



Catalyst Preparation

FHI-AC Lecture Series on Heterogeneous Catalysis

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Catalyst preparation

- Target configuration (active sites) often unknown
- Material and pressure gap in heterogeneous catalysis
- High surface area material are desired ("porous" / "nano")
- Defects play a role
- Homogeneity is an issue
- Mostly empirical (trail & error) optimization
- Application requires reproducibility, scalability and economic / environmental feasibility

Progress of catalyst synthesis

Table 1.2	Survey	of selected	catalysts	with	their	main
application	15.					

Catalyst	Applications		
Ni/SiO	Hydrogenation		
K2O/Al2O1/Fe	Ammonia synthesis		
Ag/a-Al, O1	Epoxidation		
CrO _x /SiO ₂	Polymerization		
CoMoS2/y-Al2O1	Hydrotreating		
Co/SiO ₂	Fischer-Tropsch synthesis		
Cu/ZnO/Al ₂ O ₁	Methanol synthesis		
Zeolite Y composite	Catalytic cracking		
Pt/Mordenite	Hydroisomerization of light alkanes		
V2O5/TiO2	NO _x abatement		
Pt/C	Hydrogenation; fuel cell		







C-atoms. Data from 1940 to 2010 with data points from left to right at years 1940, 1990, and 2010, respectively.





Figure 1.2 Development of hydrodesulfurization (HDS) catalyst activity over the years [6].

Catalyst Microstructures



a) Supported catalyst



b) Porous bulk catalyst



c) Skeletal catalyst







Distribute a liquid precursor *uniformly* on a pre-formed support

Form acitve phase and stabilizer by thermal treatment of a *homogeneous* solid precursor

Leach pores into a previously *homogenous* alloy

Aspects of solid state chemistry

- Reactivity of solids is limited by slow diffusion processes: Increase in temperature required for solid-solid reactions (T > 600℃)
 - Formation of thermodynamically stable compounds of high density (low porosity), high crystallographic symmetry and low variation in building blocks
 - High temperatures favour sintering
- Catalyst preparation procedures avoid high temperatures
 - Homogeneous distribution achieved by other measures than diffusion
 - Liquid phases involved
 - Catalysts are meta-stable systems



Ni-Al distribution

Outline

- Impregnation
 - Support materials
 - Key experimental parameters
 - Example: 3-way catalyst
- Co-precipitation
 - Precursor chemistry
 - Example: Methanol synthesis catalyst
- Skeletal catalysts
 - Leaching of alloys
 - Example: Raney nickel
- Hydrothermal synthesis
 - Principles of hydrothermal synthesis
 - Example: Zeolites
- Summary

Impregnation: Role of supports

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- Why (most) catalysts need to be supported: Sintering
- Support:
 - carries the active components
 - provides surface area (porosity)
 - keeps high dispersion of active component
 - is thermostable and chemically inert (?)
- Maximum active surface area of the active component desired
 - per unit weight (expensive precious metals, typical loading ~ 1 wt.-%)
 - per unit volume (base metals, oxides, typical loadings ~20 wt.-%)



Preparation of supported catalysts

- Deposition of a precursor of the active phase onto the surface of the support
 - At the solid-liquid (wet) interface
 - Impregnation
 - Deposition precipitation
 - At the solid-gas (dry) interface
 - Vapor deposition
- Transformation of the precursor into the active phase







Figure 1.5 Three generations of silica-supported cobalt catalyst. (a) 50 wt% Co/kieselguhr prepared by precipitation; (b) 18 wt% Co on silica gel obtained by calcination in air flow; and (c) 18 wt% Co on silica gel obtained by calcination in NO/He flow.

Typical supports

- Alumina
 - mostly γ -phase
 - SA up to 400 m^2g^{-1}
- Silica
 - amorphous
 - SA up to 1000 m²g⁻¹
- Carbon
 - amorphous, unstable in oxidative environment
 - SA up to 1000 m²g⁻¹
- Zeolites / ordered mesoporous materials
 - highly defined pore system
- also: TiO₂, MgO, ...



