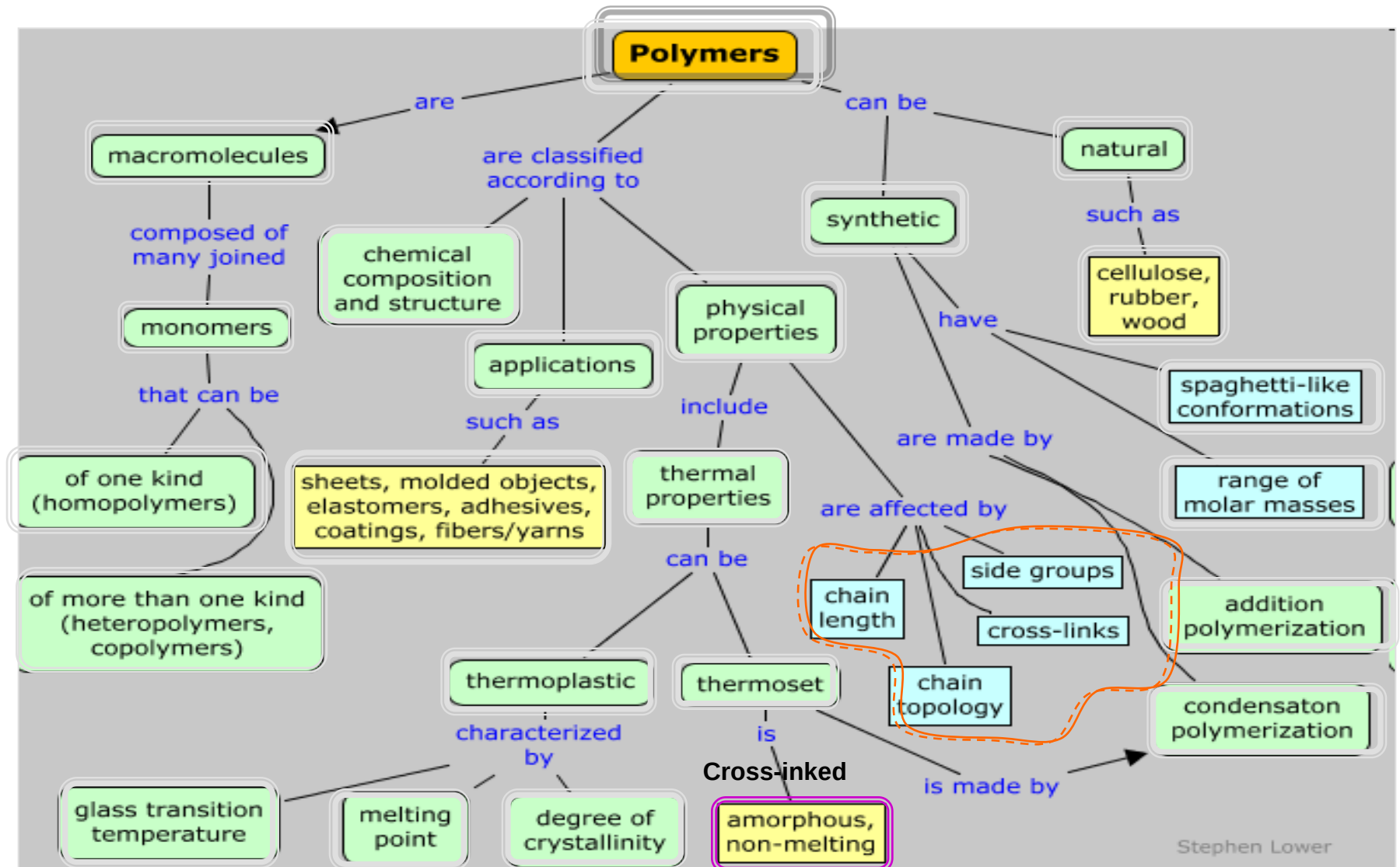


POLYMER SCIENCE AND TECHNOLOGY

Lecturer: Dr. Ljerka Kratofil Krehula, Associate Prof.
krehula@fkit.hr

POLYMERS

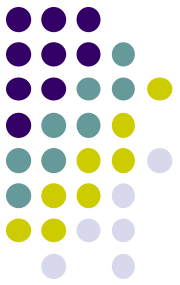
-classified by their chemical structure, properties and applications.



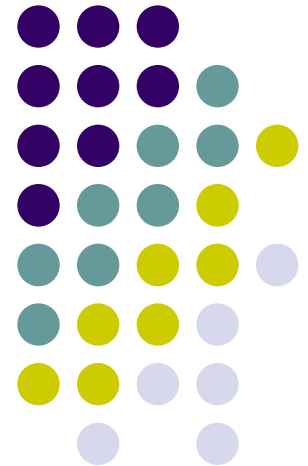
Classification of polymerization reactions:

according to:

1. mechanism of chain growth
2. media of polymerization



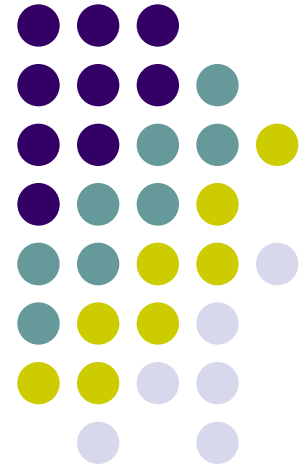
1. Polymerizations by the mechanism of chain growth



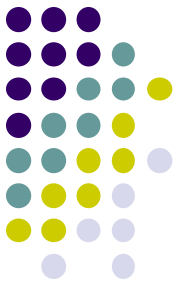
Mechanism of chain growth

RADICAL POLYMERIZATION

STEP BY STEP POLYMERIZATION



Mechanism of chain growth (*type of reaction*)



1. free-radical polymerization

addition polymerization, chain-growth of polymer

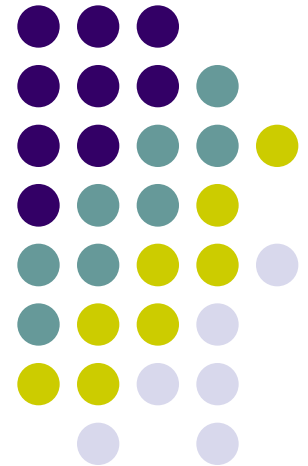
- I. initiation
- II. propagation
- III. termination

