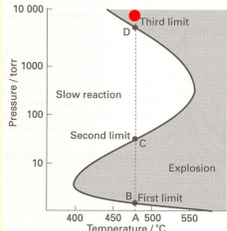
between explosion limits 2 and 3:

5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination

→ no explosion

M any species present
at the hydrogen/air explosion it is mainly N_2 O_2 H_2
but can be any other species (e.g. $\cdot\text{H}$ $\cdot\text{HO}_2$ $\cdot\text{OH}$)

1	$\text{H}_2 + \text{O}_2 \rightarrow \cdot\text{H} + \cdot\text{HO}_2$	chain initiation
2	$\cdot\text{H} + \text{O}_2 \rightarrow \cdot\text{OH} + \cdot\text{O}$	chain branching
3	$\cdot\text{OH} + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}$	chain continuation
4	$\cdot\text{O} + \text{H}_2 \rightarrow \cdot\text{OH} + \cdot\text{H}$	chain branching
5	$\cdot\text{H} + \text{O}_2 + \text{M} \rightarrow \cdot\text{HO}_2 + \text{M}$	chain termination*
6	$\cdot\text{H} \rightarrow \text{wall}$	chain termination
7	$\cdot\text{O} \rightarrow \text{wall}$	chain termination
8	$\cdot\text{OH} \rightarrow \text{wall}$	chain termination
9	$\cdot\text{HO}_2 + \text{H}_2 \rightarrow \cdot\text{H} + \text{H}_2\text{O}_2$	chain initiation*
10	$2 \cdot\text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	chain termination
11	$\text{H}_2\text{O}_2 + \text{M} \rightarrow 2 \cdot\text{OH} + \text{M}$	chain initiation
12	$\cdot\text{HO}_2 \rightarrow \text{wall}$	chain termination



above explosion limit 3

- high pressure \Rightarrow reaction step 5 produces much $\cdot\text{HO}_2$
- reaction steps 9 and 10 convert $\cdot\text{HO}_2$ to H_2O_2
- at high pressure the decomposition of H_2O_2 is fast (reaction 11) and it produces highly reactive OH radicals

→ explosion

Computer codes for the study of complex reaction systems

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Reaction kinetics simulation codes

WINPP/XPP Windows simulation code

solving systems of ODEs, DAEs and PDEs.

The user has to provide the rate equations \Rightarrow applicable for small systems only

<http://www.math.pitt.edu/~bard/classes/wppdoc/readme.htm>

KPP: Kinetic Preprocessor <http://people.cs.vt.edu/~asandu/Software/Kpp/>

production of the kinetic ODE from the reaction mechanism

numerical solution of stiff ODEs; sparse matrix routines

V. Damian, A. Sandu, M. Damian, F. Potra, G. R. Carmichael:

The Kinetic PreProcessor KPP - A software environment for solving chemical kinetics.

Comp. Chem. Eng. **26**, 1567-1579 (2002)

SUNDIALS: SUite of Nonlinear and Differential/ALgebraic equation Solvers

<https://computation.llnl.gov/casc/sundials/main.html>

MATLAB interface to the following solvers:

CVODE solves initial value problems for ordinary differential equation (ODE) systems

CVODES solves ODE systems and includes sensitivity analysis capabilities

ARKODE solves initial value ODE problems with additive Runge-Kutta methods

IDA solves initial value problems for differential-algebraic equation (DAE) systems

IDAS solves DAE systems and includes sensitivity analysis capabilities

KINSOL solves nonlinear algebraic systems.

