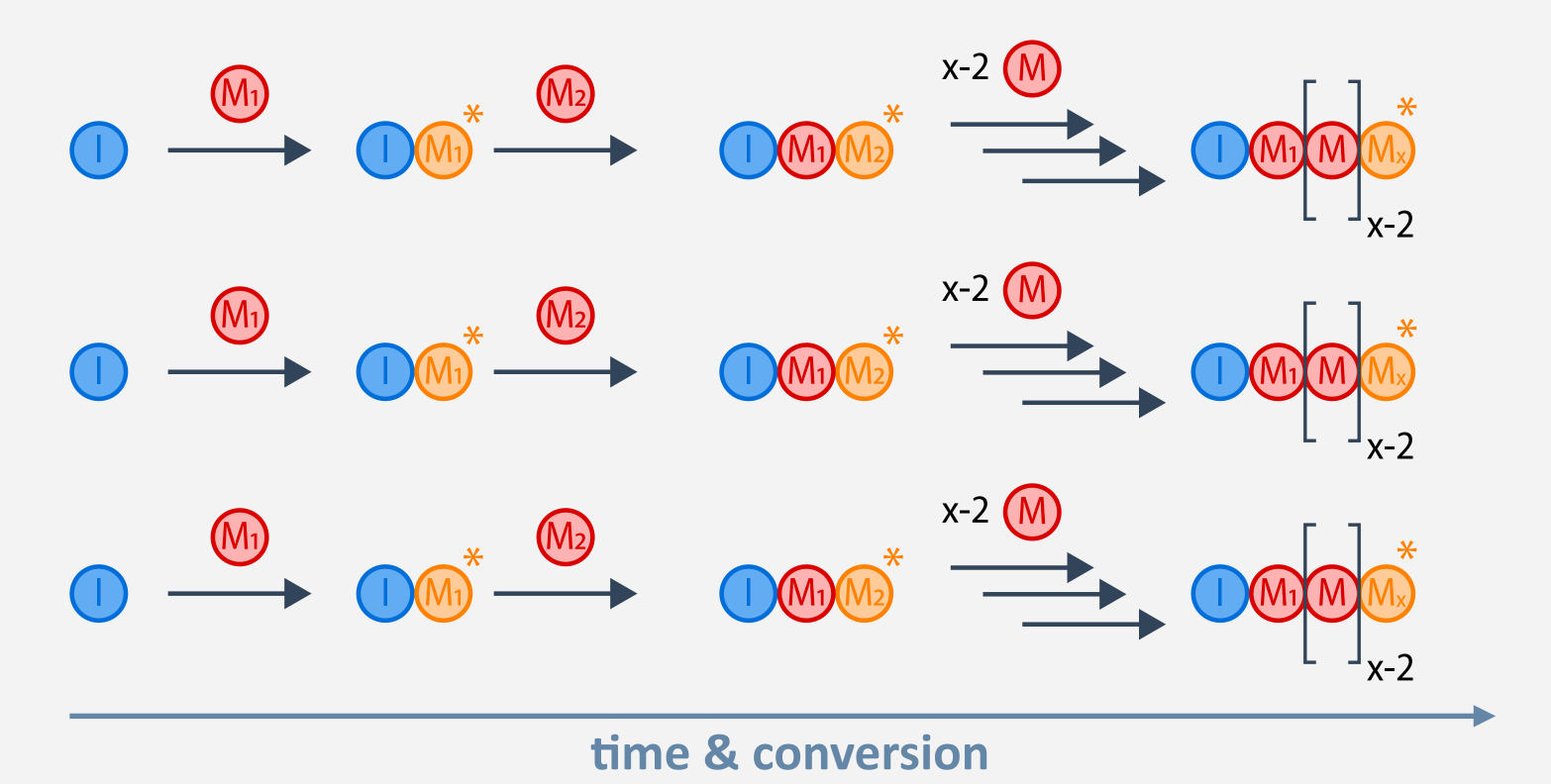
5.4 Living and Controlled Polymerizations