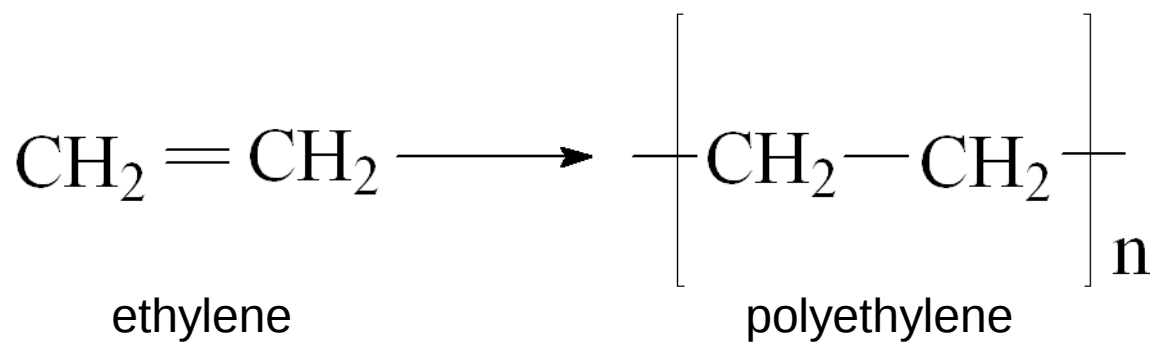
2. step by step polymerization

condensation polymerization, stepwise growth of polymer

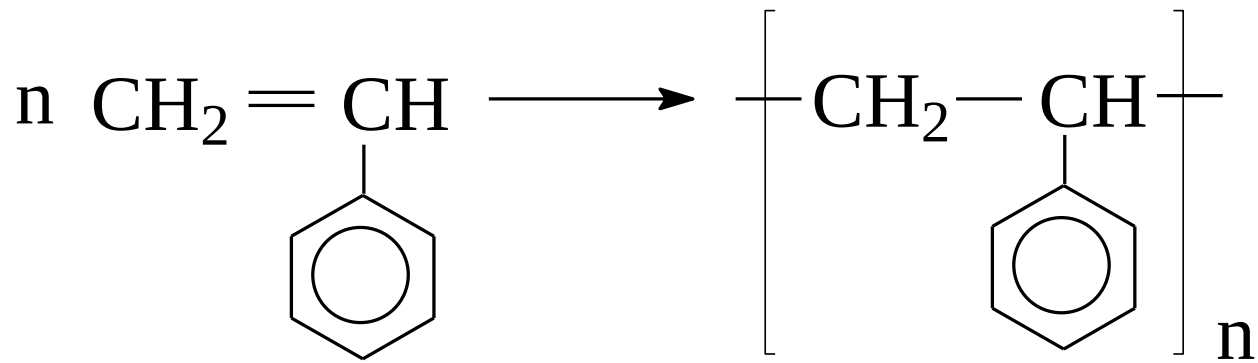
monomer + monomer $\rightarrow$ dimer
dimer + monomer $\rightarrow$ trimer
dimer + dimer $\rightarrow$ tetramer
trimer + monomer $\rightarrow$ tetramer etc.

1. FREE-RADICAL POLYMERIZATION



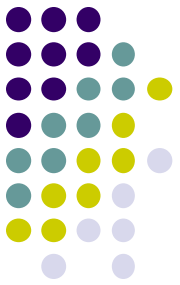


Low density polyethylene is produced by free radical polymerization.



styrene
(vinyl-benzene)

polystyrene



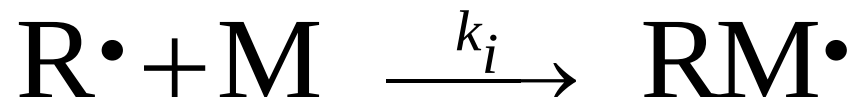
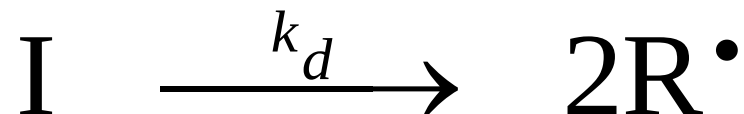
Free-radical polymerization



Three phases:

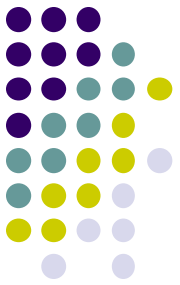
- I. initiation
- II. propagation
- III. termination

1. Initiation – initiator decomposition



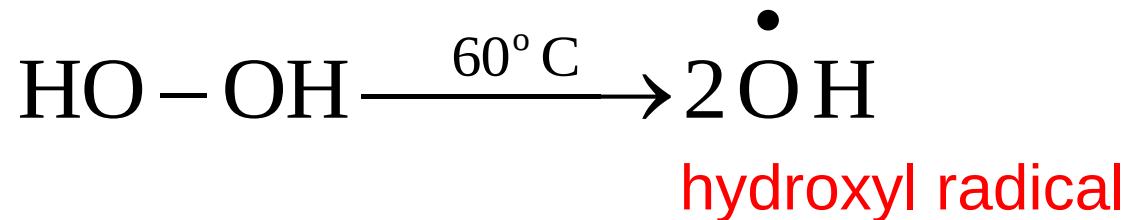
Primary active species

Initiators: **specific decomposition temperature!**

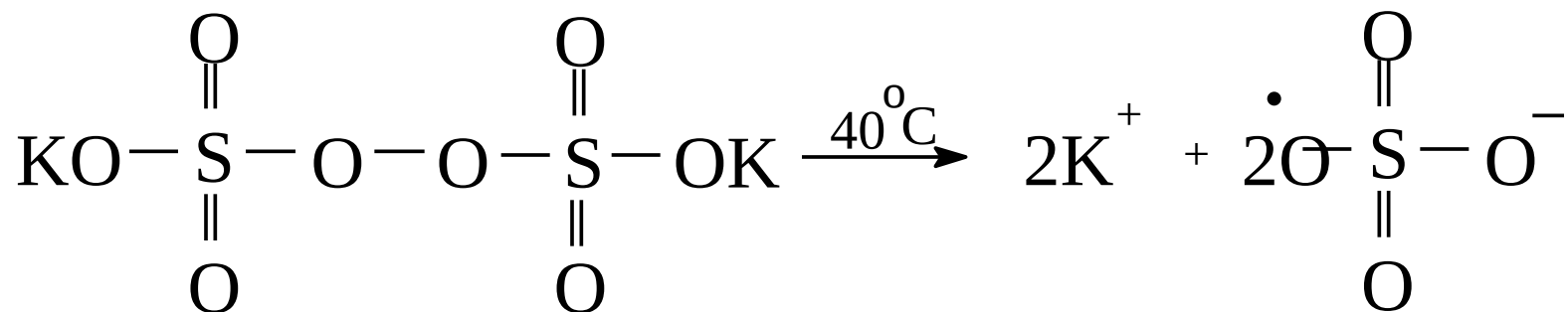


a) inorganic initiators

hydrogen peroxide, H_2O_2 (O-O bond)



potassium persulfate, $\text{K}_2\text{S}_2\text{O}_8$ (O-O bond)

