

8.10: Unsaturated Hydrocarbons

Two important families of hydrocarbons which are not found in petroleum are the **alkenes** (also called olefins) and the **alkynes** (also called acetylene). As can already be seen, there are often two names for simple organic compounds. Since 1930 the International Union of Pure and Applied Chemistry (IUPAC) has developed a systematic method for naming all organic compounds, but many of the earlier names still survive, particularly among industrial chemists. Where appropriate, both names will be given, the older name in parentheses. Alkene molecules are similar to alkane molecules, except that they contain a carbon-carbon double bond ($C=C$) and two fewer H atoms. They thus have the general formula C_nH_{2n} . Alkyne molecules contain triple bonds ($C\equiv C$) and have four H atoms less than the corresponding alkane. Their general formula is thus C_nH_{2n-2} . Compounds containing double or triple bonds are often referred to collectively as **unsaturated compounds**. Because of their multiple bonds, alkenes and alkynes are usually more chemically reactive than alkanes and aromatic hydrocarbons.

The presence of a double or triple bond in the molecule opens up many more possibilities for isomerism than is the case for alkanes. There are usually several alternative locations for the multiple bond, and in the case of a double bond there is the possibility of cis-trans isomerism. Thus while there are only *two* alkane molecules possible with *four* carbon atoms, four alkene molecules are possible:

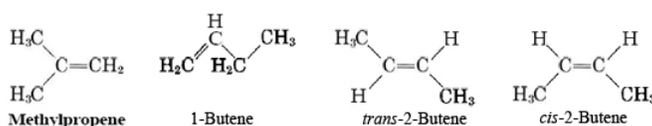


Figure 1) joining the two atoms together. In making such a model, it is necessary to bend the springs a fair amount in order to fit them into the appropriate holes in the balls. The ability to bend or stretch is characteristic of all chemical bonds—not just those between doubly bonded atoms. Thus each atom can vibrate about its most stable position. Perhaps ball-and-spring models would be more appropriate than ball-and-stick models in all cases.

The bent-bond picture makes it easy to explain several characteristics of double bonds. As noted in [Chemical Bonding - Electron Pairs and Octets](#), the distance between two atomic nuclei connected by a double bond is shorter than if they were connected by a single bond. In the case of carbon-carbon bonds, for example, the $C=C$ distance is 133 pm, while the $C-C$ distance is 156 pm. This makes sense when we realize that each bent bond extends along a curved path. The distance between the ends of such a path (the C nuclei) is necessarily shorter than the path itself.

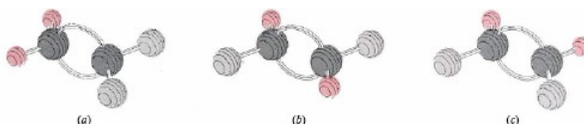


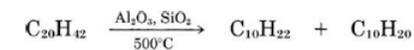
Figure 8.10.2 Ball-and-stick models for the three isomeric forms of $C_2H_2F_2$. Carbon atoms are dark gray, hydrogen atoms are light red, and fluorine atoms are light gray. Note that all three structures are different, although the molecular formula is always $C_2H_2F_2$. Structures (b) and (c) differ only because of the barrier to rotation around the carbon-carbon double bond.

Another characteristic of double bonds is that they make it difficult to twist one end of a molecule relative to the other. This phenomenon usually is called a **barrier to rotation**. Such a barrier accounts for the fact that it is possible to prepare *three* different compounds with the formula $C_2H_2F_2$. Their structures are shown in Figure 2. Structure (a) is unique because both F atoms are attached to the same C atom, but (b) and (c) differ only by a 180° flip of the right-hand $=CHF$ group. If there were no barrier to rotation around the double bond, structures (b) and (c) could interconvert very rapidly whenever they collided with other molecules. It would then be impossible to prepare a sample containing only type (b) molecules or only type (c) molecules.

Since they have the same molecular formula, (a), (b) and (c) are isomers. Structure (b) in which the two F atoms are on *opposite* sides of the double bond is called the **trans** isomer, while structure (c) in which two like atoms are on the *same* side is called the **cis** isomer. It is easy to explain why there is a barrier to rotation preventing the interconversion of these cis and trans isomers in terms of our bent-bond model. Rotation of one part of the molecule about the line through the C atoms will cause one of the bent-bond electron clouds to twist around the other. Unless one-half of the double bond breaks, it is impossible to twist the molecule through a very large angle.

Although alkenes are not present in crude petroleum, they are produced in large quantities in petroleum *refining*. Many of the hydrocarbons in crude oil have very long chains and are solids or thick syrupy liquids at ordinary temperatures because of their relatively large intermolecular forces. The most important petroleum product, gasoline, requires molecules containing from 6 to 12 carbon atoms. These can be obtained by heating the longer-chain compounds. These big molecules writhe around so fast at higher

temperatures that they “crack” or break into smaller fragments. Usually a catalyst is added to speed up the reaction, which is called **catalytic cracking**. When cracking occurs, one alkene and one alkane molecule are produced:



An alkane

An alkane

An alkene

The most important alkenes from an industrial point of view are the two simplest: ethene (ethylene), $\text{H}_2\text{C}=\text{CH}_2$ and propene (propylene), $\text{CH}_3\text{CH}=\text{CH}_2$. Almost 1.11×10^{11} kg of ethene and 6.9×10^{10} kg of propene were consumed worldwide in 2006 ^[1]. Both are used in the manufacture of **plastics**. They are also raw materials for production of detergents, antifreeze, elastics, and lubricating oils.

Among the alkynes only the two simplest are of any industrial importance. Both ethyne (acetylene), $\text{CH}\equiv\text{CH}$ and propyne (methyl acetylene), $\text{CH}_3\text{C}\equiv\text{CH}$, are used in welding and steel cutting where they are burned together with pure oxygen gas in an oxyacetylene torch. As supplies of petroleum dwindle, though, acetylene may become more important as a starting material for the manufacture of other chemicals, since it can be made from coal.

1. Thomasson, A. "CMAI Announces Completion of Olefins World Analyses; World Light Olefins Analysis and World Butadiene Analysis." Chemical Market Associates, Inc. 5 December 2006. www.cmaiglobal.com/Marketing/...WLOA_WBA07.pdf

This page titled [8.10: Unsaturated Hydrocarbons](#) is shared under a [CC BY-NC-SA 4.0](#) license and was authored, remixed, and/or curated by [Ed Vitz, John W. Moore, Justin Shorb, Xavier Prat-Resina, Tim Wendorff, & Adam Hahn](#).