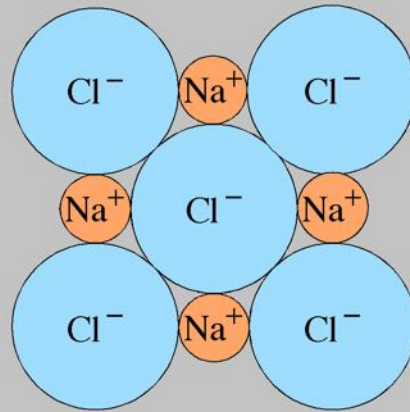
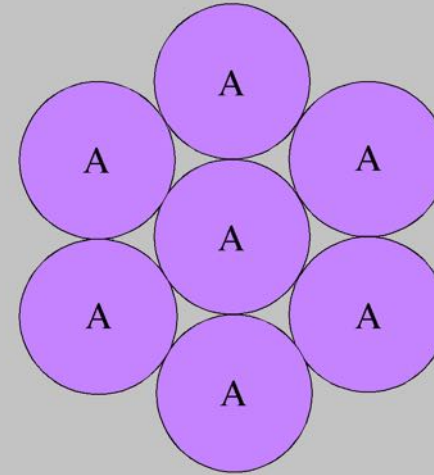


The image shows a complex, interconnected network of polygons representing a crystal lattice. The polygons are colored in shades of blue, grey, and white. A central text box with a yellow background and black border contains the text 'III. Bindungen im Kristall'. The overall background is a light blue gradient.

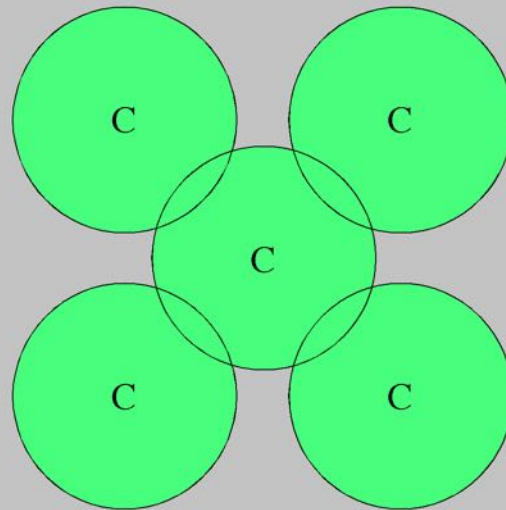
III. Bindungen im Kristall



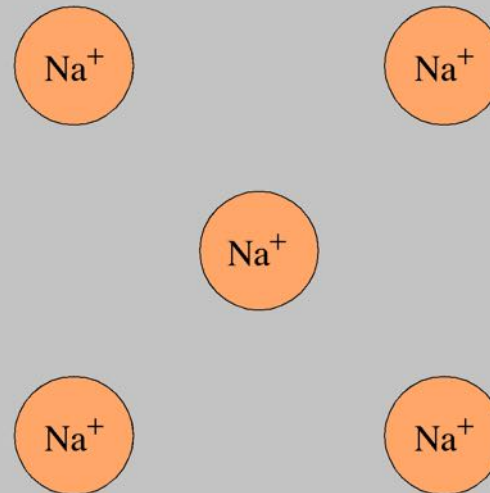
Natriumchlorid
(ionisch)



Kristallines Argon
(van der Waals)



Diamant
(kovalent)



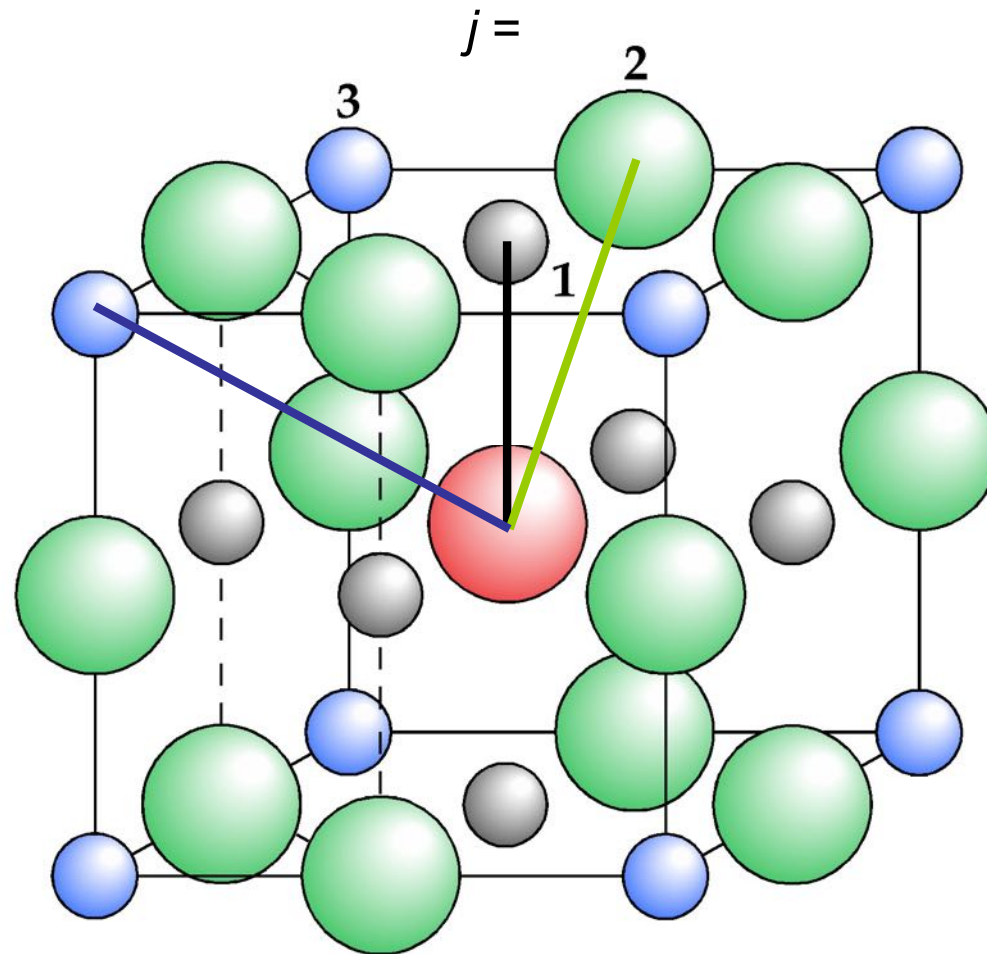
Natrium
(metallisch)

Bindungstyp	Beispiel	Bindungsenergie (eV)
Ionisch	NaCl	8.23
	LiF	10.92
Van-der-Waals	Ar	0.080
	Kr	0.116
Kovalent	Diamant	7.36
	Si	4.64
Metallisch	Na	1.13
	Fe	4.29
	W	8.66
Wasserstoff-Brücken	H ₂ O	0.52
	HF	0.30

Bindungsenergien

Angegeben ist die pro Atom notwendige Energie, um aus einem Festkörper bei 0K und 1 atm freie, neutrale Atome in ihrem Grundzustand zu bilden. Die Daten wurden von Prof. Leo Brewer in der Einheit kcal/mol angegeben, nach dem LBL-Report 3720 vom 4.Mai 1977.

