

Heterogeneous phase enzyme reactions

Advantages/disadvantages:

Advantages:

- homogeneity of the system,
- enzyme does not need previous preparation - (over isolation and purification)

Economic disadvantages:

- Enzymes are expensive, 1-10- \$/mg
- can be used only once, after reaction they are to be discarded...

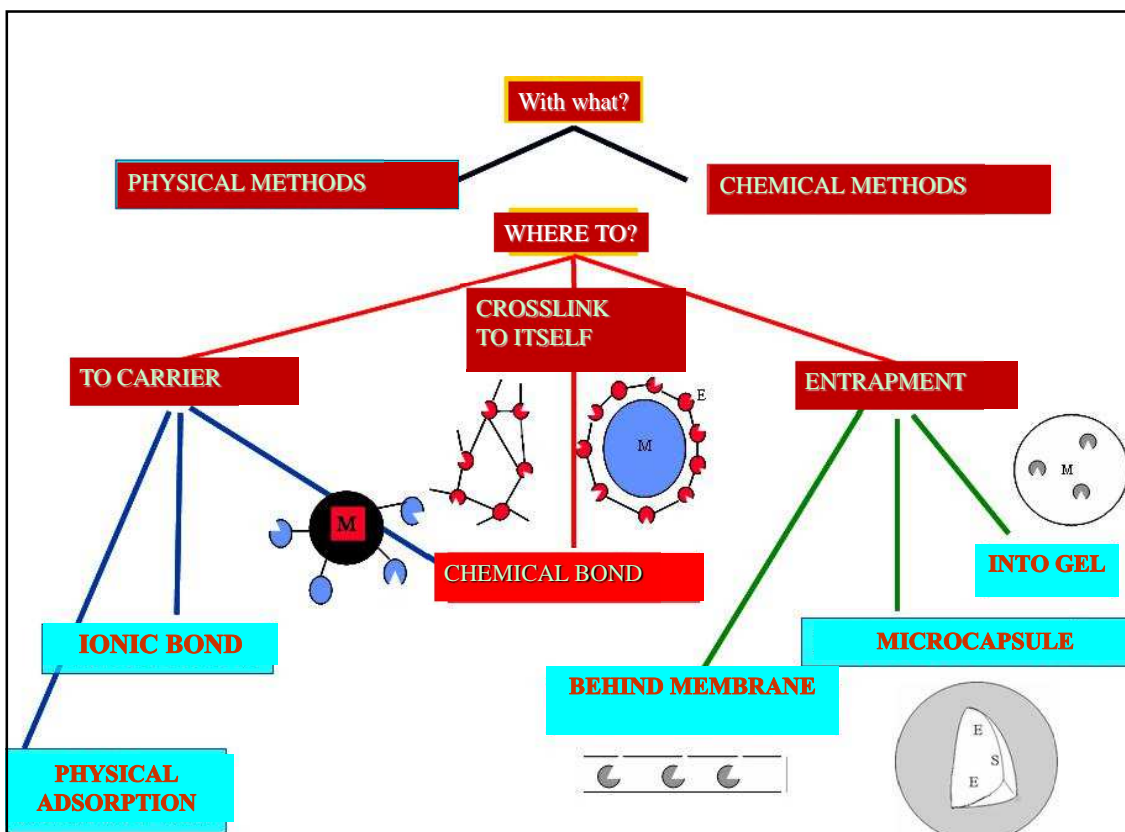
Technological disadvantage:

