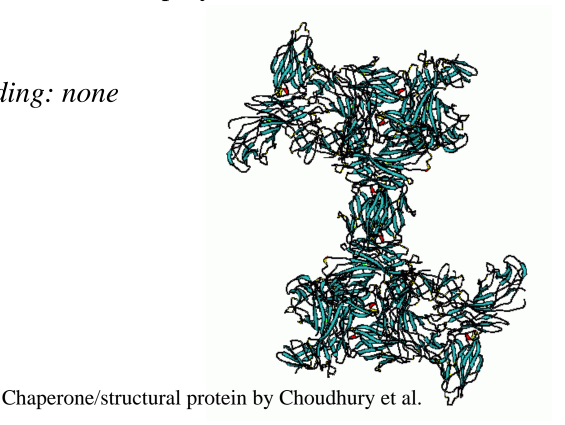
### **Chapter Outline: Polymer Structures**

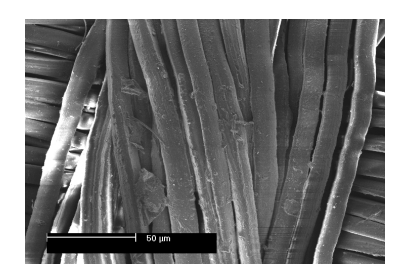
- Hydrocarbon and polymer molecules
- > Chemistry of polymer molecules
- ➤ Molecular weight and shape
- ➤ Molecular structure and configurations
- > Thermoplastic and thermosetting polymers
- ➤ Copolymers
- ➤ Polymer crystals and degree of crystallinity
- > Defects and diffusion in polymers

Optional reading: none



## **Polymers: Introduction**

- ➤ **Polymers** materials consisting of *polymer molecules* that consist of repeated chemical units (`mers') joined together, like beads on a string. Some polymer molecules contain hundreds or thousands of monomers and are often called *macromolecules*.
- ➤ Polymers may be **natural**, such as leather, rubber, cellulose or DNA, or **synthetic**, such as nylon or polyethylene.



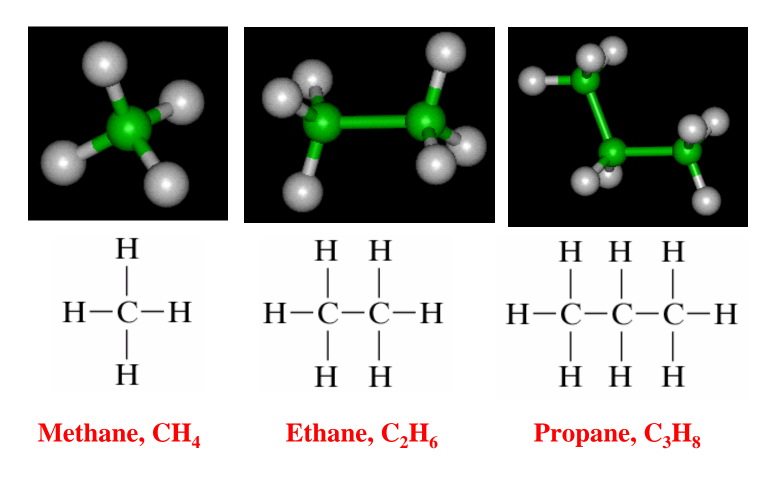
Silk fiber is produced by silk worms in a cocoon, to protect the silkworm while it metamorphoses in a moth.

Many of important current research problems and technological applications involve polymers. Living organisms are mainly composed of polymerized amino acids (proteins) nucleic acids (RNA and DNA), and other *biopolymers*. The most powerful computers - our brains - are mostly just a complex polymer material soaking in salty water. We are just making first small steps towards understanding of biological systems.

