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Term rules for simple metal clusters

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Hund's term rules are only valid for isolated atoms, but have no generalization for molecules or clusters of several atoms. We present a benchmark calculation of Al₂ and Al₃, for which we find the high and low-spin ground states ³Π_u and ²A₁' respectively. We show that the relative stabilities of all the molecular terms of Al₂ and Al₃ can be described by simple rules pertaining to bonding structures and symmetries, which serve as guiding principles to determine ground state terms of arbitrary multi-atom clusters.

The ground state terms (spin and angular momenta) of isolated atoms are determined by Hund's rules¹, which are explained by the lowering of the electronuclear attraction energy^{2–5}. For molecules and clusters, such term rules do not exist. Group theory allows us to determine the possible ^{2S+1}Ξ molecular terms, where S denotes the total spin and Ξ the symmetry species, but there is no systematic way to figure out which of these is the ground state. Direct experimental observation or quantum chemical total energy calculation is only available for a few prototype systems.

Hund's first rule of maximum spin multiplicity holds for many organic molecules^{6–8}, but not all⁹. Diatomic molecules on the other hand, tend to have spin singlet ground states with the exception of O₂ and B₂ (see e.g. refs 10–19 for diatomic molecules of main group elements). These molecules tend to have ground states that minimize the internuclear bond length, which may be associated with a lowering of the electronuclear attraction energy, but whether or not such discussion generalizes to metallic clusters remains unknown. Moreover, recent attempts to generalize Hund's rules for molecules^{8,20} or clusters^{21–23} only focus on a spin multiplicity rule, and trends or rules for Ξ (symmetry) terms remain completely uncharted territory.

Simple Al clusters emerge as the ideal model system to study term rules. Bulk Al is paramagnetic, but in low dimensional structures Al atoms may spontaneously align their spins. For example, strained quasi-1D chains of Al may exhibit ferromagnetism^{24,25}, and Al_n clusters with even n = 2, 4, 6, 8 have spin-triplet ground states^{21,26–29}. Al₃ on the other hand has spin-doublet (low-spin) and spin-quadruplet (high-spin) configurations, but which one of these is the ground state remains unresolved^{28,30–33}. The present benchmark study confirms that Al₂ has the ³Π_u high-spin ground state and unambiguously shows that Al₃ has the low-spin ²A₁' ground state. The Al₂ high-spin state is stabilized by Fermi correlation, which is not overcome by Coulomb correlation that tends to increase the stability of low-spin terms. For Al₃, however, the high-spin term has a symmetry broken geometry that preempts effective Coulomb correlation from taking place, thus un-stabilizing the high-spin term. Such symmetry lowering can debilitate high-spin terms of any multi-atom system. Moreover, for each spin state, we find a simple rule for the Ξ terms. The Ξ term with least node wavefunction is most stable, and for terms with equal number of nodes, the one with most bonds is most stable. Notice that for diatomic molecules, Ξ is the angular momentum (Ξ = Λ_(g/u)^(±)) along the internuclear axis, which can be either minimized or maximized by this rule.

Results

Al₂ has five stationary states, ¹Σ_g⁺, ¹Π_u, ¹Δ_g, ³Π_u and ³Σ_g[−], corresponding to the occupation of different molecular orbitals by two 3p electrons. Al₃ has three stationary states, ²A₁', ⁴A₂ and ⁴B₁, corresponding

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Al ₂	HF		CAS(6, 18)		CAS(6, 26)				
	E	r	E	r	E	r			
³ Π _u	-483.7824(0)	2.770	-483.8939(23)	2.714	-483.9082(13)	2.729			
³ Σ _g ⁻	-483.7700(2)	2.544	-483.8892(27)	2.474	-483.9064(16)	2.501			
¹ Σ _g ⁺	-483.7647(1)	2.996	-483.8830(21)	2.972	-483.8980(12)	2.980			
¹ Π _u	-483.7627(1) ^a	2.754	-483.8775(14)	2.745	-483.8925(8)	2.749			
¹ Δ _g	-483.7369(6)	2.619	-483.8760(28)	2.528	-483.8915(15)	2.556			
Al ₃	HF			CAS(9, 12)			CAS(9, 18)		
	E	r	θ	E	r	θ	E	r	θ
² A ₁ '	-725.6780(30)	2.607	60.0°	-725.8100(74)	2.567	60.0°	-725.8655(73)	2.530	60.0°
⁴ A ₂	-725.7007(27)	2.651	71.7°	-725.7962(63)	2.620	69.6°	-725.8554(64)	2.591	70.0°
⁴ B ₁	-725.6995(25)	2.868	54.2°	-725.7969(63)	2.784	55.8°	-725.8496(67)	2.760	55.6°

Table 1. Total energies and equilibrium structures of Al₂ and Al₃. Total energies E are given in hartree atomic unit (a.u.), and Al-Al bond r lengths in Å. For Al₃ r is the length of either leg of an isosceles triangle with vertex angle θ . Values in parenthesis are maximum errors within the virial theorem. ^aCAS(2, 2) calculation using doubly degenerate configurations in $(3p\pi_u^+)^1(3p\pi_u^-)^1$.

to the occupation of different molecular orbitals by three $3p$ electrons. Their equilibrium nuclear geometries and corresponding total energies E are shown in Table 1. Hartree-Fock (HF) calculation predicts Al₂ and Al₃ to have ³Π_u and ⁴A₂ high-spin ground states, respectively. Inclusion of Coulomb correlation by CAS-SCF (see Methods) maintains the high-spin ground state of Al₂, but stabilizes the ²A₁' low-spin ground state of Al₃. At the same time, high-spin terms of Al₂ (³Π_u, ³Σ_g⁻) and Al₃ (⁴A₂, ⁴B₁) become nearly degenerate; the energy difference between them is smaller than 0.01 a.u. The ³Π_u ground state for Al₂ is consistent with experiment³⁴, and our prediction of the ²A₁' ground state for Al₃ is corroborated by the Stern-Gerlach experiment³⁰. More importantly, the ground state of Al₂ is consistent with both Hund's first and second rules, whereas Al₃ violates both of them.

