



Atomic and molecular interactions; applications

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24 October 2013

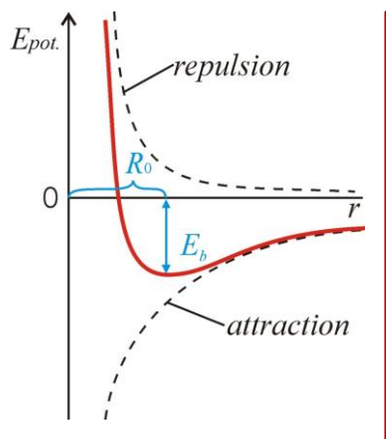
Atomic interactions

Noble gases: found in their atomic form in nature (*He, Ne, Ar, Kr, Xe, Rn*)

Other elements: form molecules held together by chemical bonds. (*e.g. H₂, HCl, H₂O, ...*) Each atom has a (more or less) fixed position in the molecule.

Atomic interactions

Bringing two atoms (ions/molecules/...) close to each other attractive and repulsive forces emerge between them resulting in a net potential energy function with an energy minimum at a certain distance:



$$E_{pot} = E_{attraction} + E_{repulsion}$$

E_{pot} : potential energy of the system
 $E_{attraction}$: E contribution of attractive forces
 $E_{repulsion}$: E contribution of repulsive forces
 r : distance of atoms
 E_b : binding energy
 R_0 : binding distance

Bond Types

Classification systems:

- Intra/intermolecular bonds
- According to the strength of bonds:
 - Strong/weak
 - Primary (covalent, metallic, ionic)
 - Secondary (dispersion, dipole-dipole, hydrogen)
- Other Bond types...



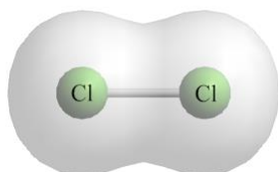
I. Covalent bond

- Atoms are held together by electrons of common orbitals, called molecular orbitals.
- Valence electrons interact the most.
- Single, double or triple bond depending on the number of shared electrons.
- Some electrostatic component may be present (see later), but not predominant.
- Strong interaction: $E_k > 1 \text{ eV}$
- ($1 \text{ eV} = 1,6 \cdot 10^{-19} \text{ J} = 96 \text{ kJ/mol} \sim 100 \text{ kJ/mol}$)

I. Covalent bond

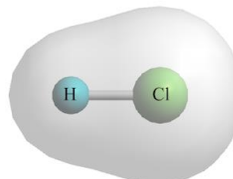
Apolar (nonpolar or homeopolar) bond

Center of (+) és (-) charges of the molecule coincides
 Binding electron(s) shared equally
 „purely covalent”
E.g. H_2 , Cl_2 , O_2 , ...



Polar (heteropolar) bond

Centers of (+) and (-) charges do not coincide
 Electric dipole is formed
 Electron cloud is polarized
 Electrostatic contribution appears
E.g.: HCl , HF , H_2O , ...

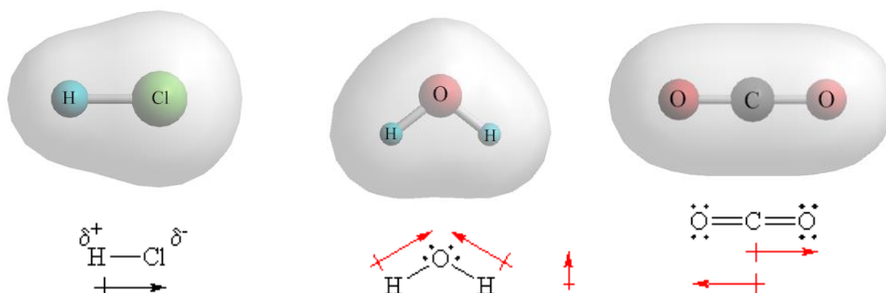


I. Covalent bond

Electric dipole moment: measure of the separation of positive and negative electric charges.

$$p = Qd$$

p: electric dipole moment
Q: charge (positive and negative)
d: distance of separation
[D, debye] ($1D = 3,34 \cdot 10^{-30} \text{ Cm}$)



I./b Metallic bond

- Atoms are held together by common electrons (delocalized orbital)
- Valence electrons form continuous electron band
- Strong interaction: $E_k > 1 \text{ eV}$
- Multi atomic systems

