

### 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



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

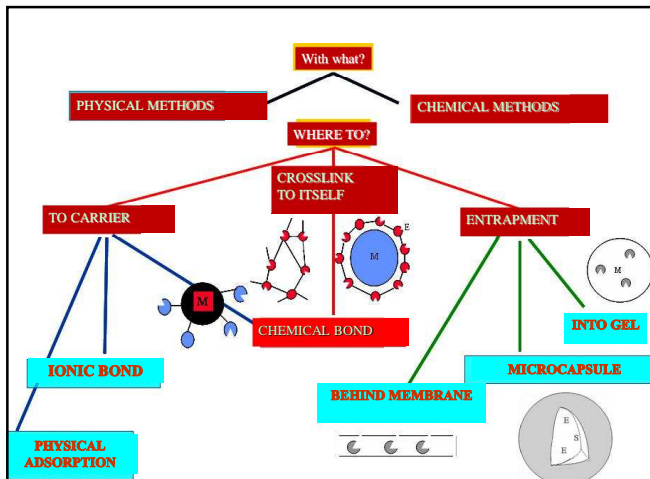
---

---

---

---

---




---

---

---

---

---

---

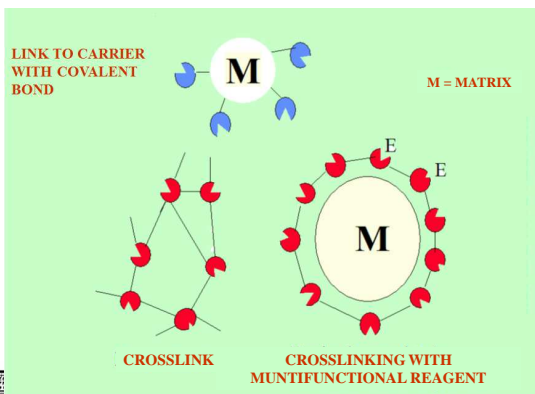
---

---

---

---

### CHEMICAL METHODS




---

---

---

---

---

---

---

---

---

---

### CHEMICAL METHODS

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


$$\text{---X} + \text{E} \longrightarrow \text{---E} + \text{X}$$

CARRIERS :

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

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

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



4

---

---

---

---

---

---

---

---


### CHEMICAL METHODS

Building of covalent bond:  
 free  $\alpha$ -,  $\beta$ - or  $\gamma$ -COOH ,  $\alpha$ -,  $\beta$  -NH<sub>2</sub> 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



5

---

---

---

---

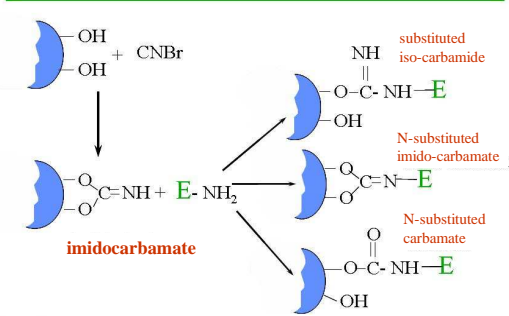
---

---


---

---

**MATRIX:** vicinal -OH groups like:  
 cellulose, Sepharose, Sephadex



The diagram illustrates the reaction of a matrix with two adjacent hydroxyl groups (-OH) and cyanogen bromide (CNBr). This reaction forms an imidocarbamate intermediate. This intermediate then reacts with an enzyme (E-NH<sub>2</sub>) to form three different covalent bonds: a substituted iso-carbamide (NH-C(=O)-NH-E), an N-substituted imido-carbamate (O=C-N-E), and an N-substituted carbamate (O=C-NH-E).