Potential energy components. Traditionally, Hund's rules have been interpreted as an energy gain due to the inter-electron repulsion potential energy V_{ee} ³⁵⁻³⁷, and more recently as an energy gain due to the electronuclear attraction V_{en} ^{2,4,5,7,8}. In order to analyze whether or not similar energy lowering mechanisms can be invoked for Al₂ and Al₃, we decompose the total energies given in Table 1 into potential energy components shown in Fig. 1. In both HF (dashed lines) and CAS-SCF (solid lines) calculations for each stationary state of Al₂ and Al₃, repulsion terms V_{ee} (red lines) and V_{nn} (blue lines; inter-nuclear repulsion) are positive and the attraction term V_{en} (purple lines) is negative. The total energies E of Al₂ and Al₃ calculated by CAS-SCF always lie lower than those calculated by HF. For both Al₂ and Al₃, upon inclusion of Coulomb correlation by CAS(6, 26) and CAS(9, 18), respectively, the individual potential energy components V_{en} , V_{ee} and V_{nn} composing V change as follows: both V_{ee} and V_{nn} increase and V_{en} decreases. The correlation energies $E^c = E^{CAS} - E^{HF}$, along with V^c , V_{en}^c , V_{ee}^c , and V_{nn}^c , defined similarly, are unique to each molecular term; $E^c < 0$ always, and for the components we find $V_{en}^c < 0$, $V_{ee}^c > 0$, and $V_{nn}^c > 0$.

For Al₂, both HF and CAS-SCF predict

$$E^{HF/CAS}({}^3\Pi_u) < E^{HF/CAS}({}^3\Sigma_g^-) < E^{HF/CAS}({}^1\Sigma_g^+) < E^{HF/CAS}({}^1\Pi_u) < E({}^1\Delta_g). \quad (1)$$

The correlation energies, however, exhibit the different trend

$$E^c({}^1\Delta_g) < E^c({}^3\Sigma_g^-) < E^c({}^1\Sigma_g^+) < E^c({}^1\Pi_u) < E^c({}^3\Pi_u) \quad (2)$$

making the excitation energies smaller. For Al₃, HF predicts

$$E^{HF}({}^4A_2) < E^{HF}({}^4B_1) < E^{HF}({}^2A_1') \quad (3)$$

and CAS-SCF predicts

$$E^{CAS}({}^2A_1') < E^{CAS}({}^4A_2) < E^{CAS}({}^4B_1) \quad (4)$$

i.e. the level ordering is altered by correlation effects. For correlation energies we find

