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# Nature of Alkali- and Coinage-Metal Bonds versus Hydrogen Bonds

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Abstract: We have quantum chemically studied the structure and nature of alkali- and coinage-metal bonds (M-bonds) versus that of hydrogen bonds between A-M and B- in archetypal [A-M-B] model systems (A, B=F, Cl and M=H, Li, Na, Cu, Ag, Au), using relativistic density functional theory at ZORA-BP86-D3/TZ2P. We find that coinage-metal bonds are stronger than alkali-metal bonds which are stronger than the corresponding hydrogen bonds. Our main purpose is to

understand how and why the structure, stability and nature of such bonds are affected if the monovalent central atom H of hydrogen bonds is replaced by an isoelectronic alkali- or coinage-metal atom. To this end, we have analyzed the bonds between A-M and B- using the activation strain model, quantitative Kohn-Sham molecular orbital (MO) theory, energy decomposition analysis (EDA), and Voronoi deformation density (VDD) analysis of the charge distribution.

#### 1. Introduction

Intermolecular interactions have been an active topic in research for many years as they are cornerstones in supramolecular chemistry.[1] Although intermolecular interactions are generally weak compared to intramolecular interactions (as shown by the lower binding energies and large equilibrium distances of the formers), accumulation of many such weak interactions may lead to strong bonding in biomacromolecules such as DNA and proteins, thus pointing out its relevance also in biochemical processes.[2]

Among the various types of intermolecular interactions, the hydrogen bond (H-bond) has received particular attention.[3] Hydrogen bonding in complexes A-H---B are stabilizing interactions between a hydrogen-bond donor, i.e., a Lewis-acidic element-hydrogen (A-H) bond, and an electron-rich atom of the hydrogen bond acceptor (B), the electron-rich site of a Lewis base. Quantum-chemical bonding analyses have shown that the stabilization associated with the H-bond emerges not only from electrostatic interactions but also from substantial HOMO-LUMO orbital interactions that go with charge transfer from the lone-pair orbital of B to the  $\sigma^*_{AH}$  orbital of A–H.<sup>[4]</sup> Proceeding from the H-bond, one can replace the central hydrogen atom by a more electronegative halogen atom which leads us to the halogen bond (X-bond) in complexes A-X...B. This type of bond is relevant, e.g., in ligand-protein interactions.<sup>[5]</sup> In the framework of quantitative molecular orbital (MO) theory, the X-bond has been shown to possess a strong covalent character provided by HOMO-LUMO interactions that are very similar in nature to those occurring in Hbonds. [6] Valence bond (VB) theory studies have identified a number of X-bonds as charge-shift bonds.[7]

In this work, we wish to analyze the nature of a less explored, yet related type of intermolecular interaction: the metal bond (M-bond) in A–M···B:

$$A-M+B \to A-M \cdots B \tag{1}$$

Here, proceeding for example from an H-bond, the central hydrogen atom has been replaced by a more electropositive, monovalent metal atom M. The first prediction of an M-bond, namely, a lithium bond, was made by Kollman et al.[8] in 1970 and experimentally verified few years later. [9] Nevertheless, the Li-bond, and M-bond in general, are much less understood than the H- and X-bond.[10]

Herein, we aim to undertake a more systematic exploration of M-bonds, their structure and stability. Our purpose is in particular to understand their nature and bonding mechanism and to compare them with H- and X-bonds. To this end, we have quantum-chemically analyzed the structure and bonding of alkali- and coinage-metal bonds versus the hydrogen bonds between A-M and B<sup>-</sup> in archetypal [A-M···B]<sup>-</sup> model systems with A, B=F, Cl and M=H, Li, Na, Cu, Ag, Au, using relativistic density functional theory at ZORA-BP86-D3/TZ2P as implemented in the ADF program.[11] Our analyses are based on the activation strain model (ASM) of chemical reactivity<sup>[12]</sup> in

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conjunction with quantitative Kohn-Sham molecular orbital (MO) theory and a canonical energy decomposition analysis (EDA).<sup>[13]</sup> Moreover, we have analyzed how the bonding affects the charge density distribution, using the Voronoi deformation density (VDD)<sup>[14]</sup> method.

# 2. Theoretical Methods

#### 2.1. Computational details

All DFT calculations were carried out by using the Amsterdam Density Functional (ADF) program. [11] The molecular orbitals (MOs) were expanded in a large uncontracted set of Slater-type orbitals (STOs), TZ2P, which is of triple- $\zeta$  quality and has been augmented with two polarization function sets: 2p and 3d on hydrogen, 3d and 4f on fluorine and chlorine.

Equilibrium structures were obtained by optimizations using analytical gradient techniques.<sup>[11]</sup> Geometries and energies were calculated at the BP86 level of the generalized gradient approximation (GGA).<sup>[15]</sup> This level of theory has been shown of adequate accuracy for the treatment of similar three-atom systems such as Halogen bonding interactions (X-bond).<sup>[7]</sup> Relativistic effects were accounted for by using the zero<sup>th</sup>-order regular approximation (ZORA).<sup>[16]</sup> Dispersion corrections were included by means of Grimme's D3 model.<sup>[17]</sup> All the minima were characterized by harmonic vibrational analysis, verifying that showed only positive definite Hessians. Thermal corrections computed at 298.15 K were found to have almost no influence on the observed trends, therefore, were not included in the discussion for clarity.

