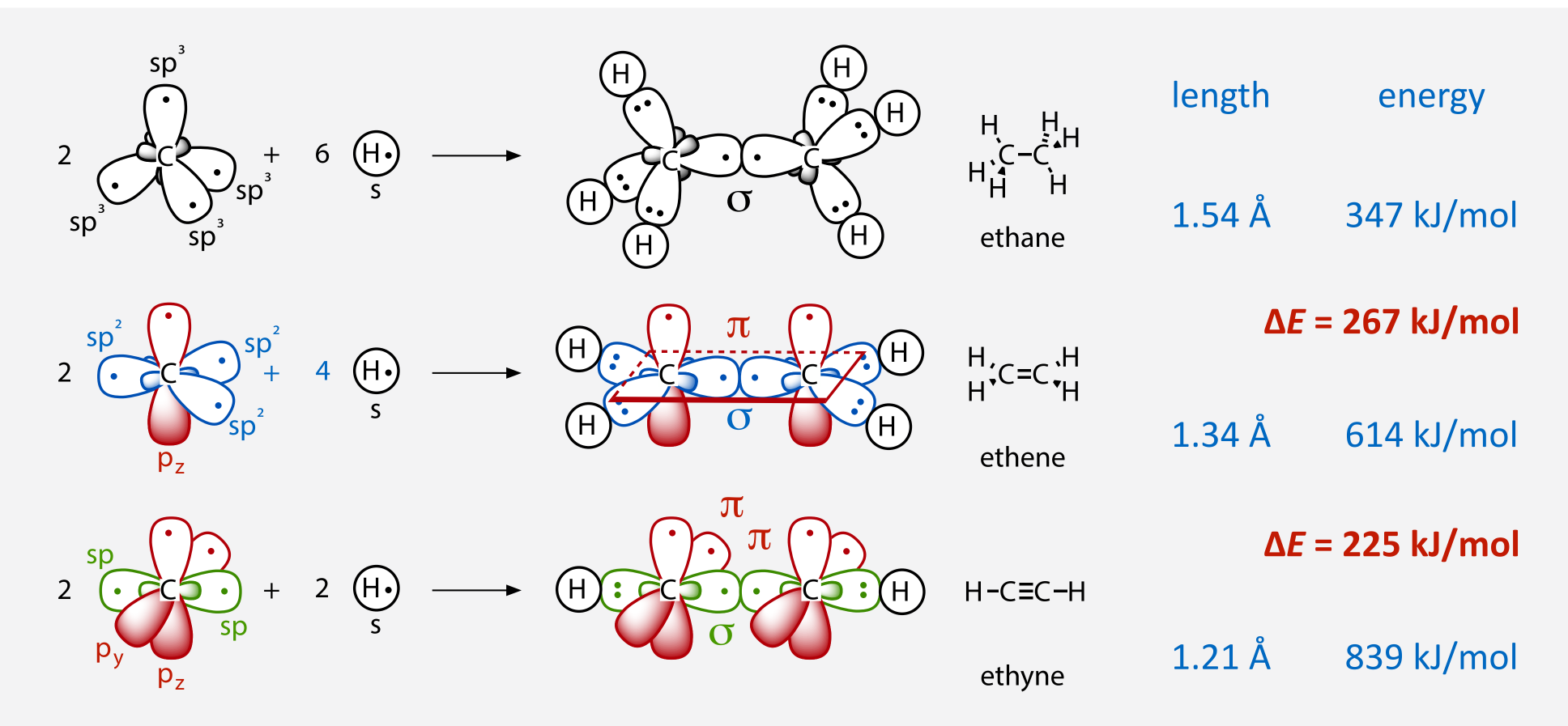
# 2.3 Formation of Multiple Bonds

# **Reading Recommendations**

- Clayden, Greeves, Warren, *Organic Chemistry*, Oxford University Press, 2<sup>nd</sup> ed., **2012**, pp 99–106.
- Jamart, Bodiguel, Brosse, Les cours de Paul Arnaud Cours de chimie organique, Dunod, 19<sup>th</sup> ed.,
  2015, pp 79–86.