$$E^c({}^2A_1') < E^c({}^4A_2) < E^c({}^4B_1). \quad (5)$$

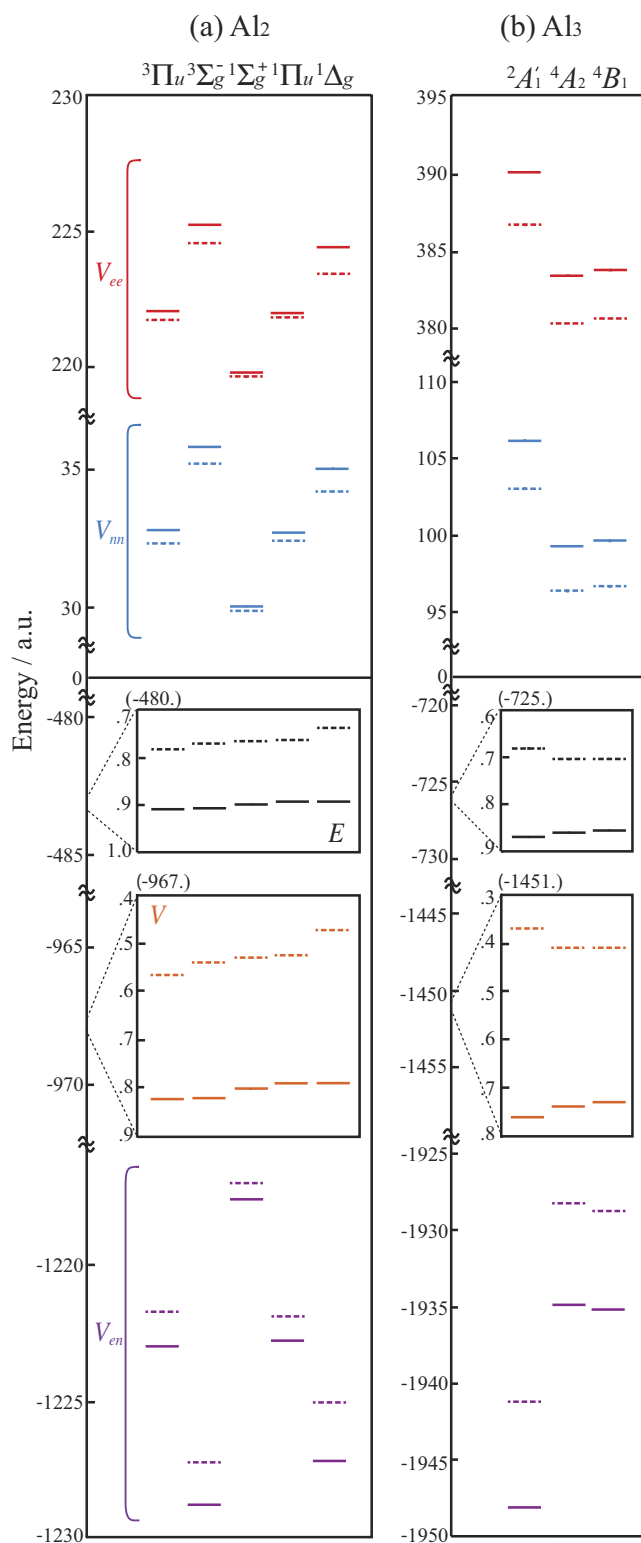


Figure 1. Energy components for each equilibrium state of Al_2 and Al_3 . HF, CAS(6, 26) for Al_2 and CAS(9, 18) for Al_3 levels are dashed lines and solid lines, respectively. E (black), V (orange), V_{ee} (red), V_m (blue), and V_{en} (purple) are given in hartree atomic units.

For both Al_2 and Al_3 , the strongest correlation effect, i.e. greatest correlation energy E^c , is observed for the ${}^1\Delta_g$ and ${}^2A_1'$ low-spin terms, respectively. For Al_3 , this correlation effect is strong enough to alter the level ordering of the molecular terms, but for Al_2 not. Thus, the relative stability of the Al_2 molecular

Al ₂		Al ₃	
³ Π _u	[3s]3pσ _g ¹ 3pπ _u ¹	² A' ₁	[3s]3pπ(a' ₂) ² 3pσ(a' ₁) ¹
³ Σ _g ⁻	[3s]3pπ _u ²	⁴ A ₂	[3s]3pπ(b ₁) ¹ 3pσ(a ₁) ¹ 3pσ(b ₂) ¹
¹ Σ _g ⁺	[3s]3pσ _g ²	⁴ B ₁	[3s]3pπ(b ₁) ¹ 3pσ(a ₁) ¹ 3pσ(a ₁) ¹
¹ Π _u	[3s]3pσ _g ¹ 3pπ _u ¹		
¹ Δ _g	[3s]3pπ _u ²		

Table 2. Electronic configurations of valence electrons of Al₂ and Al₃. [3s] represents the configurations of 3s electrons; In Al₂, [3s]=3sσ_g²3sσ_u², In Al₃, [3s] = 3sσ(a'₁)²3sσ(e')⁴ for ²A'₁, and [3s] = 3sσ(a₁)²3sσ(a₁)²3sσ(b₂)² for ⁴A₂ and ⁴B₁.

terms can be discussed based on Fermi correlation (Pauli's exclusion principle) and HF calculations, but for Al₃, Coulomb correlation included by CAS-SCF is crucial for the description of molecular terms.

For Al₂, Hund's first and second rules predict

$$E(^3\Pi_u) < E(^3\Sigma_g^-) < E(^1\Delta_g) < E(^1\Pi_u) < E(^1\Sigma_g^+), \quad (6)$$

which is valid only for the spin-triplet terms. The spin-singlet terms exhibit an opposite trend to Hund's second rule. Term stabilities have earlier been interpreted by either V_{ee}^{35-37} or $V_{en}^{2,4,5,7,8}$, which imply that total energy differences are dominated by one potential energy component V_i ($i = ee, en, \text{ or } nm$), i.e., the total energy should follow the trend of this dominant V_i . Figure 1, however, shows that

$$(\pm)V_i(^3\Sigma_g^-) < (\pm)V_i(^1\Delta_g) < (\pm)V_i(^3\Pi_u) < (\pm)V_i(^1\Pi_u) < (\pm)V_i(^1\Sigma_g^+), \quad (7)$$

which is different to Eq. (1). Here the + sign corresponds to $i = en$, and the - sign to $i = ee$ and $i = nm$. Clearly total energy trends do not follow any one particular potential energy component. Although the highest spin multiplicity (Hund's first) rule does not follow any of the potential energy components V_{en} , V_{ee} , and V_{nm} , the Ξ terms, when observed for spin-triplet and spin-singlet states individually, exhibit the following trends

$$(\mp)V_i(^3\Pi_u) < (\mp)V_i(^3\Sigma_g^-) \quad (8a)$$

$$(\pm)V_i(^1\Sigma_g^+) < (\pm)V_i(^1\Pi_u) < (\pm)V_i(^1\Delta_g). \quad (8b)$$

Note that Eq. (8a) has the opposite sign convention to Eqs (7) and (8b). Thus, for a given spin multiplicity, the potential energy components follow the same trend as total energies, but the sign may vary case by case!