# 2.2. Activation Strain Analysis

The formation of the  $D_{\infty h}$ -symmetric  $[A-M\cdots B]^-$  complex is associated with the occurrence of the new  $M\cdots B$  bond between A-M and  $B^-$  and a simultaneous stretch of the already existing A-M bond in the neutral A-M molecule (for M=H, Li, see e.g. Ref. 18). This association process has been explored by analyzing the equilibrium geometry of  $[A-M\cdots B]^-$  complex and a fictitious structure in which the A-M fragment remains as in an isolated A-M diatomic i. e. without stretching, while the  $M\cdots B$  distance is as in the equilibrium geometry of the real  $[A-M\cdots B]^-$ . [19]

The complexation reactions were analyzed using the activation strain model (ASM) for the two above-mentioned situations. Within this framework, the bond energy  $\Delta E$  associated with forming the hydrogen bond or metal bond is decomposed according to Equation 2:

$$\Delta E = \Delta E_{\text{strain}} + \Delta E_{\text{int}} \tag{2}$$

Herein,  $\Delta E_{\text{strain}}$  and  $\Delta E_{\text{int}}$  correspond to the strain and interaction energy, respectively. The strain energy  $\Delta E_{\text{strain}}$  is associated with the required energy to deform the reactants from their equilibrium geometry to the geometry they adopt in

the hydrogen- or metal-bonded complex. This term depends on the extent of deformation, caused by the interaction, as well as the rigidity of the molecules involved in the association reaction. In general, the strain is positive (destabilizing). On the other hand, the interaction term  $\Delta E_{\rm int}$  depends on the electronic structure of the associating molecules and on how they approach each other. The interaction can be further analyzed within Kohn-Sham MO conceptual framework according to our canonical energy decomposition analysis (EDA)  $^{[13]}$  as

$$\Delta E_{\text{int}} = \Delta V_{\text{elstat}} + \Delta E_{\text{Pauli}} + \Delta E_{\text{oi}} + \Delta E_{\text{disp}}$$
(3)

Herein,  $\Delta V_{\rm elstat}$  is the classical Coulombic interaction between the unperturbed charge distribution of each of the two reactants. The term  $\Delta E_{\rm Pauli}$  is the Pauli repulsions between occupied orbitals of the two fragments and is responsible for steric repulsion. The term  $\Delta E_{\rm oi}$  stands for the stabilizing orbital interaction energy, including charge transfer (such as HOMO-LUMO interactions). Finally,  $\Delta E_{\rm disp}$  constitutes a correction for dispersion interactions.

#### 2.3. Voronoi Deformation Density (VDD) Analysis

The electron density distribution is analyzed using the Voronoi deformation density (VDD) method for computing atomic charges. The VDD atomic charge ( $Q_A$ ) is computed as the numerical integral of the deformation density in the volume of the Voronoi cell of atom A, that is, the compartment of space bounded by the bond midplanes on and perpendicular to all bond axes between nucleus A and its neighboring nuclei (Eq. 4).

$$Q_{A} = \int \left[ \rho(r) - \sum_{B} \rho_{B}(r) \right] dr$$
 (4)

Here,  $\rho(r)$  is the electronic density of the molecule and  $\sum_{\mathcal{B}} \rho_{\mathcal{B}}(r)$  is the superposition of atomic densities  $\rho_{\mathcal{B}}(r)$  of a fictitious promolecule without chemical interactions where all atoms are considered neutral. Therefore,  $Q_A$  directly measures the charge that flows out of  $(Q_A > 0)$  or into  $(Q_A < 0)$  the Voronoi cell of atom A due to chemical interactions.

### 3. Results and Discussion

#### 3.1. Trends in Structure and Stability

The results of our ZORA-BP86-D3/TZ2P calculations for the selected  $[A-M\cdots B]^-$  (X=F, Cl; M=H, Li, Na, Cu, Ag, Au) complexes are collected in Table 1. Full structural details of the optimized structures and thermal corrections can be found in the Supporting Information.

All  $[A-M\cdots B]^-$  complexes form spontaneously, without a barrier, on a single-well potential energy surface. Upon formation of the new B···M bond, the A–M bond of the original diatomic expands (vide infra), thus leading to a  $D_{\infty h}$ -symmetric