The Living Nature

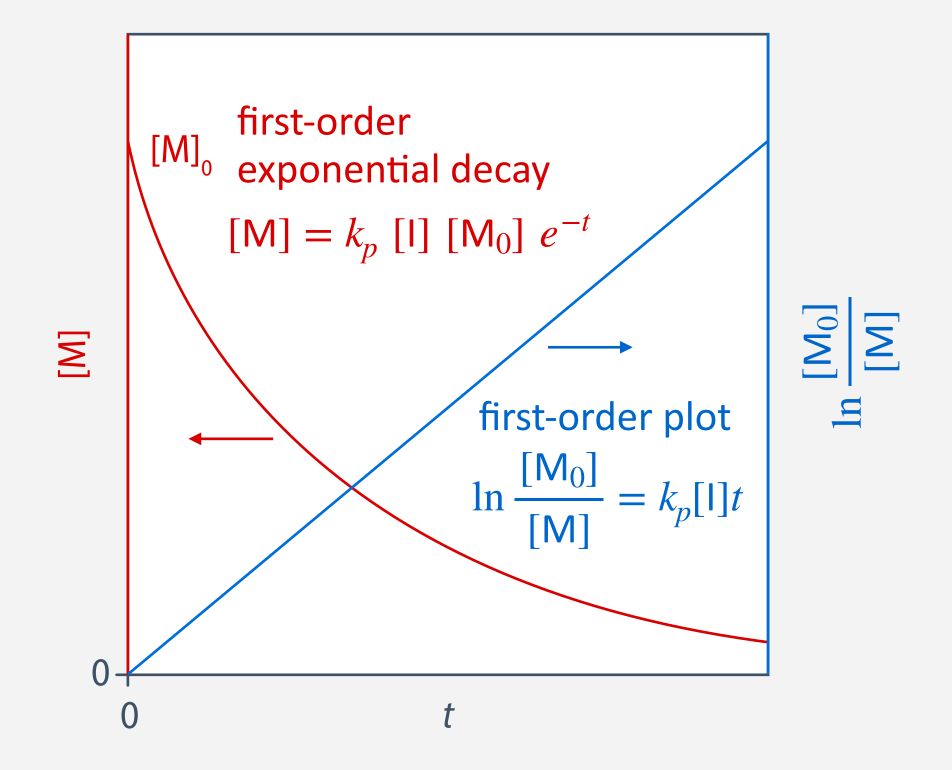
• absence of termination (no mechanism & strong electronic repulsion between active chain ends!)



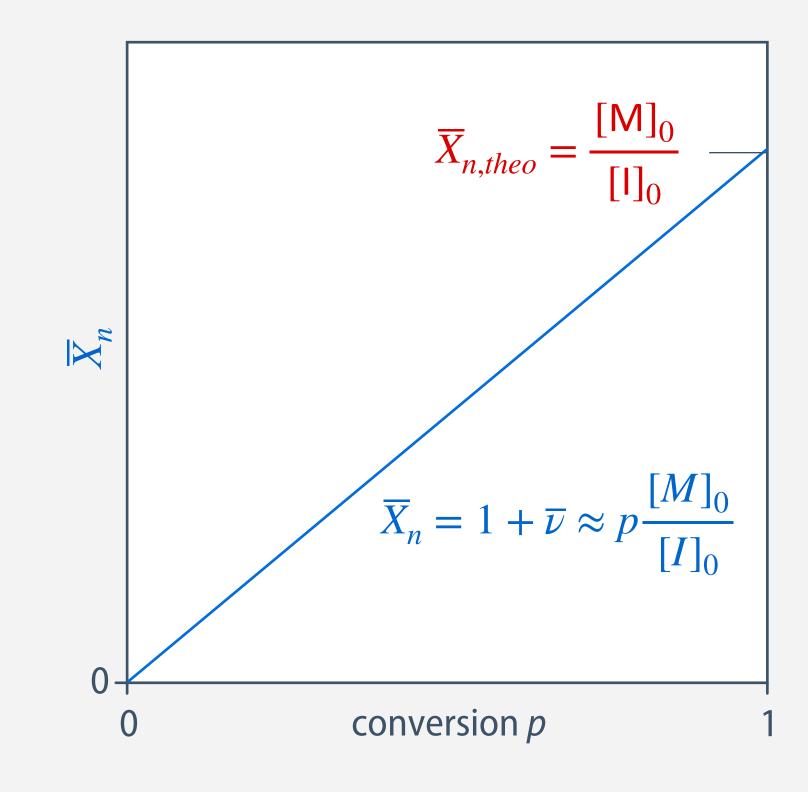
- polymers are all initiated fast and at approximately the same time; origin of (small) dispersity
- chain ends remain active after full monomer consumption (absence of impurities!)
- the polymerization can be continued with an additional feedstock of monomers (same or different)