Li 158. 1.63 37.7	Be 320. 3.32 76.5											B 561 5.81 134	C 711. 7.37 170.	N 474. 4.92 113.4	O 251. 2.60 60.03	F 81.0 0.84 19.37	Ne 1.92 0.020 0.46																												
Na 107. 1.113 25.67	Mg 145. 1.51 34.7	\longleftrightarrow kJ/mol \longleftrightarrow \longleftrightarrow eV/atom \longleftrightarrow \longleftrightarrow kcal/mol \longleftrightarrow										Al 327. 3.39 78.1	Si 446. 4.63 106.7	P 331. 3.43 79.16	S 275. 2.85 65.75	Cl 135. 1.40 32.2	Ar 7.74 0.080 1.85																												
K 90.1 0.934 21.54	Ca 178. 1.84 42.5	Sc 376 3.90 89.9	Ti 468. 4.85 111.8	V 512. 5.31 122.4	Cr 395. 4.10 94.5	Mn 282. 2.92 67.4	Fe 413. 4.28 98.7	Co 424. 4.39 101.3	Ni 428. 4.44 102.4	Cu 336. 3.49 80.4	Zn 130 1.35 31.04	Ga 271. 2.81 64.8	Ge 372. 3.85 88.8	As 285.3 2.96 68.2	Se 237 2.46 56.7	Br 118. 1.22 28.18	Kr 11.2 0.116 2.68																												
Rb 82.2 0.852 19.64	Sr 166. 1.72 39.7	Y 422. 4.37 100.8	Zr 603. 6.25 144.2	Nb 730. 7.57 174.5	Mo 658. 6.82 157.2	Tc 661. 6.85 158.	Ru 650. 6.74 155.4	Rh 554. 5.75 132.5	Pd 376. 3.89 89.8	Ag 284. 2.95 68.0	Cd 112. 1.16 26.73	In 243. 2.52 58.1	Sn 303. 3.14 72.4	Sb 265. 2.75 63.4	Te 211. 2.19 50.34	I 107. 1.11 25.62	Xe 15.9 0.16 3.80																												
Cs 77.6 0.804 18.54	Ba 183. 1.90 43.7	La 431. 4.47 103.1	Hf 621. 6.44 148.4	Ta 782. 8.10 186.9	W 859. 8.90 205.2	Re 775. 8.03 185.2	Os 788. 8.17 188.4	Ir 670. 6.94 160.1	Pt 564. 5.84 134.7	Au 368. 3.81 87.96	Hg 65. 0.67 15.5	Tl 182. 1.88 43.4	Pb 196. 2.03 46.78	Bi 210. 2.18 50.2	Po 144. 1.50 34.5	At	Rn 19.5 0.202 4.66																												
Fr	Ra 160. 1.66 38.2	Ac 410. 4.25 98.	<table border="1" style="width: 100%; border-collapse: collapse; text-align: center;"> <tbody> <tr> <td>Ce 417. 4.32 99.7</td> <td>Pr 357. 3.70 85.3</td> <td>Nd 328. 3.40 78.5</td> <td>Pm</td> <td>Sm 206. 2.14 49.3</td> <td>Eu 179. 1.86 42.8</td> <td>Gd 400. 4.14 95.5</td> <td>Tb 391. 4.05 93.4</td> <td>Dy 294. 3.04 70.2</td> <td>Ho 302. 3.14 72.3</td> <td>Er 317. 3.29 75.8</td> <td>Tm 233. 2.42 55.8</td> <td>Yb 154. 1.60 37.1</td> <td>Lu 428. 4.43 102.2</td> </tr> <tr> <td>Th 598. 6.20 142.9</td> <td>Pa</td> <td>U 536. 5.55 128.</td> <td>Np 456 4.73 109.</td> <td>Pu 347. 3.60 83.0</td> <td>Am 264. 2.73 63.</td> <td>Cm 385 3.99 92.1</td> <td>Bk</td> <td>Cf</td> <td>Es</td> <td>Fm</td> <td>Md</td> <td>No</td> <td>Lr</td> </tr> </tbody> </table>															Ce 417. 4.32 99.7	Pr 357. 3.70 85.3	Nd 328. 3.40 78.5	Pm	Sm 206. 2.14 49.3	Eu 179. 1.86 42.8	Gd 400. 4.14 95.5	Tb 391. 4.05 93.4	Dy 294. 3.04 70.2	Ho 302. 3.14 72.3	Er 317. 3.29 75.8	Tm 233. 2.42 55.8	Yb 154. 1.60 37.1	Lu 428. 4.43 102.2	Th 598. 6.20 142.9	Pa	U 536. 5.55 128.	Np 456 4.73 109.	Pu 347. 3.60 83.0	Am 264. 2.73 63.	Cm 385 3.99 92.1	Bk	Cf	Es	Fm	Md	No	Lr
Ce 417. 4.32 99.7	Pr 357. 3.70 85.3	Nd 328. 3.40 78.5																Pm	Sm 206. 2.14 49.3	Eu 179. 1.86 42.8	Gd 400. 4.14 95.5	Tb 391. 4.05 93.4	Dy 294. 3.04 70.2	Ho 302. 3.14 72.3	Er 317. 3.29 75.8	Tm 233. 2.42 55.8	Yb 154. 1.60 37.1	Lu 428. 4.43 102.2																	
Th 598. 6.20 142.9	Pa	U 536. 5.55 128.																Np 456 4.73 109.	Pu 347. 3.60 83.0	Am 264. 2.73 63.	Cm 385 3.99 92.1	Bk	Cf	Es	Fm	Md	No	Lr																	