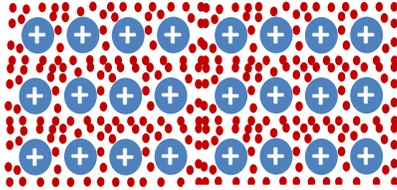
Periodic Table of the Elements

1	H	2	He
3	Li	4	Be
11	Na	12	Mg
19	K	20	Ca
27	Co	28	Ni
35	Br	36	Kr
43	Tc	44	Ru
51	Sb	52	Te
59	Pr	60	Nd
67	Ho	68	Er
75	Re	76	Os
83	Bi	84	Po
91	Pa	92	U
99	Es	100	Fm
107	Unh	108	Uno
115	Nh	116	Nl
117	Ts	118	Og

Legend:

- hydrogen (green)
- alkali metals (yellow)
- alkali earth metals (light blue)
- transition metals (orange)
- poor metals (blue)
- nonmetals (white)
- noble gases (red)
- rare earth metals (grey)

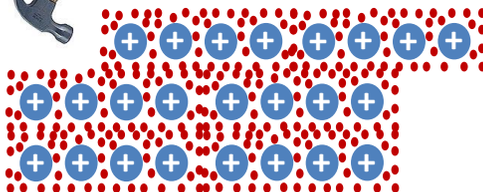
I./b Metallic bond



Metallic lattice: positively charged metal ions in a crystall lattice surrounded by a cloud of delocalized electrons.



I./b Metallic bond



Physical properties:

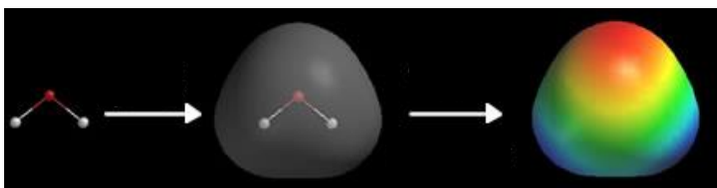
Metals are:

- Non transparent
- Ductile (shapeable)
- Good electric conductors
- etc.

II. Bond types involving electrostatic interactions

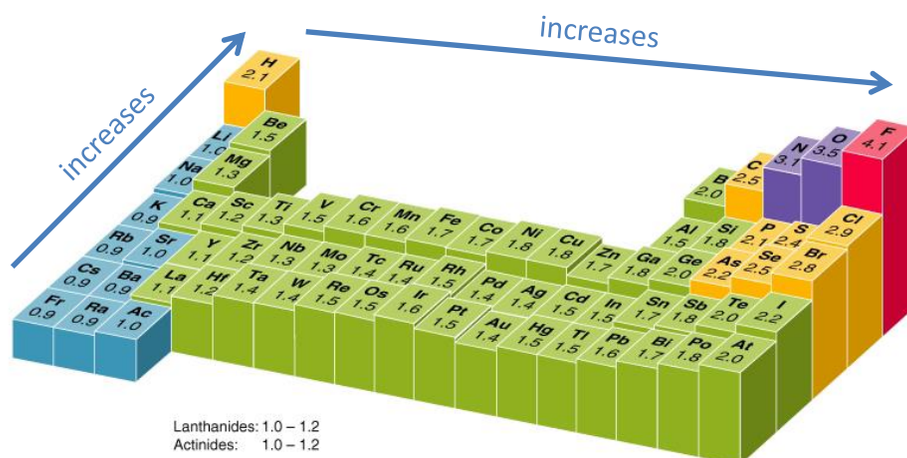
Electronegativity

- is a chemical property that describes the tendency of an atom or a functional group to attract electrons towards itself.
- The higher the associated electronegativity number, the more an element or compound attracts electrons towards it.
- Approximately proportional to the sum of ionization energy and electron affinity.
- Calculated with different methods (*Pauling, Mulliken, Sanderson...*)



II. Bond types involving electrostatic interactions

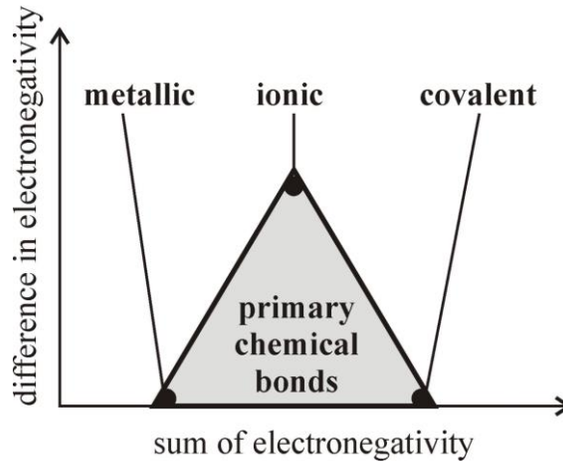
Electronegativity according to L. **Pauling** (dimensionless units)



