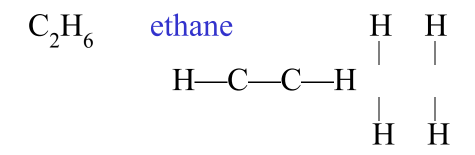
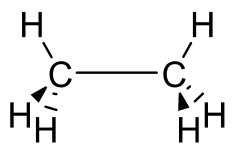
Alkanes

ALKANES

(a "family" of	f hydrocarbons)
$C_n H_{2n+2}$	CH_4
	C_2H_6
	$C_{3}H_{8}$
	C_4H_{10}
	etc.



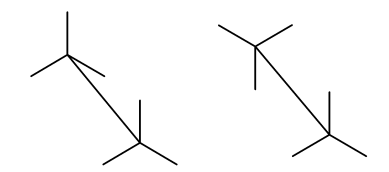
sp³, bond angles = 109.5° σ -bonds (sigma)

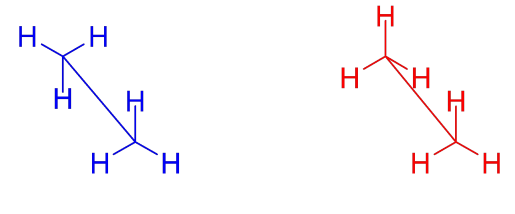


rotation about C--C (conformations)

representation:

"andiron" or "sawhorse"



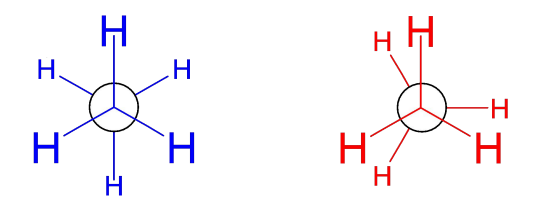


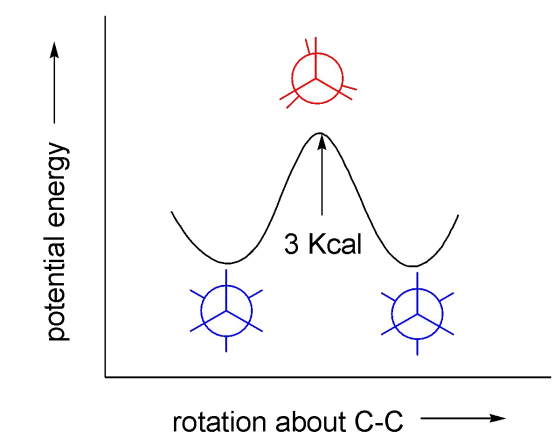
"staggered"

"eclipsed"

torsional strain: deviation from staggered.

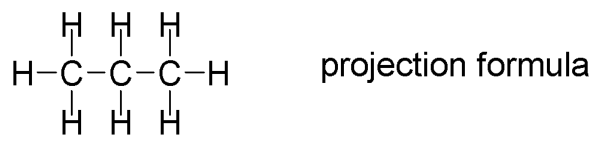
Newman projections:



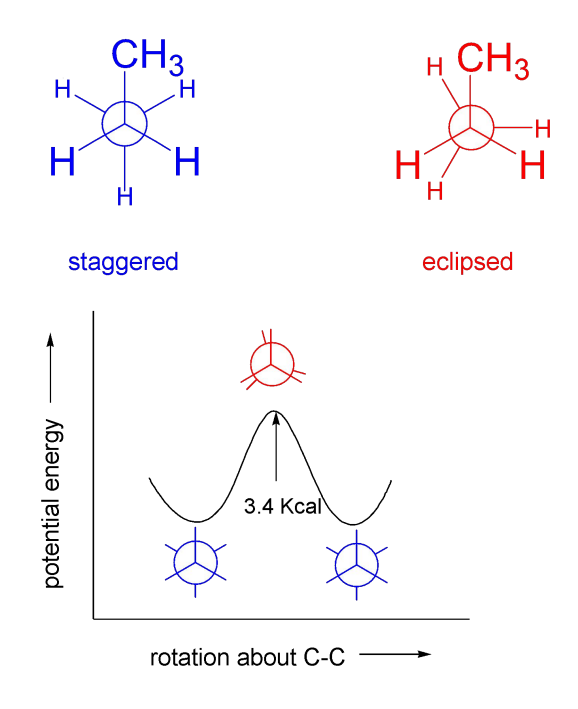


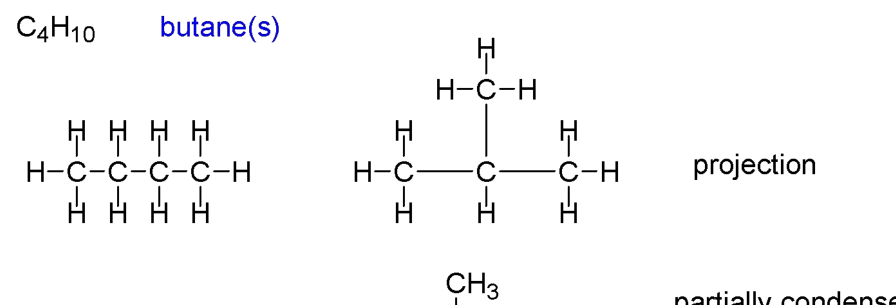
The barrier to rotation about the carbon-carbon bond in ethane is 3 Kcal/mole. The rotation is ~ "free."

C₃H₈ propane

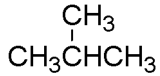


partially condensed formula $CH_3CH_2CH_3$





 $CH_3CH_2CH_2CH_3$



partially condensed





stick formulas

Two isomers of butane C_4H_{10} :

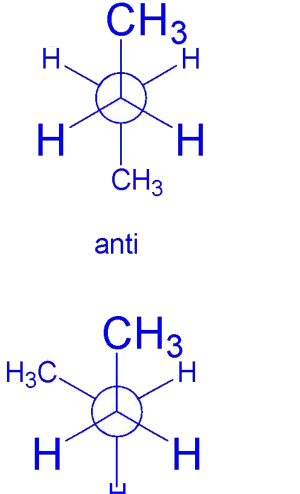
 $CH_3CH_2CH_2CH_3$ *n*-butane

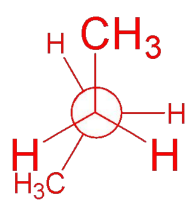
bp 0 °C mp -138 °C d 0.622 g/cc



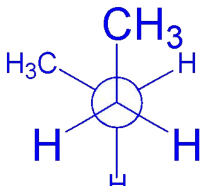
isobutane

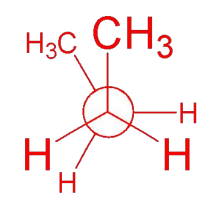
bp -12 °C mp -159 °C d 0.604 g/cc Conformations about C2-C3 in *n*-butane:





CH₃/H eclipsed

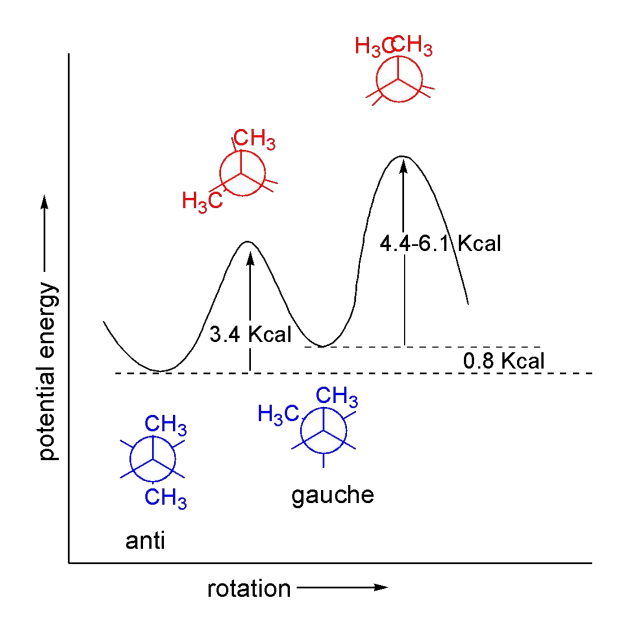


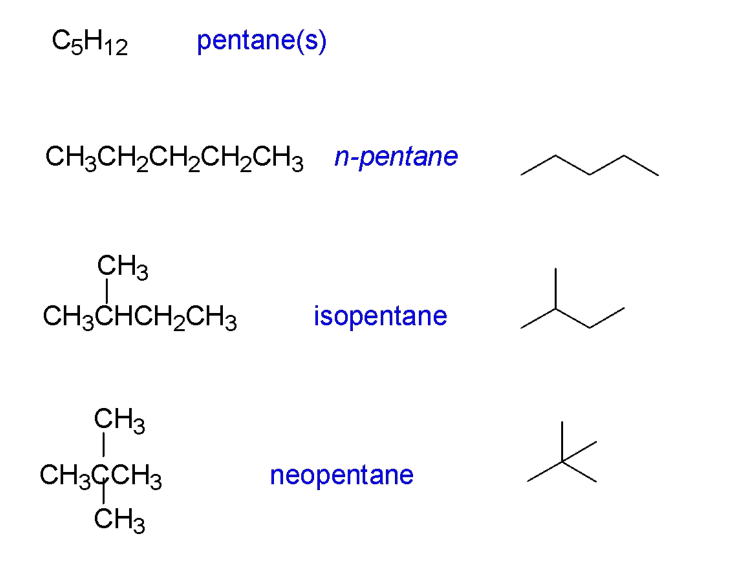


gauche

CH₃/CH₃ eclipsed

conformations about C2-C3 in *n*-butane:





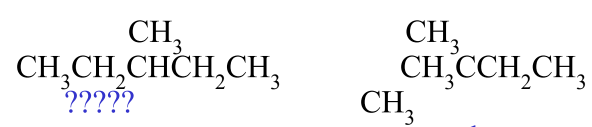
these are common, or trivial, names where a prefix is used to idicate the structure.

Alkane	name	isomers
CH ₄	methane	1
C_2H_6	ethane	1
$C_{3}H_{8}$	propane	1
C_4H_{10}	butanes	2
C_5H_{12}	pentanes	3
C_6H_{14}	hexanes	5
$C_7 H_{16}$	heptanes	9
C_8H_{18}	octanes	18
$C_{9}H_{20}$	nonanes	35
C ₁₀ H ₂₂	decanes	75
$C_{20}H_{42}$	eicosanes	366,319

each new common name requires a new prefix... hexanes C_6H_{14} common names

CH₃CH₂CH₂CH₂CH₂CH₂CH₃ *n*-hexane

CH₃ CH₃CHCH₂CH₂CH₃CH₃ isohexane



CH₃ neohexane

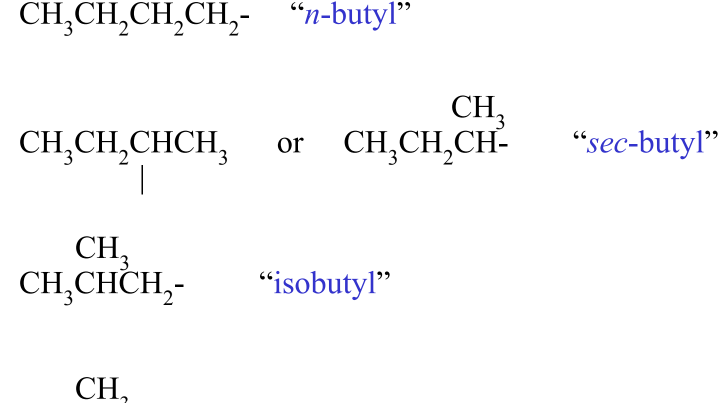
CH₃ CH₃CHCHCH₃ CH₃

IUPAC nomenclature (Geneva, 1920)

names of radicals (alkyl groups): CH₃- "methyl" CH₃Cl methyl chloride CH₃OH methyl alcohol, etc.

CH₃CH₂- "ethyl"

CH₃CH₂CH₂- "*n*-propyl" CH₃CHCH₃ "isopropyl"



CH₃ CH₃CCH₃ "*tert*-butyl"

n-propyl bromide $CH_3CH_2CH_2Br$ $BrCH_2CH_2CH_3$ CH_3 LH_2CH_2Br

isopropyl bromide

$$\begin{array}{ccc} & CH_3 & CH_3 \\ - CH_3CHCH_3 & - CH_3CHBr & - CHBr \\ Br & CH_3CHBr & - CHBr \\ - CH_3 & CH_3 \end{array}$$