Table 1. E $\langle \sigma_{\text{AM}}^*   \text{n} p_z \rangle$	Sond leng (B <sup>−</sup> )) in c	gths d <sub>AM</sub> (	In A), stretc	h of the A– ious <sup>[b]</sup> [A–A	Table 1. Bond lengths $d_{AM}$ (in A), stretch of the A–M fragment $\Delta d_{AM}$ (in $\langle \sigma_{AM}^{-1}   n \rho_{Z}(B^{-}) \rangle$ in optimized $b^{ 0}$ and fictitious $^{ 0 }$ [A–M····B] complexes.	-	A), AM···B¯ intera	action energy o	decompositi	on analysis (ir	ı kcal mol⁻'),	VDD charge a	ccumulation	ΔQ <sub>AM</sub> of tra	agment AM (i	interaction energy decomposition analysis (in Kcal mol $^-$ ), VDD charge accumulation $\Delta Q_{_{\Delta M}}$ of fragment AM (in au), and overlap
A–M	<u>-</u> 8	q	ΔЧ	q	ΛE	$\Delta E_{ m strain}$	ΔEint	AVektat	$\Delta E_{ m Pauli}$		7	ΔE <sub>oi</sub>		ΔE <sub>disp</sub>	ΔQ <sub>AM</sub>	$\langle \sigma_{AM}^*   np_z(B^-) \rangle$
				•						total	Q	В	ø	t		
F.F.	Ŀ	1.16	0.23	1.16	-53.2	19.8	-73.3	-76.3	68.1	-64.9	-58.0	6.9	1	-0.2	-0.19	0.27
							(-37.8)	(-72.9)	(83.1)	(-47.8)	(-41.5)	(-6.4)		(-0.1)	(-0.11)	(0.16)
ī		1.70	0.12	1.70	-73.7	1.8	-75.6	-87.2	25.8	-14.2	-7.2	-7.5	1	-0.1	-0.06	0.11
							(-72.0)	(-83.9)	(25.9)	(-13.9)	(-6.9)	(-6.9)		(-0.1)	(-0.06)	(0.10)
F–Na		2.06	0.11	5.06	-65.8	1.0	-66.9	-77.4	20.9	-10.3	-5.3	-4.9	ı	-0.1	-0.06	0.13
							(-64.8)	(-75.6)	(21.0)	(-10.0)	(-5.2)	(-4.9)		(-0.1)	(-0.06)	(0.13)
F_Cu		1.79	0.05	1.79	-96.9	9.0	-97.6	-124.3	71.6	-45.0	-32.7	-11.7	9.0-	-0.1	-0.20	0.26
							(-96.5)	(-123.1)	(20.8)	(-44.2)	(-32.0)	(-11.6)	(-0.6)	(-0.1)	(-0.20)	(0.25)
F–Ag		2.02	0.03	2.02	-81.8	0.1	-81.9	-105.7	9.79	-43.7	-33.9	-8.9	6.0—	-0.1	-0.23	0.24
							(-81.9)	(-105.1)	(86.8)	(-43.3)	(-33.5)	(-8.9)	(-0.9)	(-0.1)	(-0.23)	(0.24)
F-Au		1.99	0.04	1.99	-97.9	9.4	-98.3	-127.6	91.6	-62.3	-50.9	-10.4	-1.0	-0.1	-0.28	0.27
							(-97.6)	(-126.8)	(90.4)	(-61.1)	(-49.7)	(-10.4)	(-1.0)	(-0.1)	(-0.27)	(0.27)
Ŧ	<u>_</u>	1.01	0.08	1.86	-27.2	3.1	-30.4	-31.4	22.9	-21.5	-19.4	-2.1	ı	-0.3	-0.11	0.36
							(-24.5)	(-30.7)	(24.3)	(-17.8)	(-15.8)	(-2.0)		(-0.3)	(-0.09)	(0.34)
I		1.66	0.08	2.21	-49.2	1.1	-50.1	-55.6	16.0	-10.2	-6.0	-4.2	ı	-0.2	-0.08	0.24
							(-48.4)	(-53.9)	(15.9)	(-9.8)	(-5.8)	(-4.0)		(-0.2)	(-0.08)	(0.23)
F–Na		2.04	0.09	2.55	-46.3	8.0	-47.1	-54.7	14.7	-6.7	-4.0	-2.7	ı	-0.3	-0.06	0.25
							(-45.7)	(-53.6)	(14.8)	(-6.5)	(-3.8)	(-2.7)		(-0.3)	(-0.06)	(0.25)
F_Cu		1.80	90:0	2.10	-76.8	8.0	-77.7	-110.6	74.4	-41.4	-29.6	-11.2	-0.5	-0.2	-0.21	0.39
							(-76.2)	(-109.1)	(74.0)	(-40.8)	(-29.1)	(-11.1)	(-0.5)	(-0.2)	(-0.20)	(0.39)
F-Ag		2.03	0.08	2.33	-66.3	0.2	-66.5	-104.2	74.4	-36.5	-28.3	-7.7	9.0—	-0.2	-0.22	0.37
							(-66.0)	(-103.2)	(73.5)	(-36.1)	(-27.9)	(-7.6)	(-0.6)	(-0.2)	(-0.22)	(0.37)
F–Au		2.00	0.05	2.28	-81.8	0.7	-82.6	-131.3	107.2	-58.3	-47.0	-10.6	-0.7	-0.2	-0.27	0.38
							(-81.5)	(-129.6)	(105.8)	(-57.2)	(-45.9)	(-10.6)	(-0.8)	(-0.2)	(-0.26)	(0.38)
H	Ū	1.59	0.30	1.59	-29.9	17.5	-48.0	-45.5	65.0	-67.1	-62.4	-4.6	ı	-0.5	-0.25	0.34
							(-17.3)	(-45.5)	(77.8)	(-49.2)	(-44.3)	(-4.9)		(-0.3)	(-0.15)	(0.21)
C-Li		2.17	0.14	2.17	-56.9	1.6	-58.5	-62.8	18.2	-13.5	-7.8	-5.7	ı	-0.4	-0.10	0.26
							(-55.6)	(-60.2)	(18.0)	(-13.0)	(-7.4)	(-5.6)		(-0.4)	(-0.09)	(0.25)
CI–Na		2.52	0.13	2.52	-51.8	1:1	-52.9	9.09-	16.6	9.8-	-5.1	-3.4	ı	-0.4	-0.17	0.26
							(-50.8)	(-58.9)	(16.7)	(-8.2)	(-4.9)	(-3.4)		(-0.4)	(-0.07)	(0.26)
D-Cr		2.11	0.07	2.11	-76.7	8.0	-77.5	-110.9	77.2	-43.5	-31.1	-12.1	-0.4	-0.4	-0.22	0.38
							(-42.8)	(-109.6)	(76.7)	(-42.8)	(-30.3)	(-12.1)	(-0.4)	(-0.4)	(-0.22)	(0.38)
Cl–Ag		2.35	0.05	2.35	-66.9	0.3	-67.3	-102.1	73.0	-37.8	-28.9	-8.3	-0.5	-0.4	-0.23	0.36
							(-66.7)	(-101.3)	(72.3)	(-37.3)	(-28.4)	(-8.4)	(-0.5)	(-0.4)	(-0.27)	(0.36)
Cl–Au		2.31	0.07	2.31	-77.5	6.0	-78.4	-125.5	105.8	-58.3	-47.0	-10.7	9.0-	4.0-	-0.29	0.37
							(-76.7)	(-124.2)	(104.7)	(-57.0)	(-45.5)	(-10.8)	(-0.6)	(-0.4)	(-0.28)	(0.37)
[a] Compu	ited at Z	ORA-BP86	-D3/TZ2P.	[b] Results	obtained for	a fictitious	A-M…B] str	acture in whicl	h the A–M f	ragment is in	the equilibri	um geometry	of an isolat	ed diatomic	molecule (Δα	[a] Computed at ZORA-BP86-D3/T22P. [b] Results obtained for a fictitious [A—MB] structure in which the A—M fragment is in the equilibrium geometry of an isolated diatomic molecule (\( \Delta d_{MM} = 0.00 \)) while the

