

AI4Science camp USC-Columbia 7/25/24
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Data science for chemistry

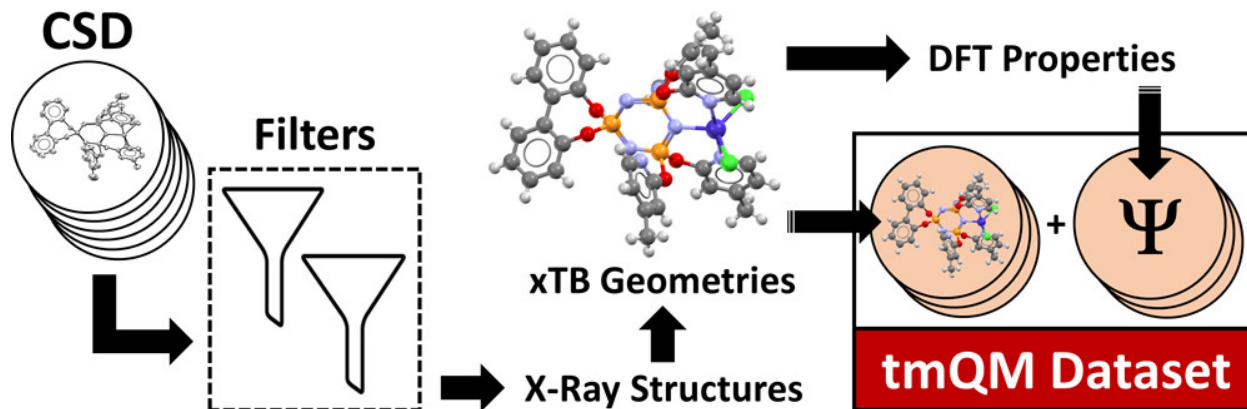
Focus on the molecules

*[some background for the rational choice
of ML descriptors]*

tmQM Dataset—Quantum Geometries and Properties of 86k Transition Metal Complexes

David Balcells and Bastian Bjerkem Skjelstad

Journal of Chemical Information and Modeling **2020** 60 (12), 6135-6146 DOI: 10.1021/acs.jcim.0c01041



- Quantum chemistry
- Electronic structure
- Machine learning

outline

- Atoms and the **periodic table**
- Electronic configurations
- Properties of elements as potential descriptors
- Molecular properties
- Chemical bonding
- **tmQM dataset**

General chemistry textbook by Stephen Lower, Ch.5

[https://chem.libretexts.org/Bookshelves/General_Chemistry/Chem1_\(Lower\)](https://chem.libretexts.org/Bookshelves/General_Chemistry/Chem1_(Lower))

Atom

Proton 'p'	m=1836.15 a.u	charge = +1
Electron 'e'	m = 1.0 a.u.	charge = -1
Neutron 'n'	m = 1838.68 a.u.	charge = 0

Atomic units:

|e-charge|=1 a.u. = 1.6×10^{-19} Coulomb e-mass = 1 a.u. = 9.11×10^{-31} kg

Distance a.u. is bohr, 1 bohr = average e-p separation in H-atom

1 bohr = 0.53 Angstrom

1 Angstrom = 10^{-10} meters = 100 picometers [pm]

Identity of an atom is defined by the number of protons

Number of neutrons defines an isotope ^2H = proton + neutron + electron

Neutral atom: $N_p = N_e$ charge = 0 sodium atom Na

Cation: remove an electron sodium cation Na^+

Anion: add an electron sodium anion Na^-

The typical size of an atom or a molecule is 1-10 Angstrom or 100-100 pm

size of H^+ = 0.00084 pm size of H = 53 pm

Atom is 'empty'

- At this scale atoms and molecules are governed by quantum mechanics (QM) and in particular electrons act as both particles and waves
- They are governed by a differential equation

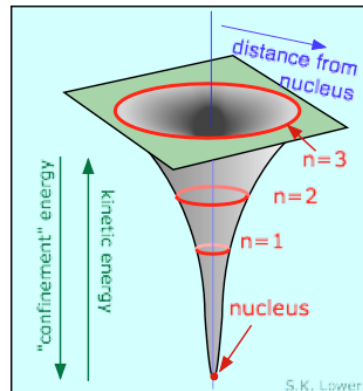
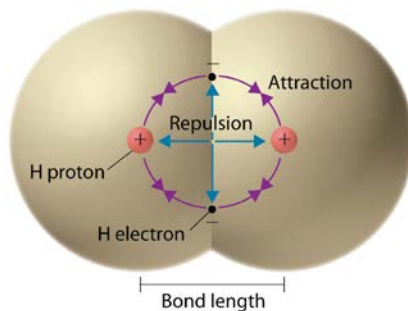
Time-independent Schrödinger equation

$$\hat{H} |\Psi\rangle = E |\Psi\rangle$$

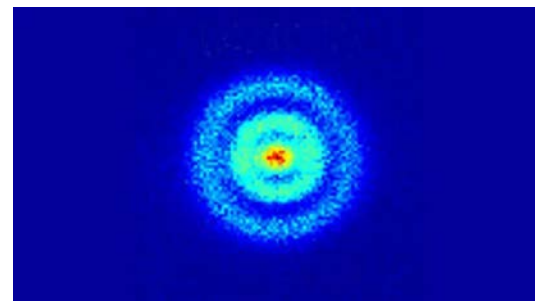
Hard!
Try ML

- Coulomb interaction between the charges
- Electrons are described by stationary probability distributions, $|\Psi|^2$, of charge -- *orbitals* -- of certain energy, E
- The energy values are discrete; orbitals have specific shapes
- Electrostatic (Coulomb) interaction between the charges

$$V \sim -1/\text{distance}$$



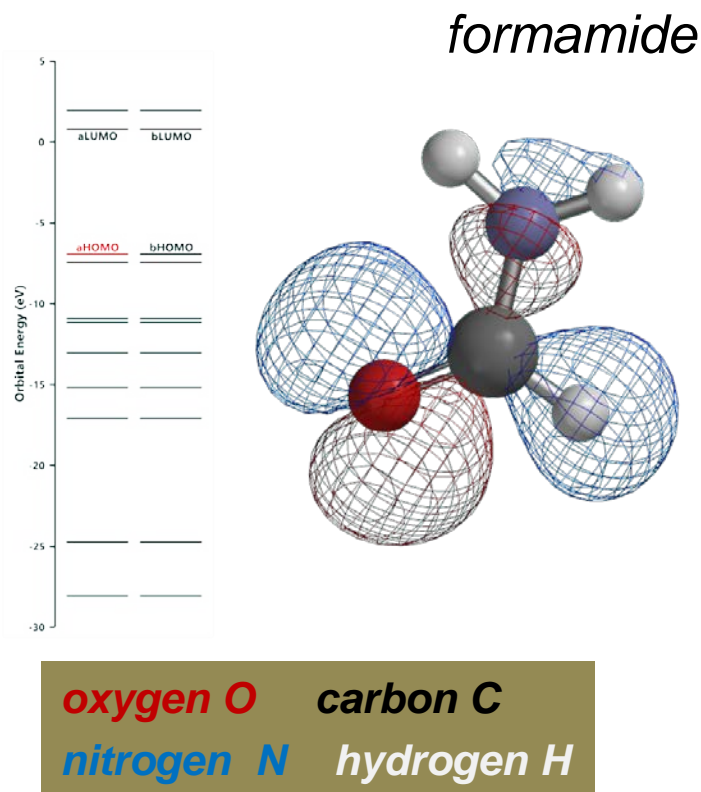
Hydrogen Atoms under Magnification: Direct Observation of the Nodal Structure of Stark States
A. S. Stodolna et al JPCL 2013



