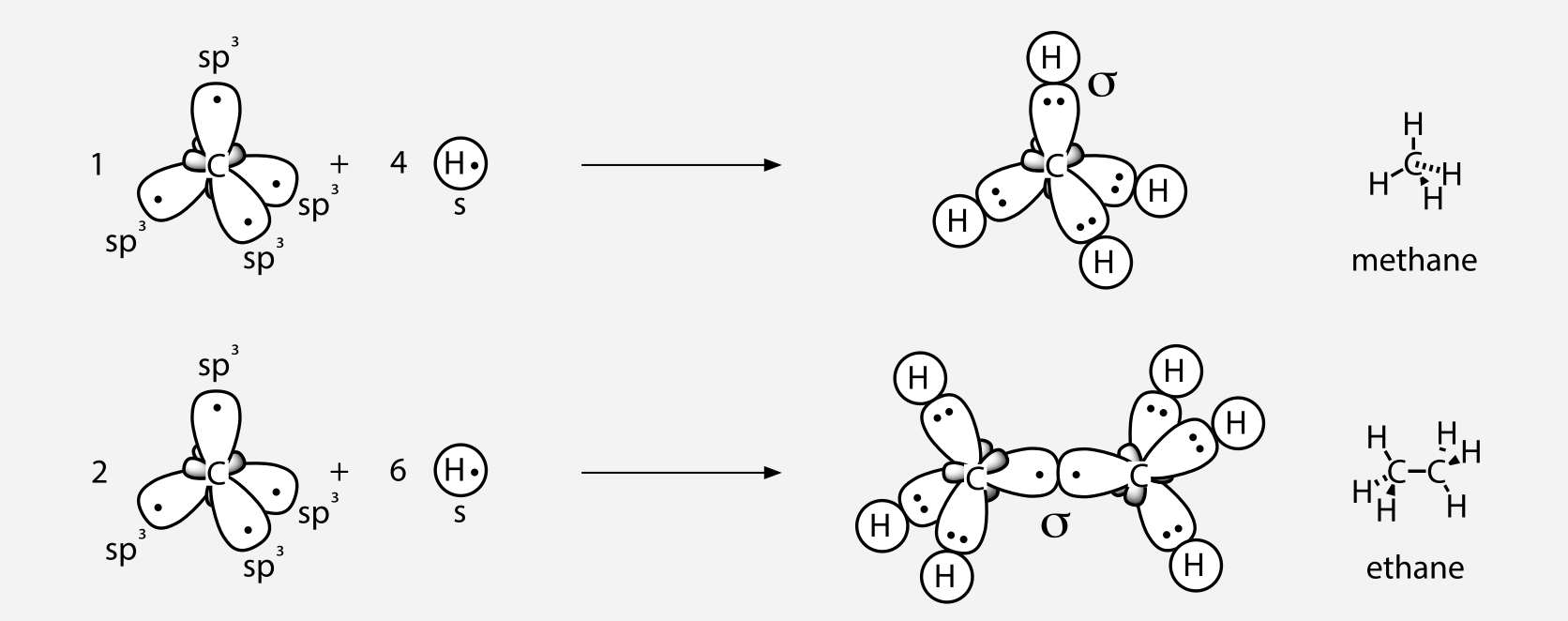
# 2.2 Formation of Single Bonds

## **Reading Recommendations**

- Clayden, Greeves, Warren, *Organic Chemistry*, Oxford University Press, 2<sup>nd</sup> ed., **2012**, pp 88–106.
- Jamart, Bodiguel, Brosse, Les cours de Paul Arnaud Cours de chimie organique, Dunod, 19<sup>th</sup> ed.,
  2015, pp 79–86.

