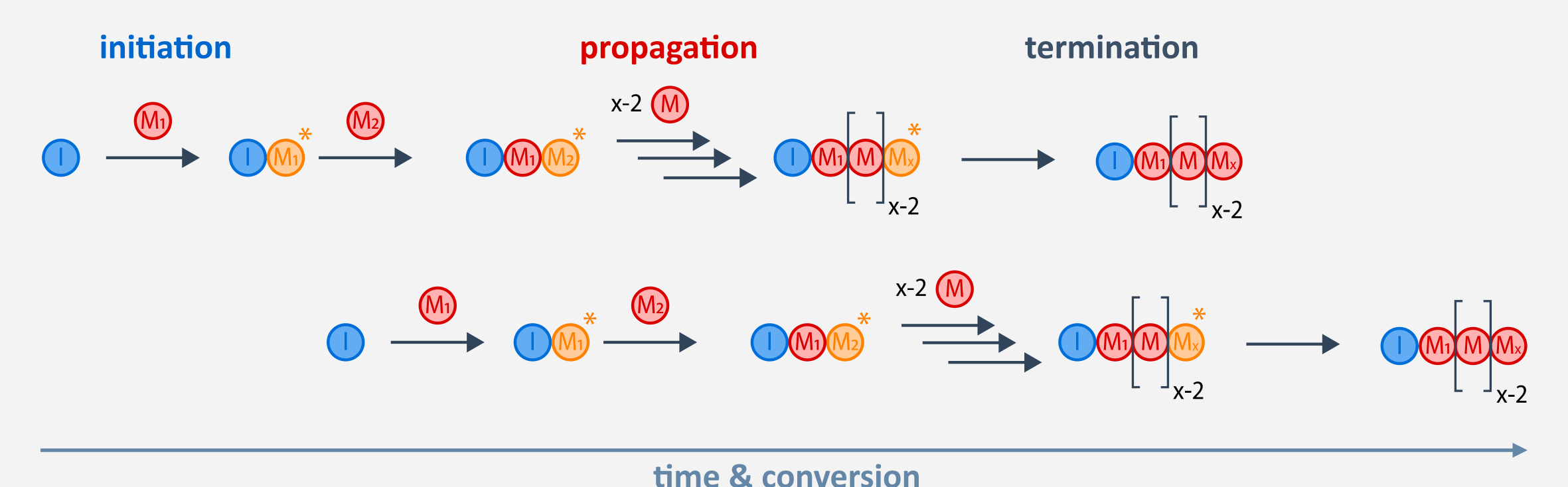
5.3 Chain-Growth Polymerization

Principal Steps of Chain-Growth Polymerizations

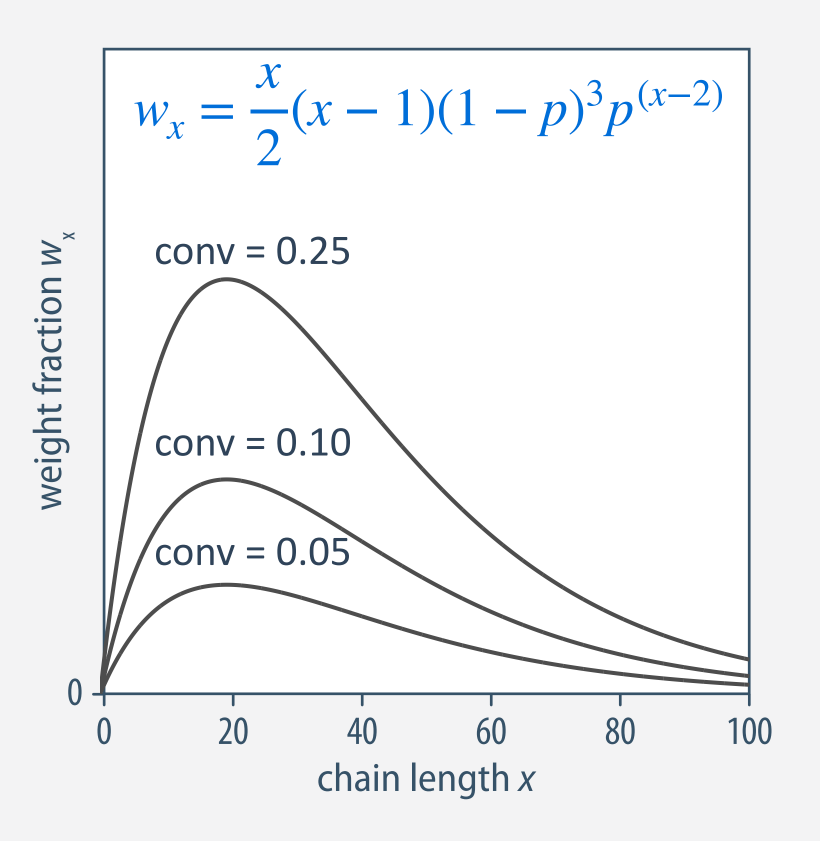
monomers add rapidly to the active center of a growing chain until that center is deactivated