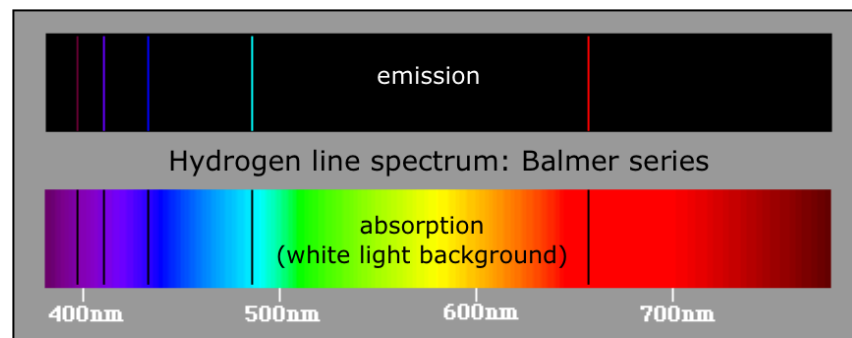
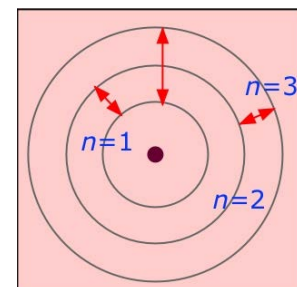
- Electrons 'occupy' energy levels
- Atomic or molecular orbitals (MO)
- HOMO = highest occupied MO
- LUMO = lowest occupied MO
- Energy level diagram

Absorption and emission of electromagnetic radiation (light) corresponds to the difference between the energy levels

A 'material design target'



H atom



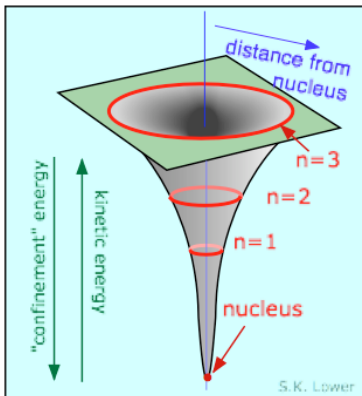
Shapes of atomic orbitals

labeled ***ns, np, nd, nf***

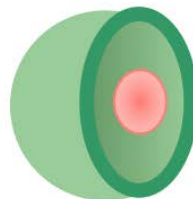
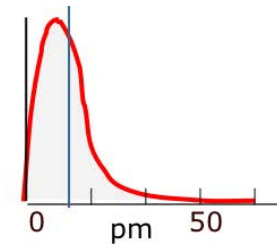
n corresponds to energy $E \sim -1/n^2$

s/p/d/f to the orbital shape

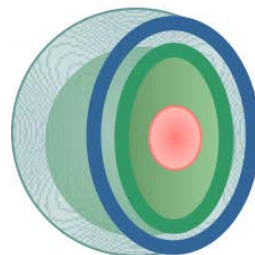
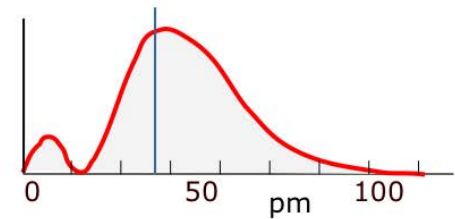
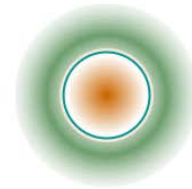
s is spherically symmetric