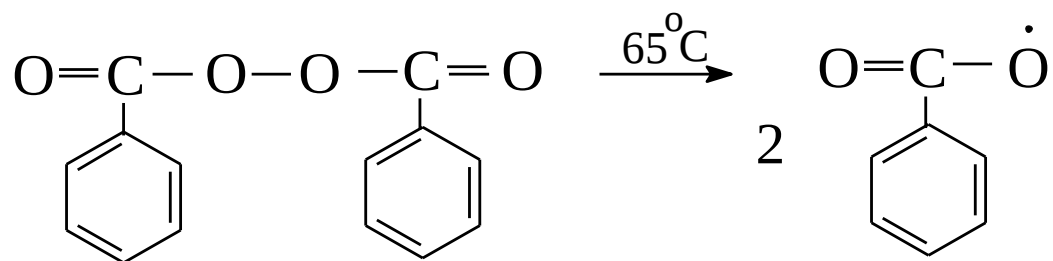


sulfate anion radical



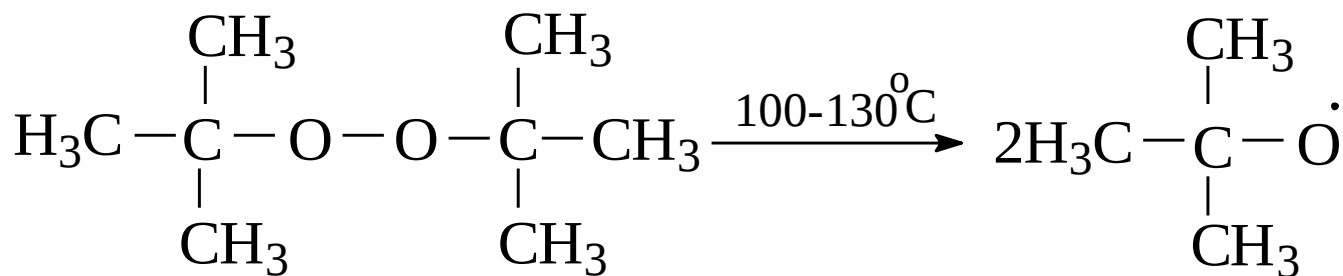
b) Organic peroxides

benzoyl peroxide, DBP (O-O bond)



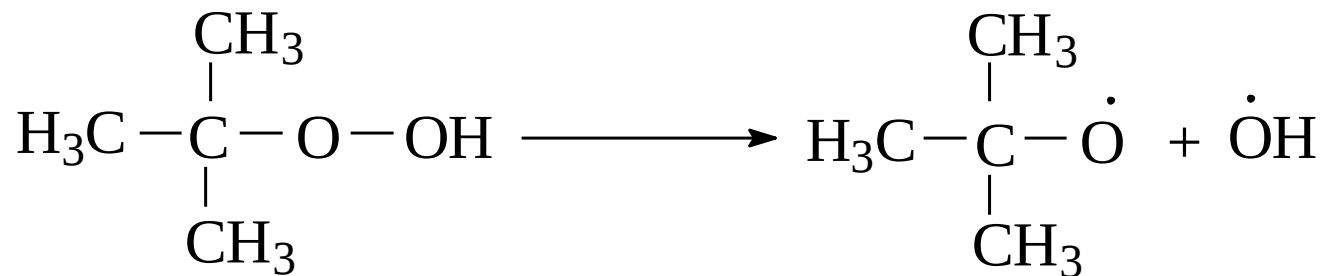
benzoyloxy radical

Di-tert-butyl peroxide (O-O bond)



t-butyloxy radical

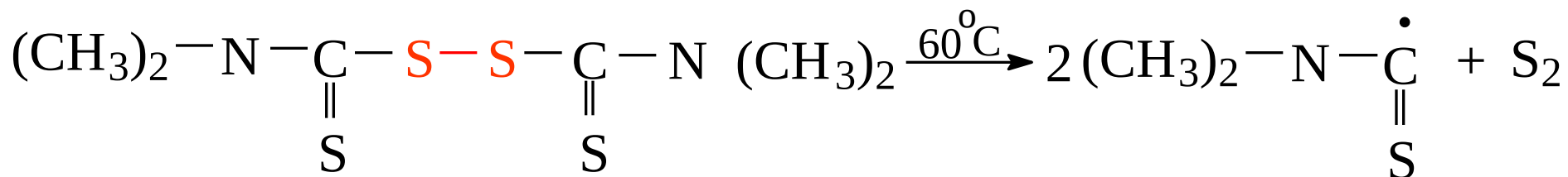
t-butylhydroperoxide (O-O bond)