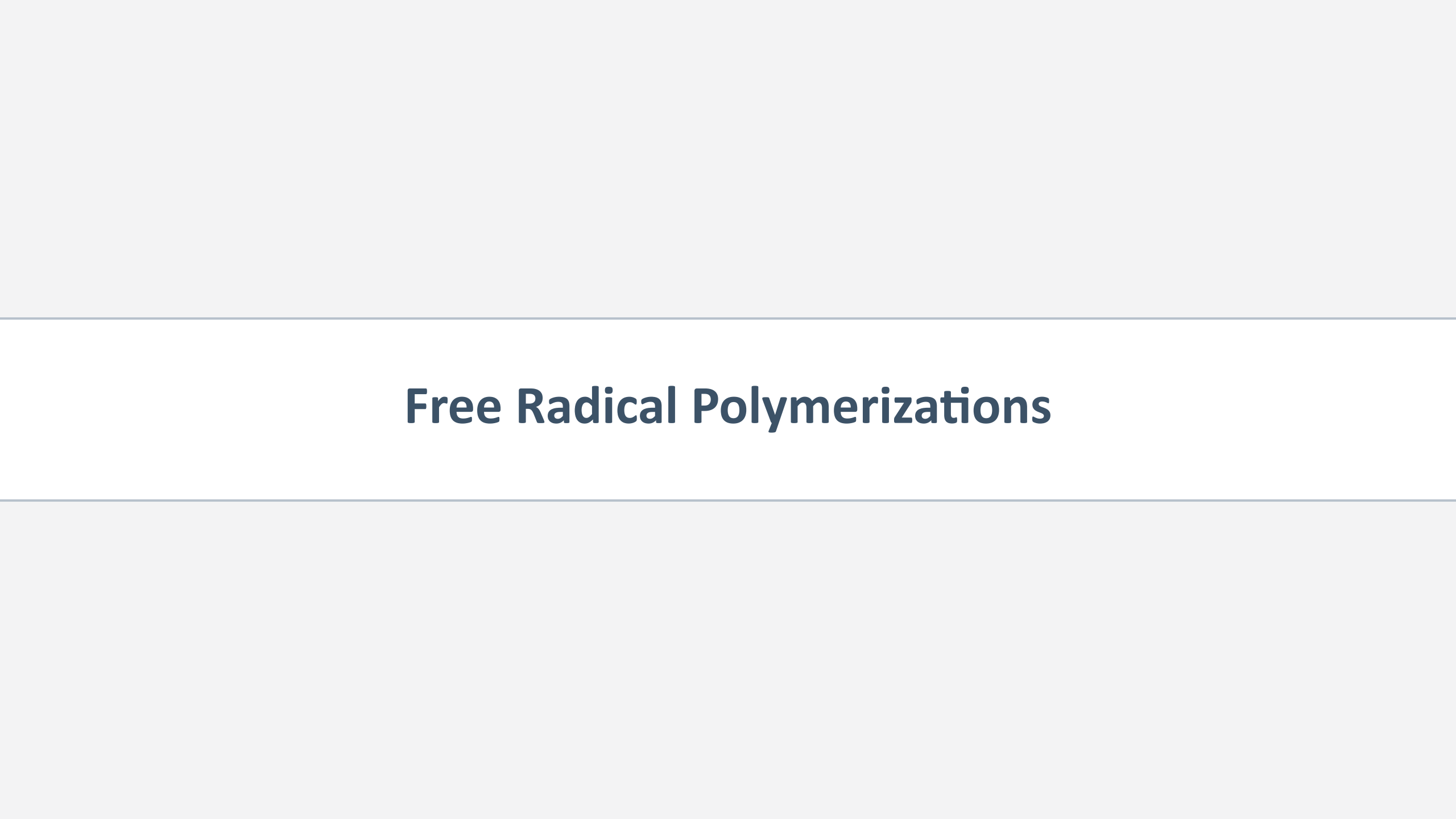
- polymerizations require an initiator that attacks a first monomer and creates an active center
- during propagation, monomers add consecutively to the active center of a growing polymer chain
- initiation continuously occurs during entire polymerization time
- termination is a stochastic event, greatly determining the molecular weight distribution

Molar Mass Distribution in Chain Grwoth Polymerizations

• Flory-Schulz-type distribution is expected (at least, in the low conversion regime)



- high molar mass polymer is formed from the beginning in a free radical polymerization
- with increasing time and conversion, the number of each species increases continuously



Initiation and Propagation in Free Radical Polymerizations

initiator decomposition (slow)

initiation (fast)

propagation (chain growth)

- reaction with a monomer instead of immediate recombination with efficiency factor $f \approx 0.3-0.8$
- initiator decomposition is a statistical process, occurring slowly throughout polymerization process
- radical life time $\tau = 0.1-10$ s, about 100–10'000 propagation steps before chain terminates

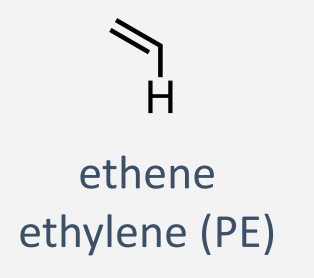
Termination Reactions in Radical Polymerizations

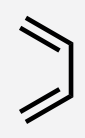
termination by radical combination

termination by disproportionation

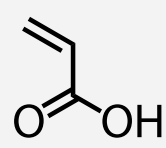
• termination by combination and disproportionation often occur both and are stochastic processes

Important Vinyl Monomers in Radical Polymerization

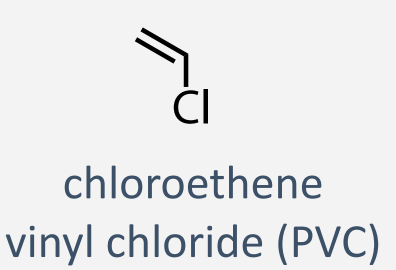


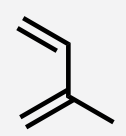


butadiene (PB)

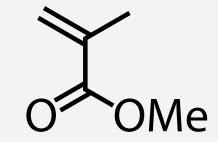


propenoic acid acrylic acid (PAA)

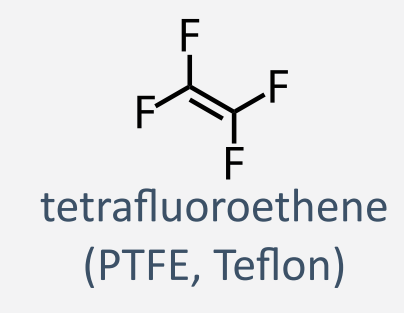


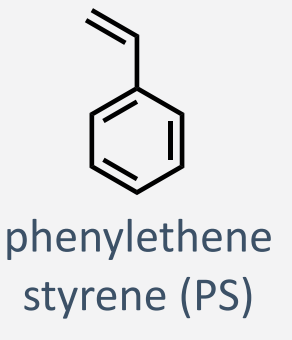


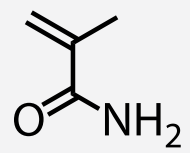
2-methylbutadiene isoprene (PI)



methyl 2-methylpropenoate methyl methacrylate (PMMA)