Fermi correlation and bond structure. Since clear term rules cannot be described based on the individual energy components discussed above, we turn our attention to the bond structures given in Table 2 for each molecular term. For Al₂, inclusion of Coulomb correlation via CAS-SCF does not alter the relative stability of the Al₂ terms, so the relative term stabilities can be understood purely based on Fermi correlation (Pauli exclusion principle). This leads to a simple description based on the bond structures of the different terms, i.e., the nodal structure of the wavefunction. Al₂ has 3pσ_g and 3pπ_u bonding orbitals, and for the spin-singlet and spin-triplet terms, the most stable Ξ term has an occupied 3pσ_g orbital, i.e., the least node configuration. The stability of the spin-triplet ³Π_u against the spin-singlet ¹Σ_g⁺ term also follows from HF theory. Starting from the nodeless ¹Σ_g⁺ wavefunction, moving one electron from the 3pσ_g into a 3pπ_u with parallel spin (forming the ³Π_u term) lowers the total energy in three steps: (i) for fixed orbitals and Al–Al bond length, V_{ee} is lowered for spin parallel electrons³⁵; (ii) relaxing the electronic orbitals lowers the total energy further; and (iii) relaxing the Al–Al bond length lowers the total energy further still. Repeating steps (ii) and (iii) obviously keeps lowering the total energy until convergence is found; these steps can be roughly associated to changes in V_{en} and V_{nm} , respectively, but as seen in Fig. 1, for Al₂ V_{ee} and V_{nm} actually increase despite the initial lowering of V_{ee} in step (i). For the Ξ terms we find that for a given spin multiplicity, the total energy increases as the number of nodes in the wavefunction increases.

Coulomb correlation and bond structure. The above discussion fails for Al₃. Inclusion of Coulomb correlation via CAS-SCF un-stabilizes the spin-quadruplet terms despite their possession of two electrons in 3pσ type orbitals (a₁ and b₂ for ⁴A₂ and two a₁s for ⁴B₁) on the Al₃ molecular plane. We analyze the effects of Coulomb correlation based on the electron density distribution change defined by

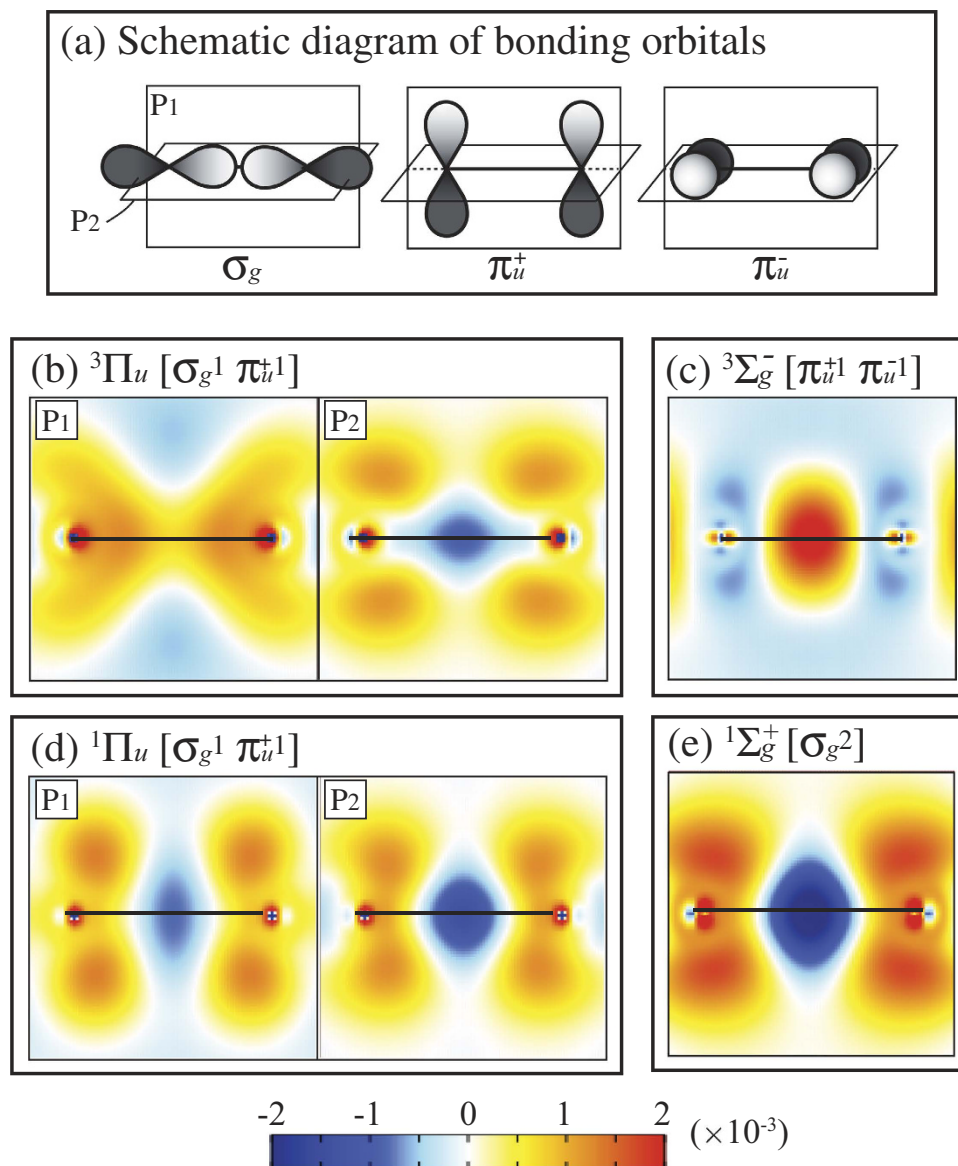


Figure 2. Schematic of bonding orbitals (a) and Coulomb correlation induced electron density distribution change ρ^c for stationary states of Al_2 (b–e). Here $\rho^c = \rho^{\text{CAS}} - \rho^{\text{HF}}$, where ρ^{CAS} is evaluated at CAS(6, 18).