### **Hydrocarbon molecules (I)**

- Most polymers are organic in their origin and are formed from hydrocarbon molecules
- Each C atom has four e<sup>-</sup> that participate in bonds, each H atom has one bonding e<sup>-</sup>

Examples of **saturated** (all bonds are single ones) hydrocarbon molecules:

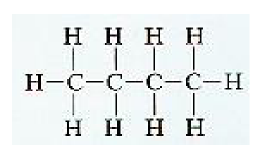


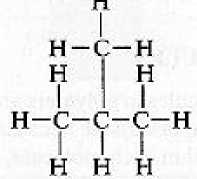
## **Hydrocarbon molecules (II)**

Double and triple bonds can exist between C atoms (sharing of two or three electron pairs). Molecules with double and triple bonds are called **unsaturated**. Unsaturated molecules are more reactive

Ethylene, C<sub>2</sub>H<sub>4</sub> Acetylene, C<sub>2</sub>H<sub>2</sub>

Isomers are molecules that have the same composition (contain the same atoms) but have different atomic arrangement. An example is butane and isobutane:





Butane  $\rightarrow C_4H_{10} \leftarrow Isobutane$ 

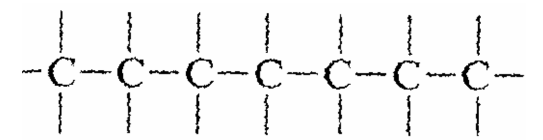
physical properties (e.g. boiling temperature) depend on the isomeric state

## **Hydrocarbon molecules (III)**

Many other organic groups can be involved in polymer molecules. In table below R represent a radical, an organic group of atoms that remains as a unit and maintains their identity during chemical reactions (e.g.  $CH_3$ ,  $C_2H_5$ ,  $C_6H_5$ )

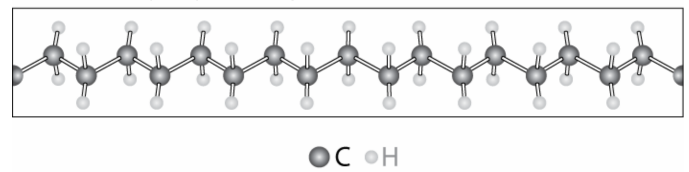
Family	Characteristic Unit	Representative Compound	
Alcohols	R—ОН	H  -  - H OH  -  -	Methyl alcohol
Ethers	R—O—R′	H H 	Dimethyl ether
Acids	R—C O	H—C—C H—O	Acetic acid
Aldehydes	R C=O	C=0	Formaldehyde
Aromatic hydrocarbons	R	OH	Phenol

## **Polymer molecules**



- Polymer molecules can be very large (macromolecules)
- Most polymers consist of long and flexible chains with a string of C atoms as a backbone
- Side-bonding of C atoms to H atoms or radicals
- > Double bonds are possible in both chain and side bonds
- ➤ Repeat unit in a polymer chain ("unit cell") is a mer
- > Small molecules from which polymer is synthesized is monomer. A single mer is sometimes also called a monomer.

polyethylene (e.g. paraffin wax for candles)



## Chemistry of polymer molecules (I)

- $\triangleright$  Ethylene (C<sub>2</sub>H<sub>4</sub>) is a gas at room temp and pressure
- Ethylene transform to polyethylene (solid) by forming active mer through reaction with initiator or catalytic radical (R·)
- (·) denotes unpaired electron (active site)