2-methylpropenoic amide methacrylamide (for PMAAm)

• vinyl monomers mostly with +M or -M substituents used for technologically relevant polymers

Examples of Radical Initiators and their Decomposition

azobis(isobutyronitrile) (AIBN)

di(tert.-butyl peroxide) (DTPO)

$$\sum_{i=1}^{\infty} o_{i}^{i}$$

dicumyl peroxide (DCPO)

dibenzoyl peroxide (DBPO)

$$2 \longrightarrow -2 CO_2 \uparrow$$

- suitable decomposition rates ($k_d = 10^{-7} 10^{-6} \text{ M s}^{-1}$) for different temperature regimes
- goal is to balance initiation and termination reaction rates to reach "steady state" conditions

Kinetics of Chain Growth Polymerization

rate of initiation

rate of propagation

rate of termination

$$R_i = \frac{d[\mathbf{R}^{\bullet}]}{dt} = 2fk_d[\mathbf{I}]$$

$$R_{i} = \frac{d[\mathbf{R}^{\bullet}]}{dt} = 2fk_{d}[\mathbf{I}] \qquad \qquad R_{p} = -\frac{d[\mathbf{M}]}{dt} = \sum_{i} k_{p}[\mathbf{M}_{i}^{\bullet}][\mathbf{M}] \qquad \qquad R_{t} = -\frac{d[\mathbf{M}^{\bullet}]}{dt} = 2k_{t}[\mathbf{M}^{\bullet}]^{2}$$

$$R_t = -\frac{d[\mathbf{M}^{\bullet}]}{dt} = 2k_t[\mathbf{M}^{\bullet}]^2$$

steady-state-conditions:

$$R_i = R_t$$

$$R_i = R_t \qquad \qquad R_p = k_p \sqrt{\frac{f k_d}{k_t}} \sqrt{[\mathbf{M}][\mathbf{I}]} \qquad \text{with } k_t = k_{tc} + k_{td}$$

with
$$k_t = k_{tc} + k_{td}$$

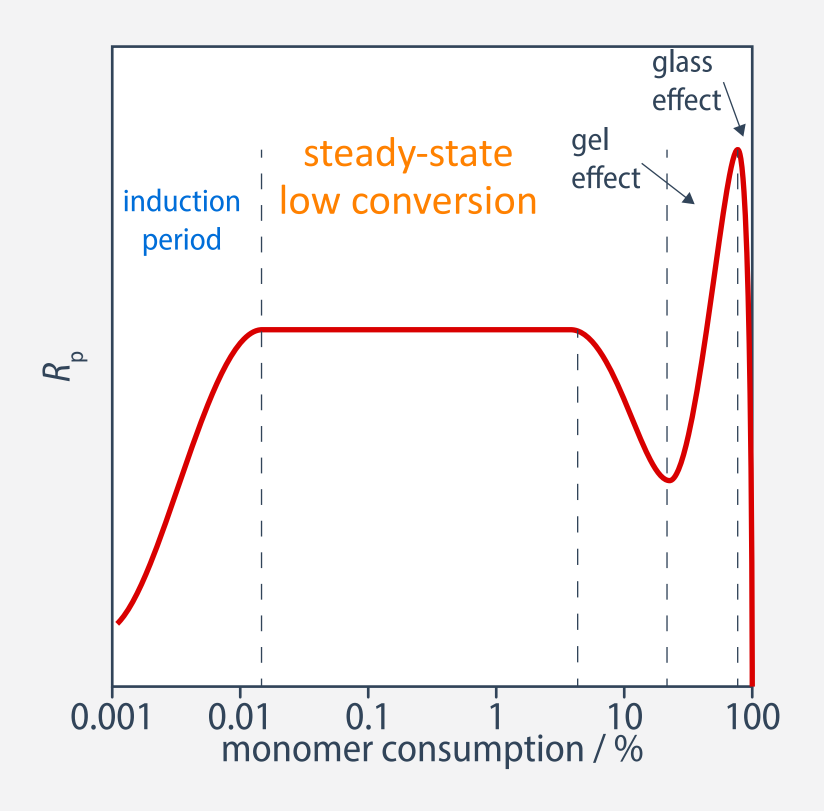
• kinetic chain length:

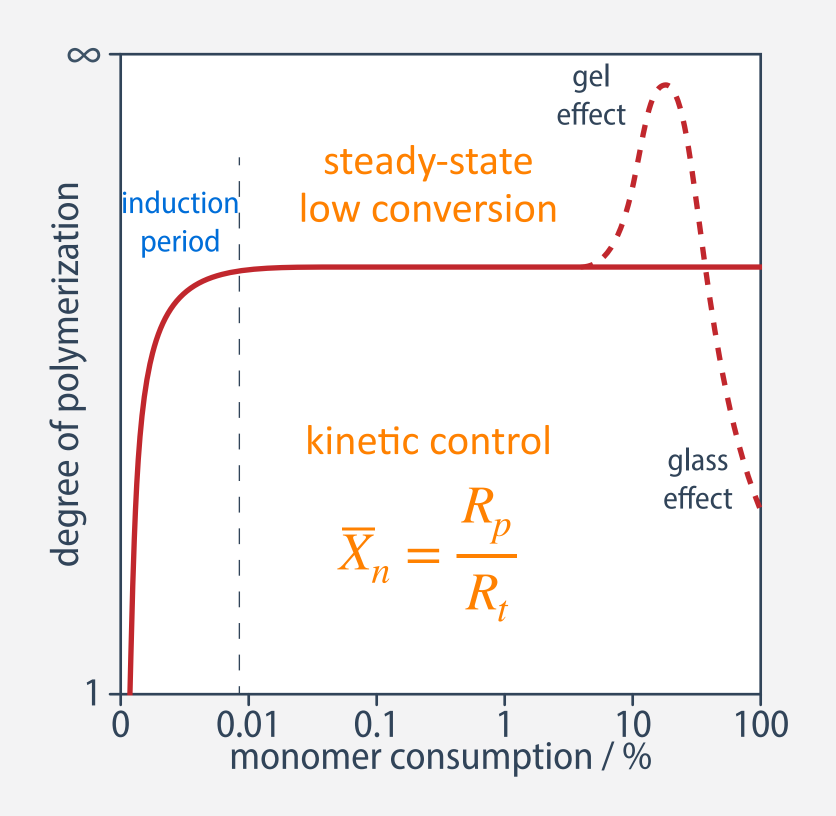
$$\overline{v} = \frac{R_p}{R_i} = \frac{R_p}{R_t} = \frac{k_p[M]}{2(fk_ik_t[I])^{1/2}} \propto \frac{[M]}{\sqrt{[I]}}$$