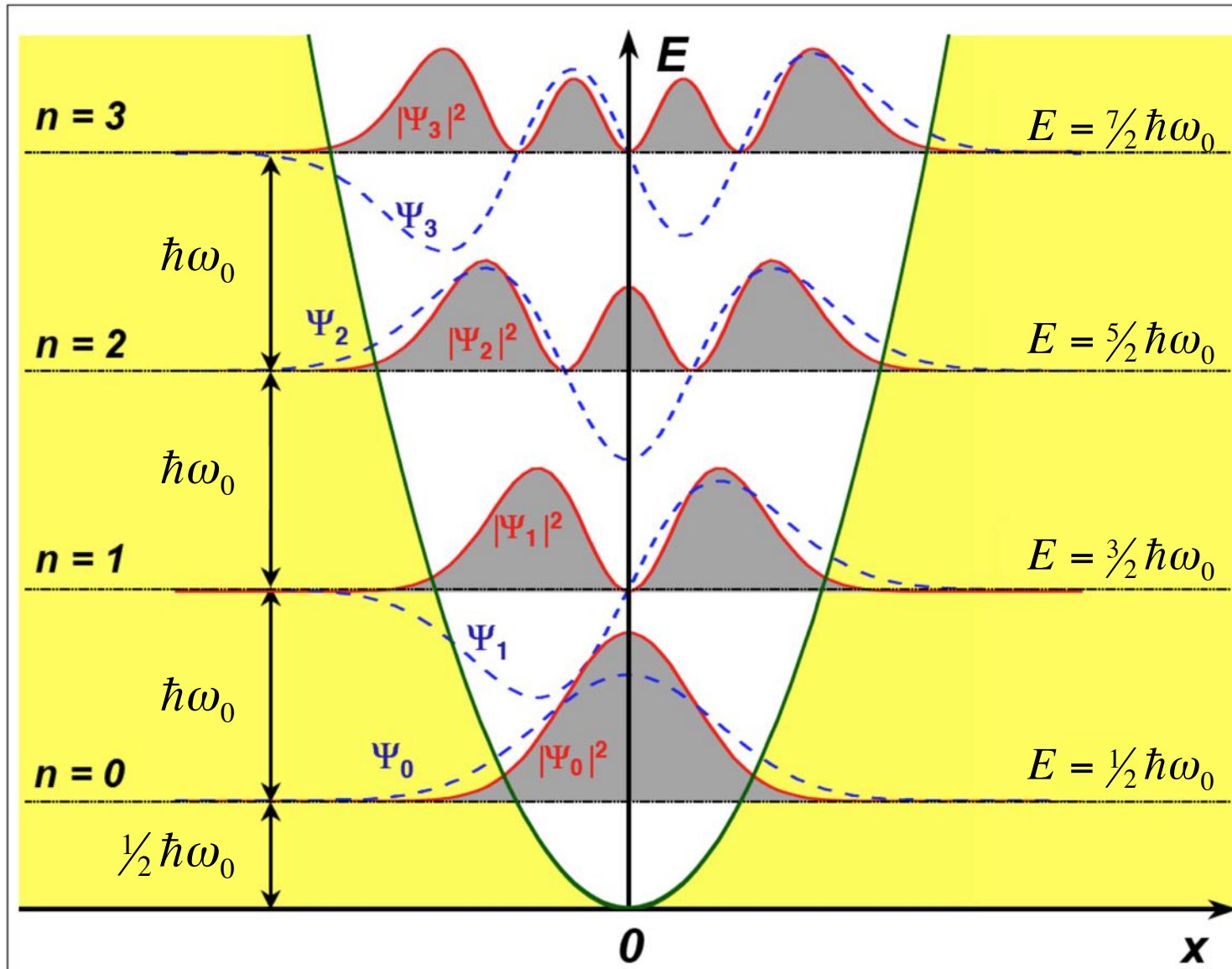


NaCl Struktur mit nächsten, übernächsten und über-übernächsten Nachbarn

$$\alpha = - \sum_j \frac{\pm 1}{p_j} = \frac{6}{1} - \frac{12}{\sqrt{2}} + \frac{8}{\sqrt{3}} - \dots = 1.748$$

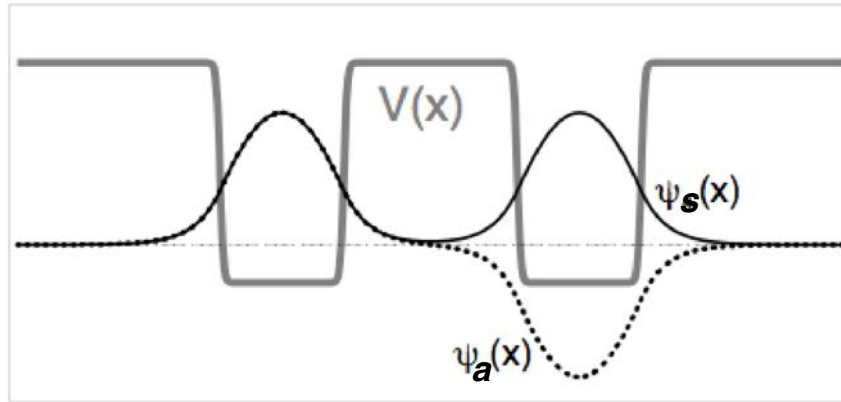
Strukturtyp	Beispiel	α
NaCl	NaCl, AgBr, EuS	1.748
CsCl	CsCl, CsBr, TlBr	1.763
Zinkblende ZnS	ZnS, CuCl, GaAs	1.638
Fluorit CaF ₂	CaF ₂ , LaH ₂ , VO ₂	5.039
Korund Al ₂ O ₃	Al ₂ O ₃ , V ₂ O ₃ , Cr ₂ O ₃	25.031
Perovskit CaTiO ₃	CaTiO ₃ , BaTiO ₃	12.377