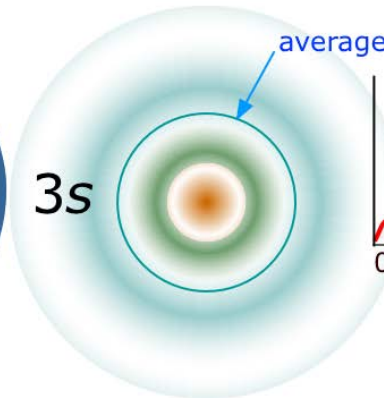
1s



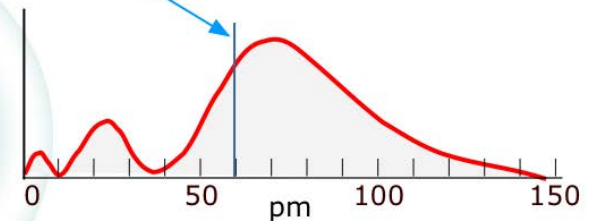
2s

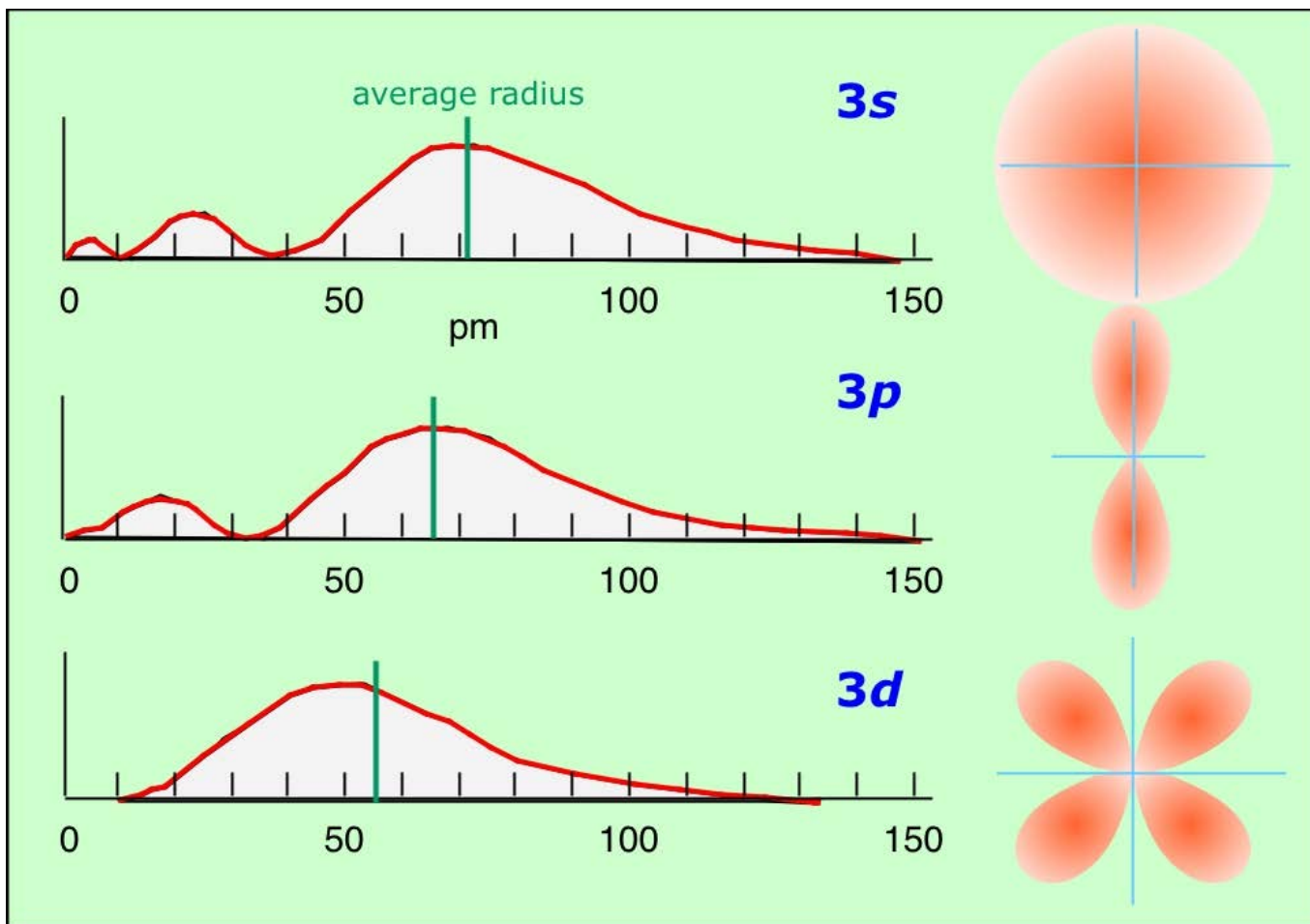


3s

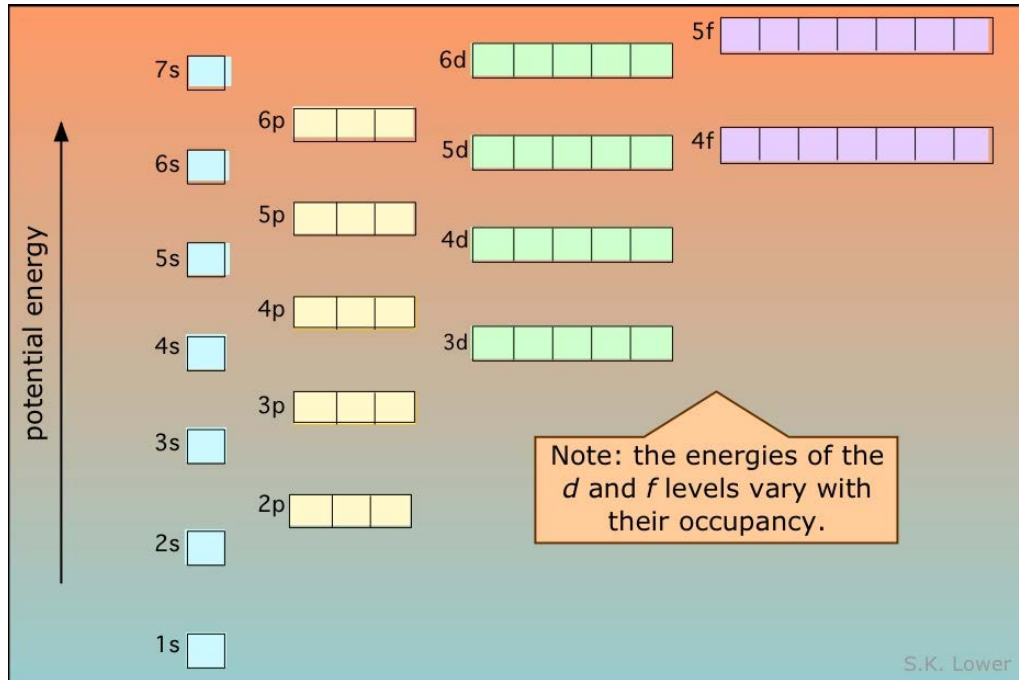


average radius





The energy level diagram in a multielectron atom



- The electrons fill up the energy levels from bottom up
- Electrons have additional property 'spin'
- Each level holds up to 2 electrons – one spin up, one spin down
- If the levels are of equal energy, the electron spins will be maximally unpaired
- The electronic structure of the highest energy shell (the highest n) -- the valence electrons -- defines chemical bonding
- Electron pairs 'make' chemical bonds (spin is important here)

Similar electronic structure in the valence shell →
 Similar chemical and physical properties →
 Periodic table of elements (Mendeleev 1869)

7s	Fr Ra	6d	Rf Db Sg Bh Hs Mt Ds 112	5f	Th Pa U Np Pu Am Cm Bk Cf Es Fm Md No Lw
6s	Cs Ba	5d	La Hf Ta W Re Os Ir Pt Au Hg	4f	Ce Pr Nd Pm Sm Eu Gd Tb Dy Ho Er Tm Yb Lu
5s	Rb Sr	4d	Y Zr Nb Mo Tc Ru Rh Pd Ag Cd		
4s	K Ca	3d	Sc Ti V Cr Mn Fe Co Cu Ni Zn		
3s	Na Mg				
2s	Li Be				
1s	H He				

energy ↑

Periods = rows

Groups = columns

Diagram of the periodic table showing the following blocks:

- s-block:** Groups 1 (IA) and 2 (IIA). Elements: H, Li, Na, K, Rb, Cs, Fr, Be, Mg, Ca, Sr, Ba, Ra.
- p-block:** Groups 13 (IIIA) to 18 (VIIA). Elements: B, C, N, O, F, Ne, Al, Si, P, S, Cl, Ar, Ga, Ge, As, Se, Br, Kr, In, Sn, Sb, Te, I, Xe, Tl, Pb, Bi, Po, At, Rn.
- d-block:** Groups 3 (IIIB) to 10 (VIII). Elements: Sc, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Y, Zr, Nb, Mo, Tc, Ru, Rh, Pd, Ag, Cd, La, Hf, Ta, W, Re, Os, Ir, Pt, Au, Hg, Ac, Rf, Db, Sg, Bh, Hs, Mt, Ds.
- f-block:** Lanthanides (Ce to Lu) and Actinides (Th to Lr).

Labels in the diagram:

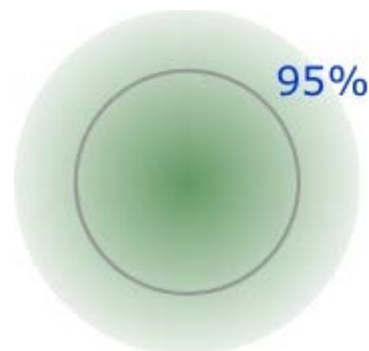
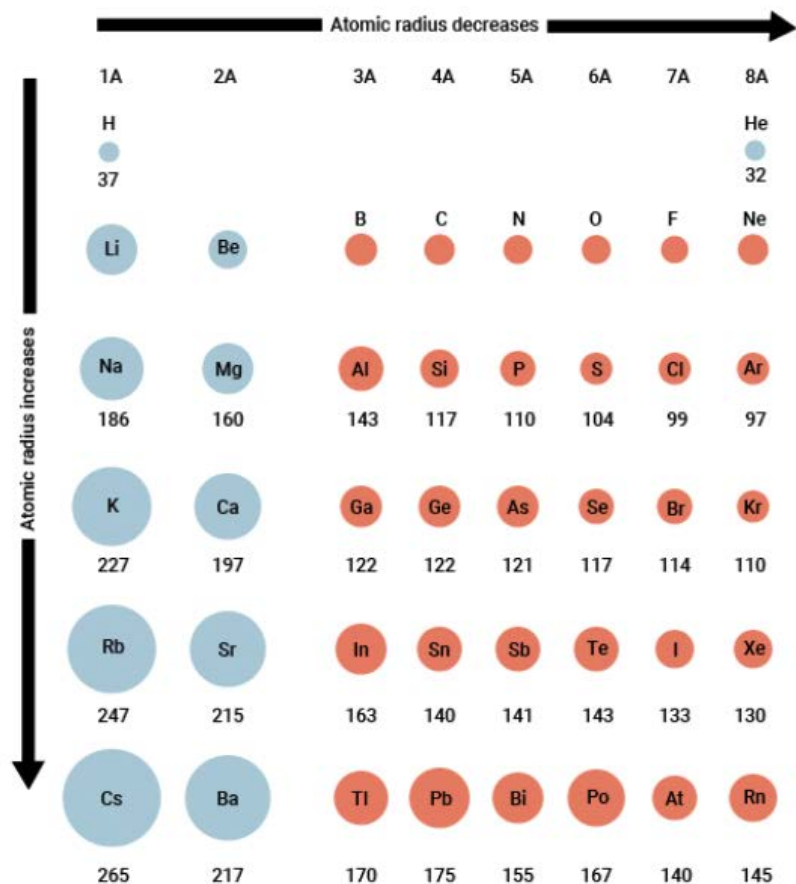
- "representative elements" (yellow box, pointing to s and p blocks)
- "main group elements" (orange box, pointing to p-block)
- s-block (red text)
- d-block (red text)
- p-block (red text)
- f-block (red text)