[a] Computed at ZORA-BP86-D3/TZ2P. [b] Results obtained for a fictitious [A—M--B]<sup>-</sup> structure in which the M--B distance is as in the equilibrium geometry of the real [A–M--B]<sup>-</sup> complex in italics and parentheses.





equilibrium geometry. The following main trends emerge from our computations: In the first place, coinage metal bonds are stronger than the corresponding alkali metal bonds and the latter are stronger than hydrogen bonds. For example, the F–M +F $^-$  bond energy  $\Delta E$  amounts to about -82, -66 and -53 kcal mol $^{-1}$  for M=Ag, Na and H, respectively (see Table 1).

Secondly, the metal bonds are consistently longer than the corresponding hydrogen bonds, by up to ca 1 Å. For example,  $d_{M\bullet\bullet\bullet}B$  in  $[F-M\cdots F]^-$  is 2.02, 2.06 and 1.16 Å, for M=Ag, Na and H, respectively (see Table 1).

Thirdly, the A–M stretch  $\Delta d_{AM}$  tends to be larger in most cases for hydrogen than for metal bonds (see Table 1). For example,  $\Delta d_{AM}$  in [F–M···F]<sup>-</sup> is 0.03, 0.11 and 0.23 Å, again for M=Ag, Na and H, respectively (see Table 1). As we will see later on, the A–M bond stretching is a consequence of charge transfer from the lone-pair orbital of B<sup>-</sup> to the antibonding  $\sigma^*_{AM}$  orbital of A–M, similar to the situation for hydrogen bonds.<sup>[4]</sup>

Lastly, both M- as well as H-bonded complexes of A–M involving the base B $^-$ =F $^-$  are more stable than the corresponding ones involving the base B $^-$ =Cl $^-$ . This is in line with the higher proton and alkali cation affinities of the fluoride compared to the chloride anion. For example, the complexation energy  $\Delta E$  of FLi, FNa and FH with F $^-$  is -73.7, -65.8 and -53.2 kcal mol $^{-1}$  which is significantly more stabilizing that the

corresponding  $\Delta E$  of -49.2, -46.3 and -27.2 kcal mol<sup>-1</sup> for complexation of the same series of diatomics with Cl<sup>-</sup> (see Table 1).

#### 3.2. Nature of the Metal Bond and Trends in Bonding

Our quantitative MO bonding analyses reveal that all metal-bonds, coinage and alkali, have in essence a similar bonding mechanism, with both electrostatic and covalent interactions, as the corresponding hydrogen bonds. The metal bonds, however, mostly go with a significantly stronger electrostatic attraction than the hydrogen bonds, in line with the more pronounced polarization in the  $A^{\delta-}\!\!\!-\!\!M^{\delta+}$  than  $A^{\delta-}\!\!\!-\!\!H^{\delta+}$  bond (see  $Q_M$  in Table 2) and because of the higher nuclear charge of the metal atoms around which the  $B^-$  lone pair extends. This is an important reason why all metal bonds, especially coinagemetal bonds, are stronger than H bonds. Dispersion interactions are essentially negligible with values of half a kcal mol $^{-1}$  at most.

Furthermore, in  $[A-M\cdots B]^-$ , the  $np_z$  lone-pair of the base  $B^-$  engages into a HOMO-LUMO interaction with the  $\sigma_{AM}^*$  of the A–M diatomic (see Figure 1). This leads to a significant covalent component, i.e., orbital interaction  $\Delta E_{\rm nir}$  in the bonding

Table 2. Bond length  $d_{AM}$  (in Å), homolytic bond dissociation energies BDE (in kcal mol<sup>-1</sup>), VDD charge  $Q_M$  of atom M (in au), frontier orbital energies ε (in eV) and overlaps of A–M molecules. (a)