n-butyl chloride

$\begin{array}{ccc} \mathsf{CH}_3\mathsf{CH}_2$

sec-butyl chloride

 $\begin{array}{c} \mathsf{CH}_3\\\mathsf{CH}_3\mathsf{CHCH}_2\mathsf{CH}_3\\\mathsf{CI}\\ \mathsf{CI}\\ \end{array} \qquad \begin{array}{c} \mathsf{CH}_3\\\mathsf{CH}_2\mathsf{CHCH}_3\\\mathsf{CI}\\ \end{array} \qquad \begin{array}{c} \mathsf{CH}_3\mathsf{CH}_2\mathsf{CHCH}_3\\\mathsf{CI}\\\mathsf{CI}\\ \end{array} \qquad \begin{array}{c} \mathsf{CH}_3\mathsf{CH}_2\mathsf{CH}_2\mathsf{CHCH}_3\\\mathsf{CI}\\\mathsf{CI}\\ \end{array} \qquad \begin{array}{c} \mathsf{CH}_3\mathsf{CH}_2\mathsf{CH}_2\mathsf{CHCH}_3\\\mathsf{CI}\\\mathsf{CI}\\\mathsf{CI}\\ \end{array} \qquad \begin{array}{c} \mathsf{CH}_3\mathsf{CH}_2\mathsf{CH}_2\mathsf{CH}_2\mathsf{CHCH}_3\\\mathsf{CI}\\\mathsf$

isobutyl alcohol

$$\begin{array}{ccc} CH_3 & CH_3 & CH_3 \\ CH_3 CHCH_2 & CH_3 CH & HO-CH_2 CHCH_3 \\ OH & CH_2 OH \end{array}$$

tert-butyl alcohol

$$\begin{array}{ccc} CH_3 & CH_3 \\ - CH_3CCH_3 & CH_3C-OH \\ OH & CH_3 \\ \end{array}$$

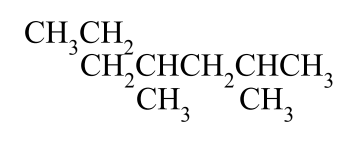
Web problems to help with naming and recognizing organic radicals:

Click here

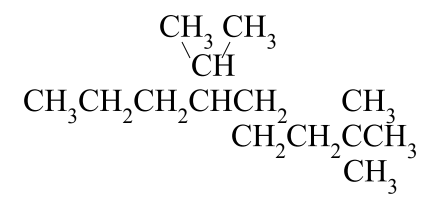
or copy and paste on the address line in your browser: http://proton.csudh.edu/structures/butyls/hwbutyls.html **IUPAC** rules for naming alkanes:

- 1. parent chain = longest continuous carbon chain \square "alkane".
- 2. branches on the parent chain are named as "alkyl" groups.
- 3. number the parent chain starting from the end that gives you the lower number for the first branch (principle of lower number).
- 4. assign "locants" to the alkyl branches.
- 5. if an alkyl group appears more than once use prefixes: di, tri, tetra, penta...; each alkyl group must have a locant!
- 6. the name is written as one word with the parent name last. The names and locants for the alkyl branches are put in alphabetic order (ignore all prefixes except iso) separating numbers from numbers with commas and letters from numbers with hyphens.

hexanes C_6H_{14} IUPAC names CH₃ CH₃CH₂CH₂CH₂CH₂CH₂CH₃ CH₃CHCH₂CH₂CH₂CH₃ (isohexane) (*n*-hexane) *n*-hexane 2-methylpentane CH₃ CH₃CCH₂CH₃ CH₃ CH₃CH₂CHCH₂CH₃ (no common name) CH₂ 3-methylpentane (neohexane) 2,2-dimethylbutane CH. CH₃CHCHCH₃ CH₃ (no common name) 2,3-dimethylbutane



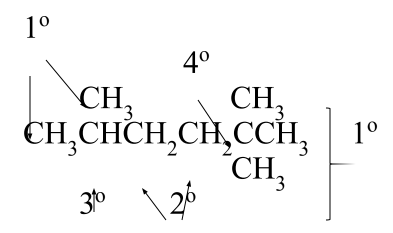
2,4-dimethylheptane



6-isopropyl-2,2-dimethylnonane

"classes of carbons"

primary carbon (1°) – a carbon bonded to one carbon secondary carbon (2°) – a carbon bonded to two carbons tertiary carbon (3°) – a carbon bonded to three carbons quaternary carbon (4°) – a carbon bonded to four carbons



classification of hydrogens, halides – hydrogens or halides are classified by the carbon to which they are attached.