II. Bond types involving electrostatic interactions

Classic bonds classified according to electronegativity:

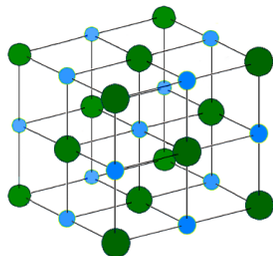
$< 0,6$ (apolar covalent) $0,6 - 2,1$ (polar covalent) $2,1 <$ (ionic)



II./a Ionic bond

- Atoms are held together by Coulombic forces between (+) és (-) point charges
- „Limiting case of heteropolar bonds”
- Formed between atoms of significantly different electronegativity (e.g.: NaCl , $\Delta EN = 3 - 0,9 = 2,1$)
- Can form between two atoms, but usually multi-atom systems.
- Long range interaction - attraction is inversely proportional to the distance (decreases slowly with it).
- Electrostatic interaction can be largely affected by other charged components (eg. dissociation in water!)
- Strong interaction ($E_b > 1 \text{ eV}$)

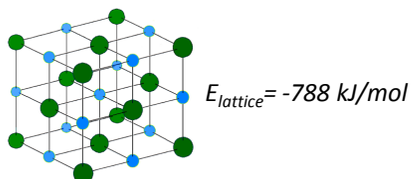
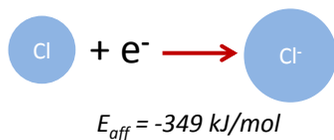
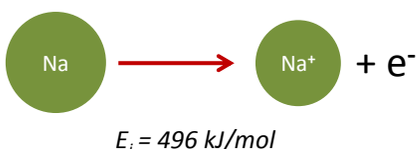
II./a Ionic bond



Ionic crystals: stoichiometric ratio of positive and negative ions are structured into a periodic crystalline structure. (e.g.: NaCl)



II./a Ionic bond



Ionization energy: is the energy *required* to remove electrons from gaseous atoms or ions.

Electron affinity: amount of energy *released* when an electron is added to a neutral atom or molecule to form a negative ion (measured in the gaseous state).

Lattice energy: measure of the strength of bonds in an ionic compound. Energy *required* to completely *separate* one mole of a solid ionic compound into gaseous ionic constituents.

II./b Dipole-dipole interaction

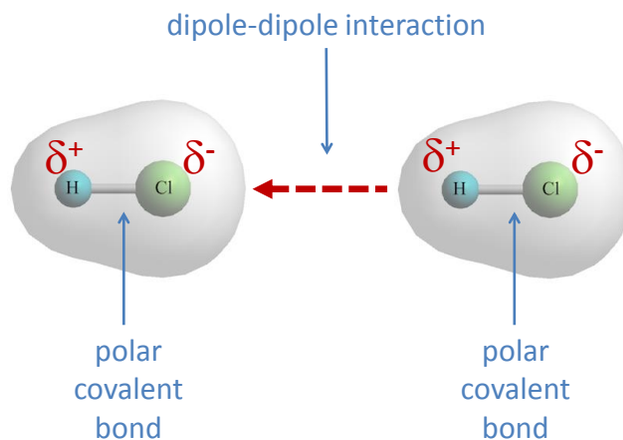
- Constant charge distribution is present in a (given part of a) molecule
- Partially (+) and (-) segments are held together by electrostatic interactions (Coulombic forces)
- Intra/intermolecular interaction.
- Weak interaction ($E_b = 0,003-0,02$ eV)
- Energy of attracting interaction between dipoles:

$$E_{\text{attraction}} = p E$$

p: dipole momentum
E: electric field strength generated by the surrounding partners

($E_{\text{repulsion}}$: stems from the repulsion of the participant's electron cloud)

