Heterogeneous phase enzyme reactions

Advantages/disadvatages:

Advantages:

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

Economic disadvatages:

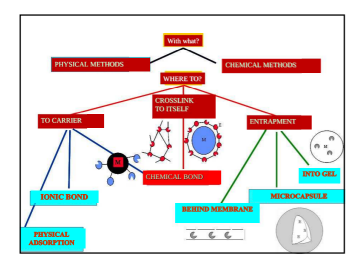
- > Enzymes are expensive, 1-10- \$/mg
- > can be used only once, after reaction they are to be discarded...

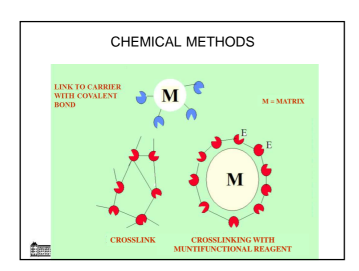
Technological disadvatage:

> Proteins contaminate products



BME Alkalmazott Biotechnológia és Élelmiszertudomány Tanszék





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

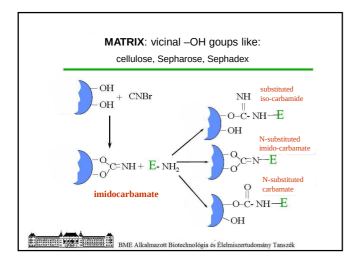
Building of covalent bond: free α -, β - or γ -COOH, α -, β -NH $_2$ groups phenyl-, OH-, SH- imidazole-groups

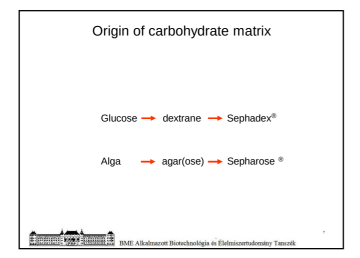
STEPS:

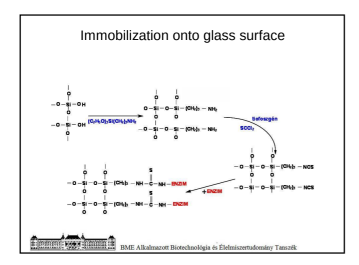
- 1. Activation of carrier (arm and reactive X-group),
- 2. Creating covalent bond between enzyme and activated carrier

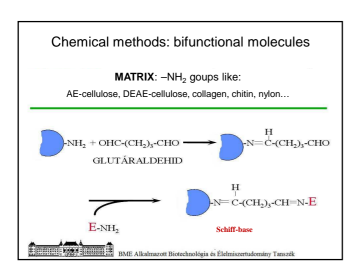
Protection of the active sites: S or analog