- Proteins contaminate products

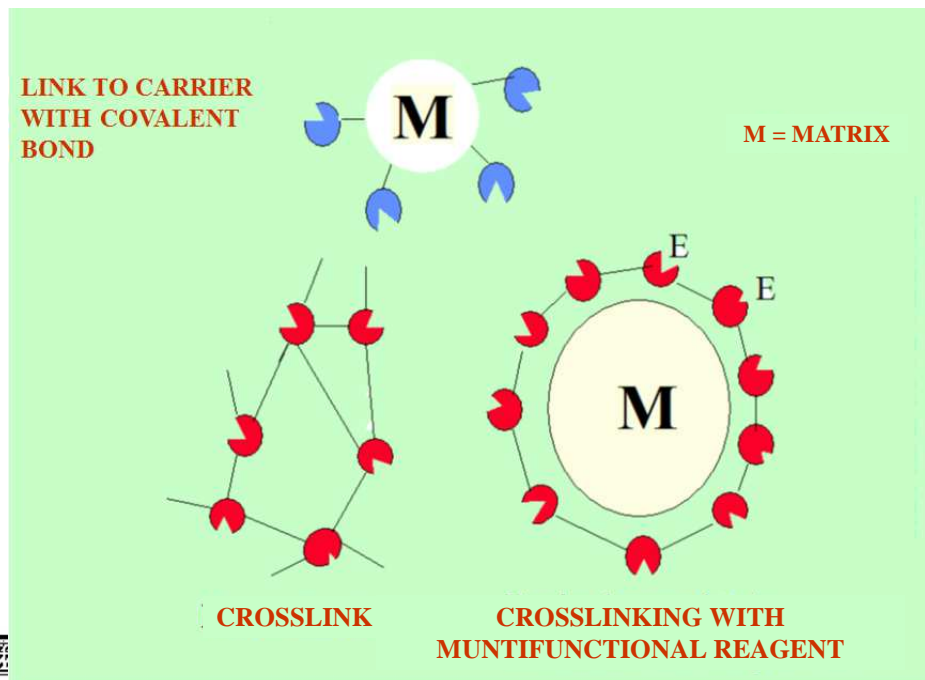


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CHEMICAL METHODS



CHEMICAL METHODS

Covalent bond between non essential amino acid sidechain(!)
and water insoluble matrix with function groups



CARRIERS :

natural polymers: agar, agarose, chitin, cellulose, collagene,....

synthetic polymer: polyurethane, polystyrene, nylon, ...

inorganics: glass, aluminium, silicagel, magnetit,...



CHEMICAL METHODS

Building of covalent bond:

free α -, β - or γ -COOH , α -, β -NH₂ groups
phenyl-, OH-, SH- imidazole-groups

STEPS:

1. Activation of carrier (arm and reactive X-group),
2. Creating covalent bond between enzyme and activated carrier.

Protection of the active sites: S or analog

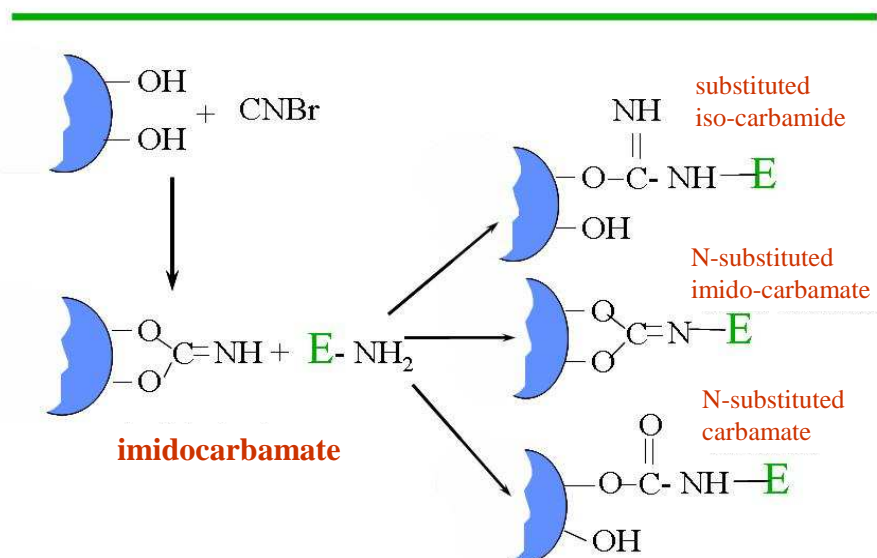


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MATRIX: vicinal -OH groups like:

cellulose, Sepharose, Sephadex



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Origin of carbohydrate matrix

Glucose → dextrane → Sephadex®

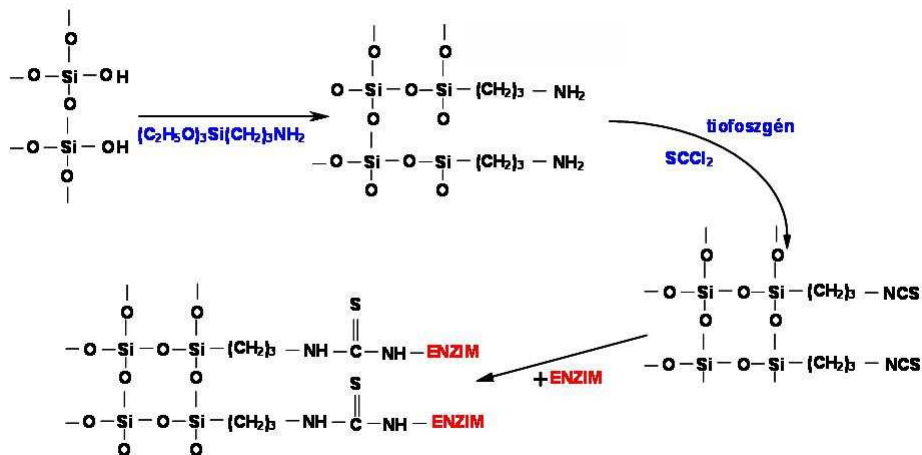
Alga → agar(ose) → Sepharose®



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Immobilization onto glass surface