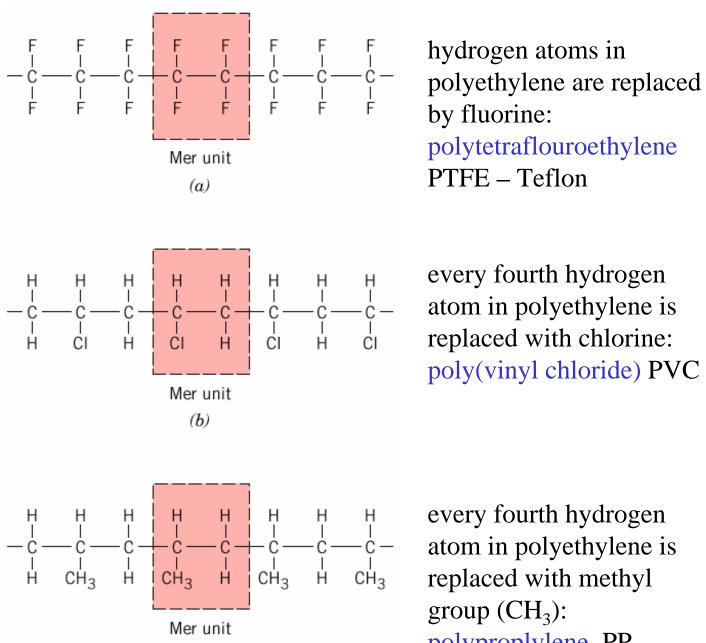
### **Polymerization:**

1. Initiation reaction: 
$$R \cdot + C = C \longrightarrow R - C - C \cdot$$
 $| \quad | \quad | \quad |$ 
 $| \quad |$ 

2. Rapid propagation ~1000 mer units in 1-10 ms:

3. Termination when two active chain ends meet each other or active chain end meet with initiator or other species with single active bond:

## Chemistry of polymer molecules (II)



polytetraflouroethylene PTFE - Teflon

every fourth hydrogen poly(vinyl chloride) PVC

group (CH<sub>3</sub>): polyproplylene PP

More examples on pp. 539-540 of the textbook

(c)

## **Chemistry of polymer molecules (III)**

- When all mers are the same, the molecule is called a homopolymer
- When there is more than one type of mer present, the molecule is a **copolymer**
- Mer units that have 2 active bonds to connect with other mers are called **bifunctional**
- Mer units that have 3 active bonds to connect with other mers are called **trifunctional**. They form three-dimensional molecular network structures

Polyethilene (bifunctional)

$$CH_2$$
 $CH_2$ 
 $CH_2$ 

Phenol-formaldehyde (trifunctional)

## Molecular weight

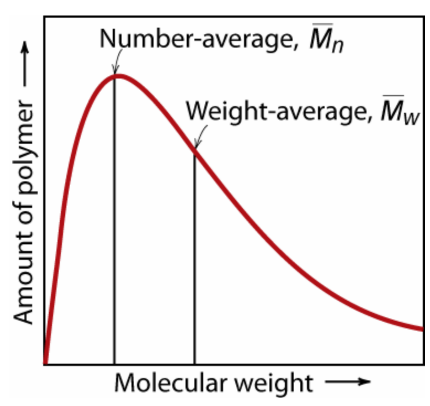
- The molecular weight (chain length) is controlled by the synthesis process: Relative rates of initiation, propagation, termination steps of polymerization
- Formation of macromolecules during polymerization results in a distribution of chain lengths and molecular weights
- The average molecular weight can be obtained by averaging the masses with the fraction of times they appear (number-average molecular weight) or with the mass fraction of the molecules (weight-average molecular weight).

number-average:

$$\overline{M}_{n} = \sum x_{i} M_{i}$$

weight-average:

$$\overline{M}_{\rm w} = \sum w_{\rm i} M_{\rm i}$$



 $w_i$  is weight fraction of chains of length i  $x_i$  is number fraction of chains of length i

# Molecular weight: Example illustrating the difference between number-average and weight-average

student	weight mass (lb)		
1	104		
2	116		
3	140		
4	143		
5	180		
6	182		
7	191		
8	220		
9	225		
10	380		

What is the average weight of students in this class:

- a) Based on the number fraction of students in each mass range?
- b) Based on the weight fraction of students in each mass range?

#### Solution:

The first step is to sort the students into weight ranges (let's use 40 lb ranges).

weight range	# of students	mean weight	number fraction	weight fraction
	$N_{i}$	$M_{i}$	$x_i$	$W_{i}$
81-120	2	110	0.2	0.117
121-160	2	142	0.2	0.150
161-200	3	184	0.3	0.294
201-240	2	223	0.2	0.237
241-280	0	_	0	0.000
281-320	0	_	0	0.000
321-360	0	_	0	0.000
361-400	1	380	0.1	0.202

$$\sum N_i = 10 \qquad \sum N_i M_i = 1881 \qquad x_i = N_i / \sum N_i \qquad w_i = N_i M_i / \sum N_i M_i$$