II./b Dipole-dipole interaction

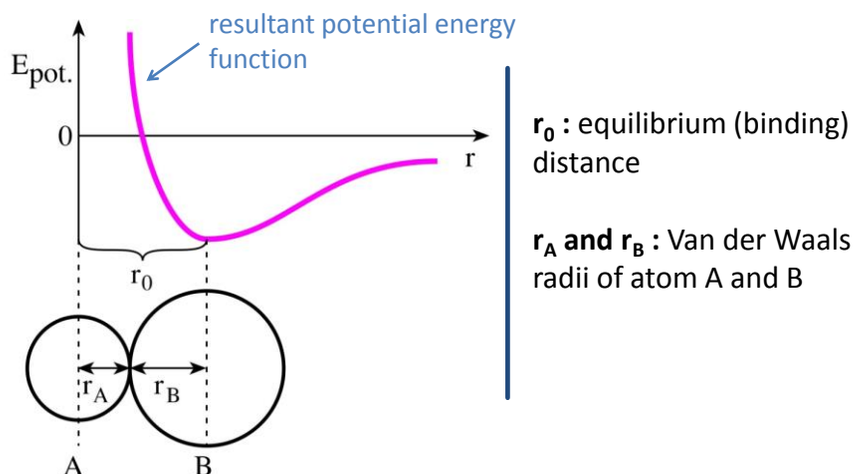


III. Van der Waals-interactions

- Sum of attractive and repulsive interactions between two apolar atoms, molecules or apolar molecular parts.
- The **attractive contribution** (also called *London-*, or *dispersion force*) is a result of temporarily created dipoles that can induce the polarization of other apolar molecule or molecular part.
- Intermolecular or intramolecular interaction.
- Important biological role: formation of organic structures.
- Weak connection ($E_b \sim 0,02 \text{ eV}$)
- *[according to other classifications Van der Waals interactions involve all types of weak electrostatic connections (permanent dipole-permanent dipole; permanent dipole-induced dipole, induced dipole-induced-dipole)]*

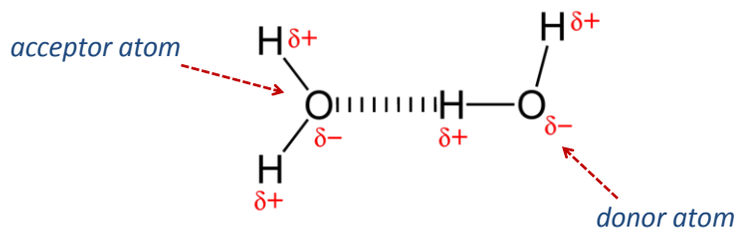
III. Van der Waals-interactions

Equilibrium distance of a Van der Waals interaction can be considered as one definition of atomic size.



IV. Hydrogen bond

- Two atoms of high electronegativity are held together with a Hydrogen-bridge.
- Primarily between **F, N, O** atoms (pillar atoms).
- Intermolecular/intramolecular interaction
- Typical bond distance: 0,23 – 0,35 nm
- Well defined structure.
- Important role in structural biology and biochemical reactions.
- Medium strong interaction (typical $E_b = 0,2$ eV)

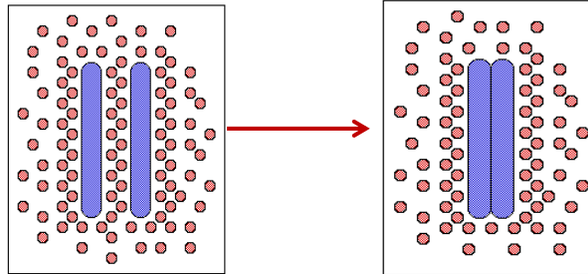


V. Hidrophobic interaction

- Appear in aqueous systems (like biological environments!)
- Apolar, hydrophobic molecules tend to aggregate (form bonds with each other) in aqueous environment in order to exclude the polar water molecules and minimize their surface area exposed to water.
- Intra/intermolecular interaction
- It has mostly entropic origin (*see later at Thermodynamics*) through reduction of highly structured water cage around the apolar surfaces. (*see Organisation of Water later*)
- Important role in structural biology and biochemical reactions.
- Weak interaction.



V. Hidrophobic interaction



Scanning Probe Microscopy (SPM)

Family of instruments used for studying surface properties of various materials.

How do they work?

Etimology and function:

Microscopy: a method being able to form image of small objects.

How small? Size of resolvable objects spans from few pm-s to several μm -s.

They are not „*scope*“-s in the classic sense of the word: They do not „*see*“ the object, they „*touch*“ it.

Scanning Probe Microscopy (SPM)

Probe:

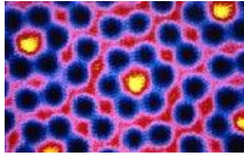
- A tiny, specifically designed component particularly sensitive to *atomic interactions*.
- The probe is brought very close to the sample surface.
- The sensed interactions can be correlated with the distance between the probe and the sample.
- Various interactions can be observed depending on the design of the probe.
- SPM methods are named after the type of atomic interaction sensed by a certain probe.