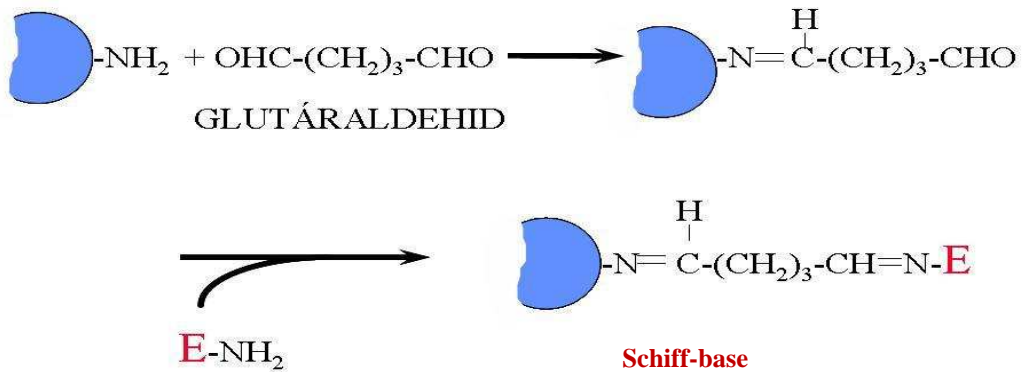


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Chemical methods: bifunctional molecules

MATRIX: -NH_2 groups like:

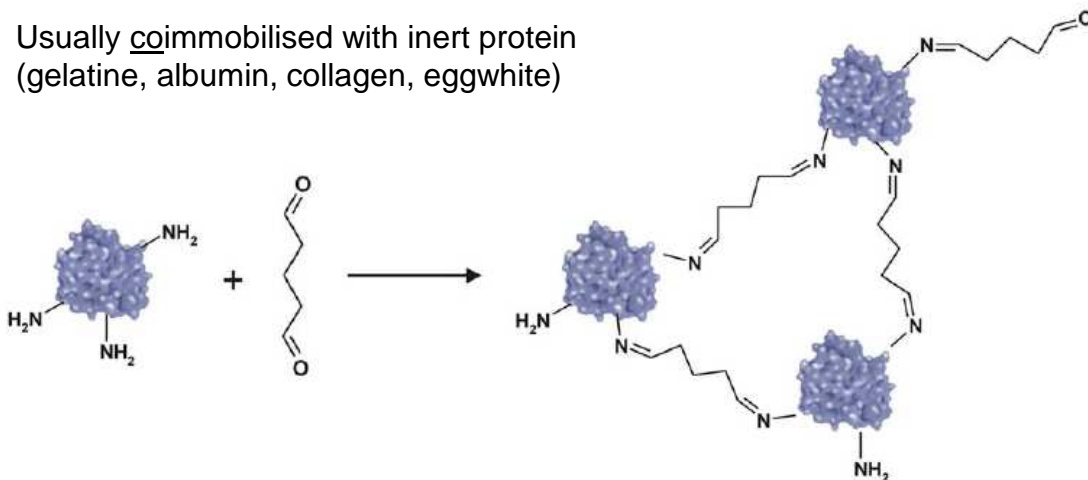
AE-cellulose, DEAE-cellulose, collagen, chitin, nylon...



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Chemical methods: **crosslinking**

Usually coimmobilised with inert protein
(gelatine, albumin, collagen, eggwhite)



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CLEC = Cross-Linked Enzyme Crystals