# Molecular weight: Example illustrating the difference between number-average and weight-average

weight range	# of students	mean weight	number fraction	weight fraction
	$N_{i}$	$M_{i}$	$x_i$	$W_{i}$
81-120	2	110	0.2	0.117
121-160	2	142	0.2	0.150
161-200	3	184	0.3	0.294
201-240	2	223	0.2	0.237
241-280	0	_	0	0.000
281-320	0	_	0	0.000
321-360	0	_	0	0.000
361-400	1	380	0.1	0.202

$$\sum N_i = 10$$
  $\sum N_i M_i = 1881$   $x_i = N_i / \sum N_i$   $w_i = N_i M_i / \sum N_i M_i$ 

$$\overline{M}_n = \sum x_i M_i = 0.2 \times 110 + 0.2 \times 142 + 0.3 \times 184 + 0.2 \times 223 + 0.1 \times 380 = 188 \ lb$$

$$\overline{M}_{w} = \sum w_{i} M_{i} = 0.117 \times 110 + 0.150 \times 142 + 0.294 \times 184 + 0.237 \times 223 + 0.202 \times 380 = 218 \ lb$$

$$\overline{M}_{w} > \overline{M}_{n}$$

## **Degree of polymerization**

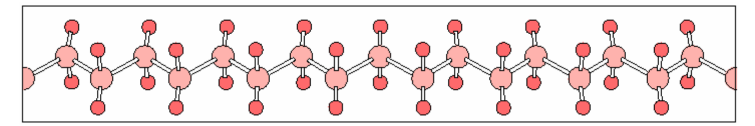
Alternative way to express average polymer chain size is **degree of polymerization** - the average number of mer units in a chain:

$$DP = \frac{\overline{M}_n}{\overline{m}}$$
 is the average molecular weight of repeat unit for copolymers it is calculated as  $\overline{m} = \sum f_i m_i$  ( $f_i$  is fraction of mer i of molecular weight  $m_i$ )

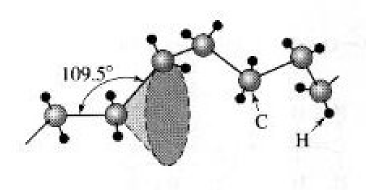
## Properties of polymers depend on molecular weight

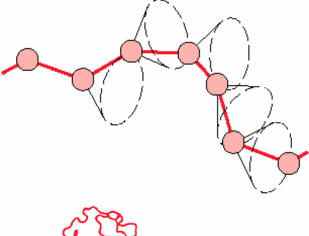
- Melting/softening temperatures increase with molecular weight (up to ~ 100,000 g/mol)
- At room temperature, short chain polymers (molar weight ~ 100 g/mol) are liquids or gases, intermediate length polymers (~ 1000 g/mol) are waxy solids, solid polymers (sometimes called *high polymers*) have molecular weights of 10<sup>4</sup> 10<sup>7</sup> g/mol

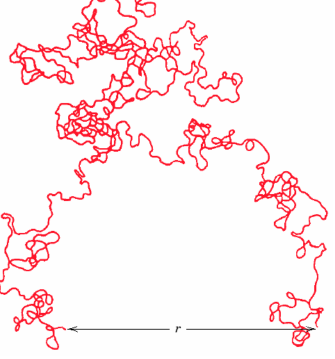
## Molecular shape (conformation)



- ➤ The angle between the singly bonded carbon atoms is ~109° carbon atoms form a zigzag pattern in a polymer molecule.
- Moreover, while maintaining the 109° angle between bonds polymer chains can rotate around single C-C bonds (double and triple bonds are very rigid).
- Random kinks and coils lead to entanglement, like in the spaghetti structure:

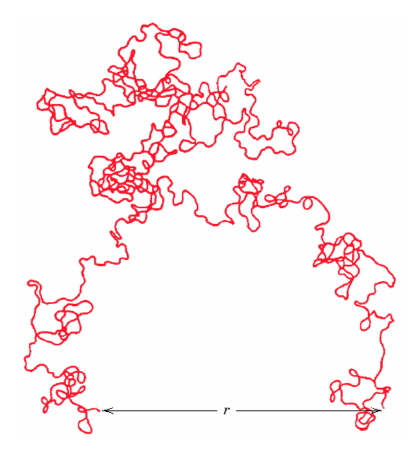






## Molecular shape (conformation)

- ➤ Molecular chains may thus bend, coil and kink
- ➤ Neighboring chains may intertwine and entangle
- Large elastic extensions of rubbers correspond to unraveling of these coiled chains
- ➤ Mechanical / thermal characteristics depend on the ability of chain segments to rotate