Scanning Probe Microscopy (SPM)

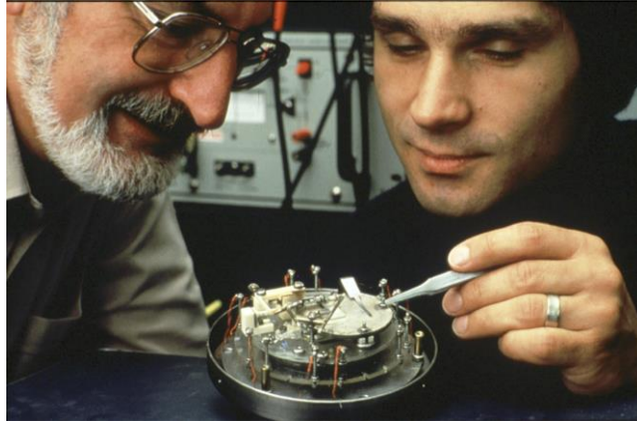
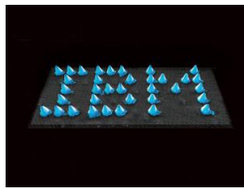
Scanning:

- A scanner controls the precise *position* (X; Y) of the probe and its *distance* (Z) from the surface to be imaged.
- The probe can be moved with pm sensitivity in X; Y; Z directions.
- The surface of region of interest (ROI) is scanned point by point during a measurement.
- (The material that enables such precise positioning is *piezoelectric ceramic*. If voltage is applied on it, the ceramic changes its geometry.)

Scanning Tunneling Microscope (STM) 1981



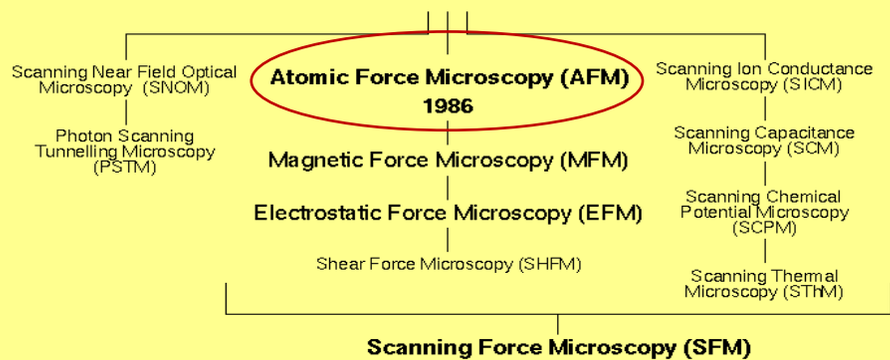
Atoms in a
silicon chip



Heinrich ROHRER and Gerd BINNING
Nobel prize: 1986

Scanning Probe Microscopy "Family Tree" (SPM)

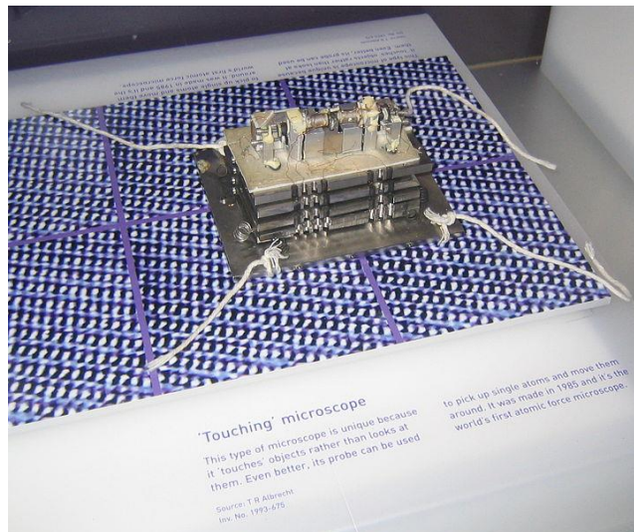
Scanning Tunnelling Microscopy (STM) 1981-2



Atomic Force Microscopy (AFM)

- The probe is a tiny, sharp tip, usually few μm -s tall, and only few nm -s in diameter at the apex.
- The tip is located at the free end of a cantilever that is usually 100-500 μm long.
- Topographic image is collected with ~ 10 pm vertical and somewhat worse horizontal resolution.
- Any surfaces (conductors, insulators and semiconductors) can be imaged.
- Works in air and in fluid environment as well.
- Usually does not require fixation or staining of the sample.
- Biological samples can be examined in their native state and physiological environment.