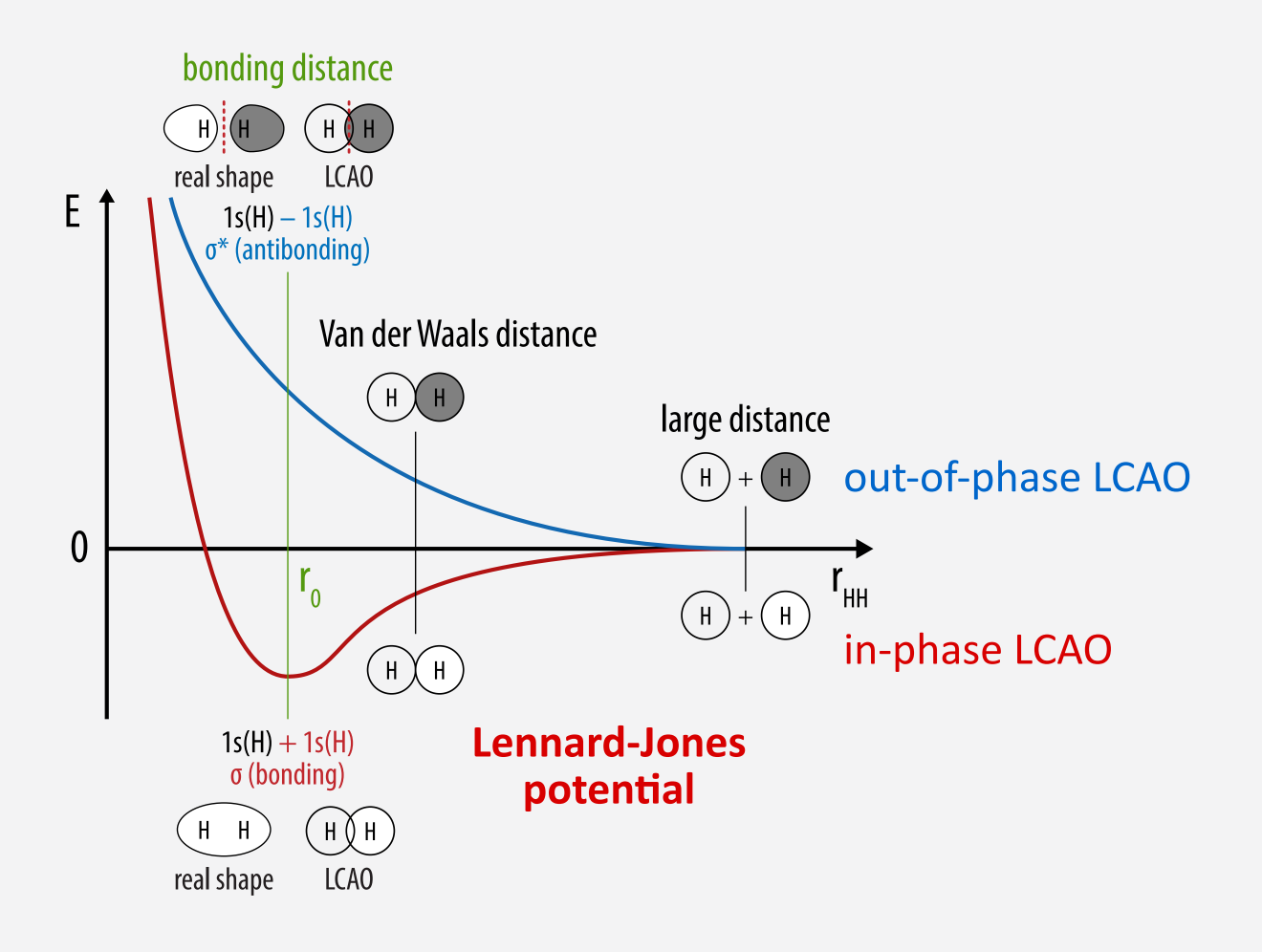
## Formation of Single Bonds (Valence Bond Model)

• according to valence bond model, covalent bonds form by pairing electrons of atomic orbitals



- in-phase combination of atomic or hybrid orbitals
- single bonds are  $\sigma$ -bonds (rotational symmetry) between sp3, sp2, sp, or s orbitals
- $\bullet$  due to rotational symmetry of the  $\sigma$ -orbital, rotation is free without breaking the bond