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CHEMKIN

Developed at the SANDIA National Laboratories, Livermore, CA, USA

CHEMKIN-I (1975-1985)

CHEMKIN-II (1985-1995)

Simulation codes: SENKIN, PSR, PREMIX, SHOCK, EQLIB

+ utility programs, data bases

FORTTRAN codes, controlled by the input files

Kee R. J., Rupley F. M., Miller J. A.
 CHEMKIN-II: A FORTRAN *Chemical Kinetics Package*
for the Analysis of Gas-Phase Chemical Kinetics
 SANDIA report No. SAND79-8009B

AnSys <https://www.ansys.com/> (formerly ReactionDesign)

Commercial codes; source code is not provided

Chemkin 3.x,

Graphical User Interface (GUI) to CHEMKIN-II

Chemkin 4.x

really new solvers, graphical interface, versatile

Chemkin Pro

Chemkin + additional utility codes (e.g. pathway plotting)

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CHEMKIN simulation codes

<https://www.ansys.com/>

CHEMKIN → CHEMKIN -II → CHEMKIN 3 → CHEMKIN 4 → CHEMKIN PRO

CHEMKIN (1975–)

classified code

CHEMKIN-II (1986–)

classified code, then freeware

since CHEMKIN 3 (1996–)

commercial code (now: Ansys)

CHEMKIN-II simulation codes:

SENKIN

spatially homogeneous reactions

PREMIX

laminar premixed flames

SHOCK

shock tube simulations

PSR

perfectly stirred reactor simulations

Options of SENKIN:

adiabatic system, constant p pressure

adiabatic system, constant V volume

adiabatic system, $V(t)$ function

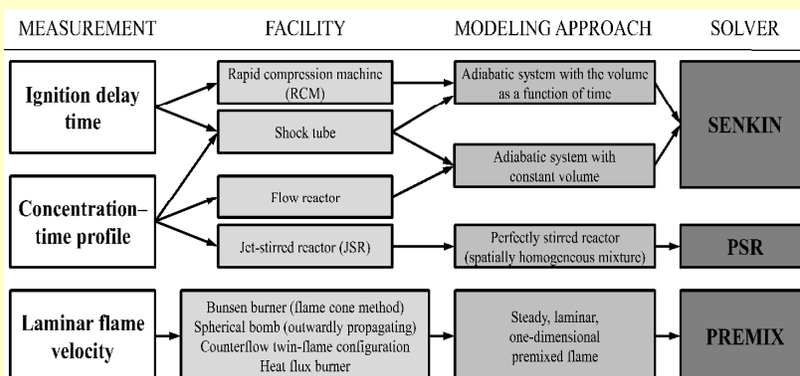
closed system, constant p , T

closed system, constant V , T

closed system, $p(t)$ and $T(t)$ function

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CHEMKIN-II simulation codes



Relation between the

- measured quantity
- experimental setup
- modelling approach
- the corresponding CHEMKIN-II simulation code

C. Olm, I. Gy. Zsély, R. Pálvölgyi, T. Varga, T. Nagy, H. J. Curran, T. Turányi:
Comparison of the performance of several recent hydrogen combustion mechanisms.
Combust. Flame, **161**, 2219-2234 (2014).

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CHEMKIN format of mechanisms

```

ELEMENTS H O N AR END
SPECIES
H2 O2 H2O H2O2 H O OH HO2 N2 AR
END
THERMO ALL
300.000 1000.000 5000.000
H2 H 2 0 0 0G 300.00 5000.00 1000.00 0 1
2.99142300E+00 7.00064400E-04 -5.63382900E-08 -9.23157800E-12 1.58275200E-15 2
-8.35034000E+02 -1.35511000E+00 3.29812400E+00 8.24944200E-04 -8.14301500E-07 3
-9.47543400E-11 4.13487200E-13 -1.01252100E+03 -3.29409400E+00 4
O2 O 2 0 0 0G 300.00 5000.00 1000.00 0 1
3.69757800E+00 6.13519700E-04 -1.25884200E-07 1.77528100E-11 -1.13643500E-15 2
-1.23393000E+03 3.18916600E+00 3.21293600E+00 1.12748600E-03 -5.75615000E-07 3
1.31287700E-09 -8.76855400E-13 -1.00524900E+03 6.03473800E+00 4
H2O H 2O 1 0 0G 300.00 5000.00 1000.00 0 1
2.67214600E+00 3.05629300E-03 -8.73026000E-07 1.20099600E-10 -6.39161800E-15 2
-2.98992E-06 3
6.96858E-04 4
H2O2 H 2O2 0 0 0G 300.00 5000.00 1000.00 0 1
4.57316E-14 2
-1.80069E-07 3
-4.62580E-04 4
H H 1 0 0 0G 300.00 5000.00 1000.00 0 1
2.50000000E+00 0.00000000E+00 0.00000000E+00 0.00000000E+00 0.00000000E+00 2
2.54716300E+04 -4.60117600E-01 2.50000000E+00 0.00000000E+00 0.00000000E+00 3
0.00000000E+00 0.00000000E+00 2.54716300E+04 -4.60117600E-01 4
...