1D- quantenmechanischer harmonischer Oszillator mit ω_0

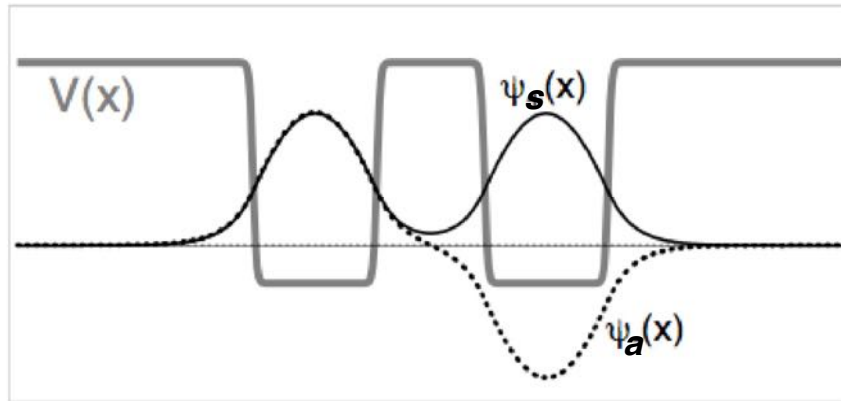


"Topfmolekül"

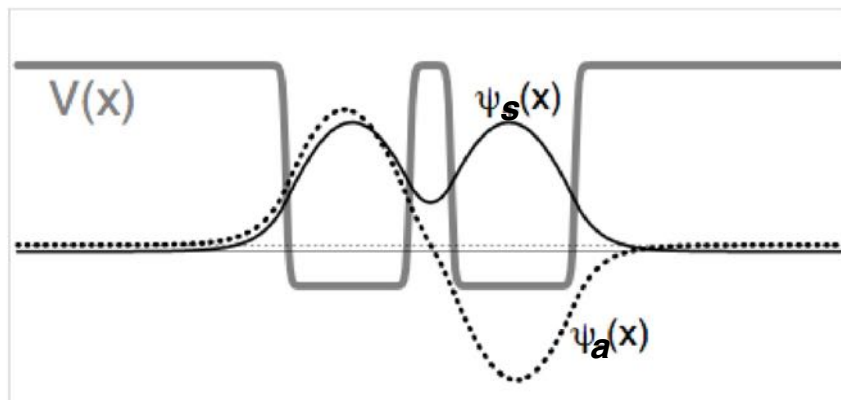
a)



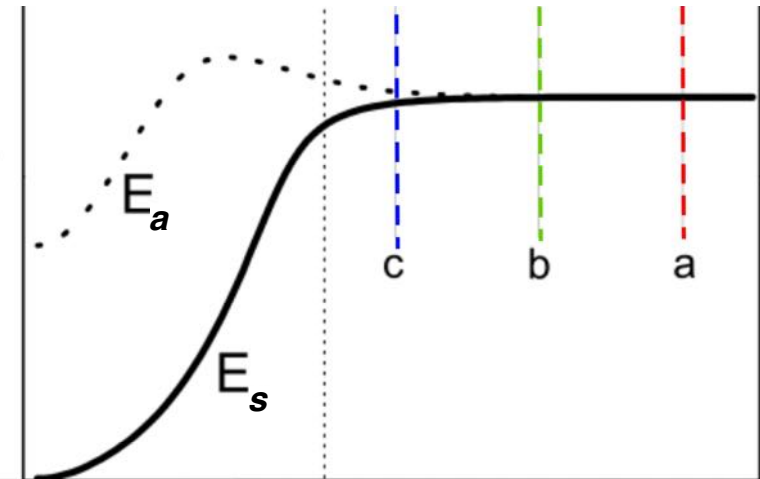
b)



c)

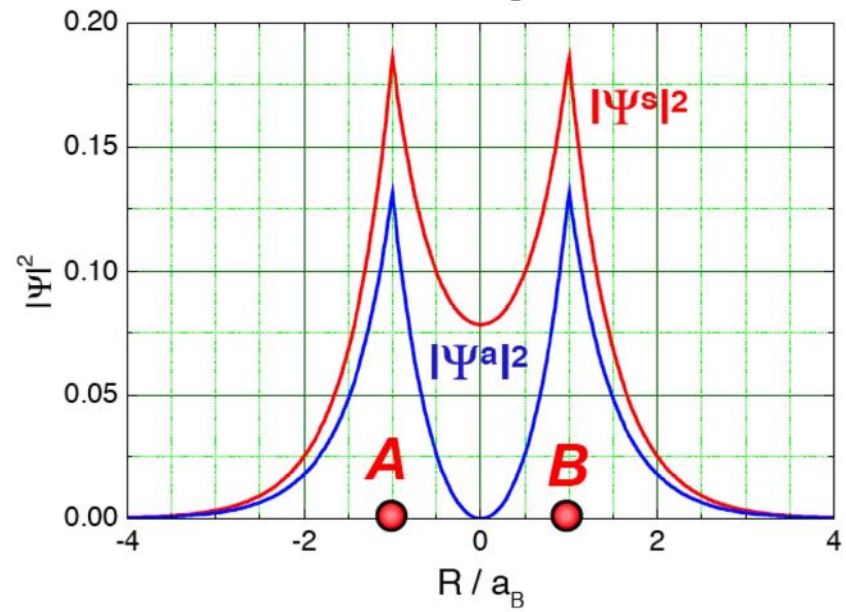
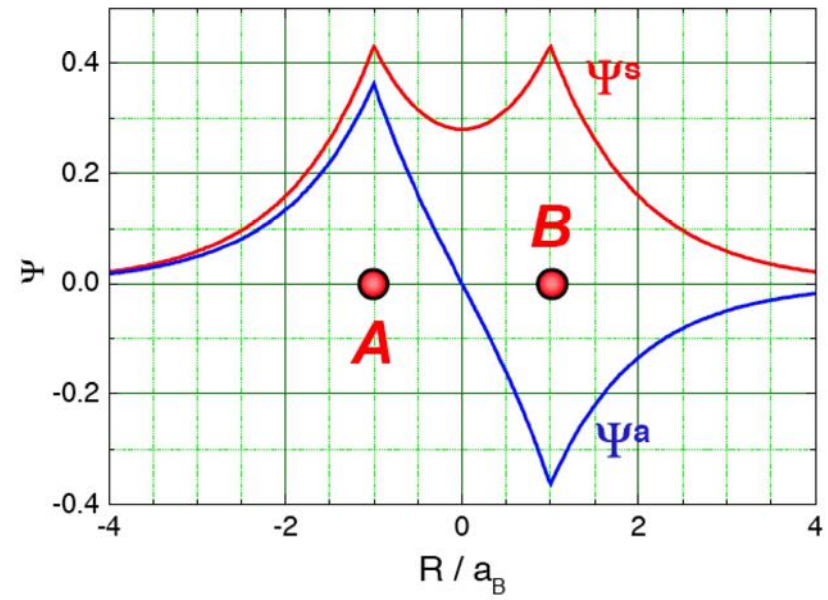