- steady state conditions required for stable polymerization, results in reaction order 0.5 for initiator
- increasing initiator concentration increases polymerization rate but results in decreased molar mass

High Conversion Effects

- gel effect: diffusion-controlled termination (auto-acceleration of propagation)
- glass effect: monomers get trapped, if the matrix becomes increasingly glassy

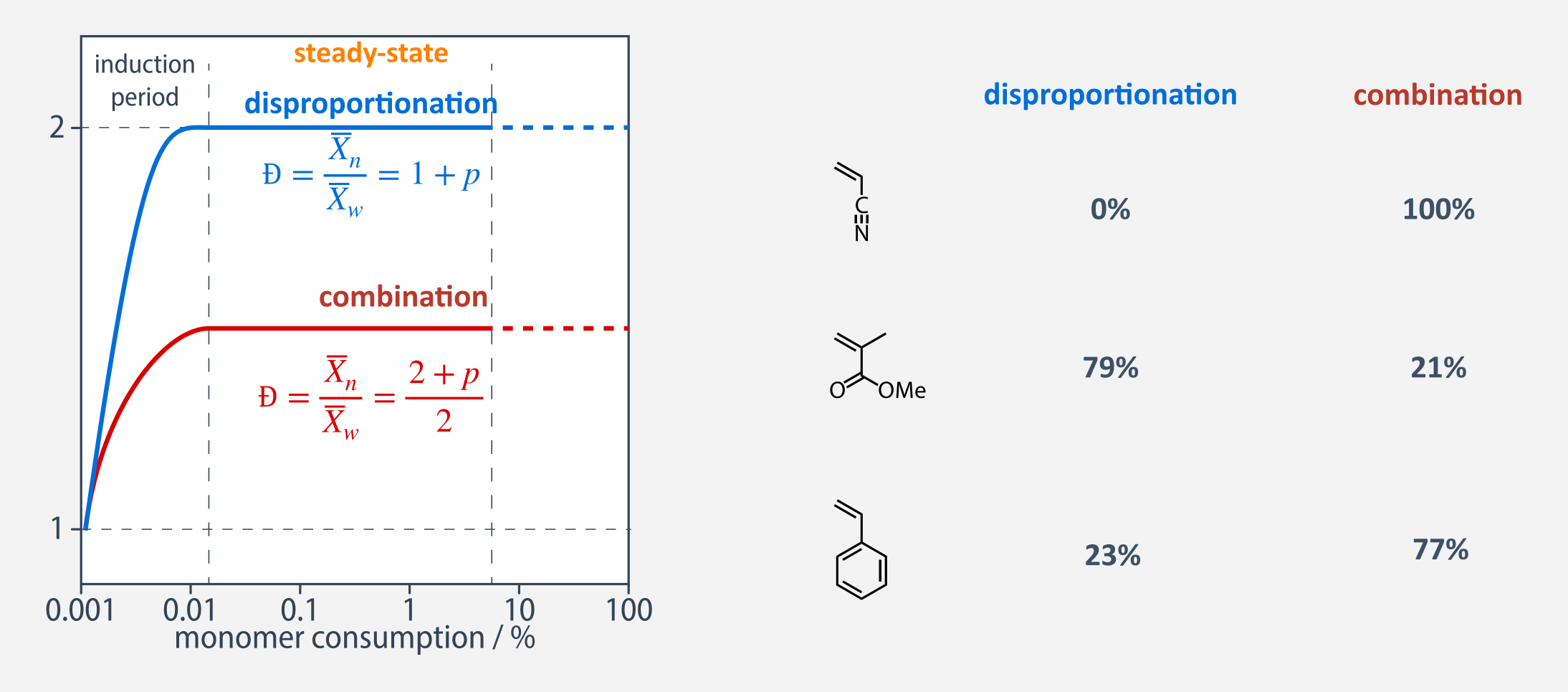




• steady-state conditions are not maintained at medium to high conversions, causing a loss of control over polymerization rate and molar mass distribution