Scanning electron microscopic view of CLEC laccase

Surface area (m²/g) 2.456

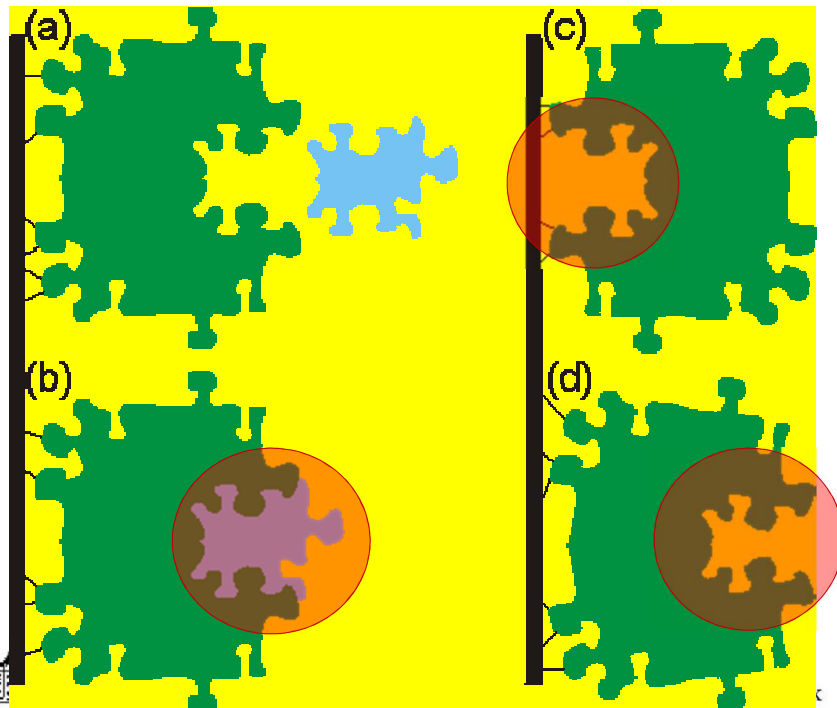
Preparation and characterization of cross-linked enzyme crystals of laccase, J. J. Roy, T. E. Abraham Journal of Molecular Catalysis B: Enzymatic 38 (2006) 31–36

Cross-linked Enzyme crystal of PNP (purine nucleoside phosphorylase)



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Possible effect of chemical immobilisation: Specific activity loss



PHYSICAL METHODS

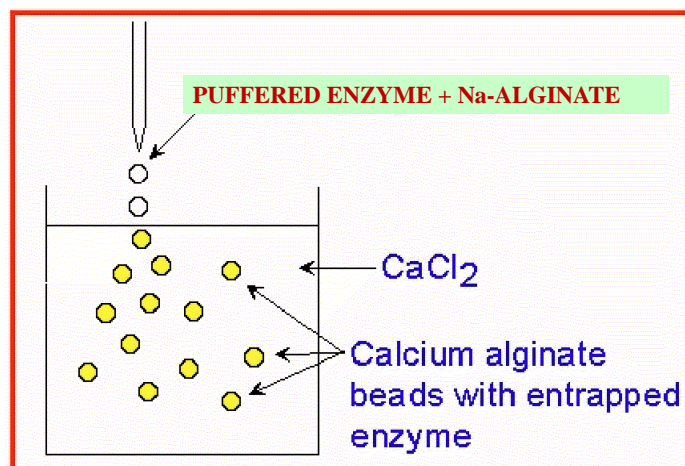
1. Adsorption e.g. on *ionexchanger resins* – nonspecific, easily desorps (pH)
2. Gel entrapment
3. Microencapsulation
4. Closing behind membrane



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ALGINATE GEL ENTRAPMENT



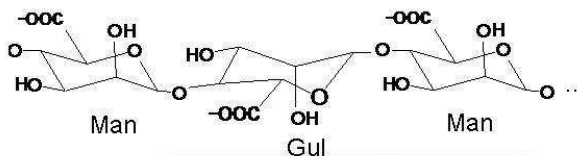
ALGINATE: poly- β -D-mannuronic acid (1 \rightarrow 4),-guluronic acid
Hydrophil colloid, linear polymer *Macrocystis pyrifera*



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Gel forming polysaccharides

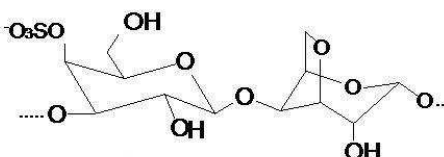
Alginate: heteropolymer of mannuronic acid and guluronic acid, 1,4-bonds



polyanionic

Solvent: water gel: Ca^{++} , Zn^{++} , Al^{3+}

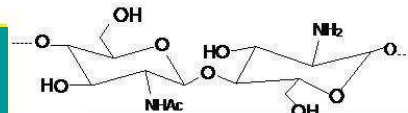
κ -carragenan: helical bio-polymer of 3,6 anhydro-galactose