Criteria for the Experimental Verification of Living Polymerizations

conversion over time



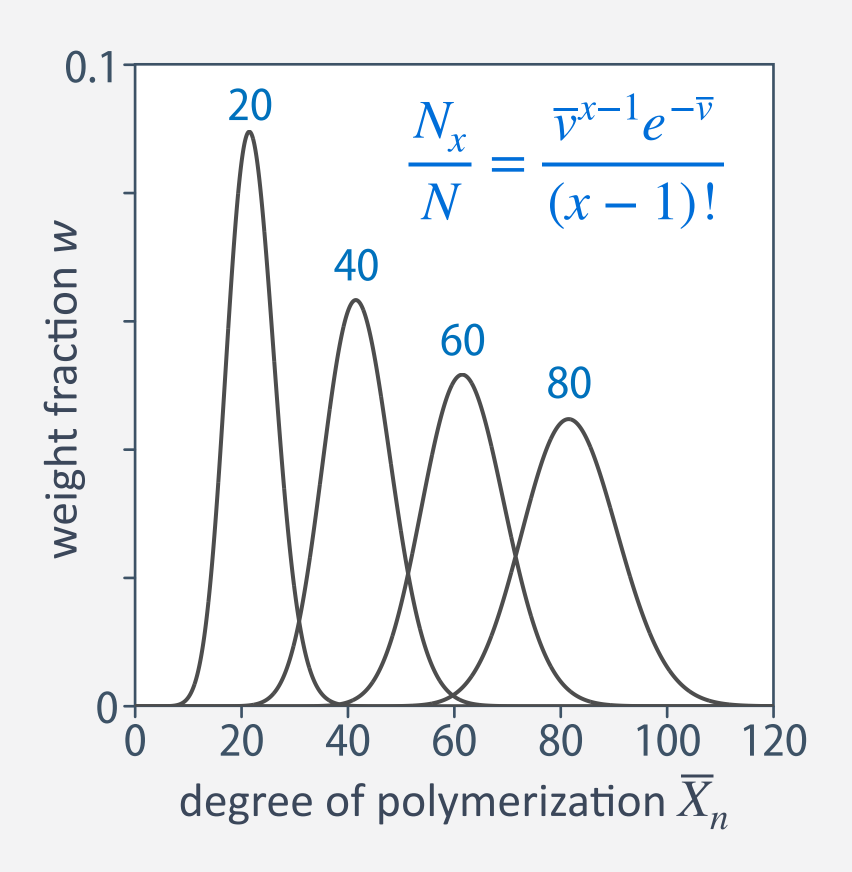
molar mass over conversion

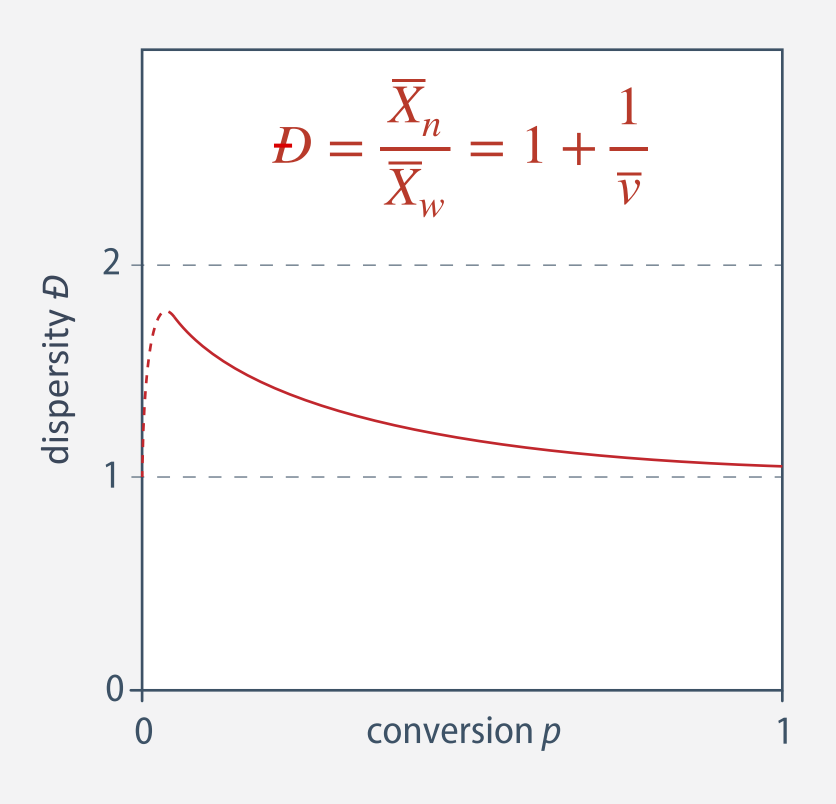


- conversion over time plot: polymerization reaction is first order in monomer concentration
- ullet molar mass over conversion plot: number-average molar mass $\overline{X}_n=1+\overline{
 u}$ linear with conversion