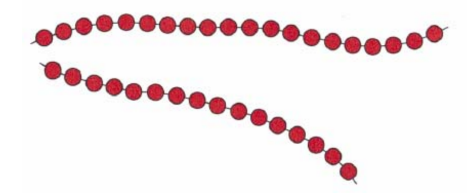


chain end-to-end distance, r

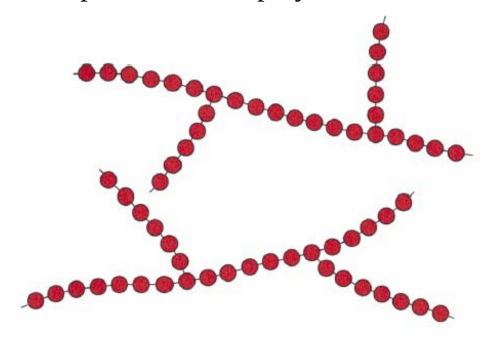
### **Molecular structure (I)**

The physical characteristics of polymer material depend not only on molecular weight and shape, but also on molecular structure:

1 Linear polymers: Van der Waals bonding between chains. Examples: polyethylene, nylon.

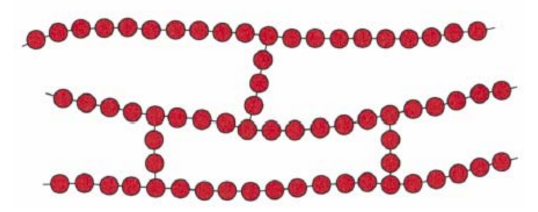


**2 Branched polymers:** Chain packing efficiency is reduced compared to linear polymers - lower density



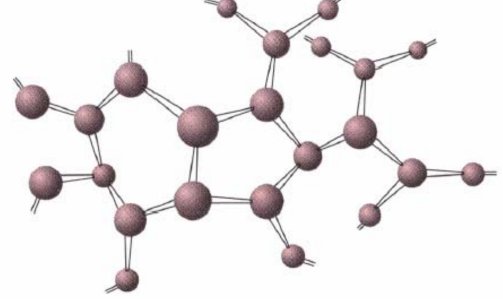
### **Molecular structure (II)**

3 Cross-linked polymers: Chains are connected by covalent bonds. Often achieved by adding atoms or molecules that form covalent links between chains. Many rubbers have this structure.



4 Network polymers: 3D networks made from trifunctional mers. Examples: epoxies, phenol-

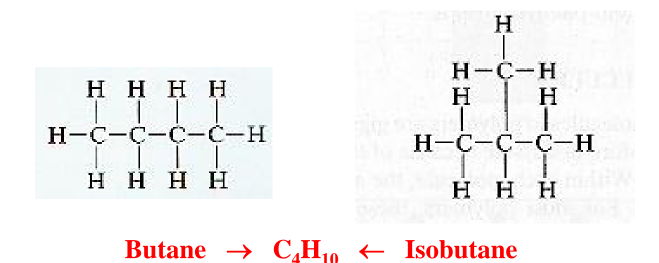
formaldehyde



#### **Isomerism**

**Isomerism**: Hydrocarbon compounds with same composition may have different atomic arrangements.

Physical properties may depend on **isomeric state** (e.g. boiling temperature of normal butane is -0.5 °C, of isobutane -12.3 °C)



Two types of isomerism in polymers are possible: stereoisomerism and geometrical isomerism

#### **Stereoisomerism**

Stereoisomerism: atoms are linked together in the same order, but can have different spatial arrangement