polyanionic

Solvent: water gel: Ca^{++} , K^{+}

chitosan: partially deacetylated N-acetyl-glucoseamin polymer

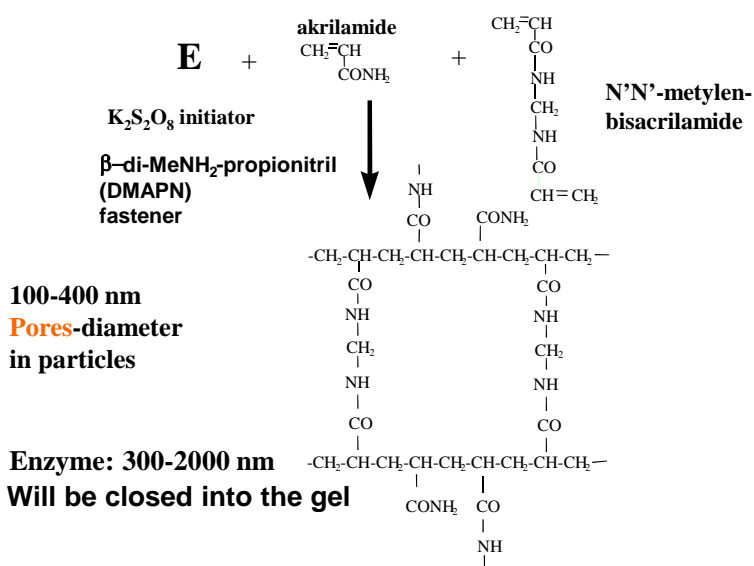


polycationic

Solvent: acetic acid, water
gel: polyphosphates, pH-change



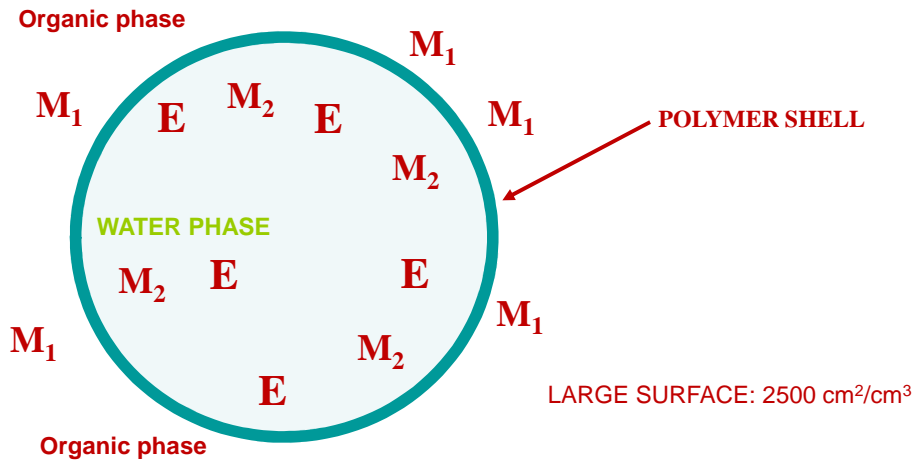
Poly-acrylamide gel entrapment



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Physical methods: microencapsulation