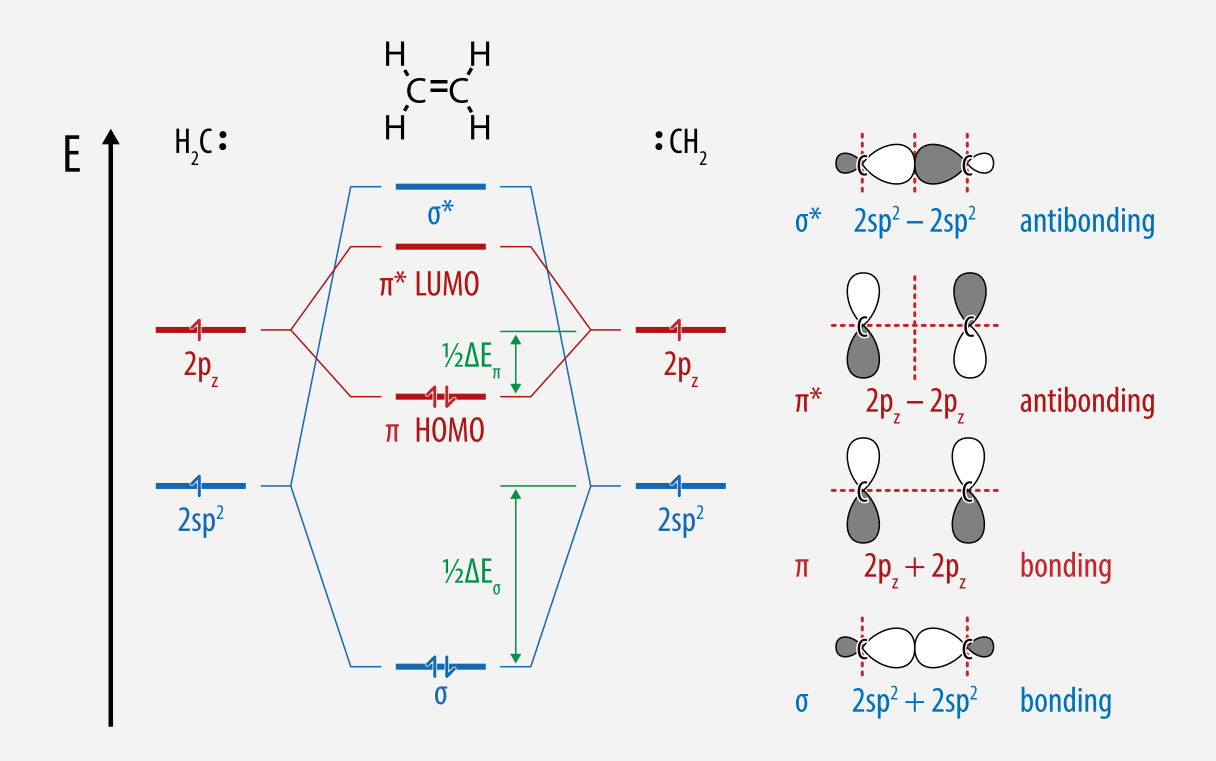
### Formation of Carbon-Carbon Multiple Bonds



- double or triple bonds are one  $\sigma$  bond plus one or two  $\pi$ -bonds, respectively
- $\bullet$   $\pi$  bonds between residual p orbitals, node plane along bond, no rotational symmetry
- $\bullet$  rotation around  $\pi$ -bonds requires breaking them, energetically too costly at r. t.