6

---

---

---

---

---

---

---

---

### Origin of carbohydrate matrix

Glucose → dextrane → Sephadex®

Alga → agar(ose) → Sepharose®




---

---

---

---

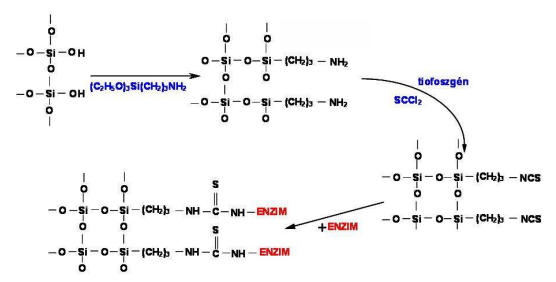
---

---

---

---

### Immobilization onto glass surface




---

---

---

---

---

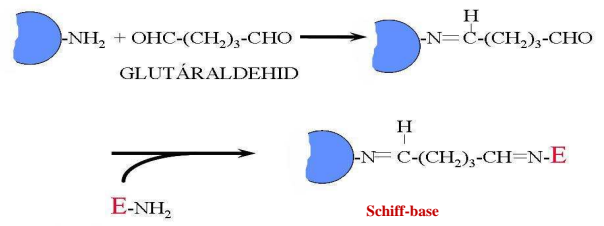
---

---

---

### Chemical methods: bifunctional molecules

**MATRIX:** -NH<sub>2</sub> groups like:  
 AE-cellulose, DEAE-cellulose, collagen, chitin, nylon...




---

---

---

---

---


---

---

---

**Chemical methods: crosslinking**

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

 BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---


---

---

**CLEC = Cross-Linked Enzyme Crystals**

Scanning electron microscopic view of CLEC laccase  
Surface area (m<sup>2</sup>/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

 BME Alkalmazott Biotechnológia és

---

---

---

---

---

---

---

---

Possible effect of chemical immobilisation: Specific activity loss

12

---

---

---

---

---


---

---

---

### PHYSICAL METHODS

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



15  
BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

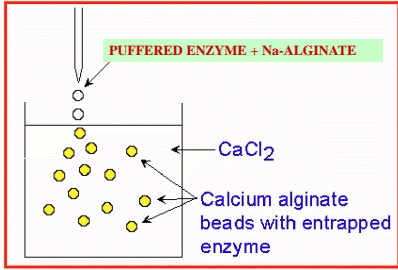
---

---


---

---

### ALGINATE GEL ENTRAPMENT



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



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

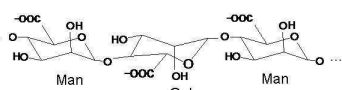
---

---

---

### Gel forming polysaccharides

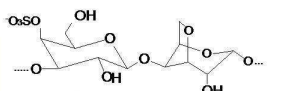
Alginate: heteropolymer of mannuronic acid and guluronic acid, 1,4-bonds



Man Gul

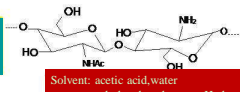
polyanionic      Solvent: water    gel: Ca<sup>++</sup>, Zn<sup>++</sup>, Al<sup>3+</sup>