- molar mass control: final number-average molar mass depends on monomer/initiator ratio

The Poisson Distribution of the Molecular Weight

kinetic analysis leads to a Poisson distribution for the molecular weight distribution:



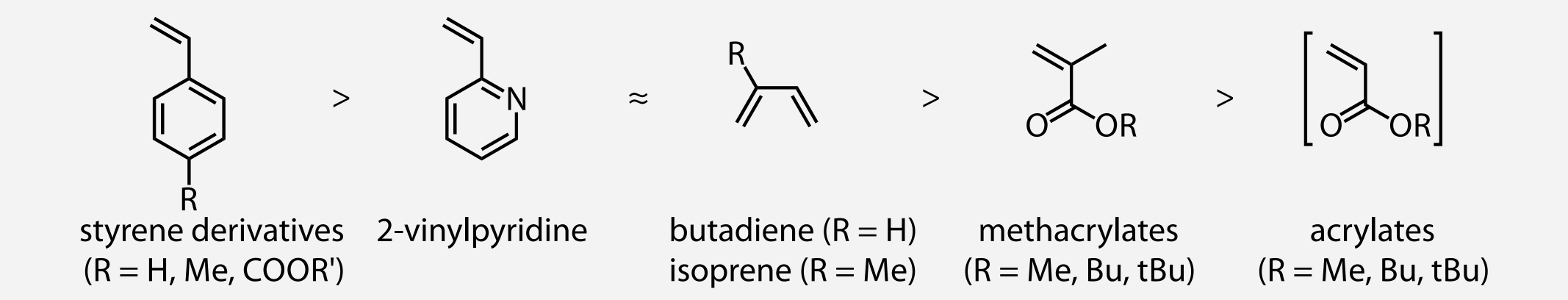


ullet number-average degree of polymerization $\overline{X}_n pprox p\, rac{[M]_0}{[I]_0}$ controlled by monomer/initiator ratio