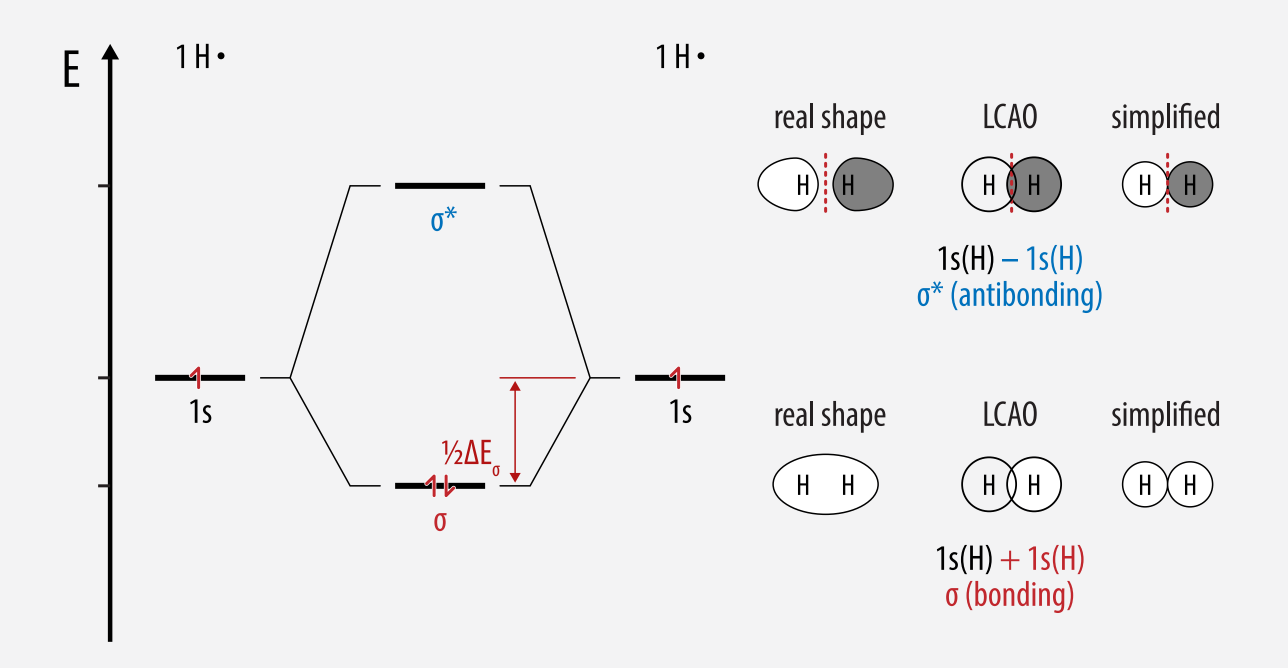
## Interaction of Atomic Orbitals upon Approach



- upon decreasing the interatomic distance, atomic orbitals interact and split energetically
- number of orbitals conserved, LCAO with one "in phase" and one "out of phase"
- energy in fact not quite conserved, overall raise of both levels due to electrostatic repulsion

## **Molecular Orbital Energy Diagrams**

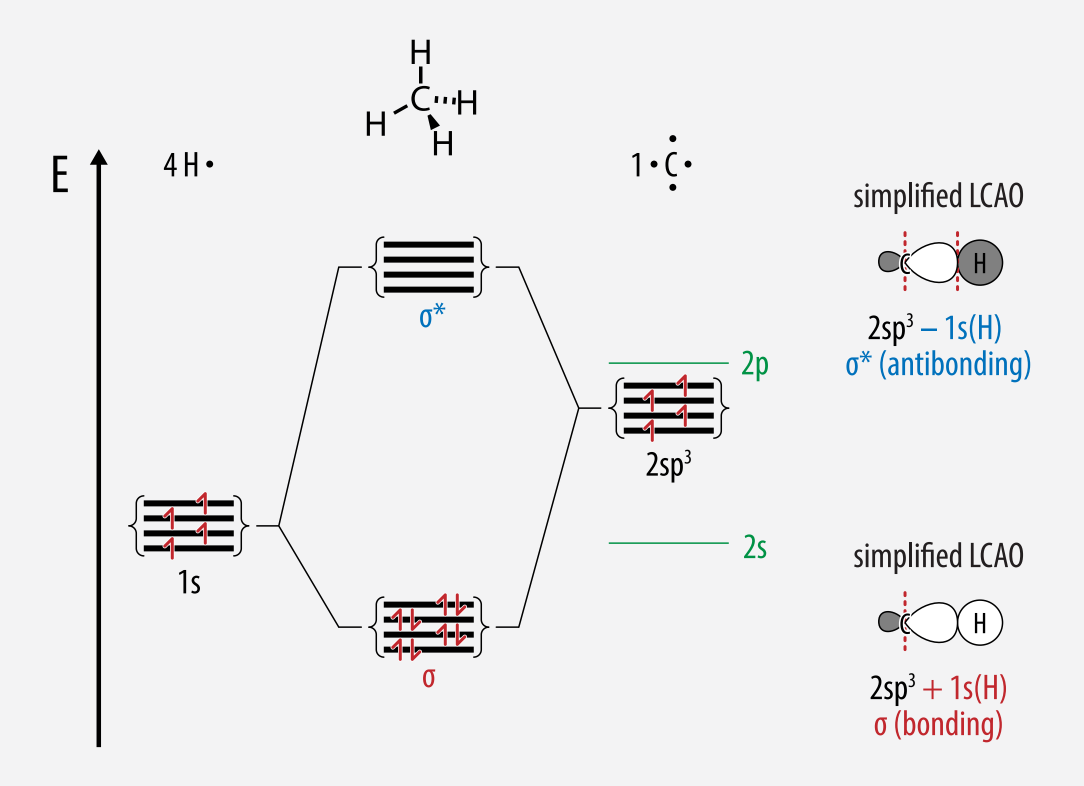
• schematic molecular orbital energy diagram for a symmetric diatomic molecule (such as H<sub>2</sub>)