Energieniveaus

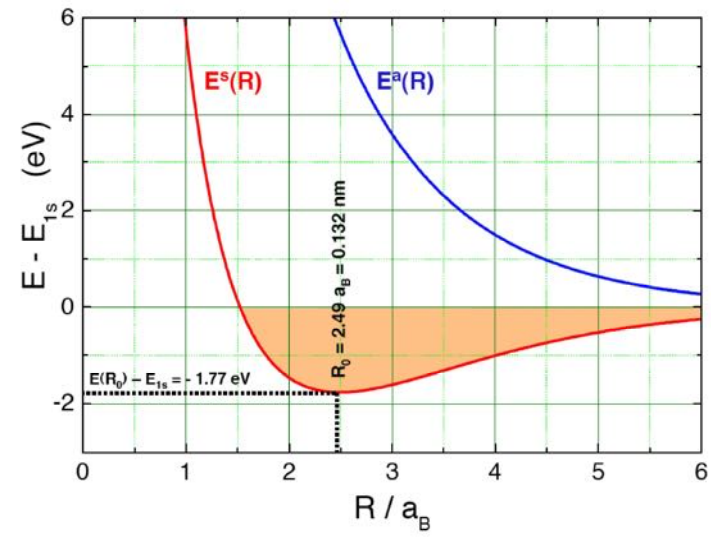


Topfabstand

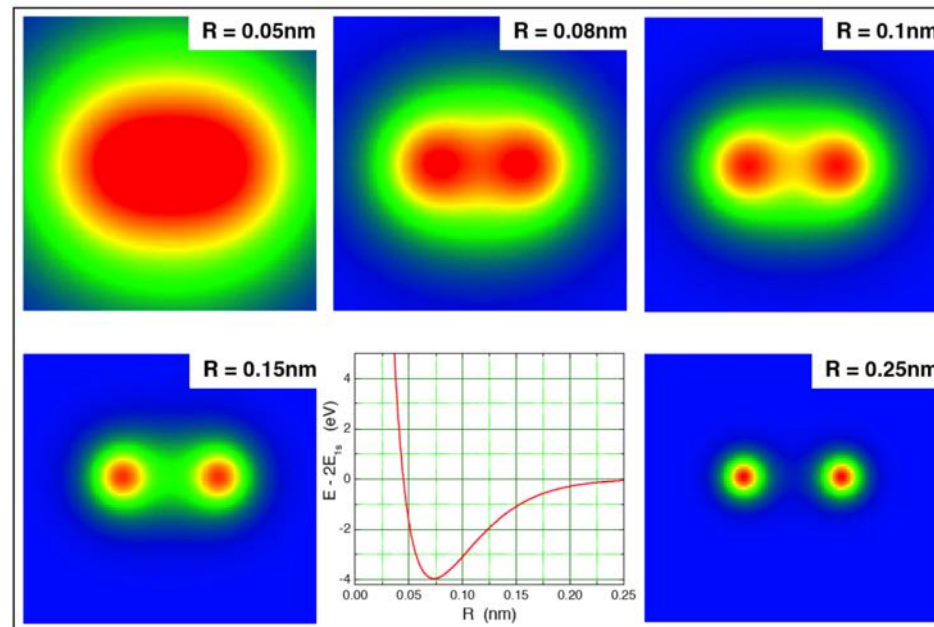
H₂⁺ Ion

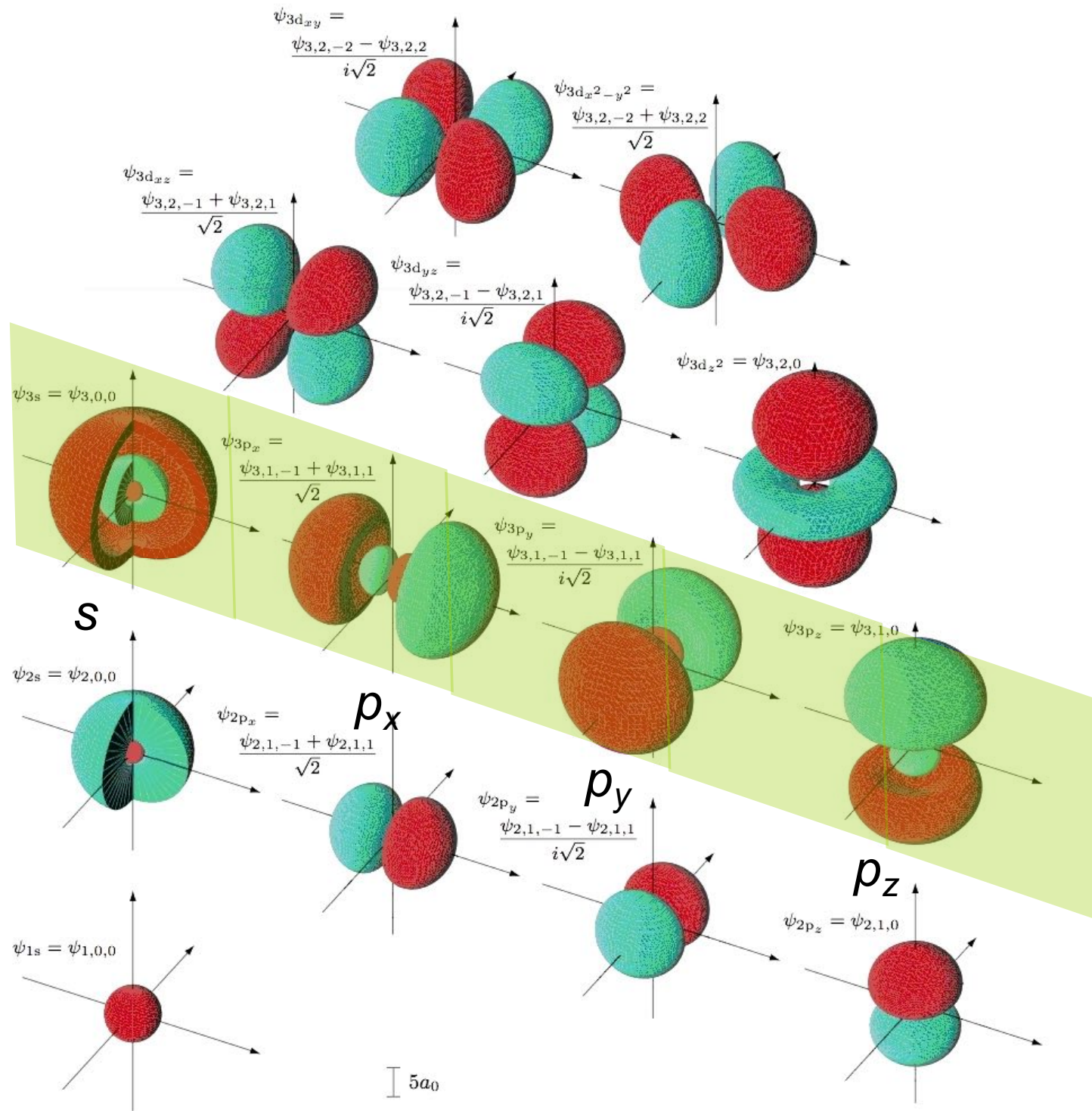


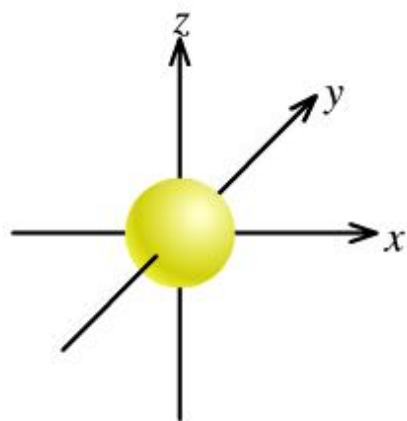
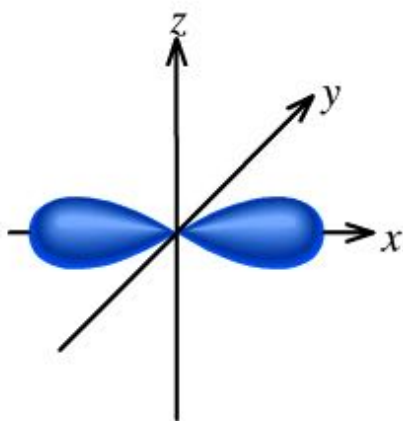
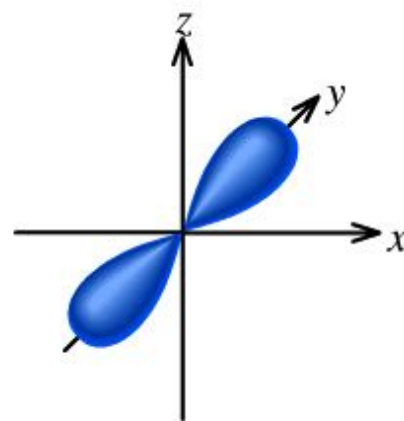
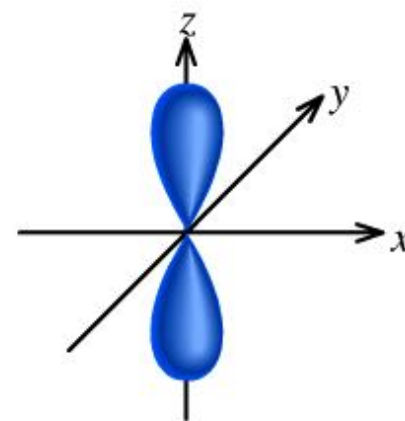
H₂⁺-Molekül