c) initiators with S-S bond

Tetramethylthiuram disulfide , TMTD

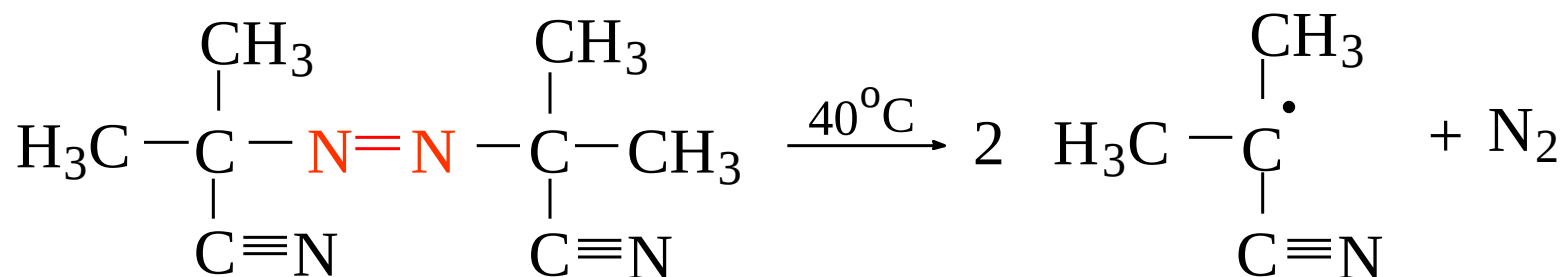
- vulcanization



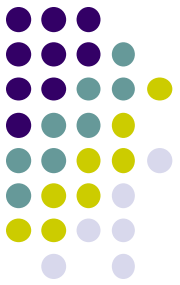


d) Initiators with N-N bond

α,α' -azobisisobutyronitrile, AIBN



2-cyano-2-propyl radical

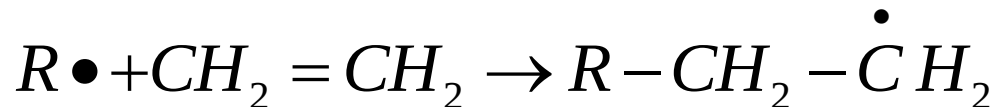
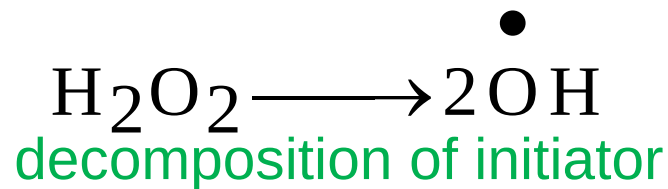
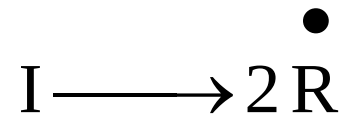


The example of

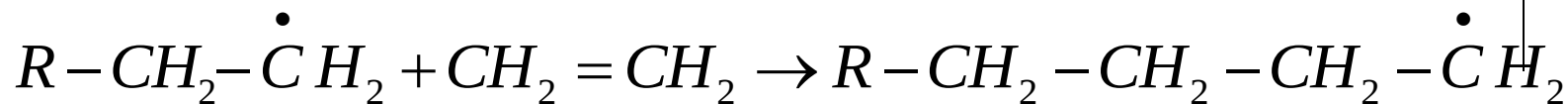
free-radical polymerization:

ethylene polymerization

1. Initiation

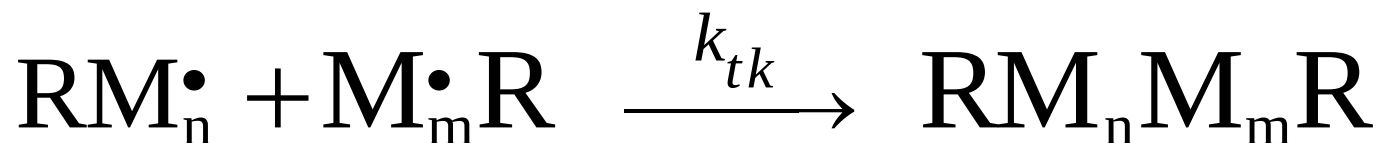


2. Propagation

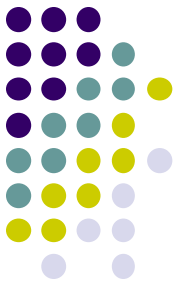
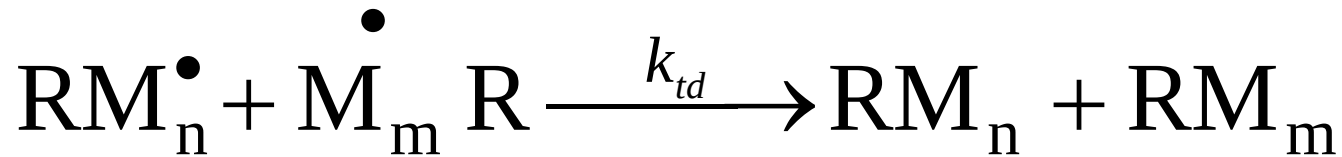


3. Termination – the end of polymerization

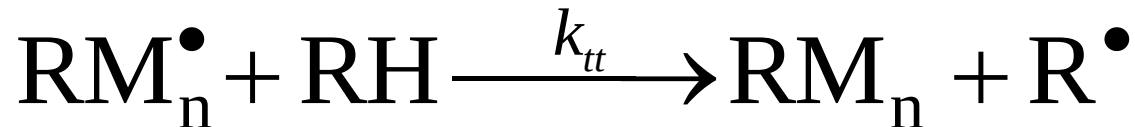
a) combination



b) disproportionation

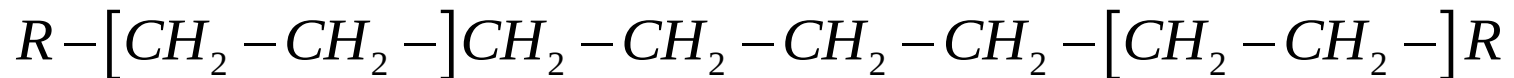
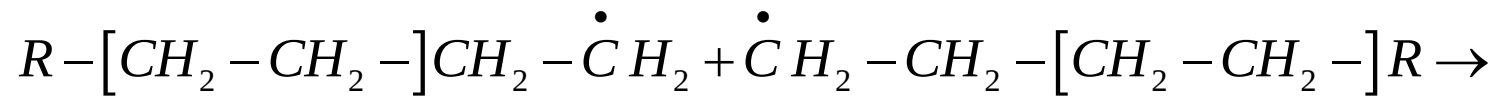
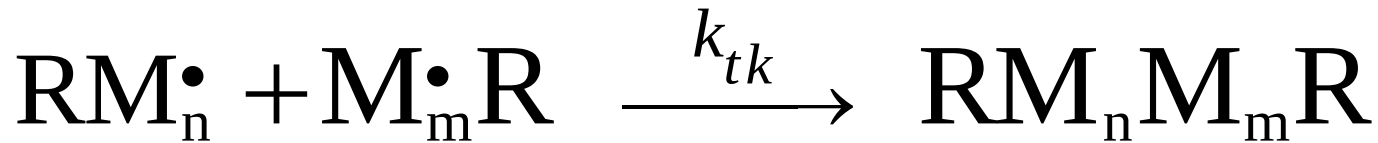


c) chain transfer

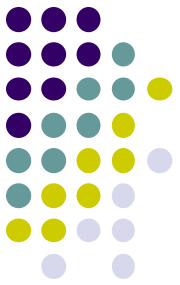


I- Initiator, $\text{R}^\bullet$ - radical, M - monomer, k_d – constant of dissociation k_i - constant of initiation, k_p - constant of propagation, k_{tk} - constant of transfer with combination, k_{td} - constant of disproportionation, k_{tt} - constant of chain transfer