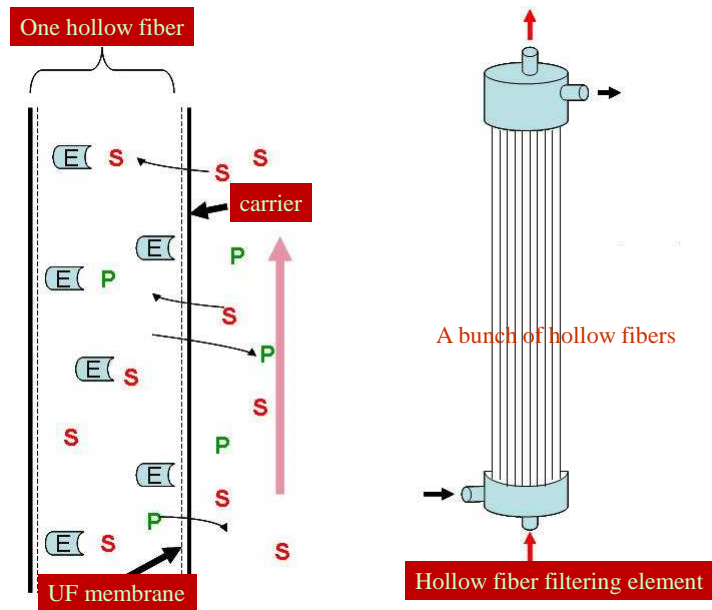
stable polymeric membranes



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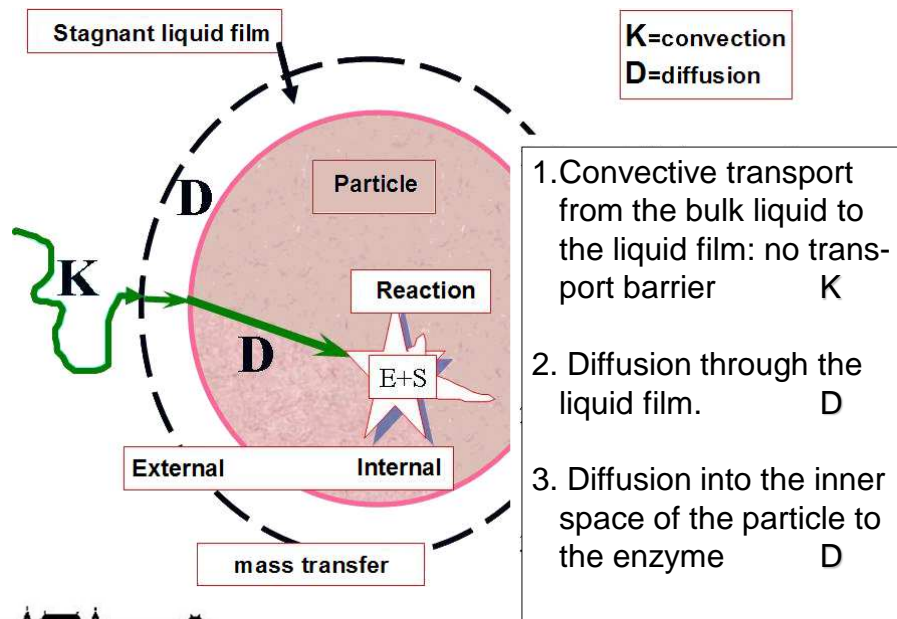
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Ultrafiltration membrane



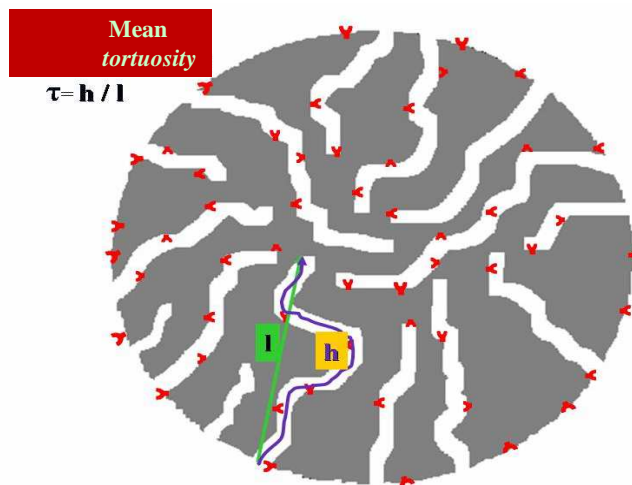
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Kinetics of immobilised enzymes



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Tortuosity



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Pros/cons about immobilised enzymes

Dissoved enzymes

- Advantages
- homogeneous system
 - no preparation needed
 - no mass transfer limitation

- Disadvantages
- expensive (1-10-50 \$/mg)
 - discarded after use
 - contamination of product
 - only batch technology