#### Molecular Orbital View of the Carbon-Carbon Double Bond

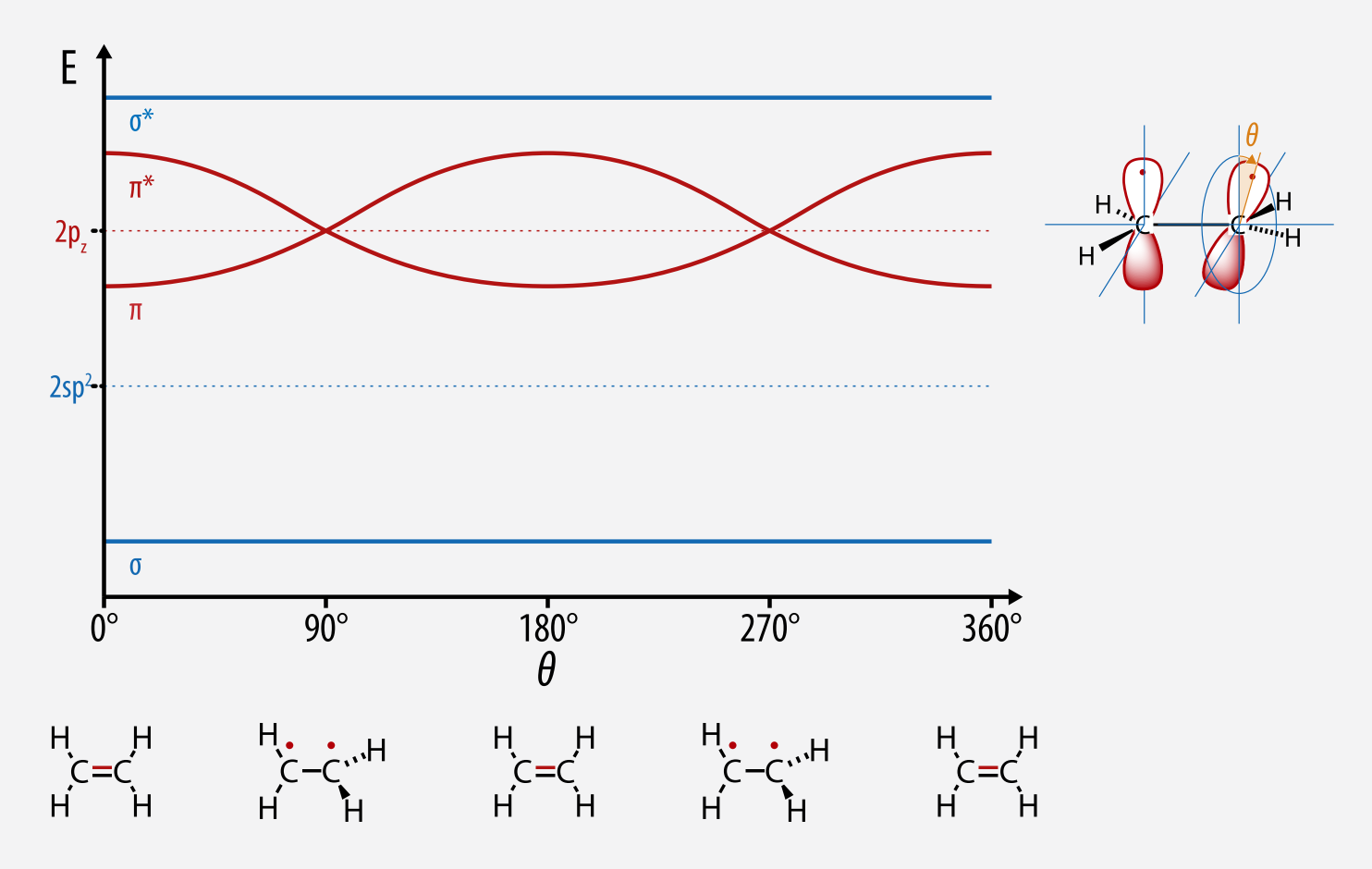
• simplified and schematic molecular orbital energy diagram of the ethene molecule



- only orbitals of matching symmetry & orientation interact, sp<sup>2</sup> with sp<sup>2</sup>, and p<sub>z</sub> with p<sub>z</sub>
- distinct  $\sigma$  bond (from two sp<sup>2</sup>) and  $\pi$  bond (from two p<sub>z</sub>) with different energy, symmetry
- chemistry ruled by highest occupied, lowest unoccupied molecular orbitals (HOMO, LUMO)
- typically  $\pi$  HOMO and  $\pi^*$  LUMO, located between  $\sigma$  and  $\sigma^*$  because  $p_z$  overlap much smaller