Synthesis of supports: Alumina

- Porous aluminas are obtained by controlled dehydration of Al(OH)₃ (Gibbsite, Bayerite) or AlO(OH) ((Pseudo-)Boehmite)
 - transition aluminas: $AI_2O_3 \cdot x H_2O$
 - active aluminas: γ and η
 - full dehydration: non-porous α Al₂O₃ (corundum)
- The non-porous hydroxides can be precipitated from AI salt solutions by increasing pH
 - Bayerite: high T and pH
 - (Pseudo-)Boehmite: low pH
- High purity aluminas are prepared by hydrolysis of Alalcoholates



Surface charge and Zeta potential





R: Partikelradius κ⁻¹: rezipr. Debyesche Abschirmlänge

Surface charge of oxide supports

- In contact with water hydroxyl groups are formed on the oxide surface (M-OH)
- M-OH may behave as Brønstedt acids or bases
 - − M-OH + OH⁻ \rightarrow M-O⁻ + H₂O
 - − M-OH + H⁺ \rightarrow M-OH₂⁺
- Surface charge depends on solution pH
- Zeta (ζ-) potential is an estimation of surface charge





Zeta potential measurement



Impregnation

- Low volume of precursor solution (usually a transition metal salt) is mixed with the support and the mixture is dried
- The whole precursor is expected to retain on the support (no washing)
- Precursor solution contains decomposable counter ions and ligands (NH_4^+ , NO_3^- , $C_2O_4^{2^-}$, ...)
- Two types
 - Pores of the support are initially filled with air: Capillary impregnation
 - Pores of the support are initially filled with liquid: Diffusional impregnation



Capillary vs. diffusional impregnation

Support of pore volume $V_{\mbox{\tiny p}}$ is in contact with the precursor solution of volume V

- V_p = V: dry or incipient wetness impregnation
- exothermal process, increase in T
 - Pre-treatment with super-heated steam
- air bubbles are trapped as the solution is sucked into the pores by capillary forces and may burst the support particles lowering the mechanical strength
 - Operation in vacuum, usage of surfactants
- filling of pores theoretically finished in seconds, but trapped air leads to delay of the process and possibly inhomogeneous distribution of the active component

- $V_p < V$: wet impregnation
- Pores of the support are pre-filled with pure solvent
- Solvent-saturated support is immersed into the precursor solution
- Concentration gradient between external precursor solution and pores leads to migration of precursor salt into the pores
- non-exothermic, but longer process

Distribution of the active phase



- If interaction is too high strong adsorption on the edge sites is observed
 - precursor concentration is locally decreased, no concentration gradient between inner particle and outer shell, migration of precursor is hindered → eggshell distribution

 Homogeneous distribution on the macroscale is not always desired

 Distribution of the active phase is determined by the interaction of the precursor and the support surface (adsorption)

Precursor = metallic complex (MC), competitor (A)



1: roughly same affinities of MC and A

- 2: high affinity of MC; no competition
- 3: high affinity of A, low affinity of MC; low A/MC ratio
- 4: high affinity of A, low affinity of MC; high A/MC ratio

Key experimental parameters I

- IEP (alumina: pH ~9, silica: pH ~2)
 - electrostatic interaction of solid oxide surface and precursor complex (anions adsorbed for pH < 9, cations for pH > 9)
 - can be regulated by varying T (IEP at higher pH at higher T), selective adsorbates (Na⁺: abrupt increase of pzc/IEP, F⁻: decrease of pzc/IEP)
- pH
 - determines the surface charge of the solid oxide
 - determines the solution species of the precursor complex (NH₃- or Pts/Ptr H₂O-complexes, monomers or oligomers, ...)
- Type of ligands
 - charge, bulkiness, stability, ...
- Competitive ions
 - blocking of edge sites forcing precursor complexes further into the pores of the support particle







Key experimental parameters II

- Solvent
 - influences the interaction of solvated complex and surface of support
- **Kinetics**
- Drying
 - if interaction of precursor and oxide surface is too low, precursor species may migrate out again as the liquid leaves the pores





(b) high drying rate to 110°C; and (c) drying at 200°C. (Data have been collected



Three-way-catalyst

Co-precipitation

- Precipitation of a mixed solution to form a precursor containing several species in a homogeneous distribution (preferably within a single phase)
- Thermal decomposition of counter ions by calcination (NO₃⁻, CO₃²⁻, NH₄⁺, C₂O₄²⁻,...),
- Sometimes further thermal treatment (activation)

Precipitation – Nucleation & growth

- Homogeneous nucleation
 - $\Delta G = \Delta G_{\text{bulk}} + \Delta G_{\text{interface}}$ $(+ \Delta G_{\text{others}})$
 - r_{critical}: size of the nucleus
 - nucleus: smallest solidphase aggregate of atoms, which is capable of spontaneous growth
 - The more nucleation is favoured compared to growth, the smaller the particles will be
 - Nucleation is temperaturedependent

Fig. 2 Simplified scheme for the formation of a solid product from solution. From the metal ions a precursor species is formed, for example by hydrolysis or raising the pH. When the concentration of the precursor species exceeds the nucleation threshold, precipitation of the product begins, consuming the precursor by nucleation and growth. New nuclei are formed only in the shaded area.