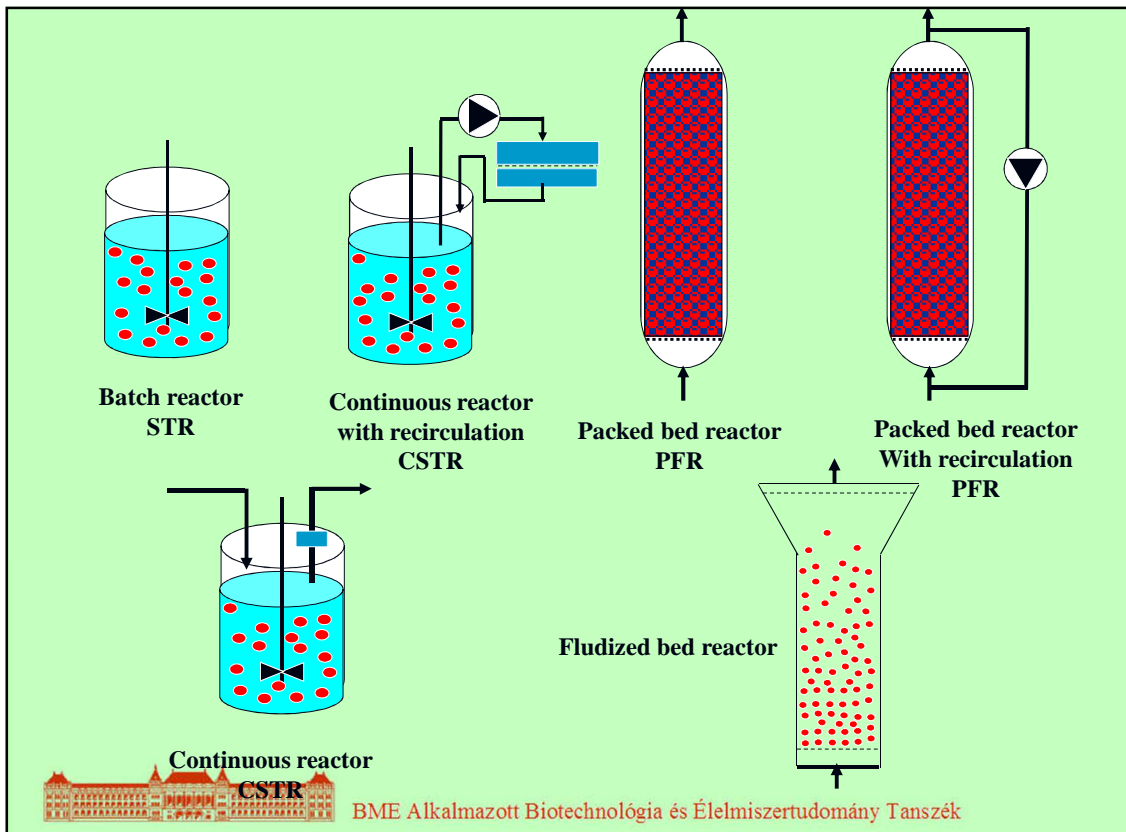
Pros/cons about immobilised enzymes

Immobilised enzymes

- Advantages
- No contamination of product
 - Easily separable
 - Possible reuse
 - Also continuous technologies
 - Easy termination
 - Increasing stability

- Disadvantages
- Expensive preparation need
 - Loss in enzyme activity
 - Diffusion barrier





Industrial application of immobilised enzymes

Aminoacilase	resolution of D,L-amino acids
Glucose-isomerase	conversion of glucose to glucose+fructose 1:1 mixture
Penicillin-amidase	preparation of 6-amino-penicilloic acid
β -galactosidase	hydrolysis of lactose to glucose+galactose
Lipase	hydrolysis and transesterification of lipids
Thermolysin	Preparation of aspartame

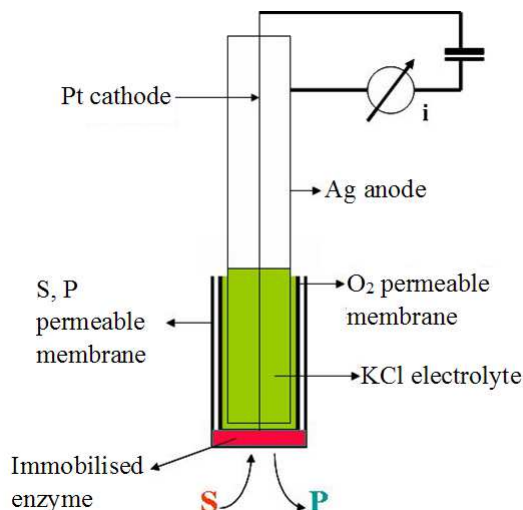
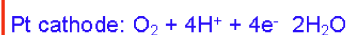
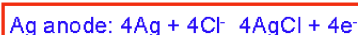


Enzyme electrode

Based on an amperometric electrode for dissolved oxygen measurement. It is covered with an enzyme producing or consuming oxygen.