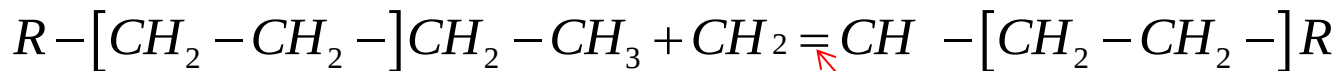
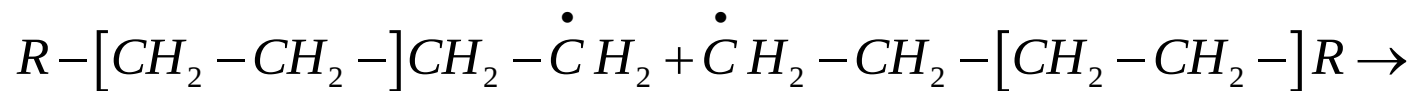
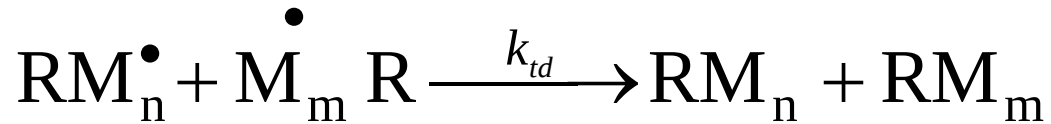
a) combination



Product: one polymer chain



b) disproportionation



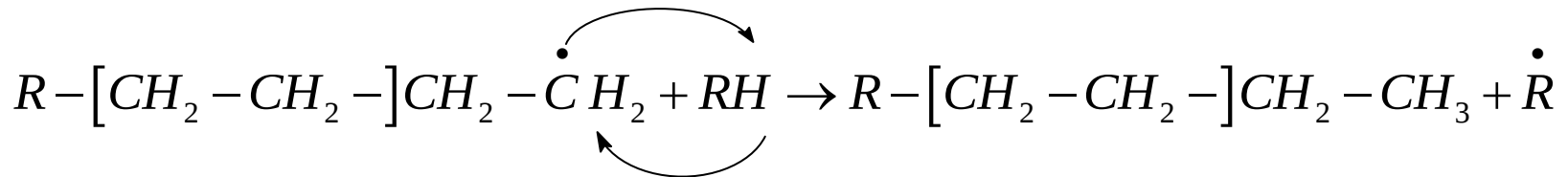
terminated polymer
chain

Double bond can be attacked
by radical
- new polymerization starts

Product: two polymer chains



c) **chain transfer** – transfer of polymerization reaction to „something else” in a reaction mixture – *in general, it is not favorable type of termination*



RH may be initiator
solvent
monomer
polymer

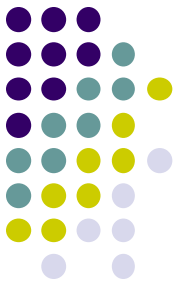
Chain transfer with initiator



Initiator forms the first radicals, then attacks monomer and forms further radicals.

Chain transfer with initiator rarely happens - *the added quantity of initiator is very small (0,1%).*

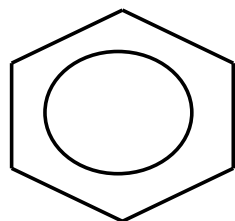
At the end of polymerization there is no initiator.



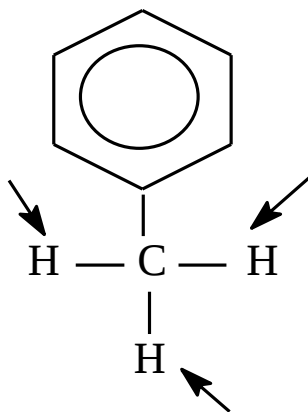
Chain transfer with solvent

Solvent is always present except at polymerization in bulk.

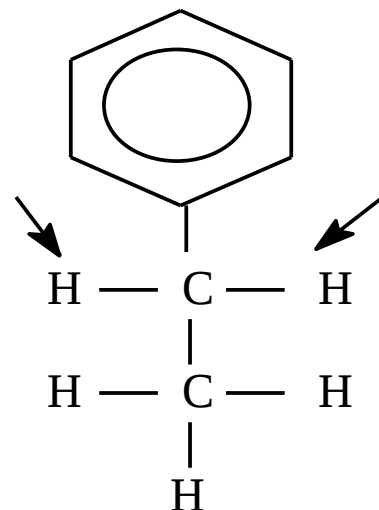
Aldehydes and ketones are very good chain transfer solvents because of easy discharge of H-atom.



benzene



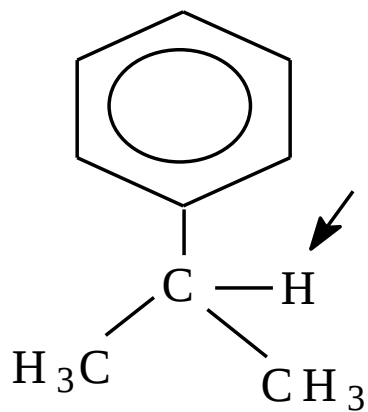
toluene



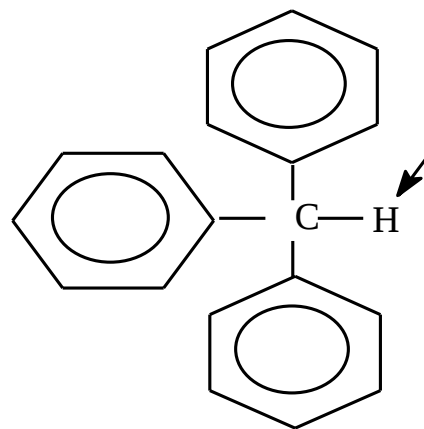
ethylbenzene

better chain transfer





cumene



tri phenylmethane

better chain transfer

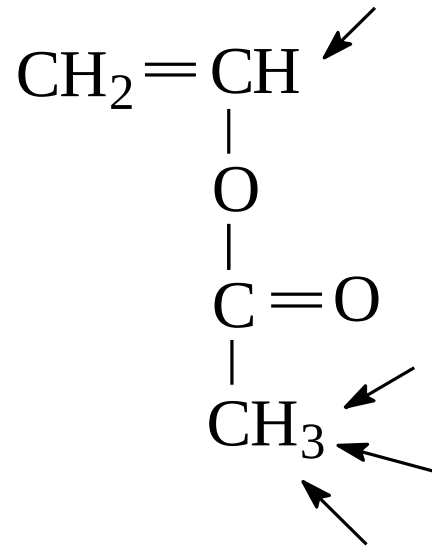




Chain transfer with monomer

Chain transfer with monomer rarely happens, because at the end of polymerization there is no monomer any more.