$\rho^c = \rho^{\text{CAS}} - \rho^{\text{HF}}$, where ρ^{CAS} and ρ^{HF} are the total electron densities calculated by CAS-SCF and HF, respectively. The crucial Coulomb correlation that alters the Al_3 term stabilities occurs at CAS(9, 12), and therefore we evaluate ρ^{CAS} for Al_2 and Al_3 using CAS(6, 18) and CAS(9, 12), respectively. The ρ^c shown in Figs 2 and 3 for Al_2 and Al_3 , respectively, are evaluated at the equilibrium nuclear configurations obtained by CAS(6, 18) and CAS(9, 12), respectively.

Al_2 . The Coulomb correlation effects are analyzed based on the bonding $3p\sigma_g$ and $3p\pi_u$ orbitals shown in panel (a) of Fig. 2. Panels (b)–(e) of Fig. 2 show the electron density differences ρ^c for the ${}^3\Pi_u$, ${}^3\Sigma_g^-$, ${}^1\Pi_u$, and ${}^1\Sigma_g^+$ terms in the planes P_1 and P_2 corresponding to the $3p\sigma_g$ and $3p\pi_u$ bonding orbitals. The blue areas indicate a depletion of electron density, and the yellow–orange–red areas an increase of electron density. For the ${}^3,1\Sigma_g^\pm$ terms, these P_1 and P_2 planes are equivalent. The ρ^c analysis is omitted for the ${}^1\Delta_g$ term, which is not correctly represented in the HF calculation.

The CAS(6, 18) calculation includes various configurations including up to $3d$ orbitals, but the essence of the Coulomb correlation effects can be described based on the mixing of the bonding $3p\sigma_g$ and $3p\pi_u$ orbitals. As shown in panels (b)–(e) of Fig. 2, the electron density distribution corresponding to orbitals occupied in HF theory (Table 2) is depleted, and increases corresponding to bonding orbitals not occupied in HF theory. For the ${}^3\Pi_u$, ${}^1\Pi_u$, and ${}^1\Sigma_g^+$ terms that in HF have an occupied $3p\sigma_g$ orbital, there is a