- energy splitting increases with atomic orbital overlap
- number of orbitals preserved but sum of orbital energies (electron density) increases
- $\bullet$  bond energy is stabilization of filled bonding orbital  $\sigma$  (due to electron delocalization)
- $\bullet$  antibonding orbital  $\sigma^*$  is energetically destabilized but remains empty

### Molecular Orbital View of the Covalent Bond in Multiatom Molecules

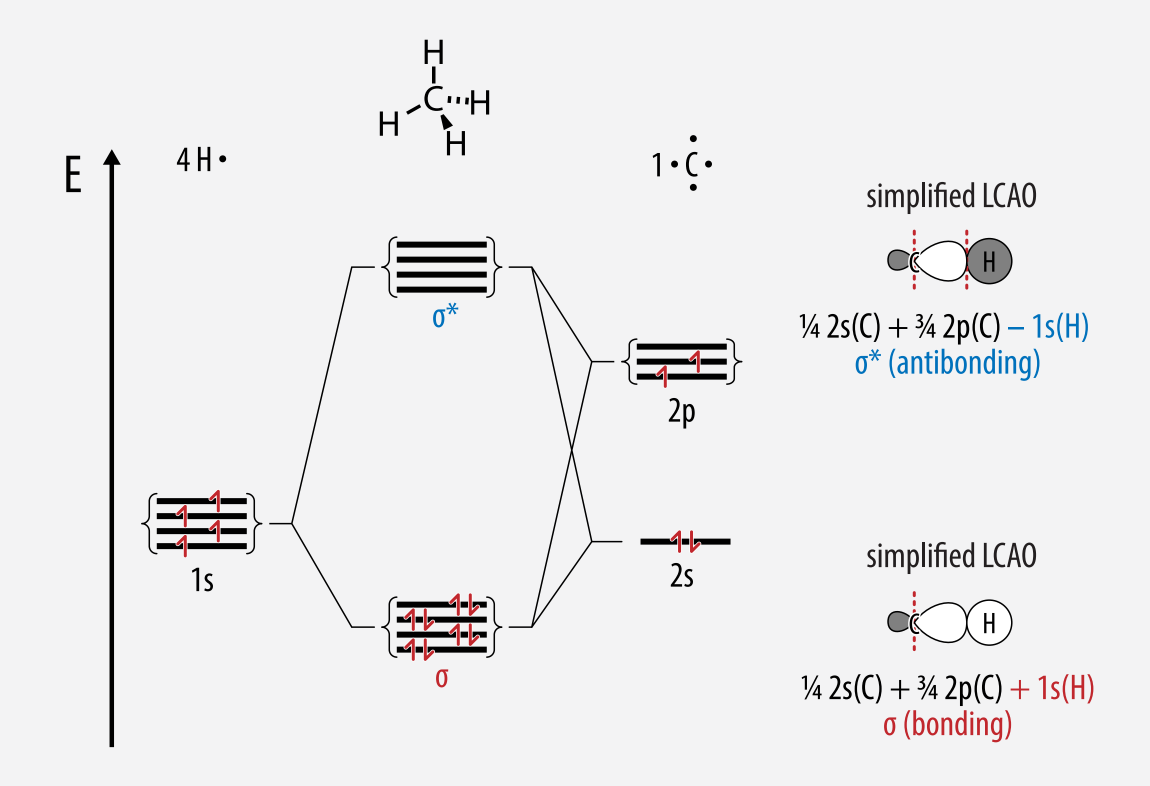
• simplified and schematic molecular orbital energy diagram of the methane molecule



- linear combination of atomic orbitals may start from LCAO hybrid orbitals
- interactions between orbitals or orbital sets of matching symmetry