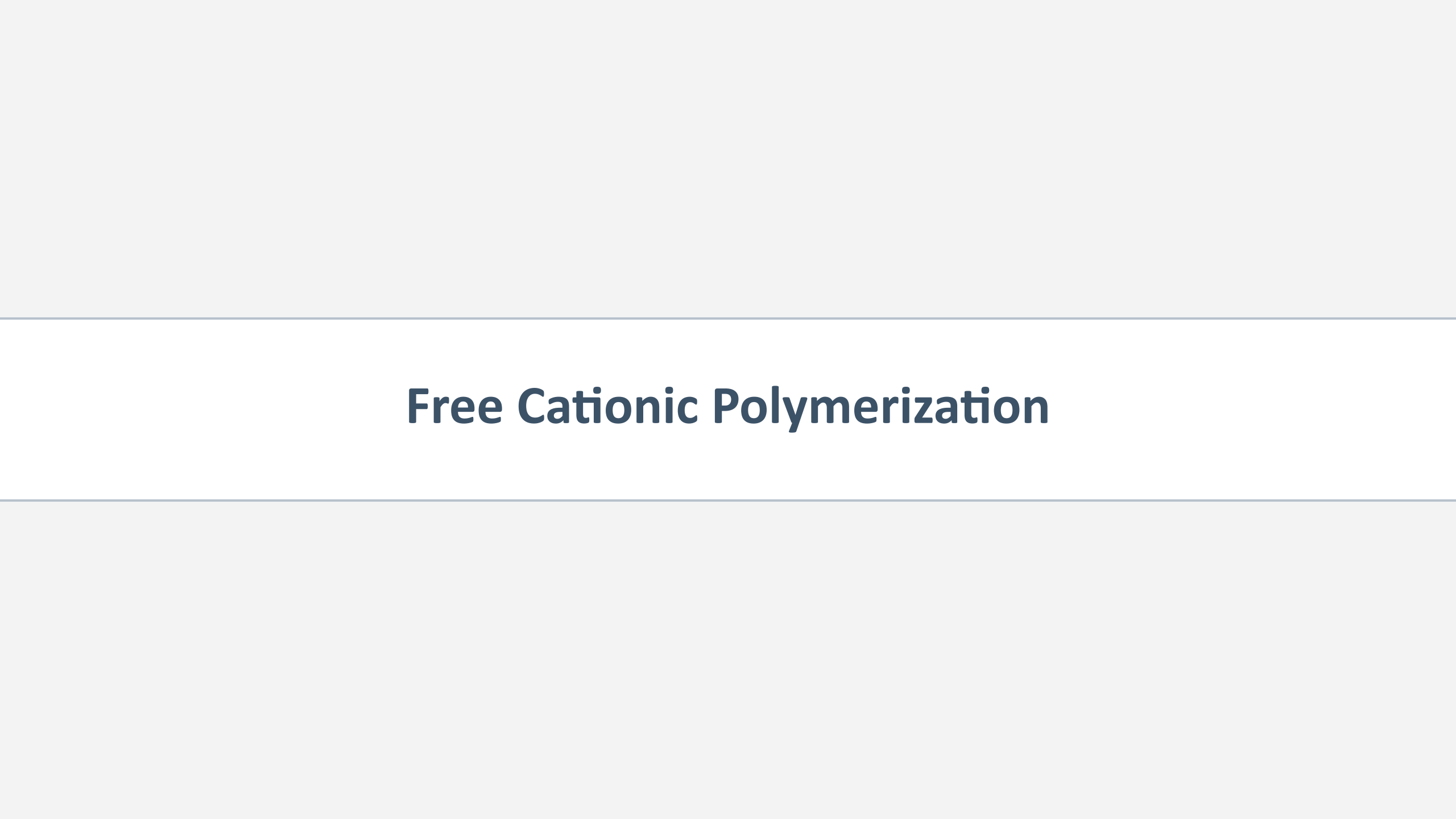
Dispersity in the Low Conversion Regime

- in reality, termination may occur via both pathways
- transfer reactions and side-reactions are not taken into consideration



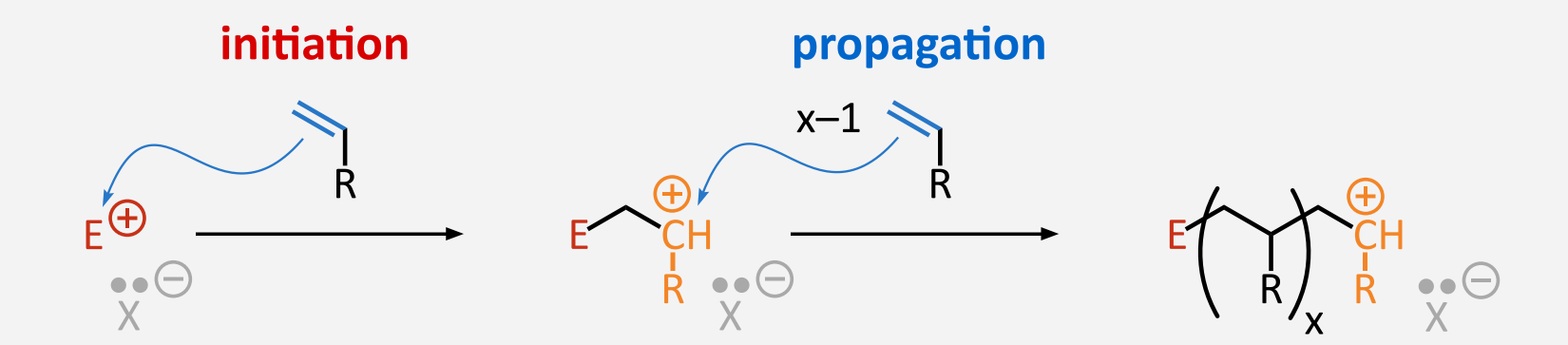
molecular weight distribution and dispersity depend on termination mechanism





Cationic Polymerization of Vinyl Monomers

• polymerization started by reaction of a vinyl monomer with a strong (cationic) electrophile



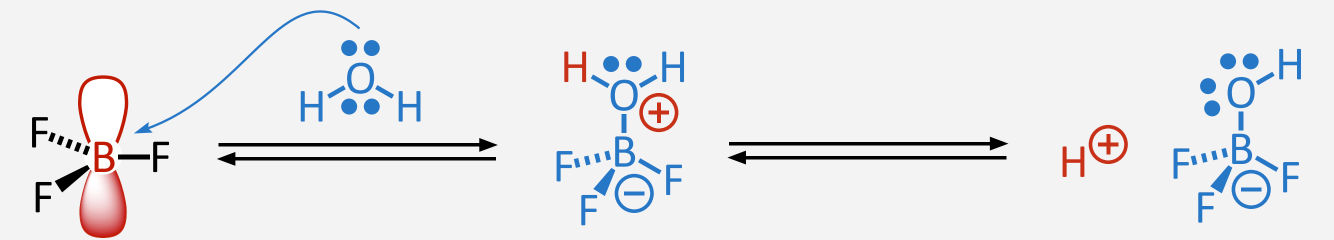
- initiation is like the first step in electrophilic addition to a double bond (of the monomer)
- reactive chain end is a (highly electron-deficient, elctrophilic, reactive) carbocationic intermediate
- monomer must have electron-donating residue R to be reactive, but also to stabilize chain end
- initiator must be sufficiently strong Lewis or Brønsted acid, hence counterion must be well stabilized
- counterion must be well stabilized / non-nucleophilic also to avoid completing the addition reaction
- cabocationic chain end and counterion are close ion pairs in organic solvents (even when not shown)
- cationic polymerization like first step of electrophilic addition, while avoiding the second step