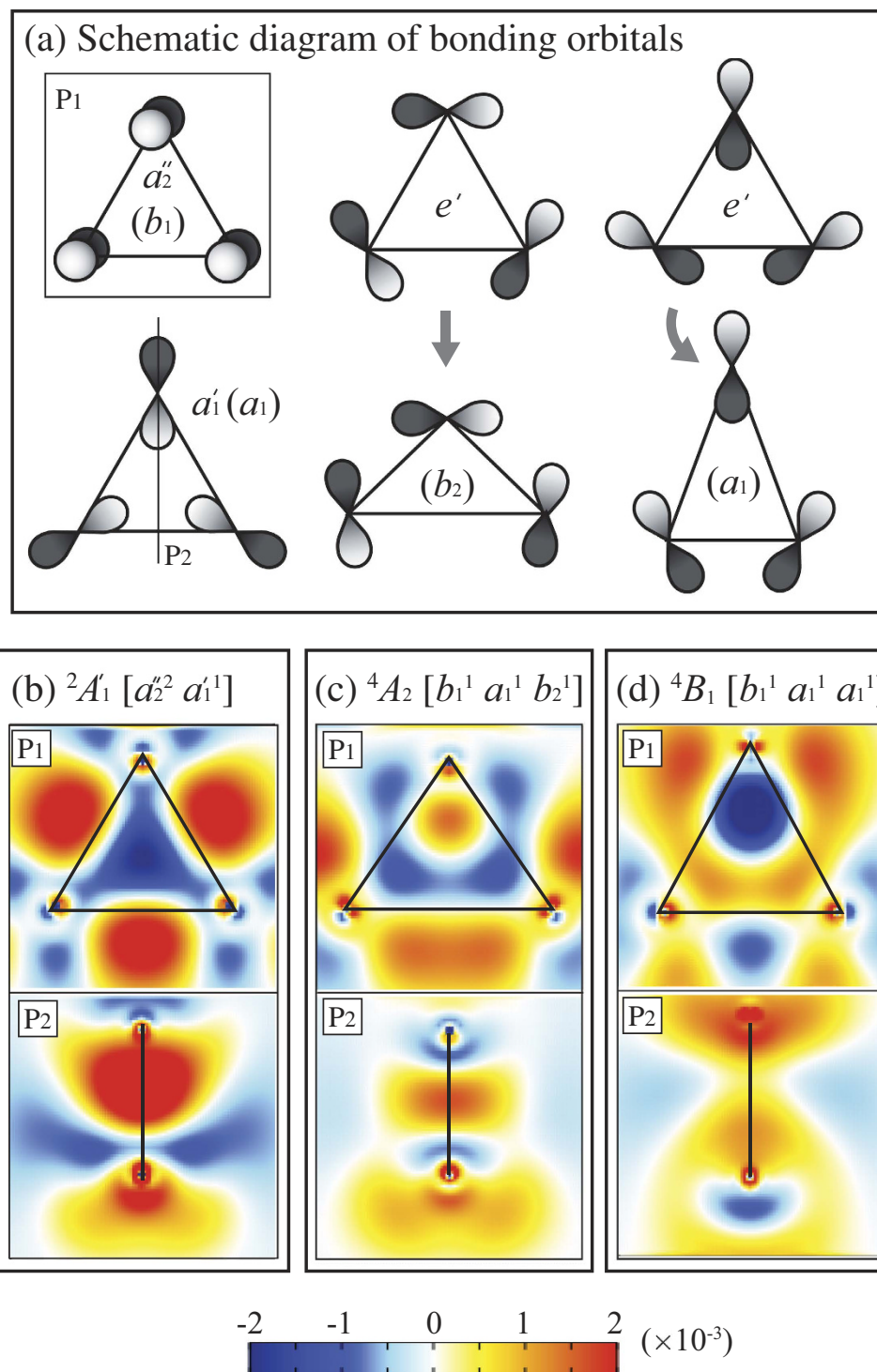


Figure 3. Schematic of bonding orbitals (a) and Coulomb correlation induced electron density distribution change ρ^c for stationary states of Al_3 (b–d). Here ρ^{CAS} is evaluated at CAS(9, 12).

depletion in ρ^c along the bond axis, and for the ${}^3\Sigma_g^-$ that in HF does not have an occupied $3p\sigma_g$ orbital, there is an increase. Likewise, ρ^c is negative in the regions corresponding to $3p\pi_u$ orbitals occupied in HF theory, and positive in the regions where the $3p\pi_u$ orbitals are not occupied in HF theory. Because all these Coulomb correlation effects essentially occur among the same set of orbitals, all of which are bonding, the effects are similar. Because the Coulomb correlation effects are similar for all terms, Coulomb correlation does not alter the relative stability of them, and the discussion above of term stability based on Fermi correlations and wavefunction nodal structure is sufficient.

Al_3 . Al_3 has the ${}^2A_1'$ low-spin ground state, against expectations from Hund's first rule or the spin-state stabilization mechanism for Al_2 pertaining to HF theory. Thus, the energy lowering effect of Coulomb correlations is different for the low-spin ${}^2A_1'$ term and the high-spin 4A_2 and 4B_1 terms. The effect of these Coulomb correlations is discussed based on the $3p\sigma$ and $3p\pi$ orbitals shown in panel (a) of Fig. 3. Panels (b)–(d) of Fig. 3 show the electron density differences ρ^e for the ${}^2A_1'$, 4A_2 , and 4B_1 terms in the plane P_1 of the nuclei of Al_3 , and its perpendicular plane P_2 , which is a reflection symmetry plane of Al_3 . Notice that the nuclei of the spin-doublet ${}^2A_1'$ term form equilateral triangle, whereas the spin-quadruplet terms 4A_2 and 4B_1 correspond to isosceles triangles. Ensuingly, the bonding orbitals for the low-spin and high-spin terms are quite different.

${}^2A_1'$ has a doubly occupied $a_{2''}$ bonding orbital, a singly occupied $a_{1'}$ orbital, and a doubly degenerate e' LUMO. The $a_{1''}$ orbital is a π bond where the plane of nuclei is a nodal plane, and the $a_{1'}$ is a σ bond with the charge density lobe in the center of the triangle. Both $a_{2''}$ and $a_{1'}$ orbitals have C_{3v} symmetry, resulting in an equilateral trimer with D_{3h} symmetry. The doubly degenerate e' LUMO corresponds to a σ bond with charge density lobes at all three sides of the triangle. The main Coulomb correlation effect is similar to what was discussed above for Al_2 . There is a depletion of electron in the regions corresponding to the $a_{2''}$ and $a_{1'}$ orbitals occupied in HF theory, and an increase in the region corresponding to the e' orbitals, as seen in Fig. 3(b).

For the spin-quadruplet terms one of the $a_{2''}$ electrons occupies either one of the e' orbitals. Individually these orbitals have the C_{2v} symmetry, yielding Jahn-Teller distorted isosceles triangles as described in Table 2. This changes the also symmetry species of the occupied $a_{2''}$ and $a_{1'}$ orbitals into b_1 and a_1 , respectively, but these orbitals still maintain their nature as π and σ bonds with similar charge density lobes as described above for the equilateral triangle. The newly formed a_1 or b_2 orbitals for the 4B_1 or 4A_2 terms, have charge density lobes at the base or legs of the triangle, respectively, as shown in Fig. 3(a). The LUMO of the 4B_1 and 4A_2 terms are b_2 and a_1 , respectively, i.e., the other one of the e' orbitals for an equilateral triangle. For the spin-quadruplet terms, the main Coulomb correlation effect is the mixing of the a_1 or b_2 orbitals, which can be seen Fig. 3(c,d) as a depletion of electron density along the legs (base) of the triangle for 4A_2 (4B_1) and the corresponding along the base (legs) of the triangle.

Because of different symmetries, the Coulomb correlations for the low-spin and the high-spin terms of Al_3 are fundamentally different. For the spin-doublet term, the main Coulomb correlation is the mixing of two occupied states and an unoccupied *doubly degenerate* state, whereas for the spin-quadruplet terms, the main Coulomb correlation is due to the mixing of one occupied and one unoccupied state. Coulomb correlation acts strongly among states nearby in energy and real space, and for the spin-quadruplet terms, the Jahn-Teller distortion imposes a severe limitation on the availability of such *nearby* states for mixing. This, combined with the fact that Coulomb correlation (even without geometrical distortions) is larger for low-spin configurations² in total stabilizes the Al_3 low-spin ground state. Thus, both Hund's first rule and the mechanism that stabilizes the high-spin ground state of Al_2 are violated because the breaking of symmetry of the Al_3 spin-quadruplet configurations reduces their Coulomb correlation. Note that Hund's maximum spin multiplicity rule is violated under exactly the opposite conditions as postulated by Kutzelnigg and Morgan²⁰.