Cu/ZnO methanol synthesis catalysts

Cation hydrolysis

Figure 7.3 Precipitation processing routes: forward (base to acid) and reverse (acid to base) precipitation, continuous precipitation, and corresponding pH profiles.

Parameters for Cu/ZnO/(Al₂O₃) preparation

"Chemical memory"

C. Baltes, S. Vukojević, F. Schüth, J. Catal. 258 (2008) 334,

Effects of ageing

The importance of precursor chemistry

The right precursor composition

Structuring of the Cu/ZnO catalysts

undefined compact shape size ca. 100 nm

thin needles size ca. 20 × 200 nm

spherical particles size ca. 10 nm

Skeletal catalysts

- Skeletal catalysts are prepared from an M-Al alloy (M: Ni, Cu, ...) by leaching of Al with sodium hydroxide (20-40 wt.-%)
- Alloys are prepared by melting the active metal and Al, quenching and crushing
- Porosity can be controlled by composition of the alloy, T of leaching
- No activation necessary

2 M-AI_(s) + 2 OH⁻ + 6 H₂O

$$\downarrow$$

2 M_(s) + 2 [AI(OH)₄]⁻ + 3 H_{2(g)}

Skeletal Cu catalysts prepared from a 50 wt.-% Cu-Al alloy

Raney nickel

- Used for hydrogenation reactions and as electrocatalyst
- 5 times more active than supported Ni catalyst used before (M. Raney, 1925)
- Ni-Al alloys (40-50% Ni) produced by dissolution of Ni in molten Al and quenching
- Phases: Ni₂Al₃, NiAl₃
- NiAl₃ is leached more rapidly
 → smaller crystallite, but
 mechanically weak
- Ni₂Al₃ is more difficult to leach → strong residual material

- BET SA 50-100 m²g⁻¹
- crystallite size and pore diameter: 2 – 6 nm
- highly pyrophoric, has to be stored under liquid (water pH~9)

Hydrothermal synthesis

- Educts and solvent (water) are heated in a close reaction vessel (autoclave)
- At T > 100°C an autogeneous pressure builds up and the properties of water change
 - viscosity decreases → higher mobility of ions
 - water self dissociation increases → hydrolysis favoured
- If 2-5 % of educts dissolve (sometimes mineralizer required) → rearrangement of building blocks (SBUs) to form new structures
- Conditions mild enough to allow crystallization of open framework structures

Zeolites

- General composition M_{x/n}(Al_xSi_vO_{2(x+v)})-z H₂O
 - M: cation of charge n
 - M: H⁺, group 1 or 2 or organic cation
- Si/Al varies between 1 and ∞
- Zeolite type obtained is determined by synthesis parameters

Zeolite A: $Na_2[(AIO_2)_{12}(SiO_2)_{12}]$ -27 H₂O

cage diameter: window diameter:

1140 pm 410 pm

Properties of zeolites

- highly defined pore structure
- application as molecular sieves
- inner surface accessible for catalysis
- ion exchange possible
- proton exchanged zeolites: solid state acids
- after heating: Lewis acids
- shape selective catalysts
- used for catalytic cracking

Hydrothermal synthesis of zeolites

- Si- and Al-sources form a mixed hydrogel
- OH⁻ as mineralizer (pH 9-13)
- Formation of polyanionic species in solution
- Nucleation and growth
 - by addition of preassembled elementary units
 - by aggregation

Template assisted synthesis

- Building blocks present in solution arrange around a structure-directing agent
- SDA acts as a template for the pore structure
- SDA can be removed by heating

Summary

- Supported catalysts can be prepared by impregnation
 - The interaction of the support surface and the precursor species in solution is the key factor for a homogeneous distribution
- Co-precipitation may provide intimate porous mixtures of active and refractory components
 - Properties of the precipitate pre-determine the final catalyst's properties, complex processes
- Skeletal catalysts exhibit high metal surface areas
 - Leaching process parameters control porosity
- Hydrothermal synthesis yields crystalline porous framework materials
 - Template assisted synthesis can lead to ordered and well defined pore structure
- (Other important preparation methods for catalysts are deposition precipitation, sol-gel processes, chemical vapour deposition, mechnano-chemical routes, solid state reactions, ...)

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