Examples of Initiators for Cationic Polymerization of Vinyl Monomers

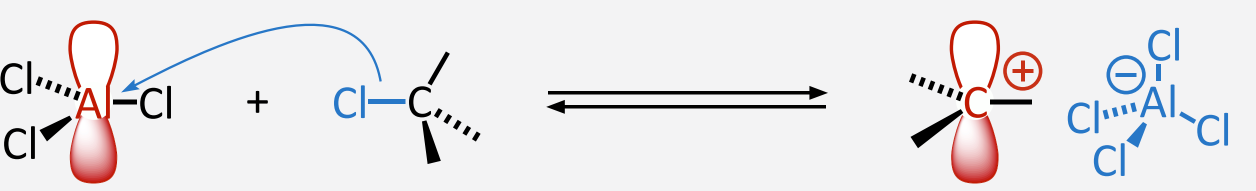
• Brønsted acids with large, resonance-stabilized, non-coordinating, non-nucleophilic counterions



- hydrogen halogenides are inefficient as initiators
- halogenide counterions are still too nucleophilic, complete addition to the double bond
- Lewis acids such as halides of metals in high oxidation states (BF₃, BCl₃, AlCl₃, TiCl₄, SnCl₄, SbCl₅, ZnCl₂)
 - two coinitiators that deliver a proton as the actual initiator

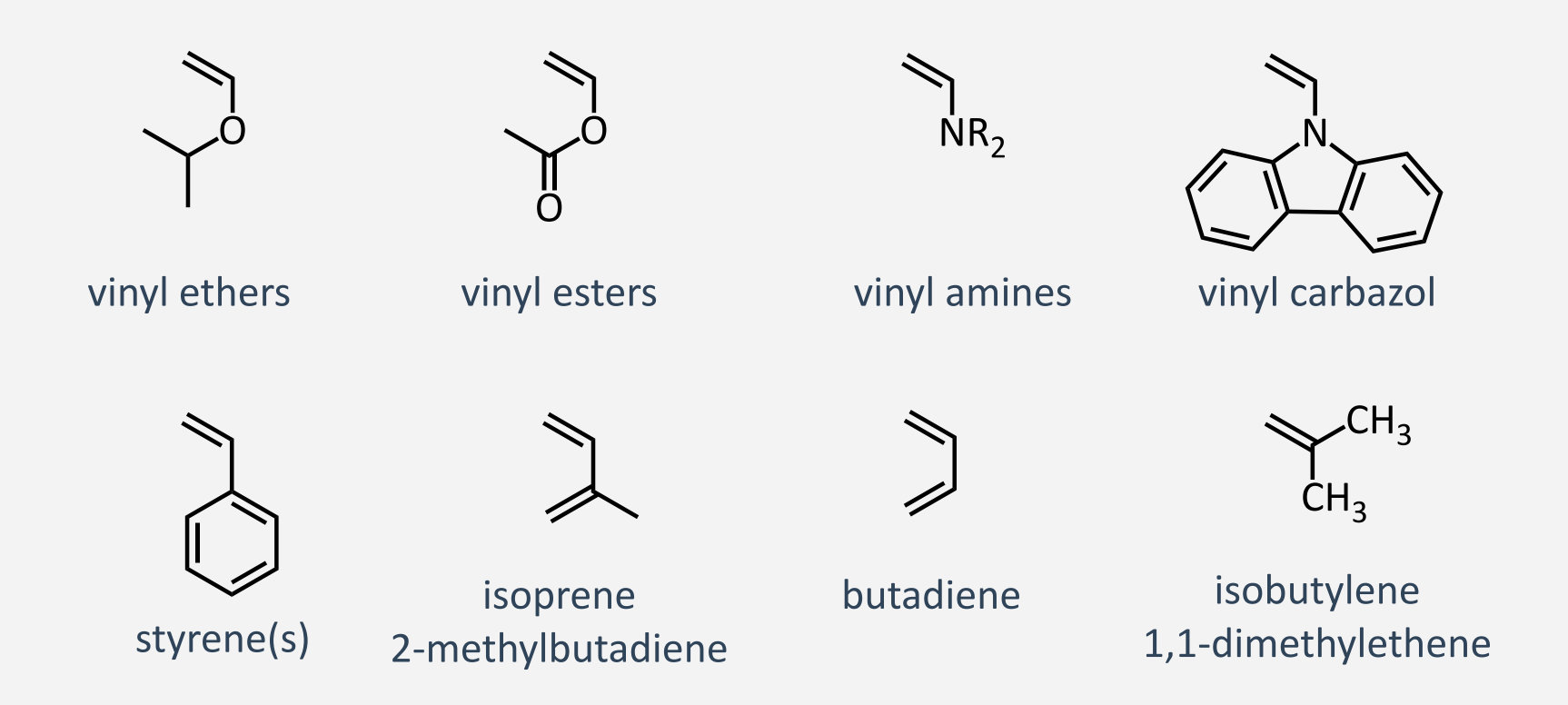


• two coinitiators, one with a good leaving group that generates a carbocation



Examples of Monomers for Cationic Polymerization of Vinyl Monomers

• +M substituents or 1,1-dialkyl substitution (stabilized and polarized by hyperconjugation)