Very reactive monomer is vinyl-acetate

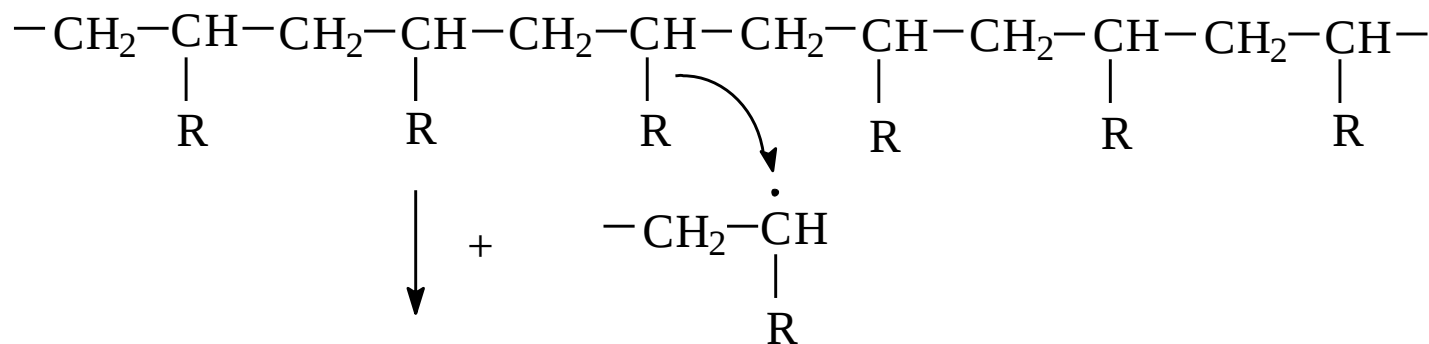


Chain transfer with polymer

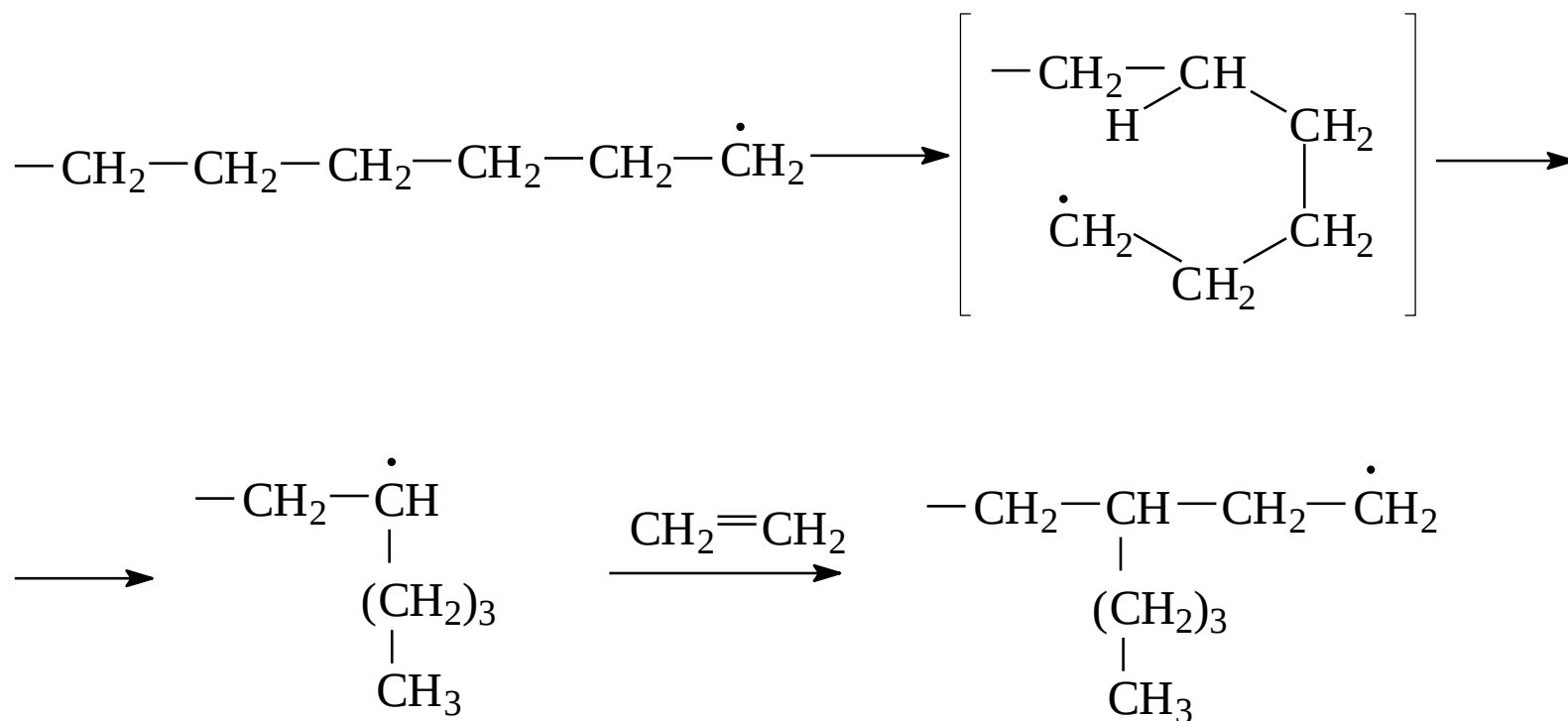


Product: branched or crosslinked polymer

It is the most common chain transfer, because at the end of polymerization there are only solvent and polymer. During a free radical polymerization, there is always a possibility for polymer branching.



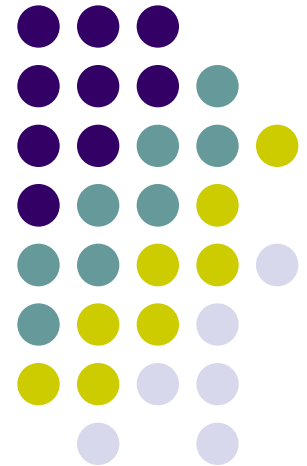
- very interesting polyethylene chain transfer:
chain reacts with its own chain – intramolecular bonding

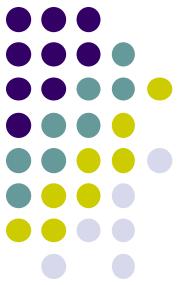


This reaction is called «**backbiting**» - it means
“to bite its own tail”.