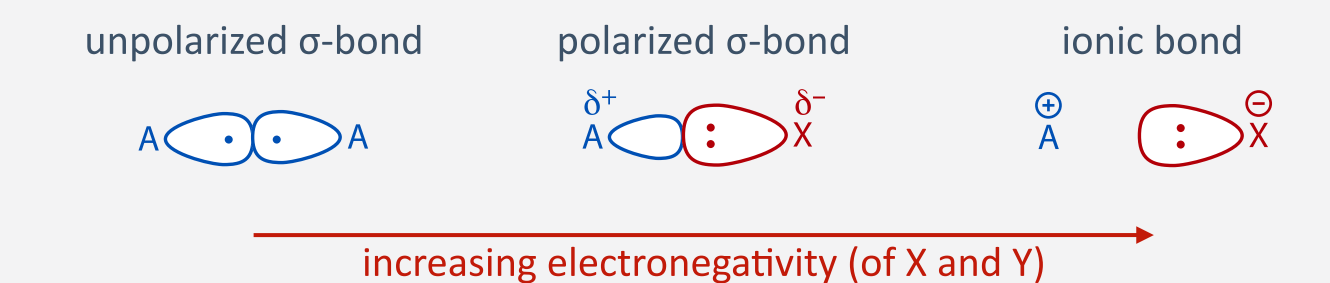
#### Molecular Orbital View of the Covalent Bond in Multiatom Molecules

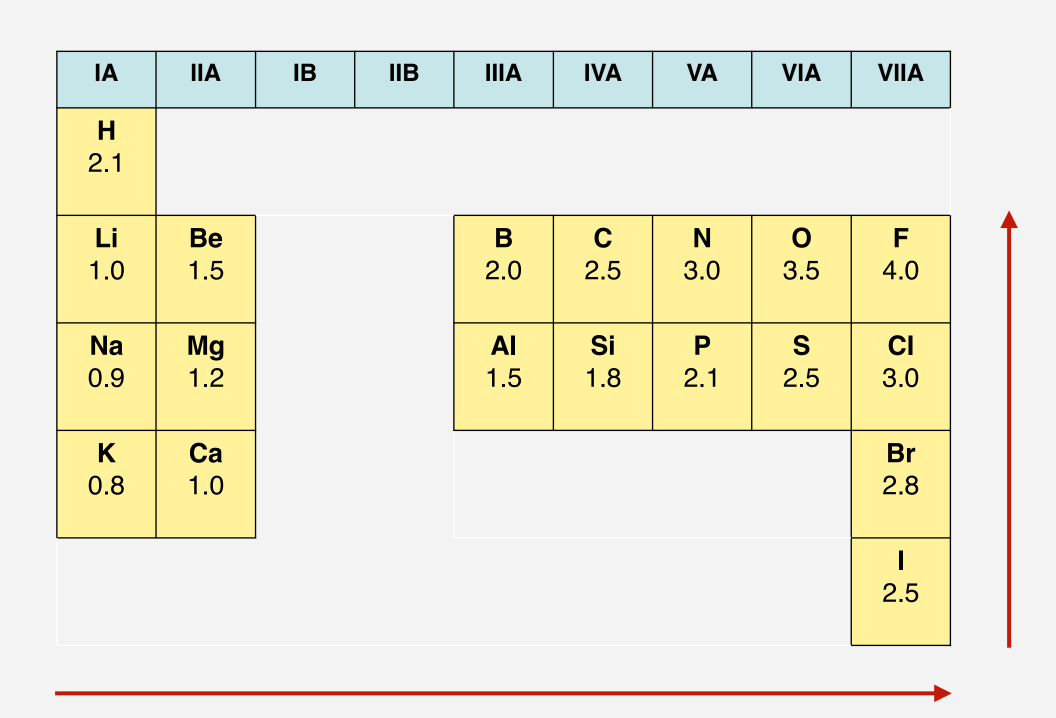
• simplified and schematic molecular orbital energy diagram of the methane molecule



- a more correct approach would start from atomic orbitals instead of hybrid orbitals
- result will be (almost) the same due to "mixing" of orbitals
- VSEPR model and hybridization are useful and valid simplifications