Larger clusters. Application of the term rules described above for other clusters is straight forward. We illustrate this generalization by predicting ground state terms for Al_4 and Al_5 . For both clusters, we consider previously described planar and pyramidal structures^{26,28,38}, incidentally, our discussion below offers a new interpretation for why planar geometries are favored against pyramidal ones³⁹. We predict ${}^3B_{1u}$ and 2B_1 ground states for Al_4 and Al_5 , respectively, well in agreement with previous works^{21,26,28}. The structures and spin multiplicities agree also with density-functional calculations^{29,38}, which however give no information of the symmetry species Ξ .

Al_4 . Al_4 can have spin-singlet, spin-triplet, and spin-quintet states due to different configurations of four $3p$ electrons, shown in Table 3. The HOMO of any of the Al_4 terms with pyramidal structure (3-fold degenerate $2t_1$ orbitals) do not form σ -type bonds, so for any spin multiplicity, the least node wavefunctions corresponds to a planar geometry. The planar Al_4 spin-quintet terms (high spin) always have at least one occupied antibonding molecular orbital, such as $2b_{3u}$, $2b_{2u}$, $2b_{3g}$, and $2b_{2g}$, whereas the spin-triplet and singlet terms ${}^3B_{1u}$, 3A_u , ${}^3B_{1g}$, and 1A_g have valence electrons occupying in bonding orbitals ($1b_{1g}$, $1b_{1u}$, and $3a_2$). Thus, Fermi correlation stabilizes the spin-triplet terms with possession of most-occupied σ -type bonding orbitals, i.e., ${}^3B_{1u}$ state. Coulomb correlation, which enhances the electron density on the nodal plane of HOMO(s), makes Al-Al bonds on the molecular plane stronger for both ${}^3B_{1u}$ and 1A_g states. As seen in stability of Al_2 's spin triplet terms, such Coulomb correlation effect cannot reverse the relative stability for ${}^3B_{1u}$ and 1A_g , and thus we predict ${}^3B_{1u}$ as the ground state term of Al_4 .

Al_5 . Al_5 can have spin multiplicities up to spin-sextet due to different configurations of five $3p$ -electrons. All pyramidal, and the planar spin-sextet terms have at least one electron in antibonding or nonbonding orbitals (see Table 4), and thus cannot be more stable than the planar spin-doublet or spin-quadruplet terms. The planar spin-quadruplet terms (intermediate spin state) only have partial bonds, such as $3b_1$,

Geometry	Electronic configuration	Schematic molecular orbitals
Planar	${}^5B_{3g/2g}$ [3s] $1b_{1g}^1 1b_{1u}^1 3a_g^1 2b_{3u/2u}^1$	
	${}^5B_{3u/2u}$ [3s] $1b_{1g}^1 1b_{1u}^1 3a_g^1 2b_{3g/2g}^1$	
	${}^3B_{1u}$ [3s] $1b_{1g}^2 1b_{1u}^1 3a_g^1$	
	3A_u [3s] $3a_g^2 1b_{1g}^1 1b_{1u}^1$	
	${}^3B_{1g}$ [3s] $1b_{1u}^2 1b_{1g}^1 3a_g^1$	
	1A_g [3s] $1b_{1g}^2 1b_{1u}^2$	
Pyramidal	5A_1 [3s] $2a_1^1 2t_2^3$	
	3T_1 [3s] $2a_1^2 2t_2^2$	
	1T_1 [3s] $2a_1^2 2t_2^2$	

Table 3. Electronic configurations of valence electrons for planar and pyramidal geometries for Al_4 . Here [3s]= $1a_g^2 1b_{3u}^2 1b_{2u}^2 2a_g^2$ for the planar, and [3s]= $1a_1^2 1t_2^6$ for the pyramidal geometries.

Geometry	Electronic configuration	Schematic molecular orbitals
Planar	4A_1 [3s] $2b_1^2 1b_2^2 4a_1^1 3b_1^1 4b_1^1$	
	4A_2 [3s] $1b_2^2 1a_2^2 4a_1^1 2b_1^1 2b_2^1$	
	2B_1 [3s] $2b_1^2 1b_2^2 4a_1^2 3b_1^1$	
	2B_2 [3s] $1b_2^2 1b_1^2 2a_2^2 2b_2^1$	
	2A_1 [3s] $2b_1^2 1b_2^2 (3b_1/2b_2)^2 4a_1^1$	
Pyramidal	4A_1 [3s] $1b_2^2 (3a_1/4a_1)^1 (2e/3e)^2$	
	4E [3s] $1b_2^2 3a_1^1 4a_1^1 (2e/3e)^1$	
	2E [3s] $1b_2^2 3a_1^2 2e^1$	
	2A_1 [3s] $2b_1^2 1b_2^2 (3b_1/2b_2)^2 4a_1^1$	

Table 4. Electronic configurations of valence electrons for planar and pyramidal geometries for Al_5 . Here [3s]= $1a_1^2 1b_1^2 2a_1^2 3a_1^2$ for the planar, and [3s]= $1a_1^2 1e_1^4 2a_1^2 1b_1^2$ for the pyramidal geometries.