	•									
					character $\sigma_{AM}^{*^{[b]}}$		$\varepsilon(\sigma_{AM}^*)^{[c]}$		$<$ n $p_{\sigma}(A) \mid ns(M) > {}^{[c]}$	
AM	$d_{AM}$	BDE	$Q_{M}$	$\epsilon(\sigma_{AM})$	$%np_{\sigma}(A)$	%ns(M)	eq	eq + 0.5 Å	eq	eq + 0.5 Å
FH	0.93	150.5	0.20	-13.55	4	48	-0.74	-4.73	0.38	0.27
FLi	1.58	145.7	0.51	-6.45	0	82	-1.31	-1.90	0.10	0.12
FNa	1.95	120.4	0.60	-5.27	2	82	-1.74	-2.25	0.10	0.11
FCu	1.74	106.5	0.37	-6.36	10	65	-4.36	-5.16	0.18	0.15
FAg	1.99	89.3	0.42	-6.91	10	75	-4.25	-4.95	0.16	0.14
FAu	1.95	80.5	0.28	-7.32	17	61	-5.24	-6.17	0.17	0.13
CIH	1.29	110.4	0.10	-11.85	22	72	-1.09	-4.52	0.48	0.32
ClLi	2.03	112.1	0.47	-6.53	0	76	-1.81	-2.32	0.22	0.20
ClNa	2.38	95.5	0.58	-5.74	3	79	-2.02	-2.47	0.19	0.16
ClCu	2.04	87.6	0.29	-6.54	13	60	-4.18	-4.96	0.29	0.22
ClAg	2.30	73.8	0.36	-7.11	13	68	-4.17	-4.87	0.26	0.19
ClAu	2.23	70.0	0.21	-7.40	22	55	-4.91	-5.82	0.28	0.20

[a] Computed at ZORA-BP86-D3/TZ2P. [b] Gross Mulliken contribution (in %) to  $\sigma_{AM}^*$ . [c] Evaluated at the equilibrium A—B bond length of AM (d<sub>AM</sub>) and at an A—M distance that has been stretched from the equilibrium geometry by 0.5 Å (d<sub>AM</sub> + 0.5 Å).

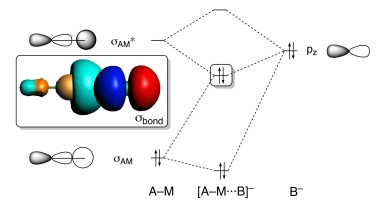


Figure 1. Schematic orbital-interaction diagram for [A-M-B] $^-$  complexes as they emerge from our quantitative MO analyses for M=H, Li, Na, Cu, Ag and Au, computed at ZORA-BP86-D3/TZ2P, with a 3D plot of the bonding  $\sigma_{AM}^* + np_z(B^-)$  combination of [F-Cu-F] $^-$ .





mechanism which reinforces the electrostatic attraction  $\Delta V_{\text{elstat}}$ between the polarized  $F^{\delta-}\!\!-\!\!M^{\delta+}$  bond and the negatively charged Lewis base F<sup>-</sup> (see Table 1). The percentage contribution of the covalent component ( $\Delta E_{oi}$ ) to the total bonding forces ( $\Delta V_{\rm elstat} + \Delta E_{\rm oi} + \Delta E_{\rm disp}$ ) is, however, smaller in the metal bonds than in the corresponding hydrogen bonds (vide infra). For example, in the strongest coinage-metal complex,  $[F-Au-F]^-$ , the gold-bond with an overall bond energy  $\Delta E_{int}$  of  $-98.3 \text{ kcal mol}^{-1} \text{ receives } -62.3 \text{ kcal mol}^{-1} \text{ of stabilization from}$ orbital interaction  $\Delta E_{\text{oir}}$  and  $-50.9 \text{ kcal mol}^{-1}$  indeed stem from the  $\sigma\text{-orbital}$  interactions  $\Delta \textit{E}_{\sigma}\text{-}$  Although this is less than the electrostatic attraction of -127.6 kcal mol<sup>-1</sup>, it is still about half the magnitude of the latter and thus of crucial importance for the stability and the nature of the bond. For comparison, in the corresponding hydrogen-bonded complex, [F-H-F]-, the overbond energy  $\Delta E_{\rm int}$  of  $-73.3 \, {\rm kcal \, mol}^{-1}$ receives  $-64.9 \text{ kcal mol}^{-1}$  of stabilization from orbital interaction  $\Delta E_{\text{oi}}$  $(-58.0\;\text{kcal}\,\text{mol}^{-1}$  of which are  $\sigma\text{-orbital}$  interactions) and -76.3 kcal mol<sup>-1</sup> from electrostatic attraction  $\Delta V_{\rm elstat}$ .

Next, after characterizing the nature of alkali- and coinagemetal bonds, we examine in more detail the computed trend in bond strengths: why does the bond strength  $\Delta E$  weaken along coinage-metal bonds, alkali-metal bonds and hydrogen bonds? Our analyses show that three phenomena in the bonding mechanism contribute significantly to this trend (see Table 1): (i) more stabilizing electrostatic attraction in the case of the metal bonds due to, among others, the aforementioned higher polarization in the  $A^{\delta-}\!\!-\!\!M^{\delta+}$  bond and, especially for the coinage-metal bonds, a higher nuclear charge at  $M_r^{[8,21,22]}$  (ii) more stabilizing orbital interactions in the coinage than in the alkali-metal bonds due to a lower orbital energy of the  $\sigma^*_{AM}$ LUMO, and thus stronger HOMO-LUMO interaction with the Lewis base, in the case of the former; and (iii) less Pauli repulsion for the alkali-metal bonds in which the occupied  $\sigma_{\text{AM}}$ bond orbital is mainly localized on A leading to little 2-orbital-4electron repulsion with the  $np_z(B^-)$  HOMO (overlap not shown in Table 1).