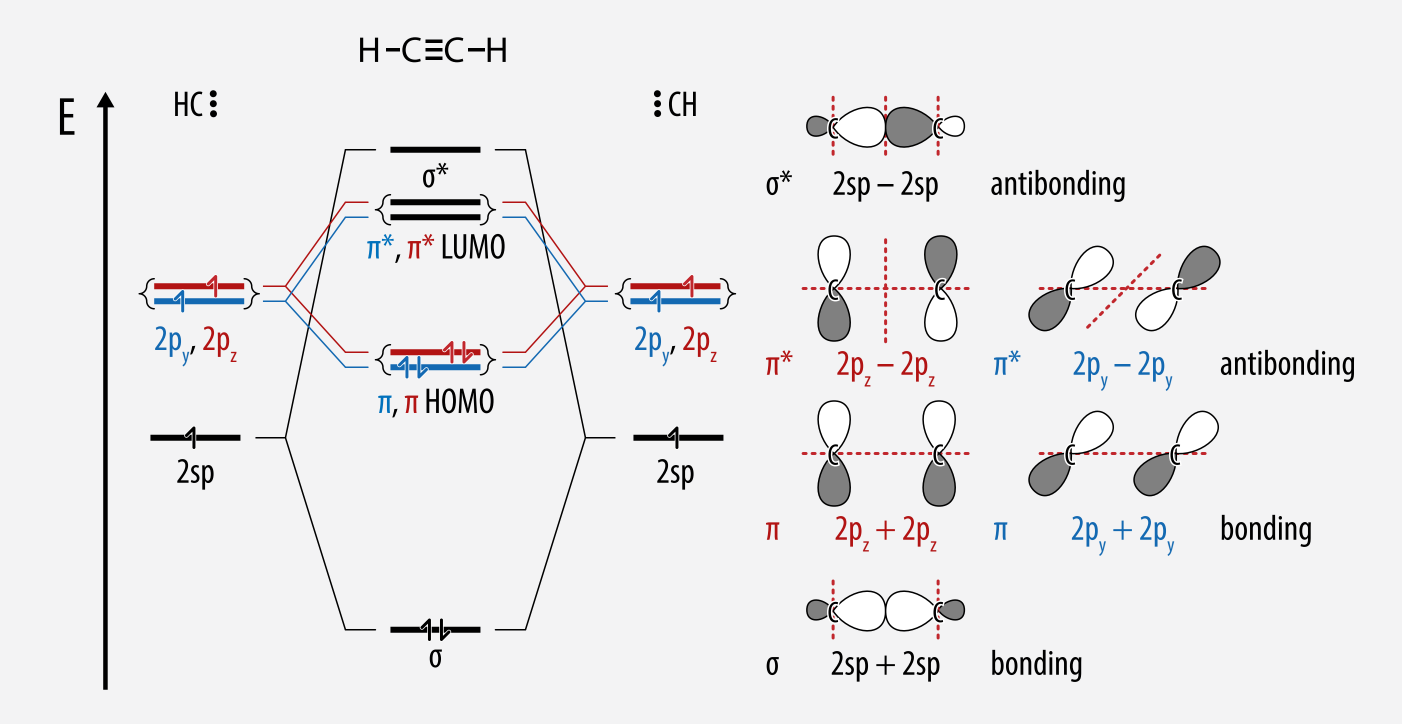
#### **Rotation Around a Double Bond**

 $\bullet$   $\sigma$  bond has rotational symmetry relative to carbon carbon bond axis, but  $\pi$  bond does not



• rotating  $\pi$  orbitals by 90° requires breaking the  $\pi$  bond ( $\approx$ 260 kJ/mol), disfavorable at r.t.

