

## 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







Equivalence ratio	
<b>fuel lean combustion</b> $CH_4+O_2$ mixture $\rightarrow CO_2 + H_2O_2$ $\varphi < 1; \lambda > 1$	$O + (O_2 \text{ remains})$
stoichiometric combustion $CH_4+O_2$ mixture $\rightarrow CO_2+H_2O$ $\varphi=1$ ; $\lambda=1$	
fuel rich combustion $CH_4+O_2 \text{ mixture} \rightarrow CO_2+H_2O$ $\varphi>1; \lambda<1$	<ul> <li>+ (CH<sub>4</sub> remains)</li> <li>In fact, no methane remains, because at high temperature methane decomposes to hydrogen and olefins!</li> </ul>
Stoichiometric ratios:	
$\begin{array}{l} \mathrm{H_2+}~0.5~\mathrm{O_2}\rightarrow\mathrm{H_2O}\\ \mathrm{CH_4+2~O_2}\rightarrow\mathrm{CO_2+2~H_2O} \end{array}$	$\varphi = \frac{n_{fuel}/n_{oxidizer}}{(n_{fuel}/n_{oxidizer})_{stoichiometric}}$
$\lambda$ : air equivalence ratio $\varphi$ : fuel equivalence ratio	(see ,, $\lambda$ sensor") $\varphi = 1/\lambda$



















	Collision efficiency parameters
М	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:	<i>e.g.</i> 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 $H_2O_2(+M) \rightarrow .OH + .OH(+M)$ :	
[M]= 5[H <sub>2</sub> O]+5.13[H <sub>2</sub> O <sub>2</sub> ]+0.8[O <sub>2</sub> ]+2.47[H <sub>2</sub> ]+1.87[CO]+1.07[CO <sub>2</sub> ]+0.67[Ar]+0.43[He]+[all <sup>1</sup> @thers]	









1	$H_{a} + O_{a} \rightarrow H + HO_{a}$	chain initiation
2	$H + O_2 \rightarrow OH + O_2$	chain branching
3	$.OH + H_2 \rightarrow .H + H_2O$	chain continuation
4	$:O + H_2 \rightarrow .OH + .H$	chain branching
5	$.H + O_2^- + M \rightarrow .HO_2 + M$	chain termination*
6	$H \rightarrow wall$	chain termination
7	$:O \rightarrow wall$	chain termination
8	$.OH \rightarrow wall$	chain termination
9	$.HO_2 + H_2 \rightarrow .H + H_2O_2$	chain initiation*
10	$2 : HO_2 \rightarrow H_2O_2 + O_2$	chain termination
11	$H_2O_2 + M \rightarrow 2 .OH + M$	chain initiation
12	$HO_2 \rightarrow Wall$	chain termination

















Reaction kinetics simulation codes		
WINPP/XPP Windows simulation code solving systems of ODEs, DAEs and PDEs. The user has to provide the rate equations ⇒ 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. <i>Comp. Chem. Eng.</i> <b>26</b> , 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:CVODEsolves initial value problems for ordinary differential equation (ODE) systemsCVODES solves ODE systems and includes sensitivity analysis capabilitiesARKODE solves initial value ODE problems with additive Runge-Kutta methodsIDAsolves initial value problems for differential-algebraic equation (DAE) systemsIDASsolves DAE systems and includes sensitivity analysis capabilitiesKINSOLsolves nonlinear algebraic systems.		

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 FORTRAN codes, controlled by the input files
Kee R. J., Rupley F. M., Miller J. A. CHEMKIN-II: A FORTRAN <i>Chemical Kinetics Package</i> <i>for the Analysis of Gas-Phase Chemical Kinetics</i> SANDIA report No. SAND79-8009B
AnSys https://www.ansys.com/ (formerly ReactionDesign) Commertial codes; source code is not provided
Chemkin 3.x,
Chemkin 4.x
really new solvers, graphical interface, versatile
Chemkin + additional utility codes ( <i>e.g.</i> pathway plotting) <sup>31</sup>

CHEMKIN simulation codes https://www.ansys.com/		
CHEMKIN $\rightarrow$ CHEMKIN -II $\rightarrow$ CHEMKIN 3 $\rightarrow$ CHEMKIN 4 $\rightarrow$ CHEMKIN PRO		
CHEMKIN (1975– ) CHEMKIN-II (1986– ) since CHEMKIN 3 (1996	classified code classified code, then freeware - ) commercial code (now: Ansys)	
CHEMKIN-II simulation codes:		
SENKIN PREMIX SHOCK PSR	spatially homogeneous reactions laminar premixed flames shock tube simulations perfectly stirred reactor simulations	
Options of SENKIN:		
adiabatic systen adiabatic systen adiabatic systen closed system, o closed system, o closed system, o	, constant <i>p</i> pressure , constant <i>V</i> volume , $V(t)$ function onstant <i>p</i> , <i>T</i> onstant <i>V</i> , <i>T</i> ( <i>t</i> ) and <i>T</i> ( <i>t</i> ) function	32







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 system	ns
<ul> <li>COSILAB www.softpredict.com</li> <li>commertial combustion simulation and mechanism analysis code</li> <li>visualization of reaction pathways</li> <li>reduction of kinetic mechanisms</li> <li>simulation of reactor networks</li> <li>two-dimensional reactors and flames</li> <li>spray and dust flames</li> </ul>	
	36





## 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 simulatons
- · 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 <i>Environmental Modelling &amp; Software</i> , <b>24</b> , 775-785 (2009)
<ul> <li>SimLab https://ec.europa.eu/jrc/en/samo/simlab</li> <li>Developed at the EC Joint Research Centre (EC-JRC) in Ispra, Italy.</li> <li>Versions up to 2.2: GUI based nice education tool</li> <li>(1) generation of random or quasi-random parameter sets</li> <li>(2) running the models (within SimLab or externally)</li> <li>(3) processing of the simulation results (FAST, Morris' and Sobol methods) visualisation of the outcome of uncertainty/sensitivity analyses.</li> </ul>

SimLab versions from 3.0: subroutine can be called from Fortran, Python, C++, or Matlab

40