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




polyanionic      Solvent: water    gel: Ca<sup>++</sup>, K<sup>+</sup>

chitosan: partially deacetylated N-acetyl-glucosamin polymer



NH<sub>2</sub> NH<sub>2</sub>

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



---

---

---

---

---

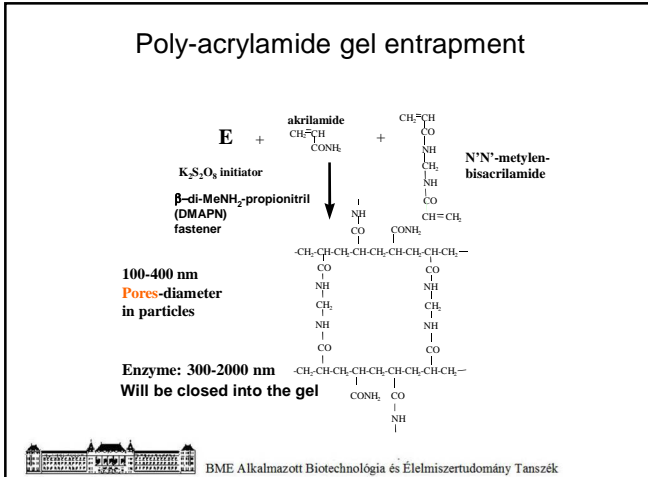
---

---

---

BME Department of Applied Biotechnology and Food Science

5




---

---

---

---

---

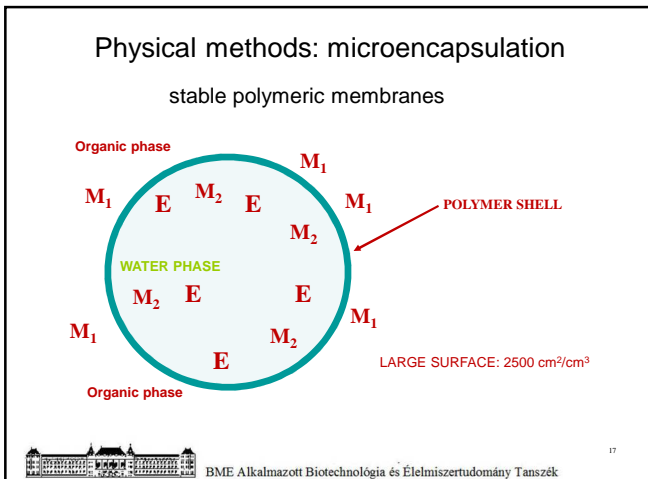
---

---

---

---

---




---

---

---

---

---

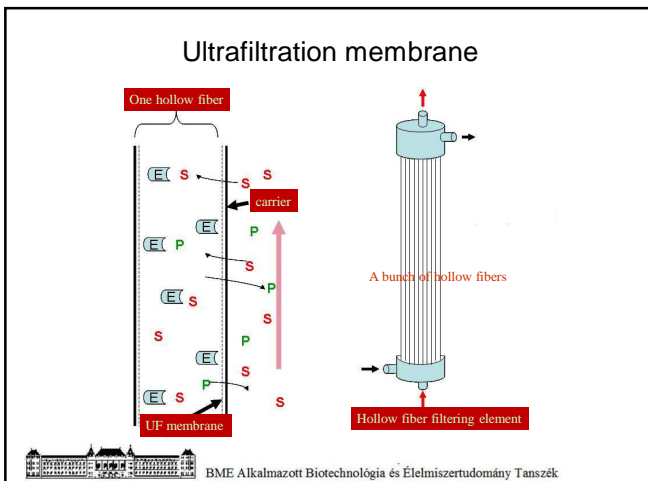
---

---

---

---

---




---

---

---

---

---

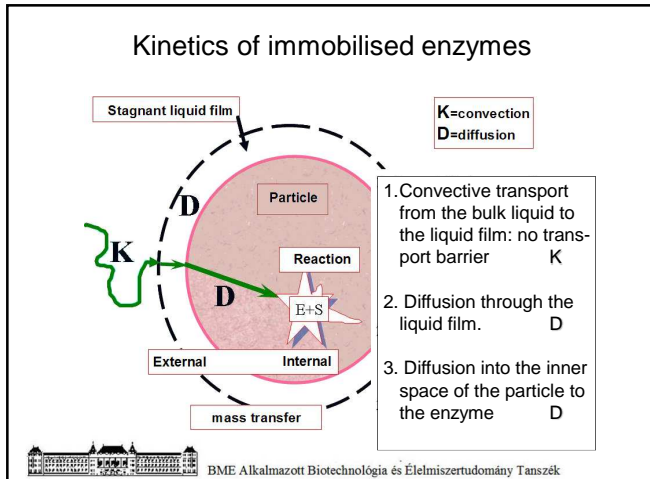
---

---

---

---

---




---

---

---

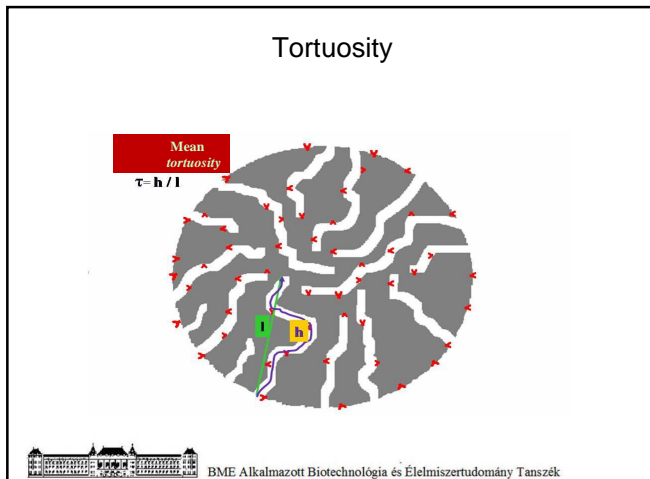
---

---

---

---

---




---

---

---

---

---

---

---

---

### 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

BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

### 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



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

22

---

---

---

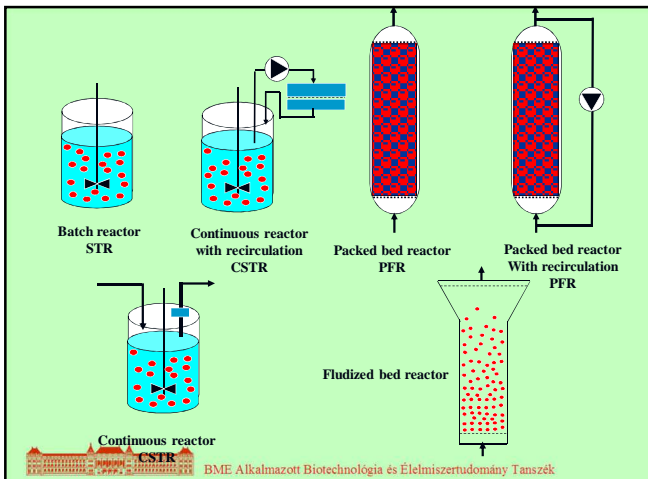
---

---

---

---

---



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

### Industrial application of immobilised enzymes

Aminoacylase	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
$\beta$ -galactosidase	hydrolysis of lactose to glucose+galactose