```

for each species, the description of the temperature dependence of H , S and c_p
NASA polynomials: 2×7 parameters

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CHEMKIN format of mechanisms

REACTIONS	MOLES	KJOULES/MOLE			
H2+O		=>	OH+H	5.120E+04	2.67 26.27
OH+H		=>	H2+O	3.534E+04	2.02 18.95
H2+OH		=>	H2O+H		
H2O+H		=>	H2+OH		
O2+H+M		=>	HO2+M	2.100E+18	-1.80 .00
N2/0.67/ O2/0.4/ H2O/0./ AR/0.28/					
HO2+M		=>	O2+H+M	1.159E+20	-1.26 211.41
				6.890E+15	.00 -8.73
				3.801E+17	-.46 202.68
O2+H		=>	OH+O	9.756E+13	.00 62.11
OH+O		=>	O2+H	1.450E+13	.00 2.94
H2O2+O		=>	OH+HO2	6.620E+11	.00 16.63
OH+HO2		=>	H2O2+O	4.073E+08	.72 77.51
H2O2+OH		=>	H2O+HO2	7.830E+12	.00 5.57
H2O+HO2		=>	H2O2+OH	4.744E+11	.45 140.59
H2O2(+M)		=>	2OH(+M)	3.000E+14	.00 202.87
N2/0.4/ O2/0.4/ H2O/6.5/ AR/0.35/					
LOW / 3.000E+17		.00	190.40 /		
TROE / 1.0000		1.00	1.00 1040.00 /		
2OH(+M)		=>	H2O2(+M)	7.230E+13	-.37 .00
N2/0.4/ O2/0.4/ H2O/6.5/ AR/0.35/					
LOW / 5.530E+19		-.70	.00 /		
TROE / 1.0000		1.00	1.00 1040.00 /		
...					
END					

Arrhenius parameters A, n, E
3rd body collision efficiencies
Arrhenius parameters of k_0
Troe-parameters for the description of p -dependence_s

Alternatives to CHEMKIN

Cantera www.cantera.org

Open source code, available from SourceForge.net
 chemical equilibrium, homogeneous and heterogeneous kinetics
 reactor networks, 1D flames

Kintecus www.kintecus.com

Excel workbook; free for academic use
 Simulation of combustion, atmospheric chemical and biological systems

COSILAB www.softpredict.com

commercial combustion simulation and mechanism analysis code

- visualization of reaction pathways
- reduction of kinetic mechanisms
- simulation of reactor networks
- two-dimensional reactors and flames
- spray and dust flames

Alternatives to CHEMKIN 2

OpenSmoke (Milano; <http://www.opensmoke.polimi.it/>)

freely available program
numerical modelling of laminar reacting flows
built on the OpenFOAM framework
homogeneous reactions, heterogeneous reactions on catalytic surfaces

FlameMaster (Aachen; <http://www.itv.rwthachen.de/downloads/flamemaster/>)
free for academic use

- homogeneous reactor and perfectly stirred reactor calculations
- freely propagating premixed flames
- steady counter-flow diffusion flames