STEP BY STEP POLYMERIZATION *condenzation* *polymerization*

- the product of the reaction are polymers and „little” molecules (water, ammonia, CO_2 , HCl , N_2 , methanol)
- polymerization in which the polymer's molecular weight increases in a slow, stepwise manner as reaction time increases.
- the polymerizations are reversible





Step by step polymerization

- *condensation polymerization*
- stepwise growth of polymer

monomer + monomer $\rightarrow$ dimer

dimer + monomer $\rightarrow$ trimer

dimer + dimer $\rightarrow$ tetramer

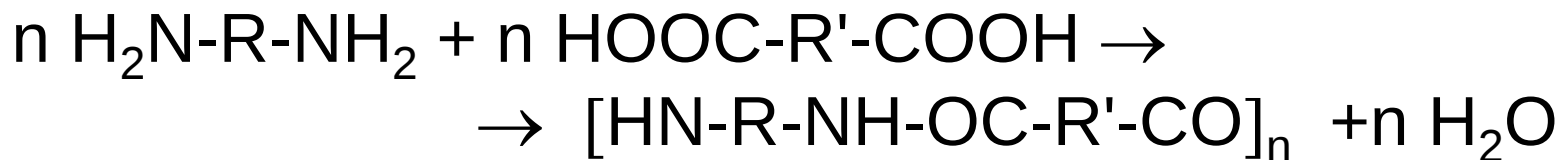
trimer + monomer $\rightarrow$ tetramer etc.

The product is **polycondenzate**

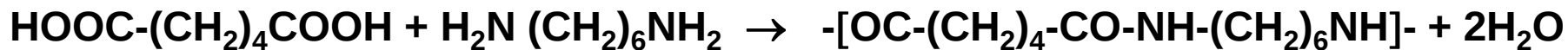
Step by step polymerization: 2 types



1. **Two different polyfunctional monomers**
- every monomer has **only one type of functional group**:



or generally:



adipic acid

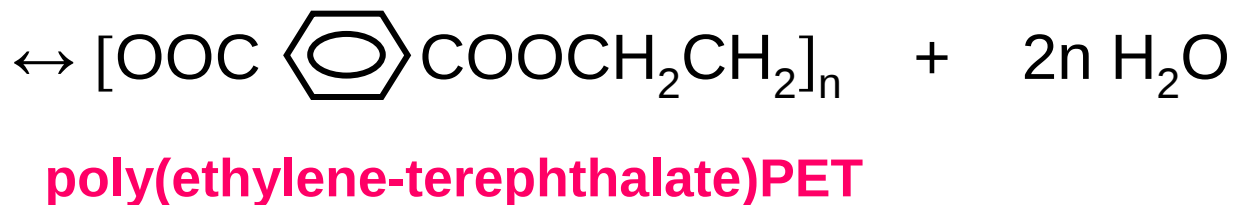
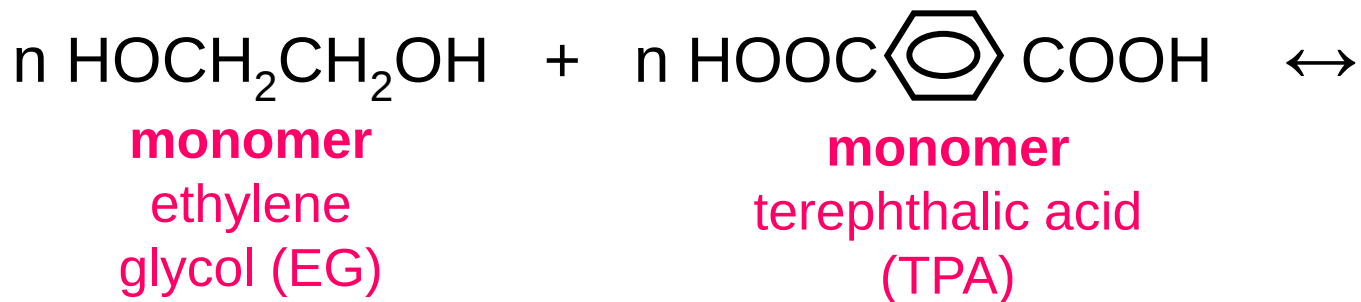
hexamethylene
diamine

nylon 6,6 - polyamide

Production of PET

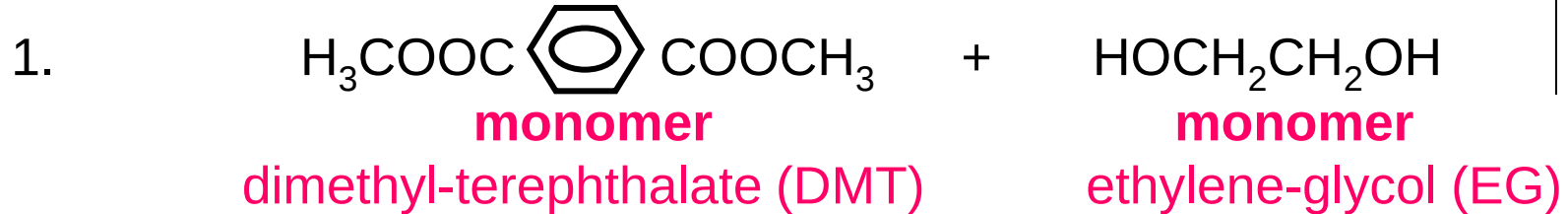
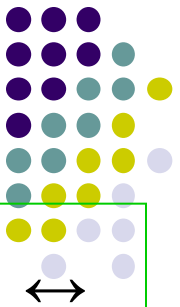


1. Direct esterification



(The same reaction is in the previous slide.)