1 Isotactic configuration: all side groups R are on the same side of the chain.

**2 Syndiotactic configuration:** side groups R alternate sides of the chain.

3 Atactic configuration: random orientations of groups R along the chain.

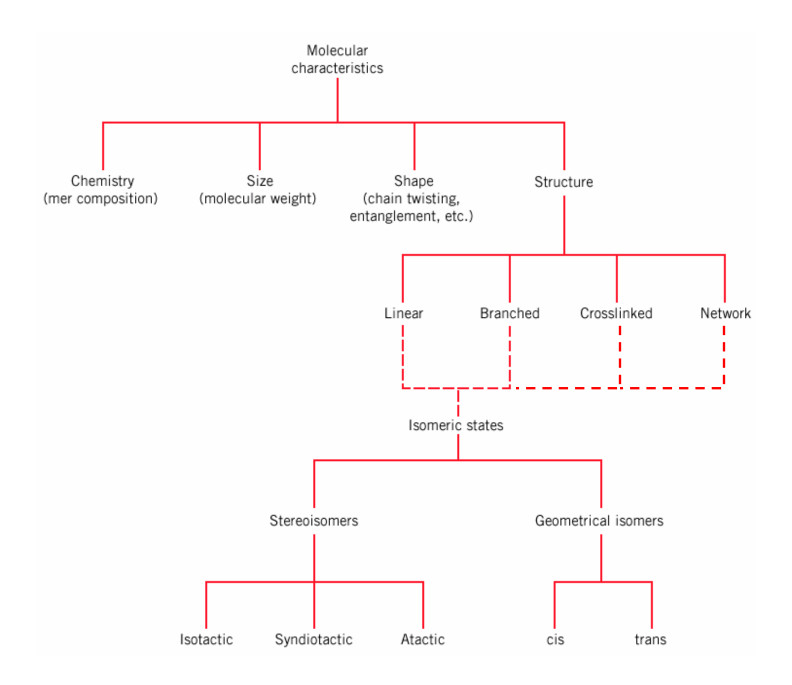
#### Geometrical isomerism

**Geometrical isomerism**: consider two carbon atoms bonded by a double bond in a chain. H atom or radical R bonded to these two atoms can be on the same side of the chain (**cis** structure) or on opposite sides of the chain (**trans** structure).

Cis-polyisoprene

**Trans**-polyisoprene

## Size - Shape - Structure classification



## Thermoplastic and thermosetting polymers

Depending on the response to temperature increase, two types of polymers can be distinguished:

(1) **Thermoplastic polymers:** soften and liquefy when heated, harden when cooled (reversible).

Molecular structure: linear or branched polymers, with secondary bonding holding the molecules together.

Easy to fabricate/reshape by application of heat and pressure

Examples: polyethylene, polystyrene, poly(vinyl chlodide).

(2) Thermosetting polymers: become permanently hard during their formation, do not soften upon heating.

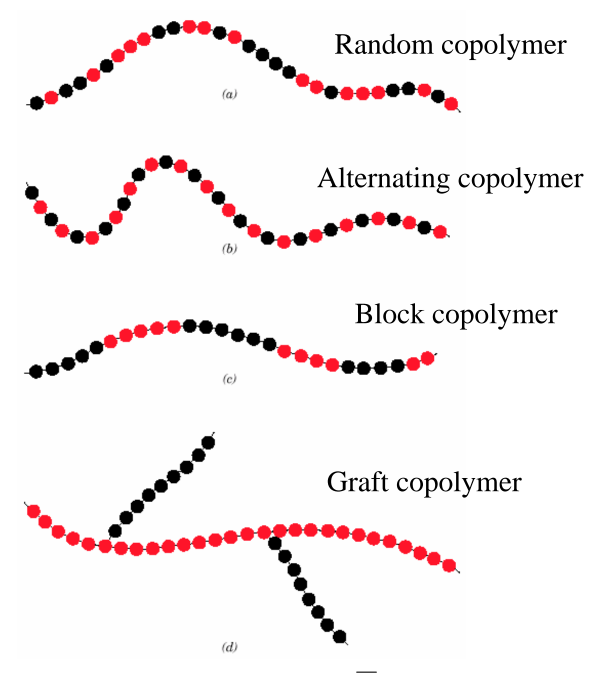
*Molecular structure:* network polymers with a large density of covalent crosslinks between molecular chains (typically, 10-50% of repeat units are crosslinked).