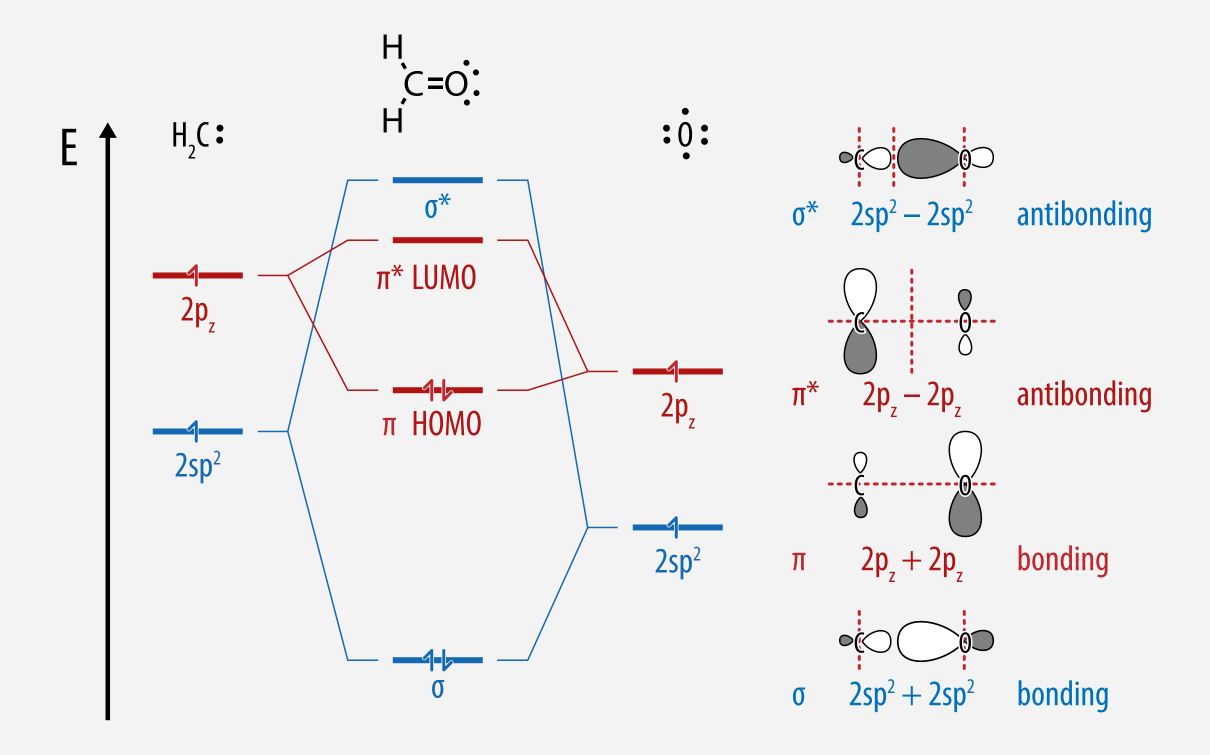
• simplified and schematic molecular orbital energy diagram of the ethyne molecule



- only orbitals of matching symmetry & orientation interact (sp with sp, pz with pz, py with py)
- distinct  $\sigma$ -bond (from two sp<sup>2</sup>) and two independent  $\pi$ -bonds (each from two  $p_z$  and two  $p_y$ )
- the two  $\pi$ -bonds are identical in energy, symmetry, just different in orientation
- chemistry dominated by  $\pi$  HOMO and  $\pi^*$  LUMO , reactivity of ethyne similar to ethene

# Molecular Orbital View of the Carbon-Oxygen Double Bond in the Carbonyl Group

• simplified and schematic molecular orbital energy diagram of the carbon-oxygen double bond



- $\sigma$  bond from sp<sup>2</sup>(C) with sp<sup>2</sup>(O),  $\pi$ -bond from p<sub>z</sub>(C) with p<sub>z</sub>(O) with different bond energies
- oxygen orbitals lower in energy due to electronegativity difference of carbon and oxygen
- $\bullet$  bonding  $\sigma$  orbital and, in particular, bonding  $\pi$  orbital "look like" oxygen orbitals
- $\bullet$  both the  $\sigma$  bond and, in particular, the more polarizable  $\pi$  bond polarized towards oxygen

## **Learning Outcomes**

- multiple bonds can be described as distinct  $\sigma$  bond plus one or two  $\pi$  bonds
  - σ bond has rotational symmetry with respect to carbon-carbon bond axis
  - $\bullet$   $\pi$  bond does not have rotational symmetry; rotation requires breaking it
- chemistry & physics dominated by frontier orbitals:  $\pi$  HOMO and  $\pi^*$  LUMO