The more stabilizing orbital interactions in the coinagemetal bonds than in the alkali-metal bonds are a consequence of the lower energy of the  $\sigma_{\text{AM}}^*$  and the larger HOMO-LUMO overlap in the former. This LUMO emerges mainly from the antibonding combination of the  $np_{\sigma}(A)$  and the ns(M), where the ns(M) of the significantly less electropositive coinage-metal atoms is already lower in energy than that of the more electropositive alkali-metal atoms, and therefore, closer in energy to np<sub>z</sub>(F<sup>-</sup>), yielding a smaller, i.e., more favorable orbitalenergy gap. For example, in the strongest alkali complex  $[F-Li-F]^-$  ( $\Delta E = -73.7 \text{ kcal mol}^{-1}$  see Table 1), the poor HOMO-LUMO overlap ( $\langle {\sigma_{{
m FLi}}}^* | np_z({
m F}^-) \rangle \! = \! 0.11$ ) implies a low orbital interaction ( $\Delta E_{oi}$ ) of  $-14.2 \text{ kcal mol}^{-1}$ ; whereas, in the strongest coinage complex  $[F-Au-F]^-$  ( $\Delta E = -97.9 \text{ kcal mol}^{-1}$ ), the HOMO-LUMO overlap is twice as large as in the lithium case (  $\langle \sigma_{FAu}^* | np_z(F^-) \rangle = 0.27$ ), thus exhibiting stronger stabilization orbital interaction ( $\Delta E_{oi} = -62.3 \text{ kcal mol}^{-1}$ ).

The energetic difference in bond strength between coinage and alkali bonds is, as already alluded to above, partially compensated by the lower Pauli repulsion of highly polarized alkali bond ( $\Delta E_{Pauli} = 25.8 \text{ kcal mol}^{-1} \text{ for } [F-Li...F]^- \text{ compared to } \Delta E_{Pauli} = 91.6 \text{ kcal mol}^{-1} \text{ for } [F-Au...F]^- \text{ see Table 1) in which the A-M HOMO } \sigma_{FAu}$  has little amplitude on the metal (see Figure S1 in the Supporting Information).

#### 3.3. A-H versus A-M stretching

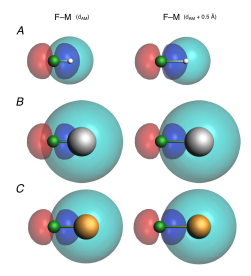
Finally, we notice that hydrogen bonds [A—H···B]<sup>-</sup> go with a substantially larger A—H expansion than the corresponding A—M expansion in metal bonds [A—M···B]<sup>-</sup>, despite the higher rigidity of the element-hydrogen bond than the element-metal bond in the A—H versus A—M molecule. For example, the diatomic F—H, with a sizeable homolytic bond dissociation energy (BDE) of ca 150 kcal mol<sup>-1</sup> (see Table 2), elongates 0.23 Å due to the complexation process (see Table 1) whereas the much weaker F—Au bond (BDE of 80.5 kcal mol<sup>-1</sup>) elongates by only 0.04 Å.

Our analyses reveal that the more substantial elongation of the A-H bond in hydrogen bonds is ultimately caused to the difference in bond-length dependence of the  $\langle np_{\sigma}(A)|ns(M)\rangle$ overlap for M=hydrogen or metal in the A-M molecule. This difference is behind an important phenomenon, namely, that the orbital energy of the  $\sigma^*_{AH}$  LUMO drops much faster upon A–H expansion than the  $\sigma^*_{AM}$  LUMO does upon A–M expansion and consequently leads to a larger gain in stabilizing orbital interaction upon A-H expansion in hydrogen bonds. For example, if we elongate the F-H bond of the hydrogen fluoride molecule from its equilibrium geometry by 0.5 Å, the  $\langle 2p_{\sigma}(F)|1s(H)\rangle$  orbital overlap significantly decreases from 0.38 to 0.27 and the orbital energy of the  $\sigma_{\text{FH}}^*$  LUMO drops significantly, from -0.74 to -4.74 eV (see Table 2), and becomes a better electron accepting LUMO for the HOMO of the F<sup>-</sup> Lewis base in [F-H--F]<sup>-</sup>. In line with this, in [F-H--F]<sup>-</sup>, the stabilizing orbital interactions  $\Delta E_{oi}$  are enhanced from -47.8 to  $-64.9 \text{ kcal mol}^{-1}$ , if we go from the fictitious situation in which the F-H molecule preserves its own, shorter equilibrium bond distance of 0.93 Å of (see Table 1). This larger gain in orbital interactions drives the A-H expansion and compensates the associated raise in strain energy.

Ultimately, this behavior could be traced to the more compact nature of the hydrogen 1s AO if compared to the metal ns AOs (see Figure 2). In the A–H molecule, the hydrogen 1s AO engages into a textbook overlap with the A np $_{\sigma}$  AO, i.e., an overlap of significant magnitude which quickly drops, as the A–H bond expands and, thus, causes the np $_{\sigma}$ (A)–1 s combination which mainly constitutes the  $\sigma^*_{AH}$  to lose its A–H antibonding overlap quickly and thus drop in energy rapidly. This can be seen in Figure 2A which shows the on-scale 3D representation of the overlapping np $_{\sigma}$ (F) and 1 s(H) atomic orbitals.