S.K. Lower

- The non-metallic elements occur only in the *p*-block;
- The *d*-block elements contain the so-called transition elements;
- The *f*-block elements go in between Groups 3 and 4 of the *d*-block.

Sizes of atoms and ions ~ 70-400 pm

In molecules both atomic sizes and charges are not well-defined



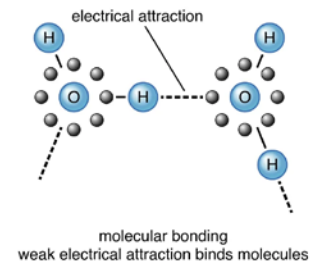
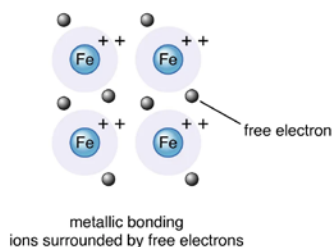
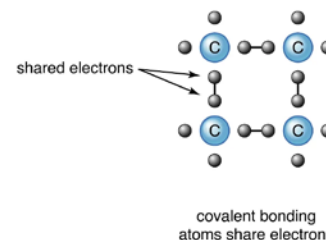
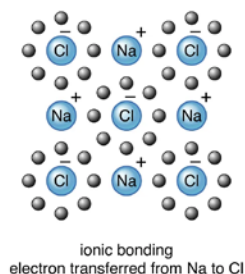
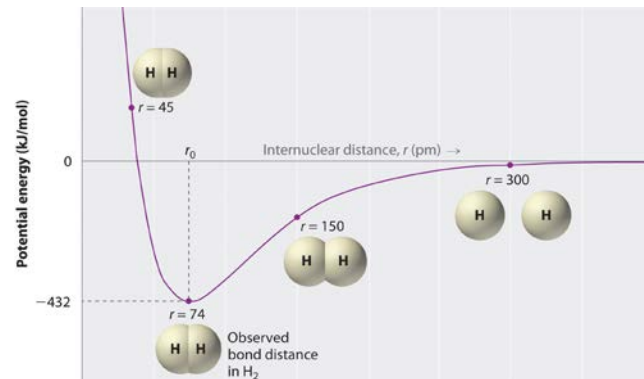
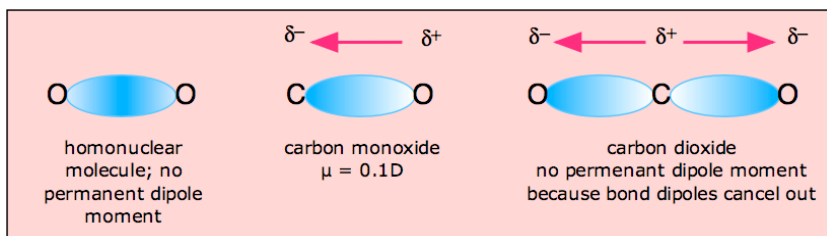
Chemical bonds

Minimum energy defines the bond distance

Shorter bonds are usually stronger

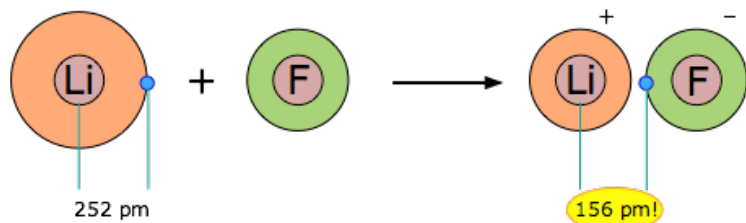
Bonding = (most often) shared electron pairs

Covalent bond (polar/non-polar)



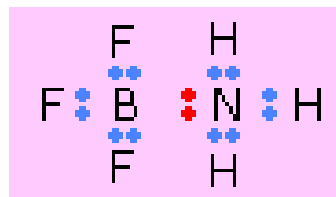
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Ionic bond



Dative or coordinate covalent bond:

both electrons come from one atom



Metallic bonding

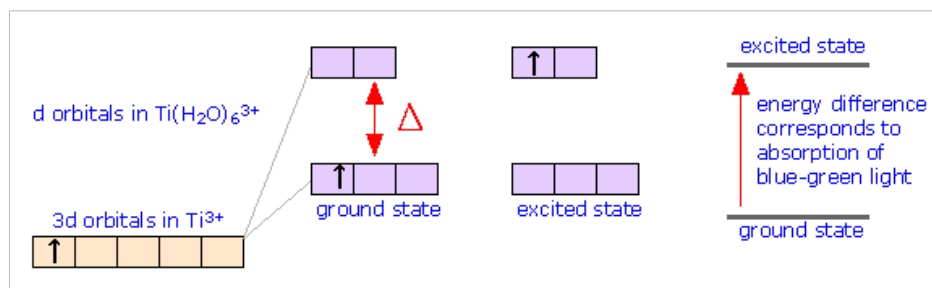
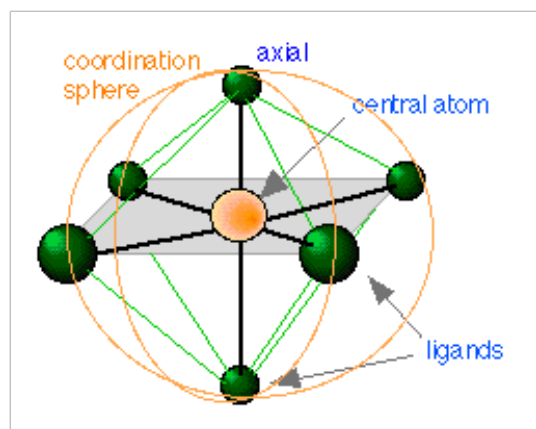
shared electrons

Hydrogen bond

partial charges

ML descriptor: number of bonds

Coordination complexes; transition metals



Fog: Color of various $\text{Ni}(\text{II})$ complexes in aqueous solution.

