## Naming Alkanes and Alkyl Groups

Table 2.1 Nomenclature and Physical Properties of Straight-Chain Alkanes					
Number of carbons	Molecular formula	Name	Condensed structure	(linear alka	
1	$CH_4$	methane	$CH_4$	are also cal <i>n</i> -alkanes,	
2	$C_2H_6$	ethane	CH <sub>3</sub> CH <sub>3</sub>		
3	$C_3H_8$	propane	CH <sub>3</sub> CH <sub>2</sub> CH <sub>3</sub>		
4	$C_{4}H_{10}$	butane	CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub>	"normol")	
5	$C_{5}H_{12}$	pentane	CH <sub>3</sub> (CH <sub>2</sub> ) <sub>3</sub> CH <sub>3</sub>	normar )	
6	$C_{6}H_{14}$	hexane	CH <sub>3</sub> (CH <sub>2</sub> ) <sub>4</sub> CH <sub>3</sub>		
7	C7H16	heptane	CH <sub>3</sub> (CH <sub>2</sub> ) <sub>5</sub> CH <sub>3</sub>		
8	C8H18	octane	CH <sub>3</sub> (CH <sub>2</sub> ) <sub>6</sub> CH <sub>3</sub>		
:	:	:	:		

Names of alkyl fragments derive from alkanes:



## **Naming Functionalized Alkanes**

CH <sub>4</sub> methane	CH <sub>3</sub> (or H <sub>3</sub> C ) methyl	—OH alcohol	CH <sub>3</sub> —OH methyl alcohol (methanol)
CH <sub>3</sub> CH <sub>3</sub> ethane	CH <sub>3</sub> CH <sub>2</sub> ethyl		$CH_3CH_2 - O \xrightarrow{O}_{acetic \ acid} O CH_3$
CH <sub>3</sub> CH <sub>2</sub> CH <sub>3</sub> propane	CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> <i>n-propyl</i> <i>iso-propyl</i> CH <sup>-</sup> CH <sup>-</sup>	esterNH <sub>2</sub>	ethyl ester (ethyl acetate) CH <sub>3</sub> CH <sup>-</sup> NH <sub>2</sub> CH <sub>2</sub> <sup>isopropyl</sup>
CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub> butane	$ \begin{array}{c} CH_3CH_2CH_2CH_2CH_2 \\ & & CH_3 \\ & CH_3 \\ & & CH_3 \\ & CH_3 \\ & CH_3 \\ & CH_2 \\ & CH_3 \\ & & CH_2 \\ & & CH_3 \end{array} \\ \end{array} $	ether	$CH_3$ $H_3C = O = C = CH_3$ $M_3C = O = C = CH_3$ $Methyl tert-butyl = CH_3$ ether (MTBE)





# **Rotation About Single Bond in Ethane**



### **Rotation About Single Bond in Ethane**



*Torsional* energy of 2.9 kcal/mol must be overcome to rotate 120°. Timescale of that happening: once every 100 femtoseconds (at room temp).

## **Rotation About Single Bond in Ethane**



### **Rotation About Single Bond in Ethane**







(Simulation performed by molecular dynamics, using standard rotational and vibrational potentials.)





## **Rotation About Single Bond in Butane**



#### Points to note:

Gauche is not that much less stable than anti; so, butane spends plenty of time in gauche conformation.

Barrier to rotation of end methyl groups (3.3 kcal/mol) is less than that of internal C-C bond (5 kcal/mol).



### Conformational Preferences Extend to Higher Alkanes

For linear alkanes, the all-anti conformation is preferred...



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For linear alkanes, the all-anti conformation is preferred...



...but not by much. Gauche conformers frequently observed.