Harder and stronger than thermoplastics, have better dimensional and thermal stability.

Examples: vulcanized rubber, epoxies, phenolics, polyester resins.

## **Copolymers** (composed of different mers)

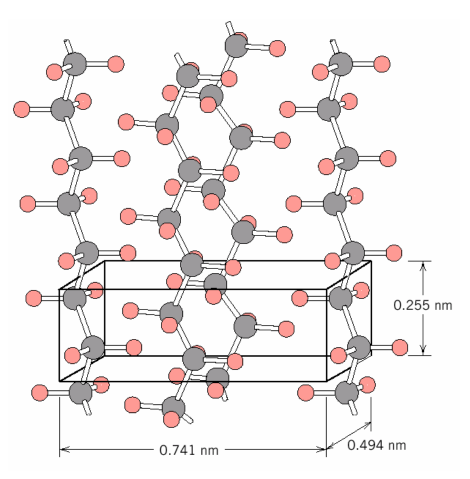
Copolymers, polymers with at least two different types of mers, can differ in the way the mers are arranged:



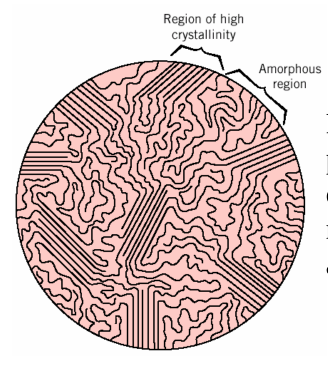
average molecular weight of repeat unit,  $m = \sum f_i m_i$ , is used in calculation of the degree of polymerization of copolymers  $(f_i \text{ is fraction of mer } i \text{ of molecular weight } m_i)$ 

## **Polymer Crystallinity (I)**

Atomic arrangement in polymer crystals is more complex than in metals or ceramics (unit cells are typically large and complex).



Polyethylene



Polymer molecules are often partially crystalline (semi-crystalline), with crystalline regions dispersed within amorphous material.

## **Polymer Crystallinity (II)**

Degree of crystallinity is determined by:

- ➤ Rate of cooling during solidification: time is necessary for chains to move and align into a crystal structure
- Mer complexity: crystallization less likely in complex structures, simple polymers, such as polyethylene, crystallize relatively easily
- Chain configuration: linear polymers crystallize relatively easily, branches inhibit crystallization, network polymers almost completely amorphous, crosslinked polymers can be both crystalline and amorphous
- ➤ **Isomerism**: isotactic, syndiotactic polymers crystallize relatively easily geometrical regularity allows chains to fit together, atactic polymers is difficult to crystallize
- Copolymerism: easier to crystallize if mer arrangements are more regular alternating, block can crystallize more easily as compared to random and graft

More crystallinity: higher density, more strength, higher resistance to dissolution and softening by heating

## **Polymer Crystallinity (III)**

Crystalline polymers are denser than amorphous polymers, so the degree of crystallinity can be obtained from the measurement of density:

% crystallinity = 
$$\frac{\rho_c(\rho_s - \rho_a)}{\rho_s(\rho_c - \rho_a)} \times 100$$

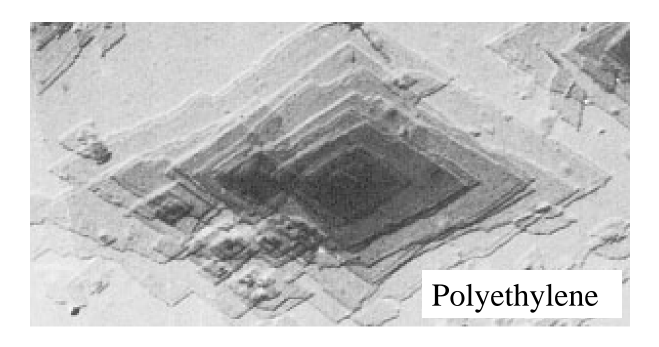
 $\rho_c$ : Density of perfect crystalline polymer