$4b_1$, and $2b_2$, which leaves only spin-doublet terms with strong σ -type bonds. Thus for Al_5 , we predict the planar 2B_1 spin-doublet term, which has the least node structure ground state.

Discussion

Our benchmark first principles calculation predicts the ${}^3\Pi_u$ (high-spin) and ${}^2A_1'$ (low-spin) ground states for Al_2 and Al_3 , respectively. Detailed analysis of potential energy components of the total energy reveal that previous interpretations, attributing atomic or molecular term stabilization to either $V_{ee}^{35,36}$ or $V_{en}^{2,5,8}$ are, in general, not valid for multi-atom systems. The relative stability of the Ξ terms for a given spin multiplicity for either Al_2 and Al_3 follows simple arguments based on bonding structures: For a given spin multiplicity the Ξ term possessing the most-occupied σ bonding orbitals (least node structure) is stabilized within the one-electron orbital picture according to Hartree-Fock (HF) theory. In addition, HF theory tends to stabilize the high-spin term due to Fermi correlation (Pauli exclusion principle). Coulomb correlation lowers the energy by mixing some of the orbitals occupied in HF theory with nearby unoccupied orbitals. For Al_2 , the Coulomb correlation effects are similar for all terms, but for Al_3 , Coulomb correlation alters the relative term stability. For Al_3 , breaking of symmetry of the the spin-quadruplet terms significantly limits the orbital mixing and energy lowering by Coulomb correlation. The high symmetry of the spin-doublet term, on the other hand, allows for mixing with *degenerate* levels followed by a much larger energy lowering by Coulomb correlation, stabilizing the low-spin ${}^2A_1'$ ground state of Al_3 . These stabilization mechanisms are not specific for Al clusters, and serve as simple term rules to determine the ground state of arbitrary multi-atomic systems. We demonstrate this predictive power by predicting ${}^3B_{1u}$ and 2B_1 ground states for Al_4 and Al_5 , respectively.

Methods

The total energy E of the ${}^{2S+1}\Xi g/u$ term of an Al_n cluster in the Born-Oppenheimer approximation is given by $\langle {}^{2S+1}\Xi g/u | \hat{T} + \hat{V}_{nn} + \hat{V}_{en} + \hat{V}_{ee} | {}^{2S+1}\Xi g/u \rangle$, where $| {}^{2S+1}\Xi g/u \rangle$ is a many-electron wavefunc-

tion and the operators \hat{O} give the electron kinetic energy, the inter-nuclear repulsion, the electronuclear attraction, and the inter-electron repulsion, respectively. The expectation values $\langle \hat{O} \rangle^{2S+1\Xi g/u}$ for each operator \hat{O} , henceforth denoted as $O^{(2S+1\Xi g/u)}$, are calculated using the GAMESS package⁴⁰. We use both Hartree-Fock (HF) method and complete active space self-consistent field (CAS-SCF) method. Our CAS-SCF many-electron wavefunctions contain configuration interactions among the 3s and 3p valence shells and empty 3d-derived orbitals: CAS(6, 26) and CAS(9, 18) for Al₂ and Al₃, respectively. CAS(n, m) stands for a CAS-SCF calculation with *n* active spaces and *m* active electrons. Atomic orbitals are expanded within the aug-cc-pVTZ basis set, and all nuclear positions are relaxed. This gives a virial ratio of $-V/T = 2.00000 \pm 0.00003$ for each molecular term $^{2S+1\Xi g/u}$.

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Author Contributions

D.Y. conducted the calculations, D.Y. and H.R. analysed the results. H.R. wrote the manuscript. Both authors reviewed the manuscript.

Additional Information

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Erratum: Term rules for simple metal clusters

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In the Methods section of this Article, all instances of 'g/u' should have been removed. The Methods section should read:

The total energy E of the $^{2S+1}\Xi$ term of an Al_n cluster in Born-Oppenheimer approximation is given by $\langle ^{2S+1}\Xi | \hat{T} + \hat{V}_{nn} + \hat{V}_{en} + \hat{V}_{ee} | ^{2S+1}\Xi \rangle$, where $| ^{2S+1}\Xi \rangle$ is a many-electron wavefunction and the operators \hat{O} give the electron kinetic energy, the inter-nuclear repulsion, the electronuclear attraction, and the inter-electron respectively. The expectation values $\langle ^{2S+1}\Xi | \hat{O} | ^{2S+1}\Xi \rangle$ for each operator \hat{O} , henceforth denoted as $O(^{2S+1}\Xi)$, are calculated using the GAMESS package.⁴⁰ We use both Hartree-Fock (HF) method and complete active space self-consistent field (CAS-SCF) method. Our CAS-SCF many-electron wavefunctions contain configuration interactions among the $3s$ and $3p$ valence shell and empty $3d$ -derived orbitals: CAS(6,26) and CAS(9,18) for Al_2 and Al_3 , respectively. CAS(n,m) stands for a CAS-SCF calculation with n active spaces and m active electrons. Atomic orbitals are expanded within the aug-cc-pVTZ basis set, and all nuclear positions are relaxed. This gives a virial ratio of $-V/T = 2.00000 \pm 0.00003$ for each molecular term $^{2S+1}\Xi$.



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