H₂-Molekül






 ψ_s

 ψ_{px}

 ψ_{py}

 ψ_{pz}

 $\psi_{sp^3_1}$

 $\psi_{sp^3_2}$

 $\psi_{sp^3_3}$

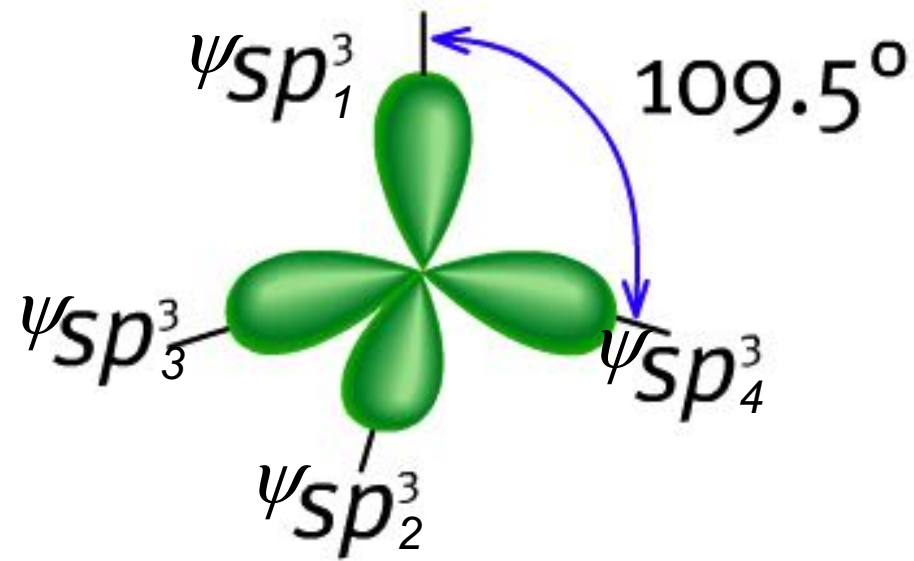
 $\psi_{sp^3_4}$

$$\frac{1}{2}(\psi_s + \psi_{px} + \psi_{py} + \psi_{pz})$$

$$\frac{1}{2}(\psi_s + \psi_{px} - \psi_{py} - \psi_{pz})$$

$$\frac{1}{2}(\psi_s - \psi_{px} + \psi_{py} - \psi_{pz})$$

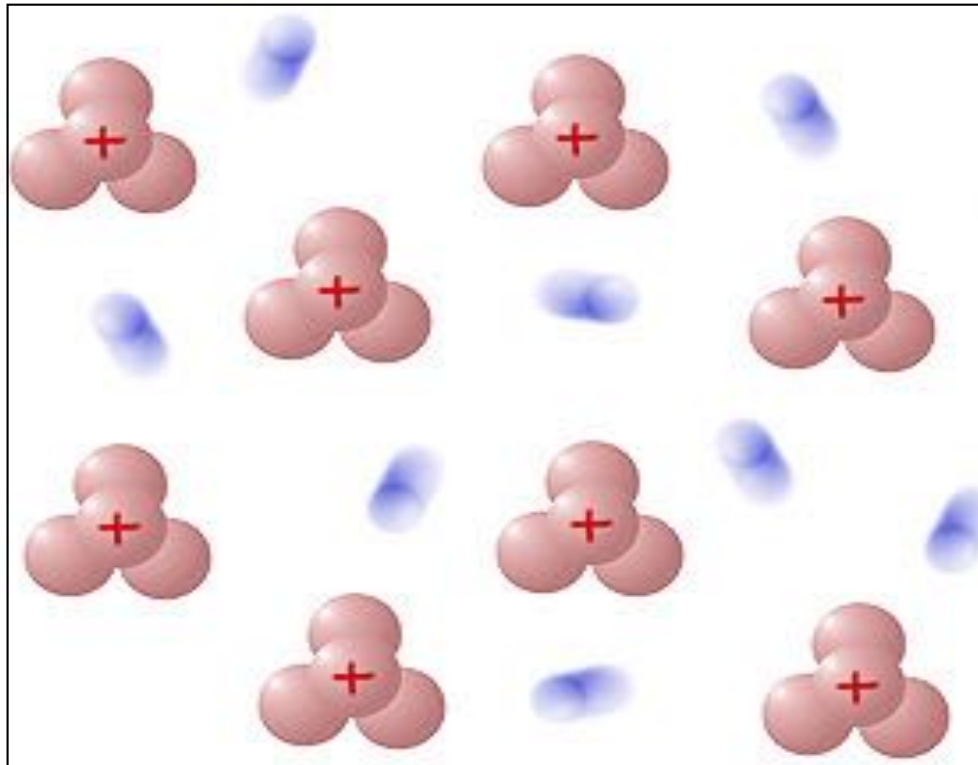
$$\frac{1}{2}(\psi_s - \psi_{px} - \psi_{py} + \psi_{pz})$$