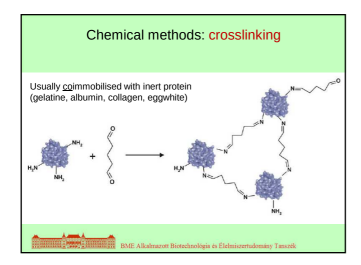


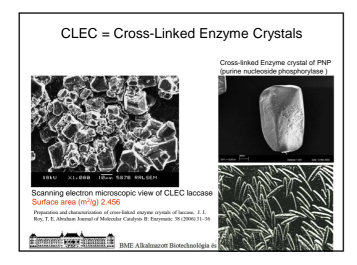


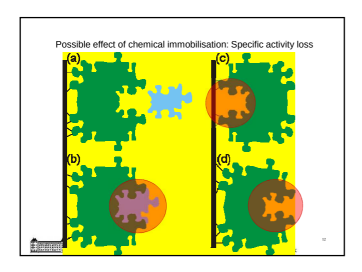




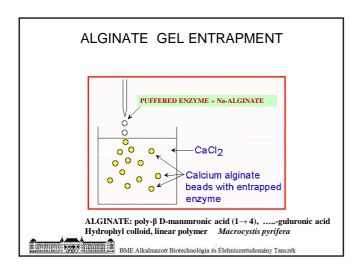


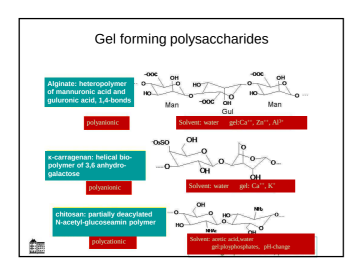


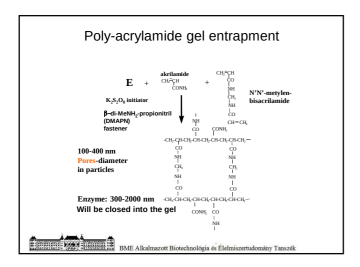


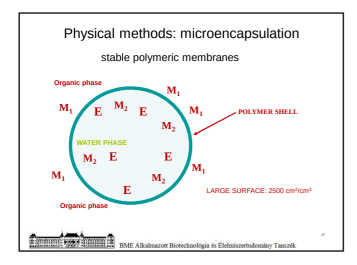


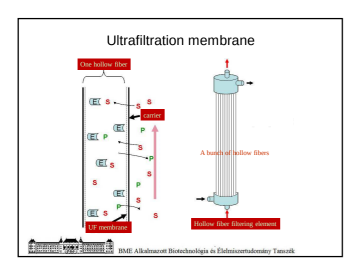
PHYSICAL METHODS 1. Adsorption e.g. on *ionexchanger resins* – nonspecific, easily desorps (pH) 2. Gel entrapment 3. Microencapsulation 4. Closing behind membrane

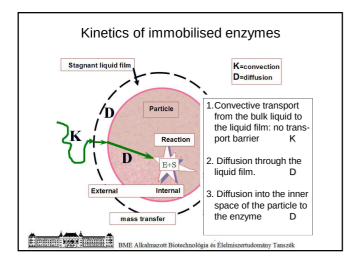


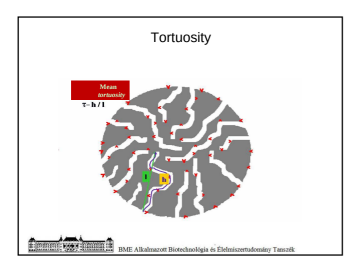


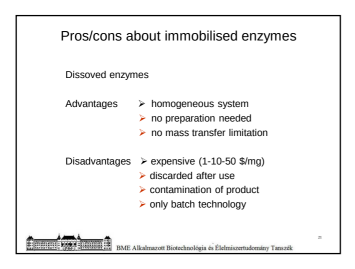


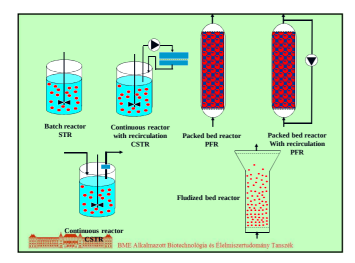


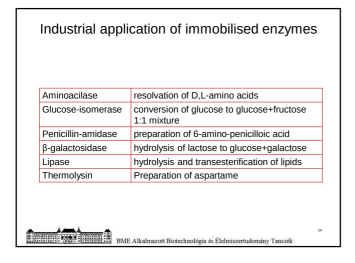


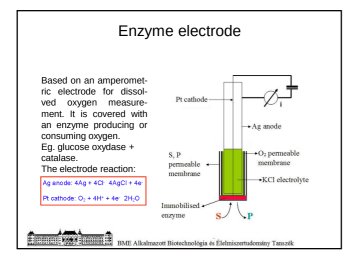


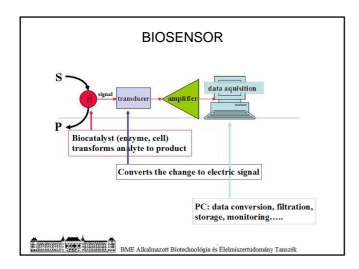












Analytical enzyme applications In these cases not the activity of enzyme is measured but the concentration of an analyt molecule. 1. Determination of S 2. Determination of I 3. Marker reactions (eg. in immunoassays) Enzyme Linked Immunosorbent Assay (ELISA) diagnostical, research purposes