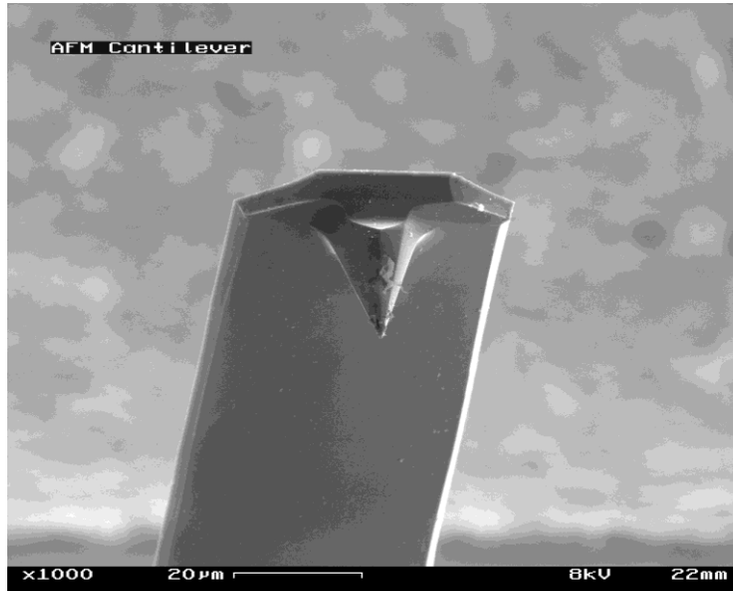
Atomic Force Microscopy



The first one. 1986.

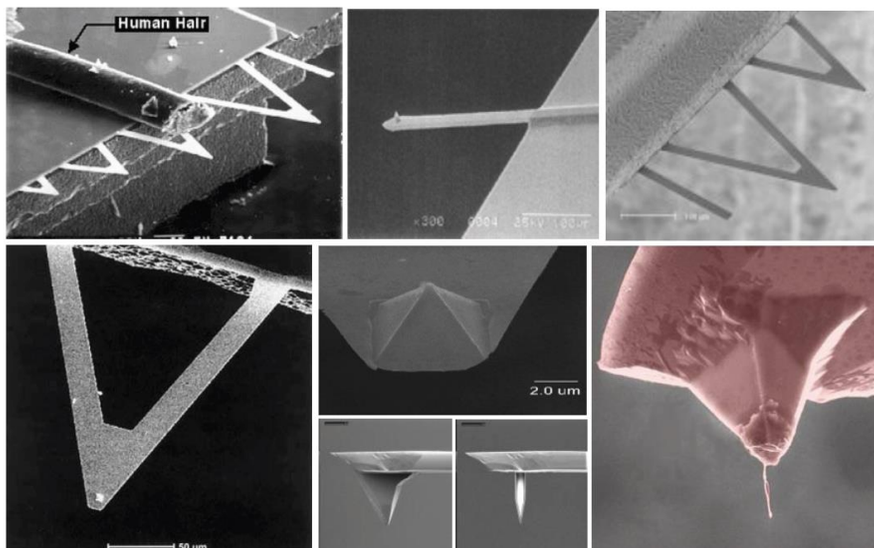
Atomic Force Microscopy

Tip at the end of a cantilever

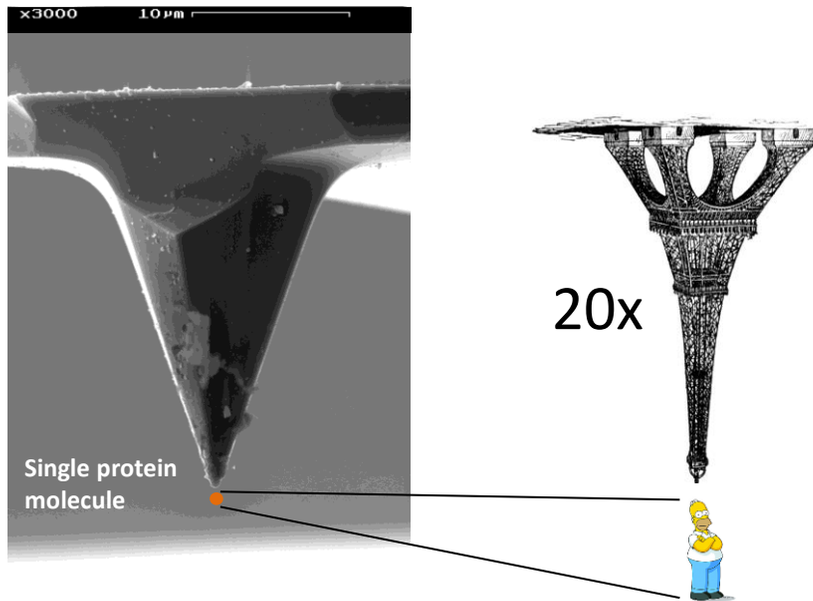


Atomic Force Microscopy

Different types of AFM cantilevers.