From left to right, hexaamminenickel(II), tris(ethylenediamine)nickel(II), tetrachloronickelate(II) and hexaaquanickel(II)

Wealth of info in the periodic table module

mendelev 0.17.0

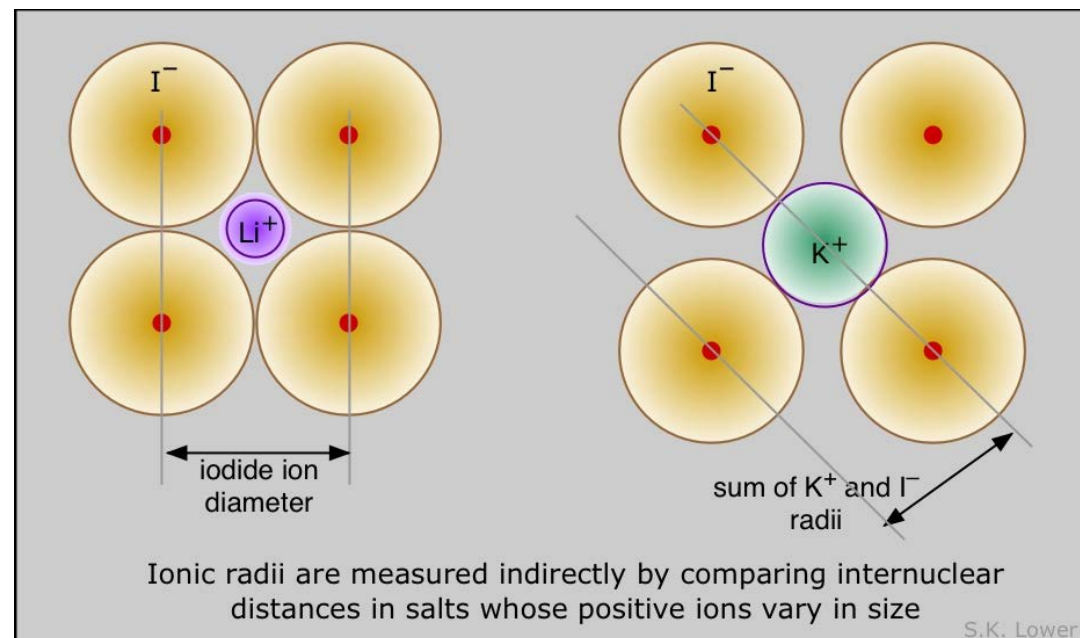
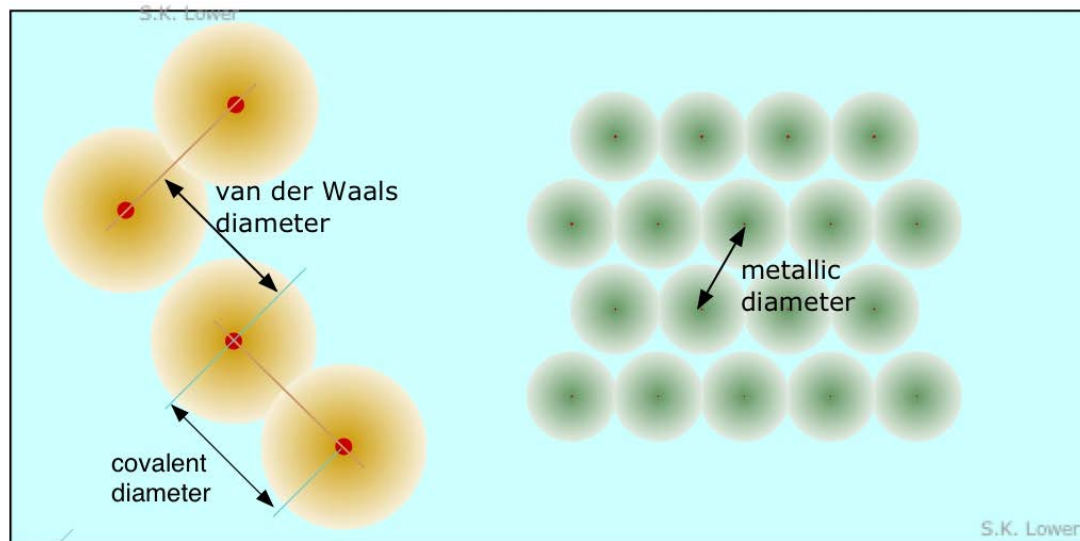
pip install mendelev

- [Data](#)

- [Basic properties](#)
- [Standardized colors schemes](#)
- [Size related properties](#)
- [Electronegativity scales](#)
- [Descriptive properties](#)
- [Physical properties](#)
- [Computed properties](#)
- [Isotope properties](#)

ML descriptor: size

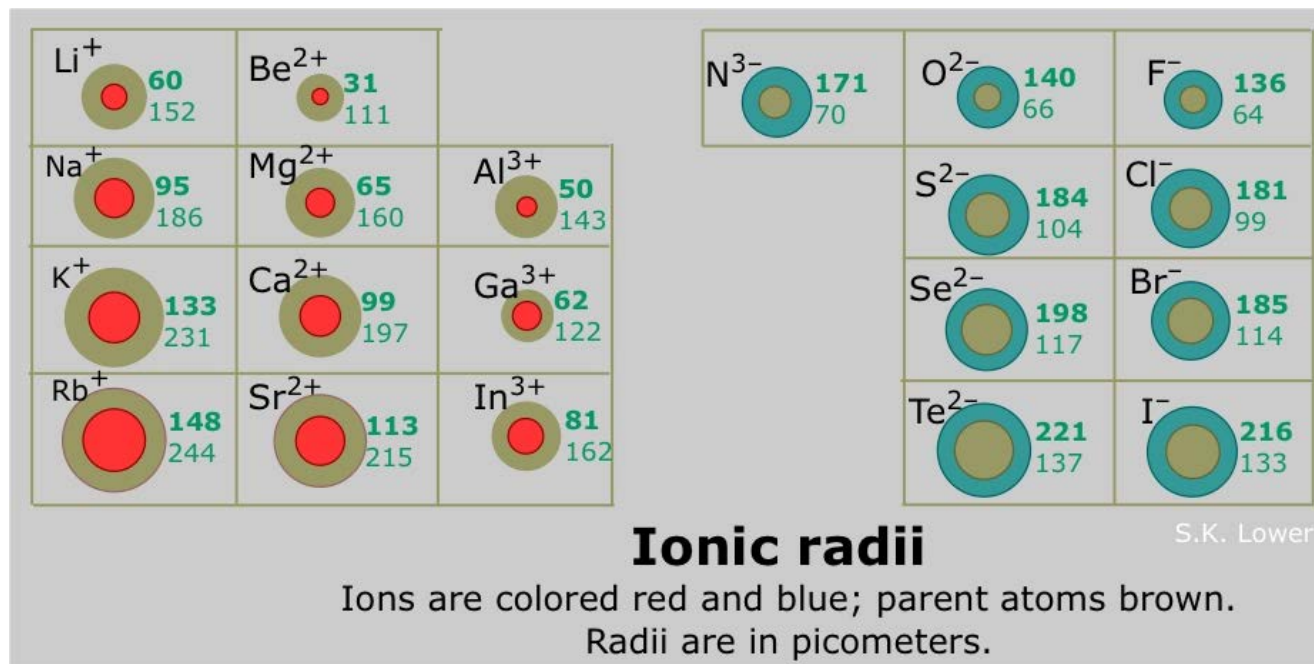
- may correlate with bonding strength, charge ...
- Choices