1° CH₃ CH₃CHCH₂CH₂CH₂CH₃ 1° 3° 2° 2° 1°

 $CH_3CH_2CHCH_3$ sec-butyl bromide 2° bromide Br

CH₃ CH₃CCH₃ *tert*-butyl chloride 3° chloride Cl alkanes, physical properties

non-polar or only weakly polar, cannot hydrogen bond \Box relatively weak intermolecular forces

lower mp/bp; increase with size; decrease with branching

@ room temperature:

 $C_1 - C_4$ are gases $C_5 - C_{17}$ are liquids $> C_{17}$ are solids

alkanes are water insoluble

alkane	<u>mp °C</u>		<u>bp °C</u>		
methane	-183		-162		
ethane	-172		-89		
propane	-187		-42		
<i>n</i> -butane	-138		0		
<i>n</i> -pentane	-130		36		
<i>n</i> -hexane	-95	69			
• • •					
<i>n</i> -heptadecane	22		292		
<i>n</i> -octadecane	28		308		
branching lowers mp/bp					

<i>n</i> -pentane	-130	36
isopentane	-160	28

fossil fuels:

natural gas

petroleum

coal

petroleum is a complex mixture of hydrocarbons

- 1. solvents
- 2. fuels

3. raw materials for chemical syntheses

separated into fractions by fractional distillation in an oil refinery

products from fractional distillation of petroleum: fraction carbons b. range natural gas below 20° $C_1 - C_4$ $C_{5} - C_{6}$ petroleum "ether" $20 - 60^{\circ}$ $C_6 - C_7$ ligroin $60 - 100^{\circ}$ raw gasoline $40 - 205^{\circ}$ $C_5 - C_{10}$ $175 - 325^{\circ}$ $C_{12} - C_{18}$ kerosine above 275° C₁₂ & up gas oil lube oil non-volaltile liquids asphalt non-volatile solids solid carbon coke

syntheses

IndustrialLaboratorylarge amounts (tons)small amounts (grams)lowest costnon-profitmixtures often okaypure substancesdedicated apparatusflexible apparatus

on exams, homework: laboratory syntheses!

Alkanes, syntheses:

- 1. (to be covered later)
- 2. Reduction of an alkyl halide
 - a) hydrolysis of a Grignard reagent
 - b) with an active metal and an acid

3. Corey-House synthesis

(coupling of an alkyl halide with lithium dialkylcopper)

2. Reduction of an alkyl halide

i)
$$R - X + Mg \square RMgX$$
 (Grignard reagent)

- ii) RMgX + H_2O \square RH + Mg(OH)X
 - SB SA WA WB

 $CH_{3}CH_{2}CH_{2}-Br + Mg \square CH_{3}CH_{2}CH_{2}-MgBr$ *n*-propyl bromide *n*-propyl magnesium bromide

 $CH_3CH_2CH_2-MgBr + H_2O \square CH_3CH_2CH_3 + Mg(OH)Br$

propane

 $\begin{array}{ccc} CH_{3} & CH_{3} \\ CH_{3}CH-Br + Mg & \Box & CH_{3}CH-MgBr \\ isopropyl bromide & isopropyl magnesium bromide \end{array}$

 $CH_{3}CH-MgBr + H_{2}O \square CH_{3}CH_{2}CH_{3}$ propane

 $CH_{3}CH_{2}CH_{2}-MgBr + D_{2}O \square CH_{3}CH_{2}CH_{2}D$ heavy water

 $\begin{array}{c} CH_{3} \\ CH_{3}CH-MgBr + D_{2}O \quad \Box \quad CH_{3}CHD \end{array}$

b) with an active metal and an acid

 $R-X + metal/acid \square RH$ active metals = Sn, Zn, Fe, etc. acid = HCl, etc. (H⁺)

 $\begin{array}{rcl} \mathrm{CH}_{3}\mathrm{CH}_{2}\mathrm{CH}\mathrm{CH}_{3} &+& \mathrm{Sn/HCl} & \square & \mathrm{CH}_{3}\mathrm{CH}_{2}\mathrm{CH}_{2}\mathrm{CH}_{3} &+& \mathrm{SnCl}_{2} \\ & & & \\ sec \text{-butyl chloride} & & n \text{-butane} \end{array}$

 $\begin{array}{c} CH_{3} & CH_{3} \\ CH_{3}CCH_{3} + Zn/H^{+} \Box CH_{3}CHCH_{3} + ZnBr_{2} \\ Br \\ tert-butyl bromide \\ isobutane \end{array}$

3. Corey-House synthesis

$R-X + Li \square R-Li + CuI \square R_2CuLi$

$R_2CuLi + R'-X \square R-R'$ (alkane)