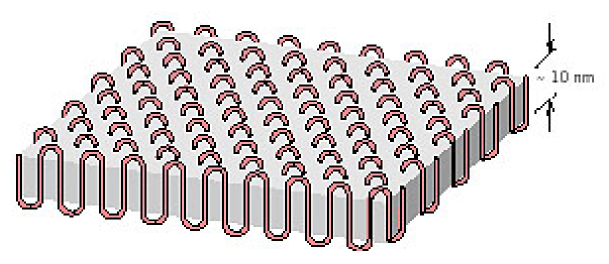
 $\rho_a$ : Density of completely amorphous polymer

 $\rho_s$ : Density of partially crystalline polymer that we are analyzing

## **Polymer Crystals (I)**

Thin crystalline platelets grown from solution - chains fold back and forth: chain-folded model

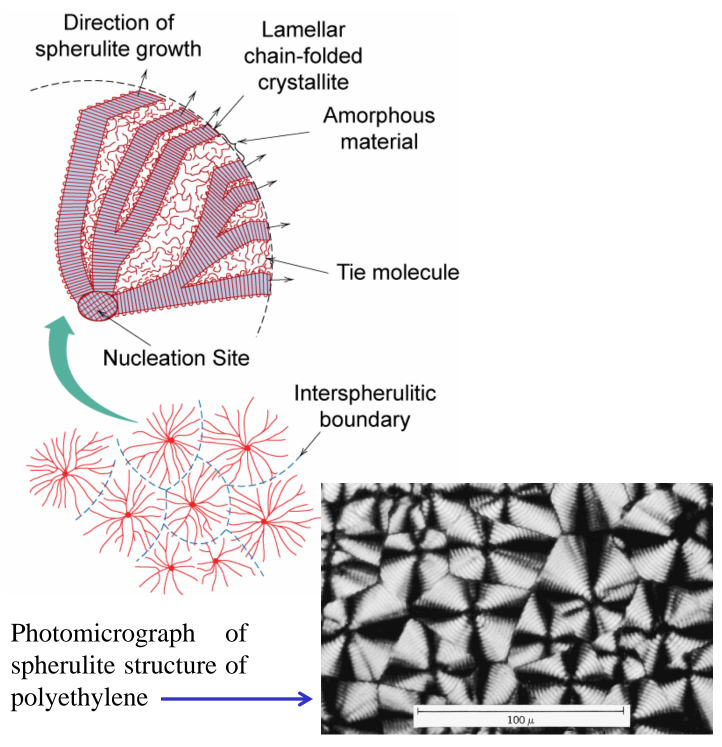




The average chain length can be much greater than the thickness of the crystallite

## **Polymer Crystals (II)**

**Spherulites:** Aggregates of lamellar crystallites ~ 10 nm thick, separated by amorphous material. Aggregates are formed upon solidification from a melted state and are approximately spherical in shape.



## **Summary**

Make sure you understand language and concepts:

- ➤ Alternating copolymer
- ➤ Atactic configuration
- ➤ Bifunctional mer
- ➤ Block copolymer
- ➤ Branched polymer
- > Chain-folded model
- > Cis (structure)
- **≻** Copolymer
- Crosslinked polymer
- ➤ Degree of polymerization
- ➤ Graft copolymer
- > Homopolymer
- > Isomerism
- ➤ Isotactic configuration
- ➤ Linear polymer
- ➤ Macromolecule
- ➤ Mer, monomer
- ➤ Molecular chemistry
- Molecular structure

- ➤ Molecular weight
- Network polymer
- ➤ Polymer
- ➤ Polymer crystallinity
- Random copolymer
- > Saturated
- > Spherulite
- > Stereoisomerism
- > Trans (structure)
- > Trifunctional mer

Five Bakers Dancing



#### Crystal

Phosphatidyl cholines taken from model lipid bilayers simulated by Heller et al., J. Phys. Chem. 97:8343, 1993.

Gel



Image by Eric Martz with RasMol