At variance, in the A–M molecules, the A n $p_{\sigma}$  AO achieves only a moderate overlap with the significantly more diffuse metal ns AO, and this overlap also changes less upon bond elongation than in the case of A–H. This situation arises in particular for the A–M systems involving alkali-metal atoms (see Figure 2). Thus, the A–M antibonding character of the  $\sigma^*_{AM}$ 





**Figure 2.** 3D plot of  $2p_{\sigma}(F^-)$  and ns(M) orbitals in A–M molecules at equilibrium distance (left) and stretched by + 0.5 Å (right) for (A) M=H, (B) M=Li and (C) M=Cu at isosurface value of 0.03.  $2p_{\sigma}(F^-)$  orbitals are represented in blue & red and ns(M) orbitals are in turquoise for clarity.

weakens more moderately and the orbital energy of this LUMO goes down to a lesser extent than in the case of the  $\sigma^*_{AH}$  LUMO. For example, if we elongate the F-Li bond of the lithium fluoride molecule from its equilibrium geometry by 0.5 Å, the  $\langle 2p_{\sigma}(F)|2s(Li)\rangle$  orbital overlap hardly changes and in fact even slightly increases, from 0.10 to 0.12 (see Table 2). The slight increase, instead of decrease, is related to the very diffuse nature of the Li 2 s AO which ranges across the F  $2p_{\sigma}$  nodal surface; this can be clearly seen in Figure 2B. Thus, F-Li elongation initially improves the overlap because unfavorable cancelation effects disappear (ultimately, if one would contribute to elongate the F-Li bond, the overlap of course decreases and goes towards zero). Altogether, the F-Li elongation by 0.5 Å has little effect on the orbital energy of the  $\sigma_{\text{FH}}^*$  LUMO which goes from -1.31 to -1.90 eV (see Table 2). As a consequence, in metal bonds [A–M···B]<sup>-</sup>, the orbital interactions  $\Delta E_{oi}$  in general hardly benefit from A–M elongation, by up to ca 1 kcal mol<sup>-1</sup> only (see Table 1).

# 4. Conclusions

Coinage-metal bonds and, to a lesser extent, alkali-metal bonds in  $[A-M\cdots B]^-$  model complexes are stronger than the corresponding hydrogen bonds in  $[A-H\cdots B]^-$ . The metal-bonds have a substantial covalent component provided by the HOMO-LUMO interaction of the occupied  $np_z$  of  $B^-$  and the empty A-M antibonding  $\sigma^*_{AM}$  that reinforces electrostatic attraction between the polarized metal-bond donor  $A^{\delta-}-M^{\delta+}$  and the anionic Lewis base  $B^-$ , similar to the bonding mechanism in hydrogen bonds, although covalency is more important in the latter. This follows from our quantitative MO bonding analyses of  $[A-M\cdots B]^-$  model complexes (M=H,Li,Na,Cu,Ag,Au;A,B=F,Cl) based on dispersion-corrected relativistic density functional theory.

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The trend in bond strengths originates from two phenomena: (i) the metal bonds have a stronger electrostatic attraction than the hydrogen bonds because of a more pronounced polarization in the  $A^{\delta-}\!\!-\!\!M^{\delta+}$  bond in the former; and (ii) coinage-metal bonds are stronger than alkali-metal bonds due to more stabilizing orbital interactions. This latter trend is the result of a lower, more favorable orbital energy of the  $\sigma^*_{AM}$  LUMO and a somewhat better more compact nature on M, leading to better overlap with the  $B^-$  np<sub>2</sub> HOMO, when M goes from the alkali metals to the somewhat more electronegative and effectively smaller coinage metals.

Finally, hydrogen bonds [A–H···B] $^-$  go with a substantially larger A–H expansion than the corresponding A–M expansion in metal bonds [A–M···B] $^-$ , despite the higher rigidity of the element-hydrogen than element-metal bond in the AH versus AM molecule. The reason for this is that the orbital energy of the  $\sigma^*_{AH}$  LUMO drops much faster upon A–H expansion than the  $\sigma^*_{AM}$  LUMO does upon A–M expansion and consequently leads to a larger gain in stabilizing orbital interaction upon A–H expansion in hydrogen bonds. This phenomenon could be traced to the difference in compactness of hydrogen (more compact) and, e.g., an alkali metal atom (more diffuse).

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# **Conflict of Interest**

The authors declare no conflict of interest.

**Keywords:** Activation Strain Model · Bond theory · DFT calculations · Hydrogen bonding · Metal bonding

- [1] a) J.-M. Lehn, Supramolecular Chemistry. Concepts and Perspectives, VCH, Weinheim, 1995; b) K. Müller-Dethlefs, P. Hobza, Chem. Rev. 2000, 100, 143–168; c) G. A. Jeffrey, An Introduction to hydrogen Bonding, Oxford University Press: New York, 1997.
- [2] a) P. Hobza, J. Sponer, Chem. Rev. 1999, 99, 3247–3276; b) J. L. Cifelli,
   C. C. Capule, J. Yang, ACS Chem. Neurosci. 2019, 10, 991–995.
- [3] a) H. Umeyama, K. Morokuma, J. Am. Chem. Soc. 1977, 99, 1316–1332; b) G. R. Desiraju, T. Steiner, The Weak Hydrogen Bond in structural Chemistry and Biology, Oxford University Press Inc.: New York, 1999; c) S. J. Grabowski, Chem. Rev. 2011, 111, 2597–2625; d) S. C. C. van der Lubbe, C. Fonseca Guerra, Chem. Asian J. 2019, 14, 2760–2769; e) S. C. C. van der Lubbe, F. Zaccaria, X. Sun, C. Fonseca Guerra, J. Am. Chem. Soc. 2019, 141, 4878–4885; f) S. C. C. van der Lubbe, C. Fonseca Guerra, Chem. Eur. J. 2017, 23, 10249–10253.