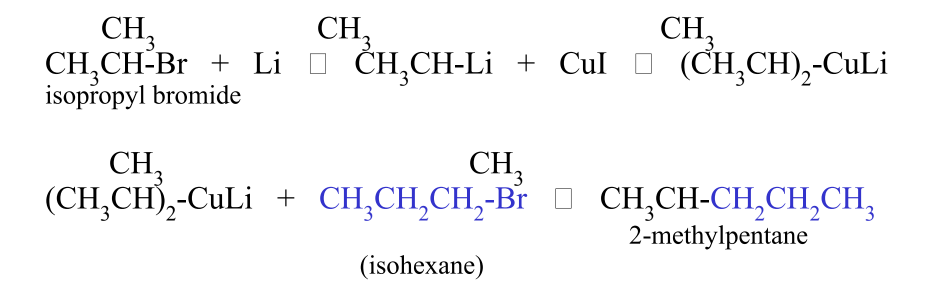
(R'-X should be 1° or methyl)

This synthesis is important because it affords a synthesis of a larger alkane from two smaller alkyl halides. note: the previous equations are not balanced:

$R-X + 2Li \square R-Li + LiX$

$2 \text{ R-Li} + \text{CuI} \square \text{ R}_2\text{CuLi} + \text{LiX}$ $R_2\text{CuLi} = \text{R-Cu}, \text{Li}^+$

 $R_2CuLi + R'X \square R-R' + RCu + LiX$



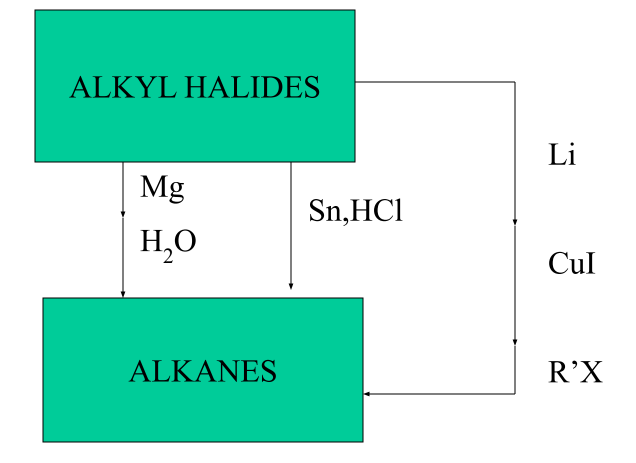
Note: the R'X should be a 1° or methyl halide for the best yields of the final product.

Alkanes, syntheses:

- 1. (to be covered later)
- 2. Reduction of an alkyl halide
 - a) hydrolysis of a Grignard reagent
 - b) with an active metal and an acid

3. Corey-House synthesis

(coupling of an alkyl halide with lithium dialkylcopper)



Reactions of alkanes:

alkane + H_2SO_4 \square no reaction (NR)

alkane + NaOH 🗆 NR

alkane + Na 🗆 NR

alkane + $KMnO_4$ \Box NR

```
alkane + H_2,Ni \Box NR
```

```
alkane + Br_2 \square NR
```

alkane + H_2O \Box NR

(Alkanes are typically non-reactive. They don't react with acids, bases, active metals, oxidizing agents, reducing agents, halogens, etc.)

Alkane, reactions:

1. Halogenation

- 2. Combustion (oxidation)
- 3. Pyrolysis (cracking)

2. Combustion

 $C_nH_{2n+2} + (xs)O_2$, flame \Box n CO₂ + (n+1) H₂O + heat gasoline, diesel, heating oil...

3. Pyrolyis (cracking)

alkane, 400-600°C \square smaller alkanes + alkenes + H₂ Used to increase the yield of gasoline from petroleum. Higher boiling fractions are "cracked" into lower boiling fractions that are added to the raw gasoline. The alkenes can be separated and used in to make plastics.