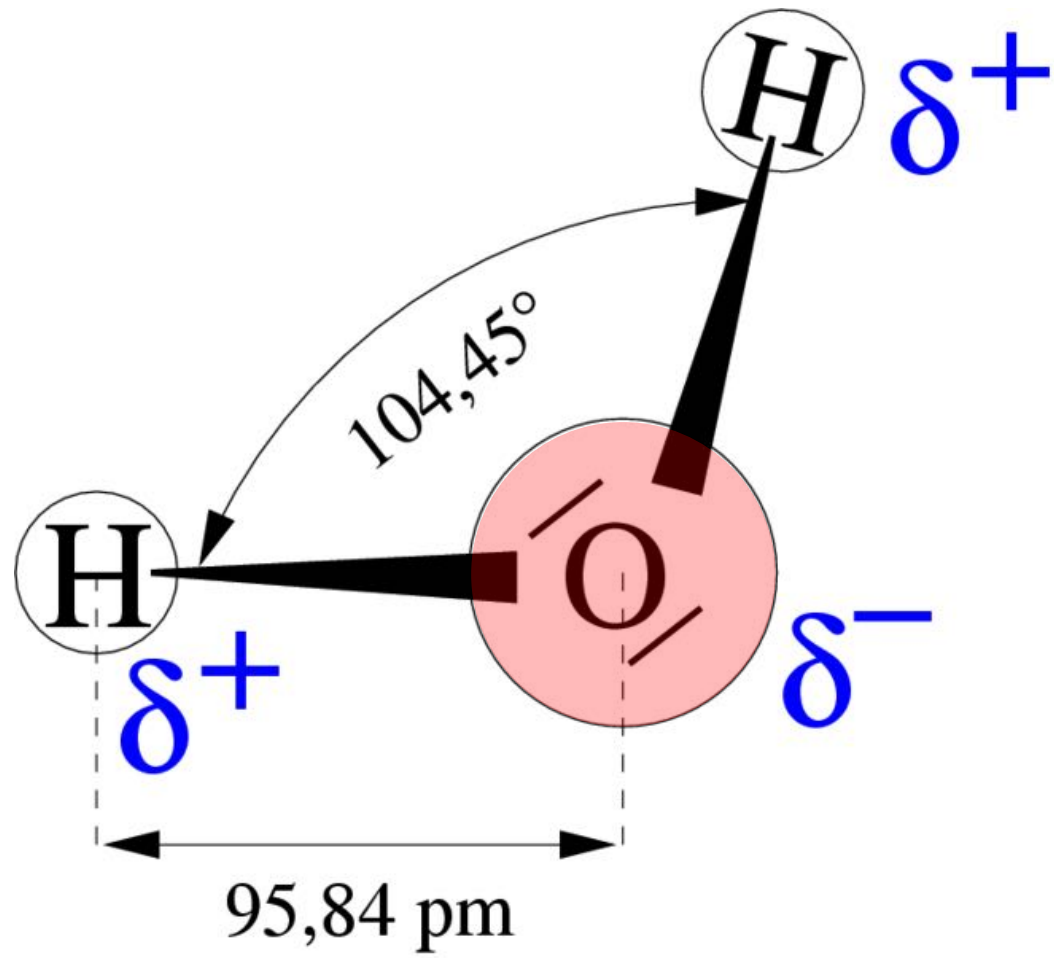
Tetraeder

Einfaches Modell eines Metalles

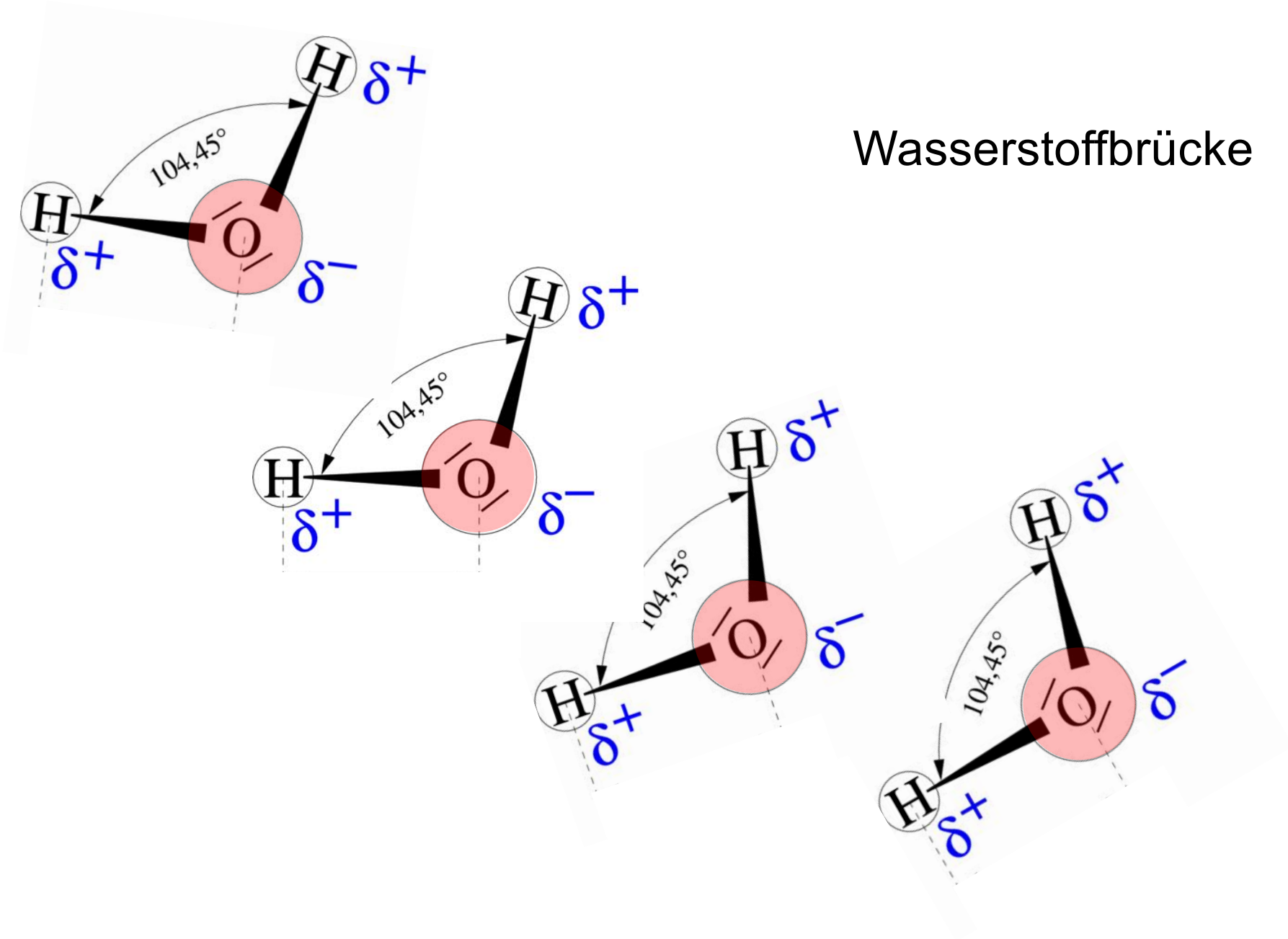