- [4] C. Fonseca Guerra, F. M. Bickelhaupt, J. G. Snijders, E. J. Baerends, Chem. Eur. J. 1999, 5, 3581–3594.
- [5] a) A. R. Voth, P. Khuu, K. Oishi, P. S. Ho, Nat. Chem. 2009, 1, 74–79; b) C.-Z. Liu, S. Koppireddi, H. Wang, D.-W. Zhang, Z.-T. Li, Angew. Chem. Int. Ed. 2018, 58, 226–230; Angew. Chem. 2018, 130, 1–6; c) E. Margiotta, S. C. C. van der Lubbe, L. Santos, G. Paragi, S. Moro, F. M. Bickelhaupt, C Fonseca Guerra, J. Chem. Inf. Model. 2020, 60, 1317–1328; d) G. Cavallo, P. Metrangolo, R. Milani, T. Pilati, A. Priimagi, G. Resnati, G. Terraneo Chem. Rev. 2016, 116, 2478–2601.
- [6] a) L. P. Wolters, F. M. Bickelhaupt, ChemistryOpen 2012, 1, 96–105; b) L. P. Wolters, P. Schyman, M. J. Pavan, W. L. Jorgensen, F. M. Bickelhaupt, S. Kozuch, WIREs Comput. Mol. Sci. 2014, 14, 523–540; c) F. M. Bickelhaupt, E. J. Baerends, N. M. M. Nibbering, Chem. Eur. J. 1996, 2, 196–207.
- [7] C. Wang, D. Danovich, Y. Mo, S. Saik J. Theory. Comput. 2014, 10, 3726–3737.
- [8] P. A. Kollman, J. F. Liebmanm, L. C. Allen, J. Am. Chem. Soc. 1970, 92, 1142–1150.
- [9] B. S. Ault, G. C. Pimentel, J. Phys. Chem. 1975, 79, 621-626.
- [10] A. B. Sannigrahi, T. Kar, B. G. Nigoyi, P. Hobza, P. v. R. Schleyer, Chem. Rev. 1990, 90, 1061–1076.
- [11] S. C. M. ADF2017.110 Theoretical Chemistry, Vrije Universiteit Amsterdam, 2017. See: www.scm.com.
- [12] a) F. M. Bickelhaupt, J. Comput. Chem. 1999, 20, 114–128; b) I. Fernandez, F. M. Bickelhaupt, Chem. Soc. Rev. 2014, 43, 4953–4967; c) L. P. Wolters, F. M. Bickelhaupt, WIREs Comput. Mol. Sci. 2015, 5, 324–343; d) F. M. Bickelhaupt, K. N. Houk, Angew. Chem. Int. Ed. 2017, 56, 10070–10086; Angew. Chem. 2017, 129, 10204–10221.
- [13] a) F. M. Bickelhaupt, E. J. Baerends Reviews in Computational Chemistry, Vol. 15; K. B. Lipkowitz, D. B. Boyd, Eds.; Wiley-VCH: New York, 2000, pp.

- 1–86; b) M. Lein, G. Frenking, *Theory and Applications of Computational Chemistry. The First 40 Years*; C. E. Dykstra, G. Frenking, K. S. Kim, G. E. Scuseria, Eds.; Elsevier: Amsterdam, **2005**.
- [14] C. Fonseca Guerra, J. W. Handgraaf, E. J. Baerends, F. M. Bickelhaupt, J. Comput. Chem. 2004 25, 189–210.
- [15] a) A. D. Becke, Phys. Rev. A 1988, 38, 3098–3100; b) J. P. Perdew, Phys. Rev. B 1986, 33, 8822–8824.
- [16] E. van Lenthe, E. J. Baerends, J. G. Snijders, J. Chem. Phys. 1994, 101, 9783.
- [17] S. Grimme, J. Antony, S. Ehrlich, H. J. Krieg J. Chem. Phys. 2010, 132, 154104.
- [18] S. A. C. McDowell, J. A. S. S. Hill, J. Chem. Phys. 2011, 135, 164303.
- [19] A. P. Bento, F. M. Bickelhaupt, J. Org. Chem. 2007, 73, 7290-7299.
- [20] a) Z. Boughala, C. Fonseca Guerra, F. M. Bickelhaupt, Chem. Asian J. 2017, 12, 2604–2611; b) Z. Boughala, C. Fonseca Guerra, F. M. Bickelhaupt, Cation Affinities throughout The Periodic Table in Advances in Inorganic Chemistry Vol. 73; R. van Eldik, R. Puchta, Eds.; Academic Press: San Diego, 2019; Chapter 6.
- [21] P. Lipkowski, S. J. Grabowski, Chem. Phys. Lett. 2014, 591, 113-118.
- [22] S. Scheiner in Lithium Chemistry, Theoretical and Experimental Overview; A. M. Sapse, P. v. R. Schleyer, Eds.; Willey: New York, 1995; p. 67–87.

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