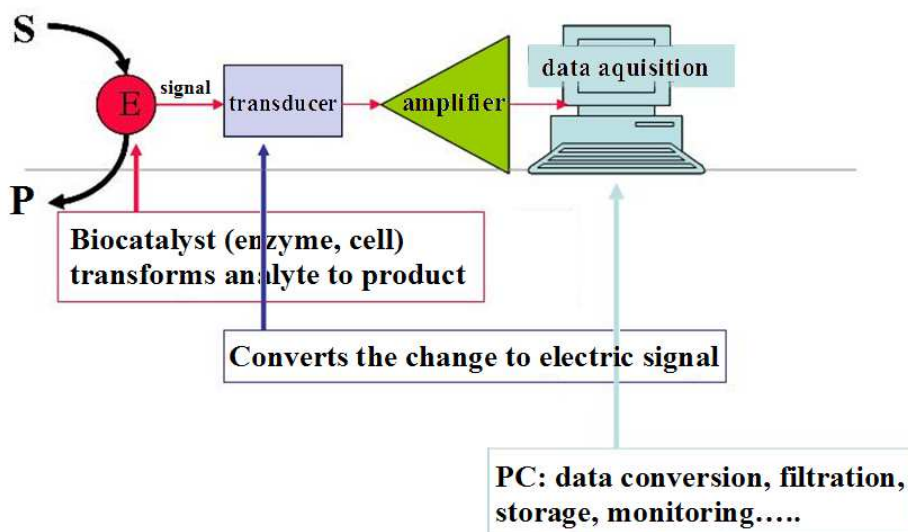
Eg. glucose oxydase + catalase.

The electrode reaction:



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BIOSENSOR



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Analytical enzyme applications

In these cases not the activity of enzyme is measured but the concentration of an analyt molecule.

1. Determination of S
2. Determination of I
3. Marker reactions (eg. in immunoassays)

Enzyme Linked Immunosorbent Assay (ELISA)
diagnostical, research purposes

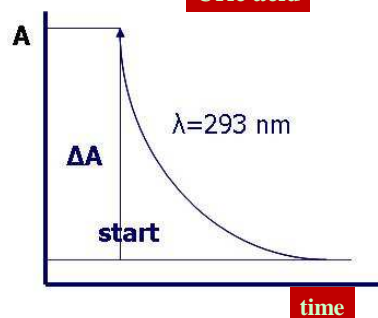
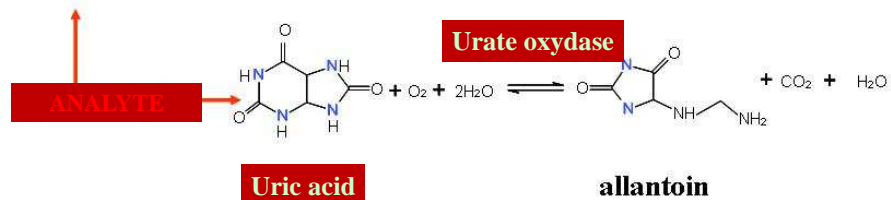


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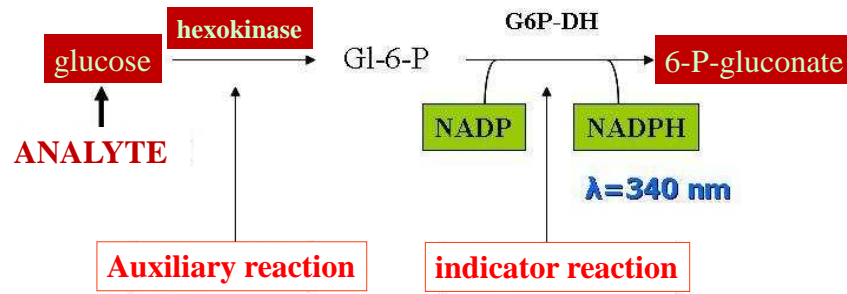
End-point measurement of substrate

The whole amount of substrate is converted – change is measured



Indicator reaction

If S and P are not observable → an enzymatic indicator reaction makes it measurable.



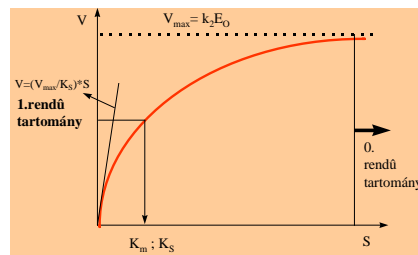
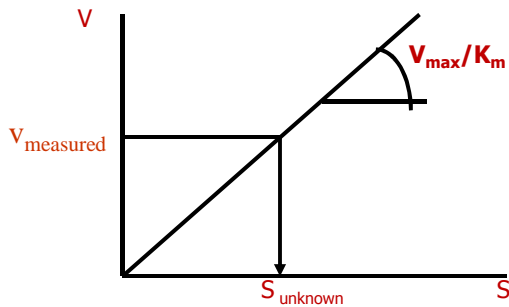
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Kinetic measurement of S

At small substrate concentrations the reaction rate changes linearly with S concentration (M-M kinetics).

If $S \ll K_m \rightarrow V \sim V_{max}/K_m \cdot S$

$\swarrow -dS/dt$
 $\searrow dP/dt$



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