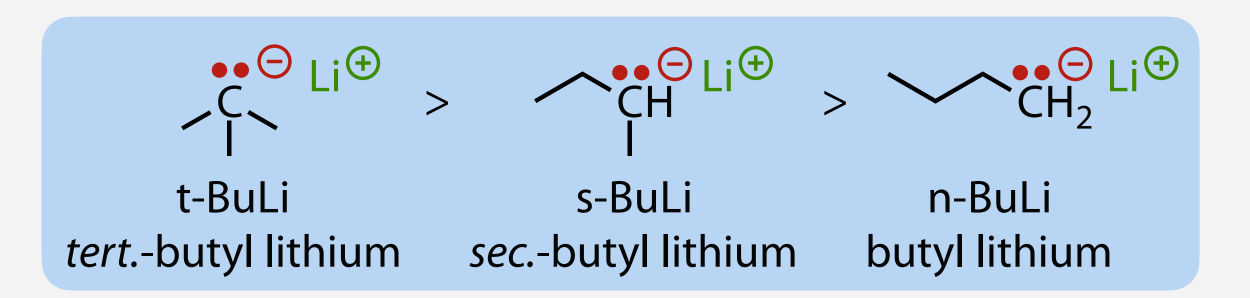
Living Anionic Polymerization

Vinyl Monomers and Initiators

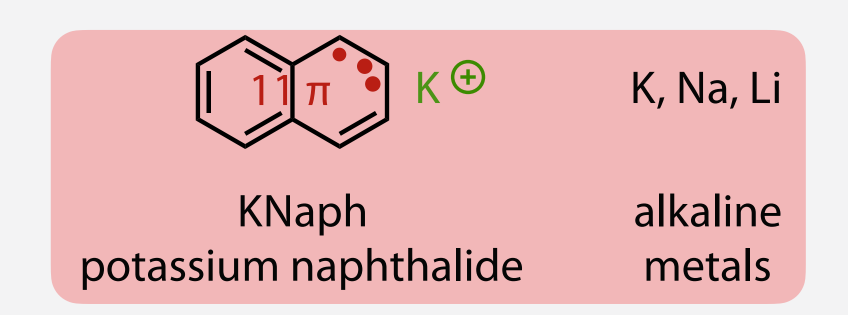
• monomers with possibility for anion delocalisation, ideally with electron-withdrawing side groups



• initiators are organoalkaline compounds or alkaline metals



strong nucleophiles for monodirectional chain growth



single-electron transfer agents for bidirectional growth

Living Anionic Polymerization of Vinyl Monomers

initiation (very fast)

- Mt⁺ R
- Mt⁺
 Nu CH

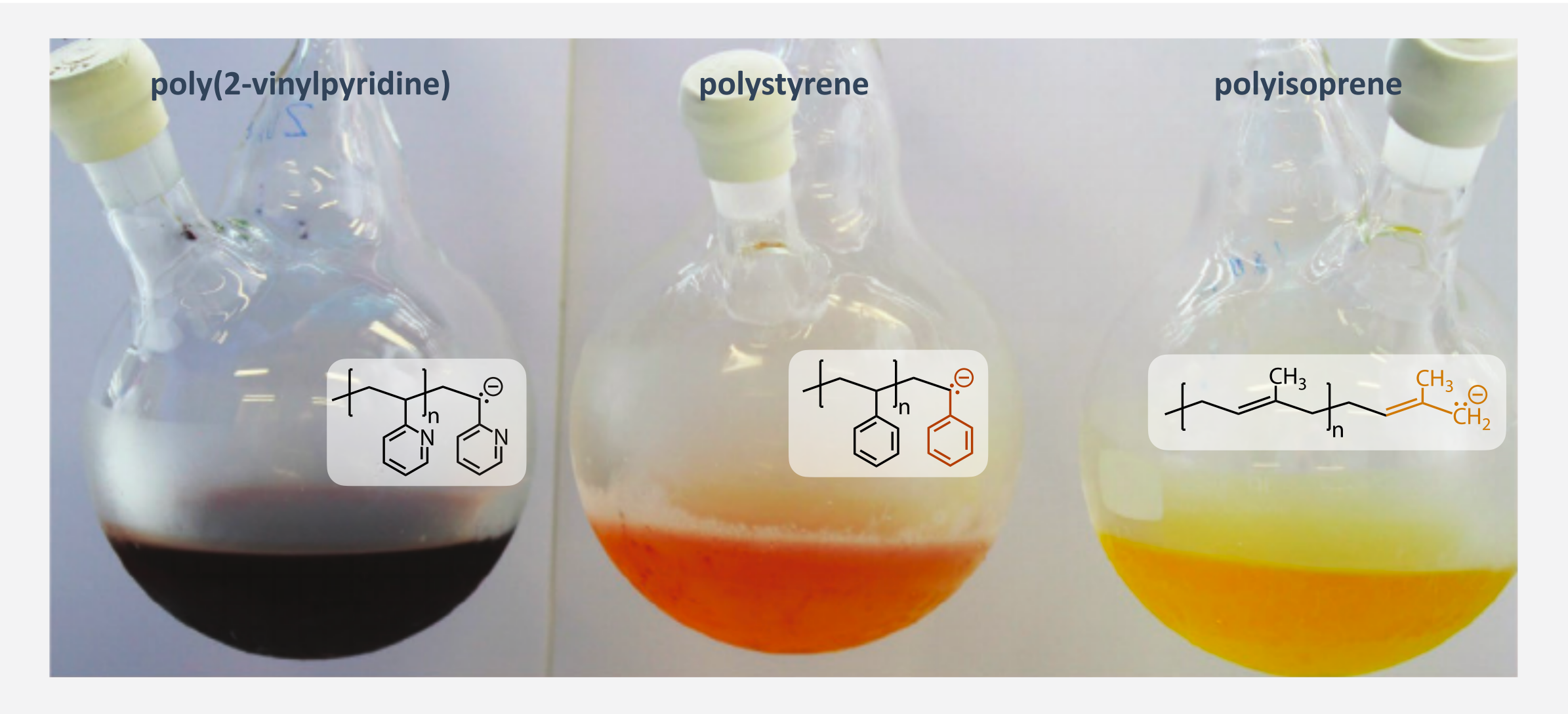
propagation (chain growth)