- electron-donating group to increase reactivity towards electrophile and bias for terminal reaction
- also necessary to stabilize the carbocationic center on the reactive chain end

Chain Transfer Reactions

• chain transfer by β-hydrogen transfer to monomer is prevalent mechanism

$$E \stackrel{H_{\beta}}{\longleftarrow} CH^{\oplus} \stackrel{K_{tr}}{\longleftarrow} E \stackrel{CH}{\longleftarrow} CH + H^{\oplus}$$

$$\downarrow R \\ \downarrow R \\$$

- individual polymer chain is deactivated but kinetic chain reaction continues
- both propagation and chain transfer are first order in monomer and in active chain end
- molar mass determined by kinetics, inverse of chain transfer constant $C_{\rm tr} = k_{\rm tr} / k_{\rm p}$
- polymerization typically performed at (very) low temperatures to increase molar mass
- β-hydrogen transfer results in double bond chain ends; polymerizable hence risk of branching
- chain transfer to monomer is principal process that limits molar mass and results in Schulz-Flory distribution, especially at reaction temperatures above r. t.

Termination Reactions

• chain termination is possible by combination with counterion (in some cases)

• different from radical polymerization, terminations are not prevalent in cationic polymerizations

Quenching

• carbocationic chain end remains reactive intermediate, is "quenched" (deactivated) by nucleophiles

- strong nucleophiles (anionic, with wekaly acidic counterions) result in irreversible termination
- weak nucleophiles that release strongly acidic coutnerion for accelerated chain transfer
- useful to tailor molar mass control

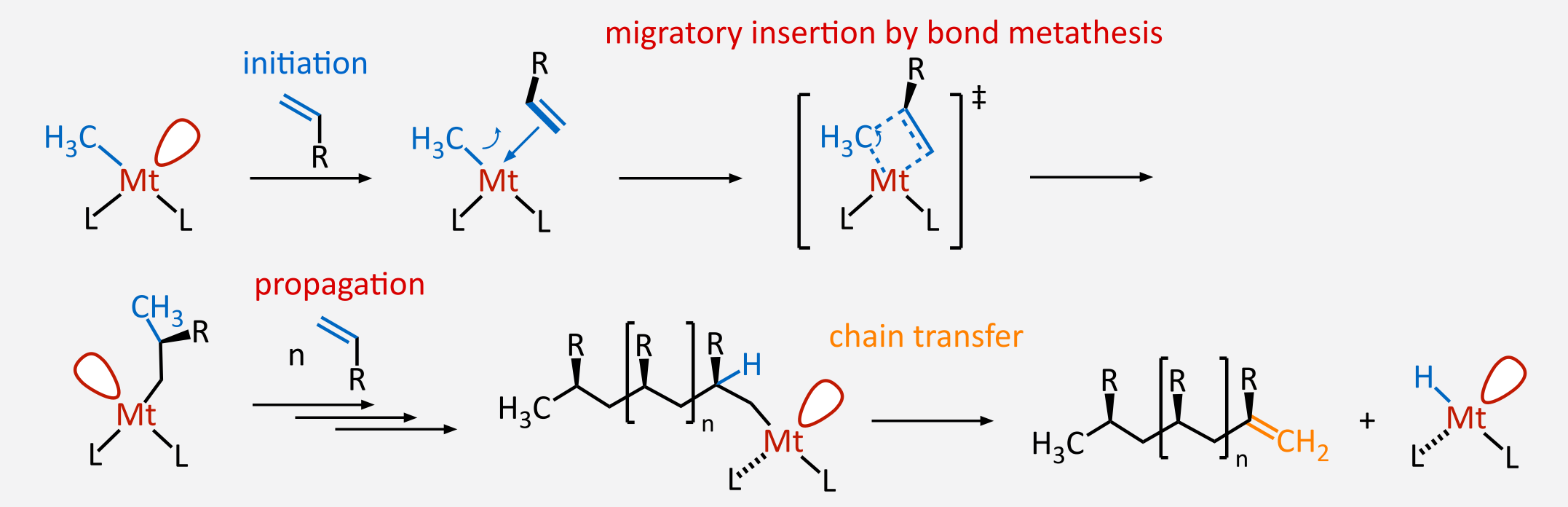
$$E \stackrel{\bigoplus}{\bigcap} H$$

- quenching generally useless for introducing end groups deliberately because of chain transfer
- cationic polymerizations require careful scavenging of inadvertent nucleophiles (including water)



Generic Mechanism of Transition-Metal-Catalyzed Coordination Polymerization

• coordination (insertion) polymerization of alkenes by transition metals with empty coordination site



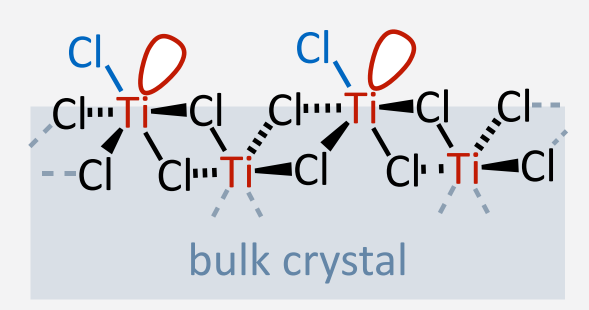
- initiation by first alkene monomer adding as ligand to the vacant coordination site (empty d orbital)
- propagation by migratory insertion of the monomer into Mt–C bond, polymer chain migrates
 - via four-electron, four-membered cyclic transition state
 - finalized by bond metathesis, generating new Mt–C & C–C bonds, reforming empty coordination site
- chain transfer by β-hydrogen elimination from last unit, generates double bond chain end