A positive ion < neutral atom because less e- electron-electron repulsion

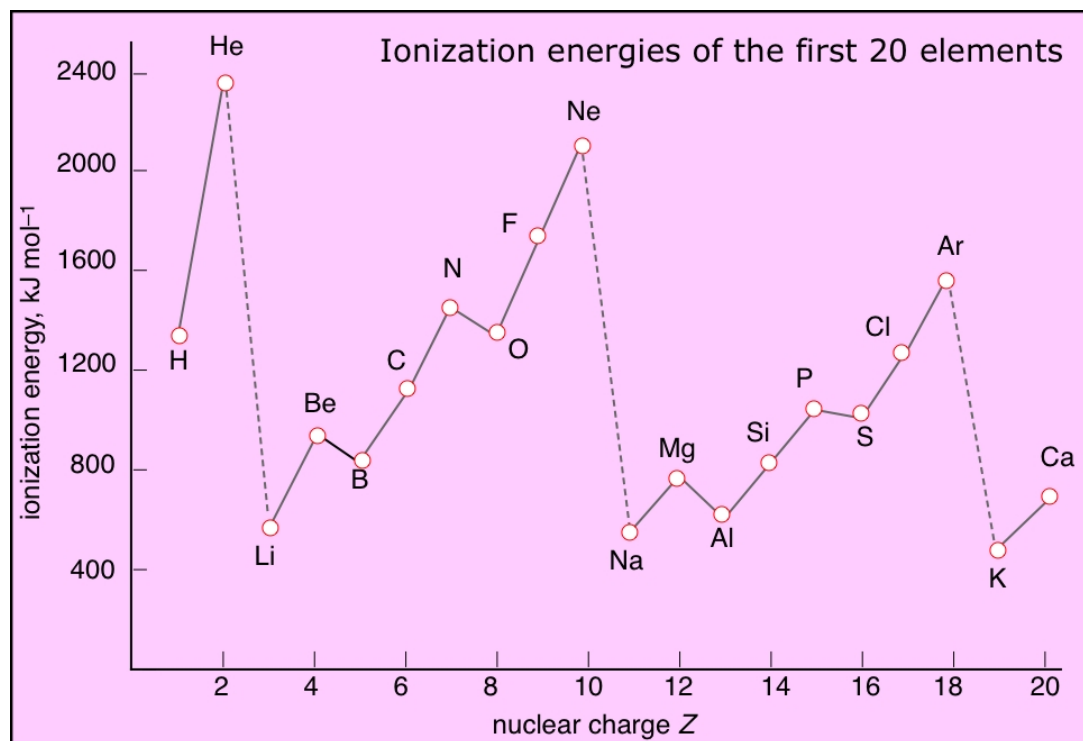
The ionic radius of Fe^{2+} is 76 pm, while that of Fe^{3+} is 65 pm

Negative ions > neutral atom because extra electron increases *electron-electron repulsion* which results in a general expansion of the atom



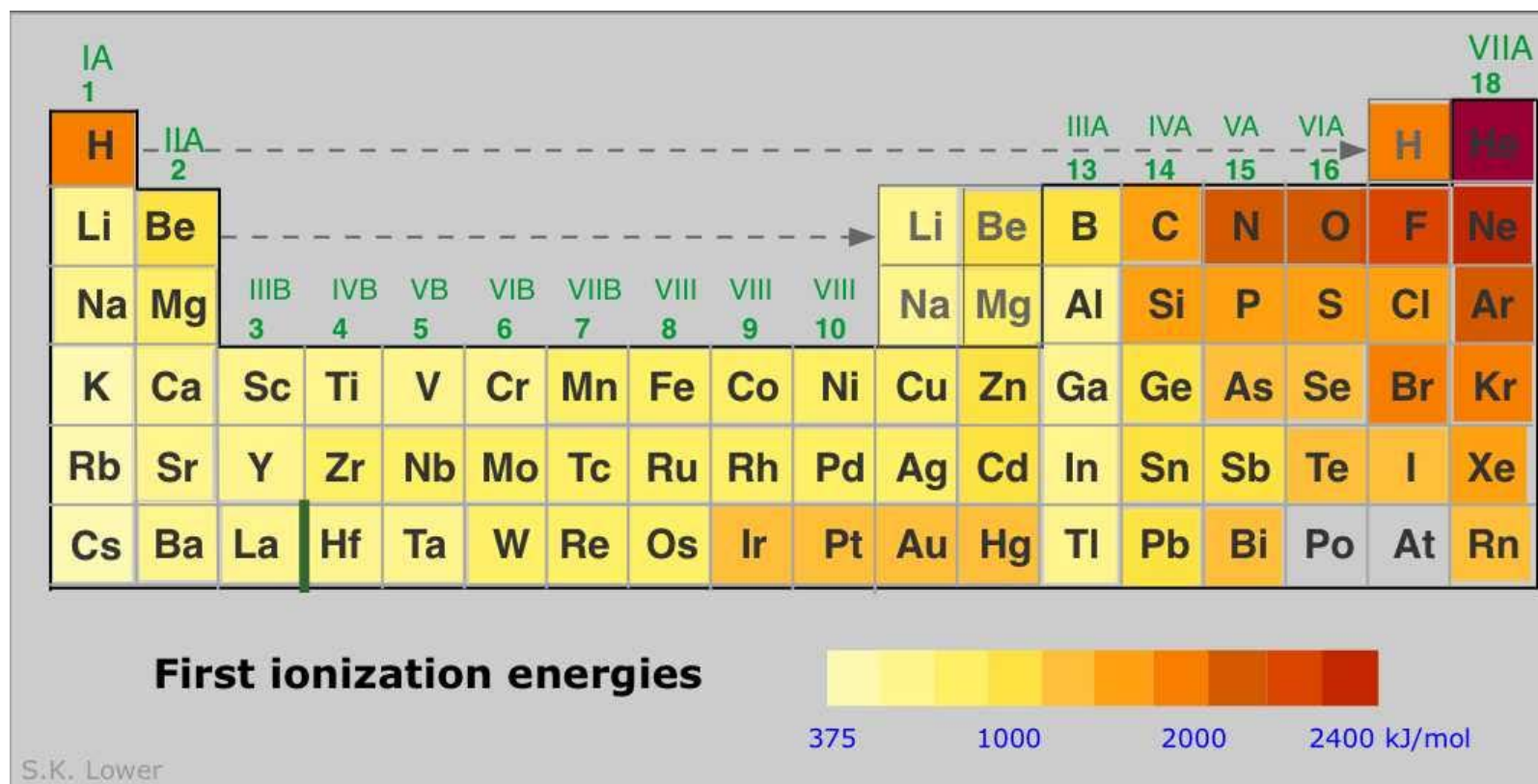
ML descriptor: Ionization Energy

This term always refers to the formation of *positive* ions. In order to remove an electron from an atom, work must be done to overcome the electrostatic attraction between the electron and the nucleus; this work is called the *ionization energy* of the atom and corresponds to the exothermic process



Periodic Trends in ion formation

- Chemical reactions are based largely on the interactions between the most loosely bound electrons in atoms
- Tendency of an atom to gain, lose or share electrons is one of its fundamental chemical properties



ML descriptor: Electron affinity

- Formation of a negative ion occurs when an electron from some external source enters the atom and become incorporated into the lowest energy orbital that possesses a vacancy (LUMO)
- Because the entering electron is attracted to the positive nucleus, the formation of negative ions is usually exothermic, i.e. the energy – **electron affinity** -- is given off