1. Halogenation

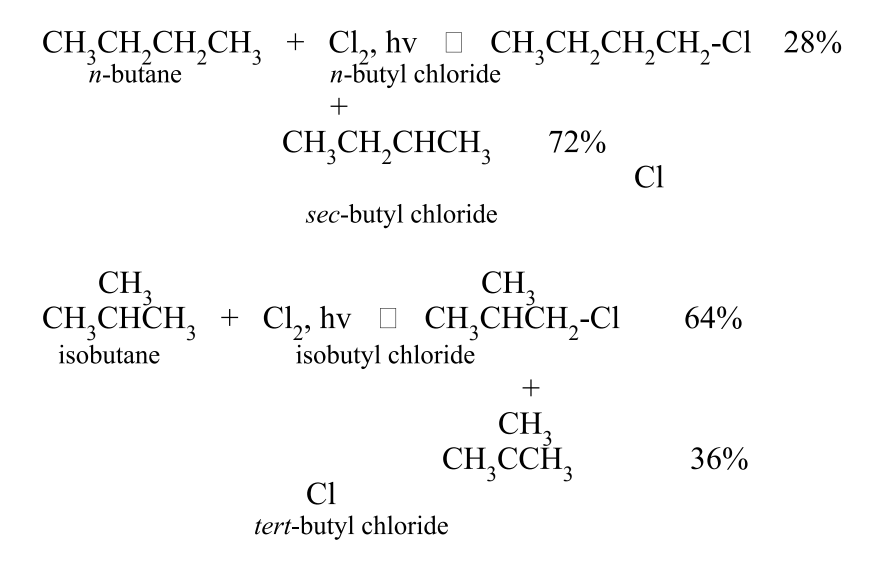
R-H + X_2 , heat or hv \Box R-X + HX

- a) heat or light required for reaction.
- b) X_2 : $Cl_2 > Br_2 \neq I_2$
- c) yields mixtures 😕
- d) H: $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_4$
- e) bromine is more selective

$CH_3CH_3 + Cl_2$, hv \Box CH_3CH_2 -Cl + HCl ethane ethyl chloride

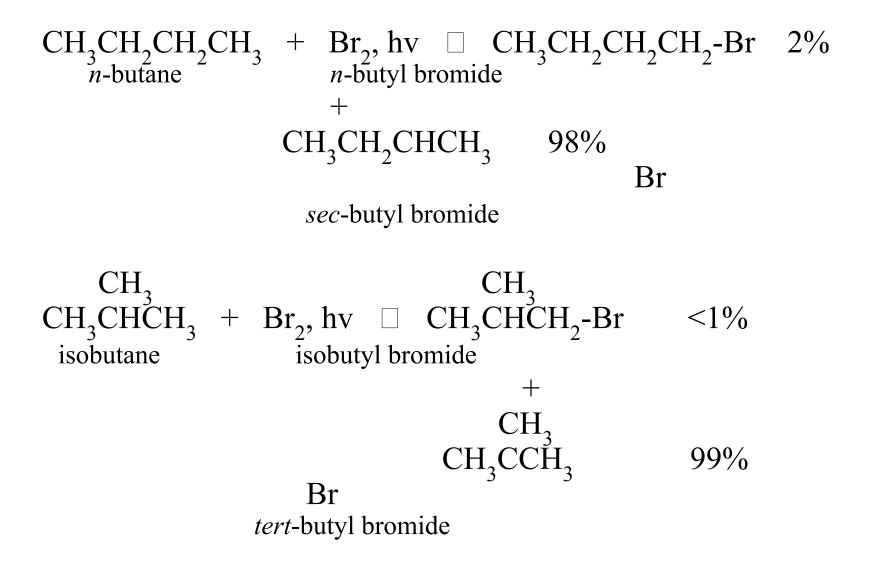
$$\begin{array}{rcl} CH_{3}CH_{2}CH_{3} + Cl_{2}, hv & \Box & CH_{3}CH_{2}CH_{2}-Cl + CH_{3}CHCH_{3} \\ propane & n-propyl chloride & Cl \\ 45\% & & & \\ 55\% \end{array}$$

gives a mixture of both the possible alkyl halides!



$CH_3CH_3 + Br_2$, hv \Box CH_3CH_2 -Br + HBr ethane ethyl bromide

$\begin{array}{rcl} CH_{3}CH_{2}CH_{3} + Br_{2}, hv & \Box & CH_{3}CH_{2}CH_{2}\text{-Br} + CH_{3}CHCH_{3} \\ propane & n\text{-propyl bromide} & Br \\ isopropyl bromide & 3\% \\ 97\% \end{array}$



In the reaction of alkanes with halogens, bromine is less reactive but more selective. Why? How? mechanism:

initiating step:

1) X—X 🗆 2 X•

propagating steps:

2) $X \bullet + R - H \Box H - X + R \bullet$

3) $R \bullet + X - X \Box R - X + X \bullet$

2), 3), 2), 3)...