Lipase	hydrolysis and transesterification of lipids
Thermolysin	Preparation of aspartame



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

24

---

---

---

---

---

---

---

---



### 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:

Ag anode:  $4Ag + 4Cl \rightarrow 4AgCl + 4e^-$   
 Pt cathode:  $O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$

BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

---

---

### BIOSENSOR

BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

---

---

### 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

BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

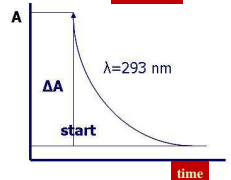
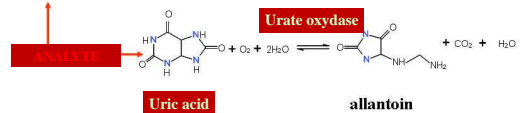
---

---

---

### End-point measurement of substrate

The whole amount of substrate is converted – change is measured




---

---

---

---

---

---

---

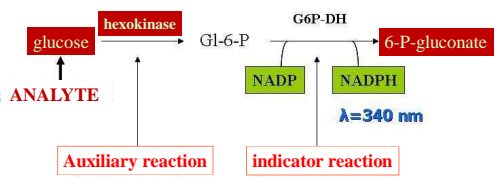
---

---

---

### Indicator reaction

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



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

---

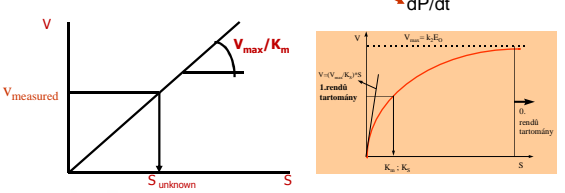
---

### Kinetic measurement of S

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

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

$-dS/dt$        $dP/dt$



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék

---

---

---

---

---

---

---

---

---

---