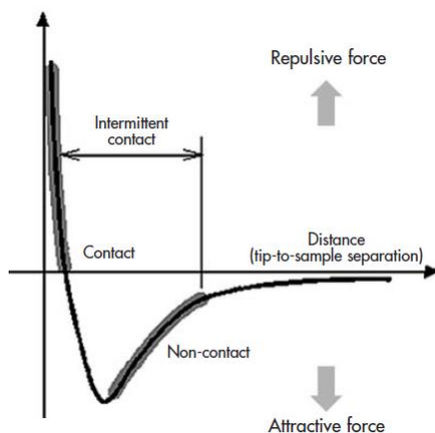


Atomic Force Microscopy



Atomic Force Microscopy

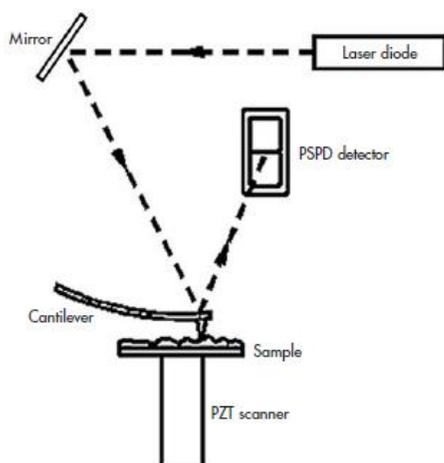
- Bringing the tip very close (few nm-s) to the sample atomic interactions arise between the very last atoms of it and the atoms of the sample.



Depending on the tip-sample distance:

- Net attraction or repulsion may occur.
- Attraction at „longer” distances.
- Repulsion at very short distances.

Atomic Force Microscopy

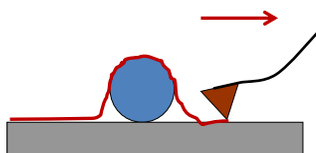


- Attractive and repulsive effects cause the cantilever to bend.
- Deflection of the cantilever is detected by an optical system.
- A laser beam is pointed at the very end of the cantilever, and reflected back to be sensed by a position-sensitive photodetector (PSD).
- Thus vertical deflection of the cantilever is amplified and can be detected with sub-Å sensitivity..

Atomic Force Microscopy

Contact mode:

- The tip is in perpetual contact with the sample surface.
- The deflection of the cantilever, i.e. the force exerted on the sample by the tip is held constant.
- A Z feedback system is utilized to maintain the deflection at a constant value (setpoint) by lifting or lowering the cantilever.
- Topography data (i.e.: height) in each X;Y point is calculated from these Z movements



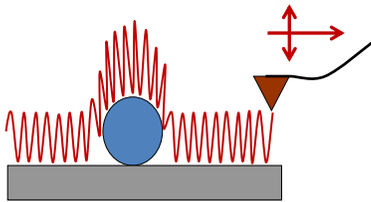
Disadvantage:

- Lateral forces exerted by the scanning tip may damage softer samples.

Atomic Force Microscopy

Oscillating mode: (Tapping mode, Non-contact mode)

- Cantilever is oscillated close to its resonant frequency.
- The tip taps the surface gently
- The amplitude of cantilever oscillation changes with surface topography.
- A Z feedback system is utilized to maintain the amplitude at a constant value (setpoint) by lifting or lowering the oscillating cantilever.
- Topography data (i.e.: height) in each X;Y point is calculated from these Z movements



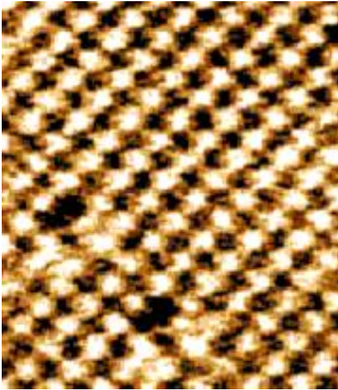
Advantage:

- Virtually eliminated lateral forces.
- Allows more gentle imaging.
- Applicable for soft samples.