#### **Polarization of Covalent Bonds**

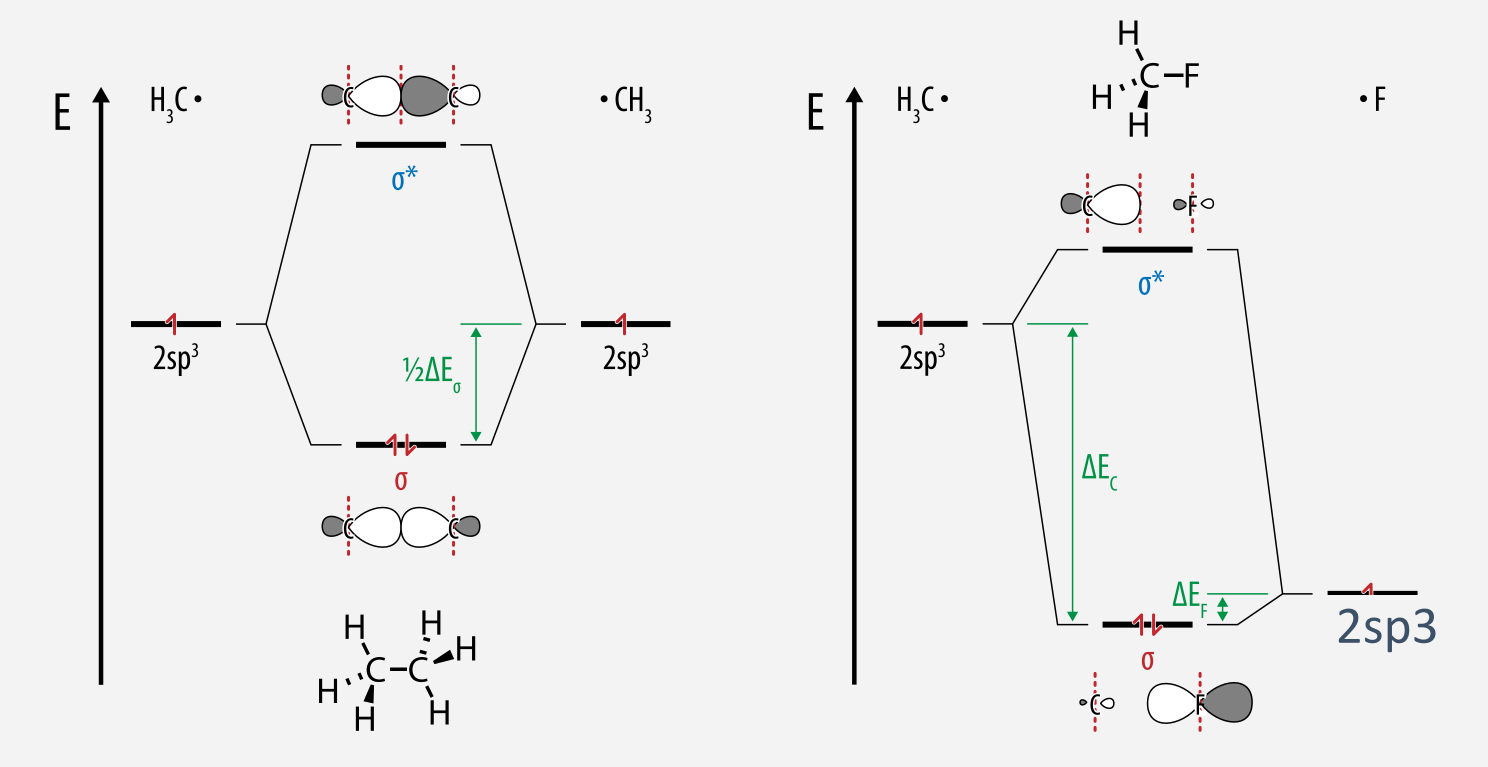




#### negative inductive Effect (-I effect)

## **Molecular Orbital Representation of Polarized Bonds**

• simplified and schematic molecular orbital diagrams of the C–C and the C–F bond



- atomic orbitals of more electronegative atoms lower in energy (higher electron affinity)
- increasing energy difference between bonding partners implies less electronic interaction
- bonding MO closer in energy to, "look more like" AO from more electronegative element
- antibonding MO closer in energy to, "look more like" AO from less electronegative element

## **Learning Outcomes**

- covalent bond can be described by linear combination of atomic orbitals
- interactions only between orbitals or orbital sets of matching symmetry
- number of orbitals preserved but sum of all orbital energies increases
- bond energy is stabilization of bonding orbital  $\sigma$  (electron delocalization)
- $\bullet$  antibonding orbital  $\sigma^*$  energetically destabilized but remains empty