IA 1	Electron affinities in kJ released per mole of mononegative ions formed																IIIA 13						IVA 14	VA 15	VIA 16	VIIA 17	VIIIA 18
H 73	IIA 2																	B 27	C 122	N -7	O 141	F 328	He -21				
Li 60	Be -19																	Al 43	Si 134	P 72	S 200	Cl 349	Ar -35				
Na 53	Mg -19	IIIB 3	IVB 4	VB 5	VIB 6	VIIB 7	VIII 8	VIII 9	VIII 10	IB 11	IIB 12	Ga 29	Ge 116	As 78	Se 195	Br 325	Kr -39										
K 48	Ca -10	Sc 18	Ti 8	V 51	Cr 64	Mn	Fe 16	Co 64	Ni 112	Cu 118	Zn -47	In 29	Sn 116	Sb 103	Te 190	I 295	Xe -41										
Rb 47	Sr	Y 30	Zr 41	Nb 86	Mo 72	Tc 53	Ru 101	Rh 110	Pd 54	Ag 126	Cd -32	Tl 20	Pb 35	Bi 91	Po 183	At 270	Rn -41										
Cs 45	Ba	La	Hf	Ta 31	W 79	Re 14	Os 106	Ir 101	Pt 205	Au 223	Hg -61																
Fr 44	Ra	Ac	Db	Jl	Rf	Bh	Hn	Mt																			

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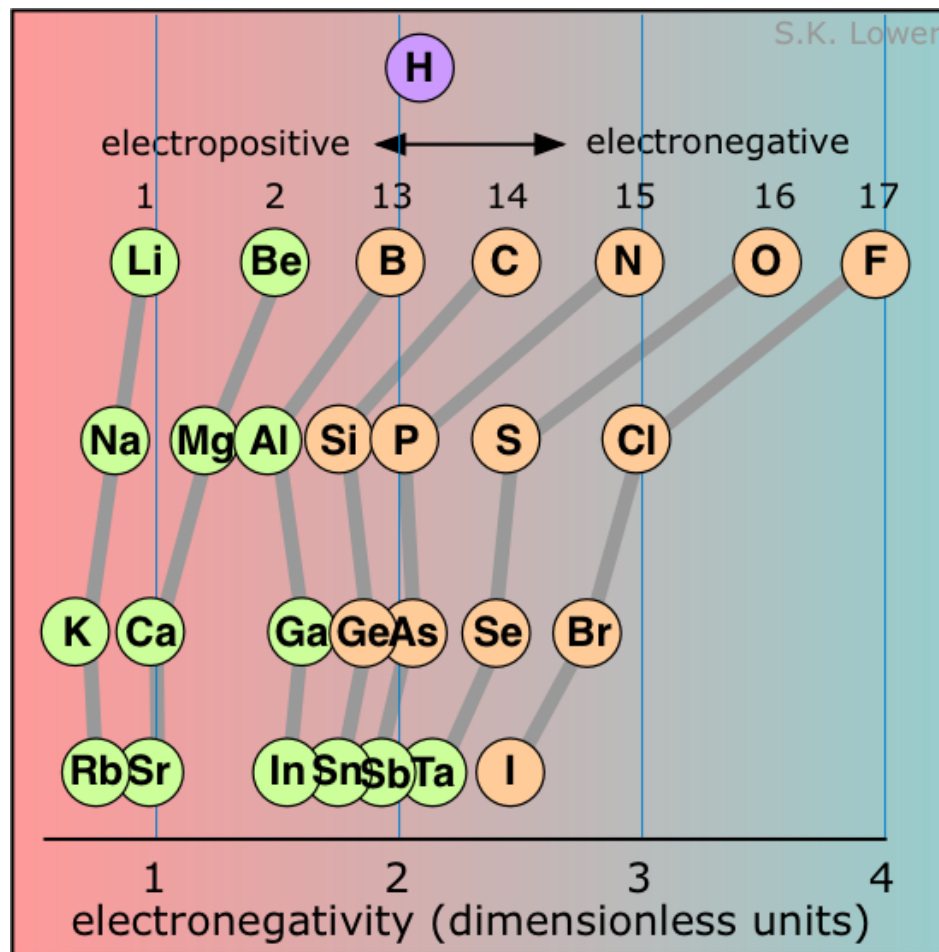
ML descriptor: Electronegativity

When two elements are joined in a chemical bond, the element that attracts the shared electrons more strongly is more electronegative

Elements with low electronegativities (the metallic elements) are said to be electropositive.

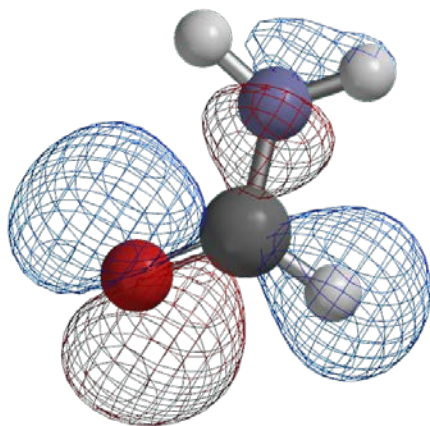
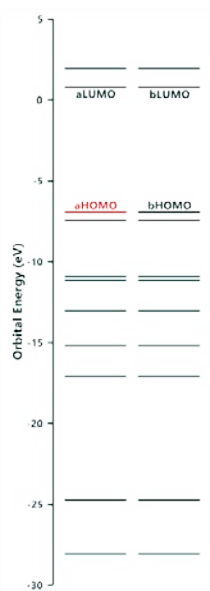
Electronegativities are properties of atoms that are chemically bound to each other, not of an isolated atom

Try the Pauling scale (other scales available)



- Computed molecular properties
- HOMO (~ionization energy)
- LUMO (~ electron affinity)
- Dipole moment (~ stacking)
- Polarizability (response to electric field)

Absorption and emission of electromagnetic radiation (light) corresponds to the difference between the energy levels



formamide

ML descriptors or design targets:

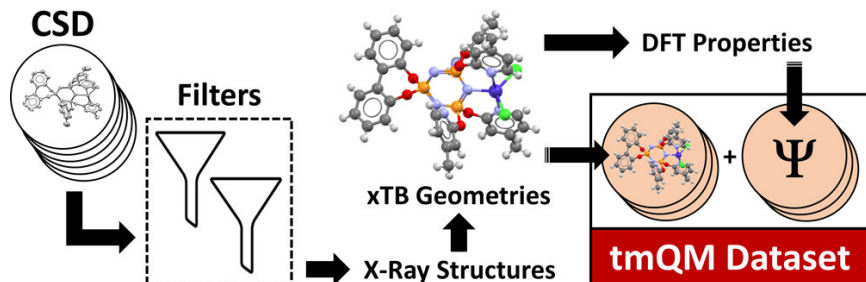
E_{HOMO} , E_{LUMO}

Energy gap = $E_{\text{HOMO}} - E_{\text{LUMO}}$

Fermi level or chemical hardness

$$\eta = (E_{\text{HOMO}} + E_{\text{LUMO}})/2$$

tmQM Dataset—Quantum Geometries and Properties of 86k Transition Metal Complexes



Computational protocol used to generate the tmQM data set. xTB = extended tight-binding; DFT = density functional theory; μ = dipole moment; α = polarizability; q = charge

- tmQM contains 86,665 complexes (we will use first 6000)
- large diversity of the TM–organic chemical space
- variety of organic ligands bound to **thirty** 3d, 4d, and 5d TMs from groups 3 to 12
- Cartesian coordinates optimized at the GFN2-xTB level
- Quantum properties computed at the DFT(TPSSh-D3BJ/def2-SVP) level
 - (i) the electronic and dispersion energies
 - (ii) metal center natural charge
 - (iii) HOMO/LUMO energies and gap
 - (iv) dipole moment
 - (v) polarizabilities