• typical catalysts are a wide variety of transition metal solids and moelcular organometallic compounds

heterogeneous catalysts

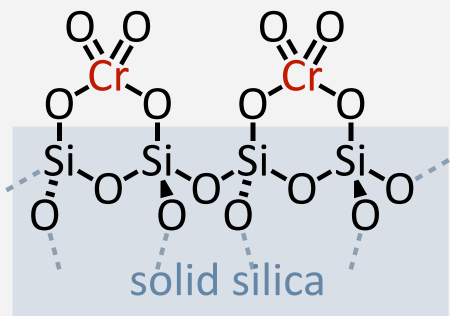
homogeneous catalysts

crystal surface



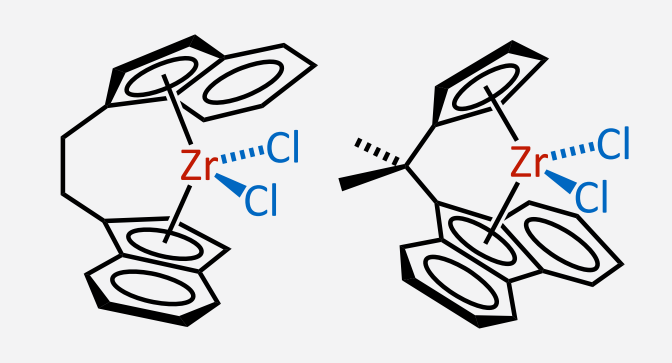
Ziegler-Natta crystalline TiCl₃ particles

particle surface

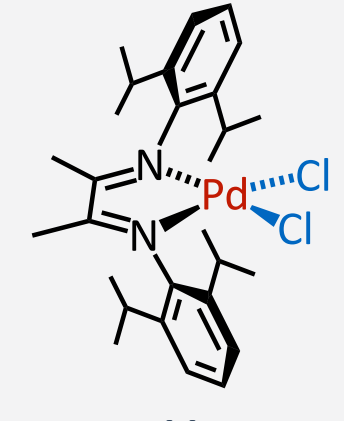


Phillips

chromium oxide
supported on silica gel



Brintzinger Zirconocenes

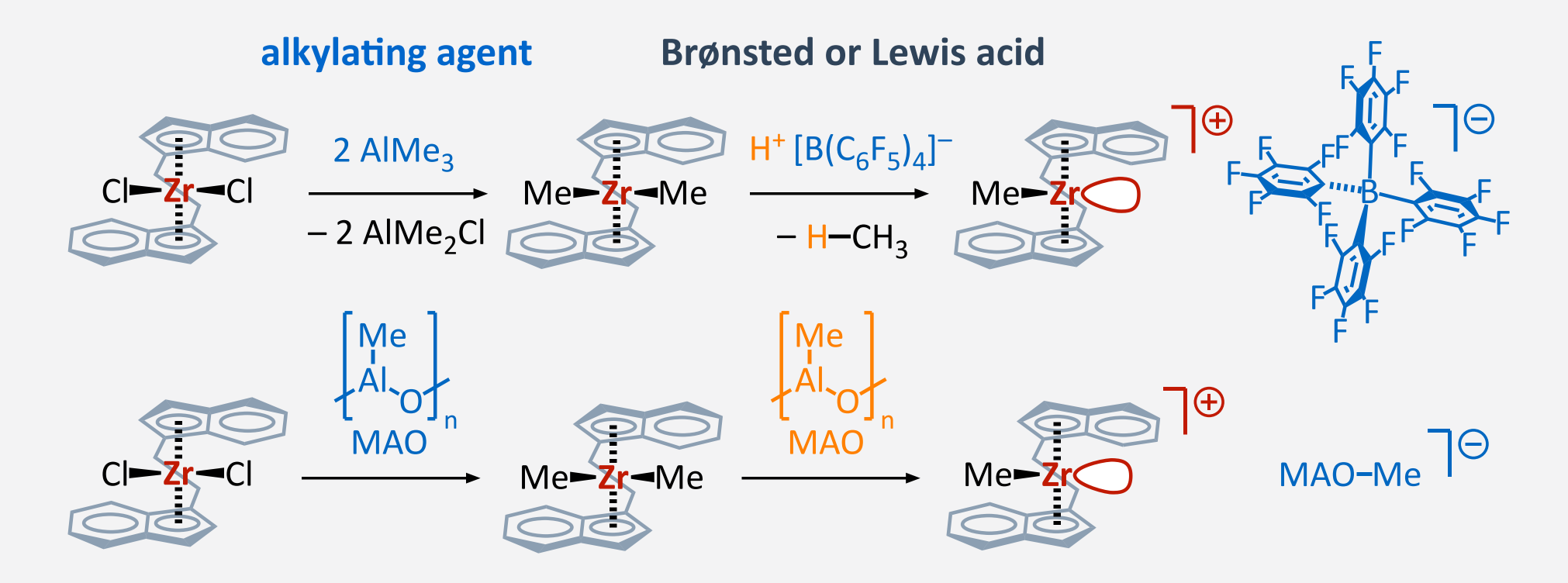


Brookhartpalladium diimines

- heterogeneous catalysts are solid particles dispersed in reaction medium, only surface is active
- homogeneous catatalysts are defined organometallic compounds, in homogeneous solution

Catalyst Activation

• catalyst activation requires generating an alkylated metal center with a free coordination site



- alkylation of stable precursors typically achieved with AlMe₃ or methylaluminoxide (MAO)
- alkylated metal center with a free coordination site is highly electron-deficient species
- very strong Lewis/Brønsted acid with weakly coordinating anion required for its formation
- MAO (clusters of about 20 Al atoms, unknown structure) does both, common in industrial processes

Example of a Ziegler-Natta Polymerization via Cossee-Arlman Mechanism

• migratory insertion means that polymer switches coordination site with every insertion step

Example of a Ziegler-Natta Polymerization

surface
$$H_3C$$
 H_3C CH_2 H_3C CH_2 H_3C CH_2 CH_2 CH_2 CH_2 CH_3 CH_3

- Cossee Arlman mechanism: monomer coordination to empty coordination site on surface metal atom
- migratory insertion into Mt–C bond, means that polymer switches coordination site at every step
- chain transfer generates terminal double bonds, can result in branches when polymerized