quenching

$$\begin{array}{c|c}
Mt^{\bigoplus} \\
Nu & \xrightarrow{E-X} \\
R & \xrightarrow{R} & \xrightarrow$$

- anionic polymerisations of vinyl monomers are initiated by strong nucleophiles
- termination reactions are absent, except for inadvertent electrophilic impurities (H₂O, CO₂)
- electrophiles serve as quenching reagents, deliberately end the reaction, introduce end groups

Living Anionic Polymerization of Vinyl Monomers



• appearance of color is an evidence of the presence and non-terminating character of living chains

Bidirectional Polymerization with Alkaline Metals and Naphthalides

alkaline metals reduce naphthalene by single-electron transfer (SET)

• SET from KNaph to a monomer results in dimerization, dianion initiator starts bidirectional chain growth

Quenching Reagents and Telechelics

• the living chain end is a strong nucleophile, end groups can be introduced by S_N and S_{AE} reactions

Thermoplastic Elastomers from BAB Triblock Copolymers

• SBCs (styrene-butadiene block copolymers) are relevant elastomer materials (like some TPUs)

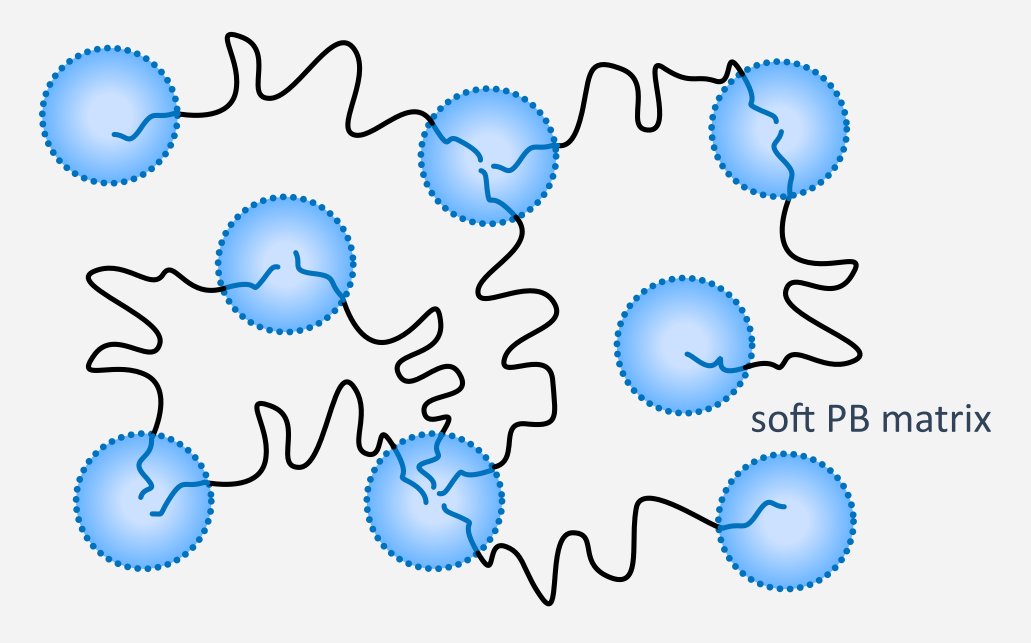
$$T_g \approx 100^{\circ}C$$

$$T_g \approx -60^{\circ}C$$

$$T_g \approx 100$$
°C

SBS rubber (Kraton™, BASF)

poly(styrene-block-butadiene-block-styrene)



glassy, hard, physical cross-links $\varnothing 10^{-8}$ – 10^{-7} m

- phase segregation (demixing) of different polymer segments in bulk material
- hard PS domains serve as physical cross-links: they melt above their T_g (reprocessable materials)