Chem1D (Eindhoven; <https://www.tue.nl/>)

flame simulations with both detailed and few-step mechanisms

laminar flames:

adiabatic, burner stabilized, ceramic burner stabilized and counterflow flames

special effects in laminar flames:

simulation of stretch, curvature, gas radiation

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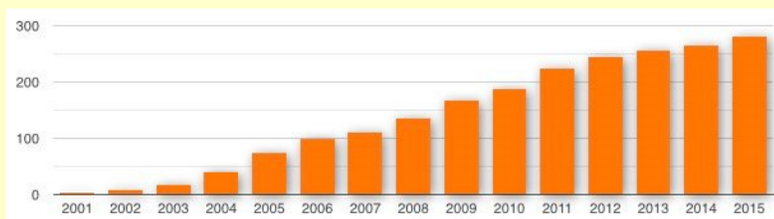
SBML

SBML: Systems Biology Markup Language <http://sbml.org/>

SBML model definition format was created
to promote the exchange of systems biology models
(similar to the role of CHEMKIN format in gas kinetics)

281 SBML-compatible software packages are available (January 2016)

The list of these simulation codes can be looked at <http://sbml.org/>



Increase of the number of SBML-based computer codes
(these include both academic and commercial codes)

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Copasi

COPASI (COMplex PATHway Simulator)

<http://copasi.org/>

Simulation and analysis of biochemical network models.

Free, support, but source code is not provided.

Homogeneous kinetic systems in interacting compartments
 Import and export of models in the SBML format (levels 1 to 3).
 Export of models in many format (XPP, C code, Latex).

- ODE-based and stochastic simulators
- stoichiometric analysis of the reaction networks
- optimization of models; parameter estimation
- local sensitivity analysis.
- time scale separation analysis
- characterization of non-linear dynamics properties (oscillations and chaos)

S. Hoops, S. Sahle, R. Gauges, C. Lee, J. Pahle, N. Simus, M. Singhal, L. Xu, P. Mendes, U. Kummer:
 COPASI — a COMplex PATHway Simulator. *Bioinformatics* **22**, 3067-3074 (2006)

Global uncertainty analysis codes

GUI-HDMR <http://www.gui-hdmr.de>

The GUI-HDMR software is based on the RS-HDMR approach, where all component functions are approximated by orthonormal polynomials using random (or quasi-random) samples. Calculation of up to second-order global sensitivity indices based on user supplied sets of input/output data. The component functions are approximated by up to 10th order orthonormal polynomials.

T. Ziehn, A. S. Tomlin: GUI-HDMR - A software tool for global sensitivity analysis of complex models
Environmental Modelling & Software, **24**, 775-785 (2009)

SimLab <https://ec.europa.eu/jrc/en/samo/simlab>

Developed at the EC Joint Research Centre (EC-JRC) in Ispra, Italy.

Versions up to 2.2: GUI based nice education tool

- (1) generation of random or quasi-random parameter sets
- (2) running the models (within SimLab or externally)
- (3) processing of the simulation results (FAST, Morris' and Sobol methods)
 visualisation of the outcome of uncertainty/sensitivity analyses.

SimLab versions from 3.0:

subroutine can be called from Fortran, Python, C++, or Matlab

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• **Respecth Collection**





RE
reaction kinetics



SPEC
spectroscopy



TH
thermochemistry

<http://respecth.hu>

reaction kinetics spectroscopy thermodynamics

- database of combustion and rate coefficient experimental data in XML format (in extended PRiMe format) for the hydrogen reactions
- code for reading and writing the XML files
- specification document of the XML data
- collection of Chemkin mechanisms
- programs for the analysis of mechanisms including *u-Limits*, *UBAC*, *JPDAP*

NEW USERS ARE WELCOME !!!

 Chemical Kinetics Laboratory, Institute of Chemistry, Eötvös University (ELTE), Budapest, Hungary 41



*Thank you for
your attention!*

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