CSD = Cambridge Structural Database
1.25M accurate 3D structures with data from X-ray and neutron diffraction

CCS = chemical compound space

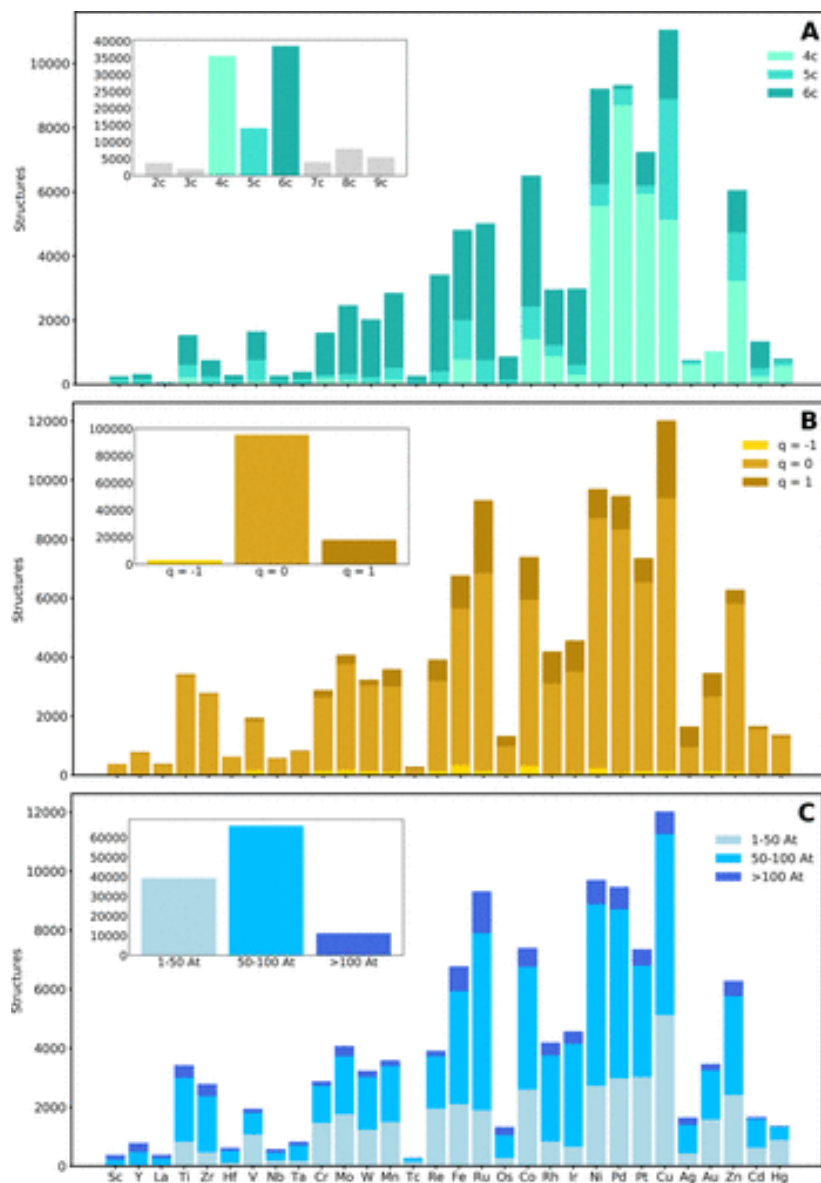
“The pairwise representations of the properties revealed unusual regions within the CCS, for example, TM complexes with large polarizabilities and wide HOMO/LUMO gaps”

**Goal: What predictive models, new descriptors and correlations can you find?
Can we interpret the results and learn new chemistry trends?**

Data set construction: Apply filters to CSD

1. Composition filter (metal elements): Excluded all structures except those containing a single transition metal (TM)
2. Composition filter (non-metal elements): Excluded all structures except those containing a minimum of one C and one H atoms. The other allowed elements: B, Si, N, P, As, O, S, Se, F, Cl, Br and I.
3. Components filter: Excluded the structure of all molecular components, except that of the metal complex (remove solvent and counterions)
4. Polymers filter: Excluded all polymeric structures.
5. Spatial coordinates filter: Excluded all structures without 3D-coordinates.
6. Disorder filter: Excluded all structures with disordered atoms.
7. Charge filter: Excluded all structures with charge higher than 1 and lower than -1

Result 116,332 structures



Distributions over the 3–5d TM series by

(A) metal node degree (MND), number of bonds to the metal

(B) molecular charge q

(C) size in number of atoms

More distributions in the paper

Using experimental structures has benefits: chemists know how to make them!

Using quantum chemistry codes optimize geometry in gas phase: apply filters to QM data

1. Calculation did not converge
 2. Computed geometry is too different from CSD (7%)
 3. Exclude species odd number of electrons
-

4. Result 88,699 molecules

Do more accurate(expensive) property QM calcs

1. Electronic and dispersion energies
 2. HOMO and LUMO energies
 3. HOMO-LUMO gap
 4. Dipole moment
 5. Metal center charge
 6. Polarizability
-

Result 88,665 molecules

Computed properties tmQM_y_tiny.csv

1	CSD_code;Electronic_E;Dispersion_E;Dipole_M;Metal_q;HL_Gap;HOMO_Energy;LUMO_Energy;Polarizability
2	WIXKOE; -2045.524942; -0.239239; 4.233300; 2.109340; 0.131080; -0.162040; -0.030960; 598.457913
3	DUCVIG; -2430.690317; -0.082134; 11.754400; 0.759940; 0.124930; -0.243580; -0.118650; 277.750698
4	KINJOG; -3467.923206; -0.137954; 8.301700; 1.766500; 0.140140; -0.236460; -0.096320; 393.442545
5	GEKVEC; -3657.137747; -0.073924; 3.044800; 1.171860; 0.138650; -0.267650; -0.129000; 266.725736
6	PIBNEV; -1184.911899; -0.132369; 2.776000; 1.926420; 0.106410; -0.151640; -0.045230; 342.341585
7	ILOJOK; -3314.579807; -0.287378; 2.947000; 1.777290; 0.092470; -0.175080; -0.082610; 726.729174

Add your own ML descriptors (analyze .xyz, use 'mendelev')

Bonds to the metal:

TM size (we've seen several radii)

bond lengths (short or long? Scale by TM radius? you decide)

number of bonds

nearest neighbor atom type

Whole structure:

size, shape, symmetry

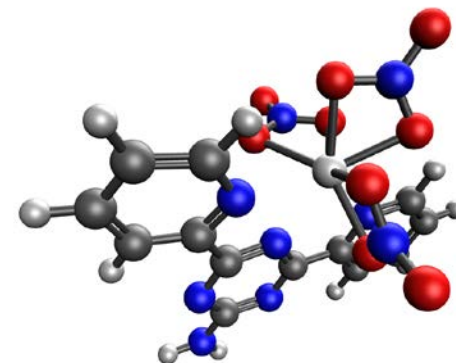
Metal identity:

number, group, period

Electronegativity, electron affinity, ionization energy

Everything can be converted to numbers!

Use *tiny* files to test your codes



Chemical hardness

$$\eta = (E_{\text{homo}} + E_{\text{lumo}})/2$$

can be easily added to .csv

Try η as your prediction target