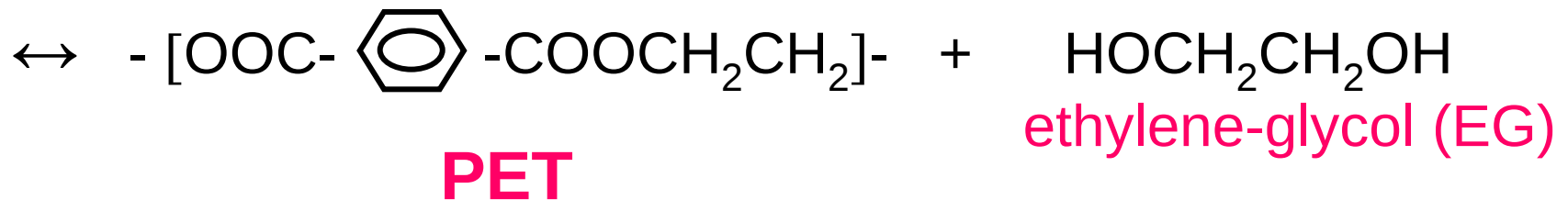
2. Ester exchange





monomer

1,4- 1,4-bis-hydroxyethyleneterephthalate (BHET)





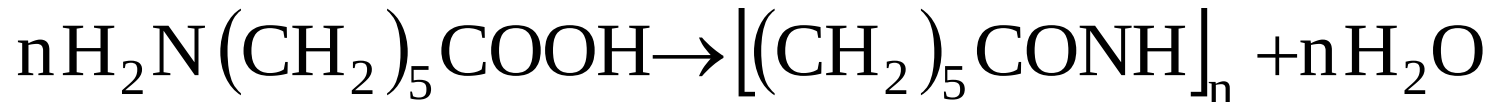
2. ***One monomer with two types of functional groups:***



or generally:

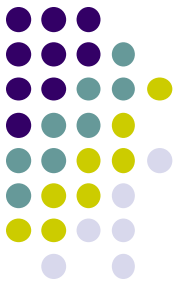


Example: polyamide



w-aminohexaacid

nylon 6, perlon - polyamide



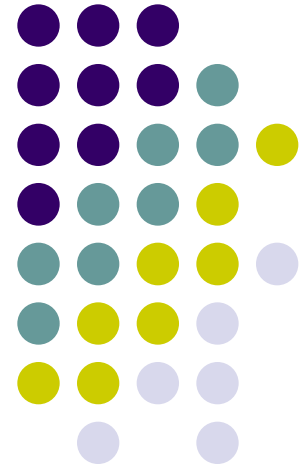
Reactions of polymerization are classified according to:

- mechanism of chain growth
- media of polymerization

Media of polymerization:

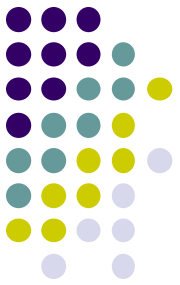
- homogeneous
- heterogeneous

2. Polymerizations by the media of polymerization



Polymerization processes –

in **gas**, **liquid** and **solid** phase
(depending on the medium)

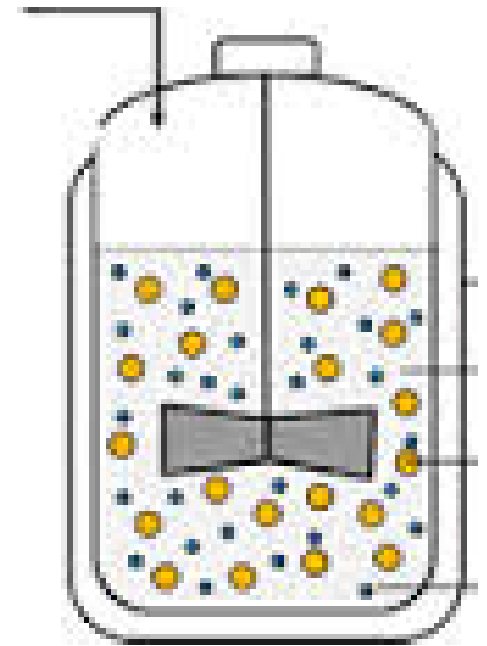


Homogeneous polymerizations:

1. Bulk polymerization
2. Solvent polymerization

Heterogeneous polymerizations :

1. Heterogeneous bulk polymerization
2. Heterogeneous solvent polymerization
3. Suspension polymerization
4. Emulsion polymerization
5. Polymerization in the gas phase
6. Interfacial polycondensations



Homogeneous polymerizations



1. Bulk polymerization

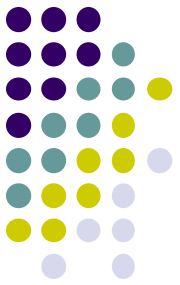
- **monomer + initiator**
- there is **no solvent**
- only **one phase**:

monomer and polymer are compatible:

unreacted monomer acts as a solvent to polymer

Initiation - in monomeric phase usually with organic peroxides,

- initiators must be **completely soluble in monomer**



Limitations of bulk polymerization:

- **inefficient heat transfer** -
high concentration of reactants -
- the control of reaction is not efficient
- high viscosity, mixing is difficult
- polymerization is not suitable for industrial processes

For industrial production

- it is necessary to control the chemical and thermal conditions of polymerization.



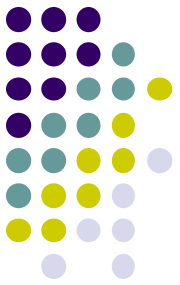
Example of self-initiated bulk polymerization:

polymerization of vinyl polymers –

long exposure to the sun (heat and UV light).

Obtained polymer – properties are not uniform
due to the uncontrolled conditions
of the polymerization reaction.

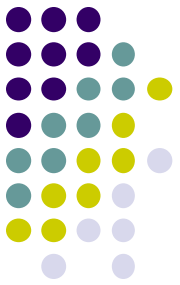
2. Solvent polymerization



- homogeneous polymerization
- **monomer + initiator + solvent**
- only **one phase**:
 - Monomer is soluble in solvent.*
 - Polymer is also soluble in solvent.*
- **solvent acts as diluent** in reaction mixture, reduces concentration of monomer

Solvent reduces concentration of reactants

- a decrease of reaction rate
- decrease of molecular weight



Solvent affects:

- molecular weight
- nature of the terminal groups of polymers

The most common solvents:
benzene, methanol, ethyl acetate...

Initiators: *substances for initiation of a polymerization* →

organic peroxides, often used:

dibenzoyl peroxide, DBP

Advantage of solvent polymerization:



- very efficient heat transfer

the heat developed during the reaction is distributed all over the system due to the presence of solvent and reduced concentration of the reactants



Disadvantages of solvent polymerization:

- low reaction rate (*low concentration of monomer*)
- obtaining low molecular weight polymer
- presence of solvent – solvent must be removed (evaporated) after polymerization