terminating steps:

- $4) \quad 2 X \bullet \Box X X$
- 5) $R \bullet + X \bullet \Box R X$

6) 2 R• \Box R—R

chlorination of propane, mechanism:

- 1) Cl—Cl 🗆 2 Cl•
- 2) abstraction of 1° hydrogen:

 $Cl \cdot + CH_3CH_2CH_3 \square CH_3CH_2CH_2 \cdot + HCl$ or abstraction of 2° hydrogen:

Cl• + CH₃CH₂CH₃ \square CH₃CHCH₃ + HCl 3) CH₃CH₂CH₂• + Cl₂ \square CH₃CH₂CH₂Cl + Cl• or CH₃CHCH₃ + Cl₂ \square CH₃CHCH₃ + Cl• •

plus terminating steps

2) abstraction of 1° hydrogen:

 $Cl \cdot + CH_3CH_2CH_3 \square CH_3CH_2CH_2 \cdot + HCl$ or abstraction of 2° hydrogen:

$$Cl + CH_3CH_2CH_3 \square CH_3CHCH_3 + HCl$$

The chloride that is produced depends on which hydrogen is abstracted by the chlorine free radical in step 2. The *n*-propyl free radical gives the *n*-propyl chloride while the isopropyl free radical yields the isopropyl chloride.

The relative reactivity in chlorination:

H: $3^{\circ}: 2^{\circ}: 1^{\circ} = 5.0: 3.8: 1.0$

The number of hydrogens (probability factor) may also be important.

 $CH_3CH_2CH_2CH_3 + Cl_2$, hv \Box $CH_3CH_2CH_2CH_2-Cl_2$ *n*-butane

C1

n-butyl chloride = (# of 1° hydrogens) x (reactivity of 1°) = 6 x 1.0 = 6.0 sec-butyl chloride = (# of 2° hydrogens) x (reactivity of 2°) = 4 x 3.8 = 15.2 % *n*-butyl chloride = 6.0/(6.0 + 15.2) x 100% = 28% % sec-butyl chloride = 15.2/(6.0 + 15.2) x 100% = 72%

$$\begin{array}{c} CH_{3} \\ CH_{3}CHCH_{3} + Cl_{2}, hv \Box CH_{3}CHCH_{2}-Cl + CH_{3}CCH_{3}\\ isobutane \end{array}$$

isobutyl chloride = (# of 1° H's) x (reactivity of 1°) = 9 x 1.0 = 9.0

tert-butyl chloride = (# of 3° H's) x (reactivity of 3°) = 1 x 5.0 = 5.0

% isobutyl = $(9.0/(9.0 + 5.0)) \times 100\% = 64\%$

In this case the probability factor outweighs the difference in relative reactivity of 1° and 3° hydrogens.

Relative reactivity in bromination:

 $3^\circ: 2^\circ: 1^\circ = 1600: 82: 1$

In bromination the relative reactivity differences are much greater than any probability differences.

isobutane + Br_2 , hv \Box isobutyl bromide + *tert*-butyl bromide

isobutyl bromide = 9 H x 1 = 9 *tert*-butyl bromide = 1 H x 1600 = 1600 % *tert*-butyl bromide = (1600/1601) x 100% = >99%

Why is relative reactivity of H: $3^{\circ} > 2^{\circ} > 1^{\circ}$? CH_3 —H \Box $CH_3 \bullet$ + $H \bullet$ $\Delta H = 104$ Kcal/mole CH_3CH_2 — $H \square CH_3CH_2 \bullet + H \bullet \Delta H = 98 K cal/mole$ 1° free radical $CH_3CH_2CH_3 \square CH_3CH_2CH_2 + H \Delta H = 98 \text{ Kcal/mole}$ 1° free radical $CH_3CHCH_3 + H \bullet \Delta H = 95$ Kcal/mole 2° free radical $CH_{3}CHCH_{3} \Box CH_{3}CCH_{3} + H \bullet \Delta H = 92 \text{ Kcal/mole}$

3° free radical

Relative reactivity in halogenation:Stability of free radicals:Ease of formation of free radicals:Ease of abstraction of H's:

 $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_4$

1. Halogenation

R-H + X_2 , heat or hv \Box R-X + HX

- a) heat or light required for reaction.
- b) X_2 : $Cl_2 > Br_2 \neq I_2$
- c) yields mixtures 😕
- d) H: $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_4$
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