Atomrümpfe

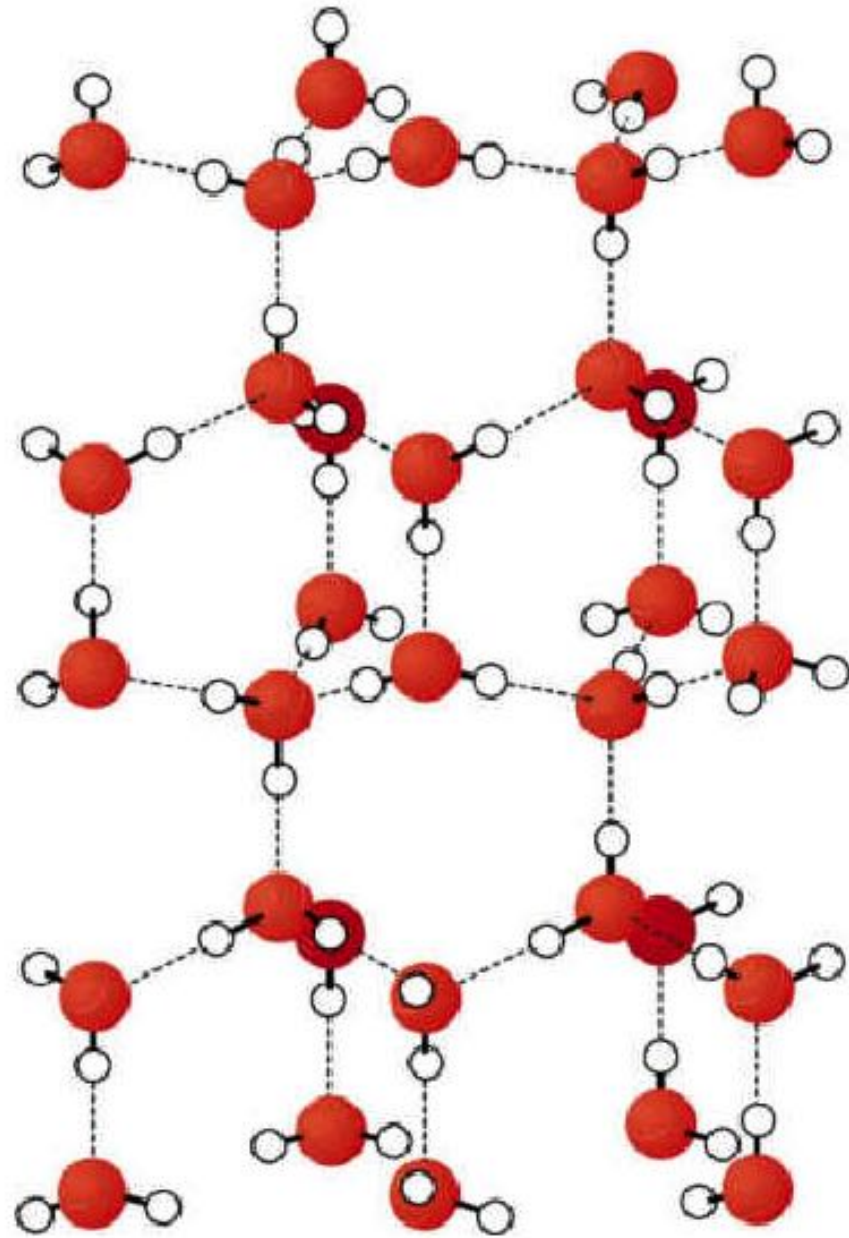


delokalisierte
äussere Elektronen



Wasserstoffbrücke





Wassereis