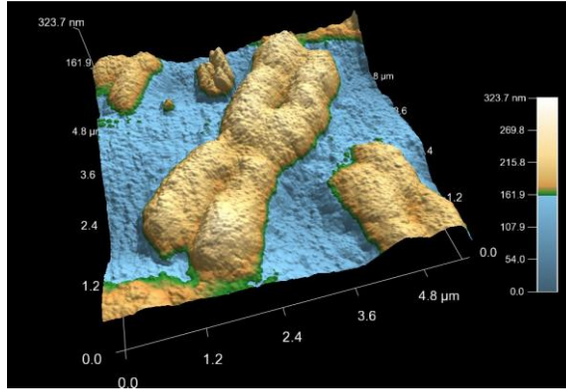
Atomic Force Microscopy

<http://www.youtube.com/watch?v=BrsoS5e39H8>

Atomic Force Microscopy Images

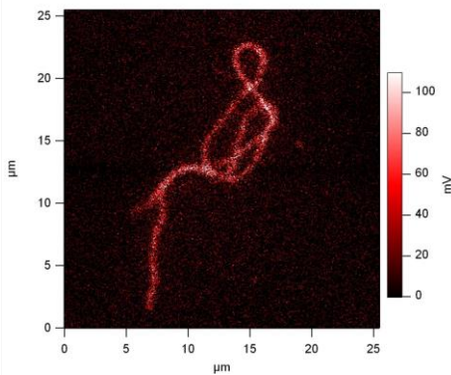


NaCl crystal surface

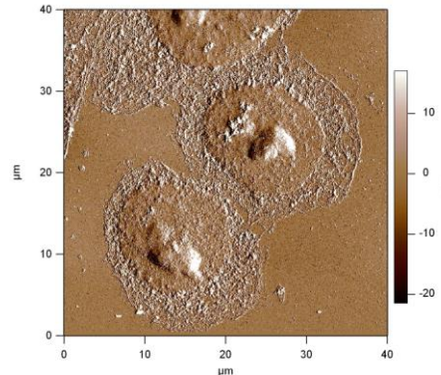


Human metaphase chromosomes

Atomic Force Microscopy Images



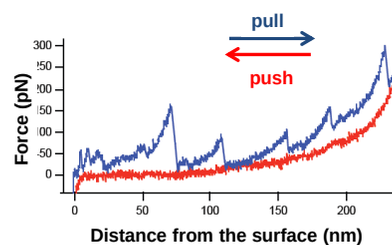
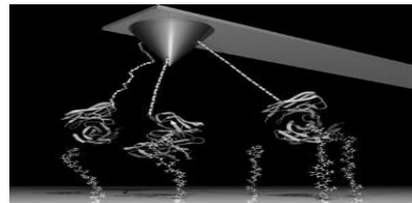
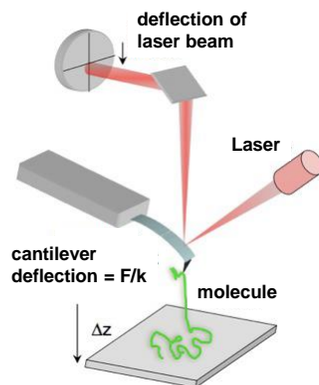
„The thinker”
a single actin polymer



HeLa cells on glass

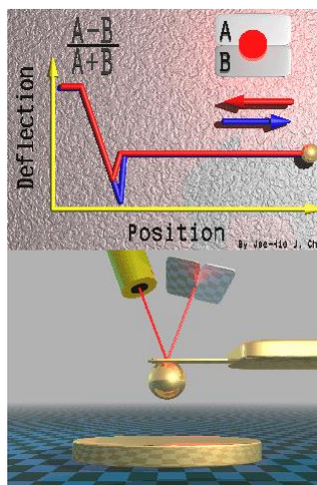
Atomic Force Microscopy

Force spectroscopy: Force-distance traces registered upon push-pull cycles of the AFM tip (movement in Z direction)
 ~ 10 pN sensitivity

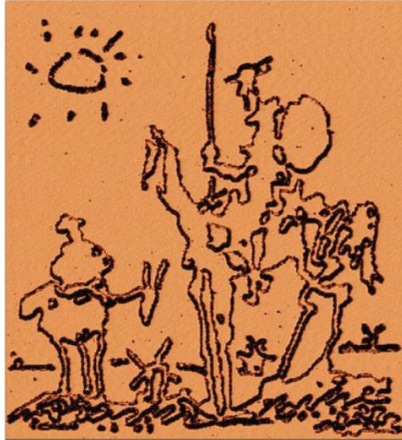


Atomic Force Microscopy

Force spectroscopy:



Thank you for your attention!



Pablo Picasso: "Don Quixote"
polikarbonát felszínbe rajzolva

← 1 μm →